Improvement of Heavy Oil Recovery in the VAPEX Process using Montmorillonite Nanoclays

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with glass beads and modified montmorillonite as the nanoclay, while they had similar porosity and permeability. The content of deposited asphaltene in swept matrixes, the propagation pattern of vapor chambers in heavy oil matrixes, and the rates of solvent consumption and oil production were determined. The results elucidated that montmorillonite changed the matrix heterogeneity and led to forming enhanced breakthroughs, to increasing the interfacial surface of vapor/bitumen and to accelerating the oil production. It was found that not only was the rate of vapor injection diminished, but the heavy oil recovery was also markedly enhanced by 30(±4)%.

INTRODUCTION

The Importance of the VAPEX Process

The resources of heavy oil in the world are more than twice those of conventional light crude oil. Saniere et al. illustrated that from the available heavy oil only a fraction is extracted by conventional methods [1]. Figure 1 shows the worldwide heavy oil resources and production.

The quota of high-viscosity oil in total worldwide oil production is constantly growing. Large reserves of high-viscosity oils are found in Canada, Venezuela, Mexico, the USA, Russia, Kuwait and China [2]. Average heavy oil production is now about 12% of total world oil production [3]. Different categories of heavy crude are usually defined according to their density and viscosity, as illustrated in Figure 2. Heavy oils, which often result from bacterial oxidation of conventional oils inside the reservoir rock, have different physical and chemical properties, generally degraded: they have much higher viscosity, higher heavy metals and higher sulfur and nitrogen contents [4]. To reduce the viscosity, solvents are frequently used to dilute heavy crudes. This is one of the most efficient methods in pipeline transportation of heavy oils. Solvents are also injected into the reservoir for well cleaning, stimulation, fracturing and, less frequently, for miscible displacement [5].

A large amount of heavy oil is still trapped in reservoirs after the traditional oil extraction, thus a number of enhanced heavy oil recovery processes have been developed for extracting the residual heavy oil [6].

Enhanced recovery processes for heavy oil are mainly thermal, solvent-based and cold methods. In thermal methods such as Steam-Assisted Gravity Drainage (SAGD), Combustion Overhead Gravity Drainage (COGD), Cyclic Steam Stimulation (CSS) and In Situ Combustion (ISC) or Toe-to-Heel Air Injection (THAI), the viscosity is reduced by heating the reservoir, but the major problem is the heat loss to adjacent formations. In cold methods such as Cold Heavy Oil Production with Sands (CHOPS), water flooding, surface mining and Gas-Assisted Gravity Drainage (GAGD), there is no solvent or thermal consumption. However, solvent-based methods such as Vapor Extraction (VAPEX) and the Solvent-Aided Process (SAP) are preferred. Recently, some interesting techniques and hybrid methods have been patented such as combined Steam And Vapor Extraction (SAVEX) and Radiofrequency-Assisted Gravity Drainage (RASD) [7].

Abu-Khader defined the asphaltene constituents as a solubility class that was precipitated from petroleum, heavy oil and bitumen by the addition of an excess of a liquid paraffin hydrocarbon [8]. Mousavi-Dehghani reported many factors which affect the asphaltene precipitation...
The Importance of Nanoclays

The advantage of nanoparticles with respect to their application is attributed to their nano-size, which strongly affects the property changes in the bulk material. Depending on the academic source, there are four main groups of clays: kaolinite, smectite, illite and chlorite. The term of nanoclay is attributed only to the smectite family, which has a unique morphology, featuring one dimension in the nanometer range. Therefore, kaolinite, illite and chlorite are not included as nanoclays.

Montmorillonite (MMT), a member of the smectite family, was discovered in 1847 in Montmorillon in the Vienne prefecture of France, more than 50 years before the discovery of bentonite in the US. MMT is 2:1 clay, meaning that it has 2 tetrahedral sheets sandwiching a central octahedral sheet. The particles are plate-shaped with an average diameter of approximately one micrometer. Individual platelet thicknesses are just one nanometer, but surface dimensions are generally 300 to more than 600 nanometers, resulting in an unusually high aspect ratio.

A characteristic of MMT nanoclays, which makes them special, is their interaction with the surrounding matrix. This special interaction is based upon their modification method and modifiers. MMTs with different structures reveal different reactivity and hence, their modifications are varied. Therefore, special characteristics of MMTs are dependent on their structure, reactivity and modifiers, which are discussed in the following.

The Structure of MMTs

As discussed above, according to Figure 3, MMT is one of the smectite group, composed of silica tetrahedral sheets layered with an alumina octahedral sheet [12]. The imperfections of the MMT crystal lattice with the isomorphous substitution induce a net surface negative charge that leads to the adsorption of metals, cations and sulfur atoms in the interlayer space and external surfaces. MMT also contains dangling hydroxyl end-groups on the surfaces and has a large specific surface area; it exhibits good adsorbability [13].

MMT’s Reactivity

Masih et al. investigated the affinity of Na⁺MMT with the modifier arsenic by preparing different types of iron species [14]. There have been other studies conducted on the cation exchange capacity of MMT [15]. Navratilova et al. studied adsorption of two cationic surfactants, hexadecyl trimethyl...
ammonium bromide and benzyl dimethyl hexadecyl ammonium chloride, onto the samples of MMT [16]. Fusova investigated the intercalation of octadecylamine into the interlayer space of MMT in relation to the quantity of added organic substance [17]. Therefore, organically modified MMTs have become an attractive class of organic/inorganic hybrids because of their potential use in a wide range of applications [18].

**Modified MMTs**

The adsorption hypothesis of modified MMT has been reported by Lin et al. [19] and Rahman et al. [20] for adsorption of Bovine Serum Albumin (BSA) protein and crude oil into the layered silicate MMT. Based upon Lin et al.'s [19] suggested mechanism, the modified MMTs is prone to adsorbing heavy oil asphaltenes and their micelles in their interlayer spaces. Therefore, this tendency of MMTs to adsorb the heavy oil asphaltenes was used to improve the rate of the VAPEX process [21]. Figure 4 illustrates schematically the asphaltene substitutions into the layered Na⁺MMT.

Based upon these methods, in the present study MMTs with d-spacing of 3.15 nm were used in order to adsorb asphaltene into the MMT's spaces. The effects of MMTs on the pattern and content of asphaltene adsorption and upgrading, the propagation pattern of the vapor chamber in a heavy oil matrix, the rates of vapor consumption and oil production, and the recovery factor were studied by comparing two systems, with and without MMTs.

**Thermal Analysis of Crude Oil**

Nowadays, much research is done to analyze the crude oil components such as asphaltenes. In this regard, thermal analysis has provided useful data [22-24]. Thermal analysis is a group of instrumental methods for evaluating different properties and the chemical composition of complex materials by heating the sample at a given rate (normally 10-50°C/min) at different pressures and compositions of the gas medium. Thermogravimetry (TG), Differential Scanning Calorimetry (DSC) and Differential Thermal Analysis (DTA) are the major methods in this field, in which certain parameters (e.g. the amount of absorbed/released heat) as a function of the temperature is recorded. Heating the heavy oil multicomponent system results in evaporation of first, light naphtha–ligroin cuts (up to 200°C), then kerosene–gasoil cuts (200-300°C), and finally, lube oil cuts (>300°C). The different vaporizability of the heavy oil components is the basis of quantitative and qualitative evaluation of the properties of a given component. Kök et al. [25] determined the Saturate, Aromatic, Resin and Asphaltene (SARA) fractions as well as their kinetic parameters in medium and heavy crude oils using thermogravimetric analysis and air atmosphere at a 10°C/min heating rate. Karacan and Kök [26] examined the pyrolysis behavior of crude oils and their fractions using DSC and TG and a 10°C/min heating rate under nitrogen atmosphere. They revealed that pyrolysis mechanisms depend on the chemical nature of the constituents. In another paper, Kök and Okandan [27] investigated the thermal characterization of crude oil using DSC. Moreover, they studied the effect of reaction temperatures and higher heat flow rates.

The aim of this work was to study the enhanced heavy oil recovery using *in situ* modified MMTs. The experiments focused on the nano-assisted dispersion, hence the setup conditions were designed in such a way that the competition phenomena such as de-asphalting were diminished. The effect of MMTs on improvement of the recovery factor and production rate was studied in two systems, one with MMTs and the other without MMTs.

### 1 EXPERIMENTAL METHOD

#### 1.1 Materials

The MMT particles were purchased from Southern Clay Products and their physical properties are summarized in Table 1. The tested bitumen was sampled from the Sarvak formation of the Kuh-e-Mond Reservoir in the south of Iran *(Tab. 2 shows its composition)*. Propane with 99.5% purity at 827 kPa (its dew point) was selected as the solvent since it
has a lower dew point pressure than methane and ethane and allows the experiments to be conducted at lower operating pressures. In addition, the amount of de-asphalting is higher with propane than with butane and pentane. Porous media was made in a sand-packed cell using glass beads.

### TABLE 1

<table>
<thead>
<tr>
<th>Properties</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity</td>
<td>1 660 kg/m³</td>
</tr>
<tr>
<td>Bulk density</td>
<td>172.9 kg/m³</td>
</tr>
<tr>
<td>X-ray diffraction d-spacing (001)</td>
<td>3.15 nm</td>
</tr>
<tr>
<td>Modifier concentration, meq/100g clay</td>
<td>125</td>
</tr>
<tr>
<td>Organic modifier</td>
<td>Dimethyl dihydrogenated Tallow quaternary ammonium</td>
</tr>
</tbody>
</table>

### TABLE 2

<table>
<thead>
<tr>
<th>Composition</th>
<th>Flashed oil</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂</td>
<td>0.00</td>
<td>mol%</td>
</tr>
<tr>
<td>CO₂</td>
<td>0.00</td>
<td>mol%</td>
</tr>
<tr>
<td>C₁</td>
<td>0.00</td>
<td>mol%</td>
</tr>
<tr>
<td>C₂</td>
<td>0.08</td>
<td>mol%</td>
</tr>
<tr>
<td>C₃</td>
<td>0.12</td>
<td>mol%</td>
</tr>
<tr>
<td>C₄ - C₅</td>
<td>7.76</td>
<td>mol%</td>
</tr>
<tr>
<td>C₆ - C₁₁</td>
<td>14.51</td>
<td>mol%</td>
</tr>
<tr>
<td>C₁₂⁺</td>
<td>77.53</td>
<td>mol%</td>
</tr>
<tr>
<td>Asphaltene content</td>
<td>32.30</td>
<td>wt%</td>
</tr>
<tr>
<td>Resin content</td>
<td>3.21</td>
<td>wt%</td>
</tr>
</tbody>
</table>

### 1.2 Methodology to Spike the MMT Nanoclays into the Glass-bead Matrix

Although many procedures have been suggested to coat the granular glass beads, the spraying method was chosen in these laboratory tests. Figure 5 presents schematically the spraying method to spike MMT particles into the glass beads. The suspension of 1.00 g modified MMT in 100 cc volatile condensate was prepared and was sprayed into the rotary mixer. The mixing time was selected to be 30 min (until the condensate was evaporated). Figure 6 shows the SEM image of coated glass beads. According to Figure 6, the MMT particles (white dots) were placed around the glass beads. The doped glass-bead particles were sieved and the required sizes (212-300 μm) were chosen. The density and the permeability of glass beads were determined to be 2 530 kg/m³ and 25-30 D, respectively.

### 1.3 Apparatus and Setup

The flow diagram of the experimental setup is schematically illustrated in Figure 7. The VAPEX cell was a three-dimensional rectangular visual model as a symbol of a reservoir cross-section with 673 × 153 × 31 mm dimensions and 3 300 cc volume, containing 5, 3 and 2 ports at the bottom, top and
on both sides, respectively. This model represents a thin slice of the reservoir (two-dimensional rectangle) that is perpendicular to the horizontal well pair. A Plexiglas window allowed the operator to monitor the shape of the swept zone.

The mass flow meter was calibrated using a nitrogen gas/gas accumulator vessel and the system was examined for leakage by nitrogen gas at 1 034 kPa pressure to pack the cell. The cell was depressurized and the cavity of the model was packed uniformly with the doped glass beads by shaking the model. The doped glass beads were transferred to the cavity of the first cell via one aperture on the side of the body and were shaken for about 60 min, simultaneously. The second cell was packed without MMTs. The absolute porosities of the two cells were calculated and found to be 36.2(±0.1)% and 36.9(±0.1)% for three replications, respectively. Subsequently, the sand-packed model was saturated with preheated bitumen at 80ºC for 12 h and was allowed to cool (to room temperature).

1.4 The VAPEX Methodology

The separator was pressurized by nitrogen gas at 827 kPa. Afterwards, propane was injected into the injection well at the top of the model. The flow rate of injected gas was measured using a calibrated mass flow meter, while the ambient temperature was set and kept at 22(±0.5)°C. The amount of live oil accumulated was monitored through the separator by a calibrated, graduated, site glass. While the oil samples were being obtained, most of the solution gas was liberated due to pressure reduction and directed to the accumulated solution gas in the vessel. After collecting the sample, the rate of free gas production was controlled by a needle valve to be 11 cc/min. During the experiment, a digital camera recorded all visible changes in the solvent chamber. The experiments were run continuously for 21 h. After the test run, the systems were depressurized; the glass beads were sampled from both the gas and bitumen chambers of the two cells. Samples were taken from 35 locations on the model by a test tube.

1.5 Chemical Analysis

The samples were weighed by a calibrated Mettler Toledo digital balance and were placed into a Lindberg furnace (Sola Basic) for about 4 h at 600°C to eliminate residual bitumen. After cooling, the glass beads were weighed again and the weight loss was measured for each sample. The weight loss represented the crude distribution at different points of both chambers. Asphaltenes were determined in the samples from thirteen locations in the vapor chamber. Then, the samples were analyzed via the ASTM D1548 standard test method and thermal analysis. For each test, three measurements were performed and Relative Standard Deviations (RSD) were calculated to be less than 3%.

1.6 Thermal Analysis

In order to run the thermal analysis, matrix samples of very low weights (of the order of 8-10 mg), which increased the quality of the thermograms, were used. The samples were heated in an air atmosphere at the rate of 10°C/min. The weight losses of samples were recorded, while the temperature was raised. After the given temperature, the weight loss on an ordinary analytical balance was determined. The operation was repeated with another sample from the matrix doped with MMTs. After that, the required curves were plotted in weight percent versus standard temperature (°C) coordinates. The DTA curve was also recorded on a scanning calorimeter and the resulting graphs were recorded similarly.

2 RESULTS AND DISCUSSIONS

2.1 Distribution of Asphaltenes and Resins

In the VAPEX experiments with and without MMTs, the asphaltene concentrations at different points of the two cells were determined and the results are presented in Figure 8.

Figure 8 shows the same pattern of asphaltene precipitation from injection to the production wells in both the nano-assisted and conventional VAPEX processes. At the oil-solvent interface, diffusion of solvent into the oil bulk led to
formation of asphaltene micelles, which tended to be adsorbed on the surrounding sides (glass beads or solid particles).

According to Figure 8, two comparative adsorption phenomena are active in the media; adsorption of asphaltene particles by MMTs and deposition of asphaltenes on the matrix surfaces, whose contents were calculated by comparing the remaining asphaltenes in both cells to be in the range of 40-48% of total precipitated asphaltenes, which were the main portion of precipitated asphaltenes.

As discussed before, the MMT crystal lattice induced a net surface negative charge (dangling hydroxyl end-groups) that led to the adsorption of metal, cation and sulfur atoms in the interlayer spaces and external surfaces. These negative charges navigate the adsorption of asphaltene as well as the penetration of resins into the interlayer spaces of the MMT's structure. Figure 9 shows the pattern of resin distribution in the swept zone.

In addition, the results showed that the asphaltene content of the produced oil was markedly lower when MMTs were used. This revealed that more asphaltenes remained in the matrix and the quality of the produced oil was enhanced. The formation damage which is caused by asphaltene deposition can be removed by washing.

Furthermore, the content of residual asphaltene in the swept zone (vapor chamber) is more than the bitumen chamber. The asphaltene distribution is not uniform in the swept zone. Its concentration around the injection well is maximum, while around the production well and both sides it is minimum. Around the injection well, the concentration gradient between solvent and bitumen increases and thereby the mass transfer is enhanced. Hence, more asphaltene will be precipitated. Far from this zone, the reduction of solvent concentration leads to a decrease in the concentration gradient and decline in the asphaltene deposition. Therefore, the precipitated asphaltene decreases markedly around the production well.

2.2 Interpretation of TG/DTA Curves

Figure 10 shows the combined TG/DTA curves of residue hydrocarbons coating the glass beads.

As the temperature rose, first the residual light cuts including light naphtha, kerosene and gasoil evaporated from the sample. This is between ambient temperatures and 200°C, which is not shown in the figure. Then, higher cuts including n-paraffins, light oil cuts, paraffin-naphthenes, naphtheneparaffins and polycyclic aromatic compounds were oxidized.
The oxidation took place without significant thermal effects. Evaporation and oxidation of these cuts were completed at 400°C. The resins, asphaltenes and carbenes were oxidized above 400, 500 and 600°C, respectively.

As shown in Figure 10, there are two vertical isothermal lines in the diagrams, which divide them into three segments including the loss of light (distillate) cuts, resins, asphaltenes and carbene-based cuts. In the figure, the reference temperature is plotted on the X-axis, the weight loss on the left ordinate, the DTA peak height on the right ordinate and the evaporation and oxidation intervals in the first line above the curves.

The left figure shows that by 486°C, the sample loses 28 percent of its weight due to the light distillate portion of the crude and oxidation of resins. At 486°C, an exothermic effect on the DTA curve appears, signaling the beginning of oxidation of the crude oil asphaltenes. Similarly, in the right figure by 495°C, the sample loses 28 percent of its weight due to the light distillate portion of the crude, without noticeable heat effects on the DTA curve. At 495°C, a sharp exothermic effect on the DTA curve depicts the beginning of asphaltene oxidation in the crude oil heavy residue.

The heat evolution continues with varying degrees of intensity until 700°C when finally the entire crude oil heavy residue in the crucible is volatilized and oxidized. The first cut (left segment) revealed the evaporation and oxidation of distillates and resins. The second cut (between isothermal lines) is related to oxidization of asphaltenes, while the loss (burning) of carbenes occurred in the third segment. The weight loss of the TG curve in this middle segment was attributed to the asphaltene content of samples. This procedure can be repeated for the other segments. The asphaltene contents of residue hydrocarbons in the conventional and nano-assisted VAPEX cells were determined to be 28 and 46%, respectively. This was confirmed by the reference method of ASTM D-1548.

2.3 Propagation of Vapor Chamber

The photographs in Figure 11 present a snapshot of the experiments after 9 h. During the VAPEX process, diffusion of solvent dilutes a thin section of bitumen and the diluted live oil drains into the production well. The solvent interacts with bitumen at the horizontal edge on the upper side of the model. The solvent chamber forms and the spreading phase begin and continue up to the end of the 21-h-long experiment.

The picture at the right side of Figure 11 shows the spreading stage using MMTs, while the picture on the left is related to the spreading stage without using MMTs. By comparing the pictures, it can be seen that in the presence of MMTs, the number of breakthroughs is more. By increasing the number of breakthroughs, the contact area between the solvent and oil was increased and the vapor diffusion rate and the production rate as well as the recovery factor were enhanced, respectively. Such a pattern continued up to the end of the test.
Figures 12 and 13 present the flow fluctuations in nano-assisted and conventional VAPEX processes, respectively. The high fluctuations represent the breakthrough formation during the experiments.

The peaks denoted with the letter b are related to breakthroughs. As can be seen from Figures 6 and 7, the numbers of breakthrough peaks in the presence of MMTs are more and their intensity is higher. Furthermore, the lower concentration of asphaltene in the oil lowers the viscosity, making the oil more susceptible to breakthrough.

### 2.4 The Recovery Factor

In Figure 14, the oil recovery results from VAPEX experiments with and without MMTs are presented. It can be seen that higher recovery was demonstrated with nano-assisted experiments; MMTs improved both the recovery factor and production rate in the VAPEX process by a factor of 30(±4)%.

Analysis of variance (ANOVA) was done by using MATLAB software (MATLAB - version 2006) on the oil production rate in simulated VAPEX processes by using a CMG simulator (CMG - version 2006 from Computer Modeling Group), for conventional and nano-assisted VAPEX processes. The results of the ANOVA test are:

- A significant difference between the nano-assisted VAPEX and the conventional VAPEX was indicated. It showed that differences in oil production between the two methods were not affected by random errors during the experiments such as uneven mixing of MMTs, the fluctuation of solvent pressure, temperature, or random asphaltene precipitation and deposition. The significant difference between the two methods elucidated that MMTs are able to change the matrix heterogeneity.
There was no difference between the simulated VAPEX and the conventional VAPEX. This confirmed the accuracy of the simulated model and revealed that the simulated VAPEX and conventional VAPEX processes were matched.

In addition, according to Darcy’s equation (Eq. 1), the production rate is dependent on the interfacial surface area, the viscosity, etc.

\[ Q = \frac{kA}{\mu} \frac{\Delta P}{L} \]

where \( Q \) denotes the flow rate of the produced oil, \( A \) presents the interfacial surface area, and \( k \) and \( \mu \) are the matrix permeability and the fluid viscosity, respectively. The term \( \Delta P/L \) shows the pressure gradient along with the length of the matrix.

In the nano-assisted VAPEX model, not only did the interfacial area increase (for the sake of breakthrough), but the heavy oil viscosity also decreased (for the sake of asphaltene transfer from the oil media into the interlayer spaces of MMTs).

The viscosity of the produced live oil was measured experimentally using an online high-pressure capillary viscometer, which is represented in Figure 7. Figure 15 depicts the viscosity of the produced oil versus the run time. As can be observed from this figure and according to the reasons mentioned above, the MMT nanoclays markedly reduced the oil viscosity.

Therefore, the production rate increased. Moreover, the specific gravity of MMTs is more than heavy oil. After adsorption of asphaltene into the MMTs (intercalation), this difference was even more, and this led to improving the live oil gravity drainage via convensional mass transfer. This led to enhancing the production rate and the recovery factor.

2.5 Solvent Consumption

Breakthrough is a dynamic phenomenon that forms in two-phase systems. The solvent chamber/bitumen system in the VAPEX process is prone to breakthrough formation. During this phenomenon, a part of the injected vapor was lost through the separator without dilution of bitumen. Therefore, the in situ solvent was considered to be distributed in two media; one, which had diffused into the heavy oil, and the other, which was trapped in the pore volumes of the swept zone. The ratio of dissolution efficiency was defined as the rate of dissolved solvent to the rate of injected solvent. The experimental results revealed that the ratio of the dissolved solvent to the injected solvent in the nano-assisted VAPEX method is more than the conventional VAPEX method. Figure 16 shows the comparative diagram of dissolution efficiency for nano-assisted and conventional VAPEX processes.

The MMT’s form enhanced breakthrough with a high aspect ratio to increase the vapor diffusion and the bitumen dilution.

Adsorption of asphaltene particles in the MMT's structure led to reducing the asphaltene deposition on the matrix surfaces. The wettability of the matrix did not shift to oil-wet conditions and the permeability remained constant.

Moreover, intercalation of asphaltenes into the MMTs reduced the oil viscosity, enhanced the gravity drainage, increased the falling rate of live oil, and expanded the interfacial area between the original oil and the vapor chamber. According to the hypothesis of surface renewal, the concentration gradient between the solvent and the heavy oil was increased. This led to increasing the solvent diffusion into the heavy oil as well as the effective consumption of the solvent.
more effective consumption of solvent (30±4%) using MMTs were directly related to the greater contact areas formed by formation of more breakthroughs, adsorption of asphaltenes into the MMT’s interlayer spaces and reduction of oil viscosity.

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REFERENCES


CONCLUSIONS

Quantitative and qualitative analysis of remaining heavy oil asphaltenes and resins in the VAPEX matrix was carried out by TG and DTA methods, respectively. The high content of precipitated asphaltenes and resins in the nano-assisted matrix revealed that asphaltenes and resins were markedly adsorbed into the MMT’s interlayer spaces. Distributions of asphaltenes and resins were not uniform in the swept zone and their concentration around the injection well was maximum, while around the production well and both sides it was minimum. Moreover, the ANOVA test showed a significant difference between the nano-assisted VAPEX and the conventional VAPEX. The difference in the oil production rate between the two methods was not affected by random errors during the experiments. It elucidated that MMTs were able to change the matrix heterogeneity. The greater oil recovery and

3 RECOMMENDATIONS FOR INJECTING THE MMT NANOCLAY

There are some restrictions in selecting the carrier to move the MMT particles from the wellhead to the bottom hole; in the case of MMTs, water is not a suitable carrier, because of the clay swelling. Reduction of carrier viscosity leads to more penetration of the carrier and nanoparticles into the reservoir media and the greater efficiency of the process. Nowadays, some technical methods, such as Coiled Tubing (CT) and a bullhead (BH), are used for acid and hydraulic oil injection into the reservoirs. Injection of nanoparticles into the reservoir can be done via both the solvent injection wells and the live oil production wells by CT or BH methods.

Figure 16
Dissolution efficiency in the presence and absent of MMTs in the VAPEX process.


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