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Hydrate Size Measurements in Anti-agglomeration at High Watercut by New Chemical Formulation

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ABSTRACT: Risk management of gas hydrates is a major challenge in offshore hydrocarbon production. Use of antiagglomerants (AAs) is an attractive option because of effectiveness at low dosage and high subcooling. The literature suggests that anti-agglomeration requires formation of water-in-oil emulsion. Our recent work has demonstrated that the process can occur without emulsion or through oil-in-water emulsion. We have shown anti-agglomeration in oil-free systems in methane hydrates and natural gas hydrates. In this work, through particle size measurements in an autoclave setup in both freshwater and brine, light is shed on the effectiveness of our chemical formulation. Hydrate particle sizes are determined by the focus beam reflectance measurement (FBRM). The results show that our formulation reduces hydrate particle size significantly and eliminates large particles. The AA formulation is a mix of three chemicals: surfactant, base, and oil (e.g., *n*-octane). *n*-Octane helps with even distribution of the surfactant in the solution. The base helps with elimination of small amounts of hydrogen ions in the aqueous solution in conditions when acid gas species are present in the natural gas. These two additives increase the effectiveness of the surfactant for anti-agglomeration. Hydrate particle size measurements show that small amounts of *n*-octane reduce particle sizes by a factor of 2. Our measurements reinforce the idea of effective anti-agglomeration to be the most feasible approach in hydrate flow assurance, with superiority over other alternatives. The formulation has many features, including viscosity reduction of slurry.

INTRODUCTION

Small gas molecules (e.g., methane, ethane, propane, carbon dioxide, and nitrogen) in oil and natural gas flowlines often



Figure 1. Schematic diagram of the (a) FBRM probe and (b) chord length during in-process measurement (modified with permission from the Mettler Toledo website).

form gas hydrates with water at low temperature and high pressure.¹⁻⁴ Such crystalline complexes raise flow assurance challenges because they may block flowlines and cause serious safety and environmental issues. Hydrate formation resulted in the failure of oil capture from the seabed in the accident in the Gulf of Mexico.⁵ The traditional method by thermodynamic inhibitors becomes less attractive in offshore oil/gas operations and productions because of the demand for large quantities of alcohols, e.g., up to 60 wt % in the aqueous phase to shift the equilibrium of the hydrate regime.³ The alternative approaches rely on changing hydrate surface properties with small chemical quantities, which are referred to as the low-dosage hydrate inhibitors (LDHIs, <2.0 wt %), including kinetic inhibitors

(KIs) and anti-agglomerants (AAs).^{1,3,6} KIs are generally ineffective at high subcooling (e.g., >10 °C), which is often the case in deepwater. AAs become a promising option because the formation of dispersed small hydrate particles allows for slurry flow.

Gas hydrate plugging is closely linked to the viscosity increase from hydrate particle agglomeration.^{7–10} In pipelines, hydrate particles stick to each other by capillary forces. These aggregates trap free liquid and gas, which results in an effective volume fraction much higher than the true hydrate volume fraction.¹¹ The large aggregates give a higher hydrate effective volume fraction than the true hydrate fraction because of liquid entrapment. AAs reduce the capillary forces, leading to a decrease in the effective hydrate volume fraction, resulting in slurry flow without plugging. Hydrate particle size measurements provide basic data to improve the understanding of hydrate slurry rheology and the anti-agglomeration mechanism.

The general belief in the hydrate literature has been that an oil phase in large quantities is required for hydrate antiagglomeration.^{1,12–15} The mixtures of oil/water/surfactant have been thought to form water-in-oil emulsion, from which hydrate particles can be dispersed in the oil phase when the conditions fall into the hydrate regime.^{1,3} Our recent work has

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demonstrated that hydrate anti-agglomeration can realize in water-in-oil emulsions, oil-in-water emulsions, and also from micelles in which there is no oil phase in both methane and natural gas systems.^{16,17}

Focused beam reflectance measurement (FBRM) is a unique tool to measure particle and droplet sizes in real time. A rotating precision optics focuses a solid-state laser beam at the interface between the probe window and the actual process, as seen in Figure 1a. In this way, the focused beam scans a circular path across the probe window, where individual particles or particle structures will backscatter the laser light to the probe and generate distinct pulses of reflected light. The pulses of backscattered light are detected by the probe and translated into chord lengths based on a simple calculation of the scan speed (velocity) multiplied by the pulse width (time); a chord length is simply defined as the straight-line distance from one edge of a particle or particle structure to another edge, as shown in Figure 1b. The lower limit of sizing measurement is 1 μ m in the probe used in this work.

The FBRM probe can provide measurement in opaque or dark media and in dense suspensions. It has been employed to study gas hydrate particles in recent years.^{12,18–20} The setup can be assembled into an autoclave to study hydrate formation and agglomeration. Turner et al. reported observation of direct transformation of water droplets to methane hydrates in crude oils.¹² The setup can also be integrated into a flow loop to monitor the flow behavior.

Particle vision and measurement (PVM) is another tool for sizing measurement. It provides the real-time imaging of particles and droplets as they exist in crystallization in a vessel or a pipeline. Sapphire lens allow the probe to work at highpressure conditions. PVM is especially useful in capturing the images of large particles.

MATERIALS AND METHODS

The sizing measurement experiments are performed in an autoclave apparatus shown in Figure 2. The effective internal volume of the



Figure 2. Schematic diagram of the autoclave reactor with FBRM and PVM probes.

autoclave is 535 mL (51.84 mm in diameter and 320 mm in depth). The water bath jacket is connected to a Huber chiller for temperature control. The stirring speed is set to 350 rpm in all of the tests. A

Table 1. Composition of Natural Gas (Mole Basis)

FBRM probe (D600X) and a PVM probe (V700S-5-K), both manufactured by Mettler Toledo, are installed in the system for sizing measurements. The particle sizes are measured right before cooling at 20 °C and at 1 °C when the hydrates are fully developed and steady state is reached. The reactor is loaded with 180 mL of liquid sample, a mixture of the additives, and water or brine (4.0 wt % NaCl), before evacuation. Both probe windows are installed below the liquid level. Then, the reactor is charged with test gas (methane at 80 bar and natural gas with compositions presented in Table 1 at 50 bar) with stirring at 20 °C. After the gas inlet valve is closed, the water bath is kept at 20 °C for 30 min with stirring before being cooled from room temperature to 1 °C at the rate of -4 or -10 °C/h. The temperature is then kept at 1 °C for a period of time, allowing the gas hydrates to further develop before the temperature ramps back to the initial temperature. A sharp pressure change indicates hydrate formation/ dissociation

The AA (from Lubrizol Corporation) contains 80-89% cocamidopropyl dimethylamine (as the effective component, shown in Figure 3), 5-10% glycerin, and small amounts of free amine and water. Glycerin and small amounts of amine and water are byproducts of surfactant synthesis. Because the concentration of these byproducts is very low (<0.05 wt %), their thermodynamic effect is expected to be negligible.

RESULTS AND DISCUSSION

In this work, we use the autoclave reactor apparatus to form gas hydrates, including methane hydrate [structure I (sI)] and natural gas hydrate [structure II (sII) because of the presence of ethane and propane]. The hydrate particle sizes are measured by the FBRM probe. We investigate the effect of AA dosage, cooling rate, salt, addition of a small amount of oil, and pH adjustment by NaOH on particle size. In the following, we will first present our results in methane hydrates and then in natural gas hydrates.

Methane Hydrate Tests. We first investigate methane hydrates in water without additive. The change of chord length distribution (CLD) is presented in Figure 4a. Ideally, there should be no chord length detected before hydrate formation when there is no surfactant added. However, gas bubbles in the system show a median size of 4.8 μ m because of strong stirring. The onset temperature of hydrate formation is in the range of 7.0-9.1 °C, and the onset pressure is in the range of 77-78 bar. The onset temperature is lower for the high rate of cooling. For example, in the blank test, the onset of the hydrate formation temperature is 9.1 $^{\circ}$ C, where the cooling rate is -4 $^{\circ}$ C/h. The onset temperature is 7.0 $^{\circ}$ C in the test of 0.50 wt % AA, and the cooling rate is -10 °C/h. Although we have some duplicate runs, we only report results from single runs. The focus of this work is on the size trends after all hydrates have been formed. The sizes reported here refer to the stable values before the temperature ramping. There is, however, a hydrate particle size change during the cooling process. Table 2 presents details of CLD, including median size, mean size (square weighted), and particle counts in different ranges. The median size of hydrate particles is 19.6 μ m measured at 1 °C when there is no AA in the solution. The number of small hydrate particles (<10 μ m) is 3241, and the number of large particles is 1869 in the range of 50–150 μ m.

| component | methane | ethane | propane | <i>n</i> -butane | isobutane | nitrogen | carbon dioxide |
|-----------|---------|--------|---------|------------------|-----------|----------|----------------|
| mol % | 80.67 | 10.20 | 4.90 | 0.753 | 1.53 | 0.103 | 1.83 |

Figure 3. Chemical structures (2D and 3D) of the main component in our AA.



Figure 4. CLD of particles before hydrate formation (blue) and hydrate particles at 1 $^{\circ}$ C (green, -4 $^{\circ}$ C/h; red, -10 $^{\circ}$ C/h) in (a) pure water, (b) 0.2 wt % AA, and (c) 0.5 wt % AA.

Panels b and c of Figures 4 depict the CLD of hydrates in 0.2 and 0.5 wt % AA at different cooling rates (e.g., -4 and -10 °C/h), respectively. The details of CLD are included in Table 2. The weighted mean size of hydrate particles is 47.9 and 49.1 μ m in cooling rates of -4 and -10 °C/h, respectively, at 0.2 wt % AA, which are significantly smaller than 100.3 μ m in the blank test. When the AA concentration increases to 0.5 wt %, the mean size of the hydrate particles is 32.0 and 32.9 μ m in low and high cooling rates. The addition of AA results in a sharp decrease in the size of hydrate particles, as shown in Figure 4 and data in Table 2. For instance, in the range of 50– 150 μ m, the counts are 1869 without AA and 383 and 198 at the AA dosage of 0.2 and 0.5 wt %, respectively. These sizes correspond to the cooling rate of -4 °C/h. At the same hydrate volume fraction, the mixture with a smaller particle size has lower viscosity.

Figure 5 illustrates an image of a large hydrate particle captured by the PVM probe. The particle size is around 180 μ m, and it has an irregular shape, which indicates it is an aggregate of several smaller hydrate particles.

We then investigate the effect of the addition of a small amount of n-octane. Oil-in-water emulsions form when 1 vol % n-octane is introduced into the aqueous phase with AA. A significant effect of *n*-octane is observed in weighted mean size and the number of large gas bubbles/droplets at the start of the cooling, as shown in Table 3. The number of particles with a size of 50–150 μ m decreases from 267 to 13 when *n*-octane is added. The weighted mean size decreases from 28.7 to 19.6 μ m, 32% smaller. Because of the much higher solubility of AA molecules in *n*-octane than in water, AA molecules distribute more evenly in water when a small amount of oil is added. As a result, a higher number of AA molecules can bind onto the hydrate particle surface to prevent agglomeration. The weighted mean size of hydrate particles decreases from 32.0 to 21.6 μ m at the cooling rate of -4 °C/h and from 32.9 to 30.1 μ m at the cooling rate of -10 °C/h. The counts of hydrate particles in the range of $50-150 \ \mu m$ drop from 198 to 14 μm at the low cooling rate and from 362 to 117 μ m at the high cooling rate.

The effect of salt addition on the hydrate particle size is also investigated. NaCl brine of 4 wt % is used in place of fresh water to perform the experiments under the same conditions as discussed above. The comparison of the particle size in brine and freshwater is presented in Table 4. The weighted mean size of hydrate particles is 3 times greater in brine than in freshwater (191.3 versus 47.9 μ m). The increase is from a large number of particles greater than 50 μ m, e.g., 1193 in the range of 50–150 μ m and 262 in the range of 150–300 μ m. The adsorption of ions on the hydrate particle surface increases the hydrate particle size. We have embarked on molecular dynamics simulations for better understanding of the salt effects at the molecular level. Results will be published in due time. The addition of a small amount of n-octane improves the effectiveness of our AA, as indicated by the small number of large particles as Table 4 indicates. In most petroleum fluids, a small amount of hydrocarbon liquids is present, so that there is no need to add *n*-octane.

Natural Gas Hydrate. We also perform four tests in natural gas hydrates. Under test conditions, natural gas forms sII hydrates, different from methane hydrates (sI). We observe foaming as we add AA in the test mixture before hydrate formation; similar observations have been reported in a recent work.¹⁷ The foam increases the gas/liquid interfacial area and reduces the concentration of the surfactant in the aqueous phase. We observe instant plugging from hydrate formation in the natural gas system. The foaming is caused by carbon dioxide in the natural gas, which decreases pH in the aqueous phase, as described by the reaction below.

$$CO_2 + H_2O \rightleftharpoons H_2CO_3 \rightleftharpoons H^+ + HCO_3^- \rightleftharpoons 2H^+ + CO_3^{2-}$$

The molecular structure of AA changes at low pH, and the AA loses effectiveness. To increase the pH of the aqueous phase to a value of above 9, NaOH is introduced into the solution.¹⁷ In

| | $AA = 0.5\%, -10 \ ^{\circ}C/h$ | 5.3 | 32.9 | 28709 | 8949 | 362 | 0 | | | | | e. | |
|----------------------|---------------------------------|-----------|--------------------------|------------|-----------|-----------|-----------|--|--|--|--|--|---|
| Cooling Rates | $AA = 0.5\%, -4 \ ^{\circ}C/h$ | 5.4 | 32.0 | 28171 | 9379 | 198 | 0 | | | Figure 5. the metha | PVM ima ne hydrat | ge of th e test wi | e hydrate pa th no AA. |
| trations at Two (| AA = 0.5%, before | 5.1 | 28.7 | 36547 | 9956 | 267 | 0 | | | the autoc The onse 11.1–13. four tests | lave test, et temper 7 °C, and . NaOH | the ini rature o d the or at a cor | itial pressur of hydrate f nset pressu ncentration |
| Various AA Concen | $AA = 0.2\%, -10 \ ^{\circ}C/h$ | 5.6 | 49.1 | 27250 | 9053 | 351 | 4 | | | test mixtu and addit presented particles of 0, 0.2, vol % <i>n</i> -o is almost wt % AA | ire for pl ion of a l in Tab are 97.1, and 0.5 ctane. W double (| f contro small an le 5. T 48.0, an wt %, re ithout <i>r</i> 68.7 ver | ol. The effe mount of <i>n</i> The weighten and 38.8 μ m espectively, <i>n</i> -octane in rsus 38.8 μ |
| Particles at 1 °C in | $AA = 0.2\%, -4 \ ^{\circ}C/h$ | 5.7 | 47.9 | 26448 | 8919 | 383 | ω | | | There than in to AA dosag are lots of detected instrumen similar te | are signi est 1. Th ge, 0% in of small by the F nt. We c st in Tab | ficantly e only test 1 a particles BRM p compare le 2 (0.2 | less large difference 1 and 0.2 wt s (<1 μ m) robe becau e the partie 2 wt % AA a |
| °C and Hydrate] | AA = 0.2%, before | 4.9 | 46.2 | 22650 | 5474 | 607 | 0 | | | We find t in the ran 50–150 explanatio compared in our ren | hat the h nge of 10 μ m. Th on for t l to meth cent wor | ydrate p $-50 \ \mu r$ he sma he lowe ane hydright k. ¹⁷ | particle num n and 198 ll particle er viscosity drates in rel |
| Formation at 20 | $AA = 0\%, -4 \ ^{\circ}C/h$ | 19.6 | 100.3 | 3241 | 5082 | 1869 | 95 | | | ■ CON In this w sizes <i>in</i> agglomer formulat | CLUSIC ork, we situ to ation mo ion can | N have us improv echanisr reduc | ed FBRM ve the un m. The re e hydrate |
| s before Hydrate | AA = 0%, before | 4.8 | 29.6 | 11615 | 2808 | 102 | 0 | | | have four particle si particle si hydrate si introduce increases | nd that a ize signifi ize redu lurries wl id in the with the | cantly i cantly i ction m nen a sn systen introdu | amount of n both fres nay explain nall amoun n. The size action of 4 |
| CLD of Particle | e and counts | (mu) | nare weighted, μm) | 10 µm |)-50 μm |)−150 µm | 50-300 µm | | | phase. Cr effects o molecula standing. complete | urrently, f salt sj r dynam The ro d. | we are pecies lics sin esults | conducting on hydrate nulations f will be p |
| Table 2. | siz | median () | mean (sq | counts <] | counts 10 | counts 50 | counts 15 | | | ■ AUT Correspo *E-mail: | HOR IN onding A abbas fire | FORM | ATION |



article aggregate (180 μ m) in

re of natural gas is 50 bar. formation is in a range of re is around 48 bar in the of 1 wt % is added in the ct of the AA concentration a-octane in the four tests is ed mean sizes of hydrate at the AA concentrations with 1 wt % NaOH and 1 the mixture, the mean size m) in the mixture with 0.5

hydrate particles in test 2 between these two tests is % in test 2. Possibly there in test 2, which are not se of the resolution of the cle sizes in test 2 to the and $-4 \,^{\circ}C/h$ cooling rate). nbers are 3634 versus 8919 versus 383 in the range of sizes may provide the v of natural gas hydrates lation to ball running times

to obtain hydrate particle derstanding of the antiesults show that our AA particle agglomeration natural gas systems. We *n*-octane reduces hydrate shwater and brine. Hydrate the low viscosity of the t of oil, such as *n*-octane, is e of the hydrate particles wt % NaCl in the aqueous g a systemic study on the e anti-agglomeration and for an improved underublished as findings are

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| Table 3. CLD of Particles before Hydrate Formation at 20 | °C and Hydrate Particles at | : 1 °C with 0.5 wt % AA, | without and with |
|--|-----------------------------|--------------------------|------------------|
| 1 vol % <i>n</i> -Octane (<i>n</i> -C ₈) | | | |

| size and counts | without <i>n</i> -C ₈ , before | without <i>n</i> -C ₈ , -4 °C/h | without <i>n</i> -C ₈ , -10 °C/h | with <i>n</i> -C ₈ , before | with <i>n</i> -C ₈ , -4 °C/h | with <i>n</i> -C ₈ , -10 °C/h |
|---------------------------------|---|--|---|--|---|--|
| median (µm) | 5.1 | 5.4 | 5.3 | 5.1 | 4.8 | 5.5 |
| mean (square weighted, μ m) | 28.7 | 32.0 | 32.9 | 19.6 | 21.6 | 30.1 |
| counts <10 μ m | 36547 | 28171 | 28709 | 37088 | 24017 | 29538 |
| counts 10–50 μ m | 9956 | 9379 | 8949 | 8864 | 5002 | 9097 |
| counts 50–150 μ m | 267 | 198 | 362 | 13 | 14 | 117 |
| counts 150–300 μ m | 0 | 0 | 0 | 0 | 0 | 2 |

Table 4. CLD of Methane Hydrate Particles at 1 $^{\circ}$ C with 0.2 wt % AA in Freshwater and 4 wt % NaCl Brine^{*a*}

| size and counts | 0.2% AA in freshwater | 0.2% AA in brine | 0.2% AA with <i>n</i> -C ₈ in brine | | | | |
|--|-----------------------|---------------------|---|--|--|--|--|
| median (µm) | 5.7 | 5.3 | 5.7 | | | | |
| mean (square weighted, μ m) | 47.9 | 191.3 | 138.8 | | | | |
| counts <10 μ m | 26448 | 18902 | 30915 | | | | |
| counts 10–50 μ m | 8919 | 5169 | 10082 | | | | |
| counts 50–150 μ m | 383 | 1193 | 67 | | | | |
| counts 150–300 μ m | 3 | 262 | 11 | | | | |
| ^{<i>a</i>} The cooling rate is -4 °C/h. | | | | | | | |

Table 5. CLD of Hydrate Particles at 1 $^\circ C$ in Natural Gas at a Cooling Rate of -4 $^\circ C/h$

| size and counts | test 1 ^a | test 2 ^b | test 3 ^c | test 4 ^d |
|---------------------------------|---------------------|---------------------|---------------------|---------------------|
| median (µm) | 6.1 | 5.7 | 9.3 | 5.9 |
| mean (square weighted, μ m) | 97.1 | 48.0 | 68.7 | 38.8 |
| counts <10 μ m | 22342 | 9655 | 14320 | 14145 |
| counts 10–50 μ m | 9137 | 3634 | 11227 | 5613 |
| counts 50–150 μ m | 1198 | 198 | 1863 | 241 |
| counts 150-300 µm | 91 | 2 | 29 | 0 |
| | | | | |

^{*a*}Test 1: 0% AA, 1 wt % NaOH, and 1 vol % *n*-octane. ^{*b*}Test 2: 0.2 wt % AA, 1 wt % NaOH, and 1 vol % *n*-octane. ^{*c*}Test 3: 0.5 wt % AA and 1 wt % NaOH. ^{*d*}Test 4: 0.5 wt % AA, 1 wt % NaOH, and 1 vol % *n*-octane.

Notes

The authors declare no competing financial interest.

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