A chelating agent system for the removal of barium sulfate scale

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Abstract
This study aimed to prepare a chelating agent SA-20 using diethylenetriaminepentaacetic acid (DTPA) and low molecular weight sodium polyacrylate raw material for the removal of barium sulfate (BaSO4) scale. The morphology of the resulting chelating agent was characterized by scanning electron microscopy. The BaSO4 scale removal ability of SA-20 was comprehensively evaluated by the dissolving ability, particle size analysis, corrosion performance, and fracture conductivity before and after the scale removal. The results showed that the SA-20 has a good BaSO4 scale removal ability due to the chelation, lattice distortion, and dispersion of DTPA and low molecular weight sodium polyacrylate.

Keywords Chelating agent · Corrosion ability · Dissolving ability · DTPA · Particle size analysis · Sodium polyacrylate

Introduction
With the development of the global oil and gas industry, higher requirements are put forward for drilling and stimulation (Luo et al. 2019). Barite, whose main constituent was barium sulfate (BaSO4), was usually added into a drilling fluid as a weighting material to increase the fluid density of a drilling fluid (Elshreef and Lashin 2016; Fang et al. 2018; Nguyen et al. 2011; Srungavarapu et al. 2018). During drilling, most drilling fluid was lost and entered the reservoir, and solid particles, such as BaSO4, were deposited in the reservoir, resulting in the damage of the reservoir (Al-Yami et al. 2010; Bozau et al. 2015; Liu et al. 2017; Mozaffari et al. 2015, 2017b; Shahbazi et al. 2006). Thus, it is important to develop a chelating system for the removal of the BaSO4 scale. At present, the development of the chelating agent mainly focused on a slightly soluble sulfate scale-dissolving agent (Egermann et al. 2002). Scale-removing agents usually had a compound formula composed of an organic complexing agent (Balanyà et al. 2018), scale inhibitor (Yan et al. 2015; Zhang et al. 2016), surface surfactant (Ratkievicius et al. 2016), and so on. Recently, the study of scale-removing agents focused on aminocarboxylate chelating agent (Uchida et al. 2001). This agent is a kind of carboxylic acid derivative with the main body of aminocarboxylic acid. It could form a stable and water-soluble chelating product easily with many metal ions and thus was widely used for the chemical treatment of slightly soluble scale, which mostly contained salts of Ca, Mg, Ba, and S (Shaw and Sorbie 2015). Besides chelation and dissolution of the scale, an aminocarboxylate chelating agent could wet, disperse, and destroy hard scale blocks, which could be changed to loose scale soil. Generally, the larger the stability constant of the aminocarboxylic acid chelating agent, the stronger the dissolution ability on the salt scale (Gál et al. 2013).

Many studies have been conducted on the blocking removal of a slightly soluble sulfate salt scale, emphasizing the use of aminocarboxylate chelating agents such as representative DTPA and EDTA (Ethylendiaminetetraacetic acid) (A.P. Abib et al. 2018). Despite many successful applications in the BaSO4 scale removal task of many oil fields, many cases were only applied in the ground pipeline at room temperature or shallow reservoir (Elkatatny 2019; Jin et al. 2014; Mahmoud et al. 2018; Wang et al. 2017). Studies on the development of a cleaning agent for deep layers under high temperature and high pressure are scarce. Thus, this study proposed a chelating system SA-20 for the removal of the BaSO4 scale.
under the aforementioned conditions. A system composed of the main agent of DTPA, which was dissolved in sodium hydroxide (NaOH) solution at a pH of 12, and the adjuvant of low molecular weight sodium polyacrylate was prepared. The performance of SA-20 was evaluated by surface morphology analysis, weight change test, particle size analysis, corrosion performance, and fracture conductivity.

**Experimental**

**Material**

DTPA, sodium polyacrylate, BaSO₄, NaOH, acetone, ethanol, and ammonium chloride (NH₄Cl) were of analytical grade and purchased from Chengdu Kelon Chemical Reagent Company of China. The main constituents of steel disks of N80 and 13Cr (About 50 10⁻³ mm³) are shown in Tables 1 and 2, respectively.

The prepare method of SA-20 is: Measuring a certain amount of distilled water into the beaker, adjust the pH value to 8 with sodium hydroxide, weighing a certain amount of DTPA (Depending on the experimental parameters), slowly add it into the distilled water, stir continuously when adding DTPA, after DTPA dissolves, add a certain amount of sodium polyacrylate (Depending on the experimental parameters), fully mix and dissolve.

**Instruments**

The following instruments were used in this study: ZF-9 electronic analytical balance (±0.1 mg, Shanghai Tetragonal Electronic Instrument Factory, China), YHG-9050A thermostatic blast dryer (± 1 °C, Shanghai Yaoshi Instrument and Equipment Factory, China), HELOS-GRADIS Laser Particle Size Meter (± 1%, SYMPATEC GMBH, Germany), scanning electron microscope (SEM, JSM-7500F produced by JEOL), and fracture conductivity instrument (Haian Petroleum Research Instruments Co., Ltd., China).

**Experimental methods**

**Surface morphology analysis**

The micromorphology of the product formed by the chelating agent system SA-20 and Ba²⁺ was characterized by SEM.

**Dissolving ability measured by weight change**

A certain amount of BaSO₄ was weighed into the reaction kettle. Then, a certain amount of SA-20 was also added to it. The reaction kettle was put into the electric drying oven with forced convection and allowed to stand for a while at a specific temperature. Finally, the residue of the reaction was filtered and dried at 80°C for 24 h. The dissolving ability was calculated using the following equation:

\[
D_R = \frac{m_p - m_n}{V} \tag{1}
\]

where \(D_R\) (g/L) is the dissolving ability of SA-20; \(m_p\) (g) is the total weight of BaSO₄ and the filter paper before the reaction; \(m_n\) (g) is the total weight of BaSO₄ and the filter paper after the reaction, filtering, and drying; and \(V\) is the SA-20 volume.

**Particle size analysis**

The particle size distribution of a certain amount of BaSO₄ was measured and put into the reaction kettle. A certain amount of SA-20 was also put into the reaction kettle. The reaction kettle was put into the electric drying oven with 4 h of chemical reaction at 150°C, and the residue of the reaction was filtered and dried at 80°C for 24 h. The particle size distribution of BaSO₄ that permeated and did not permeate the filter paper after the reaction was measured.

**Corrosion ability**

Steel disks of N80 and 13Cr were washed with acetone and ethanol, dried, and weighed. After immersing into SA-20 for 10 h at 90°C, the steel disks were taken out, washed with distilled water, acetone, and ethanol, dried, and weighed. The corrosion ability \(C_R\) was calculated using the following equation:

\[
C_R = \frac{m_p' - m_n'}{tS} \tag{2}
\]

where \(C_R\) (g h⁻¹ m⁻²) is the corrosion ability; \(m_p'\) (g) is the steel disk weight before corrosion; \(m_n'\) (g) is the steel disk weight after corrosion; and \(S\) is the surface area of the steel disk.

| Element | C  | Si  | Mn  | P  | S   | Cr  | V   | Al  |
|---------|----|-----|-----|----|-----|-----|-----|-----|
| Content (%) | 0.34 | 0.22 | 1.55 | 0.18 | 0.013 | 0.14 | 0.13 | 0.02 |
weight after corrosion; \( t (h) \) is the duration of corrosion; and \( S (m^2) \) is the surface area of the steel disk.

**Flow conductivity test before and after the removal of BaSO\(_4\)**

The flow conductivity experiment was performed using the fracture conductivity instrument. The theoretical analysis of conductivity experiment was carried out as follows: Two 7-cm-long cores with a diameter of 2.54 cm were slit along the axis. The BaSO\(_4\) particles were spread over one of the profiles of the core to simulate the blocking of BaSO\(_4\) in microfracture in the reservoir stratum (No. A1). A depth of 100 m fine screens was put first on another core. Then, the BaSO\(_4\) powder was spread over it to simulate the BaSO\(_4\) blocking in a broad fracture in the reservoir stratum (No. A2). The aforementioned cores were put into the core holder. Then, 3% NH\(_4\)Cl base solution, SA-20, and 3% NH\(_4\)Cl base solution were injected into the core holder sequentially. The change in the permeability during the injecting process was measured.

**Results and discussion**

**Surface morphology**

The morphology after the chelation of SA-20 and Ba\(^{2+}\) was characterized by SEM. The characterization result is shown in Fig. 1.

SA-20 could form a stable chelating product with Ba\(^{2+}\). The chelating ability of the main agent DTPA was strong, resulting in the chelating product with high stability constant. Multiple ligands provided multiple pairs of electrons to form coordination bonds with the metal ion Ba\(^{2+}\) at the center. The chelating product with the structure of a five or six membered ring was much more stable than the ordinary coordination compound (Mozaffari et al. 2017a).

**Dissolving ability**

**Influence of pH on the dissolving ability**

Figure 2 shows the dissolving ability of SA-20 on BaSO\(_4\) at different pH values.

The influence of pH on the dissolving ability of SA-20 was large. The dissolving ability continued to increase with increasing pH, reached the maximum at pH 12, and then slowly decreased with a further increase of pH. This was because the carboxyl group mainly existed in the form of...
–COOH in the DTPA molecule under weak alkaline condition, limiting the chelating performance with Ba\(^{2+}\). With a gradual increase in pH, H\(^+\) of the carboxyl group ionized to form –COO\(^-\), which could strongly chelate with Ba\(^{2+}\). Thus, the dissolving ability increased. However, under the strong alkaline condition with overlarge pH, metal hydroxide was formed and crystallized to influence the chelation with Ba\(^{2+}\), decreasing the dissolving ability.

**Influence of sodium polyacrylate concentration on the dissolving ability**

Figure 3 shows the dissolving ability of SA-20 on BaSO\(_4\) at different concentrations of sodium polyacrylate.

The dissolving ability of SA-20 on BaSO\(_4\) increased initially, reached a maximum, and further decreased with the increase of the concentration of the adjuvant sodium polyacrylate. This was because large amounts of negative polyacrylate ions ionized from low molecular weight sodium polyacrylate were strongly adsorbed on the surface of BaSO\(_4\) particles, which thus possessed an identical negative charge. Other ions (Na\(^+\)) with opposite charge freely diffused to the peripheral liquid medium to form a diffusion layer with charged ions, resulting in the electric double-layer structure with a certain electric potential. When the BaSO\(_4\) particles approached each other, particles with identical charge formed electric repulsion to maintain a certain distance between each other, which prevented the aggregation and flocculation of particles. Thus, the dispersion was realized, resulting in full contact between the main agent DTPA and BaSO\(_4\) particles as well as the complete reaction, which resulted in a high dissolving ability.

When a small amount of low molecular weight sodium polyacrylate was added, particles that were not surrounded by polyacrylate ion aggregated due to the electric attraction and decreased the contact area of DTPA and BaSO\(_4\) particles, resulting in incomprehensive reaction and low dissolving ability. When the amount of low molecular weight sodium polyacrylate was too large, excessive polyacrylate ions led to the changes in the ionic environment in the dispersion solution; the interface adsorption reached saturation, and excessive hydrocarbon bonds flocculated through hydrophobic association to synthesize micelles, thereby reducing the contact area between DTPA and BaSO\(_4\) particles, resulting in insufficient reaction and low dissolution capacity.

**Influence of DTPA concentration on the dissolving ability**

Figure 4 shows the dissolving ability of SA-20 on BaSO\(_4\) with different concentrations of DTPA.

At 90 °C, the BaSO\(_4\) dissolving ability of SA-20 increased initially, reached the maximum at a concentration of 6%, and decreased with the continuous increase of the DTPA concentration. At 150 °C, the BaSO\(_4\) dissolving ability of SA-20 increased initially, reached 15.87 g/L at a concentration of 6%, and then remained almost unchanged with the continuous increase of DTPA concentration due to the influence of threshold effect (Mahmoud et al. 2018). The small addition of DTPA could stably dissolve much Ba\(^{2+}\) in water, whereas the chelating performance did not increase dramatically or even decreased when exceeding a certain concentration.

**Influence of the reaction time on the dissolving ability**

Figure 5 shows the dissolving ability of SA-20 on BaSO\(_4\) at different reaction times.

At 90 °C, due to the decrease in mass transfer rate in the reaction system, the dissolving ability was mainly influenced by reaction time. With the increase in time, the dissolving ability gradually increased initially and then reached a plateau. At 150 °C, due to the fast mass transfer rate, the chelating product of the reaction could rapidly diffuse to the fluid system that was far from the BaSO\(_4\) solid surface. Thus, the
dissolving ability could reach the limit in a short time of 4 h and then remained constant afterward.

Influence of temperature on the dissolving ability

Figure 6 shows the dissolving ability of SA-20 on BaSO$_4$ at different temperatures.

With the increase in the reaction temperature, the dissolving ability of SA-20 on BaSO$_4$ increased initially, reached 14.2 g/L at 110 °C, and then increased slowly with a small change due to the low stability constant $K_{sp}$ of the chelating reaction at low temperature, which made the chelation of Ba$^{2+}$ ion in the solution system not effective. However, when the reaction temperature increased to a certain extent, not only the stability constant $K_{sp}$ but also the mass transfer rate increased, accelerating the chelating reaction and the dissolution of BaSO$_4$. However, the stability constant $K_{sp}$ did not increase without limit. Thus, the increasing change in the dissolving ability at temperatures between 150 and 170 °C was gradually small. Besides, the high temperature would not influence the dissolving ability of SA-20 due to the good stability of the blocking removal agent at high temperature.

Influence of other scales on the dissolving ability

Mg(OH)$_2$ and CaCO$_3$ were common scales in the oil fields that were easier to be dissolved than BaSO$_4$. Related experiments were conducted to study the influence of Mg(OH)$_2$ and CaCO$_3$ on the dissolving ability of SA-20. Table 3 shows the dissolving ability of SA-20 with the existence of Mg(OH)$_2$ and CaCO$_3$.

Table 3: Influence of Mg(OH)$_2$ and CaCO$_3$ on the dissolving ability of SA-20

| No. | Scale type | Moles (mol) | Dissolving moles (mol) | Dissolving amount (g) | Total dissolving amount (g) |
|-----|------------|-------------|------------------------|-----------------------|---------------------------|
| 1   | Mg(OH)$_2$ | 0.025       | 0.01122                | 0.6508                | 2.5124                    |
|     | CaCO$_3$  | 0.025       | 0.00555                | 0.5545                |                           |
|     | BaSO$_4$  | 0.025       | 0.00561                | 1.3071                |                           |
| 2   | CaCO$_3$  | 0.03        | 0.00825                | 0.825                 | 3.144                     |
|     | BaSO$_4$  | 0.03        | 0.00995                | 2.319                 |                           |

No impurity of either floccules or precipitate was observed after 15 days of the residue following the reaction at high temperature (Fig. 7). This indicated that the blocking removal agent SA-20 possessed a good precipitation inhibition ability.
The analysis of the No. 2 scale sample showed that the advantage of the higher polarity of $\text{Ba}^{2+}$ than $\text{Ca}^{2+}$ became more dramatic with only $\text{CaCO}_3$ and $\text{BaSO}_4$ and without the competition of $\text{Mg}^{2+}$; thus, $\text{BaSO}_4$ reacted with the SA-20 more easily. In the field application, it was possible to pump a plug of acid solution to dissolve acid-soluble blockage of $\text{Mg(OH)}_2$ and $\text{CaCO}_3$ before the injection of SA-20. Thus, the blocking remover SA-20 could specifically solve the blocking problem of barite.

**BaSO$_4$ particle size analysis**

The $\text{BaSO}_4$ dissolution by SA-20 could smash a large block of $\text{BaSO}_4$ scale into the small ones. Figures 8, 9, and 10 are the distribution curves of $\text{BaSO}_4$ particles before reaction, $\text{BaSO}_4$ particles that did not permeate, and those that permeated through the filter paper after the reaction, respectively. Table 4 shows cumulative particle size distributions of 10%, 50%, and 90% before and after the reaction.

The average particle size is about 2.7 $\mu$m before adding the chelating agent, the average size of barite particles that did not permeate through the filter paper is about 1.8 $\mu$m after adding the chelating agent, the average size of barite particles that permeate through the filter paper is about 1 $\mu$m after adding the chelating agent. The particle size is mainly distributed between 0.4 and 1000 $\mu$m before adding the chelating agent. The particle size of the barite particles that did not permeate through the filter paper is mainly distributed between 0.3 and 30 $\mu$m, and the particle size of the barite particles that permeate through the filter paper is mainly distributed between 0.15 and 5 $\mu$m after adding the chelating agent.

The particle size of $\text{BaSO}_4$ dramatically decreased after the reaction with SA-20. Some too small $\text{BaSO}_4$ particles could even permeate through the quantitative filter paper slowly, illustrating the excellent ability to smash $\text{BaSO}_4$ particles of SA-20. This was due to the physical and chemical adsorption of the dispersive agent sodium polyacrylate on the $\text{BaSO}_4$ particle surface, which produced charge aggregation and the electric double layer around the particle surface. The resultant repulsion between particles made them suspended and dispersed. Thus, $\text{BaSO}_4$ was dispersed and smashed. Besides, the particles dispersed in the system increase the contact area with SA-20, which greatly improves the reaction rate.

**Corrosion ability**

DTPA in SA-20 had a metal corrosion performance to some extent. The N80 and 13Cr corrosion ability of SA-20 was tested. Table 5 shows the N80 and 13Cr steel disk corrosion rate of SA-20, and Fig. 11 shows the appearance of the steel disk before and after the corrosion.

Almost no change was observed on the surface of N80 and 13Cr steel disks. The average N80 steel disk corrosion rate of SA-20 was only 0.1182 g m$^{-2}$ h$^{-1}$, whereas the average 13Cr steel disk corrosion rate was lower, only 0.0388 g m$^{-2}$ h$^{-1}$, which could fulfill the field usage requirement, with no adverse effect on tubing and casing.

**Flow conductivity ability test**

The $\text{BaSO}_4$ blocking dissolution performance of SA-20 was measured using the fracture flow conductivity test before and after $\text{BaSO}_4$ blockage dissolution in the fracture. Figure 12 shows the profile and the end surface morphology in microfracture before and after $\text{BaSO}_4$ dissolution (No. A1). Figure 13 shows the morphology of the profile and the end surface in a broad fracture before and after $\text{BaSO}_4$ dissolution (No. A2). The red line in Figs. 12 and 13 represents the...
oil and gas flow path formed by the dissolution of BaSO4. Figure 14 shows the flow conductivity change curve in microfracture before and after BaSO4 dissolution (No. A1). Figure 15 shows the flow conductivity ability change in a broad fracture before and after BaSO4 dissolution (No. A2). The y-axis represents the ratio of fracture permeability to initial permeability in the experimental process, and the abscissa represents the ratio of the cumulative volume of injected liquid to fracture volume in Figs. 14 and 15.

Figure 12 and Fig. 13 indicate that after thorough wetting, chelating, and dispersion, BaSO4 in fracture was gradually dissolved, dispersed, and brought out. A flowing passage was formed apparently by the corrosion and scouring of BaSO4 in the profile of the fracture, greatly increasing the permeability of the fracture. Figures 14 and 15 indicate that after injecting of SA-20 into A1 and A2 cores, the permeability recovery rate was 340 and 590%.

### Table 4 BaSO4 particle size before and after the reaction

| BaSO4 particle size before the reaction (μm) | BaSO4 particle size that did not permeate through the filter paper (μm) | BaSO4 particle size that permeated through the filter paper (μm) |
|-------------------------------------------|------------------------------------------------------------------------|-----------------------------------------------------------------|
| d(0.1) 1.013                              | 0.570                                                                  | 0.279                                                           |
| d(0.5) 2.706                              | 1.861                                                                  | 1.071                                                           |
| d(0.9) 8.226                              | 7.150                                                                  | 2.437                                                           |

### Table 5 N-80 and 13Cr steel disk corrosion rate of SA-20

| Steel disk type | Tested diskNo | Length (cm) | Width (cm) | Thickness (cm) | Weight before the corrosion (g) | Weight after the corrosion (g) | Corrosion rate (g m⁻² h⁻¹) | Average corrosion rate (g m⁻² h⁻¹) |
|-----------------|---------------|-------------|------------|----------------|-------------------------------|-------------------------------|-------------------------------|----------------------------------|
| N80             | 254           | 4.91        | 0.97       | 0.30           | 10.8766                       | 10.8751                       | 0.1128                        | 0.1182                           |
|                 | 998           | 4.92        | 0.98       | 0.23           | 10.7277                       | 10.7261                       | 0.1236                        |                                  |
| 13Cr            | 102           | 4.98        | 1.11       | 0.22           | 11.4511                       | 11.4506                       | 0.0354                        | 0.0388                           |
|                 | 185           | 4.94        | 1.04       | 0.38           | 11.5846                       | 11.5840                       | 0.0422                        |                                  |

Fig. 11 Appearance of steel disks before and after corrosion. a Before BaSO4 dissolution. b After BaSO4 dissolution

(a) N80 steel disk before and after the corrosion.  
(b) 13Cr steel disc before and after the corrosion.

Fig. 12 Profile and the end surface morphology in microfracture before and after BaSO4 dissolution (No. A1). a Before BaSO4 dissolution. b After BaSO4 dissolution

(a) Before BaSO4 dissolution.  
(b) After BaSO4 dissolution.
respectively, illustrating that the SA-20 system possessed good blocking removal ability. Moreover, it also illustrated that the physical properties of the fractured reservoir with low porosity would be severely influenced after the damage of the barite. The oil and gas production enhancement potential would be huge after the blocking removal procedure.

Dissolution process and mechanism

The blocking removal process of SA-20 included the softening and dissolution of BaSO₄ solid phase and the dispersion and suspension of BaSO₄ particles to form the solution by chelating function (Fig. 16). The BaSO₄ dissolution mechanism by SA-20 mainly included chelation, lattice distortion, and dispersion.

Chelating

SA-20 was composed of the main agent of DTPA and the adjuvant of low molecular weight sodium polyacrylate. Figure 17 shows eight active coordination bonds, including three Ns and five Os in each molecule of DTPA, providing eight coordination bonds around a free metal ion. Thus, it was a good metal ion chelating agent. In alkaline solution, DTPA gradually changed into a negative ion cluster of DTPA⁵⁻. The resultant Ba-DTPA chelating product

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**Fig. 13** Morphology of the profile and the end surface in a broad fracture before and after BaSO₄ dissolution (No. A2)

**Fig. 14** Flow conductivity ability changes in microfracture before and after BaSO₄ dissolution (No. A1)

**Fig. 15** Flow conductivity ability changes in a broad fracture before and after BaSO₄ dissolution (No. A2)

**Fig. 16** Dispersion, chelation, and dissolution process of BaSO₄ by SA-20
possessed high stability, whose chelating reaction equation is shown as follows:

\[
DTPA^{5-} + Ba^{2+} \rightarrow Ba - DTPA^{3-}
\]  

(3)

The stability constant \( lg(K_{sp}) \) of the reaction of \( DTPA^{5-} + Ba^{2+} = BaDTPA^{3-} \) was 8.78 (Wang et al. 1999). The adjuvant low molecular weight sodium polyacrylate (500–10,000) was electrolyte with functions of dispersion and scale inhibition. Thus, low molecular weight sodium polyacrylate was chosen as the adjuvant of SA-20 to realize both dispersion and blocking removal.

**Lattice distortion**

DTPA entered the crystal structure and distorted the lattice. The original structure changed, producing dislocation and distortion in the lattice. Figure 18 shows that the interaction between particles decreased, thus decreasing the hardness of \( BaSO_4 \) that was made easier to be dispersed in the solvent (Darjani et al. 2019, 2017).

**Dispersion function**

The adjuvant of low molecular weight sodium polyacrylate was mainly used as the dispersive agent and the scale inhibitor. The corresponding relatively high charge density produced charge aggregation on the surface of the \( BaSO_4 \) particles and the electric double layer as well as the resultant repulsion (Fig. 19).

The dispersion of particles into the fluid system of the blocking remover SA-20 increased the contact area between SA-20 and the blockage \( BaSO_4 \), realizing the acceleration of the reaction rate. The addition of the dispersive agent of sodium polyacrylate in SA-20 was very effective for blockage removal. After the reaction between the blockage remover, which contained the dispersive agent, and \( BaSO_4 \), the residue was transparent and had good dispersion effect, whereas after the reaction with the blockage remover, which did not contain the dispersive agent, the residue was in a state of muddy emulsoid with a large amount of unreacted \( BaSO_4 \) and with a poor dispersion effect (Fig. 20).

**Conclusions**

(1)SA-20 was composed of the main agent DTPA and the adjuvant low molecular weight sodium polyacrylate. \( BaSO_4 \) blocking could be effectively removed by the...
chelating function and lattice distortion of DTPA, and the dispersion function of sodium polyacrylate.

(2) The influence of pH, sodium polyacrylate concentration, and DTPA concentration on the dissolving ability of SA-20 was large, wherein the dissolving ability had a peak value. The dissolving ability increased with the increase in the reaction time and reaction temperature, though within a limit.

(3) Physical and chemical adsorption function occurred between sodium polyacrylate and BaSO₄. The electric double layer on the surface of the particles could produce repulsion between particles, causing them to be in a state of dispersion and suspension. BaSO₄ was thus dispersed and smashed. Due to the increased contact area between SA-20 and BaSO₄ particles, the reaction rate and efficiency were greatly enhanced.

(4) The steel disk corrosion ability of SA-20 was extremely low, whereas the dissolving ability of SA-20 on BaSO₄ was relatively high. The blocked BaSO₄ in the fracture could be dissolved, forming a flowing passage with high conductivity ability, which thus increased the permeability of the fracture in the core.

Fig. 20 The residue of the chemical reaction between blockage remover and BaSO₄

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