Scale inhibitor testing of new organophosphoric compounds

A new generation of organophosphorous scale inhibitors has been developed. Inhibitors are intended to prevent the deposition of sulphate and carbonate salts in surface and downhole equipment during oil production, treatment and transportation. The I type inhibitor (aminophosphonic acid) was synthesized through the Moedritzer-Irani reaction of the condensation of 4,4'- (propane-2,2-diylbis(4,1-phenylene)bis(oxy)dianiline (A diamine) with formaldehyde and phosphorous acid, the II type inhibitor by phosphorylation of glycolyzed PET flex-(bis-hydroxyethyl terephthalate) with polyphosphoric acid. The structure of the developed organophosphorus compounds was confirmed by IR spectroscopy. High inhibitory efficacy of the first and second types of reagents for the deposition of carbonate and calcium sulphates was revealed. The morphology of the solid sediments of mineral salts was studied by scanning electron microscopy (SEM). It is revealed that the inhibition of crystal formation occurs at the stage of nucleation. Embedding in the surface of crystal nucleus, the inhibitors prevent formation of crystalline phase in a solution and crystal growth.

Keywords: organophosphoric compounds, aminophosphonic acid, phosphoric acid ester, scaling inhibitors, synthesis, reaction of Moedritzer-Irani, glycolyze, testing.

Introduction

Oil production in Kazakhstan has been steadily increasing over the last several years, while the average water cut of oil wells has been increasing, for some fields, water cut exceeds 90 % [1]. As a result of increased water cutting of the extracted products, the inorganic salt scale precipitation on the surface of pumping equipment, infield equipment, in production columns, in oil collection and treatment systems increases that result in a failure of expensive equipment, capital expenditures on equipment repair, and decline in oil production [2].

The efficiency of suppression of scale inhibitors is comprehensive, taking into account physical and chemical processes, the composition of produced water, development of wells, etc. Preferred method of combating scale buildup is inhibitor protection that refers to preventative methods [3].

Inorganic compounds, different classes of organic compounds can be scale inhibitors, but the compounds based on derivatives of phosphonic and phosphoric acids have received the greatest distribution, as these compounds are capable of forming complex compounds with salt-forming cations and preventing their interaction with salt-forming anions [4].

In this paper, we present the results of a comparative evaluation of the effectiveness of protection against the deposition of sulphate and carbonate salts by new organophosphoric compounds of 2 types — aminophosphonic acid and phosphoric ester of bis-hydroxyethyl terephthalate.

Experimental

Reagents and materials. Phosphonic acid 98+% (Alfa Aesar, Karlsruhe, Germany); formaldehyde, aqueous 37 % solution (Sigma-Aldrich); 4,4'-(propane-2,2-diylbis(4,1-phenylene)bis(oxy)dianiline (diamine A) the product purity is 99.9 % by 1H NMR spectroscopy; bis-hydroxyethylterephthalate obtained by the glycolysis reaction with ethylene glycol of a secondary PET flex, the structure of the product was confirmed by FTIR spectroscopy; polyphosphoric acid 115 % (Sigma-Aldrich); sodium chloride, State Standard 233–77; hexavalent magnesium chloride State Standard 4209–77; sodium bicarbonate State Standard; calcium chloride, dehydrated State Standard 6–09–47–11–81: ethylenediaminetetraphosphonic acid (EDTA); filtering paper; distilled water; laboratory scales in accordance with State Standard 24104–01.

Synthesis of organophosphates. Aminophosphonic acid was obtained by condensation of diamine A, formaldehyde and phosphorous acid through the Moedritzer–Irani reaction. The synthesis technique is described in the paper [5].
The phosphoric ester was obtained by phosphorylation of bis-hydroxyethylterephthalate with polyphosphoric acid. The synthesis technique is described in the paper [6]. The structure of synthesized organophosphates was confirmed by IR spectral analysis using Agilent Carry 660 IR spectrometer and Bruker Tensor II Fourier IR.

Static tests for scale inhibition [7]. The evaluation of the effectiveness of the scale inhibitor aminophosphonic acid against calcium carbonate and calcium sulphate deposits under static conditions was carried out according to previously described methods. The static laboratory method consists of precipitating CaCO₃ and CaSO₄ from water, i.e. heating with or without the presence of a scale inhibitor. In our work we used imitates of formation water of the chloride-calcium and sulphate-calcium type. The salt composition that simulates carbonate deposits is Ca²⁺ = 1100 mg/L, Mg²⁺ = 380 mg/L, Na⁺ = 7500 mg/L, HCO₃⁻ = = 976 mg/L, Cl⁻ = 14023 mg/L. Two solutions are required to imitate scale formation. Solution 1 contains 3046 mg/L calcium chloride and hexavalent magnesium chloride 3177 mg/L, while solution 2 is sodium chloride 18084 mg/L and sodium bicarbonate 1344 mg/L.

The effectiveness of scale inhibition was determined for concentrations of 10, 30, 50, 70, 100, 150 or 200 ppm of solution 1 in 50 mL. This was mixed with 50 mL of solution 2. Lids were placed on the flasks, and the temperature was fixed at 75 °C for 5 hours. Then the samples were filtered and the calcium ion content determined. 10 mL of filtrate were placed in the conical flasks and 40 mL of distilled water, 2.5 mL of 20 % sodium hydroxide solution and 10–15 mg of a mixture of murexide and sodium chloride were added. Secondly, the salt composition simulating sulphate (gypsum) deposits is Ca²⁺ = 4911 mg/L, Mg²⁺ = 148 mg/L, Na⁺ = 11623 mg/L, SO₄²⁻ = 8792 mg/L, Cl⁻ = 20291 mg/L. To prepare the imitation of scale formation in this case, two further solutions are needed. Solution 3 is calcium chloride 1360 mg/L, while solution 4 contains sodium chloride 18800 mg/L, hexahydrate magnesium chloride 1240 mg/L and sodium sulphate 13000 mg/L. The same procedure was carried out with solutions 3 and 4 as for solutions 1 and 2. The content of calcium ions in each case is calculated by the formula:

\[ X = \frac{V_t \times N_t \times 20.04 \times 1000}{10}, \]

where \( V_t \) — volume of EDTA-Na₂, which went to titration, ml; \( N_t \) — molar concentration of equivalent solution of EDTA-Na₂, mol /L equivalent; 10 — volume taken for titration; 20,04 — molar mass of the equivalent of calcium ion, g/mol.

For comparison, the experiment is conducted in parallel without the addition of scale inhibitor. The effectiveness of the scale inhibitor (%) is determined by the formula:

\[ E = \frac{C_0 - C_s}{C_0} \times 100 \%, \]

where \( C_s \) — content of calcium ions in a sample that does not contain an inhibitor, mg/L; \( C_p \) — content of calcium ions in the sample containing the inhibitor, after thermostating, mg/L; \( C_0 \) — content of calcium ions in the initial solution, mg/L.

Results and discussion

Well-known foreign technologies for controlling scale precipitation of mineral salts on the surface of process equipment at petrochemical complexes are based on the treatment of water with inorganic phosphate derivatives [8]. In water treatment systems for inorganic phosphates, there are environmental limitations in the concentration due to the fact that phosphates promote excessive growth of algae, oxygen deficiency and extinction of fish [9].

One of the safe water treatment technologies is the use of organic phosphates, such as phosphoric esters, phosphonic acids. Organic phosphates contain less phosphorus, are biodegradable, have low toxicity, are not inferior in efficiency to inorganic phosphates. For their properties, these compounds are classified as complexons, their complexing properties are realized due to the presence of hydrogen atoms of phosphonic groups [10]. According to [10], the mechanism of scaling inhibition by organophosphates can be explained by the formation of complex compounds with crystal nucleus of mineral salts. Inhibition occurs due to adsorption of organophosphates on the surface of growing crystals of mineral salts, which excludes the possibility of formation of a crystalline phase in a solution. The scheme for the complex formation of the phosphoric ester with calcium ions is shown in Figure 1.
The new organophosphonates that we developed were identified and characterized [5, 6]. In order to confirm the structure, their IR spectral analysis was carried out. Figures 2 and 3 shows the IR spectra of aminophosphonic acid and phosphoric acid monoester of bis-hydroxyethylterephthalate, respectively.

![Figure 1. Scheme of complexation of the phosphoric acid ester with calcium ions](image1)

The IR spectrum showed absorption bands in the 2964 cm\(^{-1}\) region attributable to stretching vibrations in the -CH\(_3\) and -CH\(_2\) groups. The absorption band at 1590 cm\(^{-1}\) indicates the stretching vibrations of the C=C double bonds in the aromatic ring. The stretching vibrations at 1363 cm\(^{-1}\) correspond to the C-N bonds. The absorption band at 1173 cm\(^{-1}\) shows the stretching vibrations of the P=O group. The 738 cm\(^{-1}\) band describes the stretching vibrations of the P-C bond.

In the IR spectrum of the phosphoric ester of bis-hydroxyethylterephthalate, an absorption band is observed in the region of 1719 cm\(^{-1}\), which can be attributed to the stretching vibrations of the carbonyl group of the compound ester. The 2966, 2881 cm\(^{-1}\) bands are attributed to the vibrations of the CH bonds in the CH\(_2\) groups; in the region of 1504 cm\(^{-1}\), they are referred to vibrations in the aromatic ring, in the 2800 cm\(^{-1}\) region to the P-OH deformation oscillations. Fluctuations in the region of 985 cm\(^{-1}\) can be attributed to the deformation in the P-OC group, in the 1106 cm\(^{-1}\) region — to the stretching vibrations in P=O.
Based on the data of IR spectroscopic studies, the expected chemical structure of the organophosphates was obtained (Fig. 4).

![Chemical structure](image)

Propane-2,2 phenylene-diylbis(4,1)-phenylene)bis(4,1-bis(azadienyl)bis(methylene)diphosphonic acid (a)

![Chemical structure](image)

Bis(2-(phosphonoxy)ethyl)terephthalate (b)

Figure 4. Alleged chemical structure of organophosphates: phosphoric acid ester of bis-hydroxyethyl terephthalate (a) and aminophosphonic acid (b) [5, 6]

In order to identify the scale inhibition ability, organophosphonic compounds have been tested on different models of produced water. The tests were carried out in model solutions at a temperature of 75 °C and an inhibitor concentration from 10 to 200 ppm. The results of the inhibition studies are shown in Figures 5 and 6. As the figures show, the new reagents exhibit a sufficiently high inhibitory activity with respect to the CaCO₃ and CaSO₄ deposits. At the same time, it can be noted that the inhibition efficiency on CaCO₃ deposits was higher for aminophosphonic acid. The maximum protection at a concentration of 200 ppm of this reagent was 94 %, and for the phosphoric ester at the same concentration was 85 %. However, the inhibition efficiency on CaSO₄ deposits (Fig. 6) is much higher for phosphoric ester, the maximum efficiency was 100 % and reached at a concentration of 30 ppm, the maximum inhibition degree for aminophosphonic acid was 97 % at a concentration of 200 ppm.

![Figure 5](image)

Figure 5. Effectiveness of inhibitors on CaCO₃ deposits

![Figure 6](image)

Figure 6. Effectiveness of inhibitors on CaSO₄ deposits
The reason for the different efficiency of the inhibitors was elucidated in the study of scale samples by scanning electron microscopy using a JEOL JSM-649OLV instrument. The authors [11] note that CaCO₃ has the property of polymorphism, the precipitates have the same chemical composition, but they have different crystalline structures — calcite, aragonite and vaterite. Aragonite, unlike vaterite, has a more stable crystalline form [12]. The vaterite under normal conditions (25 ºC, atmospheric pressure) eventually transforms into a more stable form of calcite, and at temperatures (70–80 ºC) first transforms into aragonite, and then into calcite. Compared to calcite, aragonite and vaterite are soluble in water. This is due to the fact that aragonite and vaterite have a lower value of free surface energy as compared to calcite, which reduces their adhesion on the surface of the equipment, due to the interaction between the crystals [13]. Moreover, it was reported in [14–17] that under certain conditions occurrence of two more forms of calcium carbonate is possible: calcium carbonate monohydrate and calcium carbonate hexahydrate.

Figures 7, 8 and 9 shows micrographs of precipitation of mineral salts before and after treatment with inhibitors.

As can be seen on Figure 7, the difference in the formation of crystals is obvious. Crystals formed without an inhibitor are in the form of calcite. The phase composition of the precipitate obtained in the presence of aminophosphonate is characterized by a higher content of aragonite and vaterite. For the phosphoric ester, there are preferably more crystals in the form of vaterite. Also, from Fig. 6 for aminophosphonic acid and phosphoric ester, it follows that these compounds have complexing properties and are able to be embedded in the surface of the crystallization nucleus, as a result of which crystal growth ceases.

![Micrographs of carbonate sediments without an inhibitor and with scaling inhibitors](image)

$a$ — blank; $b$ — phosphoric acid ester (30 ppm); $c$ — aminophosphonic acid 30 ppm

Figure 7. Micrographs of carbonate sediments without an inhibitor and with scaling inhibitors

![Microphotographs of sulfate sediments](image)

$a$ — blank; after addition of phosphoric acid ester; $b$ — 5 ppm; $c$ — 10 ppm; $d$ — 20 ppm

Figure 8. Microphotographs of sulfate sediments

Figure 8a shows that the structure of deposits of sulphate sediments without an inhibitor has structures in the form of smooth needles. When phosphoric ester inhibitors were added, the destruction of gypsum deposits started (Figure 8b, 8c, 8d). Complete destruction of the deposit was achieved at a concentration of 30 ppm. A similar effect was observed for aminophosphonic acid (Fig. 9), but in this case the concentration of the reagent was higher than that of the phosphoric ester (Fig. 9a, b, c). The beginning of destruction of sulphate sediments was observed at a concentration of 10 ppm. Complete destruction was noted at 200 ppm.
Thus, it was shown that inhibitors have not only high inhibitory properties with respect to carbonate and calcium sulphate deposition, but are also capable of altering the morphology of crystals, which favorably affects the destruction of already fallen salt crystals and their removal with process fluid.

The inhibitors are highly effective for total deposits of carbonate salts (94 and 85 %) and sulphates (97 and 100 %) at a concentration of 200 ppm.

References


Тұз шогінділерінің ингибиторлары ретінде
жана фосфорлы органылық косылыстардың сынактары

Мұнай оңдіру, дайындау және тасымалдау барысында колданылатын жер бетіндегі және ұшынмадағы кондырғыларда сульфатты және карбонатты тұздардың шогінділеріне қорғағын, жаңа буныңдағы фосфорлы органылық тұз шогінділерінің ингибиторы жасап шығарылады. Бірнеше тұрлі ингибитор (аминофосфон қышқылы) Моедритцер-Иран реакциясы арқылы 4,4’-(пропан-2,2-диллиб(4,1-фенилен))бис(окси)диаминданын (диамин A) формалдегиддің және фосфорқышқылымың конденса-
циясы арқылы синтезделеді, екіншісі — гликолизденген ПЭТ флексі — полиацетафталең және полиацетатындагы бисгидроксиэтилтерефталатын фосфорлау арқылы. Жасалған фосфорлы органылық косылыс-
тарының құрылысының ИК-спектроскопия зқұралған. Сканерлесуші электронды микроскопия арқылы (РЭМ) минералды тұздардың жаттығуын морфологияға зерттеді. Кристалл тұзларының өзіндең арқылы құрылған сапының тазалуын және кристалдардың осуі бөлінеді.

Кізіл сөздер: фосфорлы органылық косылыстар, аминофосфон қышқылы, фосфор қышқылының құрделі эфирі, тұз шогінділері ингибиторлары, синтез, Моедритцер-Иран реакциясы, гликолиз, тестілеу.

Испытания новых органофосфорных соединений
в качестве ингибиторов солеотложений

Разработаны органофосфорные ингибиторы солеотложений нового поколения, предназначенные для предотвращения отложений сульфатных и карбонатных солей в наземном и скважинном оборудовании при нефтедобыче, подготовке и транспорте нефти. Ингибитор первого типа (аминофосфоновая кислота) был синтезирован по реакции Моедритцера-Иран конденсацией 4,4’-(пропан-2,2-диллиб(4,1-фенилен))бис(окси)диаминла (диамин A) с формалдегидом и фосфористой кислотой, второго — фосфорилированием гликолизированного ПЭТ флекса — бисгидроксиэтилтерефталаты полифосфорной кислотой. Структура разработанных органофосфорных соединений подтверждена методом ИК-спектроскопии. Растровой электронной микроскопией (РЭМ) изучена морфология твердых осадков минеральных солей. Выявлено, что ингибирование кристаллообразования происходит на стадии зародышобразования. Встраиваясь в поверхность зародышекристалла, ингибиторы препятствуют образованию кристаллической фазы в растворе и росту кристаллов.

Ключевые слова: органофосфорные соединения, аминофосфоновая кислота, сложный эфир фосфорной кислоты, ингибиторы солеотложения, синтез, реакция Моедритцера-Иран, гликолиз, тестирование.

References


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