Mitigation capacity of an eco-friendly locally sourced surfactant for gas hydrate inhibition in an offshore environment

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Abstract
Gas hydrate inhibition is very key and has become more sensitive as oil and gas exploration goes into deeper terrains especially deep offshore as a result of technological advancement. Use of chemicals has been the most efficient and cost effective in these areas. These chemicals add to the cost of doing oil and gas business and also cause harm to the environment; hence, research has been going on for more eco-friendly and cost-efficient inhibitors. This study takes a look at a locally sourced surfactant as one of such inhibitors. Varying weight percentages of the LSS were screened in a locally fabricated laboratory mini flow loop of 39.4 m with an internal diameter of 0.5 inch mounted on an external frame work. The various pressure plots (pressure vs. time, change in pressure vs. time, initial and final pressures vs. time) show that the LSS used in very small percentages performed better than the synthetic inhibitor methanol (MeOH) used in higher weight percentage than the LSS. The final pressures for MeOH for 1–5 wt% were 104, 111, 123, 120 and 123 psi while those of the LSS were 115, 128, 125, 127 and 131 psi, respectively, for 0.01–0.05 wt%, respectively. This means that the system with LSS had more stable pressure values than those of MeOH. Similarly, the change in pressure at the end of 120 min for MeOH was 46, 39, 27, 30 and 27 psi against 35, 22, 25, 23 and 19 psi for LSS. This was an indication that more gas was used up in the system with MeOH than in the system with LSS. The mitigation capacity of the LSS in percentage was calculated to be 69.30, 80.71, 78.07, 79.82 and 83.3% for 0.01–0.05 wt% while MeOH had values of 59.65, 65.79, 76.32, 73.68 and 76.32% for 1–5 wt%, respectively. This showed that the LSS inhibited hydrates better than MeOH in all the weight percentages considered. There is need to harness and develop the LSS for gas hydrate mitigation because it performed better than MeOH which is a known toxicant to man, terrestrial and aquatic habitat.

Keywords  Gas hydrate · Locally sourced surfactant · Methanol · Mitigation capacity · Laboratory mini flow loop
Introduction

When gas molecules referred to as guest molecules and water called the host molecule balance out via hydrogen bonding, gas hydrate is said to form (Koh 2002). Depending on the stability of guest molecules, temperature and pressure conditions (Fleischer and Janda 2013), different gas hydrate structures, namely SI, SII and SH, can be formed. Of all these structures, the prominent structures encountered in oil and gas operations are SI and SII. When water and gas molecules are present under high pressure and low temperatures, hydrate may likely form. Its formation undergoes processes such as hydrate nucleation, growth and agglomeration (Metaxas et al. 2019). Hydrate agglomeration could block subsea flowlines and pipelines which can cause safety, economic and environmental threat (Zerpa et al. 2011). Although gas hydrates can be challenging for oil and gas operators, which has led to heavy investments to prevent them (Zerpa et al. 2011), they can hold more hydrocarbon volume percentages (say 50%) or fractions in line with water content (Kelland 2006).

Examples of thermodynamic hydrate inhibitors are methanol (MeOH), ethylene glycols (MEG) and triethylene glycol (TEG).

(2) Low dosage hydrate inhibitors (LDHIs) They are so named because they are effective in low dosages of about 0.5 to 3wt of water content although some LDHIs mode of action has not been fully understood and show low degradability (Kelland 2006). LDHIs are classified into kinetic hydrate inhibitors (KHIs) and anti-agglomerants (AAs). Kelland (2006) gave an insight into LDHIs and also early simulations by Carver et al. (1995).

(a) Kinetic hydrate inhibitors (KHIs) These are water-soluble polymers which delay the onset and agglomeration of hydrates. They reduce the rate of gas hydrate formation significantly and allow for production/transportation of fluids (gas and/or oil) during a certain time frame or interval (Kelland, 2006). KHIs tend to lose their effectiveness at higher subcooling (over 120 °C) (Gao 2009).

(b) Anti-agglomerants (AAs) are surface active agents (Chua and Kelland 2013) that prevent agglomeration and if properly administered, produces a slurry of hydrates which is transportable. Anti-agglomerants (AAs) are mostly utilized for hydrate management where thermodynamic inhibition is no longer feasible (Anklam et al. 2008; Gao 2009). The use of AAs allows hydrates to form in the pipeline, but the chemical additives reduce the cohesion or clumping of the particles to one another and the adhesion of particles to the walls, such that hydrates flow as a slurry until the pipeline reaches higher temperature, lower pressure conditions, and the hydrates dissociate back into their constituent components. AAs have complex chemical compositions, but many contain surfactant-type chemical groups which are molecules with both hydrophilic and oleophilic functionalities, which allow for interaction with both the water and the oil in a pipeline (Chua and Kelland 2013; Zanota et al. 2005). An increase in salt concentrations increases their performance (Aman et al. 2015). AAs are known to be effective at high subcoolings and may possibly not slow the rate of growth but can prevent agglomeration to form large plugs. This action can prevent blockages in pipelines. Examples such as quaternary ammonium salts (QAs) are well-known AAs (Zanota et al. 2005). Most AAs are effective only at lower water cuts (< 30%).
LDHIs do not prevent the formation of hydrates completely because they do not cause shifts in hydrate equilibrium curves such that once hydrates form they cannot be removed by LDHIs (because the conditions of operation cannot be changed); therefore, a well using LDHIs is still at risk of forming hydrates and will still require THIs. The advantages of LDHIs is that they reduce OPEX and CAPEX. The disadvantage is that their application field on large scale cause important problems due to the fact that they are poor biodegradable inhibitors (Jensen et al. 2010).

Typically, LDHIs are affected by oil composition, density, viscosity, emulsion forming tendency, amount of gas present in oil (gas–oil ratio, GOR) (Aman et al. 2017). The performance of LDHIs can be drastically affected by little changes in their molecular structure. It is common practice for systematic experiments to be carried which is aimed at identifying the optimum dosage and composition of LDHIs.

Another important way of managing gas hydrate is by preventing the fluid produced from entering the hydrate stability region during flow (through front end engineering design). This can be achieved via multiphase system modeling (Charlton et al. 2018). This is done by finding out the pipeline size, subsea system pumping requirement and the insulation thickness. This is imperative due to the changing nature of produced fluid as the well gets older, making it possible for the fluid to enter hydrate stability zone, causing blockages during flow as the field matures (Mehta et al. 2002). The internal surfaces of pipelines/flowlines make for many nucleation sites and coldest point. Field trials and several experiments had suggested that deposition of hydrates often occurred on the wall/surfaces of the pipelines (Nicholas et al. 2009; Aman et al. 2014). Changes in rheology such as sloughing and jamming were indicators of hydrate plugging. To effectively prevent plug formation in offshore operations, knowledge of hydrate formation and flow behavior are key. Great efforts have been put in by researchers to get the knowledge of these key factors (Aman et al. 2010, 2011).

Farhadian et al. (2020) carried out a study on a novel strategy which had potential. This was based on waterborne polymers (which were synthesized in the absence of a catalyst and according to Farhadian et al. 2019) for hydrate and corrosion risk management and incompatibility issues. They found out that the waterborne polymers (waterborne polyurea/ureathenes, WPUUs) performed well in delaying hydrate induction by 17 times and reduced growth rate by 7.7 times when compared to water. They reported that WPUUs are promising kinetic/anti-agglomerant methane hydrate and corrosion inhibitors, which will chart a new course in hydrate and corrosion risk management. Kumar et al., (2019) in their study used a simple apparatus and short experimental times to assess morphology, phase equilibria and apparent kinetics of methane hydrate in the presence of thermodynamic hydrate inhibitor ethane-1,2-diol (MEG) and sodium chloride (NaCl) and kinetic hydrate inhibitor polyvinyl-pyrrolidone (PVP). They found out that both NaCl and MEG acted as kinetic inhibitors at the studied concentrations and concluded that in all, the obtained morphology, kinetics and phase equilibria results were reproducible and consistent and that assessment of multiple subcooling in a single experiment was possible from gradient experiments. They opined that the 3-in-1 reactor technique should be used especially in assessment of potential inhibitors at critical turnaround times or when samples are limited in quantities.

In their study, Zhang et al. (2017) used a newly built up loop, pilot scale experiments to study typical hydrate plug phenomena exploring the reasons behind them. They used a tetrahydrofuran (THF) hydrate slurry @ 50 and 100 vol% with a typical anti-agglomerant KL-1. Based on morphologies, temperature variation as well as liquid loading and an anti-agglomerant on morphology and temperature, they found out that hydrate plugs were induced by hydrate deposition around the pipeline walls but not from the bed at the bottom of the pipeline. Also that AA compressed hydrate deposition and plug by dispersing hydrate particle agglomeration. They concluded that adhesive forces among hydrate particles can be reduced by AA and that hydrate agglomeration was loosened at lower subcooling but failed at higher subcooling. A study on the effect of soy lecithin as an anti-agglomerant using cetyltrimethylammoniumbromide (CTAB) on tetrahydrofuran (THF) hydrate was studied by Saikia and Mahto (2018) using an experimental setup built in house. It was found out that soy lecithin @ 0.1 wt% and 1°C in an experimental solution increased agglomeration time by 1440 min while @ 0.01 wt% it increased it by 1000 min. They concluded that soy lecithin a bio surfactant, can be effectively used as an anti-agglomerant additive to prevent hydrate plugging in oil/gas pipelines and drilling fluid flowlines.

An anti-agglomerant showing high effectiveness at low dosage in methane/natural gas hydrate over an entire water cut range was reported by Zhao et al. (2016). They used an improved formulation in various systems at constant high pressure (100 bar of natural gas) and high cooling rate (~8 °C/h) over water cut ranges of 30–80% in a closed rocking cell system with condensate liquid and crude oil as hydrocarbon liquid phase. They demonstrated the effectiveness of improved AA formulation in an extensive set of measurements. They found that increasing salinity can decrease the dosage of base chemical in formulation significantly.

Muhammad et al. (2016) presented a work on the various water-soluble polymers used for gas hydrate inhibition, including conventional and novel polymeric inhibitors along with their limitations. The work covered relevant properties of vinyl lactam, amide, dendrimeric fluorinated and natural biodegradable polymers. A complete review of the modus operandi used to assess the performance of the polymeric...
inhibitor was given. It addressed recent developments, current and future challenges and field applications of a range of polymeric kinetic hydrate inhibitor.

Jiyeon et al. (2016) investigated the effect of 2-diamine piperazine (PZ) and hydrazine (HZ) on the formation of methane hydrate (CH$_4$) with primary focus on thermodynamic, spectroscopic and computational analysis for their potential application in natural gas hydrate production and transportation. The methane hydrate equilibrium lines were shifted to low temperatures and high-pressure zones based on the concentrations of 2-diamines used. A slower hydrate growth and longer induction time as well as significant thermodynamic hydrate inhibition effect were shown by HZ than PZ.

The use of chemicals as a means of gas hydrate inhibition has been ongoing, and these are detrimental to the environment and humans. Local materials have been reported in enhanced oil recovery (Emuchay et al. 2013; Ojo et al. 2013) and also in the pharmaceutical industries use has been made of local extracts (Ezejiofor et al. (2013). Their use has been found to be effective in the area of flow assurance as local extracts have been used as hydrate inhibitors in lieu of the synthetic ones (Elechi et al. 2018, 2019, 2020 and Okon et al. 2018). This paper is a continuation of the use of such materials as gas hydrate inhibitors.

Materials and method

The materials used for this study are water, ice blocks (facilitated the attainment of hydrate formation temperature), compressed natural gas (CNG with mass of 14.5 g, 0.5 specific gravity), conventional hydrate inhibitor, methanol and locally sourced surfactant (LSS) for gas hydrate inhibition.

The LSS is of the plant family caricaceae. It was obtained from the botanical garden of University of Port Harcourt. The extracts were obtained by manual crushing of tender leaves, sieving with the aid of a cheese cloth and then collected in air tight containers and used immediately. Phytochemical screening showed the presence of some phytochemicals present in the LSS as shown in Table 1. They have characteristics that make them good inhibitors based on the polyphenolics and hydroxyl group they contain. It contains the –OH group just like methanol (MeOH). It contains oxygen bonded to hydrogen. It is pair of atom that is commonly found in organic compounds such as sugar (carboxylic acids) and alcohols. The anion [OH–] called hydroxide consists of a hydroxyl group. IUPAC (2015) rules say that the term hydroxyl refer to the radical OH only while the functional group-OH is called hydroxyl group. Its chemical formula is written as –OH where—refers to the carbon to which the hydroxyl group is bonded.

The extract is water soluble containing naturally occurring bioactive compounds like alkaloids, saponins, flavonoids and tannins. Flavonoids are polyphenolics that are anti-coagulants that prevent coagulation and act as powerful oxidants which protect against reactive oxygen. Tannins are a heterogenous group of high molecular weight polyphenolic compounds with capacity to form reversible and irreversible complexes. They are also found in fruits, legumes and grasses. Polyphenolics are a class of chemical compounds consisting of a hydroxyl group bonded directly to an aromatic hydrocarbon group. They are otherwise called poly hydroxyl phenols and are phytochemicals. They are a structural class of mainly natural products (chemical compounds or substance produced by a living organism that is found in nature (Quideau 2011). Condensed tannins are the most abundant poly phenols found in virtually all families of plants. Leaf tissues, the epidermis, bark layers, flowers and fruits have larger concentrations of poly phenols (Quideau et al. 2011).

Alkaloids are natural products that contain heterocyclic nitrogen atoms and are basic in character. They are naturally synthesized by a large number of organisms including animals, plants, bacteria and fungi. Saponins form a stable foam in aqueous solutions such as soap hence the name ‘saponin.’ Chemically, saponins as a group include compounds that are glycosylated steroids, hypernoids and steroid alkaloids. Saponins also prevent agglomeration and are surface acting agents with bubbles and foams acting as barriers that give good stability. Fourier transform infrared spectroscopy (FTIR) showed the presence of tannins, and alkaloids (with functional groups containing nitrogen, oxygen and carbon) with aromatic rings. The presence of flavonoids, tannins, saponins and alkaloids in plant extract contributes to its ability to inhibit hydrates just like conventional alcohols methanol and glycols.

Table 1 Qualitative analysis of bioactive components of the Locally Source Surfactant

| Bioactive components | Quantity present in extract |
|---------------------|-----------------------------|
| Tannins             | + + +                       |
| Saponins            | + +                         |
| Flavonoids          | + +                         |
| Resins              | +                           |
| Alkaloids           | + +                         |

Key: + means present in small quantity, + + means present in appreciable quantity, + + + means present in abundance.

Experimentals

The experimental setup used for this work is a 39.4-m-long, 0.5-inch I.D. locally fabricated laboratory mini flow loop housed in a PVC pipe (4 inches in diameter) used for the
screening of the locally sourced and conventional inhibitors (Fig. 1).

The system is first flushed to remove debris before the actual experiments. For hydrate formation experiment, water is poured into the mixing vessel, at that point pump 3 is turned on and some quantity of water (about 435 ml) of water is let into the inner line via valve 4 (with pressure of about 25 psi), after which both pump and valve are closed. In order to build the pressure of the inner line to 150 psi, valve 1 and orifice are opened and then both are closed. Water enters into the cooling unit (in this case refrigerator) and is moved around the PVC by opening pump. Agitation and mixing of gas and water as well as circulation in the inner line is initiated by turning on pump 1. Pressures and temperatures readings are taken every 2 min for a total of 2 h. Pointers to formation of gas hydrates in this setup are: increase in loop temperature, a rapid decrease in system pressure, increase in pressure between inlet and outlet pressures (differential pressure), clogging of sample point valve and milky color of effluent). For hydrate mitigation experiments, water and inhibitor (depending on the concentration or weight percentage to be used) are poured into the mixing vessel and the experiment proceeds as earlier described.

**Results and discussions**

The performance of the local surfactant (LSS) was validated using pressure versus time and temperature versus time plots and superimposed with the pressure, temperature versus time plots for the uninhibited experiments. The performance is ascertained by the pressure value at the end of the experiment. A drastic temperature increase or pressure decline are pointers to hydrate formation for the uninhibited experiment while a regulated temperature or pressure profile is an indicator of the LSS’s ability to mitigate gas hydrate formation.

Pressure versus time for the various weight percentages of LSS and pressure versus time for uninhibited system is shown in Fig. 2. For 0.01 wt.% LSS, the pressure dropped from an initial pressure of 150 psi to 121 psi in 30 min, then to 118 psi in 60 min, 116 psi in 90 min and finally dropped to 115 psi at the end of 120 min. For 0.02 wt%, the temperature decreased to 135 psi in 30 min, 130 psi in 60 min, 129 psi in 90 min and finally to 128 psi at the end of the experiment. For 0.03 wt%, pressure dropped from 150 psi at the inception of the experiment to 133, 128, 126 and 125 psi in 30, 60, 90 and 120 min. For 0.04 wt%, the pressure decreased from 150 psi to 135, 131, 128 and 127 psi in the same time interval. Pressure decreases for 0.05 wt% in 30 min were 140 psi, then reduced to 131 in 60 min and was maintained till the end of 120 min.

For the uninhibited experiments, the pressure decreased from 150 psi to 113, 80, 40 and 36 psi for the same time frame. The various weight percentages of the LSS were able to regulate the pressure in the system, and the initial decreases were caused by the cold water flowing in the PVC pipe. Also, the gradual pressure decreases were occasioned by the formation of an impermeable layer between the fluids (gas and water) flowing in the inner line (Movareji et al. 2016). This was not the case for the uninhibited experiment where the pressure declined drastically from 150 to 36 psi at the end of 120 min. This drastic decrease was as a result of gas being used up to form hydrates due to large gas
dissolution in water which is the initial gas hydrate formation phase. This was the resultant effect of release of latent heat due to exothermic nature of gas hydrate formation (Movareji et al. 2016).

Figure 3 is the temperature versus time for 0.01–0.05 wt% locally sourced surfactant and uninhibited experiments. The temperature decreases for 0.01 wt% was from 30 °C to 20.5 °C, 10 °C, in 30 and 60 min and then to 6.5 °C in 90 min and were maintained till the end of 120 min. 0.02 wt% had temperature decreasing from 30 to 14 °C, 7.5 °C, 7 °C and 6.5 °C in 30, 60, 90 and 120 min while for 0.03 wt%, the values were from 30 °C to 13, 9, 8 and 7.5 °C, respectively, for same time interval. For 0.04 wt% had temperature decreasing from 30 °C to 16.5, 11.5 and 7 °C in 30, 60, 90 and 120 min. The values reduced from 32 °C to 18, 9.5, 7.5 and 6.5 °C in 30, 60, 90 and 120 min. In all the weight percentages, the temperature decrease was regulated and there was no drastic temperature increase which was noticed in the uninhibited experiment. After a steady decrease from 30 to 25 °C in 30 min, the temperature increased to 28, 29 and 29.5 °C in 60, 90 and 120 min. This increase led to large dissolution of methane gas in water which hastened.
agglomeration/precipitation of hydrate as critical concentrations were reached (Zhang et al. 2017).

In Fig. 3, the temperature increase was significant because from the plots, it is seen that for the inhibited experiment using 0.01–0.05 wt% of the surfactant, the temperature declined steadily and was well controlled. The sustained decrease in pressure was as a result of the inhibitors being able to regulate the temperature of the system which was observed in all the weight percentages but for the uninhibited, the temperature declined initially but after about 30 min, there was a spike which increased gradually and was sustained till the experiment ended. This sudden increase heralds the inception of gas hydrate agglomeration in the inner line owing to the exothermal nature (heat given off) of gas hydrate formation which may have been responsible for the temperature increase. As earlier stated in the discussion in Method section, one of the pointers to gas hydrate formation is a sudden spike in system’s temperature.

Gas hydrate mitigation comparison of LSS and MeOH

To assess the performance of the locally sourced surfactant (LSS) and the conventional inhibitor methanol (MeOH), various plots of pressure versus time, initial and final pressure versus time and change in pressure versus time were plotted. Furthermore, the mitigation capacity (MC) of the various weight percentages of both the local and conventional inhibitors was also calculated. It is worthy to note here that the LSS was used in low dosages of 0.01–0.05 wt% that is as low dosage hydrate inhibitor (LDHIs) while the conventional inhibitor was used as thermodynamic hydrate inhibitors (THIs) in high dosages of 1–5 wt%.

The conventional inhibitor, MeOH, was used in larger dosages of (1–5 wt%) while the LSS was used in smaller weight percentages (0.01–0.05 wt%) to see how they compare or contrast. The use of low dosages for the LSS was to ascertain the degree of their effectiveness or otherwise as a local surfactant for the study. Hence various weight percentages were compared.

The surfactant functions by adhering or adsorbing to the hydrate crystal structure via hydrogen bonding which forces the crystal to grow around and between the surfactant strands, thus preventing water from entering and stabilizing the hydrate cages. This adsorption occurs because of the interaction force between the surfactant molecules and the water molecule such that inhibition effect is attributed to the adsorption of the surfactants molecules through their functional groups on hydrate molecules. Water has very strong cohesive force which makes surface tension high, and as the surfactants adsorb (water molecules and surfactant molecules have low intermolecular forces hence decreases surface tension), they break these interactions.

The surfactant monomers disrupt the hydrogen bonding between water molecules at the surface, thus lowering the surface tension.

![Pressure versus time plot for 0.01, 0.02 and 0.05 wt% LSS and 1, 2 and 5 wt% MeOH and uninhibited experiment](image-url)
Pressure versus time plot

Figure 4 is the pressure versus time plot for 0.01, 0.02 and 0.05 wt% LSS and 1, 2 and 5 wt% MeOH. These plots are used as representatives of 0.03 and 0.4 wt% LSS and 3 and 4 wt% MeOH. The pressure dropped from 150 psi to 121 in 30 min, 118 in 60 min and then increased to 116 psi in 90 min and finally to 115 psi at the end of 120 min for the LSS. For the MeOH inhibited system, the pressure decrease for the same time interval was from 150 psi to 114, 107, 106 and 104 psi, respectively. The pressure in the LSS inhibited system was more regulated than that of the MeOH inhibited system. The pressure versus time plot for 0.02 wt% LSS and 2 wt% MeOH. Pressure decreases were 135, 130, 129 and 128 psi in 30, 60, 90 and 120 min for LSS while values for 2 wt% MeOH for same time interval were 120, 117, 114 and 111 psi. The pressure versus time plots for 0.03, 0.04 and 0.05 wt% of LSS followed the same steady pressure decrease trend when compared to the 3, 4 and 5 wt% MeOH. In the weight percentages considered, the LSS inhibited system had better pressure control than the MeOH system. This shows that the LSS performed better than the conventional MeOH even when used in weight percentages that were 0.99, 0.98, 0.97, 0.96 and 0.95 less than those of MeOH. The uninhibited scenario had rapid pressure decrease from 150 psi to 113, 80, 40 and 36 psi in the same time interval. This was as a result of more of the gas dissolving in water such that the dissolving fluid could no longer take any more gas which led to the shortening of the time it takes for hydrate to form (Bishnoi and Natarajan 1996).

Initial (P_i) and final (P_f) pressure plots

Another yard stick used in assessing the performance of the conventional and local inhibitor was the initial (P_i) and final (P_f) pressure. Basically, it is the final pressure because the initial pressure was the same for both the conventional and local inhibitors. The final pressure tells how much gas was used up or dissolved in the carrier fluid in order to form hydrates. The more the final pressure (P_f), the better the performance of the inhibitor.

The P_i and P_f versus time for 0.01–0.05 wt% LSS and 1–5 wt% MeOH and uninhibited experiment is shown in Fig. 5. At the end of the experiment (120 min), the pressure in the system using MeOH was 104 psi while that of the LSS system was 115 psi. For 2 wt% MeOH and 0.02 wt% LSS, the system with LSS had a P_f of 128 psi while that of MeOH was 111 psi. For 0.03 wt% LSS and 3 wt% MeOH, final pressure for the system using LSS was 125 psi while the system with MeOH had a value of 123 psi. For 0.04 wt% LSS and 4 wt% MeOH, the P_f for system using MeOH was 120 psi while that of the LSS was 127 psi. For 0.05 wt% LSS and 5 wt% MeOH, the values were 131 and 123 psi, respectively, at the end of the experiments. The LSS performed better than the conventional MeOH although both inhibitors inhibited gas hydrate formation better than in the uninhibited inhibited experiment where the final pressure at the end of the experiment was 36 psi meaning that almost all the gas in the system was used for gas hydrate formation. The cooling process contributed to the initial rapid pressure decrease due to large dissolution of gas in water (Movareji et al. 2016).
The change in pressure was also used to assess how the local and conventional inhibitors behaved in terms of suppression performance. The value of change in pressure tells how well the inhibitor was able to prevent gas hydrate formation. Figure 6 show the change in pressure versus time for 0.01, 0.02 and 0.05 wt% LSS and 1, 2 and 5 wt% MeOH alongside the uninhibited experiment. The change in pressure for 0.01 wt% LSS at the end of 120 min was 35 psi while that of 1 wt% MeOH was 46 psi. Change in pressure for LSS was 11 psi less than that of MeOH. For 0.02 wt% LSS, the change in pressure was 22 psi while that of 2 wt% MeOH was 39 psi which is 17 psi more than that of LSS, while 0.05 wt% LSS and 5 wt% MeOH had ΔP values of 19 psi and 27 psi, respectively, at the end of 120 min. The other weight percentages (0.03/3, 0.04/4 wt%) of LSS and MeOH showed a similar trend as the those discussed above. Generally, both inhibitors had far lesser change in pressure values than the uninhibited experiment which had a change in pressure value of 114 psi which was an indication that almost all the gas was used up in gas hydrate formation (Jensen et al., 2005). It is worthy of note that even though the concentration of MeOH was higher than that of LSS in terms of weight percentage, it performed less than the LSS. This is very remarkable and speaks volume for the unconventional compound as gas hydrate inhibitor.

**Mitigation capacity**

The performance of the inhibitors was further ascertained by calculating the rate at which they mitigated gas hydrate formation.

Mitigation capacity, MC, is given as

\[
\%MC = (1 - X)\%
\]

(1)

Mitigation factor is given as

\[X = \frac{\Delta P_{inhibited}}{\Delta P_{uninhibited}} = \frac{Pi - \Delta P_{uninhibited}}{Pi - P_{inhibited}}\]

(2)

\[\Delta P_{inhibited} = Pi - P_{inhibited}\]

(3)

\[\Delta P_{uninhibited} = Pi - P_{uninhibited}\]

(4)

where.

\[\Delta P_{inhibited}\] is the difference between initial and final pressure of the inhibited experiment.

\[\Delta P_{uninhibited}\] is the difference between the initial and final pressures of the uninhibited experiment.

\[Pi\] is the initial pressure for both the inhibited and uninhibited systems.

\[P_{inhibited}\] is the final pressure for the inhibited systems.

\[P_{uninhibited}\] is the final pressure for the uninhibited system.

The mitigation capacity (MC) versus weight % for the various weight percentages of LSS and MeOH is shown in Fig. 7. For 1/0.01 wt% MeOH and LSS, the MC values were 59.65 and 69.3%. At this weight percentage, over all the MC of LSS was 9.65% more than that of MeOH. The MC for 2 wt% MeOH was 65.79% while that of 0.02 wt% LSS was 80.70% and was able to mitigate gas hydrate formation 14.91% more than the mitigation ability of MeOH. The value of 0.03 wt% LSS was 78.07% while that of 3 wt% MeOH was 76.32%, a value that was less than that of LSS by 1.75%. For 0.04/4 wt%, the values for LSS and MeOH were, respectively, 79.82 and 73.68%. LSS mitigated gas hydrates better than MeOH by 9.65%, while for 5 wt% MeOH, the MC value was 76.32% while 0.05 wt% LSS had a MC value of 83.3%, which is greater than the value of MeOH by 6.98%.
From the analysis of the MC of the inhibitor, it is seen that the mitigation capacity for MeOH decreased after 3 wt% (which is considered the optimum weight % for MeOH). As concentration increased for MeOH, the mitigation capacity decreased at 4 wt% and then increased to the same weight percentage as 3 wt% at the end of the experiment as shown in Fig. 7. This is a case of over inhibition leading to decrease in effectiveness. For LSS, it was observed that after 0.02 wt%, the MC value decreased by 2.63% and 0.88%, respectively, after which it increased by 2.63% at 0.05 wt%. The decrease in MC could also be as a result of over inhibition, but it picked up at 0.05 wt%. 0.02 wt% is considered as optimum for LSS because above this concentration the MC dropped. Even though the MC at 0.05 wt% increased, the value was 2.63% more than the MC value at 0.02 wt% (this can be neglected). It is seen that LSS performed much better than the conventional MeOH in all the weight percentages considered.

The optimum weight fraction for LSS is considered to be 0.02 wt% because at this weight percentage, it had a better pressure drop control, had least change in pressure values and highest final pressure value when compared to the other weight percentages except for 0.05 wt%. The values gotten at 0.05 wt% are negligible such that inhibiting above that weight percentage (0.02 wt%) is not economical. The MC for 0.02 wt% LSS is higher than that of all the wt.% (1–5 wt%) of MeOH.

### Conclusions

The pressures in the system with LSS at the end of 120 min for the various weight percentages were more regulated with values of 115, 128, 125, 127 and 131 psi while those of MeOH inhibited system were 104, 111, 123, 120 and 123 psi, respectively, at the end of 120 min. For the uninhibited system, the pressure values were 113, 80, 40 and 36 psi in 30, 60, 90 and 120 min.

The change in pressure in the systems showed the LSS inhibited system to have values of 35, 22, 23 and 19 psi for 0.01–0.05 wt%, respectively, at the end of 120 min while values for MeOH were 46, 39, 27, 30 and 27 psi, respectively, for the same time frame. The MC for the various weight percentages of the LSS was 69.3, 80.7, 78.07, 79.82 and 83.3%, respectively, while values for MeOH were 59.65, 65.79, 76.32, 73.68 and 76.32% for each weight percentage considered.

From all these, it is clear that the LSS performed far better than the conventional MeOH in all weight percentages considered even though they were used in much higher dosages than the LSS. The weight percentage that is to be considered optimal for inhibition for both inhibitors is 0.02/3 wt% for LSS and MeOH, respectively. At this weight, the mitigation capacity was highest after which it began to decrease and then increased at 5 wt% (for MeOH). For LSS after this weight percentage, the MC decreased and then increased to a value of 83.3% which was gotten at the end for 0.05 wt%. The difference between the MC at 0.02 wt% and 0.05 wt% is 2.63%. This is not so significant and inhibiting above 0.02 wt% for LSS would not be economical. The LSS should be considered and harnessed for gas hydrate inhibition given its performance moreover it is more eco-friendly than methanol which is a known toxicant.

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**Fig. 7** Mitigation capacity, MC, versus weight% for 0.01–0.05 wt% LSS and 1–5 wt% MeOH
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Declarations

Conflict of interest The authors whose names are listed above certify that they have no affiliation or involvement in any organization or entity with any financial interest or non-financial interest in the subject matter or material discussed in this manuscript.

Ethical approval All authors approved final manuscript.

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