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Laboratory evaluation and numerical simulation of the alkali-surfactant-polymer synergistic mechanism in chemical flooding

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Alkali-surfactant-polymer (ASP) flooding, which can reduce interfacial tension (IFT) and the mobility ratio between oil and water phases, has been proven to be effective for enhancing oil recovery in laboratory experiments and field pilots. However, the study of interactions within alkali-surfactant-polymers for chemical flooding is neither comprehensive nor complete until now. Laboratory experiments were conducted and a corresponding numerical simulation model was established to characterize multiple component interactions during the ASP flooding process. Synergistic effects of multiple component interactions on viscosity variation, IFT reduction, and multicomponent adsorption were studied separately. ASP solution viscosity shows non-linear variation behavior with an increasing polymer concentration. Alkali decreases the molecular hydraulic radius of a polymer, and then limits its contribution to viscosity. Oil-water interfacial tension decreases with the join in of polymer which can act as an alternative effect to replace surfactant adsorbed on a mineral surface. Petroleum acid will react with alkali and produce petroleum soap to perform a synergetic action with the surfactant on IFT reduction. Adsorption fraction and diffusion rate of a surfactant will diminish due to rheology improvements caused by a polymer. Alkali can protect a surfactant from adsorption consumption by competitive adsorption. A viscosity non-linear logarithm mixing method, IFT reduction-relative permeability curve interpolation method, and a multicomponent adsorption isotherm model were developed to characterize and simulate the synergistic effects obtained by experiments. A novel ASP flooding numerical simulation model was constructed which coupled the synergistic effects simulation methods of viscosity variation, IFT reduction, and multicomponent adsorption. The numerical simulation result based on the proposed model has better agreement with experiment results compared with that of the traditional model. Validation results proved the effectiveness of the proposed model which can be used to enhance a synergistic mechanism study and field application of ASP flooding.

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Introduction

Laboratory research conducted by Lin *et al.* in 1987 indicated that oil recovery can be improved by using a combination of alkali, surfactant, and polymer. After that, ASP flooding technology which can improve mobility ratio and sweep efficiency of remaining oil has been reported on a world scale with a high proportion of success. The effect of alkali on ASP flooding was studied by Nasr-EI-Din *et al.* and they indicated that residual oil was recovered by two mechanisms: low interfacial tension and wettability reversal. An alkali agent will react with acid to generate an *in situ* surfactant and, hence, attain ultralow interfacial tension. The role of a polymer is to increase viscosity, hence reducing the mobility ratio and reaching greater volumetric

sweep efficiency.^{12,13} The surfactant component of a system is responsible for reducing IFT between oil and water phases to a level that promotes mobilization of trapped oil drops.^{14,15} Viscoelasticity theory was developed by Hou *et al.* to describe the mechanism of ASP flooding. They considered viscoelasticity rather than IFT as an evaluation index to study and optimize ASP flooding.¹⁶ Different from sandstone reservoirs, carbonate reservoirs usually are oil-wet, an unfavorable condition for ASP flooding. However, Bortolotti *et al.* and Levitt *et al.* evaluated the effects of ASP flooding on carbonate rocks in laboratory experiments. The results showed that ASP flooding was effective for enhancing ultimate oil recovery in oil-wet carbonate reservoirs.^{17,18}

Numerical simulation software was developed based on dozens of years of phase behavior, core flood, and mechanistic research on ASP flooding. The University of Texas Chemical Compositional Simulator (UTCHEM) was usually used to model ASP flooding process. 19,20 Yuan *et al.* established a 3D multifunctional compositional numerical simulator to study and predict the outcomes of ASP flooding. 21 Using commercial

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simulation software CMG to design and simulate field pilot, Moreno *et al.* and Van *et al.* indicated that it is possible to increase oil recovery dramatically by ASP flooding.^{22,23} Farajzadeh *et al.* coupled a multi-purpose dynamic reservoir simulator (MoReS) and geochemistry software program (PHREEQC) to provide a versatile tool for ASP flooding numerical simulation.²⁴

Pilot tests and field applications of ASP flooding have been carried out around the world. ASP flooding technology has been proven successful in three completed projects in North America. Vargo *et al.* reported that an ASP flooding project in the Cambridge Minnelusa field was a technical and economic success with an ultimate incremental oil recovery of 28.1%. In China, the first ASP flooding pilot was conducted in the Shengli oilfield in 1992 and the incremental oil recovery was reported to be 26% OOIP. Wang *et al.* indicated that five ASP flooding pilots were conducted in the Daqing oilfield and the oil recoveries of four of them were 20% OOIP above those obtained with water flooding. B

It should be mentioned that an alkali-surfactant-polymer will affect each other more or less rather than noninterference during a multicomponent flooding process, and those interactions are usually synergistic effects. However, a synergistic effect which can restrain or enhance ASP flooding performance hasn't attracted enough attention until now. The study of synergistic effects for ASP flooding is neither comprehensive nor complete according to our knowledge. Given the reasons mentioned

Table 1 Compositions of injection water and formation brine

	Concentration (mg L^{-1})			
Parameter	Injection water	Formation brine		
Calcium	190	683		
Magnesium	96	113		
Sodium	1192	3133		
Bicarbonate	30	161		
Chloride	2096	5473		
Sulphate	408	234		
Total dissolved solids	4011	9797		

above, our goal was to investigate an alkali-surfactant-polymer synergistic effect based on laboratory experiments, and then establish a corresponding numerical simulation model with the synergistic mechanism combined to describe the multiple component's interactions during the ASP flooding process.

2. Experimental design

2.1. Material descriptions

Artificial homogeneous cores were used in ASP flooding tests. Relative parameters of artificial cores were designed based on formation properties of the Lvda offshore oilfield in Bohai Bay in China. A total of 50 cores with dimensions of 2.50 cm diameter and 10.00 cm length were prepared. The approximate porosity and permeability of each core was 0.28 and 1750 mD, respectively. Crude oil, injection water, and formation brine used in this study were collected from the Lvda oilfield. Crude oil density and viscosity were 866 kg m⁻³ and 57.13 mPa at atmospheric pressure and a temperature of 25 °C. Compositions of injection water and formation brine are shown in Table 1. The alkali used in this study was pure sodium hydroxide (NaOH).29 Sodium dodecyl sulfate with an active content of 85.0 wt% was chosen as the surfactant in the ASP agent system. A high molecular weight polyacrylamide (HPAM, 24.0% hydrolyzed and molecular mass of 11.6 million daltons) was used as the polymer in this study.

2.2. Experimental plan

Two groups of core flooding experiments were designed as shown in Tables 2 and 3. The aim of the experiments in Table 2 was to study the characteristic of ASP flooding and then compare the performances of unitary, binary, and ternary chemical flooding. Effects of alkali and polymer on surfactant adsorption were measured by experiments and listed in Table 3.

2.3. Experimental apparatus and procedure

2.3.1. Viscosity test. The effects of alkali, surfactant, and polymer on ASP solution viscosity were measured using a HAAK RS-150H rheometer with temperature maintained at a constant $^{\circ}$ C. Concentration variations for alkali, surfactant, and

Table 2 Experimental design for ASP flooding characteristic and performance study

Initial water Core number slug volume (PV)	Turbital	ASP slug volume (PV)	Porosity (%)	Permeability (mD)	Initial oil saturation (%)	Concentration (wt%)		
	slug volume (PV)					Surfactant	Alkali	Polymer
1	0.08	0.43	28.34	1817.10	73.13	0.25	1.0	0.12
2	0.26	0.43	32.03	1573.20	73.00	0.25	1.0	0.12
3	0.92	0.43	28.72	1557.90	74.44	0.25	1.0	0.12
4	0.32	0.43	29.82	1751.26	72.86	_	1.0	_
5	0.32	0.43	28.35	1680.10	73.31	0.25	_	_
6	0.32	0.43	28.68	1682.27	71.59	_	_	0.12
7	0.32	0.43	29.45	1759.64	72.56	0.25	1.0	_
8	0.32	0.43	27.89	1823.59	73.15	_	1.0	0.12
9	0.32	0.43	28.64	1789.33	72.89	0.25	_	0.12
10	0.32	0.43	28.76	1698.25	73.21	0.25	1.0	0.12

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Table 3 Experimental design for chemical adsorption study

	Concentration	(wt%)			Concentration (wt%)		
Core number	Surfactant	Alkali	Polymer	Core number	Surfactant	Alkali	Polymei
11	0.1	_	_	31	0.1	_	_
12	0.1	_	0.08	32	0.1	0.75	_
13	0.1	_	0.16	33	0.1	1.5	_
14	0.1	_	0.24	34	0.1	2	_
15	0.2	_	_	35	0.2	_	_
16	0.2	_	0.08	36	0.2	0.75	_
17	0.2	_	0.16	37	0.2	1.5	_
18	0.2	_	0.24	38	0.2	2	_
19	0.3	_	_	39	0.3	_	_
20	0.3	_	0.08	40	0.3	0.75	_
21	0.3	_	0.16	41	0.3	1.5	_
22	0.3	_	0.24	42	0.3	2	_
23	0.4	_	_	43	0.4	_	_
24	0.4	_	0.08	44	0.4	0.75	_
25	0.4	_	0.16	45	0.4	1.5	_
26	0.4	_	0.24	46	0.4	2	_
27	0.5	_	_	47	0.5	_	_
28	0.5	_	0.08	48	0.5	0.75	_
29	0.5	_	0.16	49	0.5	1.5	_
30	0.5	_	0.24	50	0.5	2	_

polymer ranged from 0-2.0 wt%, 0-0.5 wt%, and 0-0.24 wt%, respectively.

- 2.3.2. Interfacial tension test. Interfacial tension between crude oil and chemical solution was tested by a Spinning Drop Tensiometer model 500. Solutions were aged for three hours after a crude oil and chemical solution was mixed, and then IFT was measured at 25 °C. The influences of alkali and surfactant on IFT were investigated using a range of chemical agent concentrations (alkali 0-2.0 wt% and surfactant 0-0.5 wt%).
- 2.3.3. Core flood test. A schematic of ASP core flood setup consists of six components: micro pump, core holder, tanks, pressure record system, measurement equipment, and collection flask. Pressure levels at the inlet and outlet of the core holder were measured by a pressure transducer. A computer equipped with a data acquisition system was used to automatically record pressure data. The following steps were applied:
- (1) Inserted one core into core holder and a confining pressure of 5.0 MPa was applied.
- (2) 5.0 pore volumes (PVs) of formation brine were injected into the core with an injection rate of 0.3 mL min⁻¹. Water flooding permeability was measured.
- (3) 5.0 PVs of crude oil were injected to saturate the core with an injection rate of 0.3 mL min⁻¹.
- (4) Water was injected into the core at a rate of 0.3 mL min⁻¹ to a specific volume (for coreflood tests in Table 2) or until injection volume reached 1.0 PV (for coreflood tests in Table 3). The injection pressure was recorded.
- (5) Then, 0.43 PV of ASP solution was injected into the core. Pressure data was recorded by a pressure transducer.
- (6) Injection water was injected into the core at a rate of 0.3 mL min⁻¹ until water cut reached 90.0%. The pressure data was recorded.

- (7) Repeated earlier steps for all core tests as shown in Table 2. Oil recovery and water cut were measured during the experimental steps of (4), (5), and (6).
- (8) Steps (1), (2), (4), (5), and (6) were conducted for core tests listed in Table 3. The effluent chemical agent concentrations were measured separately.

3. Performance evaluation of chemical flooding

3.1. Performance comparison of unitary, binary, and ternary chemical flooding

Enhanced oil recovery (EOR) performance of unitary, binary, and ternary chemical flooding was estimated based on experiments

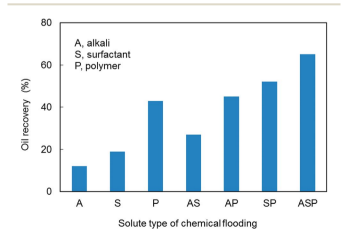


Fig. 1 Performance comparison of unitary, binary and ternary chemical flooding

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and the results are listed in Table 2 (core numbers 4 to 10). Experimental outcome documents that ternary chemical flooding (ASP) achieved the highest oil recovery compared with unitary and binary cases. This is due to the contribution of a synergistic effect within the alkali–surfactant–polymer. Nevertheless, alkali–polymer (AP) flooding only has a slight oil recovery increment compared with that of polymer flooding. This implies that not all chemical agents will have synergistic effects at all times and the synergistic effects between chemical agents are conditional. Considering the complexity of interactions within an alkali–surfactant–polymer, the synergistic mechanism of ASP flooding still needs further investigation (Fig. 1).

3.2. Characteristic study of ASP flooding

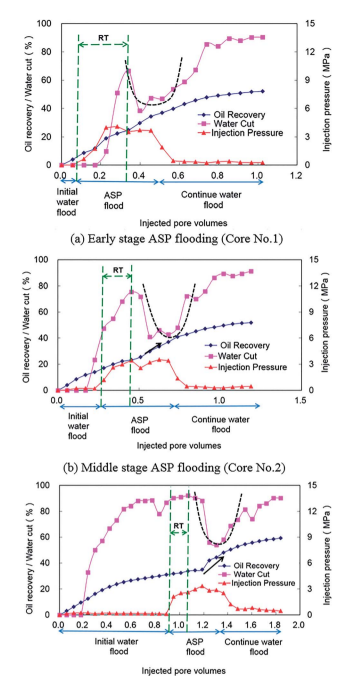
Different initial water flood slug sizes represent different development stages after which an ASP solution is injected into a core (core numbers 1, 2, and 3 as shown in Table 2). For core number 1, ASP flooding was conducted in an early stage of the development when water cut was low. As shown in Fig. 2(a), water cut increased rapidly to 66% and then dropped significantly after it achieved a critical point, which indicated that ASP flooding was effective. After that, water cut was reversed to increase slowly during the later period of ASP flooding and the continued water flooding period. The U-shape (as black dashed line shown in Fig. 2(a)) of a water cut curve during this period is a clear response characteristic of ASP flooding. Another important observation is that injection pressure is shown as a trapezoid and a high injection pressure stair during an ASP flooding period. The experimental results of corefloods 2 and 3 with ASP flooding conducted at middle and late stages of development are shown in Fig. 2(b) and (c), respectively. Water cut reverse and U shape of a water cut curve also can be observed from these two figures. The performance of injection pressure is similar to that observed in Fig. 2(a). It is also important to note that oil recovery increase accelerates gradually as a result of ASP solution injection (marked by black arrows in Fig. 2(b) and (c)). Additionally, the response time (RT), which is the interval between ASP flooding start time and water cut critical time, shortens with the delay of ASP flooding (compare Fig. 2(a)-(c)). This indicates that quick response and effectiveness can be obtained for late stage ASP flooding.

4. Synergistic mechanism investigation of ASP flooding

Generally, inhibitory and synergist actions within chemical agents will occur at the same time for multi-component chemical flooding. These interactions will affect the properties of a multi-component solution and then master the performance of chemical flooding. Hence, the interactions of an alkali–surfactant–polymer on viscosity, interfacial tension, and adsorption were studied with laboratory experiments.

4.1. Viscosity sensitivity analysis of ASP solution

The effects of alkali, surfactant, and polymer on viscosity of ASP solution were studied based on rheology experiments and the



(c) Late stage ASP flooding (Core No.3)

Fig. 2 Production performance under different stages of ASP flooding.

results are shown in Fig. 3. Alkali, surfactant, and polymer affect the viscosity of an ASP solution at same time, but the sensitivity of each chemical agent is different according to its ternary phase diagram as shown in Fig. 3. As viscosity contour lines show, viscosity increases step by step with increasing polymer concentration and crosslinking probability of polyacrylamide molecules grows exponentially with increasing polymer concentration. This leads to a non-linear variation of ASP solution viscosity and performs as the viscosity increase gradually accelerates. It is also important to note that the effect of

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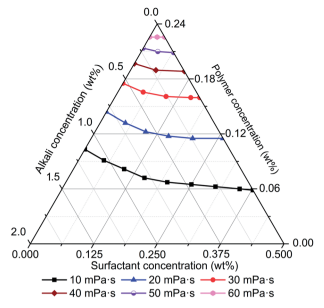


Fig. 3 Ternary phase diagram of ASP solution viscosity.

polymer on viscosity is more significant compared with that of alkali and surfactant, which indicates that polymer is the primary control factor of ASP solution viscosity.

As shown in Fig. 3, the decline of ASP solution viscosity with increase of alkali concentration is significant, and implies that alkali limits the contribution of polymer to viscosity. Polyacrylamide molecular stretches exist without alkali as a repulsive force. Alkali provides cations which can reduce the repulsive force within polyacrylamide molecules based on a charge shielding mechanism. Then, polyacrylamide molecules shrink rather than stretch, which leads to a decline of molecular hydraulic radius. Additionally, crosslinking probability of polyacrylamide molecules decreases with the reduction of molecular hydraulic radius. Finally, the viscosity of an ASP solution diminishes with an increase of alkali concentration. It is also important to note that a minor reduction of viscosity can be found with an increase of surfactant concentration. This indicates that a surfactant has slight influence on viscosity. Consequently, ASP solution viscosity is determined primarily by polymer and secondary by alkali, and the effect of a surfactant is relatively small.

4.2. Interfacial tension sensitivity analysis of ASP solution

Interfacial tension (IFT), at the petroleum-aqueous phase interface, is one of the key parameters to determine ASP flooding effectiveness. Generally, the lower the IFT, the easier an oil-water mixture occurs, hence, ultra-low IFT usually is expected for ASP flooding. The effect of polymer on IFT is insignificant when crude oil mixes with ASP solution in a vessel. But, if an ASP solution is injected into a core which is saturated with crude oil, then a polymer can reduce surfactant adsorption on the mineral surface by an alternative mechanism during the flooding process. Then, oil-water IFT will decline indirectly due to an increasing active-surfactant concentration.

Fig. 4 shows oil-water IFT as a function of surfactant and alkali concentrations. A sharp decrease can be seen after alkali joins in and the trend flattens out quickly with an increase of alkali concentration. When oil and aqueous phases come in contact, *in situ* petroleum acid HP in the oil phase and NaOH in the aqueous phase will migrate to the interface, react, and produce petroleum soap (NaP) which is a kind of *in situ* surfactant. With an increase of NaOH concentration, more petroleum soaps are produced and the density of P⁻ at the interface increases, which leads to the dropping of IFT. But limited by the content of petroleum acid in crude oil, more NaOH won't contribute to IFT reduction anymore after petroleum acid is completely converted. Hence, an IFT surface turns into a platform after a critical alkali concentration is achieved.

It is obvious that the effect of surfactant on IFT is more significant than that of alkali. The addition of surfactant into a petroleum-alkali-polymer system can influence oil-water IFT behavior and oil recovery in two ways. On one hand, the surfactant may adsorb at the oil-water interface to improve interfacial properties and then enhance the oil-water mixture. On the other hand, the surfactant may form mixed micelles with petroleum soap NaP which is produced by an alkali and petroleum acid chemical reaction, and then perform a synergetic action between surfactant and alkali. Another important observation is that IFT decreases rapidly when surfactant concentration is low, but it turns to increasing after surfactant concentration achieves 0.2 wt% (see Fig. 4). Diffusion equilibrium may be the reason for IFT turning to increasing from decreasing with the variation of surfactant concentration. Part of the surfactant may dissolve in the oleic phase rather than aqueous phase and won't contribute to IFT reduction anymore when surfactant concentration achieves critical value. Meanwhile, emulsification also can lead to the IFT increase.

It should be mentioned that all the analysis discussed above is based on laboratory experiments conducted in a vessel. Moreover, an oilfield pilot is still necessary to ensure that the synergistic mechanism of alkali–surfactant–polymer on viscosity can indeed be achieved *in situ* when the ASP solution contacts with crude oil in a porous media during the flooding process.

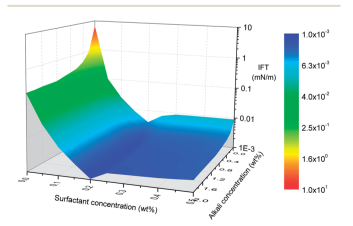
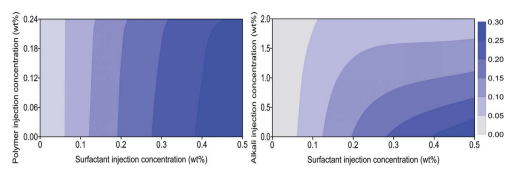


Fig. 4 Effects of alkali and surfactant on IFT.

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- (a) Effect of polymer on surfactant adsorption
- (b) Effect of alkali on surfactant adsorption

Fig. 5 Surfactant adsorption concentration contour plot during SP/AS flooding process.

4.3. Adsorption investigation of ASP flooding

Physical reactions between chemical agents with sandstone mineral also will affect the performance of ASP flooding. Adsorption, entrapment, retention, and trapping are the main reasons that lead to the loss of chemical agents, and adsorption is the leading cause for chemical consumption.³⁰⁻³² Chemical adsorption behavior and the effects on the synergistic mechanism of ASP flooding were determined through coreflood experiments.

In Fig. 5(a), the abscissa and ordinate are surfactant and polymer concentrations of a surfactant-polymer (SP) solution injected into a core. The concentration of surfactant which adsorbed on mineral surface of the core can be obtained by detecting effluent surfactant concentration after SP flooding. Surfactant adsorption concentration scatter data were measured by experiments with core numbers 11 to 30 listed in Table 3. Then a surfactant adsorption concentration contour plot was drawn based on obtained scatter data, and the result is shown in Fig. 5(a). It can be observed that surfactant adsorption concentration increases with increasing surfactant injection concentration when polymer injection concentration is constant. The interval between two adjacent contours enlarges with increasing surfactant injection concentration, which denotes that the increasing speed of surfactant adsorption concentration will gradually slow down. This implies that surfactant adsorption will tend to a dynamic plateau with mounting surfactant injection concentration. It also can be observed that surfactant adsorption concentration diminishes slightly with increasing polymer injection concentration for the same surfactant injection concentration. A polymer has a disadvantage on competitive adsorption compared with surfactant. As a result, it has a slight effect on surfactant adsorption under low concentration. Nevertheless, the viscosity of a SP solution will increase with increasing polymer concentration. Rheology improvement of a SP solution will decline surfactant diffusion rate in a core.33 Then surfactant adsorption concentration has a significant decrease under high surfactant and polymer concentrations.

Fig. 5(b) shows a surfactant adsorption concentration contour plot of alkali-surfactant (AS) flooding based on experimental results from core numbers 31 to 50 listed in Table 3. It can be observed that the effect of alkali on surfactant adsorption is more significant compared with that of polymer (see

Fig. 5(a) and (b)). First, alkali is usually a sacrificial agent to replace surfactant from adsorbing on a mineral surface to keep chemical adsorption equilibrium. Also, it can protect a surfactant from adsorption consumption and enhance its contribution to IFT reduction. This is an alternative mechanism between alkali and surfactant. Second, alkali will increase AS solution's pH and lead a negatively charged mineral surface to become more electro-negative. Then, the repulsion force between mineral surface and anionic surfactant is enhanced which reduces the proportion of surfactant to be adsorbed directly on mineral surface. Third, crude oil usually contains petroleum acid which can react with alkali to produce a water-soluble surfactant. This produced petroleum surfactant will occupy adsorption positions of injected surfactant on a mineral's surface, and then reduce its adsorption loss.34 Hence, dramatic reduction of surfactant adsorption concentration can be seen in the presence of alkali, and the reduction decelerates gradually with increasing alkali injection concentration (see Fig. 5(b)). Surfactant adsorption concentration will convert from an abrupt drop to a smooth decline when alkali injection concentration rises to a certain degree. This indicates that more alkali will have less contribution to surfactant adsorption reduction when adsorption capacity of the mineral is approached or achieved. Additionally, the alternative mechanism of a surfactant with alkali is usually used to reduce the dosage of surfactant and enhance oil recovery in oilfield applications, which ensures better economic performance of ASP flooding as a surfactant is usually more expensive than alkali.

5. Numerical simulation development of synergistic mechanism

In this part, numerical simulation methods were developed to characterize the ASP flooding synergistic mechanism which had been obtained from experiments. Simulations of viscosity mixing, IFT reduction, and multicomponent adsorption were studied separately.

5.1. Viscosity non-linear logarithm mixing calculation

In formula (1) of a traditional linear logarithm mixing rule, mass fraction w_i acts as a weight factor during multi-component viscosity mixing calculations.³⁵

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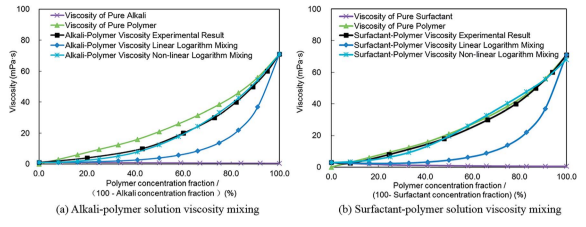


Fig. 6 Alkali-surfactant-polymer multi-component viscosity mixing calculation methods.

$$\ln \mu = \sum_{i=1}^{n} w_i \ln \mu_i \tag{1}$$

The viscosity mixing calculation of ASP solution in this study can be converted as:

$$\ln \mu = w_A \ln \mu_A + w_S \ln \mu_S + w_P \ln \mu_P \tag{2}$$

where μ is viscosity of ASP solution, mPa s; w_A , w_S and w_P are mass fractions of alkali, surfactant, and polymer components, and μ_A , μ_S and μ_P are viscosities of pure alkali, surfactant, and polymer solutions separately, mPa s.

Based on the linear logarithm mixing method shown in formula (2), the viscosity of an alkali-polymer solution can be calculated (see Fig. 6(a)). It can be observed that the calculated viscosity by the linear logarithm mixing method doesn't match as well with the experimental results. A similar phenomenon also can be observed from the comparison of the surfactantpolymer solution viscosity mixing result with experimental data (see Fig. 6(b)). This denotes that the viscosity linear logarithm mixing method, which doesn't consider interactions within alkali, surfactant, and polymer, is neither comprehensive nor complete. To solve this problem, a multivariate regression analysis was conducted according to the consequence of experiments we had already obtained (as shown in Fig. 3). The regression result, which is a new multi-component viscosity non-linear logarithm mixing rule, is put forward in formula (3). The effects of alkali and surfactant on the viscosity of polymer are considered by an exponential term which involved alkali and surfactant concentrations. The corresponding affect factors can be measured and matched based on the experimental results in part 4.1 of this paper.

Consider formula (3):

$$\ln \mu = w_{A} \ln \mu_{A} + w_{S} \ln \mu_{S} + c \times e^{(c_{A}w_{A} + c_{S}w_{S})} \times w_{P} \ln \mu_{P}$$
 (3)

where c is normalized factor; c_A and c_S are affect factors of alkali and surfactant on the viscosity of polymer.

The normalized summation of alkali and polymer (Fig. 6(a)) or surfactant and polymer (Fig. 6(b)) mass fractions as 100%, and the viscosity non-linear logarithm mixing result is shown in Fig. 6. The outcome documents that the viscosity calculation result based on non-linear logarithm mixing method matches very well with experiment results compared with that of the linear logarithm mixing method. Hence, the non-linear logarithm mixing method was employed to replace the traditional linear logarithm mixing method to conduct the multi-component viscosity mixing calculation in ASP flooding numerical simulations.

5.2. IFT reduction-relative permeability curve interpolation

During ASP flooding process, oil-water phases IFT reduction with the join in of chemical agent usually reflects on the variation of capillary number and relative permeability curve. Formula (4) is usually employed to describe the relationship of capillary number with IFT:

$$N_{\rm c} = \frac{\mu v}{\delta_{\rm S}} \tag{4}$$

where N_c is capillary number; μ is fluid viscosity, mPa s; ν is fluid velocity, m s⁻¹; and δ_S is IFT with the existing of surfactant, $mN m^{-1}$.

IFT will be affected by surfactant and alkali at the same time according to the experimental results discussed before. As shown in formula (4), capillary number is the function of surfactant only, which is inaccurate because the effect of alkali is ignored. Then, a formula which considers the effects of surfactant and alkali on IFT at the same time can be improved as:

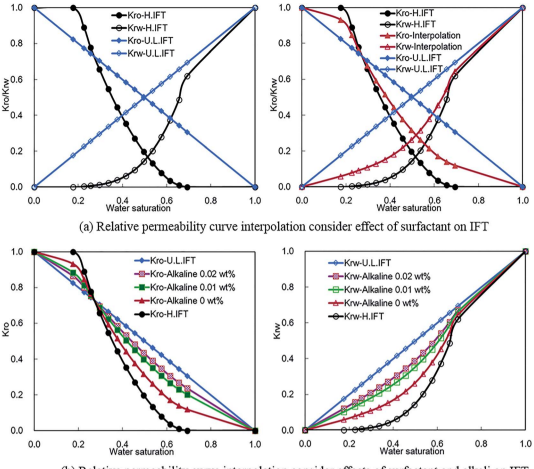
$$N_{\rm c} = \frac{\mu \nu}{j_{\rm A} \delta_{\rm S}} \tag{5}$$

$$j_{\mathbf{A}} = j_0 + a \times \mathbf{e}^{w_{\mathbf{A}}/b} \tag{6}$$

where j_A is IFT reduction factor due to the appearance of alkali; w_A is mass fractions of alkali, wt%; j_0 , and a and b are fit

IFT changes with the variation of surfactant and alkali concentrations, then the corresponding capillary number can

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(b) Relative permeability curve interpolation consider effects of surfactant and alkali on IFT

Fig. 7 Relative permeability curve interpolation methods, (note that Kro and Krw represent oil and water phase relative permeability separately; U.L.IFT and H.IFT are abbreviations of ultra-low and high interfacial tension separately).

be calculated by formula (5). Improved water phase relative permeability for different surfactant and alkali combinations can be calculated according to the interpolation method shown in formula (7).36 The oil phase relative permeability curve interpolation method is similar to that of the water phase:

$$k_{\text{rw}} = k_{\text{rwh}} \times \left\{ 1 - \left[\frac{\lg(N_{\text{c}}) - d_{\text{h}}}{d_{\text{l}} - d_{\text{h}}} \right]^{I_{\text{w}}} \right\} + k_{\text{rwl}} \times \left[\frac{\lg(N_{\text{c}}) - d_{\text{h}}}{d_{\text{l}} - d_{\text{h}}} \right]^{I_{\text{w}}}$$
(7)

$$d_{\rm h} = \lg(N_{\rm curr}) \tag{8}$$

$$d_{\rm l} = \lg(N_{\rm c_{\rm ILL,IFT}}) \tag{9}$$

where k_{rw} is water phase relative permeability after interpolation; k_{rwh} and k_{rwl} are water phase relative permeability under high and ultra-low IFT conditions separately; $d_{\rm h}$ and $d_{\rm l}$ are data interpolation parameters under high and ultra-low IFT conditions separately; $N_{c_{\rm H.IFT}}$ and $N_{c_{\rm U.L.IFT}}$ are capillary numbers under high and ultra-low IFT conditions separately; and $I_{\rm w}$ is a curvature and interpolation parameter of the water phase.

Oil-water phase IFT is high for water flooding without alkali, surfactant, and polymer. The corresponding oil and water phase relative permeability curves are shown by Kro-H.IFT and Krw-H.IFT separately in Fig. 7. IFT will reduce to an ultra-low level, and oil can mix with water completely when surfactant (or surfactant and alkali) reach to a critical concentration. Then, oil and water phase relative permeability curves will convert to straight lines (see Kro-U.L.IFT and Krw-U.L.IFT in Fig. 7). According to a traditional interpolation method, the relative permeability curve for actual ASP flooding can be interpolated based on the two sets of original relative permeability curves shown in left chart of Fig. 7(a). The interpolation results of Krointerpolation and Krw-interpolation can be observed in Fig. 7(a) (corresponding surfactant concentration is 0.1 wt%). It should be mentioned that the interpolated relative permeability curve only varies with surfactant concentration. So, it can't characterize the synergistic effects of surfactant and alkali on IFT reduction for ASP flooding.

For different alkali concentration conditions (surfactant concentration is fixed as 0.1 wt%), corresponding oil and water phase relative permeability curves can be interpolated based on the improved interpolation method, and the results are shown

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in Fig. 7(b). The interpolated relative permeability curves describe the effects of surfactant and alkali on oil-water phase filtration behavior. It can be observed that oil and water phase relative permeability capacities increase significantly with the join in of surfactant (Kro-H.IFT to Kro-Alkali 0 wt% and Krw-H.IFT to Krw-Alkali 0 wt%). The effect of alkali on relative permeability decreases gradually with the increase of alkali concentration. It is interesting to note that the effect of surfactant on relative permeability improvement is dominant compared with that of alkali. These phenomena are in good agreement with experimental results shown in Fig. 4. This implies that the improved relative permeability curve interpolation method is more appropriate for ASP flooding.

5.3. Multicomponent adsorption characterization

Generally, adsorptions of alkali and polymer usually obey a Langmuir adsorption isotherm. By contrast, surfactant adsorption is affected by alkali and polymer at the same time according to experimental results shown in Fig. 5. So, surfactant adsorption can't be characterized by a Langmuir adsorption isotherm model anymore. The effect of polymer on surfactant adsorption is slight enough compared with that of alkali that we neglected it in this study. Hence, surfactant adsorption analysis can be simplified into surfactant and alkali double factors. Multivariate regression was conducted based on the experimental result shown in Fig. 5(b). A multicomponent adsorption isotherm model which considers the effects of alkali and surfactant at the same time was constructed and the corresponding formulas are shown as follows:

$$S_{\rm ad} = g + h \times e^{E_{\rm A}} + m \times e^{E_{\rm S}} + n \times e^{E_{\rm A} + E_{\rm S}}$$
 (10)

$$E_{\rm A} = -e^{\frac{a_1 - w_{\rm A}}{a_2}} \tag{11}$$

$$E_{\rm S} = -\mathrm{e}^{\frac{s_1 - w_{\rm S}}{s_2}} \tag{12}$$

where S_{ad} is surfactant adsorption fraction; w_A and w_S are mass fractions of alkali and surfactant, wt%; a_1 , a_2 , s_1 and s_2 are attribute constants depending on the properties of alkali and surfactant; and g, h, m and n are fit coefficients.

This multicomponent adsorption isotherm model can be used to characterize surfactant adsorption phenomenon under alkalisurfactant-polymer multicomponent adsorption condition during an ASP flooding process. Fig. 8 shows surfactant adsorption contour lines which are plotted based on the proposed model. It can be observed that the differences between proposed model simulation result (Fig. 8) and experimental results (Fig. 5(b)) are insignificant. This validates that the accuracy of our proposed model is acceptable so now it can be used to conduct multicomponent adsorption numerical simulations.

Numerical simulation model construction and validation

A static simulation core (as shown in Fig. 9(b) and (d)) which has the same properties as our experimental core was established (see Fig. 9(a) and (c); the details are listed in Table 2). A

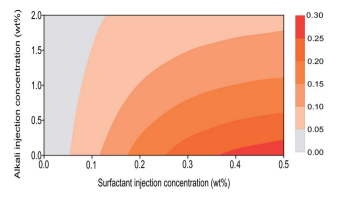


Fig. 8 Surfactant adsorption contour plot based on multicomponent adsorption isotherm model.

total of 2100 grids were designed for each cross section (see Fig. 9(a) and (b)). Fig. 9(c) and (d) show the experiment core and corresponding simulation core from a longitudinal view. A cylindrical grid system was employed to simulate plane radial flow in the core during a flooding process. It can be observed that the simulation core was divided into 50 segments lengthways. There is a total of 105 000 (21 \times 100 \times 50) grids in the simulation core after meshing. Based on an established simulation core, a novel ASP flooding numerical simulation model was constructed, which integrated the created viscosity nonlinear logarithm mixing method, IFT reduction-relative permeability curve interpolation method, and multicomponent adsorption isotherm model. To validate the effectiveness of this proposed novel numerical simulation model, experiments with core numbers 1, 2, and 3 (as shown in Table 2) were repeated using this numerical simulation.

Numerical simulation results based on our proposed model and a traditional model are shown in Fig. 10, and the corresponding experimental results collected from Fig. 2 also were supplied for comparison. It can be observed that all water cut curves in Fig. 10 show a U-shape during ASP flooding process, which validates the analysis results of the characteristic study. Compared with a traditional model, water cut simulated by our proposed model declines more significantly and also is much closer to experimental results for early, middle, and late stages of ASP flooding. This indicates that our proposed model accurately can characterize filtration behavior of ASP flooding. This is due to the proposed model being developed using the numerical simulation methods of viscosity mixing, IFT reduction, and multicomponent adsorption, and then enhancing the characterization of synergistic effects within an alkali-surfactant-polymer during ASP flooding process. In brief, comparing results validated the effectiveness and advantage of our proposed model for ASP flooding simulations.

Furthermore, our proposed model can be used to enhance the study of ASP flooding because it can overcome the problem with experiments in which it is impossible to measure all parameters exactly. On one hand, it can be used to study filtration behavior in a core and cooperate with lab experiments to reveal an unknown mechanism of ASP flooding. On the other hand, it can also expand to an oilfield scale to conduct

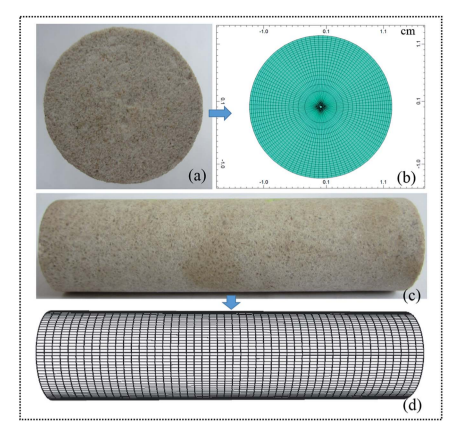


Fig. 9 Experiment core with corresponding static simulation core.

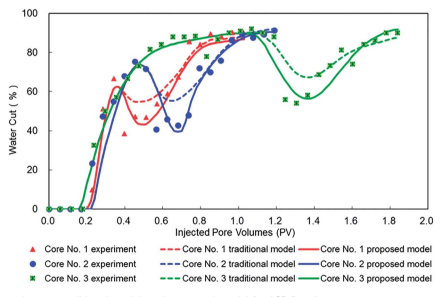


Fig. 10 Comparison of experiments, traditional model, and proposed model for ASP flooding.

development optimization and production prediction, and then enhance an ASP flooding filed application.

7. Conclusions

(1) EOR capacity of ternary chemical flooding is better than that of unitary and binary due to alkali-surfactant-polymer synergistic

effects. Water cut reverse, high injection pressure, and U shape of a water cut curve are response characteristics of ASP flooding.

(2) ASP solution viscosity presents non-linear variation behavior due to polymer molecules crosslinking probability growing exponentially. Alkali will diminish the molecular hydraulic radius of polyacrylamide and then limit its contribution to viscosity. A novel viscosity non-linear logarithm mixing simulation method was created to couple the multiple effects of alkali–surfactant–polymer on viscosity.

- (3) Diffusion equilibrium, reverse dissolution, and emulsification of surfactant will affect IFT reduction performance. Alkali can perform a synergetic action with surfactant on IFT reduction by producing *in situ* petroleum soaps. An improved relative permeability curve interpolation method which considers the effects of surfactant and alkali on IFT was developed to characterize their influences on filtration behavior.
- (4) ASP solution rheology improvement caused by polymer will restrain diffusion rate of a surfactant and then diminish its adsorption. Alkali can protect a surfactant from adsorption by pH improvement and an alternative mechanism. A multicomponent adsorption isotherm model was established to simulate adsorption behavior in an alkali–surfactant–polymer multicomponent environment.
- (5) A novel ASP flooding numerical simulation model was constructed which coupled developed synergistic mechanism characterization methods of viscosity non-linear logarithm mixing, relative permeability curve interpolation, and multicomponent adsorption. Comparison results validated the effectiveness of this novel model, and it can be used to enhance mechanism studies and field applications of ASP flooding.

Nomenclature

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ASP	Alkali–surfactant–polymer
IFT	Interfacial tension
m D	10^{-3} um^2

mD $10^{-3} \mu m^2$ wt% Weight percent PV Pore volume MPa 10^6 Pa

Conflicts of interest

There are no conflicts to declare.

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References

1 F. F. J. Lin, G. J. Besserer and M. J. Pitts, *J. Can. Pet. Technol.*, 1987, **6**, 54–65.

- 2 J. J. Sheng, Asia-Pac. J. Chem. Eng., 2014, 9, 471-489.
- 3 K. M. Ko, B. H. Chon, S. B. Jang and H. Y. Jang, *J. Ind. Eng. Chem.*, 2014, **20**, 228–233.
- 4 M. Sedaghat, O. Mohammadzadeh, S. Kord and I. Chatzis, *Can. J. Chem. Eng.*, 2016, **94**, 779–791.
- 5 B. Huang, W. Zhang, R. Xu, Z. Shi, C. Fu, Y. Wang and K. Song, *Energies*, 2017, **10**, 951.
- 6 Y. Ma, J. Hou, F. Zhao and Z. Song, *RSC Adv.*, 2018, **8**, 8269–8284.
- 7 H. A. Nasr-El-Din, B. F. Hawkins and K. A. Green, J. Pet. Sci. Eng., 1992, 6, 381–401.
- 8 Z. Tong, C. Yang, G. Wu, H. Yuan, L. Yu and G. Tian, *SPE/DOE Improved Oil Recovery Symposium*, Tulsa, Oklahoma, 1998, SPE 39662.
- 9 W. Pu, C. Yuan, W. Hu, T. Tan, J. Hui, S. Zhao, S. Wang and Y. Tang, *RSC Adv.*, 2016, **6**, 50640–50649.
- 10 M. Nedjhiouia, N. Moulai-Mostefa, A. Morslia and A. Bensmailib, *Desalination*, 2005, **185**, 543–550.
- 11 T. Guo, Y. Li, Y. Ding, Z. Qu, N. Gai and Z. Rui, *Energy Fuels*, 2017, 31, 10479–10489.
- 12 W. Pu, R. Liu, B. Li, F. Jin, Q. Peng, L. Sun, D. Du and F. Yao, *RSC Adv.*, 2015, 5, 88002–88013.
- 13 H. Gong, H. Zhang, L. Xu, K. Li, L. Yu, Y. Li and M. Dong, *RSC Adv.*, 2017, 7, 39564–39575.
- 14 Z. Liu, X. Yue, J. Hou and L. Zhang, SPE Asia Pacific Oil and Gas Conference and Exhibition, Melbourne, Australia, 2002, SPE 77876.
- 15 P. Shen, J. Wang, S. Yuan, T. Zhong and X. Jia, *SPE J.*, 2009, 237–244.
- 16 J. Hou, Z. Liu, S. Zhang, X. Yue and J. Yang, J. Pet. Sci. Eng., 2005, 47, 219–235.
- 17 V. Bortolotti, P. Macini and F. Srisuriyachai, *SPE Asia Pacific Oil and Gas Conference and Exhibition*, Jakarta, Indonesia, 2009, SPE 122499.
- 18 D. B. Levitt, S. Dufour, G. Pope, D. Morel and P. Gauer, International Petroleum Technology Conference, Bangkok, Thailand, 2012, IPTC 14915.
- 19 H. Mohammadi, M. Delshad and G. A. Pope, SPE Reservoir Eval. Eng., 2009, 518–527.
- 20 L. E. Zerpa, N. V. Queipo and J. L. Salager, SPE/DOE Fourteenth Symposium on Improved Oil Recovery, Tulsa, Oklahoma, 2004, SPE 89387.
- 21 S. Yuan, P. Yang, Z. Dai and K. Shen, *International Meeting on Petroleum Engineering*, Beijing, PR China, 1995, SPE 29904.
- 22 R. Moreno, L. Anselmi, D. Coombe, C. Card and I. Cols, Petroleum Society's Canadian International Petroleum Conference, Calgary, Alberta, Canada, 2003, SPE 2003–199.
- 23 S. L. Van and B. H. Chon, J. Ind. Eng. Chem., 2016, 38, 200–210.
- 24 R. Farajzadeh, T. Matsuura, D. van Batenburg, H. Dijk and B. V. Rijswijk, *SPE Enhanced Oil Recovery Conference*, Kuala Lumpur, Malaysia, 2011, SPE 143671.
- 25 K. Wyatt, M. J. Pitts and H. Surkalo, *SPE Eastern Regional Meeting*, Lexington, Kentucky, USA, 2002, SPE 78711.
- 26 J. Vargo, J. Turner, B. Vergnani, M. J. Pitts, K. Wyatt, H. Surkalo and D. Patterson, SPE Reservoir Eval. Eng., 2000, 3, 552–558.

Paper

27 H. L. Chang, Z. Q. Zhang, Q. M. Wang, Z. S. Xu, Z. D. Guo, H. Q. Sun, X. L. Cao and Q. Qiao, *JPT*, *J. Pet. Technol.*, 2006, 58, 84–89.

- 28 D. Wang, J. Cheng, J. Wu, Z. Yang, Y. Yao and H. Li, *SPE Asia Pacific Improved Oil Recovery Conference*, Kuala Lumpur, Malaysia, 1999, SPE 57288.
- $29\,$ S. Huang and M. Dong, J. Can. Pet. Technol., 2004, 43, 56–61.
- 30 J. Shen, Y. Bai, Q. Yin, W. Wang, X. Ma and G. Wang, *J. Ind. Eng. Chem.*, 2017, **56**, 82–89.
- 31 M. Chahardowli, R. Farajzadeh and H. Bruining, *J. Ind. Eng. Chem.*, 2016, **38**, 50–60.
- 32 R. Farajzadeh, A. Ameri, M. J. Faber, D. W. van Batenburg, D. M. Boersma and J. Bruining, *Ind. Eng. Chem. Res.*, 2013, 52, 13839–13848.
- 33 D. Li, M. Shi, D. Wang and Z. Li, SPE International Symposium on Oilfield Chemistry, Woodlands, Texas, USA, 2009, SPE 121598.
- 34 S. Liu, D. L. Zhang, W. Yan, M. Puerto, G. J. Hirasaki and C. A. Miller, *SPE J.*, 2008, 1–16.
- 35 D. Camilleri, A. Fil, G. A. Pope, B. A. Rouse and K. Sepehrnoori, *SPE Reservoir Eng.*, 1987, 433–440.
- 36 M. R. Todd, J. K. Dietrich, A. Goldburg and R. G. Larson, *The Fifth Symposium on Improved Methods for Oil Recovery*, Tulsa, Oklahoma, USA, 1978, SPE 7077.