EXPERIMENTAL STUDY OF THE RHEOLOGY AND STABILITY BEHAVIOUR
OF SURFACTANT STABILIZED WATER-IN-OIL EMULSION

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EXPERIMENTAL STUDY OF THE RHEOLOGY AND STABILITY BEHAVIOUR OF SURFACTANT STABILIZED WATER-IN-OIL EMULSION

A THESIS APPROVED FOR THE MEWBOURNE SCHOOL OF PETROLEUM AND GEOLOGICAL ENGINEERING

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Dedication

I dedicate this work first to God almighty for his love, provision and protection. My deepest gratitude goes also to my parents, brothers and my fiancée for their indefatigable love and support throughout my life. Thank you all for being there for me always.
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Abstract

Formation water almost always comes out with crude oil production. Under extreme mixing and high turbulence provided by pumps, valves, chokes and pipelines in the production system, water and oil can mix to form highly stable and viscous emulsions. These emulsions create flow assurance and separation problems in deep water production facilities because of the increasing viscosity with water fraction. In order to predict the changes in viscosity and stability with time it is important to understand the flow behavior of these emulsion in pipelines. It is also important to understand how the acid and bases interact with the other surface active components to stabilize emulsion. This thesis focuses on characterizing the behavior of surfactant stabilized crude oil emulsion with regards to stability, viscosity, total acid and base number. The outcome of this study will be used to study the flow of emulsions in pipelines under different conditions.

A reproducible emulsification protocol was established. Experimental data showed that span85 surfactant produces very stable emulsion with the crude oil. Increasing the surfactant concentration only stabilizes the emulsion up to certain amount before additional amount destabilizes the emulsion. Adding surfactant at low concentration reduces the viscosity of the emulsion by as much as 75% when compared to emulsion without any surfactant. Viscosity of the emulsion plays a significant role in emulsion stabilization. Viscosity profile with increasing total acid number shows a minimum beyond which the viscosity increases with increasing total acid number. For a given water fraction, there is a range of total acid number where the acid stabilizes the emulsion. Outside of this range the acid destabilize the emulsion.
CHAPTER 1

INTRODUCTION

1.1 Importance of oil and gas in energy generation

Energy is fundamental to human existence. From day to day we make use of energy without even thinking about it. Gasoline, diesel and compressed natural gas are all used in our automobiles for transportation, cooking our food, heating our homes and commercial building and also for generating electricity. Petroleum oil and gas also provide basic raw materials for plastics, paints fertilizer, insecticides, soaps industries etc. Providing raw materials to these industries is very important to modern day industrial growth. It is perhaps the single most substance consumed in our modern society (Speight, 2007). Energy from fossil fuels such as crude oil and gas supply over a half of the world’s total energy demand. The trend in energy use is expected to increase worldwide as the population continues to grow (Petrecca, 2014). The United States Energy Information Administration, (EIA, 2016) energy outlook projects that the world’s energy consumption will grow by 48% from 2012 to 2040 as shown in Figure 1.1. Over the next two decades nearly all of these growth is forecasted to come from developing world (Wolfram et al, 2012).

1.2 Oil and gas Recovery mechanisms

Oil and gas is produced today in almost every part of the world. From a few barrels per day to several millions of barrels in shallow 20-meter-deep reservoir to 3000-meter-deep water reservoirs (Devold, 2009). Production usually takes place either onshore or offshore. Offshore productions can either be from shallow or deep water fields.
Figure 1.1: Projected world energy consumption by 2040 (source: US EIA International Energy Outlook 2016)

Today, most oil and gas extraction still takes place onshore. Nonetheless, a considerable amount of gas and oil is already produced offshore. Offshore oil extraction currently accounts for 37 per cent of global production. At present, 28 per cent of global gas production takes place offshore – and this is increasing (Cluster of Excellence).

Onshore and offshore oil exploration each has its own unique challenge. One thing that is common with exploration is that well has to be drilled. Oil and gas are extracted at first using the natural energy of the reservoir in what is called primary recovery mechanism. Sometimes multiple wells are drilled in order to recover the oil and gas economically. With time the natural energy of the reservoir is depleted hence the need for secondary recovery process such as water flooding, gas flooding, steam flooding, water alternating gas flooding etc. Secondary recovery processes sometimes might mean drilling secondary wells to inject water, steam or gas into the reservoir. This is to sustain or increase average reservoir pressure for constant production rate. Secondary recovery techniques sometimes may involve placing pumps (electric submersible pump)
downhole to provide additional energy to pump the oil to the surface. It may also involve gas injection to provide artificial lift of the oil to the surface. Primary and secondary recovery methods including water flooding or reinjection of produced natural gas, produce on the average about one-third of the original oil in place (OOIP). However, by applying the tertiary recovery (commonly called Enhanced Oil Recovery, EOR), production could reach 40 to 60% of oil in the reservoir (Abubaker et al, 2015).

1.3 Emulsion Challenges associated with crude oil production

Emulsion constitute a major challenge in all aspect of crude oil production and processing. One of the biggest flow assurance problems associated with crude oil production is the formation of stable and viscous water in oil emulsion. This is particularly true for deep water production. Emulsion is usually not a problem in the initial stage of the life of a reservoir because the amount of produced water is so little. With time however the amount of the produced water increases significantly with a corresponding decrease in oil production as shown in Figure 1.2. The figure shows that as a field matures, produced water becomes the biggest fraction of production. Produced water is water trapped in underground formations that is brought to the surface along with oil or gas. It is by far the largest volume waste stream associated with oil and gas production (John A. Veil et al, 2004). Produced water together with crude oil form emulsion when they flow through valves, chokes or pumps. They are stabilized by the presence of natural surfactant present in the oil. Emulsion formation can either occur in the reservoir or during production and transportation to the surface separation facilities. High viscosity associated with water-in-oil emulsion is the root cause of the flow assurance problems. In the reservoir, viscous emulsion can block the
reservoir pore spaces reducing permeability and sweep efficiency. During transportation, viscous emulsion causes very significant pressure drop and eventual loss of production. Emulsion poses very serious flow assurance problems in deep water reservoirs where different wells are commingled together before they are transported in subsea pipelines to separation facilities thousands of miles away. Emulsion also causes a lot of operational problems such as tripping of separators equipments, production of off-specification crude oil, increased chemical cost of separating emulsion and corrosion of pipelines (Kokal, 2005)

1.4 Research Problem Statement

Production of crude oil almost always comes with the production of water. Usually the amount of water at the early stage of production is small but with time the amount of produced water increases as oil rate declines as shown in Figure 1.2. Formation water when produced with crude oil through the production system can form unstable, stable or very stable emulsion. These emulsions form because of a combination of factors such
as the presence of the crude oil, water, shearing energy and surface active substances present in the crude oil. Several factors are known to influence the stability and viscosity of the emulsion such as the presence of heavy polar fractions, viscosity of the base oil, temperature, degree of mixing, pH, salinity, droplet size and droplet size distribution. Several researchers have attributed the stability of emulsion to the presence of asphaltenes and resin with little consideration given to the presence of inorganic acid and bases also present in the crude oil. This research attempts to correlate the acid and base number with viscosity and stability. Also, several emulsion studies have been carried out using external surfactants to stabilize oil emulsion in situations where the oil does not form stable emulsions itself. Little attention has been given to understanding the stability and viscosity behavior of these emulsions using different surfactant.

### 1.5 Research Scope and Objectives

The objective of these research is to study the stability and rheology behavior of surfactant stabilized water-in-crude oil emulsions. This is important because when studying the flow of emulsion in reservoirs and pipelines natural surfactant present in the crude influence the stability and rheological properties. To be more specific, first an emulsification procedure will be developed and the effectiveness of different surfactant in stabilizing an emulsion will be investigated. The most effective surfactant is investigated further at a specific water fraction and later at different water fractions. Also, the effect of changing surfactant concentration on the emulsion viscosity, stability, total acid and base number is investigated. Finally, attempt is also made to relate the total acid and base number to the emulsion relative viscosity and stability.
All of the emulsions to be investigated will be water-in-oil emulsion. Dead oil samples at low pressure with no dissolved gases will be used to prepare the emulsion. Deionized water is used throughout the experiment. External surfactants will be used to stabilize the emulsion except in the base case for comparison. All emulsion emulsions will be prepared at room temperature (22-25°C) and atmospheric pressure (14.7psi)
CHAPTER 2
THEORECTICAL BACKGROUND

2.1 Emulsion

Emulsions are colloids in which one phase is dispersed in another phase. The dispersed phase is called the internal phase while the other phase is called the external or continuous phase. In order to have an emulsion one of the two liquid phases must be dispersed in the other as small droplets. The droplet of the dispersed phase is suspended in the continuous phase because of the resistance of the droplets to coalesce and separate. The resistance to separate (stability) is usually because of the presence of agents at the interface of the two phases. Emulsions are commonly encountered in our daily life such as in milk, yoghurt, mayonnaise, butter, cream, paints, pharmaceutical etc. To prepare an emulsions oil, water, surfactant, and energy are needed (Tadros, 2013). Each of these components play a significant role in the formation, type, stability and properties of the emulsion. Emulsions are generally classified as either macro, micro or nano emulsions depending on the particle size of the dispersed phase. When the dispersed droplets are larger than 0.1μm, the emulsion is a macroemulsion (Schramm, 1992). Microemulsion have dispersed particle size less than 10nm and are thermodynamically stable. Commonly encountered emulsions are macroemulsions which are thermodynamically unstable but kinetically stable. They are thermodynamically unstable because the contact between the two phases i.e. water and oil molecule is unfavorable hence they will always break down over time. This break down results in free energy needed to create the interfacial tension (Tadros, 2009).
Figure 2.3: Process of forming an Emulsion: water, oil, emulsifier and mixing are needed

2.1.1 Types and Classification of Emulsions

Emulsions are generally classified as macro and micro emulsions depending on the droplet size of the dispersed phase. There are four different types of emulsion depending on which of the two phases is the dispersed phases. The different types are water-in-oil emulsion (W/O – water droplet dispersed in oil phase), oil-in-water emulsion (O/W-oil droplet dispersed in water), water-in-oil-in water (W/O/W) and oil-in water-in-oil (O/W/O). The different types are shown in Figure 2.2. The last two are called multiple emulsion because an emulsion is dispersed in another phase. There is also another type of emulsion called Pickering emulsion. These are emulsions stabilized by the adsorption of small solid particles at the oil-water interface. Whether the emulsion is W/O or O/W depends on the contact angle of the adsorbed particle.

2.1.2 Mechanism of Emulsification

For a stable emulsion to be formed, four components are need: oil, water, surfactant and agitation (or mixing). Oil and water when mixed separate creating an
Forming an emulsion is not spontaneous, energy in the form of mixing is needed to break down the interface and allow one of the two phases to form dispersed droplets. Adding surfactants helps to lower the interfacial energy and allow the creation of smaller droplets. Mixing can be done simply by shaking the mixture with hand or by high speed stirrers. In order to form nanoemulsion, a very high amount of energy is needed.

2.2 Role of Surfactant in Emulsification

Surfactants are very important in the formation of an emulsion. They are surface active agents because they are adsorbed at the oil water interface where they lower the interfacial tension and are therefore able to minimize the energy required to create emulsions. They also form interfacial films around the droplets. These interfacial films aid the stability of the emulsion by acting as barriers to coalescence, flocculation and settling of the droplets. Surfactants are able to perform their function because of the nature of their chemical structure. Surfactants are amphiphilic meaning they consist of
both a non-polar hydrophobic tail (water hating) and a polar hydrophilic head (water loving) as seen in Figure 2.5. The head and tail of the surfactant align themselves in the phases in which they are soluble. The surfactant molecules align themselves at the interface until the critical micelle concentration (CMC). At CMC, the surfactant molecule with a little mixing begin to form micelles around the dispersed droplets as shown in Figure 2.6

![Figure 2.5: Typical Structure of a Surfactant (Petroleum Engineering Handbook)](image)

There are two different classes of surfactant based on their behavior in aqueous solution: Ionic (anionic and cationic) and Nonionic. Anionic Surfactants are dissociated in water in an amphiphilic anion, and a cation (which in general is an alkaline metal or a quaternary ammonium). Examples include lauryl sulphate, alkylbenzene sulfonates, soaps etc. Cationic Surfactants dissociates in water into an amphiphilic cation and an anion (mostly halogen). Non-ionic surfactant do not dissociate in water. Their hydrophilic group are either esters, alcohol, phenol, ether or amide (Salager, 2002)
Figure 2.6: Surfactant molecules align themselves along the oil-water interface (a) 
Formation of micelles around oil or water droplet (b) (Henríquez, 2009)

2.2.1 Hydrophilic Lipophilic Balance (HLB)

The HLB of a surfactant indicate the relative proportion and size of the hydrophilic to the lipophilic group. In the past selecting surfactant for a particular purpose was done by conducting lots of experimental trial and error i.e. empirical. The HLB is a semi empirical method developed by W.C. Griffin (Griffin, 1949) to help in selecting surfactants. Every surfactant is given an HLB number based on the hydrophilic and lipophilic group it has. An emulsifier that is lipophilic in character is given a low HLB number (below 9) while surfactants that is hydrophilic in nature is given a high HLB number (above 11). The HLB is number between 0 and 20. Surfactant with low HLB number are more oil soluble and tend to form water-in-oil emulsions while those with high HLB number are more water soluble and tend to form oil-in-water emulsion (ICI Americas Inc, 1984) see Table 2.1
Table 2.1: Different HLB number of Surfactants and their Application

<table>
<thead>
<tr>
<th>HLB range</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>3–6</td>
<td>W/O emulsifier</td>
</tr>
<tr>
<td>7–9</td>
<td>Wetting agent</td>
</tr>
<tr>
<td>8–18</td>
<td>O/W emulsifier</td>
</tr>
<tr>
<td>13–15</td>
<td>Detergent</td>
</tr>
<tr>
<td>15–18</td>
<td>Solubilizer</td>
</tr>
</tbody>
</table>

This is in accordance with the Bancroft rule which states that the phase in which the surfactant is more soluble is usually the continuous phase. HLB values are additive. The HLB value of a surfactant mixture is the weighted average of the HLB values for each component as shown below

\[ HLB = x_1 HLB_1 + x_2 HLB_2 \]

The HLB number required to emulsify a given oil depends on the nature of the oil. Different oils have predetermined required HLB number as shown in Table 2.2. The HLB of the surfactant are chosen to match the required HLB of the oil for good emulsification. Required HLB of any oil can be determined experimentally.
Table 2.2: Required HLB for Emulsion of different Oils

<table>
<thead>
<tr>
<th>Oil</th>
<th>W/O emulsion</th>
<th>O/W emulsion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paraffin oil</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>Beeswax</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Linolin, anhydrous</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>Cyclohexane</td>
<td>—</td>
<td>15</td>
</tr>
<tr>
<td>Toluene</td>
<td>—</td>
<td>15</td>
</tr>
</tbody>
</table>

2.3 Breakdown Processes in Emulsions

Most emulsions are only kinetically stable and therefore separate into their different phases in a number of different ways such as sedimentation, creaming, aggregation and coalescence as illustrated in Figure 2.7. Sedimentation and creaming occur in emulsion if there is a significant difference between the density of the dispersed and continuous phase. These two physical instability mechanisms result from external gravitational or centrifugal forces pulling the dispersed phase either upward or downward. When the dispersed phase is less dense than the continuous phase (as in O/W emulsion) the droplet separates to the top (creaming) while if the dispersed droplets are denser (as in W/O) the droplets settles to the bottom in a sedimentation process (Pena, 2004). The rate of sedimentation of droplets is governed by stoke’s law at low concentration.

2.4 Water-in-Crude Oil Emulsion Stability Factors

Water-in-crude oil emulsions are the most common type of emulsion encountered in the oilfield. Over the years several researchers have done a lot of work to understand the factors responsible for their formation, stability and rheology. Bottle test, dielectric, rheology and nuclear magnetic resonance methods have been used extensively to study the behavior of emulsion. Opawale A. et al (2013) carried out a number of experimental
laboratory test to study the effect of shear and emulsifier on the stability of oilfield emulsions at different water fractions. Asphaltenes, reservoir fines and inorganic solids were used as emulsifiers. They found that increasing the water fraction, asphaltene and shearing energy lead to the formation of tight emulsion.

Abdurahman et al (2008) observed in their experiment that high resin to asphaltene ratios decreases the water-in-oil emulsion stability. They also conclude that emulsion stability increases with increasing surfactant concentration and decreases with increasing temperature.

Silset (2008) showed that emulsion stability showed a temperature dependence according to Arrhenius equation. He also observed a very strong correlation between emulsion stability and viscosity; an increase in viscosity causes an increase in stability.

**Figure 2.7 : Emulsion Destabilization Mechanisms (Tadros, 2013)**
(Henríquez, 2009) studied water in oil emulsion using paraffin oil and a mixture of sorbitan monooleate (span 80) and nonylphenol ether (Arkopal N 040) as surfactant. The surfactant stabilizes the emulsion in a wide range of water content. He observed that the highest stability of emulsion at the highest surfactant concentration (10%).

### 2.5 Role of Heavy Polar Fractions in Emulsion Stabilization

Emulsion are stabilized by surface active components present in the crude oil. These components are heavy polar fractions which includes asphaltenes, carboxylic organic acids, bases, and fine inorganic particles (Kilpatrick, Water-in-Crude Oil Emulsion Stabilization: Review and Unanswered, 2012). Some of these components are identified by carrying out a Saturates, Aromatic, Resin and Asphaltenes (SARA) analysis on the crude oil. Several authors have attributed the formation of very stable emulsion to the presence of asphaltenes. Asphaltenes are complex molecules soluble in toluene, benzene and ethyl acetate but insoluble in low molecular weight n-alkanes (Kokal and Sayegh., 1995). Michell and Speight (1973) found that asphaltenes exist as colloidal suspension and aids stabilization by the presence of resin adsorbed on their surface. Asphaltenes with resin adsorbed on their surface form micelles around their dispersed droplets. These micelles acts as rigid films exhibiting steric repulsion as shown in Figure 2.6.
Strassner (1968) demonstrated through a series of experiment that removing asphaltene from crude oil resulted in formation of mobile film and unstable emulsion. Adding the asphaltene resulted in the formation of rigid films and very stable emulsion.

Eley et al (1998) found out that the degree of emulsification depended on the ratio of aromatic component to aliphatic component. According to them the tendency to form very stable emulsion decreases with increasing aromatic content in the crude oil.

McLean and Kilpatrick (1997) showed that the stability of water-in-crude oil emulsions was related to the asphaltene precipitation point. The most stable emulsions occurred when the asphaltenes were on the verge of precipitation.

Sjöblom et al (1990) adsorped the interfacial active fraction in ten different Norwegian continental shelf crude oils and found out that all of the adsorped crude were incapable of stabilizing water-in-crude oil emulsions. They measured stability by observing separation of water over time.
Kilpatrick (2012) in his review of water in oil emulsion stabilization noted that asphaltene is a complex mix of chemically distinct molecules many of which actually are not soluble in toluene and destabilize emulsion. According to him many of the molecules in crude oil are acidic and can ionize at the oil-water interface lowering the interfacial tension and stabilize the emulsion. Some of these acids are simply carboxylic acids, naphthenic acids and some are fused aromatic acids.

2.6 Total Acid and Base Number (TAN & TBN)

Asphaltene is a complex mix of different compounds, it consists of a continuum of high molecular weight compounds with different functional groups. Present in crude oil are also acid and bases functional groups. The interaction between the different functional asphaltene and acid and base functional group is central to emulsion stability (Tanja et al, 2005). In addition to the other heavy polar fractions found in crude, some crude oils are known to also contain significant amount of acids and bases. These acids are in the form of what is called naphthenic acids. Although the term “naphthenic acids” originally implied that the acids contained naphthenic rings, today it comprises cyclic, acyclic, and aromatic acids in crude oils (Brient et al, 1995). They are a complex mixture of compounds with diverse molecular weight and structures. Recently there has been an increasing interest in studying how these acids and bases interact with other crude oil heavy fractions to stabilize emulsions. The complex interactions of asphaltenes and naphthenic acids is an area of research that is getting a lot of attention. Pål V. Hemmingsen et al (2006) carried out an experimental in which they washed a North Sea acidic crude oil to reduce the total acid number using liquid-liquid extraction. The original and washed crude oil were characterized using electrospray ionization
Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR MS) and by Fourier transform infrared spectroscopy. Each of the washed crude was used in forming an emulsion and the stability was measured using the critical electric field method. They observed that reducing the acid component in the crude increases the interfacial tension and also increases the water-in-oil stability. They concluded that acidic fractions in crude oil has a solubilizing effect on asphaltene hence removing them causes the asphaltene to be more surface active.

Tanja Barth et al (2005) found a very strong correlation between the total base number and asphaltene content of twenty crude oils. They suggested that there could be a very strong interaction between acids and bases in asphaltene emulsion stability.

2.7 Emulsion Rheology and Viscosity Measurement

2.7.1 Rheometry

Rheometry refers to the measurement of the rheological properties of a fluid. Rheology is defined as the study of the deformation and flow of fluids under the influence of an applied shear stress. Stress is an internal reaction to an external force. Viscosity is a very important rheological property of a fluid that measures the internal resistance to stress or it is a measure of the fluid friction. Viscosity usually cannot be measured directly, forces, torque and rotation speed are measured and then converted to viscosity models. Viscosity in a very simple term is a ratio of shear stress to shear strain.

Viscosity can be measured either by using capillary tube viscometers or rotational viscometer.

Capillary tube viscometers work on the principle of fully developed laminar flow described by the Hagen-Poiseuille equation. Glass capillary viscometer are the simplest
types. They have two bulbs connected in a U-shaped geometry. The kinematic viscosity is given by the equation below

\[ \vartheta = \left( \frac{4n}{3n + 1} \right) \left\{ \left( \frac{\pi g(z_1 - z_2)}{128L\nu_o} \right) D^4 \right\} (\Delta t - K\Delta t) \]

For glass viscometer \( n = 1 \), for large L/D ratio viscometer, the last correction term \( K \) is negligible (Leblanc et al, 2000). Therefore, the kinematic is proportional to the efflux time. Every glass capillary viscometer comes with a constant equivalent to the second term in bracket. They are simple, inexpensive but for low viscosity.

Rotational viscometers are the most widely used viscometer because they offer the ability to change the shear rate and also work continuously at given rate. In rotational viscometer the fluid is sheared continuously between two surfaces. Rotational viscometers can either be stress or rate controlled. In stress controlled viscometer a constant torque is applied to generate rotation which can be converted to a shear rate. In rate controlled viscometer, a constant shear speed is applied to generate a measurable torque (Silset, 2008). There are different configurations of the rotational viscometer depending on the geometry of the rotating and stationary plates. The most common are the concentric cylinder, cone and plate and parallel plate as shown in Figure 2.9. The rotational viscometer offers the advantage that they are easy to use, they have a wide range of viscosity they can measure, can be used for nonsettling suspension and they can be adopted for use at high temperature and pressure. The disadvantage with this type of viscometer is that prolong shearing of the fluid can sometimes alter the properties of the fluid that is measured. Temperature increase due to prolong shearing is another major disadvantage (Pal et al, 1992)
2.7.2 Emulsion Rheology

Emulsion exhibit both newtonian and non-newtonian behavior. A fluid is said to be Newtonian when it obeys Newton's law of viscosity that is the shear stress is proportional to the shear rate. In another form we say the viscosity is independent of the shear rate. If the viscosity is a function of the shear rate, then the emulsion is non-newtonian. Non newtonian fluid can be broadly categorized into time-independent and time-dependent non newtonian fluid. Time independent non-newtonian fluid can either be pseudoplastic (shear thinning) or dilatant (shear thickening) as shown in Figure 2.8. Time dependent non-newtonian have viscosities both a function of shear rate and shear rate history. They are broadly categorized either has thixotropic or rheopectic fluid.

The viscosity of an emulsion depends on a number of factors such as the viscosity of the continuous phase, volume fraction of the dispersed phase, temperature, shear rate (for non-newtonian), pressure, droplet size and distribution of the dispersed phase. Several
researchers have found out that the viscosity of an emulsion can be significantly higher than the viscosity of the constituent crude oil and water (Rønningsen, 1995; Johnsen, 2003; Kokal, 2003). Hans Petter Rønningsen (1995) investigated the apparent viscosity of eight different North Sea crude oils emulsion and found that the viscosity increases linearly with water fraction at low water cut. At higher viscosity, he found that the viscosity increases more rapidly. Most of the emulsion he investigated were moderately non-newtonian even at high water cut.

(Mohammed, 2009) created water-in-oil emulsion using crude oil from Gulf of Mexico, 3.5wt% brine and no external surfactant. He found that at 50% water fraction, the emulsion viscosity decreased tenfold from 7°C to 80°C. Also the viscosity of the emulsion increases by almost two order of magnitude with higher water cut, the highest stable emulsion was also observed at 60% water cut. Pal (2000) in his study of several oil-in-water emulsion using stress controlled rheometer. The emulsion exhibit newtonian behaviour at low to moderate water fraction. At higher water fraction, the

Figure 2.10: Rheological Behavior of Fluids (Kumari, 2016)
emulsion exhibit shear thinning behavior. He also observed that the viscosity of the shear thinning emulsion is strongly affected by the droplet size.

2.8 Phase Inversion

Phase inversion refers to the changing of an emulsion dispersed phase to a continuous phase. For example, a water-in-oil emulsion can undergo a phase inversion to oil-in-water emulsion. This change can occur with time or a change in condition of the emulsion. Phase inversion is usually accompanied with the formation of multiple emulsion (Tadros, 2009). There are two types of phase inversion: transition inversion and catastrophic inversion. Catastrophic inversion occurs when the water fraction in an emulsion is increased up to a point such that the droplet packing is close to the critical packing factor. Transition inversion occurs when conditions such as the surfactant, salinity or temperature changes.

2.9 Emulsion Viscosity Correlations

There are several correlations that have been developed by researchers to predict the viscosity of emulsion. A lot of these correlations are empirical or semi-empirical correlations. Several factors affect the viscosity of emulsion such as: viscosity of the continuous phase, water fraction, temperature, shear rate, nature and concentration of the emulsifying agent, droplet size and distribution and presence of solids in the crude oil. It is almost impossible to incorporate all of these factors into one correlation so several correlations only consider the effect of water fraction. A few take into account the effect of water fraction, temperature and shear rate. Correlations for predicting the viscosity of emulsions are usually represented in term of relative viscosity (Rønningsen, 1995). Relative viscosity is the viscosity of the emulsion divided by the
viscosity of the base oil. One of the most basic correlation for predicting emulsion viscosity was proposed by Einstein (1906) when he studied suspension of nondeformable solid particle at low concentration. He expressed the relative viscosity linearly with volume fraction of the dispersed phase as follows:

\[ \eta_r = 1 + 2.5\phi \]  

(1)

At higher water fraction, emulsion viscosity exhibits a non-linear behavior with water fraction. (Richardson, 1933) proposed an exponential dependence of viscosity on water fraction as follows:

\[ \eta_r = e^{K\phi} \]  

(2) where K is Richardson constant which is dependent on the system. Broughton and Squires (1937) introduced another system correction factor (A) into the Richardson equations as follows

\[ \eta_r = Ae^{K\phi} \]  

(3)

\[ \ln \eta_r = \ln A + K\phi \]  

(4)

(Rønningsen, 1995) extended the constant A and K to be a linear function of temperature as shown below

\[ \ln \eta_r = k_1 + k_2T + k_3\phi + k_4T\phi \]  

(5) where T is the temperature, \( \phi \) is water volume Percent (not fraction), \( k_1, k_2, k_3 \) and \( k_4 \) are shear rate dependent (values given in the literature). The correlation was developed using eight different North Sea crude without the addition of any emulsifier. This correlation is oil type specific and it is the only correlation that does not require experimental tuning. It is also noticed that although being developed for gas-free systems, the correlation is valid also for live oil multiphase systems at pressures up to 100 bars (Zahra, 2012)
(Mooney, 1951) developed a correlation from studying viscosity of monodisperse spheres:

\[ \ln \eta_r = \frac{2.5 \phi}{1-\lambda \phi} \quad \ldots \ldots \ldots \ldots (6) \]

where \( \lambda \) is a constant to be used to tune experimental data. This correlation is sensitive to the turning point (Johnsen & Ronningsen, 2003).

(Pal, 2000) using a large set of experimental data developed a two parameter emulsion viscosity correlation as shown below

\[ \eta_r^{-2/5} \left[ \frac{2 \eta_r + 5K}{2+5K} \right] = 1 - K_\phi \phi \quad \ldots \ldots \ldots \ldots (7) \]

Where K is the ratio of dispersed phase viscosity to continuous phase viscosity, \( \phi \) is the water fraction of the dispersed phase. \( K_\phi \) is takes into account the effect of adsorbed surfactant. In absence of any data use 1.35 (inverse the water fraction equivalent to maximum packing factor, 0.74) (Zahra, 2012). This correlation is also very sensitive to the turning point. The correlation was developed using a combination of polymer thickened oil-in-water emulsion, mineral oil-in-water emulsion, oil-in-water containing milk fat, heavy oil-in-water emulsion and petroleum oil-in-water emulsion.

Pal & Rhodes (1989) developed a correlation to predict the viscosity of both water-in-oil and oil-in-water emulsion. They presented both the empirical correlation and the correlation model. They observed good agreement between the model and the correlation. The oil used to form their emulsions are bayol-35/CCl\(_4\) oil mix, shell vitrea 220, Diesel No.2 and Furnace No.2. The surfactant used were Triton X100 (0.5-1\%vol), Span80 (1.5-2\%vol), Span 85 (2\%vol). All emulsions were prepared at 23°C-29°C. Their correlation is presented below:
\[ \eta_r = \left( 1 + \frac{\phi/K_2}{1.1884 - \phi/K_2} \right)^{2.5} \]  

\( \Phi \) is the dispersed phase volume fraction. \( K_2 \) is the dispersed phase concentration at which the relative viscosity is equal to 100. \( K_2 \) accounts for system variation and is used as a fitting parameter. It is an experimental value at a high dispersed phase volume fraction.

Almrit et al (2011) noted that Ronningsen corelation does not reduce to a relative viscosity of one (that of the base oil) when the water fraction is zero. Their correlation also put into consideration the effect of oil type

\[ \ln \eta_r = \phi \ast (c_1 + c_2 T) \ast \left[ \ln(API) \right]^{c_3 + c_4 T} \]  

c_1-c_4 were defined at low shear rate (<100s\(^{-1}\)) and high shear rate (>100s\(^{-1}\)).
CHAPTER 3
EXPERIMENTAL PROCEDURE

3.1 Experimental Techniques

In this chapter, I will discuss the experimental setups as well as the workflow technique I have employed in carrying out these research. I will also discuss the materials used. The first stage in these research was to create stable synthetic emulsions in the lab with or without the use of an external surfactant. In order to create stable and reproducible emulsion it is important to have an emulsion mixing procedure. Different mixing methods were investigated and reported. Crude oil emulsion without any external surfactant was created and checked for stability and viscosity. A number of surfactants were then used to stabilize the emulsion while observing their effect on viscosity. A surfactant scan was carried out to identify the best surfactant in terms of stability and viscosity. After identifying the best method to create the emulsions, the method was tested for repeatability by creating multiple emulsion using exactly the same method and measuring the stability and viscosity. Stability measurements were carried out using both bottle test and centrifuge. Viscosity test was done using capillary viscometer and Fann concentric cylinder rotational viscometer. The total acid number (TAN) and total base number (TBN) of the crude were measured using Metrohm oil titrino plus potentiometric titrator. The effect of adding external surfactant was investigated on the TAN and TBN and their associated emulsion stability and viscosity.

3.2 Materials

The main crude oil used in this research was obtained from a wellhead in Midland Texas (Texas crude oil). A second crude oil sample (Crude oil A) was also used in some cases for comparison. Mineral oil obtained in the lab was also used during the initial
stage of the research to create emulsions. The water used in preparing the emulsions is deionized water. Bulk properties of the Texas crude, mineral oil and heptane are presented in Table 3.1. Density was measured using a pycnometer while viscosity was measured using capillary viscometer. The following surfactants were used: Span80, Span 85, Tween 80, TritonX100 and Tergitol. They were purchased from Sigma Aldrich. The properties of the surfactants are presented in Table 3.2. All surfactants are non-ionic. There was a need for surfactant because the available crude oil does not naturally form stable emulsion.

<table>
<thead>
<tr>
<th>Name</th>
<th>Density (g/cc)</th>
<th>API</th>
<th>Viscosity (cP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Texas Crude oil</td>
<td>0.884</td>
<td>27</td>
<td>19</td>
</tr>
<tr>
<td>Crude oil A</td>
<td>0.849</td>
<td>33</td>
<td>60</td>
</tr>
<tr>
<td>Mineral oil</td>
<td>0.867</td>
<td>-</td>
<td>32</td>
</tr>
<tr>
<td>Deionized water</td>
<td>1.000</td>
<td>10</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 3.1: Bulk Properties of Fluids Used at 25°C

<table>
<thead>
<tr>
<th>Name</th>
<th>ID</th>
<th>Chemical name</th>
<th>Molecular Weight (g/mol)</th>
<th>Density, g/cc @25°C</th>
<th>Viscosity, cP @ 25°C</th>
<th>HLB</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Span80</td>
<td>S80</td>
<td>Sorbitan monooleate</td>
<td>428.62</td>
<td>0.986</td>
<td>1000-2000</td>
<td>4.3</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Span 85</td>
<td>S85</td>
<td>Sorbitan trioleate</td>
<td>957.52</td>
<td>0.956</td>
<td>200-300</td>
<td>1.8</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Tween 80</td>
<td>T80</td>
<td>Polyoxyethylene sorbitan monooleate</td>
<td>1310</td>
<td>1.076</td>
<td>375-480</td>
<td>15</td>
<td>TCI America</td>
</tr>
<tr>
<td>Triton X100</td>
<td>TX100</td>
<td>Octylphenol Ethoxylate</td>
<td>625</td>
<td>1.061</td>
<td>240</td>
<td>13.5</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Tergitol</td>
<td>TERG.</td>
<td>Alcohol Ethoxylate</td>
<td>642</td>
<td>1.006</td>
<td>60</td>
<td>13.3</td>
<td>Sigma Aldrich</td>
</tr>
</tbody>
</table>

Table 3.2: Properties of the Surfactants
3.3 Experimental Work flow

Figure 3.1 gives the emulsification and measurement procedure taken throughout this research.

![Experimental Work Flow](image)

**Figure 3.1: Experimental Work Flow for emulsification and data gathering.**

In first stage, the surfactant is mixed with the crude oil. Viscosity, TAN and TBN of the oil are measured after mixing. Water is then gradually added while mixing to form emulsion. The emulsion viscosity is measured using Fann 35 rotational viscometer and/or capillary viscometer. Portions of the emulsions are also used for stability test as well as TAN and TBN. Details of each step is further discussed in the following sections.

3.4 Emulsification Protocol

Emulsions were formed by adding a known amount of oil (crude or mineral) with surfactant (agent-in-oil method). This method was used because most of the available surfactant were soluble in oil. Water was then added slowly while stirring the mixture. Oil and water fractions are quoted as volume percentages but in doing the oil and water
measurement their corresponding weight was used. Surfactant volume is a percentage of the total emulsion volume. Two methods were employed in creating the emulsion; batch method and continuous method. In the initial stage all emulsions were prepared using the batch procedure. In the batch procedure, each water fraction was prepared separately (measuring out each water and oil weight separately). In the continuous method, the lowest water fraction emulsion was prepared first. To prepare the next water fraction additional water was added to make up any volume of emulsion removed. This method was used when investigating emulsions at different water fraction. Continuous method helps to minimize the amount of crude oil used and also reduce waste emulsion.

3.5 Mixing Method

Five mixing methods was used in these research, most of which were either ineffective or time consuming. Methods used were handshaking, magnetic stirrer, mechanical 3-blade stirrer, VWR mini-incubating shaker and IKA UltraTurrax T18 mixer. Figure 3.2 shows the hand mixing and magnetic stirrer. Hand mixing simply involves shaking the oil and water mixture to form the emulsion. The mixing is done for a set time. Although this method produced stable emulsion it was difficult to reproduce. Also the method requires a lot of human effort to sustain continuous mixing over a long period.

Magnetic stirring involves placing the mixture on a magnetic stirrer and dropping a magnetic stirring bar in the mixture. The stirring bar are polytetrafluoroethylene (PTFE) coated magnetic cylindrical bars, so it does not interact with the liquids. Figure 3.3 shows the 3-blade mechanical stirrer and the VWR mini-incubating shaker. The mechanical stirrer is an Arrow 6000 model. It had three equally spaced 1” blade.
Figure 3.2: Left (Magnetic stirring bar) and Right (Hand shaking)

Figure 3.3: Picture of the 3-blade stirrer (left) and VWR mini-incubating shaker (right)

The mixing speed range was between 0-6000rpm. The blades were aligned to be at the center of the mixing container. Mixing with the mechanical 3-blade stirrer was limited to 1200rpm for 15minutes because of splashing.
The mini-incubating was purchased from VWR. It is a variable speed shaker with an orbit radius of 3mm, it also had a temperature and time adjustment control. The speed range was between 100-1200rpm. 125mL square bottles were used in each of the four mixing spot on the shaker. The shaker operates by vibrating about it orbit. Mixing with this method was carried out at 24°C at 1200rpm for 15minute.

The IKA UltraTurrax T18 is a digital high speed dispersing tool capable of producing emulsion with droplet size ranging from 1-10μm (when using dispersion shaft model S-18-19G). The speed range is from 500-25,000rpm and can mix up to 1.5L. **Figure 3.4** shows the picture of the mixer as well as the emulsion mixing setup. The dispersion is placed such that it offsets from the center to avoid vortex and air entrapment during mixing.

![Image of T18 mixer and schematic](source: IKA works manual)
A minimum distance of 10mm was maintained between the bottom of the mixing vessel and the bottom of the dispersing tool. The mixture is drawn axially into the dispersion head and then forced radially out through the rotor/stator arrangement as shown in Figure 3.5. High shear produced between the rotor and stator gap creates turbulence to provide optimum dispersion. Total emulsion volume prepared were 80cc, 150cc and 170cc depending on the measurements to be taken and the size of the mixing vessel. It was ensured that the fluids cover a hole at the bottom of the dispersing tool at all time.

Mixing was done at two speeds; 5000rpm for 30 minutes and 20,000rpm for 2 minutes.

![Figure 3.5: Operating principle of the dispersing tool of the IKA T18 mixer (source: IKA Works manual)](image)

### 3.6 Stability Test

Two methods were used to study the stability of emulsion; bottle test and centrifuge test. The popular bottle test involves putting sample of the emulsion in a container and allowing it to separate under gravity. The amount of water and oil separated over time is used to determine if the emulsion is stable or not. In this research, 40mL of emulsion samples were kept in a 50mL centrifuge bottle and allowed to separate over a period of 120 hours. Measurement of the oil and water separated were taken every 24 hours.
Centrifuge test is very similar to the bottle test except that in this case a centrifuge is used to speed up the rate of separation. Centrifuge used in this research is a Dynac Model No 420063 shown in Figure 3.6. The centrifuge has a 4-place 50mL horizontal head that can hold 4 centrifuge bottle at the same time. The speed is from 1000rpm at speed setting 24 to 2790rpm at speed setting 100. 12mL of the emulsion samples were put in 15mL centrifuge bottle and the speed setting set to 50. The centrifuge was allowed to run at time intervals 2, 3,5 (6 times) and 10 minutes for a cumulative of 45 minutes. After each time interval, the emulsion was taken out to record the volumes of oil and water separated.

Separation was used as a measure of stability. Separation was defined as the volume of oil and water separated divided by initial emulsion volume expressed as a percentage. The lower the separation percentage, the more stable the emulsion.

Figure 3.6: Dynac Centrifuge
3.7 Viscosity Measurement

Rheological measurements were carried out using a Cannon Fenske routine capillary viscometer (Figure 3.7) and Fann 35 rotational viscometer.

3.7.1 Capillary Viscometer

The capillary viscometer was used extensively in this research because it operates at a very low shear rate and does not change the properties of the emulsion through excessive shearing. Capillary viscometer size to be used were chosen based on the observed viscosity to be measured (each viscometer has a viscosity range and a viscometer constant). The larger arm of the capillary viscometer was filled until the large bulb was half filled (approximately 12mL).

Figure 3.7: Cannon Fenske Routine Capillary Viscometer
A suction bulb was used to draw the emulsion up the smaller bulb until it goes past the upper etched mark. A stop watch is then used to take the time it takes the emulsion to flow from the upper etched mark to the lower mark (the efflux time). The efflux time is multiplied by the viscometer constant to get the kinematic viscosity (in cSt). The dynamic viscosity (cP) is calculated by multiplying the kinematic viscosity by density of emulsion. Density of emulsion was calculated as weighted fraction of the density of the constituent oil and water. The concentration of the surfactant does not significantly affect the density of the emulsion therefore it was not included in the calculation.

Experimental measurement confirms the calculated density. If the calculated viscosity falls outside the range of that viscometer, the next higher viscometer size was used to measure the viscosity again.

It is important to note that the efflux time was taken in a single pass down the viscometer. Repeat measurements were only taken after washing the viscometer using a procedure described in the next section

3.7.2 Washing of the Capillary Viscometer

In order to ensure reproducibility of viscosity, it was important to have a procedure for washing the glass viscometers so that the wettability of the glass surface does not change with each measurement. The procedure used for washing involves; first the sample used is poured out into a waste container, then toluene is sucked up through the bulbs several times to dissolve any remaining oil. Tap water is thereafter passed through the viscometer until any remaining toluene is cleaned off. Finally, acetone is used to dissolve any remaining water before the viscometer is dried in a stream of dry air.
3.7.3 Fann 35 Rotational Viscometer

Newtonian and non-Newtonian behavior of some of the emulsions were measured using Fann 35 rotational viscometer with a cup containing 350mL of the emulsion sample as shown in Figure 3.8. It is a couette rotational viscometer with a stationary bob (B), rotating rotor (R), and a torsion spring (F). The fluid is sheared in the annulus space by rotor and this produces a torque on the bob that is transmitted through the torsion spring to a dial. An R1-B1-F1 combination was used in this research. The shear gap was 0.117cm. The viscometer operates at 6 different speeds (3,6,100,200,300 and 600rpm). The dial reading at speed 300rpm corresponding directly to the newtonian viscosity at that shear rate. Viscosity at other shear rate was calculated by multiplying the dial reading with some constants available in the operating manual.

Figure 3.8: Side and Front View of Fann 35 Rotational Viscometer
Before any measurement was taken the viscometer was checked with a 50cP (at 25°C) viscosity standard. To measure the viscosity, the dial reading was taken at all six speed starting with the lowest. Time effect was investigated by first operating at each speed for 2 minutes (total of 12 minutes for all six speeds) and then repeating the measurements again starting from the lowest speed for another 12 minutes and so on up to 60 minutes.

3.8 TAN and TBN Measurements

TAN and TBN correspond to the mass of potassium hydroxide (KOH) needed to neutralize the acids contained in 1 g of the crude oil. Whereas the TAN uses a base as the titrant (KOH), the TBN uses an acid (perchloric acid) as the titrant. The amount of acid used in the TBN is converted to an equivalent KOH mass. Both TAN and TBN are expressed in mgKOH/g.

Acid and base fraction present in the crude oil and its emulsion were measured using a Metrohm autotitrator (model 848) oil tritino plus with a solvotrode electrode shown in Figure 3.9. The setup consists of the control unit on which the exchange unit slides on. There are two exchange units: one for TAN and the other for TBN. Each exchange unit has its own separate titrant bottles. The stirrer and the electrode are mounted on an adjustable stand. Two dosing units are also available for adding the solvent.

Standard ASTM D664 and D2896 procedure were followed for TAN and TBN measurement respectively. The titrator was configured to follow these procedures. TAN measurement makes use of the Monotonic Equivalence point titration method (MET). In MET titration, the reagent is added in a constant volume step. Before carrying out any TAN measurement, the TAN of the solvent is first determined and stored in the control unit memory. This volume is
Figure 3.9: Metrohm Oil Tritino 848 Autotitrator for TAN and TBN

subtracted from the titrant volume of any measurement made to correct for the solvent used to dissolve the sample. To carry out TAN measurement, 10g of the sample (crude oil or emulsion) in a 150mL glass beaker was dissolved with 100mL of TAN solvent. TAN solvent is a mixture of toluene (50%) and water (0.5%) dissolved in Isopropyl alcohol (49.5%-IPA). This solvent was also obtained from Metrohm as a standard solution. The solvent dissociates the acid in the crude oil or emulsion. The mixture was titrated against the titrant which is a standard alcoholic KOH solution (0.1mol KOH/L IPA) obtained from Metrohm. The mixture is continuously stirred and the electric potential detected by the electrode. The end point is determined by plotting the electric potential against titrant volume and by using some endpoint recognition criteria (ERC; calculated using Fortuin method) to generate a cumulative plot of ERC against titrant
volume. The titrant volume at the highest peak on the cumulative ERC plot gives the endpoint volume. Typical result plot for TAN is shown in Figure 3.10

**Figure 3.10: Typical TAN plot showing Endpoint Recognition Criteria**

TBN measurement is carried out in a similar way except that the method used is the Dynamic Endpoint Titration method (DET). In DET method the reagent is added in variable volume steps. The volume steps vary as a function of the slope of the curve. The TBN solvent is a 50%-50% mixture of toluene and glacial acetic acid. The TBN solvent was prepared in the lab. The titrant for TBN is a 0.1mol/L of perchloric acid in glacial acetic acid. To ensure reproducibility, at least two measurements were made for both the texas crude oil and crude oil A.
CHAPTER 4
RESULTS AND DISCUSSION

This chapter presents and discusses the results of the research

4.1 Mixing method

Four different mixing techniques were used in this research: handshaking, magnetic stirring, VWR incubating mini shaker, mechanical 3-blade stirrer and IKA Ultra-Turrax mixer. Details of their operation have been given in the previous chapter. The effectiveness of each method was tested in terms of stability of the emulsions. The stability results for hand mixing, magnetic stirrer and mechanical stirrer are presented in Figure 4.1. The emulsions used were prepared using 60% of texas crude oil, 40% deionized water, 7% Triton X100 mixed for 15 minutes.

Figure 4.1: Effect of mixing method on emulsion stability; mixing methods used in this case are handshaking, magnetic stirrer and mechanical stirrer

The best method is the method that results in the most stable emulsion after undergoing the centrifuge stability test for 45 min at 1600 rpm. Figure 4.1 shows the handshaking to be the most effective. Magnetic and mechanical stirrer emulsion showed almost 100%
separation after a few minutes. The reason for this is because hand mixing allows the bulk of the fluid to be agitated while magnetic and mechanical stirrer can only produce localized mixing because of the orientation of their blade. They only mix the fluids at the level of their blade orientation in the emulsification vessel. Sometimes after mixing an oil layer could be seen at the top. Hand mixing, even though effective can only be used to mix very small amount of sample and requires a lot of effort to sustain mixing for long period hence it use was discontinued. VWR mini incubating shaker could not produce stable emulsion because it only vibrates. The orbit radius was too small to provide complete mixing of the fluids. There were four spots on the mixer and each spot produced very different emulsion in viscosity and stability i.e. the emulsions were not homogenous. The IKA UltraTurrax T18 mixer is a digital high speed mixer that was able to produce homogenous and repeatable emulsions depending on the speed. The effectiveness of the mixer was tested at two different speeds of 5000rpm for 30min and 20,000rpm for 2min. Figure 4.2 shows the viscosity of the emulsions prepared using span 85 and span80 surfactant and speed 5000rpm and 20,000rpm respectively. The emulsion water fraction was 30%. The emulsions were prepared in duplicate to test repeatability. It can be seen that operating at 5000rpm created emulsion with different viscosity just after mixing. Also, the viscosities were not repeatable after 48hours. There was also a very significant increase in temperature (10°C-15°C) when operating at 5000rpm for 30min. The emulsion also was not completely homogenous because different portion of the
emulsion consistently had different viscosities. However, operating at 20,000rpm for 2 minutes produced homogeneous and repeatable emulsion as shown by the overlapping viscosities of the emulsions. Operating at this condition was able to create turbulence to ensure proper mixing, the temperature increase was also low (about 5°C increase).

Going forward, all emulsion was prepared using the IKA UltraTurrax T18 mixer at 20,000rpm for 2 minute. It is important to note at this point that all emulsions were prepared at room temperature (25°C) and atmospheric pressure.

4.2 Stability testing

Stability is defined as the resistance of the emulsion to separation. In this research separation is used as a criterion to define whether an emulsion is stable or not.

Separation was defined as the total volume of oil and water separated ($V_s = V_o + V_w$) divided by the total volume of sample (Figure 4.3). Low separation percentage means the emulsion was stable while high separation percentage meant the emulsion was unstable. For the purpose of this research stable emulsion was defined as
Figure 4.3: Span85 stabilized emulsion and stability measurements

that with less than 10% separation after 45 minutes of centrifugation. Stability testing
was done using two independent methods. The bottle test involves physical observation
of oil and water separated from the emulsion over 5 days. Centrifuge test also involved
physical observation of the emulsion after been centrifuged at 1600rpm at time interval
between 1-45 minutes. Figure 4.4 shows the stability result for both the centrifuge and
the bottle test.

Figure 4.4: Stability result for both centrifuge (left) and bottle test (right). All emulsions were prepared using 70% oil, 30% deionized water and 1% surfactant
For comparison both tests were done at room temperature. Figure 4.4 shows that both methods have the same stability profile i.e. the span85 is more stable than TX100 and blank emulsion. Based on the research conditions, these is considered to be good reproducibility. However, there seems to be much faster separation in the centrifuge than the bottle test because of the applied centrifugal force. Bottle test is the ideal test because the emulsions are allowed to separate naturally under gravity but it takes several days before results are obtained. Since the profile are similar, going forward the stability of the emulsions will be tested using centrifuge.

4.3 Viscosity Measurement

Viscosity measurement was done using capillary viscometer extensively. In some cases, rotational viscometer was used to investigate newtonian and non-newtonian behavior of emulsion. Capillary viscometer measures the efflux time for the emulsion to move past two etched marks on the glass viscometer. The efflux time is multiplied with the viscometer constant to get the kinematic viscosity.

4.3.1 Reproducibility of Viscosity Measurement

![Figure 4.5: Viscosity of Texas crude oil and its emulsion taken in duplicate or triplicate showing high reproducibility of viscosity measurement. All emulsions were prepared using 70% oil, 30% deionized water and 1% surfactant](image)
Figure 4.5 shows that viscosity measurement using capillary viscometer is highly reproducible. The emulsions were prepared using 70% vol texas crude oil, 30% deionized water and 1% surfactant. Blank is the emulsion of the crude oil without any surfactant. The average standard deviation in viscosities for each sample is ±0.45cSt for texas crude oil based on three measurements, ±0.62cSt for blank emulsion based on two measurements, ±0.63cSt for TX100 emulsion based on two measurements and ±1.15 for span 85 emulsion based on three measurements. The efflux time was measured in a single pass through the viscometer. Each repeated measurement was taken after washing the viscometer using the standard procedure discussed in the previous chapter.

4.3.2 Newtonian and Non-newtonian behavior

The rheology of both TX100 stabilized mineral oil emulsion and Span 85 stabilized texas crude oil emulsion was studied using Fann 35 concentric cylinder rotational viscometer. The viscosities were measured both as a function of shear rate and time. Figure 4.6 shows the viscosity of the pure mineral oil as well as 70% mineral oil and 30% deionized water stabilized with 1% TX100. There was no appreciable change in the viscosity of the mineral oil with shear rate hence it is non-Newtonian. However, the viscosity of the emulsion reduces with shear rate (pseudoplastic or shear thinning behavior). Figure 4.5 also shows an overlapping of viscosity plot at different time interval indicating that the emulsion behavior is time-independent. This means that the viscosity of the emulsion is independent of the shear history i.e it is neither rheopectic or thixotropic.
Figure 4.6: Viscosity of 70% mineral oil - 30% deionized water and 1% TX100 emulsion using Fann 35 rotational viscometer. The Viscosity has been plotted as a function of shear rate and time.

Result for the effect of changing shear rate on the texas crude oil emulsion is presented in Figure 4.7. The two points to the left represent CV-Initial and CV-Final (CV means capillary viscosity). The initial capillary viscosity reading was taken immediately after emulsion preparation while the final reading was taken using the Fann 35 sheared sample.

Figure 4.7: Viscosity of 70% texas crude oil - 30% deionized water and 1% Span 85 emulsion using Fann 35 rotational viscometer and capillary viscometer. The Viscosity has been plotted as a function of shear rate and time. CV means Capillary Viscosity.
It can be seen from Figure 4.7 that the slope of the crude oil line and the emulsion are the same. The texas crude oil and its emulsion are therefore both Newtonian (viscosity does not change very much with shear rate) and time independent (viscosity plot overlap at different time intervals) at 30% vol. water fraction and 1% surfactant concentration. Because the emulsion is non-newtonian the viscosity measured using the Fann 35 and capillary viscometer are very close despite shearing the sample for about an hour.

4.3.3 Comparison between Capillary and Rotational Viscometer

Table 4.1 shows the capillary viscosity for both the mineral and texas crude oil emulsions before and after undergoing Fann 35 viscosity measurement. Shear rate data for the capillary viscometer was derived by interpolating between efflux time data provided by the manufacturer. It can be seen from the table that capillary viscometer is a low shear rate viscometer. For the mineral oil, the initial capillary viscosity is between the lowest shear rate viscosity on the Fann 35 (500cP and 750cP at 10s\(^{-1}\) and 5s\(^{-1}\) respectively). Final capillary viscosity shows a drop in viscosity indicating that prolong shearing affects the viscosity of the mineral oil emulsion. This supports the earlier conclusion that the emulsion exhibits a shear thinning behavior.

Table 4.1: Capillary Viscosity (CV) of both Mineral oil and Texas crude oil emulsions measured before and after shearing in a Fann 35 Viscometer

<table>
<thead>
<tr>
<th></th>
<th>Mineral oil</th>
<th>Texas crude oil</th>
</tr>
</thead>
<tbody>
<tr>
<td>shear rate (s(^{-1}))</td>
<td>viscosity, cP</td>
<td>shear rate (s(^{-1}))</td>
</tr>
<tr>
<td>CV-Initial</td>
<td>13.2</td>
<td>586.1</td>
</tr>
<tr>
<td>CV-Final</td>
<td>18.5</td>
<td>390.5</td>
</tr>
</tbody>
</table>
Capillary viscometer measurement for the Texas crude oil emulsion only show a small change (5.3cP) before and after shearing further indicating that the emulsion is Newtonian. Capillary viscometers can be used in place of rotational viscometer provided the emulsion viscosity is not significantly affected by shearing.

### 4.4 Effect of various Surfactant on Emulsion Stability and Viscosity

#### 4.4.1 Effect of different Surfactants on Emulsion Stability

The effect of six different surfactants on emulsion stability was evaluated. Figure 4.8 shows the stability result for both the Texas crude and mineral oil emulsion. All emulsions were prepared using 70% oil, 30% deionized water, 1% surfactant mixed at 20,000 rpm for 2 minutes. The highest stability was obtained with span 85 for the Texas crude oil (less than 10% separation) while the highest stability was obtained with Triton X100 for the mineral oil (less than 5% separation). Blank emulsion for mineral oil separated much faster than the crude oil emulsion because of the absence of any surface active agent. Span 85-R1 is a duplicate emulsion of span 85 used for reproducibility test. The figure shows the span 85 emulsion is highly reproducible. Figure 4.9 shows the picture of the emulsion taken after centrifugation. Blank and span80 Texas crude oil emulsion showed clear oil separation at the top. There was little oil separation with span85. The boundary of separation was not so clear for the Tween 80 emulsion; personal judgement was used in this situation. In addition, there was some water also separating at the bottom.
**Figure 4.8:** Centrifuge stability result for crude oil emulsion (upper) and mineral oil emulsion (lower). All emulsions were prepared using 70\% oil, 30\% deionized water, 1\% surfactant mixed at 20000rpm for 2minutes.

It was a lot easier to read separating volumes for the mineral oil. The lower HLB surfactant emulsion (span80 and span85) had oil separating out while the high HLB surfactant (Twee80 and TX100) had water separating out or no separation happening at all.

Further studies were also carried out to investigate the quantitative increase in stability by adding span85. **Figure 4.10** shows the effect of adding 1\% span85 to texas crude oil.
Figure 4.9: Texas crude oil and mineral oil emulsion pictures taken after centrifugation

(TCO) and crude oil A (CR. A). Emulsion of crude oil A is more stable than texas crude oil because of the higher viscosity of crude A. Higher continuous phase viscosity reduces droplet diffusion. Lower droplet diffusion reduces droplet coalescence rate. The higher stability could also be a result of compositional difference in heavy polar fractions in the crude oils. That information is not available right now. The figures show that after 45minute of centrifugation addition of just 1% span85 increases the stability of the texas crude oil by as much as 94% while addition of the same amount of surfactant to crude oil A increase the stability by as much as 84%.
Figure 4.10: Stability result for texas crude oil and crude oil A emulsion showing increased stability due to addition of just 1% span85 surfactant. All emulsions were prepared using 70% oil, 30% deionized water, 1% surfactant mixed at 20000 rpm for 2 minutes.

4.4.2 Effect of different Surfactant on Emulsion Viscosity

Figure 4.11 and Figure 4.12 shows the capillary viscosity for different surfactant stabilized emulsion. The mineral oil viscosity was higher than the viscosity of the crude oil. Also mineral oil emulsion had higher viscosity than the crude oil emulsion for the same surfactant. The viscosity of all emulsions were higher than the viscosity of the base oil. This holds true for both the texas crude oil and the mineral oil. Higher viscosity than that of the base oil suggests all emulsions were water-in-oil emulsion. Span 80 emulsions had very high viscosity outside of the range of available capillary viscometer. It should be mentioned that the viscosity was taken using different viscometer sizes but it was ensured that the viscosity value falls within the range specified for that viscometer.
Figure 4.11: Effect of different Surfactant on Crude oil Emulsion Viscosity. All emulsions were prepared using 70% oil, 30% deionized water, 1% surfactant mixed at 20000rpm for 2 minutes.

![Viscosity Chart for Crude Oil Emulsions](image)

Figure 4.12: Effect of different Surfactant on Mineral oil Emulsion Viscosity. All emulsions were prepared using 70% oil, 30% deionized water, 1% surfactant mixed at 20000rpm for 2 minutes.

![Viscosity Chart for Mineral Oil Emulsions](image)

4.5 Effect of Water fraction on Stability and Viscosity

A wide range of water fraction was used in creating the emulsions in this study. Figure 4.13 shows the relative viscosity and stability of texas crude oil emulsions as a function...
of water fraction. Relative viscosity has been plotted on the primary axis while stability has been plotted on the secondary axis. All emulsions were prepared using 1% span85 surfactant at 20,000rpm for 2minutes. As is shown in Figure 4.12, the relative viscosity of the emulsion increases sharply with increasing water cut up to the inversion point. At the inversion point the viscosity of the emulsion drops as the emulsion changes from water-in-oil to oil-in-water emulsion. The inversion point for the emulsion at this condition is between 80-90% ±5%. Getting the exact inversion point will be difficult because that will involve very small water fraction increment between 75-90% water fraction. It can be observed from the plot that at 70% water fraction, the relative viscosity of the emulsion increases by as much as a magnitude of 49.

![Graph showing effect of water fraction on emulsion stability and viscosity.](image)

**Figure 4.13:** Effect of water fraction on emulsion stability and viscosity. All emulsions were prepared using 70% texas crude oil, 30% deionized water, 1% span85 mixed at 20000rpm for 2minutes at 25°C.

The higher viscosity observed with increasing water fraction agrees with what other researcher have found (Mohammed (2009), Kokal (2005), Kokal & Álvarez (2003), Almrit et al (2011), Rønningsen (1995) etc). The increase in viscosity is because of
droplet interaction. Distance between droplets decreases as water fraction increase. Reduced inter droplet distance increases the van der waal forces of attraction between the droplets hence a higher resistance to flow. Figure 4.13 also shows that the stability of the emulsion is closely related to viscosity. Since all water fractions were prepared at the same surfactant concentration, increased stability can only be a result of the increasing viscosity of the emulsion. From 10-40% water fraction the stability increases with increasing viscosity because according to stoke’s law, the rate of settling (separation) reduces as the viscosity of the emulsion increases. However, as the inversion point is reached the droplets begin to coagulate and form droplets with a larger radius. This increased droplet radius overrides the effect of increasing viscosity hence the emulsion becomes unstable and invert.

4.5.1 Time Effect on Emulsion Stability

In order to study the flow of emulsion in pipelines it is important to have emulsion that do not change properties such as viscosity and stability over time. This is to ensure that the pipe flow results are consistent and reproducible. The emulsion might also be prepare at a certain time and then studied at another time. It is therefore important to know how the properties change over time. Since the most stable emulsion was produced using span85 it was used to form emulsion at different water fractions. The emulsions were allowed to settle in a bottle test and their viscosity measured every 24hrs for 48hours. Figure 4.14 is a picture of the emulsion just after preparation and 48 hours after. It can be observed that the color of the emulsion changes from dark brown at low water fraction to light brown at high water fraction. With time the emulsion begins to break and the color also begins to change. Each emulsion was prepared in
duplicate. One was used for viscosity measurement the other was used for stability.

Viscosity measurement after 24 and 48 hours was carried by first removing the top oil layer. Samples used for viscosity measurement were taken from the middle (emulsified layer only).

**Figure 4.15** shows the variation in viscosity for each water fraction over 48 hours.

There is little to know variation in the viscosity (and stability) of the emulsion at low water fractions (0-50%). The average standard deviation in viscosity over 48 hours is ±0.69cP for 10%, ±5.38cP for 20%, ±1.13cP for 30%, ±5.62cP for 40%, ±5.79cP for 50%. However, at higher water fraction (60% & 70%) the emulsion breaks down much faster resulting in lower viscosity. The average standard deviation in viscosity is higher; ±32.55cP for 60% and ±163.57cP for 70%.

![Figure 4.14: Emulsion pictures at different water fraction immediately after preparation (upper) and 48 hours after (lower)](image_url)
Figure 4.15: Viscosity as a function of water fraction and time. All emulsions were prepared using texas crude oil, 1%vol span85 mixed at 20,000rpm for 2minutes

4.6 Effect of Surfactant Concentration on Viscosity, TAN and TBN of crude oil

Before investigating how increasing surfactant concentration affects the emulsion viscosity and stability, it is important to see what changes takes in the properties of the crude oil when surfactant is added. This investigation was carried out by mixing different concentration of surfactant with crude oil and measuring their effects on viscosity, TAN (total acid number) and TBN (total base number). Figure 4.16 shows the increase in viscosity of the crude oil by increasing surfactant (span 85) concentration from 0.5-5%vol. The viscosity of the crude did not change significantly, there was only a 19% increase in crude oil viscosity by adding 5%vol of span 85. TAN and TBN measurements were done twice for both the texas crude and crude A. Table 4.2 shows the result for two runs of TAN and TBN for texas crude oil and crude A. The results are generated associated with a standard deviation of less than 2%
between runs. The variation is larger in the TBN than the TAN. TBN values for both crudes are almost the same while texas crude oil has more acids than crude A.

**Table 4.2: TAN and TBN of the Crude Oils**

<table>
<thead>
<tr>
<th>Crude oil</th>
<th>TAN (mgKOH/g) Run1</th>
<th>TAN (mgKOH/g) Run2</th>
<th>TBN (mgKOH/g) Run1</th>
<th>TBN (mgKOH/g) Run2</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCO</td>
<td>0.431</td>
<td>0.470</td>
<td>1.00</td>
<td>1.04</td>
</tr>
<tr>
<td>Crude A</td>
<td>0.118</td>
<td>0.101</td>
<td>1.06</td>
<td>1.03</td>
</tr>
</tbody>
</table>

**Figure 4.17** shows the effect of increasing surfactant concentration on the TAN and TBN. There was no consistent trend observed in the TBN, the value measured were consistently close to that of the texas crude after adding different surfactant concentration. For the TAN however, there was an increase with increasing surfactant concentration. It is important to note that the surfactant does not contain any acid. The TAN setup could not detect any titration endpoint for the surfactant.

**Figure 4.16**: Effect of increasing surfactant concentration on the viscosity of texas crude oil emulsion
Effect of increasing surfactant concentration on the TAN and TBN of Texas crude oil and crude oil A

Figure 4.17: Effect of increasing surfactant concentration on the TAN and TBN of Texas crude oil and crude oil A

4.7 Effect of Surfactant Concentration on Viscosity & Stability of Emulsions

Different concentrations of the surfactants were added to the Texas crude oil. Emulsions were formed and subjected to stability, viscosity, TAN and TBN measurements. At a particular concentration of the surfactant, emulsions of different water fractions were prepared using a continuous process. The lowest water fraction was prepared first and subsequent fractions were prepared by adding extra water to account for the amount of emulsion taken out for viscosity, stability, TAN and TBN measurement. Figure 4.18 and 4.19 shows the result of the water fractions against the relative viscosity and stability respectively for changing surfactant concentration. One of the first that can be observed from Figure 4.17 is that at very low water fraction (0-10%) the effect of surfactant is not very significant except at 5% surfactant concentration. The viscosity profile matches exactly at 0.5% and 1% surfactant concentration. Adding surfactant
Figure 4.18: Relative viscosity at different water fraction for changing surfactant concentration. All emulsions were prepared using Texas crude oil, deionized water and mixed at 20000 rpm for 2 minutes.

lowers the viscosity of the emulsion. However not all concentration of surfactant lowers the viscosity. Adding 0.5% -3% lowers the viscosity of the emulsion when compared to the emulsion viscosity at 0% span85. For instance, at 40% water fraction the viscosity was lowered by as much as 75% just by adding 0.5% span85, 55% by adding 2% span85 and 29% by adding 3% span85. It becomes clear from here that the highest reduction in emulsion viscosity occurs at the lowest surfactant concentration. Reduction of the emulsion viscosity by adding surfactant was also observed by Almrit et al (2011) where they observed that at 20% and 40% water fraction, the emulsion viscosity reduces as they increased alkaline surfactant concentration. This discovery is very important during alkaline surfactant injection as a tertiary oil recovery mechanism. At 5% span85 concentration (and still at 40% water cut) there was a drastic increase in emulsion viscosity. The viscosity increased by as much as 354% over the viscosity at 0% span85 emulsion.
In summary, there is a certain range of surfactant concentration that reduces the viscosity of the emulsion. Observe also that at 2% and 3% vol span 85 concentration the viscosity profile starts to undergo a downward trend at 60% and 50% water fractions respectively. This downward trend indicates that the emulsion at this concentration is starting to invert. Figure 4.18 confirms this finding. Inversion point is usually associated with instability of the emulsion. At 2% and 3% vol, span85 stabilizes the emulsion at all water fraction except at 60% and 50% water cut where the emulsion becomes unstable. This is important because reduction in viscosity and inversion from water-in-oil to oil-in-water emulsion can be take advantage of in pipeline flow.

Figure 4.18 also shows that span85 stabilizes the emulsion at 0.5% - 3% vol concentration. However, at 5% vol concentration the emulsion becomes unstable. This happens to be true for water fraction between 10%-50%.
To confirm if this trend is consistent across different types of crude oil, crude oil A with a different API and viscosity was used. The result is presented in Figures 4.20 and 4.21.

**Figure 4.20**: Relative viscosity against water fraction for crude oil A.

The trend observed with texas crude oil is consistent with what is observed with crude oil A. Adding span85 reduces the viscosity at 1%vol and 2.25%vol. There was also an inversion at 50% water cut at 2.25%span85 concentration.

**Figure 4.21**: Stability of span85 stabilized crude oil A emulsion at different water fraction.
**Figure 4.22** below shows the picture of the texas crude emulsion stabilized with span85 surfactant at 0.5%vol and 5%vol. The pictures were taken after subjecting the emulsion to centrifugation. The emulsions were much stable at 0.5% than at 5%vol span85.

![Images of emulsion samples](image)

**Figure 4.22**: Pictures of texas crude oil emulsion after 45minutes in a centrifuge. On the left is 0.5%vol span85 and on the right is 5%vol span85 concentration

### 4.8 Effect of TAN and TBN on Emulsion Viscosity and Stability

In Figures 4.17 through 4.21 surfactant addition affects both the stability and viscosity of emulsion. The effect is dependent on the concentration of the surfactant. Surfactant must therefore be changing some internal properties of the emulsion. Figure 4.16 shows the TAN of the crude oil increasing with surfactant addition hence it becomes necessary to investigate how the changing surfactant concentration affects the total acid and base number of the emulsion. This section also present graphical correlation between these parameters, viscosity and stability. **Figure 4.23** presents the TAN of emulsion at different water fractions.
The emulsion TAN reduces linearly with increasing water fraction. This appears to be logical given the fact that increasing the water fraction causes a decrease in the amount of surfactant and crude oil. All TAN measurements appear to eventually converge at the same TAN value of 100% water. There was no titration endpoint detected when the TBN of the emulsion was measured. Figure 4.24 summarizes the effect of changing TAN on the relative viscosity of Texas crude oil emulsion. From the figure, the effect of TAN is not so significant at low water fraction (10%). At medium water fraction (20-50%), the viscosity goes through a minimum at low TAN and then shows an upward increase with TAN. The initial drop in viscosity with increasing TAN appears to be more significant at medium to higher water fraction. The zigzag trend seen at 60% water fraction is due to the inversion at 2 and 3%vol surfactant concentration. The decrease and eventual increase in relative viscosity may be explained by the fact that the presence of a small amount of acid in the emulsion lowers the internal friction and interfacial tension of the emulsion thereby lowering the viscosity. At higher TAN however, the effect of the bulk viscosity of the surfactant itself causes the viscosity of
the emulsion to increase. This conclusion is subject to further studies to ascertain if adding the surfactant actually lowers the interfacial tension.

![Figure 4.24: Relative viscosity against TAN at different water fraction for texas crude oil emulsion.](image)

Similar to Figure 4.24 is Figure 4.25 that shows the stability behavior of the emulsion with increasing TAN. One common trend is observed; the emulsion is only stabilized within a certain TAN range. For example, at 10% water fraction, the range is between 0.47-0.79mg KOH/g. The range tends to become smaller with increasing water fraction. There is a range of surfactant concentration within which the associated increase in total acid number increases stability. Acids in crude oil does stabilize emulsion within a particular acid concentration after which any additional increase in concentration destabilize emulsions. A possible explanation for this observation could be the effect the total acid number has on the mobility of the stabilizing film on the water droplet. Within the range that the emulsion is stable the acid interact with other surface active component in the emulsion to form highly stable non-mobile film around the water droplet. At higher TAN the stabilizing films interact and becomes mobile increasing the
tendency for the droplet to coalesce. Other researchers have also reported the effect of mobile and non-mobile stabilizing film on emulsion stability.

Figure 4.25: Effect of TAN on the stability of Texas crude oil emulsion at different water fractions.
CHAPTER 5
CONCLUSIONS AND RECOMMENDATIONS

5.1 Conclusion

In this study, the rheology and stability of surfactant stabilized emulsions were studied. First a surfactant scan was carried out to determine the best surfactant for stable emulsion formulation. The Stability of the emulsion was investigated using bottle and centrifuge test while viscosity was carried out using capillary and rotational viscometer. TAN and TBN of the emulsion were also studied to correlate them with stability and viscosity changes.

The following conclusions about 70%-30% oil-water texas crude oil and mineral oil emulsions using 1% vol surfactant concentration were made:

- An emulsification protocol was developed that produced homogenous and reproducible emulsions. Mixing with IKA T18 homogenizer at 20,00rpm for 2minutes produced emulsions that could be reproduced.
- Among the surfactants investigated Span85 produced the most stable emulsion with texas crude oil while TritonX100 produced the most stable emulsion with mineral oil. They had less than 10% separation after 45minutes in a centrifuge.
- All surfactants investigated produced emulsion with viscosities higher than that of the base oil i.e. all produced water-in-oil emulsion
- The stability of the texas crude oil increased by as much as 94% just by adding 1% span85 while the stability of crude A increased by 84% by adding the same amount of surfactant.
- The texas crude oil emulsion was Newtonian while the mineral oil emulsion was non-Newtonian. Both had time-independent viscosity behaviour.
The following conclusions about 1% span85 texas crude oil emulsion at different water fractions were made:

- Viscosity of the emulsion increases with increasing water fraction up to the inversion point where the viscosity reduces significantly and the emulsion inverts from water-in-oil to oil-in water. Relative viscosity at 70% water fraction increases by as much as a factor of 37 over the base oil.

- The inversion point for the crude at 1% span85 is between 80%-90% ±5%.

- Stability of the emulsion increases with increasing relative viscosity. However, as the inversion point is approached the emulsion becomes less stable despite increasing viscosity.

- All emulsions were stable for 48 hours with little variation in viscosity except at high water fractions (>60%).

The following conclusions about the effect of increasing surfactant concentration on the viscosity of the crude oil, TAN and TBN were made:

- Adding a maximum of 5% vol span85 to the crude oil only increases the viscosity by 19%.

- TAN of the crude oil increases with increasing surfactant concentration while there was no regular trend observed with the TBN.

Finally, the following conclusions about the effect of changing surfactant (span 85) concentration on the TAN/TBN, viscosity and stability of emulsion were made:

- The TAN of the emulsion decreases with increasing water fraction.
• The relative viscosity of the emulsion is reduced by adding between 0.5-3% span85. At 40%, the relative viscosity of emulsion with no surfactant was 19.73 while the relative viscosity with just 0.5% span85 was 4.94 (75% reduction).

• Relative viscosity is independent of surfactant concentration at low water fraction (<10%) except at 5% span85 concentration.

• At 2 and 3% surfactant concentration there was an observed lowering of the inversion point to 60% and 50% respectively. At the observed inversion point the emulsion became unstable.

• At a fixed water fraction, increasing the TAN reduces the viscosity initially thereafter the viscosity begins an upward with increasing TAN i.e. the viscosity goes through a minimum with increasing TAN.

• Emulsion are only stabilized within a particular TAN range. Outside of this range the acids destabilizes the emulsion. The range reduces with increasing water fraction.
5.2 Recommendation

The following further studies are recommended

- Investigate if other stability and viscosity measurement method give the same result. Determining the interfacial boundary of separation can sometimes be subjective. Sometimes the separation boundary is not even distinct.

- Study the effect of changing TAN on the surface and interfacial tension.

- Investigate the flow of stable crude oil emulsion and surfactant stabilized crude oil emulsion in pipeline.

- Develop correlation to predict emulsion viscosity at higher water fraction.

- Repeat the stability and viscosity study using brine instead of deionized water.
References


