Research Article

Synthesis of a Novel Filtrate Reducer and Its Application in Water-Based Drilling Fluid for Ultra-High-Temperature Reservoirs

Dongyu Qiao,1,2 Zhongbin Ye,1,3 Lei Tang,1 Yiping Zheng,2 Xindong Wang,2 and Nanjun Lai1,3

1School of Chemistry and Chemical Engineering of Southwest Petroleum University, Chengdu, 610500 Sichuan, China
2Engineer Technology Research Institute, CNPC Xibu Drilling Engineering Company Limited, Urumqi 830001, China
3Oil & Gas Field Applied Chemistry Key Laboratory of Sichuan Province, Chengdu, 610500 Sichuan, China

Correspondence should be addressed to Zhongbin Ye; yezb@swpu.edu.cn and Nanjun Lai; lainanjun@126.com

Received 11 September 2021; Accepted 1 October 2021; Published 29 October 2021

Academic Editor: Jinjie Wang

Copyright © 2021 Dongyu Qiao et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

The high-temperature stability and filtration property controlling of ultra-high-temperature water-based drilling fluids is a worldwide problem. To resolve this problem, a high-temperature-resistant quaternary copolymer (HTRTP) was synthesized based on molecular structure optimization design and monomer optimization. The physical and chemical properties were characterized by infrared spectroscopy, thermal weight, and spectrophotometry, and their temperature and salt resistance was evaluated in different drilling fluids, combined with adsorption, particle size analysis, and stability test. The results show that the thermal stability of HTRTP is very strong, and the initial temperature of thermal decomposition is above 320°C. The salt resistance of HTRTP is more than 162 g/L, and the calcium resistance is more than 5000 mg/L, which is equivalent to the foreign temperature-resistant polymer DCL-a, and is superior to the domestic metal ion viscosity increasing fluid loss agent PMHA-II for drilling fluids. It has excellent high-temperature resistance (245°C) and fluid loss reduction effect in fresh water base mud, fresh water weighted base mud, saturated brine base mud, and composite salt water base mud, which is better than foreign DCL-a (245°C) and domestic PMHA (220°C). The adsorption capacity of HTRTP on clay particles is large and firm, and the adsorption capacity changes little under the change of chemical environment and temperature. Both before and after HTRTP aging (245°C/16 h), the permeability of filter cake can be significantly reduced and its compressibility can be improved. By optimizing the particle size gradation of the drilling fluid and enhancing the colloid stability of the system, HTRTP can improve the filtration building capacity of the drilling fluid and reduce the filtration volume. The development of antithermal polymer provides a key treatment agent for the study of anti-high-temperature-resistant saline-based drilling fluid.

1. Introduction

The fluid loss additive, also known as the fluid loss control preparation and water loss additive, is an important drilling fluid treatment agent to ensure the stability of drilling fluid performance, reduce the loss of harmful fluids to the formation, stabilize the well wall, and ensure the well diameter rule [1, 2]. At present, the following categories of fluid loss agents are used at home and abroad: (1) modified lignite [3]; (2) modified starch [4]; (3) modified cellulose [5]; (4) modified resin [6]; and (5) olefin monomer polymers [7]. With the development of the depth of petroleum exploration and development, the drilling depth keeps deepening and the bottom hole temperature is getting higher and higher, which puts forward higher requirements for the control of drilling fluid filtration and rheological properties [8, 9]. Commonly used natural and natural modified fluid loss reducers could not meet the needs of deep and ultradepth well drilling. Polymer fluid loss additive not only reduces fluid loss but also regulates rheological properties [10, 11]. It is an indispensable key treatment agent for high-temperature and ultra-high-temperature water-based drilling fluids [12, 13]. Since
the 1970s, the new synthetic polymer drilling fluid treatment agent which is resistant to high temperature and electrolyte pollution has been studied at home and abroad and has been applied in deep and ultradeep well drilling [14, 15]. With excellent performance of resistance to high temperature and salt resistance, the treatment agent has the following several kinds: Oseh et al. [16] on synthesizing a polypropylene-silica nanocomposite (PP–SiO₂) NC using hot–emulsion sol–gel method as a viscosity modifier and filtration control agent for WBMs. PP–SiO₂ NC is spherical, with a particle size distribution between 80 and 390 nm, finely dispersed; a stronger resistance from 430°C to 485°C; and better filtration control and rheological properties, and plastic viscosity is reduced by 22.7%. Chang et al. [17] have developed a new environmentally friendly fluid loss agent nano-LS (nano-LS) with good thermal stability, salt tolerance, and calcium resistance. The water-based drilling fluid prepared with it maintains its stability at 200°C and can control the water loss at 7.5 mL. The high-temperature- and salt-resistant polymer DCL-a, developed by Philips and Chevron, can effectively reduce the high-temperature and high-pressure filtration of drilling fluid. The temperature resistance is more than 230°C, and it has strong salt and calcium resistance. At present, the thermal resistance of polymer filtrate reduction agents at home and abroad still cannot meet the requirements of ultra-high-temperature filtration control in deep and ultradeep well drilling [18, 19]. Therefore, this paper carried out the synthesis and characterization of high-temperature- and salt-resistant quaternary polymer, evaluation of filtration loss performance and stability in driling fluid, and analysis of its action mechanism [20, 21].

2. Experimental

2.1. Material. Anhydrous sodium carbonate, 2-methyl-2-acrylamide propanesulfonic acid (AMPS), N-vinylpyrrolidone (NVP), N, N-diethylacrylamide (DEAM), and dimethylallylammonium chloride (DMDAAC) are all industrial products for the industrial pilot amplification production [22, 23]. The initiator and sodium hydroxide were analytical grade pure. Drilling fluid with metal ions increased sticky fluid loss agent PMHA from the Xinjiang oil field (high-temperature-resistant polymer DCL-a, foreign ConocoPhillips company).

IRAffinity-1 Fourier Transform Infrared Spectrometer (Shimadzu Corporation, Japan), HTG-1 thermogravimetric analyzer (Beijing Hengliu Scientific Instrument Factory), Model 722 UV-Vis Spectrophotometer (Shanghai Precision Instrument Co., Ltd.), ZNS2 medium pressure filter loss instrument (Qingdao Haitongda Special Instrument Co., Ltd.), and ZNN-D6 model six-speed rotary viscometer (Qingdao Haitongda Special Instrument Co., Ltd.) were the materials used.

2.2. Synthesis of Quaternary Copolymer

2.2.1. Synthesis Principle

(1) Competitive Rate of Polymer. The reactivity rate is the ratio of the rate constants of homopolymer (self) growth and copolymer (cross) growth. The reactivity ratios of the designed high-temperature-resistant quaternary copolymers can be estimated using the - equation proposed by Alfrey and Price. The calculated values of the reactivity ratios of the four monomers (Table 1) show that the product of the reactivity ratios between the monomers is less than 1, indicating that the nonideal copolymerization occurs in this synthesis [24].

(2) Structural Formula. According to the principle of polymerization reaction, the possible structural formula of HTRTP is shown in Figure 1.

2.2.2. Synthesis. The reaction monomers (AMPS : DEAM : DMDAAC : NVP = 6 : 3 : 3 : 1) were added in the four-point flask, and then, the reaction monomers were added in turn. The pH value of the system was adjusted to 7.0 with 30% NaOH solution, and the temperature was raised to 60°C. The initiator of 0.1% of the total monomer was added, and the viscous liquid product was obtained by stirring for 4 h. The product was dried and crushed at 105°C to obtain HTRTP in powder form.

2.3. Characterization

2.3.1. Composition and Molecular Weight. Acetone was used as precipitant, and HTRTP was purified and dried by solution precipitation. The structure of HTRTP was analyzed by Fourier transform infrared spectroscopy (FT-IR) with KBr.

The “one-point method” was used to determine the intrinsic viscosity of the polymer (fluid loss additive), and the viscosity average molecular weight of the polymer was estimated by the Mark-Houwink empirical formula [25]. The calculation is as follows:

\[ M = \sqrt[\frac{2}{k_c}]\left(\frac{2}{(\tau/\tau_0)} - 1\right) \ln \left(\frac{\tau}{\tau_0}\right) \]

where C is solute concentration, g/mL; M is the molecular weight of the solute; and K and α are constants related to the determination conditions and polymer structure, and the value of C is 3.684 × 10⁻², and the value of α is 0.646.

2.3.2. Resistance to Temperature. In N₂ atmosphere, the purified HTRTP was subjected to thermogravimetric analysis on a thermogravimetric analyzer at a temperature of 29°C-800°C and a heating rate of 5°C/min [26].

With 0.2 mol/L NaCl solution as solvent, 0.2% HTRTP and DCL-a solutions were prepared, and their intrinsic viscosities after aging at different temperatures for 8 hours were measured by an ohmmeter.

2.3.3. Ability to Antisalt. The salting-out resistance of high-temperature-resistant quaternary copolymer was tested and evaluated by UV-visible photometry [27]. In making polymer aqueous solution (PMHA, DCL-a, and HTRTP), the concentration is 0.5%. Add different amounts of distilled water and 180 g/L sodium chloride solution to a test tube
2.4. Evaluation of Filtration Loss Reduction Effect

2.4.1. Evaluation of Fluid Loss Reduction Effect in Fresh Water Weighted Base Slurry. The experimental base slurry is composed of 400 mL tap water + 1.2 g anhydrous sodium carbonate + 6% secondary bentonite + 5% evaluation soil + 30% barite, respectively, to the early hydration good base paste to add 1% of the sample evaluation (PMHA, DCL-a, HTRTP), stirring evenly, into the high-temperature aging tank, respectively, under 220°C and 245°C hot rolled out after 16 h, measuring the rheological property and filtration.

The suspension was agitated vigorously and then left to stand for 20 min to allow the adsorption process of the quadripolymer to complete at given temperature. The suspension was subsequently centrifuged for 30 min at a speed of 1000 rpm using a KUBOTA 3500 centrifuge. The concentration of the unadsorbed quadripolymer, $c_0$, located in the clear centrifuged supernatant was measured via absorbance at wavelengths of 390 nm and 330 nm for polymer solution using an ultraviolet spectrophotometer (722, Shanghai Yuanxi Instruments Co., Ltd., China). Applying material balance, the adsorption of polymer on clay particles can be calculated via

$$\tau = \frac{\nu(c_0 - c_1)}{G},$$

(2)

where $\tau$ is the adsorption rate, $\nu$ is the flow rate, $c_0$ is the initial concentration of polymer, $c_1$ is the concentration of polymer in the supernatant, and $G$ is the rate of mass transfer.

2.4.2. Effect of Fluid Loss Reduction in Saturated Salt-Water-Based Slurry. The experimental base slurry is composed of 400 mL tap water + 1.2 g anhydrous sodium carbonate + 6% secondary bentonite + 10% evaluation soil + 30% sodium chloride, respectively, to the early hydration good base paste to add 1.5% of the sample evaluation (PMHA, DCL-a, HTRTP), stirring evenly, into the high-temperature aging tank, respectively, under 220°C and 245°C hot rolled out after 16 h, measuring the rheological property and filtration.

2.5. Research on Mechanism of Action

2.5.1. Adsorption Characteristics on Clay Particle Surface. A known amount of polymer solution was added to 4 wt% calcium montmorillonite (purified with 15 wt% H₂O₂ suspension and then diluted with 50 mL distilled water. The initial concentration of polymer was recorded as $c_0$. The suspension was agitated vigorously and then left to stand for 20 min to allow the adsorption process of the quadripolymer to complete at given temperature. The suspension was subsequently centrifuged for 30 min at a speed of 1000 rpm using a KUBOTA 3500 centrifuge. The concentration of unadsorbed quadripolymer, $c_1$, located in the clear centrifuged supernatant was measured via absorbance at wavelengths of 390 nm and 330 nm for polymer solution using an ultraviolet spectrophotometer (722, Shanghai Yuanxi Instruments Co., Ltd., China). Applying material balance of the polymer, the adsorption of polymer on clay particles can be calculated via

$$\tau = \frac{\nu(c_0 - c_1)}{G},$$

(2)
where \( \tau \) is the mass of polymers absorbed onto specific weight calcium montmorillonite, mg/g; \( v \) is volume of solution, L; \( c_0 \) and \( c_1 \) are polymer concentration before and after polymer absorbed onto calcium montmorillonite, mg/L; and \( G \) is the mass of calcium montmorillonite, g.

2.5.2. Influence on Filter Cake Quality. The effect of its addition in fresh water base slurry on the permeability of filter cake was investigated. After measuring the API filtration loss, carefully remove the substrate slurry on the surface of the filter cake \([29]\), replace it with distilled water, measure the volume of water for 30 min, measure the thickness of the filter cake with a steel plate ruler, and calculate the permeability \( K \) of the filter cake according to the following formula:

\[
K = 0.175vh \times 10^{-6},
\]

where \( K \) is the permeability of mud cake, \( \mu m^2 \); \( v \) is the effluent volume, mL; \( h \) is the thickness of mud cake, mm; and 0.175 is the conversion factor (0.07 for high temperature and high pressure).

The effect of several treatment agents on the compressibility of base mud was investigated by using the “two-stage water loss method” which is simple, feasible, and reliable. If other factors are set to be constant, the compressibility of filter cake can be expressed by a simple proportional relationship of filtration rate at 500 psi and 100 psi:

\[
R = \frac{V_{500}}{V_{100}},
\]

in which \( R \) is the compressibility coefficient of the filter cake and \( V_{500} \) and \( V_{100} \) are static filtration loss under pressure difference of 500 psi and 100 psi, respectively.

2.5.3. Particle Size and Specific Area Measurements of Calcium Clay. The suspensions were prepared by mixing deionized water, 4% (w/v) calcium montmorillonite, and 0.3% (w/v) viscosity breaker (XY-28, sulfonated tannin and quadripolymer separately), stirring for 20 minutes at a high speed of 10,000 rpm and aging for 24 hours at room temperature. Aging experiments of suspensions were carried out in a XGRL-4A-type rolling oven through hot rolling at 220°C (XY-28) or 245°C (sulfonated tannin and quadripolymer) for 16 hours. Concentrated NaOH (10 mol/L) and HCl (6 mol/L) solutions were used to change pH so as not to cause excessive dilution of samples. The size distribution and specific area of clay particles in suspension were determined using a Bettersize2000 laser particle analyzer.

2.5.4. Influence on Stability of Fresh Water Base Slurry Colloid. Turbiscan, an infrared scanning dynamic stability tester, can quantitatively evaluate the stability of dispersion systems such as colloid, foam, and suspension. The effects of 0.1%, 0.2%, and 0.3% HTRTP on the colloidal stability of 3% bentonite slurry before and after aging (245°C/16 h) were investigated.
3. Results and Discussion

3.1. Characterization

3.1.1. Composition and Molecular Weight. It can be seen from Figure 2 that the stretching vibration absorption peak at 3450 cm⁻¹ is N-H; 3060 cm⁻¹ is the absorption peak of stretching vibration of C-H bond of -CH₂ in the ring. 2960 cm⁻¹ is the characteristic absorption peak of C-H bond of -CH₃. 2920 cm⁻¹ is the absorption peak of stretching vibration of C-H bond of -CH₂ in chain. At 1730 cm⁻¹, there is a stretching vibration absorption peak with C=O. The characteristic absorption peaks of -C(O)NH- are at 1660 cm⁻¹ and 1540 cm⁻¹. At 1520 cm⁻¹, there is the vibration absorption peak of N-H bond deformation. The absorption peak of bending vibration of C-H bond of -CH₂ is at 1460 cm⁻¹. At 1300 cm⁻¹, =C=N- stretching vibration absorption peaks. The characteristic absorption peaks of -SO₃⁻ are at 1190 cm⁻¹ and 1040 cm⁻¹. The results show that

| Table 3: Experimental results of antisalting out of PMHA. |
|----------------------------------------------------------|
| Concentration of Cl⁻ (g/L)   | 0  | 90  | 108 | 126 | 144 | 162 |
| Solution transmission ratio of PMHA | 98.8 | 98.6 | 98.3 | 97.9 | 95.1 | 94.3 |
| Solution transmission ratio of DCL-a | 99.0 | 98.9 | 98.7 | 98.5 | 98.3 | 98.0 |
| Solution transmission ratio of HTRTP | 98.3 | 98.2 | 98.1 | 98.0 | 97.7 | 97.4 |

| Table 4: Experimental results of antisalting out of PMHA. |
|----------------------------------------------------------|
| Concentration of CaCl₂ (mg/L) | 0  | 500 | 1500 | 2500 | 4000 | 5000 |
| Solution transmission ratio of PMHA | 98.8 | 98.4 | 98.0 | 96.2 | 95.4 | 94.5 |
| Solution transmission ratio of DCL-a | 99.0 | 99.0 | 98.8 | 98.6 | 98.4 | 98.1 |
| Solution transmission ratio of HTRTP | 98.3 | 98.2 | 98.1 | 98.0 | 97.8 | 97.5 |

| Table 5: Comparison of fluid loss controlling of fluid loss agents in fresh water slurry. |
|----------------------------------------|
| Experimental slurry | Test conditions | AV (mPa.s) | PV (mPa.s) | YP (Pa) | API (mL) |
| 3-1 basic slurry | Room temperature | 8.5 | 3 | 5.5 | 24.0 |
| Room temperature after 220°C/16 h | 7 | 3 | 4 | 34.0 |
| Room temperature after 245°C/16 h | 8.5 | 8 | 0.5 | 41.0 |
| 1% PMHA | Room temperature | 53 | 3 | 50 | 8.4 |
| Room temperature after 220°C/16 h | 4 | 1 | 3 | 12.4 |
| Room temperature after 245°C/16 h | 11 | 10 | 1 | 20.6 |
| Basic slurry + 1%PMHA | Room temperature | 40 | 28 | 12 | 7.8 |
| Room temperature after 245°C/16 h | 17 | 14 | 3 | 11.0 |

| Table 6: Comparison of fluid loss controlling of fluid loss agents in weighted fresh water slurry. |
|----------------------------------------|
| Experimental slurry | Test conditions | AV (mPa.s) | PV (mPa.s) | YP (Pa) | API (mL) |
| 3-2 basic slurry | Room temperature after 220°C/16 h | 35 | 16 | 19 | 16.0 |
| Room temperature after 245°C/16 h | 38 | 26 | 12 | 21.0 |
| 1% PMHA | Room temperature after 220°C/16 h | 33.5 | 27 | 6.5 | 8.4 |
| Room temperature after 245°C/16 h | 45 | 35 | 10 | 6.6 |
| 1% DCL-a | Room temperature after 245°C/16 h | 41 | 32 | 9 | 6.0 |

| Table 7: Comparison of fluid loss controlling of fluid loss agents in saturated brine slurry. |
|----------------------------------------|
| Experimental slurry | Test conditions | AV (mPa.s) | PV (mPa.s) | YP (Pa) | API (mL) |
| Basic slurry | Room temperature after 220°C/16 h | 18 | 8 | 10 | 124 |
| Room temperature after 245°C/16 h | 17.5 | 6 | 11.5 | 143 |
| 1.5% PMHA | Room temperature after 220°C/16 h | 16.5 | 9 | 7.5 | 49 |
| Room temperature after 245°C/16 h | 40 | 19 | 21 | 24 |
| 1.5% DCL-a | Room temperature after 245°C/16 h | 45 | 24 | 21 | 18 |
the molecular chain of the product has the predicted group of molecular structure design, which verifies the correctness of the molecular structure design and the correctness of the HTRTP synthesis reaction scheme.

It can be seen from the test results in Table 2 that the molecular weight of HTRTP is about 1.04 million, which is smaller than that of high-temperature-resistant polymer DCL-a, and its influence on drilling fluid viscosity will be smaller than that of DCL-a.

3.1.2. Temperature Resistance. It can be seen from the thermal difference curve in Figure 3 that the thermal decomposition process of HTRTP can be divided into many stages, of which there are two main pyrolysis stages. The first stage occurs between 320 and 360 °C with a weight loss rate of 20%-25%, which is mainly the decomposition of carboxylic acid group 9 and sulfonic acid group on the side chain. The second stage, which occurs above 400 °C, is mainly the decomposition of the backbone chain of polymerization. The total weight loss rate of these two stages is 52.41%. It can be seen that the high-temperature-resistant quaternary copolymer HTRTP has strong thermal stability.

As shown in Figure 4, with the increase of aging temperature, the intrinsic viscosity of HTRTP decreased more slowly than that of DCL-a, indicating that HTRTP had better temperature resistance than DCL-a in aqueous solution.

3.1.3. Resistance to Salting Out. It can be seen from Table 3 that when the Cl− concentration of PMHA increases from 126 g/L to 144 g/L, the transmittance of PMHA solution changes suddenly, indicating that the solution begins to become cloudy; that is, the antisalting out ability of PMHA (calculated by Cl−) is between 126 g/L and 144 g/L. At the Cl− concentration of 162 g/L, the transmittance of DCL-a solution and HTRTP solution still has no obvious change, indicating that the solutions have not become turbid; that is, the salting-out resistance of DCL-a and HTRTP (calculated by Cl−) is greater than 162 g/L.

It can be seen from Table 4 that when the concentration of CaCl2 increases from 1500 mg/L to 2500 mg/L, the transmittance of PMHA solution changes suddenly, indicating that the solution begins to become cloudy; that is, the calcium resistance of PMHA (calculated by CaCl2) is between 1500 mg/L and 2500 mg/L; with the increase of CaCl2 concentration, the transmittance of DCL-a solution and HTRTP solution has no significant change. When CaCl2 concentration reached 5000 mg/L, the transmittance did not change

![Figure 5: Effect of temperature on adsorption quantity of DCL-a and HTRTP.](image)

![Figure 6: Effect of pH value on adsorption quantity of DCL-a and HTRTP.](image)
obviously, which indicated that the solutions did not become turbid; that is, the calcium resistance of DCL-a and HTRTP (calculated by CaCl₂) was greater than 5000 mg/L.

3.2. Evaluation of Fluid Loss Reduction Effect

3.2.1. Evaluation of Fluid Loss Reduction in Fresh Water Base Slurry. As shown in Table 5, the API filtration loss of HTRTP was the lowest before and after aging, and the viscosity changed the least before and after aging. The filtration loss of fresh water base slurry treated with 1% HTRTP decreased from 41 mL to 11 mL after aging at 245°C/16 h at high temperature. 1% PMHA was processed in fresh water (calculated by CaCl₂) was greater than 5000 mg/L. That is, the calcium resistance of DCL-a and HTRTP obviously, which indicated that the solutions did not become turbid; that is, the calcium resistance of DCL-a and HTRTP (calculated by CaCl₂) was greater than 5000 mg/L.

3.2.2. Effect of Fluid Loss Reduction in Fresh Water Weighted Base Slurry. As can be seen from Table 6, API filtration loss of experimental slurry after aging is low, among which the experimental slurry treated by 1% HTRTP has the lowest filtration loss, which is 6.0 mL. It was shown that in fresh water increased base slurry, HTRTP fluid loss effect is better than PMHA and DCL-a.

3.2.3. Effect of Fluid Loss Reduction in Saturated Salt-Water-Based Slurry. It can be seen from Table 7 that, compared with the base slurry, the API filtration loss of the aging experimental slurry treated with filtration reduction agent was significantly reduced, among which the experimental slurry treated with 1.5% HTRTP had the lowest filtration loss, which was 18.0 mL. It is shown that the filtration reduction effect of HTRTP is better than that of PMHA and DCL-a in saturated salt-water-based slurry.

3.3. Study on Mechanism of Action

3.3.1. Adsorption Properties on the Surface of Clay Particles

(1) Influence of Temperature. As can be seen from Figure 5, at the same temperature, with the increase of HTRTP and DCL-a concentration, their adsorption capacity on clay particles increased. With the increase of temperature, the adsorption amount decreased. It can also be seen from the figure that both HTRTP and DCL-a have different adsorption isotherms at different temperatures. The reason is that adsorption is an exothermic reaction, and increasing temperature is conducive to equilibrium in the direction of desorption. In addition, when the temperature rises, the thermal movement of clay particles intensifies, which is not conducive to adsorption.

(2) Influence of pH. As can be seen from Figure 6, when pH is 8, the adsorption capacity of the two polymers on the surface of clay particles reaches the maximum. When pH is higher or lower than this value, the adsorption capacity decreases. Within the test range, the higher the pH value, the lower the adsorption capacity. The reason is that when the pH is 8, the clay particles are fully dispersed and the negative charge on the surface of the particles is moderate. When pH becomes lower, it is not conducive to the full dispersion of clay particles, which can be used for adsorption of small surface area; when the pH value is higher than this value, more OH⁻ is adsorbed on the surface of clay particles, which further enhances the electronegativity of the surface. The electrostatic action is not conducive to the adsorption of polymer molecules. As can be seen from the figure, not only is the adsorption capacity of HTRTP greater than that of DCL-a, but with the increase of pH, the adsorption capacity of HTRTP on the surface of clay particles decreases more slowly than that of DCL-a. This is because certain cationic groups are introduced into HTRTP molecules, which can weaken the influence of the increase of electronegativity on the surface of clay particles on adsorption.
near clay particles, compressing their diffusion electric double layer; the ζ potential decreases; the repulsive force weakens; the agglomeration of clay particles increases; and the surface area decreases. In addition, the addition of sodium chloride weakens; the agglomeration of clay particles increases; and the surface area decreases. It can be seen from Figure 7 that with the increase of HTRTP dosage, the adsorption amount of HTRTP decreases. This is because, after sodium chloride addition, the adsorption amount of HTRTP decreases. It can be seen from Table 8 that the compressibility coefficient of experimental slurry before aging is lower than that after aging, indicating that high temperature will cause the compressibility reduction of the drilling fluid filter cake. The compression coefficient of experimental slurry with filtrate loss additive before and after aging is lower than that of base slurry, which indicates that improving the compressibility of filter cake is also one of the action mechanisms of filtrate loss additive. The filter cake added with HTRTP has the lowest compressibility coefficient, which is consistent with the best filtration loss reduction effect.

3.3.2. Influence on the Quality of Filter Cake. It can be seen from Figure 8 that there is a good linear relationship between filter cake permeability and API filtration amount. With the increase of HTRTP addition, API filtration amount decreases before and after experimental slurry aging, and the permeability of filter cake also decreases. It can be seen from Table 8 that the compressibility coefficient of the experimental slurry before aging is lower than that after aging, indicating that high temperature will cause the compressibility reduction of the drilling fluid filter cake. The compression coefficient of experimental slurry with filtrate loss additive before and after aging is lower than that of base slurry, which indicates that improving the compressibility of filter cake is also one of the action mechanisms of filtrate loss additive. The filter cake added with HTRTP has the lowest compressibility coefficient, which is consistent with the best filtration loss reduction effect.

3.3.3. Effect on Particle Size Distribution of Fresh Water Base Slurry. As can be seen from Figure 9, after aging, the proportion of particles with smaller particle size in the experimental slurry decreases, while the content of particles with larger particle size increases; both particle size and distribution become larger, which is one of the reasons for the increase of filtration loss after aging. The mean particle size of different suspensions was 1.7 μm, 21.5 μm, 7.5 μm, and 17.6 μm, respectively (tagged in Figure 9). After adding HTRTP, both before and after aging, the particle size distribution of experimental slurry is wider and the grading is more reasonable, so the filtration loss of experimental slurry is reduced.

3.3.4. Effect on Stability of Fresh Water Base Slurry Colloids. As can be seen from Figure 10, the colloidal stability of the experimental slurry was significantly better than that of the base slurry after the addition of HTRTP, and the colloidal stability of the experimental slurry increased with the increase of the amount of HTRTP. After aging at 245°C/16 h, high temperature promoted the hydration and dispersion of bentonite. In the experiment, it was measured that the thickness of the upper “clear liquid” of the aging base slurry was 0.30 mm after standing for 24 h, 0.1 mm when adding 0.1% HTRTP, 0.06 mm when adding 0.2% HTRTP, and 0.04 mm when adding 0.3% HTRTP. Combined with the previous particle size test results, it can be seen that for the base slurry with low bentonite content, the contribution of adhesive protection of HTRTP to filtration loss reduction is greater than that after aging. After aging, the mechanism
of HTRTP reducing filter cake permeability by improving particle size grading in the system and enhancing the compressibility of filter cake is more important.

The action mechanism of high-temperature-resistant quaternary copolymer HTRTP is as follows:

(1) Temperature resistance mechanism: the molecular structure difference is the main factor that determines the temperature resistance of the fluid loss agent. High-temperature-resistance quaternary copolymer HTRTP molecule main chain is C-C bond and has the internal structure of high temperature resistance; C-S and C-N bonds are used between the main chain and the side chain, and the thermal stability of the main chain and the side chain is strong.

(2) Mechanism of salt resistance and calcium resistance: the hydrophilic group is the sulfonic acid group (-CH2SO3-), which has strong hydration ability and can thicken the hydration film on the surface of clay particles with strong salt resistance. The sulfonic acid group does not react with Ca2+ to form precipitation, which has a strong anticalcium ability. In addition, the side groups introduced into the HTRTP chain (e.g., -CONHC(CH3)2CH2SO3-) are large and rigid, and the chain is not easy to curl in electrolyte solutions (salt solutions).
(3) Adsorption mechanism: in addition to the phthalamine group (-CON(CH₂CH₃)₂), the cationic adsorption group (-CH₃N(CH₃)₂CH₂) is also introduced into the HTRTP molecular chain. -CON(CH₂CH₃)₂ is adsorbed by hydrogen bond, which has strong adsorption stability and is not easy to be desorbed under temperature change. -CH₃N(CH₃)₂CH₂ is chemisorbed through static ion bonds, and the adsorption is stronger.

(4) Mechanism of filtration loss reduction: firstly, HTRTP is adsorbed on the surface of clay particles to form an adsorption layer, which prevents flocculation of clay particles from becoming larger. At the same time, it stabilizes the fine particles separated by stirring so as not to become larger. So it can improve clay particle grading in drilling fluid. It has excellent protective adhesive effect, forming thin and compact filter cake, reducing filtration loss. Secondly, HTRTP increases the degree of hydration by adsorbing on clay particles, so that the hydration film on the surface of clay particles is thickened to enhance the compressibility of the filter cake and reduce the permeability of the filter cake, thus reducing the filtration loss. Thirdly, the molecular size of HTRTP is just within the range of colloidal particles, and the molecular chain wedges into the pores of the filter cake or curls into a ball to block the micropores of the filter cake, making the filter cake thin and compact, thus reducing the filtration loss. Finally, HTRTP is a high molecular weight polymer that is added to the drilling fluid to increase the viscosity of the filtrate and thus reduce filtration loss.

4. Conclusions

The high-temperature-resistant quaternary copolymer HTRTP was synthesized by molecular structure optimization and monomer optimization. The viscosity average molecular weight of HTRTP was about 1 million, which had less effect on the viscosity of drilling fluid than DCL-α. HTRTP molecular structure has good thermal stability, temperature resistance in drilling fluid up to 245°C, salt resistance and calcium resistance, and good compatibility with fresh water and salt water experimental base slurry. The performance of fluid loss reduction, temperature resistance and salt and calcium resistance in drilling fluid is better than that of foreign temperature resistance polymer DCL-α and domestic similar products. The adsorption capacity of HTRTP on clay particles is higher than that of foreign anti-temperature polymer DCL-α. In the case of chemical environment and temperature change, the adsorption amount changed little and was still higher than that of DCL-α, and the adsorption was firm. Before and after aging (245°C/16 h), HTRTP can significantly reduce the permeability and improve the compressibility of filter cake. Improving the particle size grading in drilling fluid system and enhancing the colloidal stability of the system are the important mechanisms of high-temperature-resistant quaternary copolymers. The quad-element polymer reducer was suitable for fresh water-based drilling fluid and low-mineralized saline drilling fluid.

Nomenclature

AMPS: 2-Acry-lamido-2-methyl-1-propane sulfonic acid
API: American Petroleum Institute
AV: Apparent viscosity
DEAM: N, N-Diethyl acrylamide
DMDAAC: Dimethyl diallyl ammonium chloride
FT-IR: Fourier transform infrared spectroscopy
NVP: N-Vinyl pyrrolidone
PV: Plastic viscosity
TGA: Thermogravimetric analysis
YP: Dynamic shear forces.

Data Availability

All data and research results are listed in the manuscript.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

Acknowledgments

This work was financially supported by the Open Foundation of State Key Laboratory of Oil and Gas Geology and Exploitation, Chengdu University of Technology (PLC20180103), and Opening Project of Oil and Gas Field Applied Chemistry Key Laboratory of Sichuan Province (YQKF202010).

References

[1] A. R. Ismail, W. R. Wan Sulaiman, M. Z. Jaafar, I. Ismail, and E. Sabu Hera, “Nanoparticles performance as fluid loss additives in water based drilling fluids,” Materials Science Forum, vol. 864, no. 1, pp. 189–193, 2016.
[2] A. Nasiri, M. J. AMERI Shahraibi, M. A. Sharif NIK, H. Heidari, and M. Valizadeh, “Influence of monoethanolamine on thermal stability of starch in water based drilling fluid system,” Petroleum Exploration and Development, vol. 45, no. 1, pp. 167–171, 2018.
[3] H. Zhong, G. Shen, Z. Qiu et al., “Minimizing the HTHP filtration loss of oil-based drilling fluid with swellable polymer microspheres,” Journal of Petroleum Science and Engineering, vol. 172, pp. 411–424, 2019.
[4] X. Ma, M. Yang, and M. Zhang, “Synthesis and properties of a betaine type copolymer filtrate reducer,” Chemical Engineering and Processing-Process Intensification, vol. 153, article 107953, 2020.
[5] F. Dugonjic-Bilic and J. Plank, “Polyelectrolyte complexes from polyethylene imine/acetone formaldehyde sulfole polycondensates: a novel reagent for effective fluid loss control of oil well cement slurries,” Journal of Applied Polymer Science, vol. 121, no. 3, pp. 1262–1275, 2011.
[6] X. Yang and Z. Wang, “Molecular design of polymer treatment agent used for drilling fluid,” *Fine and Specialty Chemicals*, vol. 18, pp. 14–18, 2010.

[7] Z. Y. Chang, D. L. Breeden, and M. J. Mcdonald, “The use of zinc dialkyldithiophosphate as a lubricant enhancer for drilling fluids particularly silicate-based drilling fluids,” in *SPE International Symposium on Oil Field Chemistry*, The Woodlands, Texas, USA, 2011.

[8] X. Bai, Y. Yang, D. Xiao, X. Pu, and X. Wang, “Synthesis, characterization, and performance evaluation of the AMI/AMPS/DMDAAC/SSS quadripolymer as a fluid loss additive for water-based drilling fluid,” *Journal of Applied Polymer Science*, vol. 132, no. 14, article 41762, 2015.

[9] J. Su, Q. Chu, and M. Ren, “Properties of high temperature resistance and salt tolerance drilling fluids incorporating acrylamide/2-acrylamido-2-methyl-1-propane sulfonic acid/N-vinylpyrrolidone/dimethyl dialyl ammonium chloride quadripolymer as fluid loss additives,” *Journal of Polymer Engineering*, vol. 34, no. 2, pp. 153–159, 2014.

[10] P. W. Livanec, G. P. Perez, and J. P. Deville, “Methods of preventing emulsification of crude oil in well bore treatment fluids,” 2012, U.S. Patent No. 9,051,508.

[11] J. B. Wen, H. J. Luo, and Z. Q. Long, “Emulsification behaviors of crude oil-water system and its quantitative relationship with exergy loss rate,” *Journal of Petroleum Science & Engineering*, vol. 176, pp. 502–508, 2019.

[12] X. I. N. Haipeng, Z. E. N. Jianguo, Z. O. U. Jianlong, Z. H. A. O. Baohui, and S. U. N. Fuquan, “Laboratory study on comb-like polycarboxylic acid friction reducer,” *Drilling Fluid and Completion Fluid*, vol. 34, pp. 72–76, 2017.

[13] G. Duan, G. Huang, A. Li, Y. Zhu, and Y. Gong, “A study of supramolecular polarization of comb-like polycarboxylate admixtures synthesized with polyoxyethylene macromolecules,” *Journal of Molecular Liquids*, vol. 174, pp. 129–134, 2012.

[14] Q. Jiang, G. Jiang, C. Wang et al., “A new high-temperature shear-tolerant supramolecular viscoelastic fracturing fluid,” in *IADC/SPE Asia Pacific Drilling Technology Conference*, Singapore, 2016.

[15] W. Li, J. Liu, X. Zhao et al., “Development and screening of additives for biodiesel based drilling fluids: principles, strategies and experience (Conference Paper),” in *Proceedings-SPE International Symposium on Oilfield Chemistry*, Texas, USA, 2019.

[16] J. O. Oseh, M. N. A. Mohd Norddin, I. Ismail, A. O. Gbadamosi, A. Agi, and H. N. Mohammed, “A novel approach to enhance rheological and filtration properties of water-based mud using polypropylene-silica nanocomposite,” *Journal of Petroleum Science and Engineering*, vol. 181, article 106264, 2019.

[17] X. F. Chang, J. S. Sun, Z. Xu et al., “Synthesis of a novel environment-friendly filtration reducer and its application in water-based drilling fluids,” *Colloids and Surfaces A: Physicochemical and Engineering Aspects*, vol. 568, pp. 284–293, 2019.

[18] X. Liu, B. Xie, Y. Gao et al., “Development of low toxicity and high temperature polymer drilling fluid for environmentally sensitive offshore drilling,” in *IADC/SPE Asia Pacific Drilling Technology Conference and Exhibition*, Bangkok, Thailand, 2018.

[19] L. Long, X. Xianguang, Z. Jinzhi et al., “Application of innovative high-temperature high-density oil-based drilling fluid technology in the efficient exploration and development of ultra-deep natural gas resources in West China,” in *International Petroleum Technology Conference*, Bangkok, Thailand, 2016.

[20] H. Shen, K. Lv, X. Huang et al., “Hydrophobic associated polymer-based laponite nanolayered silicate composite as filtrate reducer for water-based drilling fluid at high temperature,” *Journal of Applied Polymer Science*, vol. 137, no. 5, 2020.

[21] D. V. Kosynkin, G. Ceriotti, K. C. Wilson et al., “Graphene oxide as a high-performance fluid-loss-control additive in water-based drilling fluids,” *ACS Applied Materials and Interfaces*, vol. 4, no. 1, pp. 222–227, 2012.

[22] H. Shen, K. Lv, X. Huang et al., “Hydrophobic-associated polymer-based laponite nanolayered silicate composite as filtrate reducer for water-based drilling fluid at high temperature,” *Journal of Applied Polymer Science*, vol. 137, no. 18, article 48608, 2020.

[23] H. A. Craddock, “Oilfield chemistry and its environmental impact,” *John Wiley & Sons*, pp. 21–110, 2018.

[24] G. Pan, J. Chen, C. Zhang et al., “Combined technology of weak gel flooding assisting thermal huff and puff enhances oil recovery for offshore heavy oil field,” in *SPE Annual Technical Conference and Exhibition*, Dubai, UAE, 2016.

[25] M. X. Li, S. C. Xie, H. Zhang, X. G. Wang, and Q. Z. Jin, “Effect of mechanical stirring rate on the texture, rheological property and microstructure of gel-like oil,” *Journal of Food Science and Biotechnology*, vol. 32, pp. 393–398, 2013.

[26] R. Jain, T. K. Mahito, and V. Mahito, “Rheological investigations of water based drilling fluid system developed using synthesized nanocomposite,” *Korea Australia Rheology Journal*, vol. 28, no. 1, pp. 55–65, 2016.

[27] K. K. Chandan and G. P. Karmakar, “Drilling fluid waste treatment using polysaccharide-grafted copolymers,” *The APPEA Journal*, vol. 59, no. 1, pp. 34–46, 2019.

[28] P. Dejtaradon, H. Hamidi, M. H. Chuks, D. Wilkinson, and R. Rafati, “Impact of ZnO and CuO nanoparticles on the rheological and filtration properties of water-based drilling fluid,” *Colloids and Surfaces A: Physicochemical and Engineering Aspects*, vol. 570, pp. 354–367, 2019.

[29] B. Xie, X. Zhang, Y. Li, W. Liu, and M. Luo, “Application a novel thermo-sensitive copolymer as a potential rheological modifier for deepwater water-based drilling fluids,” *Colloids and Surfaces A: Physicochemical and Engineering Aspects*, vol. 581, article 123848, 2019.