Performance of a novel green scale inhibitor

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Abstract: Many aspects of oilfield scale inhibition with green scale inhibitors (SIs) have remained untouched. For instance, the discharge of large amounts of produced water containing various types of hazardous chemicals, such as SIs into the environment has become a major concern. Instead, environmental regulators encourage operators to look for greener SIs. In this study, the performance of a green SI was investigated using PHREEQC simulation. For a specific case study, two brines are considered to mix incompatibly to estimate the critical mixing ratio that has the highest tendency to scaling. Subsequently, for 50/50 mixing ratio as the critical value, the optimal dosage of SI and its performance in the presence of two different rocks were investigated such that 450 mg/L SI would be considered as optimal value. Moreover, the simulated results show that more SI adsorption on calcite would be predicted, compared to dolomite.

1 Introduction

In the oil and gas industry, one of the primary production problems is mineral deposition resulting from the water-flooding, incompatible water mixing, and/or hydro-fracturing processes that are applied to maintain sustainable hydrocarbon production in oil, gas, or gas-condensate fields [1]. Scaling is the deposition of inorganic salts due to supersaturated conditions when their solubility limit is exceeded. It can cause immediate and severe technical problems such as the reduction in production capacity, reservoir choking, accelerated corrosion and flow loss, or blockage on flow lines and equipment [2], which can influence both production and economic aspects of the process. Generally, the associated expenses due to scaling were estimated at some billion dollars per year in major developed industrialized countries [3].

Therefore, effective scale management is of great practical importance for oilfield development planning which will dominate production viability [4]. The utilization of scale inhibitors (SI) in oilfields is widespread due to their impact on the mitigation of scale formation as an efficient solution to enhance oil recovery (EOR) and prevent formation damage.

In general, the oil industry, like other industries, suffers from the environmental consequences of using hazardous chemicals to inhibit scale formation. The selection of environmentally friendly inhibitors which should be non-toxic, biodegradable, and show no bioaccumulation [1], and their appropriate dosage are two crucial factors to get the optimum
inhibitory action which must be achieved by simulating the well [5]. Although both types of phosphonate and polymeric SIs have been investigated experimentally [6-10], nevertheless, well-known polymeric environmentally friendly scale inhibitors, like poly aspartic acid (PASP) [11-14], polyepoxysuccenic acid (PESA) [15-17], and carboxymethyl inulin (CMI) [18-21], are gaining increased attention. Moreover, natural organic molecules and plant extracts like the green SIs [13,22-24] in this study, were assessed as brand-new SIs.

It is evident from the above that the applied study of green scale inhibitor would help to keep the environment safe and mitigate the scaling phenomenon in porous media in the oil industry. Accordingly, the present work endeavors to investigate a green SI for an Iranian oilfield water-flooding process through simulation by PHREEQC to evaluate its performance. Although data on complex interactions between SIs and brines are still lacking, this study nevertheless examines the impact of the presence of a green SI in an incompatible water mixing process in the absence and presence of rocks using PHREEQC based on their pH dependency. In other words, this work presents a brand-new approach to a novel green scale inhibitor performance through PHREEQC simulation.

2 Modeling

As a major operational problem, i.e., scale formation, which is recognized as a significant cause of formation damage [25], has frequently been observed in various oil reservoirs and interconnected production equipment in Iranian oilfields, specifically near the wellbores. Employment of an efficient green SI would be a perfect solution to overcome formation damage of various types, as well as to safeguard the environment in the water-flooding process.

Scale potential and specific SI performance for a particular case study (Table 1) have been investigated using PHREEQC simulation, as a pH-redox equilibrium geochemical model [26] to estimate the saturation ratio of the minerals that may precipitate, as well as the mass of deposits. Two brines are ionically characterized for expected simultaneous reactions (1-3), which will be taking place as they mix to reach equilibrium [27].

$$\text{Ca}^{2+} + 2\text{HCO}_3^- \rightarrow \text{CaCO}_3 \downarrow + \text{CO}_2 + \text{H}_2\text{O} \quad (1)$$

$$\text{Mg}^{2+} + \text{Ca}^{2+} + \text{CO}_3^{2-} \rightarrow \text{MgCa(CO}_3)_2 \downarrow \quad (2)$$

$$\text{Ca}^{2+} + \text{SO}_4^{2-} \rightarrow \text{CaSO}_4 \downarrow \quad (3)$$

Firstly, the critical mixing ratio of the two brines is examined to find out the maximum amount of precipitated particles at 90°C and atmospheric pressure [21,28,29] necessary to simulate incompatible mixing process at high-temperature standard static tests. Secondly, the critical mixing ratio test is conducted to assess the various dosage of the SI to evaluate the optimum dosage of the SI. Finally, the impact of 10 grams [29] of dolomite and calcite rock’s presence is studied to determine the SI adsorption capacity for each rock.

| Ion (ppm) | Formation Water | Seawater |
|-----------|-----------------|----------|
| [HCO$_3^-$] | 2274 | 183 |
| [SO$_4^{2-}$] | 665 | 2880 |
| [Cl$^-$] | 137114 | 26625 |
| [Ca$^{2+}$] | 14840 | 750 |
| [Mg$^{2+}$] | 3086 | 1350 |
| [Na$^+$] | 67102 | 19550 |
| [K$^+$] | - | 410 |
| pH | 6 | 7.5 |

Table 1. Ionic composition of investigated brines
The aforementioned simulation steps were performed with the help of PHREEQC which handles geochemical reactions at the equilibrium state. After the definition of both formation water and seawater in terms of ionic composition, temperature, pressure, and pH, and mixing these two brines, the equilibrium state of the mixing process was considered. Subsequently, the results were obtained the mole and mass variations of each participant. It should be noted that the simulations used experimental pH values after adding different doses of the green SI to seawater. In other words, based on the pH-dependent nature of the simulation software, it is intended to manipulate the pH values of the SI-added-seawater to study the SI performance in an incompatible mixing process.

3 Results and discussion

3.1 Scale Potential

To study mineral scale formation at high temperatures which simulates reservoir conditions, incompatible mixing of two brines was performed to estimate the potential of scale formation versus different mixing ratios of the brines. Figure 1 demonstrates that the highest scaling tendency occurs around 50% formation water which will be considered as the critical mixing ratio due to the maximum amount of predicted precipitates (gr) per one liter of water produced after mixing.

![Graph](image)

Fig. 1. Determination of scale potential

Moreover, both carbonated and sulfated particles were observed in Figure 1, confirming a complex multi-scale system that will be subsequently treated through the utilization of scale inhibitors. It is noteworthy that these results will reliably resemble experimental standard test conditions for a vessel [28].

3.2 Scale Inhibitor’s assessment

The predicted particle mass and inhibition efficiency are two predominant parameters for studying the green scale inhibitor performance. Scale inhibition efficiency is usually calculated [30] based on the change of calcium ion concentration (Eq. 4) as well as particle mass using Eq. (5) proposed by [31]:

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1. CaCO3
2. CaSO4
3. CaMg(CO3)2
4. total amount of precipitated particles

Mixing ratio of formation water

Mass of particles (g)

-2 0 2 4

0 0,5 1 1,5
\[
\%I = \left( \frac{[Ca^{2+}]_{in} - [Ca^{2+}]_{non-in}}{[Ca^{2+}]_{i} - [Ca^{2+}]_{non-in}} \right) \times 100
\]

\[
\%I \text{ (Std. ave. mean)} = \frac{m_0 - m}{m_0} \times 100,
\]

where \(\%I\), \([Ca^{2+}]_{in}\), \([Ca^{2+}]_{non-in}\), \([Ca^{2+}]_{i}\), \(m_0\), and \(m\) indicate percentages of inhibition, soluble calcium concentration of the inhibited sample, soluble calcium concentration of the non-inhibited sample, initial soluble calcium concentration, mass of non-inhibited particles, and mass of inhibited particles, respectively.

Accordingly, Figure 2 shows that increased dosage of the scale inhibitor into the seawater increases the inhibition efficiency until it shows a plateau above 450 mg/L. It can be concluded that 450 mg/L would be the appropriate dosage for utilization as the optimum dosage of the SI in order to minimize the mass of particles since it is observed in the blue curve. The finding would be justified by the fact that the addition of the SI reduces pH value due to its acidic functional groups’ presence in the SI’s chemical structure which may result in less precipitation [32].

3.3 Impact of Rock Presence

It should be noted that calcite and dolomite rocks are considered in this work due to two reasons: i) carbonated reservoirs account for more than 50% of the world’s hydrocarbon reservoirs [33] most common Iranian scales (calcium carbonate and calcium sulfate) generally result from the complex surface interactions between brines and carbonated rocks [34]. Consequently, in this part, the brines were exposed to ten grams of calcite and dolomite specifically to assess the SI performance.

According to Fig. 3a, the negative value of the mass of the dolomite particles indicates dissolution phenomena, as expected in the presence of more dolomite in the solid surface, although positive value of the mass of other particles indicates the precipitation mechanism. Similar findings are observed in Fig. 3b relating to dissolution of calcite in the presence of
calcite rock. One can unambiguously conclude that after coming to equilibrium, the dominant mechanism is precipitation due to the positive “net of particles” bar.

![Fig. 3a. Precipitation/dissolution mechanism in the presence of dolomite rock](image)

![Fig. 3b. Precipitation/dissolution mechanism in the presence of calcite rock](image)

Moreover, Fig. 4 illustrates the impact of two different rocks on the SI performance. More particles will be formed in the presence of the rock due to the dissolution phenomenon of calcium and/or magnesium ions compared to conditions where no rock is present.

Additionally, a comparison of the simulation results between dolomite and calcite shows that because of Mg^{2+} in dolomite rock, the SI would not be more effective for inhibi-
tion than calcite rock. In other words, it seems that the dolomite rock interferes with the SI adsorption compared to calcite [36].

![Graph showing predicted mass of particles in the absence and presence of rocks](image)

**Fig. 4.** Predicted mass of particles in the absence and presence of the rocks

## 4 Conclusions and recommendations

In this study, the performance of a green scale inhibitor is investigated using an incompatible mixing process and in the presence of rock based on the geochemical PHREEQC model. The main results of this work are as follows:

- The critical mixing ratio of formation and seawater is 50/50 according to the maximum amount of the predicted mass of particles at 90°C and atmospheric pressure.
- The optimum SI dose for the defined system is 450 mg/L, because after this dose an asymptotic behavior for inhibition efficiency is observed.
- Although the presence of rock affects SI performance in such a way that more particles are formed compared to the condition of no rock, nonetheless, the adsorption of the SI on calcite is greater than on dolomite. In other words, the SI would be more efficient in the presence of calcite.
- Since PHREEQC is based on equilibrium conditions and reaching equilibrium is not conceivable in reality, experimental tests, as well as characterization analyses such as SEM/EDX are highly recommended to elucidate the SI mechanism. It should be noted that achieving qualitative similar trends between experimental and simulated results would be reasonable and sufficient, consistent with a complex geochemical system in the presence of a chemical like a scale inhibitor.

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