Occurrence of naphthenate deposition in crude oil production field offshore Niger Delta

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Abstract

Crude oil and produced water samples obtained from ten wells in an offshore field, Niger Delta, were analyzed, in order to determine the occurrence of naphthenates deposition in the field. Total acid number (TAN) and °API of the crude oil samples, pH and metal ions concentrations of the produced water samples were determined. The results revealed that TAN values ranged from 0.47 to 1.01 mgKOH/g with pH of 6.9–8.9, which were above established threshold. The metal ions concentrations especially for Ca++ and Na+ were relatively high. These imply a high possibility of metal-naphthenate precipitation in the oil production facilities in this field.

Keywords Naphthenate · Crude oil · Produced water · Metal ions · Niger delta

Introduction

Naphthenate deposit or scale is presently one of the production chemistry challenges in the upstream crude oil sector with occurrence in different sites, both onshore and offshore, across the globe (Igwebueze et al. 2013). In recent years, there has been a worldwide increase in the production of acidic crudes that are rich in naphthenic acids (NAs) (Oliveira et al. 2013). Geochemical methods have proven useful in determining the presence of these acids in crude oil, which when present implies high potential or possibility of two major problems that are significant in process control as well as flow assurance in crude processing plants; naphthenic acid corrosion (NAC), and formation of naphthenates (naphthenic acid salts) which can either precipitate and form organic deposits (which block process lines and vessels thereby immensely contributing to flow assurance difficulties) or form interfacial active salts (soaps) that are prone to stabilized emulsions (Oliveira et al. 2013). Therefore, determining the presence of naphthenic acids in crude oil is the first step in detecting a portending deposit nucleation sequence and tackling these two aforementioned problems. For example, deposition of calcium naphthenate (CaN) occurs when tetraprotic acids (tetra-acids with molecular weight above 1200 Da) are present in crude oil (Igwebueze et al. 2013).

Meanwhile, it is a known fact that the naphthenic acid phase behavior is a function of produced water composition and pH. Both the tetraprotic acids and other naphthenic acids present in crude oil may react with the cations present in produced water at certain pH values and some specific set of conditions. For example, if there is sufficient calcium and the pH of produced water increases above pH 6 (due to depressurization); there will be a competition of the different naphthenic acids to precipitate as scale (Meredith et al. 2000; Igwebueze et al. 2013). Cation size and hydrated water number have been used to explain the selectivity toward the diveralent ions (Sundman et al. 2010). The reaction between the tetraprotic acids and produced water cations is very fast. These acids are surface active, and thus react to form a network with the calcium ions in the produced water (Nordgard et al. 2010). This network is very sticky, insoluble in oil or water and settles out as organic deposits. A specific set of conditions are necessary for eventual formation of these deposits (Arla et al. 2007). Sodium naphthenate is formed when naphthenic acid reacts with Na⁺ in the produced water. Rousseau et al. (2001) described the naphthenate appearance mechanism as being dependent on the pH increase that occurs during crude oil production, due to carbon dioxide...
(CO$_2$) being evolved from the brine leading to an increase in produced water pH (Rousseau et al. 2001; Goldszal et al. 2002; Dyer et al. 2003; Shepherd et al. 2005). Calcium naphthenate formation has been observed at brine pH greater than about 6 (Goldszal et al. 2002; Rousseau et al. 2001; Turner and Smith 2005) whilst sodium carbonate/eamulsion has been seen at slightly higher brine pH, that is, ~8.5 (Gallup et al. 2005; Turner and Smith 2005). It has also been reported through several experiments that at low produced water pH values precipitation did not occur, as metal naphthenates do not precipitate when the solution pH value is below 6, with a buffered system (Laredo et al. 2004).

Naphthenates or organic deposition is known to be a big challenge in oil fields worldwide. The Niger Delta crude oils are not free of naphthenic acids, at the right pH conditions, the interaction of the naphthenic acids with cations in produced water would lead to formation of naphthenates. However, data on naphthenates deposition in Niger Delta oil fields are scarce. Therefore, in this preliminary study, oil and produced water samples from Field L02, a production field in the offshore Niger Delta, were evaluated in order to determine the level of risk for naphthenates deposition in this field.

**Geology of the study area**

The Niger Delta Basin is situated in the southern part of Nigeria where it is bordered by the Atlantic Ocean and lies between longitude 5°E–8°E and latitude 3°N–6°N (Evamy et al. 1978). The basin has a land area of about 141,639 km$^2$ and sedimentary sequence of 39,600 ft. thickness (Knox and Omatola 1989). It spans in an east–west direction from southwest Cameroun to the Okitipupa Ridge. The basin is framed by older mega tectonic elements such as the Calabar Flank which is the subsurface continuation of the Oban Massif borders at the eastern part. At the western part, the basin is truncated by the Dahomey Basin. The northwestern part is flanked by the Benin Flank which is a subsurface continuation of the West Africa shield. The basin is framed by the Abakaliki anticlinorium (fold belt) in the northeastern part (Evamy et al. 1978). The Delta has an arc shape (Short and Stauble 1967) and it is a high—energy constructive locate delta (Whiteman 1982). The basin is a major geological feature of significant petroleum exploration and production in Nigeria (Whiteman 1982), and it is presently leading oil province in Africa (Reijers 2011). It is the most important sedimentary basin in Nigeria from the view of size, thickness of sediments and economics as its petroleum reserve provides the largest percentage of foreign exchange earnings in the country. Three major cycles of sedimentation have been established in the Niger Delta as well as other parts of the southern Nigerian Sedimentary Basins (Short and Stauble 1967; Murat 1972). They are; the Lower Cretaceous to Santonian Cycle (oldest), Campanian to Paleocene Cycle, and Paleocene/Lower Eocene to Recent cycle (youngest). The third sedimentary cycle, commencing in the Paleocene/Early Eocene, is responsible for the main part of the growth of the delta. The Niger Delta oil province with its commercial oilfields is confined to the area covered by a thick sequence of rocks belonging to the youngest (Tertiary) sedimentary cycle. The area that forms the scope of this study is within the shallow water blocks, located offshore in the southeastern coast of Nigeria. Samples used in this study were recovered from four different comingled samples representing localized blends of all wells in each of the four drilling platforms in the field: L02-D1 (A), L02-D2 (B), L02-D3 (C), and L02-D5 (D). Representative sample of the total blend of all of the aforementioned was also obtained at the comingling flow line, labeled E. For the sake of necessary analysis of produced water in this study, samples were importantly recovered from certain wells with high water cut. All these were within the Field L02.

**Materials and methods**

**Samples**

Crude oil samples and produced water samples used in this study were obtained from Field L02 reservoir streams in the Niger Delta (Fig. 1). This crude oil samples were obtained from each of the drilling platforms in the field. The field has four drilling platforms with a total of 39 oil producing wells. Each platform produces from relatively different reservoirs in the same field. Total blend of all the platform productions was also assayed in this study.

**Determination of total acid number (TAN) of oil samples**

Standard solution of alcoholic KOH was prepared by adding 6 g of KOH to 1 L of anhydrous isopropyl alcohol and gently boiled for 10 min to effect solution. The solution was allowed to stand for 2 days and then filtered through a fine-sintered glass funnel. The solution was stored in a chemical resistant bottle and protected from CO$_2$ with a guard tube containing soda lime. Standardization was achieved with 0.2 g potassium hydrogen phthalate and 80 mL CO$_2$-free water.

Potentiometric titration method (Fig. 2) was employed to obtain the total acid number (TAN) of the oil samples. 125 mL of alcoholic KOH solution was dispensed into the titration vessel and 3 g of crude oil sample was added into the vessel, electrode was immersed in the mixture. KOH delivery tube was also inserted into the titration vessel. Titration was effected until the end point was reached.
Precautions were taken to flush out all bubbles from the KOH delivery tube. Also, after the titration, the electrode and burette tip were rinsed with solvent, then with water for 5 min, then again with solvent. Blank titration was carried out on the alcoholic KOH solution without sample.

**Determination of cations content of produced water samples**

Standard solutions of the calcium, sodium and magnesium ions were prepared from their stock solutions. The stock solutions were diluted with deionized distilled water to prepare each working solution for calibration. The concentrations of the cations in the produced water samples were determined by atomic absorption spectrometry. AA500 atomic absorption spectrometer by PG Instrument was employed. AA500 is equipped with both Flame Atomizer and Graphite Atomizer. The AAS is equipped with automatic 8 hollow cathode lamp turret controlled and optimized by the AAWin3 software. A deuterium-arc lamp was used for background correction. A universal autosampler was mounted on the front of the AA500 instrument. The analysis was performed according to the PG Instrument manual.

Spectrophotometric method was used for measurement of ferrous ion with the use of HACH spectrophotometer DR 2800. Method 255 measurement was carried out at 510 nm. The FerroVer iron reagent is a powder pillow which when dissolved in 25 mL of the sample (contained in a cuvette) converts all soluble iron and most insoluble forms of iron in the sample to soluble ferrous ion. The ferrous ion reacts with phenanthroline indicator in the reagent to form an orange color in proportion to the iron concentration. The result is measured at 510 nm in the HACH 2800 spectrophotometer.

**Results and discussion**

**Geochemical implication of total acid number of crude oil**

The total acid number (TAN) measurement has been used as an indication of the acid composition of crude oil. It has been established that the acidity of crude oils is mainly due to the presence of significant quantities of naphthenic acids.
acids (Passade-Boupat et al. 2012). The total acid number of crude oil samples from Field L02 ranges from 0.47 to 1.01 (Table 1). These relatively high TAN values suggest that the naphthenic acid content of the crude oil samples can intrinsically induce organic deposition. The results showed that there is no linear correlation between the values of TAN and °API (Table 1) because it is expected that TAN value increases with decrease °API.

Mohammed (2010) reported that most of the calcium naphthenate-forming crude oils have higher TAN (0.20–2.50 mg KOH/g), whereas the sodium carboxylate/emulsion forming crude oil samples showed low (0.40–0.41 mg KOH/g) to moderate TAN(0.48–0.50 mg KOH/g), although some calcium naphthenate forming crude oil may have a low TAN. A reasonable guide was established, that is, the higher the TAN value then the lower the API gravity, and subsequently, the tendency of that crude oil sample to form calcium naphthenate deposits during oil production. Although this is not always the case, as there are some crude oil samples with low TAN and API still form calcium naphthenate. Therefore, formation of calcium naphthenate or sodium carboxylate/emulsion is much more associated with the particular acid composition in the crude. Since TAN values for the crude oil samples in this study range from 0.47 to 1.01 mg KOH/g and only samples A and J have TAN values less than 0.50 mg KOH/g, thus it suggests that these oil samples would induce calcium naphthenate formation, which is usually a solid organic deposit with the attendant consequences. This result indicates that Field L02 would have naphthenate deposition and its associated challenges.

Effect of pH and metal ions concentration on naphthenate deposition

The pH of produced water samples from Field L02 range from 6.9 to 8.9, which are above the threshold value of 6.0 (Table 1). The values of TAN in crude oil samples in Field L02 indicate the presence of naphthenic acids in the oil samples (Table 1). Produced water composition is known to play a significant role in the basic naphthenate formation mechanism. Therefore, the pH of the produced water samples can induce the formation of naphthenates in this field, in other words, the pH of Field L02 production system, as measured, presents a good condition for metal naphthenate precipitation.

The profile of metal ions concentrations of the produced water samples from Field L02 (Table 1) reveals that the levels of Ca++, Na+, Mg++ and Fe++ range from 18.28 to 557.97, 78.30 to 11,380, 2.22 to 810 and 0.28 to 13.48 mg/L, respectively. Ca++ and Na+ have average high values; Both Ca++ and Na+ have been implicated in naphthenate formation that resulted organic deposit in production facilities. Calcium naphthenate (CaN) formation has been observed at produced water with pH value greater than 6 (Rousseau et al. 2001; Goldszal et al. 2002; Turner and Smith 2005) while sodium naphthenate (NaN) has been observed at produced water with pH value of

| Sample | TAN (mg KOH/g) | °API | pH | pH* | Ca++ (mg/L) | Mg++ (mg/L) | Na+ (mg/L) | Fe++ (mg/L) |
|--------|----------------|------|----|-----|-------------|-------------|------------|-------------|
| A      | 0.49           | 37.6 | 8.2| NA  | 97.26       | 2.22        | 105.2      | 0.83        |
| B      | 0.54           | 35.6 | 7.9| NA  | 22.18       | 3.02        | 89.91      | 1.15        |
| C      | 0.66           | 40.1 | 8.7| NA  | 26.81       | 2.74        | 98.46      | 13.48       |
| D      | 0.50           | 40.1 | 6.9| NA  | 130.4       | 2.38        | 78.3       | 3.14        |
| E      | 0.58           | 39.8 | 8.2| 7.4 | 112.48      | 2.89        | 99.92      | 1.48        |
| F      | 0.61           | 32.2 | 8.9| NA  | 69.97       | 2.79        | 100.5      | 0.35        |
| G      | 0.50           | 35.7 | 8.6| NA  | 18.28       | 2.79        | 101.8      | 2.66        |
| H      | 0.60           | 38.4 | 8.1| NA  | 101.8       | 1.99        | 82.18      | 2.77        |
| I      | 1.01           | 36.8 | 8.2| NA  | 557.9       | 3.11        | 94.64      | 3.18        |
| J      | 0.47           | 42.3 | 8.4| NA  | 318         | 810         | 11,380     | 0.28        |

*pH* pH measured after acetic acid treatment

TAN total acid number

\[
\text{TAN} = \frac{\text{Base volume} \times \text{base concentration} \times \text{baseM.W}}{\text{Crude oil sample weight}}
\]

where the base concentration is the molar concentration of the base and baseM.W is the molecular weight of the base used.

The mathematical expression for the total acid number (TAN) calculation after the titration was carried out as:

\[
\text{TAN} = \frac{\text{Titrant consumption} - \text{Solvent blank} \times \text{Molarity of titrant} \times \text{Factor on titrant} \times \text{Molar mass of KOH}}{\text{Sample weight}}
\]

\[
(56.106)
\]

Table 1 Geochemical properties of crude oil and produced water samples of Field L02, offshore Niger Delta
about 8.5 (Gallup et al. 2005; Turner and Smith 2005). Therefore, with pH range of 6.9–8.9 and relatively high concentrations of calcium and sodium ions, Field L02 is at a risk of CaN and NaN deposition (Fig. 3). Compared with the concentrations of calcium and sodium ions, the concentrations of magnesium and ferrous ions in produced water samples in this field, though in high pH environment (Fig. 4) is not likely to induce naphthenate formation. Organic deposit (CaN) that results from the reaction Ca\(^{++}\) and naphthenic acid are known to be hard solids while deposit (NaN) from the reaction of Na\(^{+}\) and naphthenic acid is quite soft and also form emulsion at lower concentrations. In essence, the relatively high concentrations of these ions would influence naphthenates precipitation in the production facilities of Field L02. Igwebueze et al. (2013) identified calcium naphthenate (CaN) scale as the major component in the solids in an offshore field in the Niger Delta. Calcium ion concentration in the produced water in the offshore field that witnessed heavy metal-naphthenate deposits was 250 mg/L, and if compared with certain wells (D. E. I and J) in Field L02 with Ca\(^{++}\) concentrations of 130.40, 112.48, 557.97 and 318 mg/L, respectively (Table 1), Field L02 is therefore at high risk of organic deposition. Hence, high concentration of Ca\(^{++}\) and Na\(^{+}\) in Field L02 produced water (brine) can aid formation of metal-naphthenate deposits that could impede production of crude oil in this field.

Impact of acetic acid treatment on pH of produced water was investigated; a sample of produced water was taken at well E in Field L02. The result shows the pH after acetic acid treatment was 7.40 which is lower compared with the pH of the sample before acetic acid treatment (8.2). The lowering of pH indicates that the acetic acid treatment is effective and favorable to inhibition of metal-naphthenate deposition.

Conclusions

The values of total acid number of the crude oil samples obtained in Field L02 indicate high acidic composition which is indicative of naphthenic acid content. The pH values of the produced water samples were all above the threshold value of 6.0. The concentrations of metal ions in the produced water samples from Field L02 are relatively high especially the concentrations of Ca\(^{++}\) and Na\(^{+}\), coupled with the high pH values, can aid the formation of metal-naphthenate formation. The results of these parameters indicate occurrence of metal-naphthenate precipitation at the present operating conditions for Field L02 production. Field L02 seems to be experiencing slight deposition already which has the potential to grow into bulk deposition if not properly
addressed. Acetic acid treatment was effective to mitigate formation of metal naphthenate deposition.

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Conflicts of interest/Competing interests No conflict of interest.

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Fig. 4 Cross plot of pH versus concentrations of Mg++ and Fe++ to evaluate the influence of these ions on naphthenate formation

![Cross plot of pH versus concentrations of Mg++ and Fe++](image-url)
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