Initiation and inhibition of pitting corrosion on C-steel in oilfield-produced water under natural corrosion conditions

S. Abd El Wanees, M.M. Kamel, S.M. Rashwan, Y. Atef, M.G. Abd El Sadek

Chemistry Department, Faculty of Science, Zagazig University, Zagazig 44519, Egypt, email: s_wanees@yahoo.com (S. Abd El Wanees)
University College of Umluj, University of Tabuk, Tabuk, Saudi Arabia, email: s_nasr@ut.edu.sa
Chemistry Department, Faculty of Science, Suez Canal University, Ismailia, Egypt, email: madhet_kamel@yahoo.com (M.M. Kamel)
Research Laboratory, Egyptian Chemistry Administration, Cairo, email: jasy9@yahoo.com (Y. Atef)
Research and Chemicals Department, General Petroleum Company, GPC, Nasr City, Cairo, email: msadek.2004@gmail.com (M.G. Abd El Sadek)

Received 22 July 2021; Accepted 4 December 2021

ABSTRACT

The initiation and inhibition of the localized pitting corrosion on the C-steel surface in oilfield-produced water are investigated by pitting corrosion current measurements. The current-time curves are characterized by the appearance of an induction period, $\tau$, followed by a rapid rise in the pitting corrosion current till reaches the limiting value, $I_L$, due to propagation of pitting corrosion. The $I_L$ value reaches 5.0 $\mu$A cm$^{-2}$ in 99% diluted oilfield-produced water and 260 $\mu$A cm$^{-2}$ in pure oilfield-produced water. The presence of HPO$_4^{2-}$, WO$_4^{2-}$, and MoO$_4^{2-}$ anions suppresses the initiated current by elongation of the induction time, $\tau$. Such anions acted as inhibitors towards the pitting corrosion of the C-steel. The inhibition efficiency, $\eta$, of the studied inhibitors increases in the following order: Na$_2$WO$_4 < Na$_2$MoO$_4 < Na$_2$HPO$_4$. The inhibition mechanism is assumed to take place through an adsorption process obeying Langmuir’s model. The thermodynamic parameters for the adsorption process $K_{ads}$ and $\Delta G_{ads}$ are calculated and discussed.

Keywords: Oilfield-produced water; C-steel; Pitting corrosion; Inorganic passivator; Adsorption

1. Introduction

Nearly all oil production wells have chloride and sulfate salts mixed with the production water or emulsified with crude oil causing a dangerous type of pitting corrosion for steel equipment utilized in oil and gas production. Equipment prepared from C-steel can be attacked easily by aggressive anions which are naturally contaminated the formation water. The presence of chloride and sulfur ions which are abundant in the underground water can easily severe corrosive problems in frameworks such as equipment and pipelines in processes of recycling and reuse of produced water that leads to considerable economic losses.

Several corrosion inhibitors are added to oilfield-production water during petroleum output for controlling the destructive effect of the corrosive anions which are mixed with the produced water environment. A broad of corrosion inhibitors are utilized to reduce the corrosion of the steel in the various corrosive media [1–8]. Generally, most of the used inhibitors are organic and inorganic materials, which generally form a protective layer on the metal surface. The unused inhibitors are released into the environment as an aqueous waste and thus affect the living organisms due to the toxic properties of some of the used chemicals. The use of inorganic inhibitors as an alternative
to organic compounds is based on the possibility of degradation of organic compounds with time and temperature [9,10].

Various inorganic salts are utilized as effective inhibitors or passivators for various metals and metal alloys in different electrolytes [11–17]. As a result of the poisonous effect of most of these ions, it limited the toxic use of these anions. Therefore, safe anions without any polluted effect were chosen as inhibitors such as Na$_2$HPO$_4$, Na$_2$MoO$_4$, and Na$_2$WO$_4$ towards the corrosion of C-steel in oilfield-production water.

In our study, the influence of the oilfield-produced water obtained from North East Sannan, NES-4, in the Western Desert, Egypt, in a depth of 2,415–2,435 m on the induced pitting corrosion of C-steel was studied. The investigation was carried out by using a simple cell constructed early for measuring the localized pitting corrosion current under natural corrosion conditions [13,18–20].

2. Experimental

2.1. Materials and the electrolytic cell

The used electrodes were prepared from carbon steel C1018. The elemental analysis of the steel samples was estimated, as weight percent, by Tabin Institute for Metalurgical Studies (TIMS), Helwan–Cairo, Table 1.

The C-steel electrodes were reinforced to Pyrex-glass pipes using Araldit resin leaving a free cross-sectional surface area is 0.04 cm$^2$. Before each run, the surface of the steel electrodes was mechanically abraded using various grades of polished papers, rinsed with acetone, and finally washed with doubly distilled water.

Due to localized pitting corrosion on most metals and steel surfaces, the pitting corrosion current can be measured using a simple electrolytic cell that is described early, Fig. 1 [13,18–20]. The cell was made of a double wall borosilicate glass with a volume of 100 cm$^3$ solution and comprising a C-steel electrode, A, as an anode. The cathode was a similar C-steel electrode with the same surface area and elemental composition, B. The cathode was enclosed in a borosilicate glass tube ending with a fine porosity fritted centered glass disc G$_4$, C, and included in the cell container. A copper wire was used for connection between the steel electrodes using a nano-amperemeter device (Siemens type N-5536).

2.2. Corrosive electrolyte, oilfield-produced water

Generally, deep oilfield-produced water is liberated in the reservoir rocks before drilling. Such water utilized in our study was gained from one of the wells of the general petroleum company, GPC, Egypt, in the Western Desert, North East Sannan, NES-4 in a depth of 2415–2435 m underground mixed with crude oil. The chemical composition and physical properties of the produced water are given in Table 2.

Table 1
The elemental analysis of utilized steel electrodes, C1018, in wt.%

<table>
<thead>
<tr>
<th>Element</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>0.60</td>
</tr>
<tr>
<td>Si</td>
<td>0.17</td>
</tr>
<tr>
<td>P</td>
<td>0.046</td>
</tr>
<tr>
<td>Mn</td>
<td>0.50</td>
</tr>
<tr>
<td>C</td>
<td>0.12</td>
</tr>
<tr>
<td>Fe</td>
<td>Bal 98.564 mass %</td>
</tr>
</tbody>
</table>

Table 2
Physical and chemical characterization of the oilfield-produced water

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>7.10</td>
</tr>
<tr>
<td>Cl$^-$, ppm</td>
<td>41,477</td>
</tr>
<tr>
<td>SO$_4^{2-}$, ppm</td>
<td>244</td>
</tr>
<tr>
<td>Cl$^-$/SO$_4^{2-}$</td>
<td>170.1</td>
</tr>
<tr>
<td>Ca$^{2+}$, ppm</td>
<td>2,525</td>
</tr>
<tr>
<td>Mg$^{2+}$, ppm</td>
<td>122</td>
</tr>
<tr>
<td>Conductivity, mS cm$^{-1}$</td>
<td>95.2</td>
</tr>
<tr>
<td>Sulfur, %</td>
<td>0.2021</td>
</tr>
<tr>
<td>Density, at 25°C, g cm$^{-3}$</td>
<td>1.03293</td>
</tr>
<tr>
<td>Total dissolved salts (TDS), mg L$^{-1}$</td>
<td>47,600</td>
</tr>
</tbody>
</table>

Fig. 1. Electric circuit and electrolytic cell used for pitting corrosion current measurement. A and B are identical C-steel electrodes, C is a fritted centered glass disc G$_4$ and nA is a nano amperometer.
The other set was conducted with 50% of oilfield-produced water mixed with different additives of the used inorganic salts, Na$_2$HPO$_4$, Na$_2$MoO$_4$, and Na$_2$WO$_4$. Each run was done with freshly prepared electrodes and with a new portion of the examined solution. This was carried out by using the same percent of diluted oilfield-produced water mixed with various additives of the investigated salts. The produced pitting corrosion current was tabulated against the immersion time. To ensure the repeatability of the results, each experiment was duplicated and the data were nearly similar. The average value of the produced current was taken.

The utilized inorganic salts were of analytical grade quality. The examined electrolytes were prepared by using double-distilled water. Experiments were done at room temperature, 25°C ± 0.1°C, excluding those investigated to the various temperatures. The cell temperature was monitored by using an ultra-programmable thermostat (USA). Scanning electron microscopy of some of the investigated steel electrode samples was done using a JEOL Scanning Microscope, JSM–T100 (Japan).

3. Results and discussion

3.1. Influence of dilution factor on the oilfield-produced water

The corrosion current density produced when the C-steel electrode was immersed in oilfield-produced water was followed against the immersion time till reaching the limiting current, $I_l$. The curves of Fig. 2 show the behavior recorded in solutions of various diluted oilfield-produced water (polluted by Cl$^-$ and S ions). It is noteworthy to see that, with more diluted oilfield-produced water, 1.0% FW (fewer amounts of the aggressive Cl$^-$ and S ions, very little corrosion current density (5.0 μA cm$^{-2}$) appears after an induction period extends to 140 min. The current starts to decay due to the passivation of meta-stable pits [21–26]. Such highly diluted oilfield-produced water contains relatively small amounts of aggressive ions (Cl$^-$ and S) that are not sufficient to initiate the pitting corrosion [13,25].

At this stage, the passivating anions naturally present in the examined electrolytes such as OH$^-$ and CO$_3^{2-}$ displaced the attacking ions such as Cl$^-$ and S ions at the pit surface causing the decay in the localized pitting current [27]. As the dilution percent of the oilfield-produced water is lowered (an increase in the amount of Cl$^-$ and S ions), the corrosion current starts to increase with decreasing the induction time, $\tau$. In the case of 85.0% FW, the induction time is lowered to 25 min with pitting corrosion current, $I_l$, reaching 195 μA cm$^{-2}$, while with pure oilfield-produced water, $\tau$ was shortened to ~20 min with increasing in the $I_l$ to reach 260 μA cm$^{-2}$. The appearance of such a high current is an indication of the presence of an electrochemical reaction at the electrode surface [22–26]. Generally, literature has been confirmed that nucleation, growth, and passivation of metastable formed pits on the C-steel surface occur during the formation of metastable pits [28–32]. Some of the formed pits are stable whereas most of them are passivated again. As the rate of pit passivation is lower than that of oxide film destruction, meta-stable pits are supposed to be formed with an increase in the pitting corrosion current [32,33]. Various models about the growth of metastable pits are suggested [33–37]. Early, Williams indicates that the meta-stable pits have a very restricted existence depending on the concentration of the corrosive anions in the oilfield-produced water [34,35]. Subsequently, the pits have been passivated and the initiated current cutout to flow.

However, the increase in the corrosive ions content (Cl$^-$ and S ions) in the oilfield-produced water by decreasing the dilution ratio causes the pitting current to outflow with a decrease in the induction period, $\tau$ till reaching the limiting value, $I_l$. This interval is usually accompanied by the initiation of the localized pitting corrosion [27,28,38,39] and could be related to the time taken by the Cl$^-$ and S ions either to penetrate the passive oxide film [40] or to compete with the passivating anions OH$^-$ and CO$_3^{2-}$ to adsorb on the corroding steel surface [41]. Obvious pits are monitored by screening the surface of the steel electrode, under the scanning electron microscopy (SEM), after running the experiment in 50% diluted oilfield-produced water, Fig. 3. The existence of the Cl$^-$ and S ions give rise to the consistency of intelligible pits. The formed pits take various shapes (fine and moderate in size) and some of them are surrounded by the corrosion products that formed on the electrode surface.

The dependence of the induction time, $\tau$, on the amount of the destructive anions, $C_{FW}$ % (Cl$^-$ and S ions), in oilfield-produced water can be depicted in Fig. 4. The data of such curve performs the logarithmic relation between the two parameters, that is obeying the relation [13,23–26]:

$$\log \tau = \zeta - \lambda \log C_{FW}$$

(1)

where $\zeta$ and $\lambda$ are constants that depend on the electrolyte composition and investigated electrode kind. The values of the constants $\zeta$ and $\lambda$ are 173 min and -0.45 log min/decade, respectively.

However, the steady rise in the pitting current could confirm the propagation of the localized pitting corrosion or be attributed to the rise in the number of active pits that are formed on the steel electrode surface with the harmonizing
raise in the total attacked area. The current density eventually attains the limiting value, $I_L$, which depends on the amount of the aggressive anions in the diluted oilfields-produced water, $C_{FW}$ % (Cl$^-$ and S ions). The attainment of the limiting current signifies that the pitting corrosion of the passive steel electrode takes place at the bottom of the pits formed, whereas their number stays constant [13,23–26]. The rise of the limiting current value upon increasing the $C_{FW}$ % (Cl$^-$ and S ions) implies a rise in the number of the initiated pits, that is, a rise in the anodic area [23]. Fig. 5 confirms the variance of the limiting-pitting corrosion current density with the $C_{FW}$ % (Cl$^-$ and SO$_4^{2-}$) on a double logarithmic scale, at 25°C.

## 3.2. Influence of inorganic passivators

Generally, some inorganic salts are utilized as inhibitors to control the induced localized pitting corrosion current of C-steel in oilfield-produced water. The effect of the inorganic passivators is characterized by their impact on each induction time and the produced corrosion current. Fig. 6 depicts the variance in the pitting corrosion current density ($\mu$A cm$^{-2}$) with the immersion time of C-steel in oilfield produced water (50% FW) containing different additions of $\lambda'$ is 0.71 log $\mu$A cm$^{-2}$ decade$^{-1}$. The constant $\zeta'$ symbolizes the logarithm of limiting current in case of the existence of 1.0% of aggressive ions. Early studies confirmed that Eq. (2) was deducted theoretically based on the competitive adsorption of the two counteracting anions on the surface of the investigated metal [18,27]. Theoretical derivation indicated that $\lambda'$ should refer to the value $\alpha_1 n_1 / z_1$, where $\alpha_1$ represents the transfer coefficient of the anodic reaction, $n_1$ is the number of electrons produced in the oxidation process while $z_1$ represents the valence of the destructive ions. Generally, $\alpha_1$ takes values between 0.7 and 0.3 for reactions controlled by transfer across a simple energy barrier. The variation in $\alpha_1$ values depends on the nature of the reaction that occurs on the electrode surface beside the kind and concentration of the attacking ions [18,27]. Such type of anodic reaction is controlled by a one-electron transfer [18,27]:

$$\text{Fe}^{2+} \rightarrow \text{Fe}^{3+} + e \quad (3)$$

where $n_1 = 1$ and $z_1$ for Cl$^-$ = 1 (and for SO$_4^{2-}$ or S$^2-$ = 2) the term $\alpha_1 n_1 / z_1$ supposes the values 0.3 as a minimum border and 0.7 as a maximum limit. In our study, the obtained value of 0.71 for the slope of the relationship is mostly in conformity with an anodic reaction of the steel electrode controlled by one-electron transfer [18,27]. The obtained value reaches the upper limit due to the sharing of SO$_4^{2-}$ or S$^2-$ as a divalent aggressive anion besides Cl$^-$ anions.

### 3.2. Influence of inorganic passivators

Generally, some inorganic salts are utilized as inhibitors to control the induced localized pitting corrosion current of C-steel in oilfield-produced water. The effect of the inorganic passivators is characterized by their impact on each induction time and the produced corrosion current. Fig. 6 depicts the variance in the pitting corrosion current density ($\mu$A cm$^{-2}$) with the immersion time of C-steel in oilfield produced water (50% FW) containing different additions...
of Na₂MoO₄ at 25°C. Comparable data were gained when Na₂HPO₄ and Na₂WO₄ are added (curves are not shown).

Screening of the data of Fig. 6, signalizes that the existence of inorganic passivators gives rise to elongating the induction time accompanied with the initiation of the localized pitting corrosion and suppressing the current accompanied the pit propagation. Such behavior explains that the added inorganic passivators act as effective inhibitors towards the localized pitting corrosion of the C-steel immersed in the oilfield-produced water. More addition of the inorganic salts decreased the pitting corrosion currents to reach the minimum value confirming the passivation of the initiated formed pits.

The data of Fig. 7 depicts the variance of the logarithm of the incubation time, τ, required for the initiation of pitting against the log $C_{inh}$. Straight-lines are gained satisfying the relation [13]:

$$\log \tau = \zeta'' - \lambda'' \log C_{inh}$$

where $\zeta''$ (minute unit) and $\lambda''$ (minutes/decade) are constants that depend on the solution composition and the metal under test.

Fig. 8 shows the variance of the logarithm of the limiting pitting corrosion current density, $I_L$, with the logarithm of the amount of the inorganic passivators, $C_{inh}$. Straight-line relationships are gained obeying the relation:

$$\log I_L = \gamma - \delta \log C_{inh}$$

where $\gamma$ and $\delta$ are constants that depend on the kind of the used inhibitor and the metal under test, Table 3. The constant $\gamma$ resembles the limiting current value, $I_L$, at $C_{inh}$ equal to a molar concentration. The obtained value $\gamma$ decreases in the sequence Na₂WO₄ > Na₂MoO₄ > Na₂HPO₄, which is congruent with the strength of the inhibition influence of the used salts against the initiation of pitting corrosion. It is clear that Na₂HPO₄ is the more effective inhibitor and Na₂WO₄ is the minimal.

The SEM micrograph of the C-steel samples after the period of the experiment when immersed in 50% diluted oilfield produce water containing 0.001 M Na₂HPO₄ is shown in Fig. 9, at 25°C. The micrograph indicated that the
existence of NaHPO$_4$ reduces the number of pits in comparison with that obtained with the oilfield-produced water (free from inhibitor), Fig. 3.

The amplitude of inhibition towards the pitting corrosion of C-steel by the added salt could be estimated from the values of the surface coverage, $\theta$, and the inhibition efficiency, $\eta$ %. The values of $\theta$ and $\eta$ % are deduced from the limiting current values in the inhibitor-free oilfield produced water, $I_{L,\text{free}}$, and in the existence of the inhibitor, $I_{L,\text{inh}}$, according to Eqs. (6) and (7) [13,20].

$$\theta = 1 - \frac{I_{L,\text{inh}}}{I_{L,\text{free}}}$$

(6)

$$\eta % = (\theta)100$$

(7)

The calculated values of $\theta$ and $\eta$ % at various amounts of the added salts are listed in Table 4. Screening of such table reveals the following:

- Each of $\theta$ and $\eta$ % values increases with increasing the concentration of the added salt.
- The inhibition efficiency $\eta$ % increase in the sequence: $\text{Na}_2\text{WO}_4 < \text{Na}_2\text{MoO}_4 < \text{NaHPO}_4$, which is the consequence of the reduction in the values of the $I_L$ values of Fig. 6 and similar ones.

### 3.3. Mechanism of inhibition

The presence of the inorganic salts decreases the localized pitting corrosion current density of C-steel in oilfield-produced water according to the order NaHPO$_4$ > Na$_2$MoO$_4$ > Na$_2$WO$_4$. This sequence is attributed to the rise in the ability of the corresponding anions to adsorb on the metal surface forming a protective layer that inhibits the localized pitting corrosion. HPO$_4^{2-}$ is the more effective anion to tolerate localized pitting corrosion. The inhibition influence of the HPO$_4^{2-}$ is of interest. Phosphate anions are rich with oxygen atoms and are theorized as oxygen-dependent inhibitors. HPO$_4^{2-}$ ions can diffuse to the metal surface consuming most of the oxygen in their oxidation [42]. From another point of view, the charge on the steel surface beside the protonation state of phosphate ions inside the solution facilitates the adsorption of phosphate on the C-steel surface. The adsorption of such anions may be incorporated into the passive film to make it efficient for passivity against the destructive effect of aggressive ions naturally present in the oilfield-produced water [37]. Another way of inhibition effect of the HPO$_4^{2-}$ anions could be attributed to the formation of a precipitate of FeHPO$_4$ on the C-steel surface that isolates the metal from the aggressive anions [38]. From another point of view, the existence of NaHPO$_4$ with higher concentrations behaves as an anodic inhibitor [28,38], that produces an insulating film on the C-steel surface formed from a mixture of $\gamma$-Fe$_2$O$_3$ and FePO$_4$·2H$_2$O [39,40].

The mechanism of corrosion inhibition of the C-steel by molybdate anions was discussed before [41,43]. It is reported that the inhibition influence could be attributed to the reduction of hexavalent molybdenum ions into trivalent ions with the formation of molybdenum dioxide (MoO$_3$) which is inserted into the passive layer formed on the surface of the C-steel electrode [44]. In addition, the anodic and cathodic reaction that occurs in neutral aerated or basic solution (pH $\geq$ 7) can be represented by the following equations:

$$\text{Fe} \rightarrow \text{Fe}^{2+} + 2e^-$$

(I)

$$\text{O}_2 + \text{H}_2\text{O} + 4e^- \rightarrow 4\text{OH}^-$$

(II)

The dissolved $\text{O}_2$ can react with surface Fe atoms according to the equation:

$$2\text{Fe} + \text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{FeOH}^+ + 2\text{OH}^-$$

(III)

The produced FeOH$^+$ ions are unstable that can further be transformed into relatively stable oxide films Fe$_2$O$_3$ [43]. The gradual accumulation of corrosion products on the C-steel surface forms a passive film that isolates the C-steel surface from the corrosive media and reduces the rate of pitting corrosion [45,46]. On the other hand, it cannot
be neglected that the MoO$_4^{2-}$ ions are adsorbed electrostatically on the electrode surface anions forms a protective film on the steel surface [47].

The inhibiting action of tungstate is discussed frequently [11–14]. It was suggested that the primary adsorption of the WO$_4^{2-}$ ions on the steel surface is accompanied by the formation of a passive iron tungstate, FeWO$_4$ that blocks the active sites on the metal surface [48] preventing pitting corrosion. However, the partial reduction of the WO$_4^{2-}$ ions [49,50] to give lower valence tungsten oxide is believed to be incorporated into the passive film. Other authors [51,52] assumed that tungstate ions oxidize ferrous ions to a higher valence ferric state that facilitated more passivity of steel. However, Abd El Kader et al. [53] argued that the inhibition by tungstate necessitates the simultaneous presence of O$_2$ in solution and the formation of an ordered arrangement of the two species on the steel surface

### 3.4. Adsorption isotherm

Generally, the corrosion inhibition of most inhibitor molecules is directly associated with an adsorption process [54]. Such an adsorption process could follow various models of adsorption isotherms, like Langmuir, Freundlich, Temkin, and Frumkin [55]. The surface coverage, θ can be deduced from the limiting current values, $I_L$, without and with the additions of inhibitors, as shown from Eq. (6) [18].

The obtained values of θ for all used salts confirmed the Langmuir model, Fig. 10. Such a model can follow the relation [56]:

$$\frac{\theta}{1-\theta} = K_{ads} C_{inh}$$

$$\frac{C}{\theta} = \frac{1}{K_{ads}} + C_{inh}$$

where $K_{ads}$ represents the adsorption–desorption equilibrium constant, and $C_{inh}$ is the equilibrium concentration of the added salts. Fig. 10 depicts straight lines relations between $C_{inh}$ vs. $C_{inh}$ for all the utilized salts with slopes very near to one, confirming correlation coefficients very near to one, Table 5. The adsorption–desorption equilibrium constants, $K_{ads}$ are deduced from the values of the adsorption–desorption equilibrium constant, $K_{ads}$ values may be considered as a measure of the power of the adsorption operation between the utilized inorganic anions and the C-steel surface [57]. The $K_{ads}$ are found to be 6,849; 5,444 and 3,891 mol$^{-1}$ for Na$_2$HPO$_4$, Na$_2$MoO$_4$, and Na$_2$WO$_4$, successively. Such a conclusion confirms that Na$_2$HPO$_4$ is more effective than Na$_2$MoO$_4$, followed by Na$_2$WO$_4$. The standard free energy of adsorption, $\Delta G_{ads}$, can be deduced from the $K_{ads}$ values of according to the relation [58]

$$K_{ads} = \frac{1}{55.5} \exp \left(\frac{-\Delta G_{ads}}{RT}\right)$$

where 55.5 represents the molar concentration of water, $R$ is the universal gas constant and $T$ is the temperature in the kelvin scale. The calculated values of $\Delta G_{ads}$ are $-31.83$, $-31.26$, and $-30.44$ kJ mol$^{-1}$ for Na$_2$HPO$_4$, Na$_2$MoO$_4$, and Na$_2$WO$_4$, successively. This is proportionate with the higher inhibition efficacy of Na$_2$HPO$_4$ followed by Na$_2$MoO$_4$, and Na$_2$WO$_4$. The negative signs of $\Delta G_{ads}$ confirm the spontaneity of the adsorption process on the C-steel surface with the constancy of the adsorbed film on the C-steel surface. The obtained values of $\Delta G_{ads}$ are consistent with a chemisorption process due to chemical bond formation between the used anions and the metal surface [59–61].

### 3.5. Effect of temperature

The influence of temperature on the pitting corrosion current of oilfield-produced water was investigated without and with the various additions of inorganic salts. The curves of Fig. 11 represent the $I$-$t$ curves for C-steel in 50% diluted oilfield-produced water at various temperatures. On the other hand, Fig. 12 depicts the $I$-$t$ curves for C-steel in 50% diluted oilfield-produced water containing $1 \times 10^{-4}$ M Na$_2$WO$_4$ at various temperatures. Similar curves are gained with other used salts Na$_2$MoO$_4$, and Na$_2$HPO$_4$ (curves not shown). The induction period in all examined solutions (without and with inhibitor) preceding the pitting corrosion process decreases with temperature followed by an increase in the pitting corrosion current till reaching a constant value, $I_c$, due to propagation of

<table>
<thead>
<tr>
<th>Type of salt</th>
<th>$K_{ads}$, mol$^{-1}$</th>
<th>$-\Delta G_{ads}$, kJ mol$^{-1}$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na$_2$HPO$_4$</td>
<td>6,849</td>
<td>31.83</td>
<td>0.998</td>
</tr>
<tr>
<td>Na$_2$MoO$_4$</td>
<td>5,444</td>
<td>31.26</td>
<td>0.999</td>
</tr>
<tr>
<td>Na$_2$WO$_4$</td>
<td>3,891</td>
<td>30.44</td>
<td>0.997</td>
</tr>
</tbody>
</table>

Fig. 10. Langmuir adsorption isotherm, $C_{inh}/\theta$ vs. $C_{inh}$ for all used salts.
pitting corrosion [34]. Such behavior could be attributed to the increase in the rate of corrosion of C-steel with temperature due to the rise in the rate of penetration of the corrosive species (Cl\(^{-}\) and S ions) to the passive oxide film with temperature [62]. The Limiting pitting corrosion current density, \(I_L\), of C-steel in oilfield-produced water devoid of and containing \(1 \times 10^{-4}\) M of different inorganic salts is plotted against the temperature in Kelvin values, Fig. 13. The data of this figure confirms that the \(I_L\) is increased with the temperature in the case of aggressive and inhibitive solutions.

The data of the curves of Figs. 11–13 reveals that:

- The rise in the temperature reduces the induction period, \(\tau\), and increases the limiting corrosion current, \(I_L\).
- In the aggressive and inhibitive solutions, the increase in the temperature rise the pitting corrosion current density due to the enhancement of the oxidation process (Fe \(\rightarrow\) Fe\(^{3+}\) + 2e\(^{-}\)).
- Various additives of Na\(_2\)HPO\(_4\), Na\(_2\)MoO\(_4\), and Na\(_2\)WO\(_4\) reduce the pitting corrosion current and \(I_L\) values although the rise in the pitting corrosion current with temperature.
- The inorganic salts reduce the \(I_L\) values following the sequence: Na\(_2\)HPO\(_4\) > Na\(_2\)MoO\(_4\) > Na\(_2\)WO\(_4\).

Broadly, increasing the temperature for pitting corrosion enhances the initiation of pitting corrosion by reducing the induction period, \(\tau\), required pit formation, and enhancing the pit propagation by increasing the limiting pitting corrosion current density, \(I_L\). The Arrhenius equation can be used to deduce the activation energy, \(E_a\). Such an equation can be represented by the following relation [63]:

\[
\log I_L = \log A - \frac{E_a}{2.303RT}
\]  

where the rate of reaction can be represented by \(I_L\) values, \(E_a\) is the activation energy of the pitting corrosion process, \(A\) is the Arrhenius constant, \(R\) is the ideal gas constant and \(T\) is the absolute temperature.

Fig. 14 depicts the variation of the logarithm of \(I_L\) vs. \(1/T\) for 50\% FW without and with various inorganic inhibitors (\(1 \times 10^{-4}\) M). Straight-line relations are obtained confirming the Arrhenius equation. The slope of each line can be used to deduce the activation energy, \(E_a\), required for pitting corrosion for all investigated electrolytes, Table 6 and correlation
coefficient, $R^2$, and correlation coefficient, $R^2$. The calculated value of $E_a$ for the corrosion of C-steel in inhibitor-free oilfield-produced water is less than that obtained in presence of the added salts. Such behavior could be attributed to the increase in the energy barrier of the inhibitor being higher than those in the oilfield-produced water free of any additives. These anions tolerate the pitting corrosion of C-steel in oilfield-produced water due to competing with the aggressive anions (Cl$^-$ and S) enhancing the oxide film repair and lowering the oxide film destruction [64]. The computed $E_a$ accompanying the pitting corrosion increases in the order: 50% FW $<$ Na$_2$MoO$_4$ $<$ Na$_2$WO$_4$ $<$ Na$_2$HPO$_4$. These additives were utilized as effective inhibitors towards the pitting corrosion of steel in calcium hydroxide [65] and zinc in sodium sulfate solutions [17].

Other thermodynamic parameters, entropy ($\Delta S^*$) and enthalpy ($\Delta H^*$) can be deduced when using transition state equation [66,67]:

$$\log \frac{I_{pit}}{T} = \log N_A - \frac{\Delta S^*}{2.303R} - \frac{\Delta H^*}{2.303RT}$$  \hspace{1cm} (13)

where $N$ is the Avogadro’s number, $R$ is the universal gas constant, and $h$ is the Plank’s constant. Plotting of $\log I_{pit}/T$ vs. $1/T$ for 50% FW without and with the various inorganic inhibitors (1 $\times$ 10$^{-4}$ M) gives straight-line relations, Fig. 15. The slope of the gained plots represents the value of $\Delta H^*/2.303R$ while the intercept is equal to $(\Delta S^*/2.303R)+\log R/N_A$. The values of $\Delta H^*$ and $\Delta S^*$ are computed and tabulated in, Table 6. The activation adsorption entropy $\Delta S^*$ is determined to be $-144.75$ J mol K$^{-1}$ for the inhibitor-free 50% FW and are $-177.2$, $-168.6$, and $-164$ J mol K$^{-1}$ for the inhibitive solution Na$_2$MoO$_4$, Na$_2$WO$_4$, Na$_2$HPO$_4$, respectively. The negative values of $\Delta S^*$ in the case of inhibitor-free oilfield-produced water and with the inorganic salts confirm that the activated complex in the rate-determining step represents an association rather than dissociation. Such attitude is due to the reduction in the disordering that takes place [67–69]. The positive values of $\Delta H^*$ confirm that the transition state (the activated complex) is an endothermic process.

### 4. Conclusions

The pitting corrosion current density for C-steel in the oilfield-produced water without and with the additions of Na$_2$MoO$_4$, Na$_2$WO$_4$, and Na$_2$HPO$_4$, was followed with the immersion time. The obtained data indicated that:

- The existence of an induction period preceding the pit formation. The increase in the corrosion currents depends

### Table 6

<table>
<thead>
<tr>
<th>Type of solution</th>
<th>$E_a^*$ kJ mol$^{-1}$</th>
<th>$\Delta H^*$ kJ mol$^{-1}$</th>
<th>$-\Delta S^*$ J mol$^{-1}$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>50% FW</td>
<td>23.51</td>
<td>21.40</td>
<td>169</td>
<td>0.992</td>
</tr>
<tr>
<td>50% FW + Na$_2$WO$_4$</td>
<td>27.98</td>
<td>25.40</td>
<td>165</td>
<td>0.997</td>
</tr>
<tr>
<td>50% FW + Na$_2$MoO$_4$</td>
<td>30.55</td>
<td>28.00</td>
<td>162</td>
<td>0.999</td>
</tr>
<tr>
<td>50% FW + Na$_2$HPO$_4$</td>
<td>32.50</td>
<td>30.16</td>
<td>159</td>
<td>0.998</td>
</tr>
</tbody>
</table>

Fig. 14. The Arrhenious plots, variation of the logarithm of $I_{pit}$ vs. $1/T$ for 50% FW without and with 0.0001 M of the various inorganic salts.

Fig. 15. Transition state plots, variation of the logarithm of $I_{pit}/T$ vs. $1/T$ for 50% FW without and with 0.0001 M of the various inorganic salts.
on the percent of dilution of the oilfield-produced water, the concentration of the inhibitive anions, as well as, the temperature of the solution.

- The limiting pitting corrosion current density, \( I_p \), decreases with the amount of the added inhibitive anions following the equation: \( \log I_p = \zeta + \lambda' \log C_{inh} \), where \( \zeta' \) and \( \lambda' \) are constants.
- The induction period, \( \tau \), required for the initiation of pitting increases with increasing the amount of the added salts, \( C_{inh} \).
- The inhibition efficiency is reduced in the order: \( \text{Na}_2\text{HPO}_4 > \text{Na}_2\text{MoO}_4 > \text{Na}_2\text{WO}_4 \).
- The activation energy accompanied by the pitting corrosion is increased in the order: \( \text{Na}_2\text{WO}_4 < \text{Na}_2\text{MoO}_4 < \text{Na}_2\text{HPO}_4 \).

### Disclosure statement

The authors declare no potential conflict of interest in preparing this article.

### References


