RHEOLOGY CHARACTERIZATION OF WATER-IN-OIL EMULSIONS

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ABSTRACT
One of the main problems of the oil industry is hydrate formation. Hydrates of natural gas are crystalline solids, formed by the association of water molecules and molecules of certain gas in an organized structure. Its occurrence is a natural concern for the Oil & Gas Industry, given the potential risk of flow assurance. Nevertheless, ever-challenging scenarios encourage the search for risk assessment methodologies seeking to identify conditions where hydrates will form, but with little possibility of plugging. The present study aims to develop a water-in-oil emulsion capable of giving hydrate at ambient pressure. This work analyzes the effects of water and surfactant concentration in viscosity, storage, and dissipation modulus in a model water-in-oil emulsion. Rheological data are obtained for several combinations of emulsion components, and their mechanical behavior is related to hydrate formation.

KEYWORDS
emulsions; rheology; hydrates; rheometry

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1. INTRODUCTION

Brazil is a major oil producer, with most of its production, about ninety-five percent (95%), derived from the sea (ANP, 2018). Faced with this frontier, major challenges arise daily in oil production from the sea, especially deep and ultra-deep water depths. Shallow, deep, and ultra-deep water depths are defined as those up to 300m, between 300m and 1000m, and above 1000m, respectively (Ferreira Filho, 2016).

This production is drained through pipes called production lines. A common problem that can occur due to particular low temperature and high-pressure, characteristics of areas where these production lines operate, is the emergence of hydrates. On average, the seabed temperature in the Atlantic Ocean is approximately 4°C (Marins et al., 2010).

In addition to the conditions of low temperature and high pressure, other variables may influence hydrate formation, such as composition of the fluid flowing into the system, agitation, contact surface for crystal formation, possibility of agglomeration, and salinity (Sloan & Koh, 2008; Koh et al., 2011).

The hydrate formation can cause clogging of the production lines, generating unscheduled interruptions in operations, for example, during drilling and completion of wells (Santos, 2006). Besides clogging of production lines, hydrate formation can cause equipment damage and compromise the security of the facility. Similarly, in many cases, it is necessary to completely disrupt drilling operations or production to remove these hydrates, which generate major financial losses (Buzatoiu et al. 1999; Baptista, 2007).

Hydrates of natural gas are crystalline solids, formed by the association of water molecules and molecules of certain gas in an organized structure (Sloan & Koh, 2008). Its occurrence is a natural concern for the Oil & Gas Industry. Nevertheless, ever changing scenarios encourage the search for risk assessment methodologies to identify conditions where hydrates will form, but with little possibility of plugging. In this context, the understanding of hydrate rheology is a key element. The first step in comprehending hydrate rheology is evaluating the rheological properties of a model fluid.

In this work, we analyze the effects of water and surfactant concentration on viscosity, storage, and dissipation module in a water-in-oil emulsion. Rheological data obtained for various combinations of the components of the emulsion and their mechanical behavior is related. This study has the differential of deepening the knowledge of this phenomenon through rheological analyses, following a method that is based on rheological and image analyses to evaluate data of water-in-oil emulsions that can generate hydrates at ambient pressure.

2. MATERIALS AND METHODS

2.1 Emulsion preparation

Water-in-oil emulsions with water volume fraction (or water cut, as commonly denoted in the oil industry) ranging from 20 to 40 vol % are prepared using Shell Morlina S2 BL 10 and Shell Morlina S2 B 150, with reported kinematic viscosities of 10 mm² s⁻¹ and 150 mm² s⁻¹ at 40 °C, and density of 0.881 g·cm⁻³ and 0.887 g·cm⁻³ at 15 °C, respectively. The choice of oils and surfactants in the study was due to the ease of obtaining them from suppliers and for presenting physicochemical characteristics of interest in this study.

In all cases, emulsifier agents are composed of a nonionic–anionic surfactant mixture. Two surfactant concentrations are studied and reported here: 1% wt and 5% wt surfactant blend with respect to the total emulsion system. Deionized water is used as the aqueous phase in all emulsions. The composition of the surfactant mixture is 90% wt and Sorbitan monooleate (Span 80, MW 428.61 g/mol, Sigma Aldrich) and 10% wt sodium di-2-ethylhexylsulfosuccinate (AOT, MW 444.56 g/mol, Fisher Scientific). Their molecular structures are shown in Figure 1.

This surfactant mixture is suitable to produce oil continuous emulsions. This is supported by Bancroft’s rule, stating that, in any emulsion, the external (continuous) phase is the one that solubilizes the surfactant (Bancroft, 1913; 1915).

Furthermore, the major component of the surfactant mixture, Span 80, is a surfactant with a hydrophilic lipophilic balance (HLB) value of 4.3. The HLB is a measure of the surfactant
hydrophilicity, which is defined as 1/5 the weight percent of ethylene oxide in the molecule (Griffin, 1949; 1954).

Generally, surfactants with HLB values between 4 and 6 will produce water-in-oil emulsifiers (Griffin, 1949). The emulsions are prepared by dissolving the surfactant mixture in mineral oil by use of a magnetic stirrer and low heating (~50°C). For all water cuts, the mixture is stirred with a high-speed homogenizer at 8000 rpm for 10 min, while water is slowly added during the first minute. After preparation, the emulsions are poured into 200-mL glass flask (Delgado-Linares, 2013).

2.2 Interfacial tension measurements

In this study, we used the Tensiometer - Lauda model platinum ring for measuring the interfacial tension of water–mineral oil interfacial tension (IFT). All measurements were made using the Du Nouy ring of the method (McClements, 1999). First, 20 mL of water was placed in a 50 mL beaker. Then, the ring was dipped approximately 1 cm into the water. Following, 20 ml of oils with surfactants were placed carefully on the water. At last, the measurements were performed.

2.3 Emulsion characterization

With the purpose of observing the relationship between viscosity and water cut, viscosity measurements were performed at ambient pressure on the emulsions by use of a Anton Paar rheometer Physica MCR301 and Physica MCR501. In this study, Cross Hatched geometry was used with a gap size of 1 mm for Shell Morlina S2 B 150 emulsions, and Double Gap – DG geometry for Shell Morlina S2 BL 10 emulsions.

According to Mezger’s work (Mezger, 2015), the choice of geometry is due to some factors related to the fluid to be studied, such as viscosity, whether it has suspended particles, density, amount of available sample, and if its water based or oil based. When the fluid has a very low viscosity, the most suitable geometry is that of concentric cylinders (couette), where the fluid will have a larger contact area with the geometry, implying a higher sensitivity. However, some precautions should be taken, such as sedimentation or particulate fluids that may interfere with the results (Mezger, 2015).

On the other hand, such difficulty is not found in parallel plate and grooved plate geometry, but for such geometries, a correction should be applied, since the shear rate is not constant throughout the geometry (Mezger, 2015).

Viscosity measurements are conducted at room temperature (~20°C). Measurements are performed at shear rates ranging from 0.01 s\(^{-1}\) to 100 s\(^{-1}\). In each test, 6 viscosity points per decade on a log scale were taken at steady state. The marking of each time point from the previous test was obtained using the steady shear rate (Constant Shear Rate) test. This test was set to run for 10 minutes, and the lowest rate of shear was applied from 0.01 s\(^{-1}\) to determine how long the emulsions enter steady state.

2.4 Parallel plates correction

The geometry of parallel plates rheometer is used widely for a variety of materials such as polymer solutions, suspensions, dispersions, and emulsions. An advantage over the cone and plate, or concentric cylinder geometries is the ease of varying the gap, which is decisive in the case of systems with particles or droplets.

However, unlike the cone and plate, or concentric cylinder geometries, the strain rate is not constant along the radius. Thus, both
measured voltage and viscosity must be corrected (Figure 2). The equation is the Weissenberg-
Rabinowitsch dependence of shear rate of non-Newtonian fluids correcting stress (Macosko, 1994). The rheometers (Phyica 301 and 501) used in this study measured viscosity and apparent shear stress, and it was necessary to make a correction.

Equations to correct the shear stress and viscosity:

\[ \tau (R) = \frac{M}{2\pi R^s} \left[ 3 + \frac{d\ln M}{d\ln \dot{\gamma}_R} \right] \]

(1)

\[ \eta (\dot{\gamma}_R) = \frac{\tau_R}{\dot{\gamma}_R} = \frac{\eta_0}{4} \left[ 3 + \frac{d\ln M}{d\ln \dot{\gamma}_R} \right] \]

(2)

Where: \( \tau \) = shear stress; \( M \) = torque; \( \dot{\gamma} \) = shear rate; and \( \eta \) = viscosity.

3. RESULTS AND DISCUSSIONS

3.1 Interfacial tension and droplet size

The calculation of each IFT oil phase with the respective concentration of surfactants was done using equation (3), since all other parameters of this formula are tabulated.

\[
\sigma_l = a_0 + \left( \frac{a_1 + a_2 + a_3}{\rho_L \rho_{ar}} \right)^{0.5} \sigma_l + a_4 \tau_L + a_5 \frac{g_0 \rho_{ar}}{g_1} \frac{a_6}{a_7} + a_6 \sigma_l + a_7 T_L + a_8
\]

(3)

Where:

\( \sigma_l \) = surface / interfacial tension of the liquid corrected in mN/m;

\( \sigma \) = surface tension indicated on the multimeter in V;

\( T_L \) = liquid temperature in °C;

\( \rho_L \) = specific mass of the liquid in g/cm³;

\( \rho_{ar} \) = specific mass of air in g/cm³;

\( g_0 \) = gravity acceleration at the calibration site in m/s²;

\( g_1 \) = gravity acceleration at the place of use of the tensiometer m/s²;

\( a_0, a_1, a_2, a_3, a_4, a_5, a_6, a_7, \) and \( a_8 \) are constants, and their respective values are:

\( a_0 = 0.7250 \)
\( a_1 = 0.00016853 \ g/m^3 \)
\( a_2 = 0.039955 \ g.m^3.V^{-1} \)
\( a_3 = 0.012798 \)
\( a_4 = 0.417473 \ mN/m \)
\( a_5 = 98.972922 \ mN . m^{-1}.V^{-1} \)
\( a_6 = 0.763219 \ mN .m^{-1}.V^{-1} \)
\( a_7 = -1.235590 \ mN .m^{-1}.°C^{-1} \)
\( a_8 = 24.841177 \ mN/m \)

The results of Figure 3 show that in all oily phases studied, the interfacial tension value tends to stabilize just before 1%, that is, with concentrations higher than this it no longer promotes significant changes in the interfacial tension value. This was the reason for the choice of 1% and 5% wt of surfactant concentration for the tests.

The mean droplet size for the emulsions with both 1% and 5% wt of surfactant is in the range 2 – 10 μm diameter, at room temperature (~20°C), for all water cuts studied in this work. These observations are in good agreement with other reported data for water-in-crude oil emulsions (Orciuch et al., 2012; Majid et al., 2015; Kolotova et al., 2017). In all cases, the droplet size is relatively similar for all water cuts. We also observed that the variation in droplet size for
emulsions with 5% wt surfactant is similar to that with 1% wt surfactant. Figures 4 and 5 show examples of microscopic images of the appearance of the emulsions at 20%, 30%, and 40% water cuts 1 day after their preparation for emulsions with 1% and 5% wt surfactant.

3.2 Viscosity results

3.2.1 Shell Molrina S2 BL 10

Viscosity profiles for emulsions prepared with 1% and 5% wt of surfactant measured at ambient pressure in temperature of 20°C are presented in Figure 6. The apparent viscosity of the emulsions increases with the water cut, for Shell Molrina S2 BL 10. This increase in apparent viscosity is due to the increase in packing of the water droplets.

Since the viscosity of the emulsion is proportional to the viscosity of the external phase, the viscosity of the emulsion will deviate further from the viscosity of mineral oil Shell Molrina S2 BL 10 as the water cut increases. Furthermore, shear-

Figure 3. Interfacial tension as a function of surfactant concentration.

Figure 4. Shell Molrina S2 BL 10 sample microscope images of water-in-oil emulsions: 1% wt surfactant with: (a) 20 water vol %, (b) 30 water vol %, and (c) 40 water vol %; 5% wt surfactant with: (d) 20 water vol %, (e) 30 water vol %, and (f) 40 water vol %.
thinning behavior was observed for both surfactant concentrations. This behavior is typical of complex dispersed systems such as human blood, food emulsions, and even water-in-crude oil emulsions; and can be attributed to, in emulsion cases, the rupture of clusters or droplet aggregates, followed by droplet alignment under shear (Izidoro et al., 2009; Cherrya & Eaton, 2013; Ariffin et al., 2016).

Thus, the shear-thinning effect is more significant at higher water cuts and lower temperatures because the system becomes highly packed and the droplet–droplet interactions increase (Ariffin et al., 2016).

Furthermore, viscosity profiles at room temperature (~20 °C) show that, for 20% and 30% vol water cuts, the viscosity of the emulsion is relatively similar for both surfactant concentrations (Figure 7). However, for 40 vol % water cuts, the viscosity of the emulsion is slightly larger for the system with 1 wt % surfactant. The viscosity is related to the size and packing (concentration) of water droplet in the emulsion, and it is known that the increase in droplet–droplet interactions increases the viscosity of the emulsion.

A very different result was obtained in the Stress Sweep tests for all studied water cuts: 20%, 30%, and 40% for the system with 5% wt.

Figure 5. Shell Morlina S2 B 150 sample microscope images of water-in-oil emulsions: 1% wt surfactant with: (a) 20 water vol %, (b) 30 water vol %, (c) 40 water vol %; 5% wt surfactant with: (d) 20 water vol %, (e) 30 water vol %, and (f) 40 water vol %.

Figure 6. Viscosity profiles at various water cuts: (a) emulsion prepared with 5% wt surfactant at 20°C, and (b) emulsion prepared with 1% wt surfactant at 20°C.
surfactant. The elastic or storage modulus $G'$ measured was either 0 (zero) or presented very small values of the order of $10^{-3}$. Therefore, Figure 8(a) shows only the values of the viscous or loss modulus $G''$. This result indicates that the emulsions found are fluids with purely viscous behavior, i.e. without elasticity.

A close result of emulsions prepared with 5% surfactants was obtained in the Stress Sweep tests (Figure 8b). The difference was the 40% water cut, which showed elasticity, since the values obtained for the elastic or storage modulus $G'$ were significant.

Such behavior can be justified by the low viscosity of the oil, coupled with the fact that the emulsions have a very small droplet size and water cuts are not so expressive as to cause the elastic modules to appear in the prepared emulsions.
3.2.2 Shell Morlina S2 B 150

Viscosity profiles for emulsions prepared with 1% and 5% wt surfactant measured at ambient pressure in temperature of 20°C are presented in Figure 9. As can be seen, the apparent viscosity of the emulsions decreases significantly with the water cut, for Shell Morlina S2 B 150 and 1% wt of surfactants, and for 5% wt for surfactants this decrease is milder.

This reduction in viscosity due to phase inversion, as the water cut increased interaction of the surfactant with the oil, was not sufficient to maintain the emulsion as water in oil. This became more evident in the concentration of surfactant at 1% wt. Emulsions for this oil also present a shear-thinning behavior, except for the emulsion with 40% water cut and 1% wt surfactants. The behavior of the emulsion with 40% water cut and 1% wt surfactant is best modeled by a power law fluid. This type of model presents a potential relationship without the presence of a yield stress, thus, it is simpler than other rheological models.

To verify that the prepared emulsions are water-in-oil, a drop test was performed. The drop test is a quick and simple way to determine if an emulsion is oil-in-water (O / W) or water-in-oil (W / O). A few drops of emulsion are poured into a test tube or beaker containing water and into another containing the oil used to make the emulsions. If the emulsion droplets get dispersed in the water sample and not in the sample oil, the emulsion is characterized as O / W. Likewise, an emulsion is deemed W / O when the droplets are dispersed in this oil and remain intact in the test tube or beaker containing the aqueous phase. Figure 9 presents the results of the drop test.

A very interesting result was found after the drop test on emulsions prepared with Morlina S2 B 150 and 5% surfactants. Only the emulsion prepared with a 20% water cut did not dissolve in water (Figure 10a), suggesting that the other emulsions would have reversed their phase, that is, they could be oil-in-water emulsions. Rheological tests for Morlina S2 B 150 with 1% wt surfactant (Figure 11) pointed to a viscosity reduction with increased water cut for emulsions prepared with 1% surfactant, indicating a probable phase inversion. In the drop test this result was confirmed because all emulsions, when dripped in water, dissolved (Figures 10d, 10e, and 10f).
Unlike the previous oil, the Shell Morlina S2 B 150 presented very different results. As the water cut was increased, the variation between the results followed the same direction, that is, they increased more and more. This variation is due to the phase reversal. The higher the water cut, the emulsion became less viscous as the oil droplets were in smaller concentration.

An expected result was obtained in the Stress Sweep tests for the tests from 5% wt surfactants (Figure 12a), in all studied water cuts, 20%, 30%, and 40%. The measured elastic or storage modulus G' was above the viscous, or loss modulus G'' up to a certain stress. This confirms the yield curve behavior, where a yield stress was well defined for each water cut, approaching from 10Pa (Figure 12a).

Emulsions prepared with 1% wt of surfactants showed elastic or storage modulus G' and viscous or loss modulus G'' within the evaluated stress range. However, the 40% water cut test revealed a small region where the elastic or storage modulus...
G' value is greater than the viscous or loss modulus G". The yield stress for the 20% and 30% water cuts was approximately 7PA, showing a viscoelastic behavior for both surfactant concentrations.

4. CONCLUSIONS

The results for the oils were analyzed and contrasted. The Shell Morlina S2 BL 10 presented very low viscosity (~ 0020 Pa.s), whereas Shell Morlina S2 B 150 displayed high viscosity (~ 0.528 Pa.s). From visual tests, one can observe that the emulsions prepared with the Shell Morlina S2 BL 10 were more stable, at surfactant concentrations of 1% and 5%, used by weight, of the emulsions prepared with Shell Morlina S2 B 150. While the first oil gave stable emulsions for more than a week, the second stable emulsions provided stability for only two days, despite the strong lipophilic interaction between surfactant molecules.

The time to perform the rheological tests was up to two days, so the stability of the emulsions was not relevant to interference factor in these results.

This study took into consideration the size of droplets presented, obtained by means of microscopy images, a factor that also influences the stability of emulsions. The emulsions prepared with the Shell Morlina S2 BL 10 showed an average droplet size between 0.5 and 2 μm, whereas the emulsions prepared with Shell Morlina S2 B 150 presented droplets with size between 3 and 10 μm, noting that the only water-in-oil emulsion with Shell Morlina S2 B 150 is prepared with 5% by weight of surfactant and 20% water cut.

Viscosity measurements showed dependency on the water cut in both oils having a shear thinning behavior. In addition, Shell Morlina S2 B 150 showed a wide variation in results between the two concentrations of surfactants used in this study.

Stress Sweep tests showed that emulsions prepared with Shell Morlina S2 BL 10 did not have elasticity, while emulsions prepared with Shell Morlina S2 B 150 showed elasticity, with well-defined linear viscoelastic regions, suggesting a yield stress.

5. REFERENCES


