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# Identification of interaction, surface species and anticorrosion potency for adsorption of myrrh extract on 330 stainless steel in 1 M HCl solution

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Steel and its alloys are essential materials in various industries, but they quickly react with the surrounding environment, especially humid ones, which leads to their destruction and corrosion on a large scale. Replacing synthetic organic inhibitors with plant-extracted or natural molecules has recently been a major challenge in terms of environmental and industrial aspects. In this report, the inhibitory dominance, adsorption of myrrh extract (MRE) as a green corrosion inhibitor on 330 stainless steel (SS) in 1.0 M HCl solution at the temperature range of 25 to 55 °C were explored. Weight loss (WL) measurements showed apparent inhibition efficiency (% IE) of MRE of up to 93.6% at 55 °C. The % IE values increased sharply with increasing inhibitor concentration and medium temperature. Adsorption investigations confirmed that the extract molecules were strongly adsorbed on the SS surface and were consistent with the Langmuir and Frumkin models. Polarization monitoring showed that the MRE behaves as a mixed-type inhibitor. EIS was also used to quantitatively evaluate the efficiency of the passive layer on the SS surface. All thermodynamic ( $\Delta H$ ,  $\Delta S$ ,  $\Delta G$ ) and kinetic ( $K_{ads}$ ,  $E_a$ ) parameters were calculated and analyzed. XPS was used to inspect the surface chemistry of SS and the nature of MRE-SS bonds. AFM demonstrated that the SS sample in the blank solution appears to be severely damaged and has the highest roughness of about 382.6 nm, compared to that obtained in the presence of MRE (145.61 nm). Various theoretical and computational methods were used to predict the performance of the examined inhibitor. All experimental findings of the different techniques are in excellent agreement with each other and with theoretical speculations. The outcomes of this study are thought to have some bearing on the sensible development of a potent inhibitor against the corrosion of metals in acidic environments.

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## 1 Introduction

Corrosion is a natural phenomenon that significantly affects the structural integrity of concrete and steel structures.<sup>1–8</sup> When infrastructures in sectors like petrochemicals and petroleum, which are vital to the world economy, are impacted, this becomes a problem.<sup>9</sup> The “cancer of metals” is a term used to describe metal corrosion,<sup>10</sup> which always leads to financial losses and security risks.<sup>11</sup> Implementing mitigation techniques has grown more crucial due to the significant impact of metal infrastructure loss due to corrosion during the past few decades. Cathodic protection,<sup>11–13</sup> anticorrosion coatings,<sup>14</sup> electroplating, and inhibitors<sup>15–20</sup> are a few methods of preventing corrosion. Because of their cheap operating costs and wide variety of applications, corrosion inhibitors have become

the most practical and efficient way among the available approaches.<sup>21–27</sup> Nevertheless, conventional commercially used corrosion inhibitors including chromates, phosphates, and nitrates are hazardous to the environment and their surroundings.<sup>28</sup> Therefore, conventional corrosion inhibitors do not meet current environmental protection standards.<sup>29</sup> In order to overcome these obstacles, research into corrosion inhibitors has shifted in recent years towards the development of green corrosion inhibitors.<sup>30–36</sup> Researchers are searching for more ecologically benign compounds, also referred to as natural or green inhibitors, which include pharmaceuticals<sup>31–36</sup> and plant extracts,<sup>37</sup> are thought to be safe materials free of heavy metals.<sup>38</sup> Since the plant extracts are usually sourced sustainably and are less harmful to the environment and human health than conventional corrosion inhibitors, they are referred to as “green” extracts. These consist of amino acids, alkaloids, polyphenols, and frequently widely disseminated plant extracts with little economic value, such as agricultural waste and by-products of agro-industrial operations. Recent studies have shown that extracts of the *Cuminum cyminum* extract are effective in protecting metals/alloys in acidic

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Table 1 Chemical ingredients of MRE<sup>40,44</sup>

Compounds	Structure	Formula	Percentage (%)
(1) Cyclohexene, 4-ethenyl-4-methyl-3-(1-methylethenyl)-1-(1-methylethyl)-, (3- <i>R-trans</i> )-		C <sub>15</sub> H <sub>24</sub>	0.3342
(2) Longifolene-(V4)		C <sub>15</sub> H <sub>24</sub>	4.0480
(3) Caryophyllene		C <sub>15</sub> H <sub>24</sub>	0.2976
(4) 1,4-Methanoazulene, decahydro-4,8,8-trimethyl-9-methylene-, [1 <i>S</i> -(1 <i>à</i> ,3 <i>áá</i> , 4 <i>à</i> ,8 <i>áá</i> )] <sup>-</sup>		C <sub>15</sub> H <sub>24</sub>	1.047
(5) 1,6-Cyclodecadiene, 1-methyl-5-methylene-8-(1-methylethyl)-, [ <i>s</i> -( <i>E,E</i> )] <sup>-</sup>		C <sub>15</sub> H <sub>24</sub>	0.3498
(6) Naphthalene, 1,2,3,5,6,7,8,8 <i>a</i> -octahydro-1, 8 <i>a</i> -dimethyl-7-(1-methylethenyl)-, [1 <i>S</i> -(1 <i>à</i> ,7 <i>à</i> ,8 <i>áá</i> )] <sup>-</sup>		C <sub>15</sub> H <sub>24</sub>	0.3874
(7) Benzofuran, 6-ethenyl-4,5,6,7-tetrahydro-3,6-dimethyl-5-isopropenyl-, <i>trans</i> -		C <sub>15</sub> H <sub>20</sub> O	26.630
(8) 1 <i>H</i> -Cycloprop[ <i>e</i> ]azulene, decahydro-1, 1,7-trimethyl-4-methylene-, [1 <i>aR</i> -(1 <i>áá</i> ,4 <i>áá</i> ,7 <i>à</i> ,7 <i>áá</i> ,7 <i>bà</i> )]		C <sub>15</sub> H <sub>24</sub>	0.5777
(9) Meso-hydrobenzoin		C <sub>14</sub> H <sub>14</sub> O <sub>2</sub>	0.3437



Table 1 (Contd.)

Compounds	Structure	Formula	Percentage (%)
(10) 9(1 <i>H</i> )-Phenanthrenone, 2,3,4,4 <i>a</i> ,10,10 <i>a</i> -hexahydro-4 <i>a</i> -methyl-, <i>cis</i> -		C <sub>15</sub> H <sub>18</sub> O	49.410
(11) Benzenemethanol, 3-methoxy- <i>a</i> -phenyl-		C <sub>14</sub> H <sub>14</sub> O <sub>2</sub>	13.790
(12) Cyclohexanemethanol, 4-ethenyl- $\dot{\alpha},\dot{\alpha}$ , 4-trimethyl-3-(1-methylethenyl)-, [1 <i>R</i> -(1 $\dot{\alpha},3\dot{\alpha},4\dot{\alpha}$ )]-		C <sub>15</sub> H <sub>26</sub> O	0.6306
(13) Testosterone		C <sub>19</sub> H <sub>28</sub> O <sub>2</sub>	2.154

environments.<sup>39</sup> Noteworthy those flavonoids, alkaloids, polyphenols and others contain electron-rich atoms such as S, O, N and halogens as well as double bonds and aromatic rings, in their structures, which are frequently prevalent in plant extracts. These electron-rich sites allow them to be strongly adsorbed on the metal surface by donating lone pairs of electrons from heteroatoms and  $\pi$ -electrons of rings to the metal, preventing corrosive agents from reaching the metal surface and hindering the corrosion process, thus protecting it.

It's worth noting that myrrh trees are readily available in many countries, are inexpensive, and are abundant in production. In particular, MRE consists of various chemical compounds (listed in Table 1) that have been considered essential oils.<sup>40,41</sup> Interestingly, the percentage of this plant's extract reaches 15%. In fact, one kg of myrrh costs about 5–10 \$, yielding 150 grams of extract, which is much less expensive (about one-tenth) than many other green inhibitors such as polysaccharide,<sup>42</sup> and *Ceratonia siliqua* L. seed oil.<sup>43</sup> Since 1 kg of myrrh raw material yields about 150 grams of extract which is enough to prepare 600 liters of 250 ppm MRE inhibitor. From an industrial perspective, its use as an anti-corrosion agent for metals/alloys is of economic and industrial value, given its effectiveness and low cost. The main challenge is its environmental compliance, as it is a low-toxic, biodegradable, and sustainable alternative. In contrast, traditional inhibitors often pose environmental and health risks due to their high toxicity.

In this work a new green corrosion inhibitor extracted from myrrh plant was developed and applied to protect 330 SS in aqueous HCl for the first time at a temperature range of (25–55 °C). A number of theoretical and computational methods have been used to predict the effect and performance of this inhibitor as well.

## 2 Experimental

### 2.1 Materials

330 SS samples were used for all the corrosion measurements of this study. The chemical composition of 330 SS (% wt) is: Fe = 42, Ni = 36, Cr = 19, Mn = 2, Si = 1.13, C = 0.080, P = 0.040, and S = 0.030. Analytical reagent-grade HCl was diluted with deionized water to prepare 1.0 M HCl (blank) as a corrosive electrolyte stock. The process of creating a myrrh-based inhibitor (MRE) by extracting myrrh in methanol directly from its powder as explained in detail earlier.<sup>44</sup> In short, after the myrrh gum was completely dried, it was ground well into a fine powder. The powder was then dissolved in methanol and the solution was left for five days, filtering the solution, and then removing the methanol through distillation at 40 °C. The remaining concentrated material is then dried, making MRE ready for use in the corrosion experiments. Table 1 shows the main MRE components with their structures and proportions.<sup>40,44</sup> To prepare the stock solution of the inhibitor, 3.0 g of freshly prepared MRE was dissolved in 1 L of distilled water and



Table 2 Corrosion data obtained from WL measurements at different temperatures after 24 hours

Inh. conc. (ppm)	$\Delta w$ ( $\text{mg cm}^{-2}$ )	CR ( $\text{mg cm}^{-2} \text{min}^{-1}$ )	$\theta$	% IE	$\Delta w$ ( $\text{mg cm}^{-2}$ )	CR ( $\text{mg cm}^{-2} \text{min}^{-1}$ )	$\theta$	% IE
25 °C					35 °C			
Blank	43.61	2.423	—	—	44.19	2.455	—	—
50	23.11	1.283	0.470	47.0	19.00	1.056	0.570	57.0
100	15.77	0.876	0.638	63.8	13.44	0.747	0.696	69.6
150	12.84	0.713	0.706	70.6	11.80	0.656	0.733	73.3
200	11.41	0.634	0.739	73.9	9.41	0.523	0.787	78.7
250	9.77	0.542	0.776	77.6	7.09	0.394	0.840	84.0
45 °C					55 °C			
Blank	44.91	2.495	—	—	45.77	2.54	—	—
50	15.79	0.877	0.648	64.8	10.39	0.577	0.773	77.3
100	11.08	0.615	0.753	75.3	8.91	0.495	0.805	80.5
150	10.32	0.573	0.770	77.0	8.61	0.478	0.812	81.2
200	8.14	0.452	0.819	81.9	6.49	0.361	0.858	85.8
250	5.08	0.282	0.887	88.7	2.95	0.164	0.936	93.6

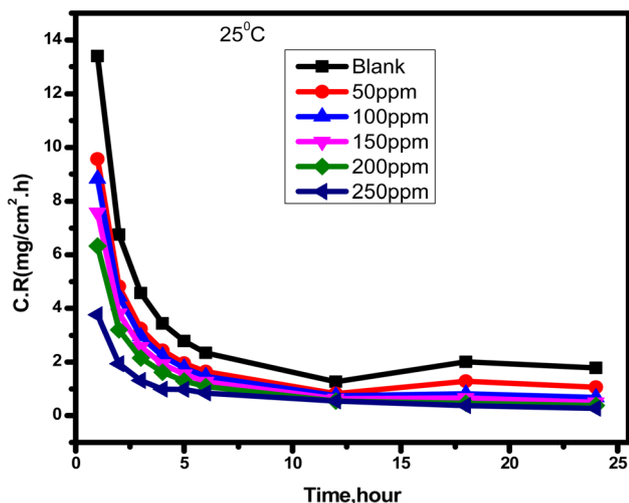


Fig. 1 Variation of CR of SS with time in 1.0 M HCl solution and in the presence of different concentrations of MRE at 25 °C.

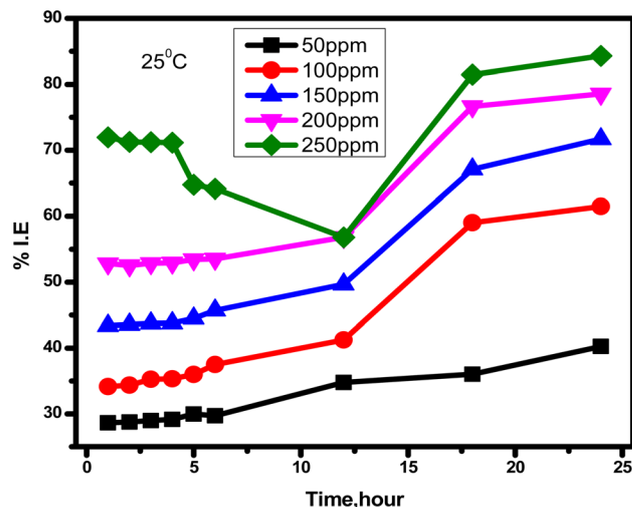


Fig. 2 Variation in % IEs of different concentrations of MRE in the corrosion of SS in 1.0 M HCl solution at 25 °C.

thus a concentration of 3000 ppm was ready. To perform inhibition tests, different concentrations of MRE (50–250 ppm) were prepared by dilution with distilled water.

Various experimental and computational techniques were used in this work. The experimental techniques are weight loss (WL), potentiodynamic polarization (PDP), open-circuit potential (OCP), electrochemical frequency modulation (EFM), linear sweep voltammetry (LSV) and electrochemical impedance spectroscopy (EIS). For surface analyses, atomic force microscopy (AFM) and X-ray photoelectron spectroscopy (XPS) were used. The computational prediction techniques are molecular dynamics (MD) simulations, density functional theory (DFT) calculations, molecular electrostatic potential (MESP), Mulliken atomic charges (MAC), and Fukui indices (FI).

## 2.2 Experimental techniques

**2.2.1 Chemical measurements.** Because the size homogeneity is essential for reducing experimental result variability, especially in surface-dependent investigations like corrosion inhibition. For WL tests, SS sheets with dimensions of 1 mm × 2 cm × 2 cm were used and therefore have the same exposure area (4 cm<sup>2</sup>) to the corrosive environment. The SS specimens were successively polished using several grades of sandpaper in order to provide a smooth and uniform surface. After polishing, they were washed several times with distilled water and then acetone to remove any oils, grease, and organics that may be present on their surface. Finally, the cleaned SS samples were air-dried to remove any moisture that would accelerate deterioration. The samples were kept dry in a desiccator to protect



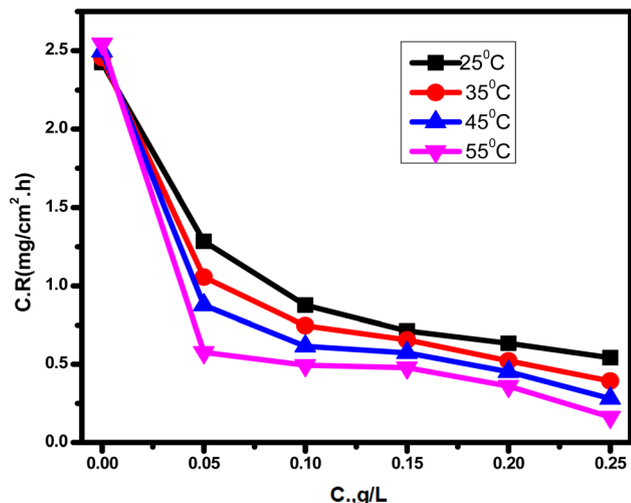


Fig. 3 Variation of CR of SS with MRE concentration in 1.0 M HCl solution at different temperatures.

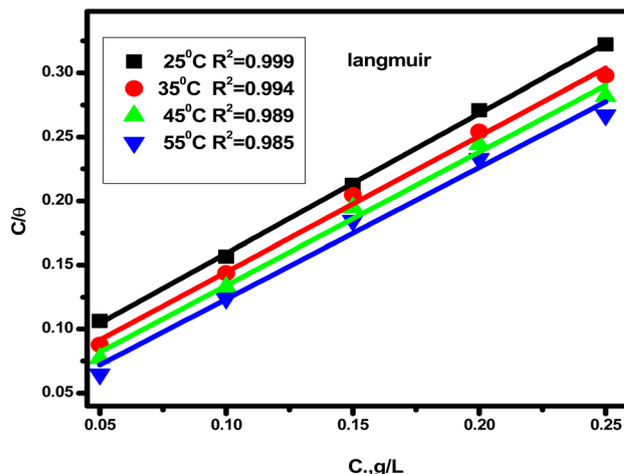


Fig. 5 Langmuir isotherm for adsorption of MRE molecules onto a SS surface in 1.0 M HCl solution at different temperatures.

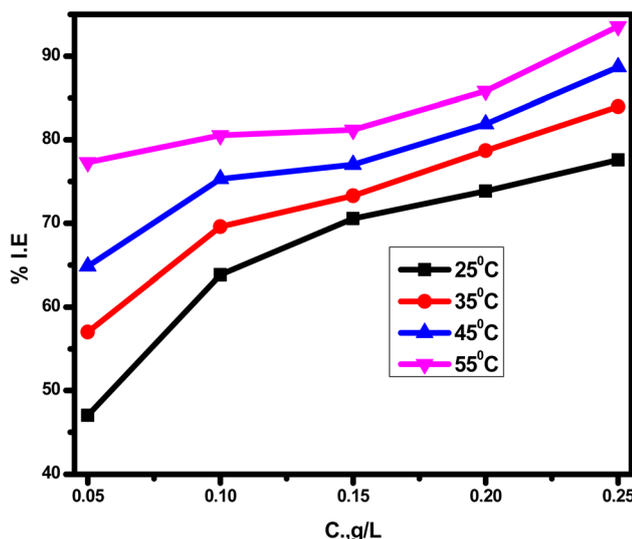


Fig. 4 Variation of % IE of MRE with its concentration in 1.0 M HCl solution at different temperatures.

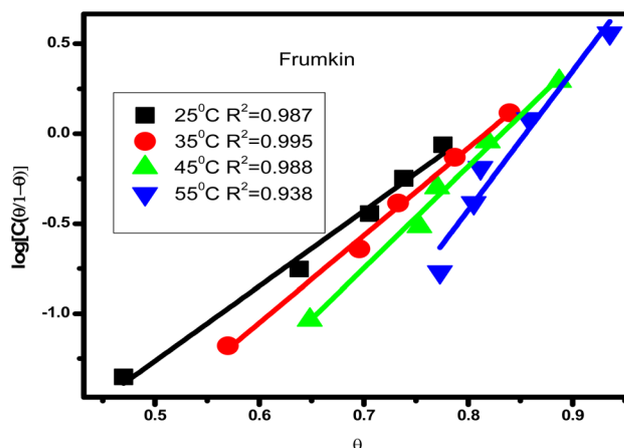


Fig. 6 Frumkin isotherm for adsorption of MRE molecules onto a SS surface in 1.0 M HCl solution at different temperatures.

them from moisture and any other potential contaminants to ensure they remained in the best possible condition and ready for accurate and reproducible investigations in corrosion experiments. The values of corrosion rates (CR) for SS in the absence and presence of MRE were calculated by:

$$CR = \Delta w/A \times t \quad (1)$$

where,  $t$  is the immersion period (min),  $A$  is the specimen's exposed surface area ( $\text{cm}^2$ ), and  $\Delta w$  is the mass loss ( $\text{mg cm}^{-2}$ ). The IE (%), and the surface coverage ( $\theta$ ) were estimated by:<sup>45</sup>

$$\% \text{ IE} = [1 - (CR_{\text{inh}}/CR_{\text{uninh}})] \times 100 = \theta \times 100 \quad (2)$$

where,  $CR_{\text{uninh}}$  and  $CR_{\text{inh}}$  are the CR of SS in the uninhibited and inhibited media, respectively.

**2.2.2 Electrochemical measurements (ECMs).** For conducting all ECM, a Gamry instrument (PCI 300/4) potentiostat/galvanostat/ZRA was employed. This device has a Gamry framework system with ESA 400. For experimental running, the device is computer-controlled using three data collection software based on the behavioral ECM method: DC105 software for PDP, EFM140 software for EFM, and EIS300 software for EIS measurements. For plotting and fitting data, Echem Analyst 6.03 software was used. A traditional three-electrode cell was used: a saturated calomel electrode (SCE) as the reference electrode, a Pt gauze as the counter electrode and a square SS sheet ( $1.0 \text{ cm}^2$ ) as the working electrode (WE). The chamber was set up to maintain a consistent and measured temperature throughout each test. Before performing ECMs, the OCP is first



Table 3 Parameters obtained from Frumkin's and Langmuir's study

Langmuir						
Temp. (°C)	Log°K <sub>ads</sub>	R <sup>2</sup>	−ΔG <sub>ads</sub> (kJ mol <sup>−1</sup> )	ΔH <sub>ads</sub> (kJ mol <sup>−1</sup> )	ΔS <sub>ads</sub> (J mol <sup>−1</sup> K <sup>−1</sup> )	
25	1.30	0.999	17.34	24.07	138.9	
35	1.41	0.994	18.60		138.5	
45	1.53	0.989	19.93		138.4	
55	1.69	0.985	21.57		139.1	
Frumkin						
Temp. (°C)	Log K <sub>ads</sub>	R <sup>2</sup>	α	−ΔG <sub>ads</sub> (kJ mol <sup>−1</sup> )	ΔH <sub>ads</sub> (kJ mol <sup>−1</sup> )	−ΔS <sub>ads</sub> (J mol <sup>−1</sup> K <sup>−1</sup> )
25	3.35	0.987	4.80	29.09	101.64	438.6
35	3.97	0.995	5.60	33.68		439.4
45	4.71	0.988	6.52	39.30		443.5
55	4.91	0.938	8.90	41.78		437.0

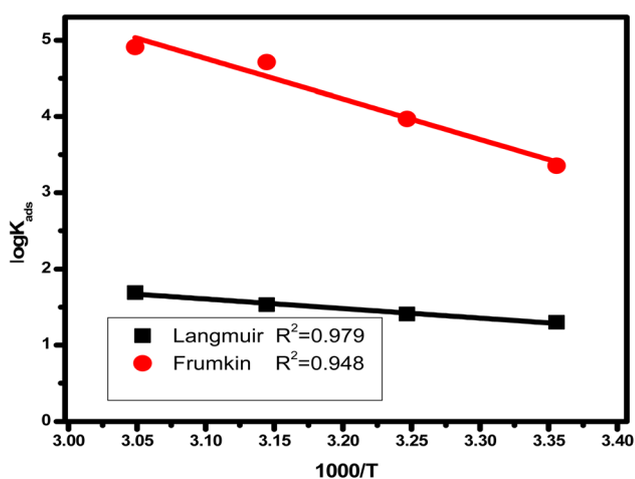


Fig. 7 van't Hoff plots for adsorption of MRE molecules onto SS surface in 1.0 M HCl solution.

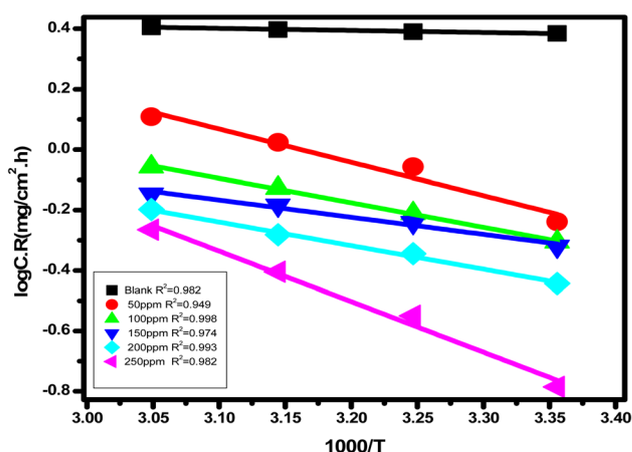


Fig. 8 Arrhenius plots for adsorption of MRE molecules onto SS surface in 1.0 M HCl solution at different MRE concentrations.

measured until it stabilizes over a period of 1800 s, as it serves as the starting point for the potential scan.

**2.2.2.1 OCP.** The change of OCP or  $E_{OC}$  vs. immersion time of SS in 1.0 M HCl solution without and with different concentrations of MRE at 25 °C over a period of 1800 seconds was examined.

**2.2.2.2 PDP.** In polarization tests, a voltage of −1.5 to 1.5 V is applied to the WE at a sweep rate of 0.1 mV<sup>−1</sup> with a step increase of ±250 mV. All experiments were performed 3 times in freshly prepared solutions to check reproducibility. The % IE, and  $\theta$  were calculated by:<sup>46</sup>

$$\% \text{ IE} = [(1 - i_{\text{corr(inh)}})/i_{\text{corr(uninh)}}] \times 100 = \theta \times 100 \quad (3)$$

Current density ( $i_{\text{corr}}$ ) was determined by extrapolating the Tafel slopes ( $\beta_a, \beta_c$ ) in the absence and presence of the inhibitor and was denoted by  $i_{\text{corr(uninh)}}$  and  $i_{\text{corr(inh)}}$ , respectively. The Tafel slopes were determined by extrapolating the linear region

Table 4 Activation parameters obtained from Arrhenius's and transition state plots in the corrosion of SS in 1.0 M HCl solution in the presence of different MRE concentration

Inh. conc. ppm	$E_a^*$ (kJ mol <sup>−1</sup> )	−ΔH* (kJ mol <sup>−1</sup> )	−ΔS* (J mol <sup>−1</sup> K <sup>−1</sup> )	LogA
Blank	1.30	1.15	170.32	0.61
50	10.87	13.50	185.84	1.59
100	14.93	17.53	190.74	2.18
150	15.52	18.12	192.22	2.42
200	24.96	23.69	195.34	3.48
250	32.02	34.61	206.52	4.85

of the anodic and cathodic curves in a potential range of ≈ ±50–100 mV relative to  $E_{\text{corr}}$ .

**2.2.2.3 EIS.** An identical cell configuration was used for EIS measurements via the ESA400 Gamry framework system. AC



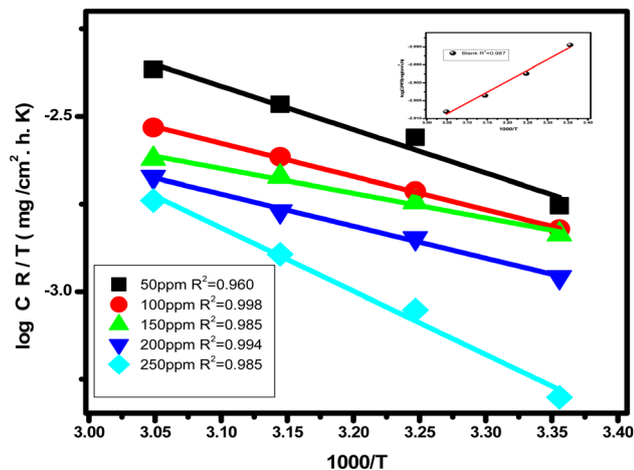


Fig. 9 Transition state plots for adsorption of MRE molecules onto SS surface in 1.0 M HCl solution at different MRE concentrations.

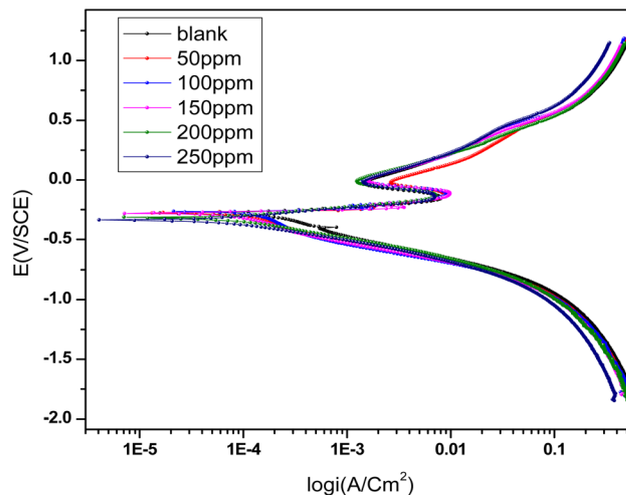


Fig. 11 PDP plots for SS in 1.0 M HCl solution and in the presence of different concentrations of MRE at 25 °C.

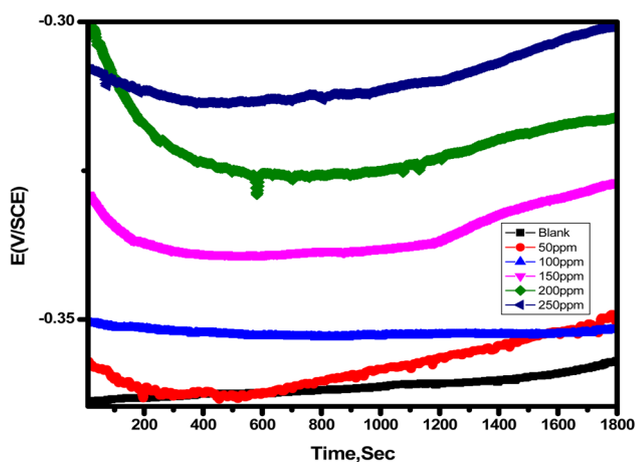


Fig. 10 OCP (potential vs. time) plots for SS in 1.0 M HCl solution and in the presence of different concentrations of MRE at 25 °C.

signals with a peak-to-peak amplitude of 5.0 mV were applied at the OCP over a frequency range of 100 kHz to 0.5 Hz. The Gamry Echem Analyst program was utilized to match all impedance data to the relevant equivalent circuit.

**2.2.2.4 EFM.** EFM tests were conducted using two sine waves at frequencies of 2.0 and 5.0 Hz and a potential disturbance signal of 10.0 mV amplitude. Three factors were taken into account when choosing the frequencies of 2.0 and 5.0 Hz:<sup>46</sup> Tafel slopes (cathodic =  $\beta_c$  and anodic =  $\beta_a$ ),  $i_{corr}$ , and causation factors CF2 and CF3 were computed using the largest peaks.<sup>47</sup>

**2.2.3 Surface analyses.** SS samples were immersed in blank solution and in the presence of MRE inhibitor (250 ppm) for 1 day. Afterwards, the samples were taken out, rinsed thoroughly with deionized water, dried, and then ready for surface examination.

**2.2.3.1 AFM.** Using AFM analyses, it is possible to envisage the effect of corrosion and inhibition on SS surfaces. It was

conducted with a  $\text{Si}_3\text{N}_4$  probe (MLCT model; Bruker) in contact mode.

**2.2.3.2 XPS.** XPS is a powerful tool for screening and identifying the species present on the SS surface, which greatly helps in better understanding the inhibition mechanism and interactions. Thus, the nature of the chemical bond on its surface was systematically determined. It was performed with a K-alpha model instrument from Thermo Fisher Scientific, USA, using monochromatic Al K X-alpha radiation with an energy range of -10 to 1350 eV under pressures of  $10^{-9}$  mbar, with a spot size of 400  $\mu\text{m}$ , with full spectrum pass energy of 200 eV and a narrow spectrum pass energy of 50 eV.

## 2.3 Computational investigations

**2.3.1 DFT calculations.** Using DFT calculations derived from quantum chemical theories, possible inhibition mechanisms can be mathematically elucidated. It was implemented by the Dmol<sup>3</sup> module within Material Studio (version 7.0) to examine the full optimization of the geometry of the investigated MRE inhibitor. Undoubtedly, the electronic configurations and molecular structure of the extract reflect its effectiveness and behavior in the inhibition strategy. Using DFT computation, the quantitative parameters estimated are HOMO and LUMO molecular orbitals, dipole moment ( $\mu$ ), electronegativity ( $\chi$ ), absolute hardness ( $\eta$ ), softness ( $\sigma$ ), and electron affinity index ( $\omega$ ).<sup>48</sup>

**2.3.2 MD simulation.** MD simulations were used to theoretically investigate the possible interactions between MRE molecules and the SS surface. A Fe(110) surface with dimensions of (32.27 Å, 32.27 Å, 50.18 Å) consisting of 6 atomic layers within a  $10 \times 10$  supercell featuring a 25 Å vacuum along the C-axis, within an HCl medium, was used. In comparison to other Fe surfaces, Fe(110) has the highest stabilisation and is a density packed surface. These experimental parameters were consistent with earlier studies.<sup>49</sup> The inhibitor molecules were



Table 5 Parameters obtained from PDP measurements (Fig. 11) at 25 °C

Inh. conc. (ppm)	$-E_{\text{corr}}$ (mV)	$R_p$ (ohm)	$i_{\text{corr}}$ ( $\mu\text{A cm}^{-2}$ )	$\beta_a$ (mV dec $^{-1}$ )	$-\beta_c$ (mV dec $^{-1}$ )	$\theta$	% IE
Blank	279.0	9.3	2499.0	58.2	241.0	—	—
50	275.3	15.7	1414.0	49.0	233.3	0.121	12.1
100	266.0	17.5	1023.2	49.8	263.4	0.299	29.9
150	280.6	41.1	633.3	50.8	205.0	0.523	52.3
200	315.3	46.9	546.4	77.7	150.3	0.655	65.5
250	336.2	53.5	487.2	85.3	143.3	0.805	80.5

modeled and optimized through the Forcite software, after which the adsorption energy was calculated using the adsorption locator module. The simulation environment was maintained at a constant 25 °C using a Brandsen thermostat inside the NVT assembly, with a time step of 1.0 fs and a total simulation period of 300 ps under the COMPASS force field.<sup>50</sup>

**2.3.3 MESP.** MESP is an important computational method for accurately predicting and interpreting various aspects of interaction. Electrostatic potentials are depicted by color gradients at different locations on the electron density surface.

**2.3.4 MAC and FI.** The potential donor and acceptor sites of the active centers of MRE molecules can be further identified by applying MAC and FI methods.

## 3 Results and discussion

### 3.1 Experimental assessments

**3.1.1 WL.** WL experiments were implemented for SS samples in blank solution (HCl 1.0 M) and in the presence of different concentrations of MRE (50–250 ppm) at different temperatures (25–55 °C). The corresponding estimated corrosion parameters ( $\Delta w$ , CR,  $\theta$  and % IE) are displayed in Table 2. Surprisingly, the %IE values increased sharply with increasing

temperature (Fig. 4), indicating the possibility of chemical bond formation between MRE molecules and the SS surface. Fig. 1 shows plots of CR versus immersion time (1–24 h) at 25 °C. Meanwhile, the curves for temperatures from 35 to 55 °C are shown in the SI File (SF), see Fig. S2. It can be observed that the CR of SS decreased with prolongation of time and increase of concentration of inhibitory extract. While the %IE values increased significantly with increasing MRE concentration (Fig. 2). The variation in CR of SS at different temperatures is shown in Fig. 3. Fig. S1 in SF shows WL measurements versus immersion time.

**3.1.2 Kinetics and thermodynamics.** Based on the structure of MRE extract (Table 1), Their molecules are expected to be easily adsorbed on the SS surface and thus replace the H<sub>2</sub>O molecules adsorbed in the aqueous medium (blank), so the adsorption property here is carried out by the substitution process as follows:<sup>51,52</sup>



In order to elucidate the mechanism of MRE inhibition, several isothermal models have been investigated, namely Frumkin, Temkin, Langmuir, Flory–Huggins, Freundlich, and

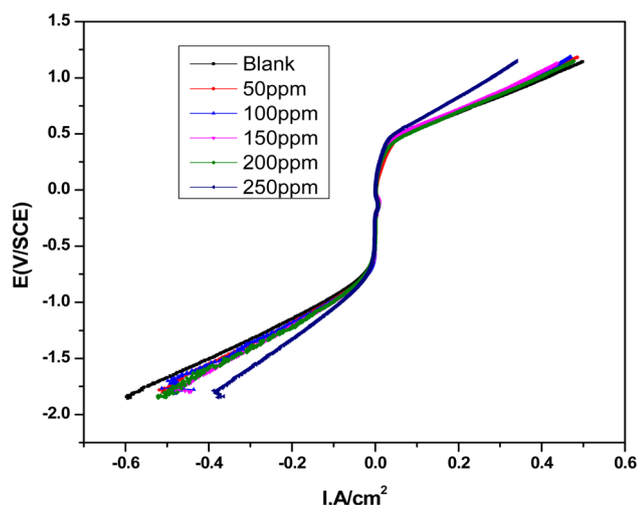


Fig. 12 LSV curves for SS in 1.0 M HCl solution and in the presence of different concentrations of MRE at 25 °C.

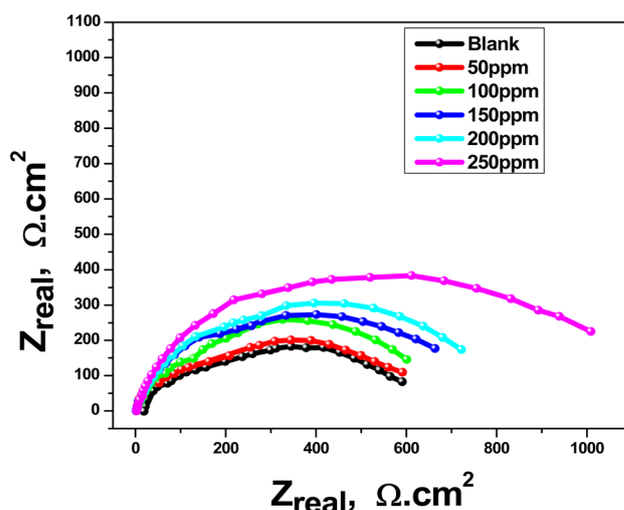


Fig. 13 Nyquist plots for SS in 1.0 M HCl solution and in the presence of different concentrations of MRE at 25 °C.



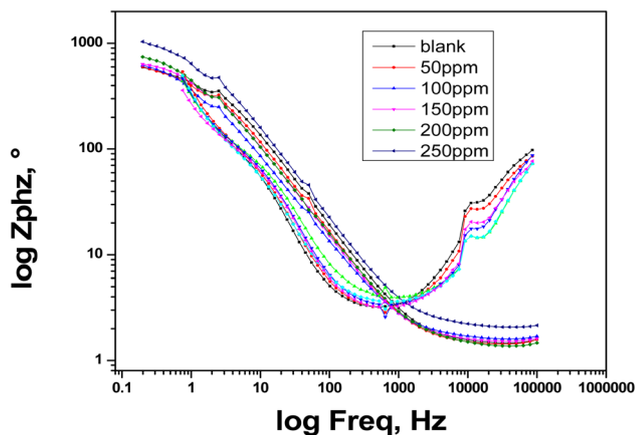


Fig. 14 BP plots for SS in blank solution (HCl 1.0 M) and in the absence and presence of different concentrations of MRE (50–250 ppm) at 25 °C.

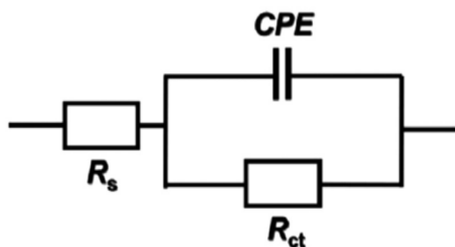


Fig. 15 Equivalent circuit proposed to fit the EIS experimental data.

Henry. Fig. 5 and 6 represent the best fit between the studied models, confirming that the MRE adsorption strategy can be understood through the Langmuir and Frumkin curves. Eqn (5) was used for the Langmuir model, and eqn (6) for the Frumkin model as follows:<sup>53</sup>

$$C_{\text{inh}}/\theta = (1/K_{\text{ads}}) + C_{\text{inh}} \quad (5)$$

$$\log(C_{\text{inh}} [\theta/(1 - \theta)]) = \log K + 2\alpha\theta/2.303 \quad (6)$$

$C_{\text{inh}}$  = MRE concentration, and equilibrium constant for adsorption =  $K_{\text{ads}}$ . Meanwhile, the Temkin, Flory–Huygens,

Henry, and Freundlich isotherms are shown in the SF file (see Fig. S4).

Returning to Fig. 5, plotting  $C_{\text{inh}}/\theta$  versus  $C_{\text{inh}}$  gives straight lines with slopes approaching one, intercepts equal to  $(1/K_{\text{ads}})$ , and correlation coefficient ( $R^2$ ) values equal to one. The  $K_{\text{ads}}$  values were then calculated and summarized in Table 3, where the increase of their values with increasing temperature confirms the improvement of the adsorption and bonding strength of MRE molecules on the SS surface in agreement with the WL results.<sup>53</sup> For the Frumkin isotherm, plotting the  $\log(C_{\text{inh}} [\theta/(1 - \theta)])$  versus  $\theta$  also gives straight lines with slopes approaching one (Fig. 6).

Then, the free energy of adsorption,  $\Delta G_{\text{ads}}$ , was calculated using eqn (7):<sup>54–56</sup>

$$K_{\text{ads}} = 1/55.5e^{-\Delta G_{\text{ads}}/RT} \quad (7)$$

where, 55.5 = concentration of water ( $M = \text{mol L}^{-1}$ ),  $R$  = gas constant ( $8.314 \text{ J mol}^{-1} \text{ K}^{-1}$ ), and  $T$  (K). The plot of  $\log K_{\text{ads}}$  versus  $1000/T$  gives straight lines from which the values of  $\Delta G_{\text{ads}}$  were determined and inserted in Table 3. Once the value of  $\Delta G_{\text{ads}}$  is known, the kind of adsorption can be suggested. Both chemical and physical adsorption kinds are possible. For chemical adsorption,  $\Delta G_{\text{ads}}$  should be  $> -40 \text{ kJ mol}^{-1}$ , whereas for physical adsorption, it should be  $< -20 \text{ kJ mol}^{-1}$ .<sup>57,58</sup> Examining the data in Table 3, we notice that the  $\Delta G_{\text{ads}}$  values are temperature dependent and all are negative. These negative values confirm the spontaneity of the adsorption process. By Frumkin's study, at temperatures below 45 °C, the values were in the range  $-40 > \Delta G_{\text{ads}} > -20 \text{ kJ mol}^{-1}$  indicating a mixed (physical and chemical) nature of the adsorption. Interestingly, at temperatures above 45 °C, the  $\Delta G_{\text{ads}}$  value was greater than  $-40 \text{ kJ mol}^{-1}$ , confirming the formation of a chemical bond between the adsorbed MRE molecules and the SS surface, *i.e.*, it is purely chemisorption, which is in agreement with the WL and kinetic studies described before.

The enthalpy of adsorption ( $\Delta H_{\text{ads}}$ ) is computed by Van't Hoff equation,<sup>59</sup>

$$\log K_{\text{ads}} = -\Delta H_{\text{ads}}/(2.303 \times RT) + \text{constant} \quad (8)$$

when  $\log K_{\text{ads}}$  is plotted against  $1000/T$  (Fig. 7), it gives slopes equal to  $-(\Delta H_{\text{ads}})/(2.303 \times R)$ , and hence the  $\Delta H_{\text{ads}}$  values were calculated (Table 3). It can be observed that the  $\Delta H_{\text{ads}}$  values are positive indicating that the inhibition process is endothermic with chemisorption dominating.<sup>60,61</sup>

Table 6 Parameters obtained from EIS investigations at 25 °C

Inh. conc. (ppm)	$R_{\text{ct}}$ ( $\Omega \text{ cm}^{-2}$ )	$R_{\text{S}}$ ( $\Omega \text{ cm}^{-2}$ )	$10^6 Y^0$ ( $\mu\Omega^{-1} \text{ S}^n \text{ cm}^{-2}$ )	$10^3 n$	$10^4 C_{\text{dl}}$ ( $\text{F cm}^{-2}$ )	$\theta$	% IE	$\chi^2$
Blank	530.5	1.47	374.1	880	2.78	—	—	0.00032
50	599.5	1.43	295.2	875	2.44	0.115	11.5	0.00042
100	635.5	1.59	275.3	826	2.21	0.165	16.5	0.00015
150	650.6	1.52	221.6	862	1.96	0.185	18.5	0.00054
200	768.8	1.37	198.2	837	1.62	0.310	31.0	0.00037
250	982.1	1.43	174.2	812	1.31	0.459	45.9	0.00024



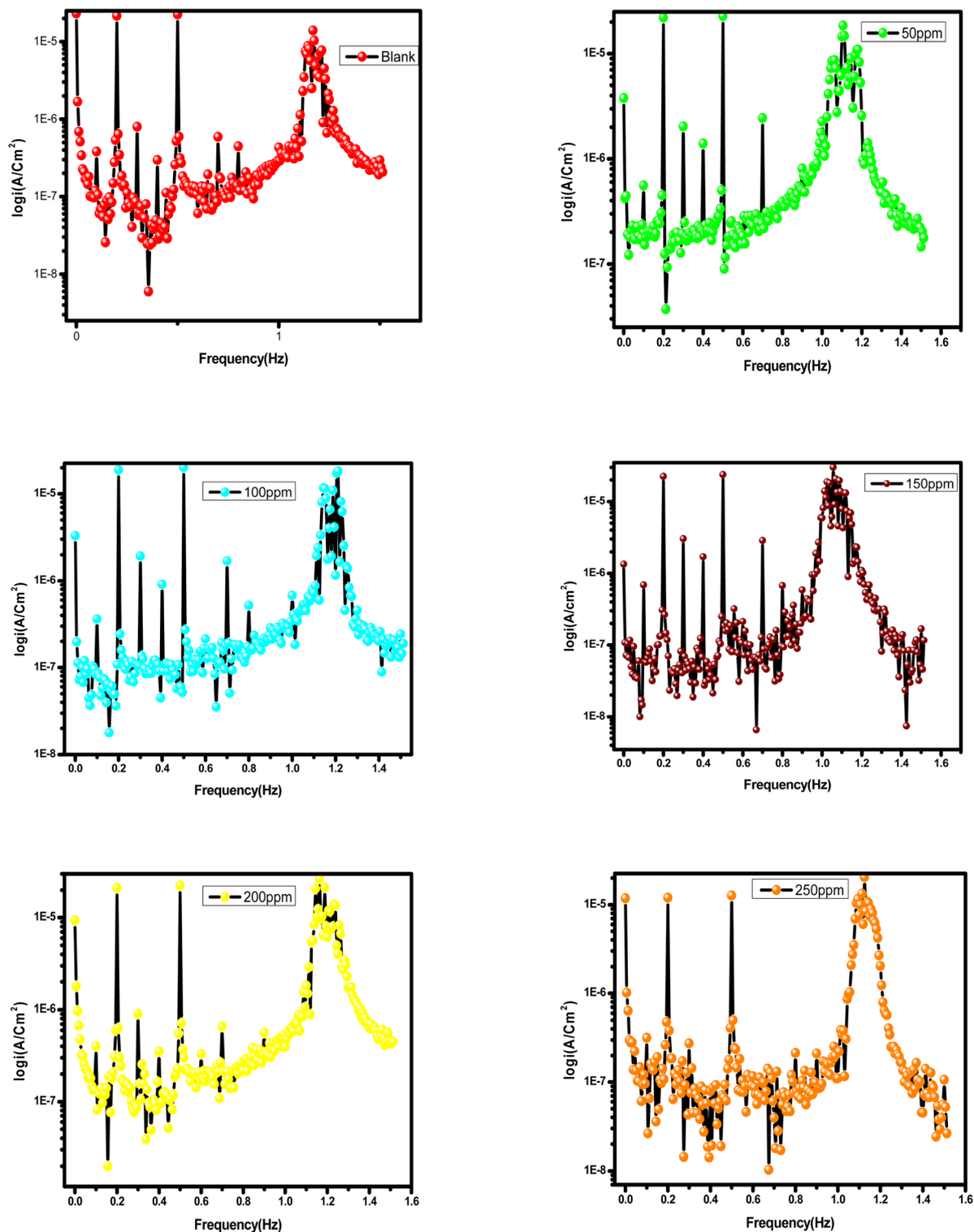


Fig. 16 EFM spectra for SS in 1 M HCl solution and in the presence of different concentrations of MRE at 25 °C.

The entropy of adsorption ( $\Delta S_{\text{ads}}$ ) is calculated by the following equation,<sup>53</sup>

$$\Delta G_{\text{ads}} = \Delta H_{\text{ads}} - T\Delta S_{\text{ads}} \quad (9)$$

Positive  $\Delta S$  values (Table 3) indicate increased randomness of the system due to the preferred replacement of  $\text{H}_2\text{O}$  molecules (desorption) by MRE molecules (adsorption).<sup>62</sup> These findings confirm the strong and preferential adsorption of MRE



Table 7 Parameters obtained from EFM investigations (Fig. 16)

Inh. conc. (ppm)	$i_{\text{corr}}$ ( $\mu\text{A cm}^{-2}$ )	$\beta_{\text{a}}$ ( $\text{mV dec}^{-1}$ )	$-\beta_{\text{c}}$ ( $\text{mV dec}^{-1}$ )	CF-2	CF-3	$\theta$	% IE	CR mpy
Blank	39	79.09	130.10	2.19	3.086	—	—	17.82
50	21	48.98	54.11	1.911	3.27	0.461	46.1	9.60
100	18.73	46.67	65.55	2.222	3.57	0.519	51.9	8.56
150	17.46	58.61	88.2	2.27	2.95	0.552	55.2	7.98
200	13.27	38.03	41.5	2.23	3.08	0.659	65.9	6.60
250	9.207	47.64	49.47	1.955	3.13	0.763	76.3	4.21

molecules followed by the formation of a protective inhibitory layer on the SS surface, acting as anticorrosion barrier.

The Arrhenius equation was used to calculate the activation energy ( $E_{\text{a}}^*$ ),<sup>63</sup>

$$\log CR = (-E_{\text{a}}^*/2.303 RT) + \log A \quad (10)$$

where,  $A$  is the Arrhenius constant. The Arrhenius plot shown in Fig. 8 gives straight lines, from their slopes the values of  $E_{\text{a}}^*$  were calculated and are listed in Table 4.

From the transition state equation,<sup>64</sup> the activation enthalpy ( $\Delta H^*$ ) and activation entropy ( $\Delta S^*$ ) are calculated as follows:

$$\ln\left(\frac{CR}{T}\right) = \left(\ln\frac{R}{Nh} + \frac{\Delta S^*}{R}\right) - \frac{\Delta H^*}{R} \frac{1}{T} \quad (11)$$

where,  $h$  = Planck's constant and  $N$  = Avogadro's number.

When  $\log CR/T$  was plotted against  $1000/T$  (Fig. 9), straight lines were obtained. From the slopes and intercepts, the values of  $\Delta H^*$  and  $\Delta S^*$  were calculated, respectively (Table 4).<sup>65</sup>

Interestingly, adding 250 ppm of MRE (the inhibitor) to the blank solution (only 0.1 M HCl) resulted in a sharp increase in the  $E^*$  value from 1.30 to 32.02  $\text{kJ mol}^{-1}$  (Table 4). Concurrently, the  $\Delta H^*$  value changed from  $-1.15$  to  $-34.61$   $\text{kJ mol}^{-1}$ . These results confirm that the addition of MRE forms a protective inhibitory layer on the SS surface, acting as a corrosion-resistant barrier.

### 3.1.3 Electrochemical measurements

**3.1.3.1 OCP.** Fig. 10 shows the variation in OCP of SS in blank solution (1.0 M HCl) and in the absence and presence of different concentrations of MRE (50–250 ppm) at 25 °C over a period of 1800 s. In the blank experiment, OCP values moved slightly toward positive values and quickly stabilized. Whereas when MRE was added, the OCP values first shifted towards the negative direction and then to the positive direction with stabilization, indicating that the anodic centres of the SS surface are the active adsorption sites of the extract molecules.

**3.1.3.2 PDP.** Tafel curves for SS in blank solution (HCl 1.0 M) and in the presence of different concentrations of MRE are depicted in Fig. 11. PDP test showed that the addition of MRE inhibitor effectively resisted SS surface corrosion by simultaneously suppressing the anodic (metal dissolution) and cathodic (e.g., hydrogen evolution or oxygen reduction) reactions.<sup>66,67</sup> This may be due to the fact that inhibitor molecules

are adsorbed onto the SS surface, blocking the oxidation/reduction sites that drive corrosion. As listed in Table 5, the sharp drop in the  $i_{\text{corr}}$  value from 2499.0 (blank) to 487.2  $\mu\text{A cm}^{-2}$  (250 ppm MRE) is evidence for this action. As a result, a significant increase in %IE of up to 81% occurred, confirming the adsorption of MRE molecules onto the SS surface, thus blocking the active corrosion centers. Table 5 summarizes the calculated PDP parameters: %IE, anodic and cathodic Tafel slopes ( $\beta_{\text{a}}$  and  $\beta_{\text{c}}$ ), corrosion current densities ( $i_{\text{corr}}$ ) and corrosion potentials ( $E_{\text{corr}}$ ). It has been reported<sup>46,66</sup> that an inhibitor is categorized as either anodic or cathodic type if the change in  $\Delta E_{\text{corr}} > \pm 85$  mV and it is a mixed-type if  $\Delta E_{\text{corr}} < \pm 85$  mV relative to  $E_{\text{corr}}$  value of the blank experiment. Its maximum value obtained here was 57 mV, confirming that MRE behaved as a mixed-type inhibitor. However, the preponderant anodic reaction control is indicated by the little change in  $E_{\text{corr}}$  values towards a positive route as MRE concentration increased. Additionally, there was no discernible change in the values of both  $\beta_{\text{c}}$  and  $\beta_{\text{a}}$ , suggesting that the two processes (anodic SS oxidation or corrosion and cathodic hydrogen reduction or evolution) were hindered without altering the mechanisms underlying the corrosion response.<sup>65–67</sup> The obtained anodic branches indicated limiting current densities at potentials positive of about 0 mV vs. SCE, evidence of the passivation nature of the 330 SS in the acidic medium. Hence, the effect of the inhibitor is measured mainly depending on the values of corrosion current densities ( $i_{\text{corr}}$ ) near  $E_{\text{corr}}$ . The MRE is effective in inhibiting the corrosion process as the reduction in  $i_{\text{corr}}$  value with increasing the MRE concentration. Moreover, %IE values enhanced in tandem with the MRE concentrations. Increasing the MRE content improved the polarization resistance due to the preferred adsorption of MRE on the SS surface as shown in the LSV plots (Fig. 12).<sup>68</sup>

**3.1.3.3 EIS.** The interactions at the SS interface boundary under various circumstances, both before and after addition of myrrh extract, were examined using EIS. EIS improved understanding of the mechanisms underlying corrosion inhibition by offering vital insights into the electro-resistive and capacitive characteristics of the substrates.<sup>69</sup> The obtained EIS spectra were introduced as Nyquist (Fig. 13) and Bode (Fig. 14) plots. As shown in Fig. 13, the study analyses the Nyquist plots (NQs) for SS in aqueous HCl (1.0 M). In the examined corrosive



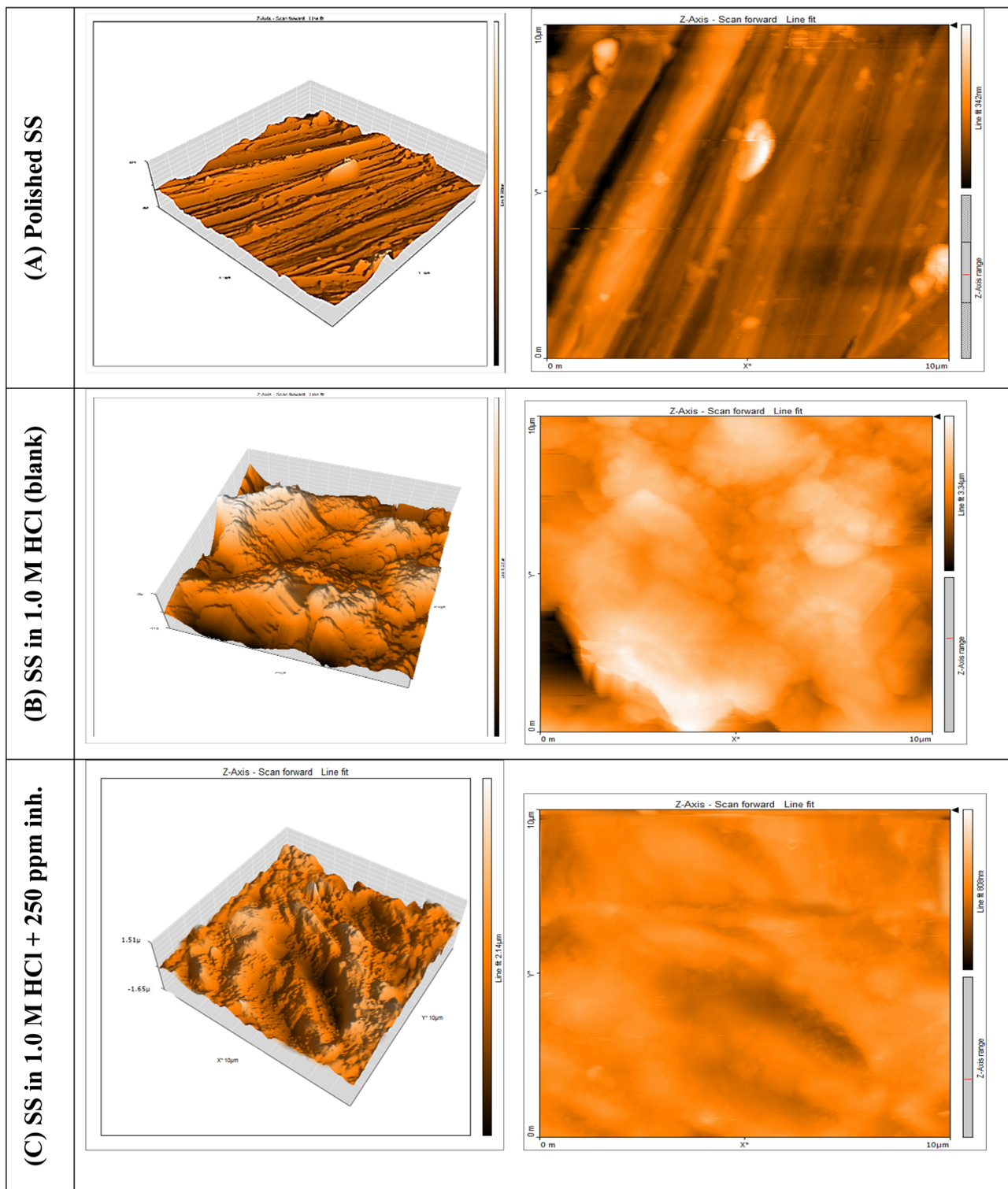


Fig. 17 AFM images of SS samples: (A) polished, (B) blank (immersed in 1.0 M HCl solution only), and (C) after adding 250 ppm of myrrh extract. Duration of corrosion/inhibition experiments = 24 h.

environment, MRE was discovered to improve the impedance response of SS; the NQs successfully demonstrated this behaviour. The semicircle size in the Nyquist plots significantly expanded in response to a rise in MRE concentration,

suggesting the formation of a shielding barrier that lessens the SS contact with the corrosive environment.<sup>70</sup> Instead of the anticipated perfect circles, the Nyquist plots revealed the existence of non-ideal, twisted loops. Usually, this variation is



Table 8 Data obtained from AFM analysis (Fig. 17)

Surface	$S_q$ (nm)	$S_a$ (nm)	Max peak to valley height (nm)
Polished	40.02	28.22	373
1.0 M HCl (blank)	501.89	382.60	3560
1.0 M HCl + 250 ppm MRE	164.52	145.61	1097

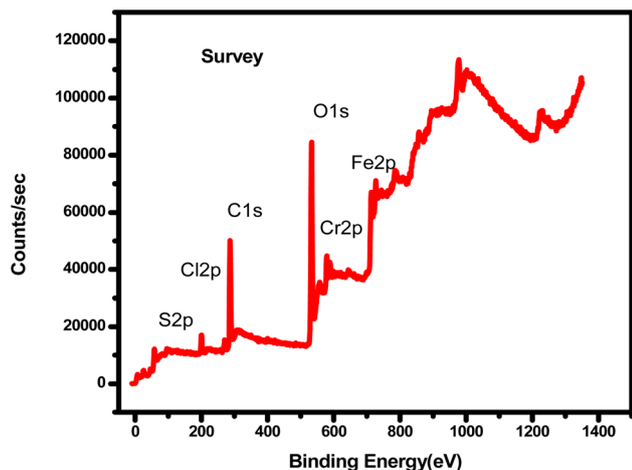


Fig. 18 XPS survey of the inhibited SS substrate in 1.0 M HCl with 250 ppm myrrh extract at 25 °C.

attributed to the SS surfaces' imperfections and heterogeneous properties.<sup>71</sup>

In this investigation, an appropriate equivalent circuit which was selected for the fitting of the acquired experimental findings in order to assess the electrochemical characteristics of the SS surface is shown in Fig. 15. Fitting the obtained experimental EIS data into the equivalent-circuit model produced the impedance values displayed in Table 6. A one-time constant in the Bode phase (BP) is depicted in Fig. 14. The figure demonstrates the fit of the line using the parametric circuit model and presents the impedance and phase data for SS exposed to HCl (1.0 M) at 25 °C as Bode plots. The majority of EIS spectra used in corrosion experiments have well-defined time constants for two capacitive semicircles or a low capacitive loop in the BP patterns. As can be shown, the corresponding CPE (constant phase element) models (see Fig. 14) are well fitted by the BP plots (Fig. 14) for myrrh extract. The  $Z_{CPE}$  value of the fitted circuit is calculated by:<sup>72</sup>

$$Z_{CPE} = 1/Y_0(j \times \omega)^n 11$$

$R_p$  = polarization resistance,  $R_s$  = solution resistance,  $n$  = CPE exponent and  $Y_0$  = CPE constant. The value of  $n$  provides an insight into the inhibitor's nature. This value for the majority of organic inhibitors is between 0.8 and 1.0. On the other hand, with an ideal circuit,  $n$  will be 1.0. It was used in place of the circuit's best match to the experimental results. By replacing

CPE with the values of double-layer capacitance ( $C_{dl}$ ), such values for this circuit can be found:<sup>73</sup>

$$C_{dl} = [Y_0(R_p)^{1-n}]^{1/n} \quad (12)$$

A significant decrease in  $C_{dl}$  values brought on by the addition of the inhibitor suggests that metal-inhibitor adsorption has replaced metal-water adsorption. At the metal/corrosive medium interface, this leads to a decrease in the local dielectric constant and/or an increase in the thickness of the electric double layer. Additionally, the Bode plots (Fig. 14) showed only one phase maximum with a single-time constant; this could be the case when the charge transfer process takes place at the Fe-electrolyte interface. The significant adsorption of metal and inhibitor may be the reason for the progressive increase in the maximum phase angle at intermediate frequency when myrrh concentration rose. Additionally, the phase angle measurements showed surface roughness resulting from structural and interfacial origins, deviating somewhat from the ideal corrosive system (values of roughly 80).<sup>74</sup> By computing the values of % IE and  $\theta$ , one may ascertain the degree of adsorption of MRE on the SS surface:<sup>75</sup>

$$\% \text{ IE} = [(R_{p(\text{inh})} - R_{p(\text{uninh})})/R_{p(\text{inh})}] \times 100 \quad (13)$$

$$\theta = (R_{p(\text{inh})} - R_{p(\text{uninh})})/R_{p(\text{inh})} \quad (14)$$

where,  $R_{p(\text{uninh})}$  and  $R_{p(\text{inh})}$  are the values of polarization resistance in the uninhibited and inhibited solutions, respectively. The values of  $R_p$  and % IE listed in Table 5 further demonstrated the adsorption of extract molecules on the SS surface. It was discovered that when the concentration of MRE increases, so did the measured values of  $R_p$  of the extract and, consequently, the values of %IE. The results demonstrate that, in 1.0 M HCl solution, the investigated MRE exhibited a good inhibitory action against the SS corrosion.

**3.1.3.4 EFM.** The EFM method can determine the current response using two sine waves of different frequencies for metal corrosion as shown in Fig. 16.<sup>76</sup> As reported earlier, the causality factors (CF-2 and CF-3) determine the validity of the EFM data and should be 0–2.0 and 0–3.0, respectively.<sup>76</sup> Table 7 illustrates the causation factors for the inhibitor are extremely near to the known values. Consistent with all the above measurements, with increasing MRE concentration, the CR rate of SS in this harsh environment decreased and an efficiency of up to 76% was achieved.

**3.1.4 Surface analysis.** Inspection the morphology, topography and species exist on a solid surface is an important and necessary step to fully understand the mechanism of action of an inhibitor in a corrosion experiment. Fig. 17 displays three AFM micrographs of SS specimens. Fig. 17a shows a polished, clean, and dried SS sample that appears uniform and smooth with a slight roughness of 28.217 nm (Table 8). To evaluate the surface after corrosion and inhibition experiments, the SS surface was checked after immersion for 24 h in HCl solution (1.0 M) in the absence (B) and presence of MRE inhibitor at a concentration of 250 ppm (C). The SS sample in the blank



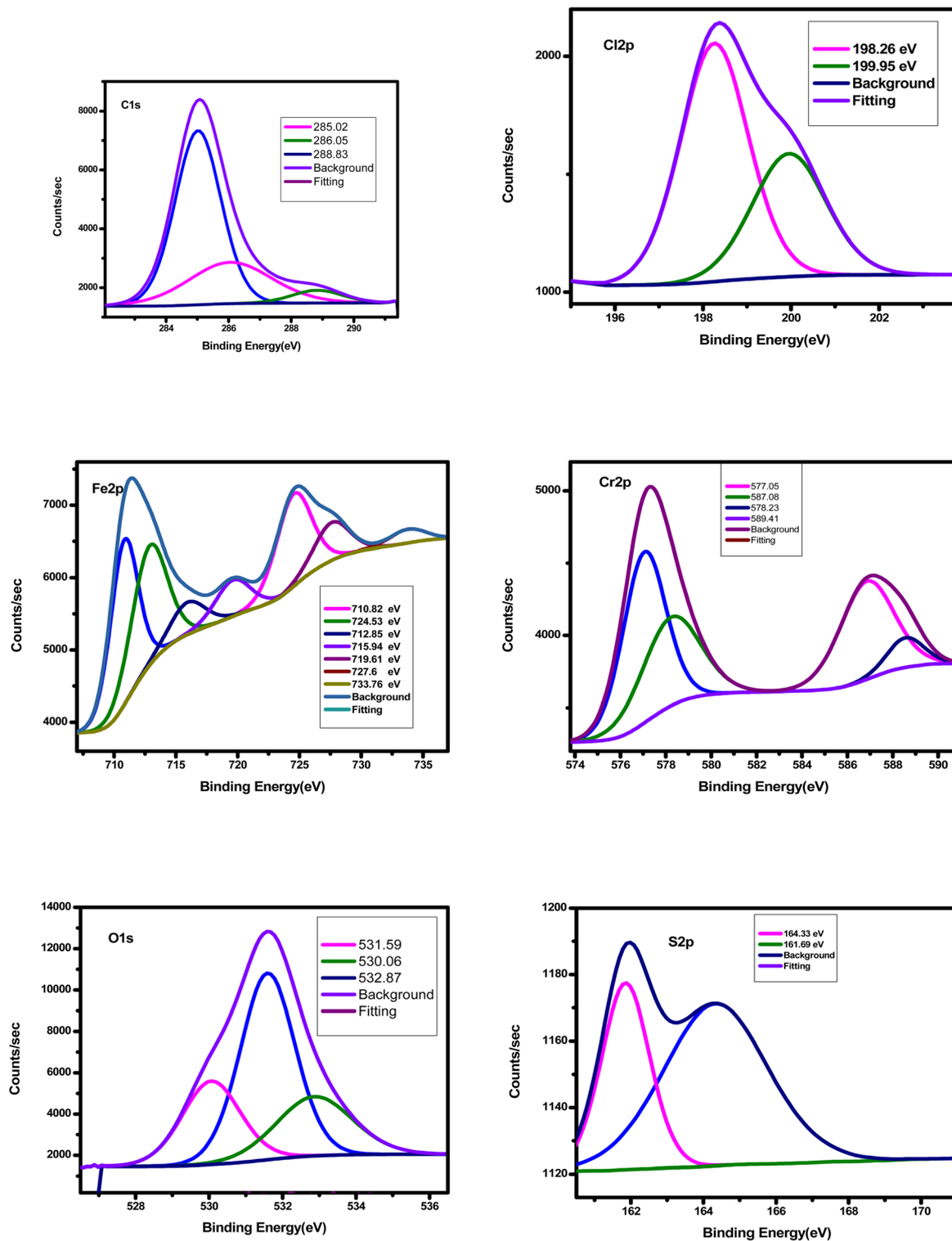


Fig. 19 High resolution XPS spectra of the inhibited SS surface (*i.e.* after immersion in 1.0 M HCl with 250 ppm of MRE at 25 °C).

experiments appears to be severely damaged and has the highest roughness of about 382.6 nm, compared to that obtained in the presence of MRE of 145.61 nm (Table 8). These results also demonstrated the effect of adsorption of MRE

molecules on the SS surface, as they blocked most of the active corrosion sites.

The chemical composition of surfaces has not been investigated so far. Therefore, XPS was used to determine the



**Table 9** Species present on the surface of the inhibited SS surface (*i.e.* after immersion in 1.0 M HCl with 250 ppm of MRE at 25 °C)

Species	BE (eV)	Components
C 1 s	285.02	C–Cl, C–C, C–H
	286.05	
	288.83	
Fe 2p	710.82	Fe <sub>2</sub> O <sub>3</sub> , Fe2p <sub>1/2</sub> of Fe <sup>3+</sup> , FeOOH, Fe <sub>2</sub> O <sub>3</sub> , Fe 2p <sub>3/2</sub> of Fe <sup>2+</sup> , FeCl <sub>3</sub> , ferric compounds satellites
	724.53	
	712.85	
	715.94	
	719.61	
	727.60	
Cl 2p	198.26	FeCl <sub>2</sub>
	199.95	
O 1 s	531.59	Metal oxide, hydroxide, FeO and Fe <sub>2</sub> O <sub>3</sub>
	530.06	
	532.87	
S 2p	164.33	Sulfide
	161.69	
Cr 2p	577.05	Chromium oxide, chromium chloride, chromium hydroxide
	587.08	
	578.23	
	589.41	

surface composition and chemical bonding of MRE molecules on the surface. An XPS survey of SS in 1.0 M HCl with 250 ppm MRE at 25 °C would reveal the surface elemental composition and chemical states after the MRE has potentially formed a protective layer on the SS surface. As shown in Fig. 18, the XPS scan shows peaks corresponding to the elements present in the SS substrate (*e.g.*, Fe, Cr, and S), and any elements of MRE with an emphasis on essential species such as C and O. The binding energies of these elements would help determine how the MRE molecules have adsorbed and chemically bonded to the steel surface, indicating the nature of the protective layer that mitigates corrosion. Fig. 19 displays the high resolution XPS spectra of the inhibited sample showed characteristic peaks that were recognized as C 1s, Cl 2p, Fe 2p, O 1s, Cr 2p and S 2p. The complying assignments and binding energies (BE, eV) for each peak constituent are provided in Table 9. Spectra of C 1 s may be attributed to C–H–, C–C–, and C–Cl bonds at 285.02, 286.05, and 288.83 eV for samples that were maintained with MRE inhibitor.<sup>77,78</sup> The appearance of a chlorine peak on the SS surface treated with MRE in 1.0 M HCl solution is caused by the interaction of chloride ions with the positively charged SS surface.<sup>79</sup> The Cl 2p spectra of the inhibited SS samples showed two peaks for Cl 2p<sub>3/2</sub> at 198.26 and additional peaks for Cl 2p<sub>1/2</sub> at 199.95 eV, indicating the presence of a Fe–Cl bond in FeCl<sub>3</sub>.<sup>80</sup> Seven peaks were seen in the Fe 2p XPS spectra of the inhibited SS samples. The origins of these peaks were determined to be Fe 2p<sub>3/2</sub> of Fe<sup>2+</sup> at 710.82 eV, Fe 2p<sub>3/2</sub> of Fe<sup>3+</sup> at 712.85 eV, Fe 2p<sub>3/2</sub> satellites of Fe<sup>2+</sup> at 719.61 eV, and Fe 2p<sub>1/2</sub> of Fe<sup>2+</sup> at 727.6, 733.76 eV. Furthermore, the high-resolution O 1s spectra of the SS samples treated with the MRE inhibitor display three distinct

peaks. The first peak, designated as O<sup>2–</sup> at 530.06 eV, is associated with the oxygen atoms that are coupled to Fe<sup>2+</sup> and Fe<sup>3+</sup> in the FeO and Fe<sub>2</sub>O<sub>3</sub> oxides.<sup>81</sup> At 531.59 and 532.87 eV, the second and third peaks are attributed to OH<sup>–</sup> and might be connected to Fe<sup>3+</sup> in FeOOH.<sup>82,83</sup> In fact, chromium was found in myrrh extract at a very low concentration of 0.001148% (11.48 ppm) by ICP-MS analysis of the inorganic constituents of MRE.<sup>84</sup> The chromium signal appearing in the XPS spectrum of a myrrh-coated SS substrate likely originates from the SS itself, which is a Cr-rich alloy (Cr = 19%), and its presence is confirmed by the signal at specific binding energies in the spectrum. Thus, Cr 2p is linked to peaks with energies of 577.05, 587.08, 578.23, and 589.41 eV in Cr-oxide, Cr-chloride, and Cr-hydroxide. However, the sulfur signal in the XPS spectrum of the MRE-coated SS substrate may come either from the MRE or from the SS itself. Returning to the data shown in Table 1,<sup>40</sup> and the ICP-MS/GC-MS analyses of the inorganic–organic ingredients of MRE,<sup>84</sup> the possibility that the S-signal came from the MRE extract is excluded as it is not present in its constituents. Interestingly, recent FT-IR (Fourier transform infrared) analyses of the Al surface inhibited by methanolic myrrh extract in 1 M HCl showed no evidence of S-containing groups.<sup>85</sup> Therefore, the S-signal is likely to originate from the SS substrate. The MRE inhibitor's adsorption on the SS surface in 1.0 M HCl solution is supported by these XPS study results.

### 3.2 Computational investigations

DFT is a very useful calculation for predicting the inhibition strength of different molecules such as MRE.<sup>86,87</sup> Then, the electrical properties of MRE were calculated and presented in Tables 10 and S1, S2 of the SI File. Given the fact that  $E_{\text{HOMO}}$  is associated with electron donating capacity, increasing its values would enhance the effectiveness of the inhibitor.<sup>88,89</sup> It is very interesting that MRE molecules occupy the highest value of HOMO, which would be the reason for the large inhibitory activity (Table 10). Meanwhile, the low  $E_{\text{LUMO}}$  value indicates that the inhibitor molecules can locate another negative charge on the SS surface, as the capacity to accept electrons is essential to  $E_{\text{LUMO}}$ . The inhibitors under study have comparatively high LUMO values and high  $E_{\text{HOMO}}$  values. The 13 MRE molecules studied were found to be highly effective because they interacted while acting as donors. The metal functions as a donor to the inhibitor molecules, lowering the inhibitor effectiveness while increasing the metal reactivity, even if the inhibitor molecules with a modest  $E_{\text{HOMO}}$  value decrease metal reactivity.  $E_{\text{HOMO}} - E_{\text{LUMO}}$  showed that the evaluated inhibitor chemicals had the best corrosion inhibition (Fig. 20, S5 and S6). Significant inhibitory action was shown by the high softness and low hardness values of the inhibitor compounds with a high  $E_{\text{HOMO}}$  and a low energy gap. The  $\chi$  values of MRE molecules may provide insight into the nature of the bond between them and the SS surface.<sup>90</sup> Iron atoms will create chemical connections by acquiring electrons from inhibitor molecules, according to the research on the inhibitory effects of inhibitor molecules intended as iron-inhibitors (Tables 10, S1 and S2). According to the  $\chi$



Table 10 Parameters computed by DFT for neutral MRE molecules

Parameter	Compound				
	(1)	(2)	(3)	(4)	(5)
$-E_{\text{HOMO}}$ (eV)	-5.005	-5.031	-5.153	-5.087	-5.018
$-E_{\text{LUMO}}$ (eV)	-0.179	0.145	-0.367	0.114	-0.214
$\Delta E$ (eV)	4.826	4.890	4.790	4.973	4.800
$\eta$ (eV)	2.410	2.450	2.395	2.49	2.400
$\sigma$ (eV) <sup>-1</sup>	0.414	0.408	0.418	0.402	0.416
Pi (eV)	-2.590	-4.890	-5.520	-4.973	-5.230
$\chi$ (eV)	2.590	4.890	5.520	4.973	5.230
Dipole moment, $\mu$	1.157	0.4177	1.043	0.7814	1.2400
Molecular area ( $\text{\AA}^2$ )	261.279	232.595	241.841	229.501	262.693
$\omega$ (Electrophilicity index)	1.390	4.960	6.360	4.970	5.700
$\omega^+$ (Electro-accepting power)	0.397	0.382	0.510	0.370	0.417
$\omega^-$ (Electro-donating)	2.990	2.970	3.270	2.870	3.030
$\varepsilon$ (Nucleophilicity index)	0.716	0.201'	0.157	0.032	0.175
$\Delta E$ back donation	-0.603	-0.613	-0.599	-0.623	-0.600
$\Delta N_{\text{max}}$ (e)	0.350	-0.136	-0.256	-0.139	-0.197

Parameter	Compound				
	(6)	(7)	(8)	(9)	(10)
$-E_{\text{HOMO}}$ (eV)	-5.046	-4.695	-5.175	-5.896	-5.295
$-E_{\text{LUMO}}$ (eV)	0.146	-0.205	0.146	-1.302	-2.042
$\Delta E$ (eV)	4.900	4.490	5.030	4.590	2.550
$\eta$ (eV)	2.450	2.250	2.520	2.300	1.280
$\sigma$ (eV) <sup>-1</sup>	0.410	0.400	0.397	0.435	0.781
Pi (eV)	-4.900	-4.900	-5.030	-7.198	-7.340
X (eV)	4.900	4.900	5.030	7.198	7.340
Dipole moment, $\mu$	0.7915	1.6092	2.150	3.6220	5.555
Molecular area ( $\text{\AA}^2$ )	249.88	254.63	250.150	232.910	234.920
$\omega$ (Electrophilicity index)	4.900	5.340	5.020	11.260	21.045
$\omega^+$ (Electro-accepting power)	0.383	0.074	0.391	0.133	0.210
$\omega^-$ (Electro-donating)	3.483	2.761	3.050	4.906	6.170
$\varepsilon$ (Nucleophilicity index)	0.204	0.187	0.199	0.089	0.048
$\Delta E$ back donation	-0.613	-0.563	-0.630	-0.575	0.320
$\Delta N_{\text{max}}$ (e)	-0.126	-0.137	-0.148	-0.634	-1.195

Parameter	Compound		
	(11)	(12)	(13)
$-E_{\text{HOMO}}$ (eV)	-5.047	-5.227	-5.135
$-E_{\text{LUMO}}$ (eV)	-0.935	0.020	-1.903
$\Delta E$ (eV)	4.110	5.027	3.230
$\eta$ (eV)	2.060	2.510	1.620
$\sigma$ (eV) <sup>-1</sup>	0.485	0.398	0.617
Pi (eV)	-5.982	-5.207	-7.038
X (eV)	5.982	5.207	7.038
Dipole moment, $\mu$	4.0833	2.7910	8.3783
Molecular area ( $\text{\AA}^2$ )	249.595	260.61	290.039
$\omega$ (Electrophilicity index)	8.690	5.400	15.290
$\omega^+$ (Electro-accepting power)	0.937	0.335	2.273
$\omega^-$ (Electro-donating)	3.928	3.016	5.792
$\varepsilon$ (Nucleophilicity index)	0.110	0.185	0.065
$\Delta E$ back donation	-0.515	-0.628	-0.405
$\Delta N_{\text{max}}$ (e)	-0.413	-0.185	-0.851

value, the inhibitor works well even when it has a low value. The investigated inhibitor molecules work better, as indicated by the  $\Delta N$  value in Tables 10, S1 and S2, which demonstrates that

they transfer more electrons to the Fe atoms on the metal surface. Another distinct parameter displays in Tables 10, S1 and S2 is dipole moment ( $\mu$ ). The IE indicates that a large  $\mu$



indicates substantial corrosion inhibition potency, even if prior research has not established a significant association between  $\mu$  and IE. Lowering the  $\mu$  value was proven to boost corrosion IE in many experiments. Since the evaluated inhibitor's  $\mu$  value is low, it indicates that the metal surface is coated more firmly. Because they expand the area in which they may interact with the SS surface, larger MRE molecules have a stronger inhibitory impact. In terms of the nucleophilicity index ( $\epsilon$ ), we found that the neutral form of the MRE components had greater numerical values than the protonated form. As a result, MRE components in their neutral state exhibit strong nucleophile, indicating that the neutral form can help electrons adhere to the SS surface. Recently, the electrical donor capacity ( $\omega^- = (3I + A)^2/16(I - A)$ ) and the electrical receiver capacity ( $\omega^+ = (I + 3A)^2/16(I - A)$ ) have been used to assess PPE's capacity to transmit or receive electrical charge.<sup>91</sup> The positive attributes of hardness ( $\eta$ ) and, consequently, the negative attributes of ( $E_{BD}$ ) (back donation energy =  $-\eta/4$ ) are pivotal indicators of and charge transfer/electron donation dynamics from iron PPE molecules.

The adsorption locator module has been identified as the most suitable structure (adsorbent) within the corrosion-simulating fluid (comprising 250 H<sub>2</sub>O molecules, along with 10 H<sub>3</sub>O<sup>+</sup> and 10 Cl<sup>-</sup> ions) for the adsorption of MRE (adsorbate) onto the Fe(110) crystal. The methodologies for the adsorption of MRE molecules, in both neutral, protonated and vacuum forms, are depicted in Fig. 20, S7 and S8, respectively, illustrating the top and lateral perspectives of the Fe(110) substrate. Tables 11, S3 and S4 enumerate the results obtained from the MD simulation concerning deformation, solid adsorption, and the overall energetic states. The calculations aim to ascertain the  $E_{total}$  pertinent to the substrate and MRE by aggregating molecular energies derived from solid adsorption alongside distortion energies. The energy quantified during the detachment of a single adsorbate particle from the adsorbent substrate is represented by  $dE_{ads}/dN_i$ . Their values associated with the MRE components exceed those related to water molecules, suggesting that MRE exhibits a stronger adsorption affinity than water. Consequently, it is anticipated that the MRE components will adhere to the SS surface, thereby forming a passive layer that safeguards the metallic substrate against acidic corrosion. According to Tables 11, S3 and S4, along with prior empirical investigations,<sup>92,93</sup> the components of the MRE demonstrate elevated adsorption energies and exhibit commendable performance within the solution matrix. A meticulous examination indicates that the molecular configuration of the local molecule is oriented parallel to the SS surface.<sup>94</sup> To enhance the contact area or surface coverage efficacy of SS, the components of the MRE are expected to adopt this specific adsorption behaviour. Rather than facilitating undesirable attacks by Cl<sup>-</sup>, the expanded SS/myrrh contact zone will restrict substrate accessibility.<sup>95,96</sup> It is hypothesized that electron transfer from the SS surface to the MRE components transpires in a retrograde manner, thereby reversing the reaction, particularly in the presence of a charged protonic component. Theoretical analyses substantiate that MRE components exhibit a strong adsorption affinity for the SS surface, resulting in the formation of durable adsorbent

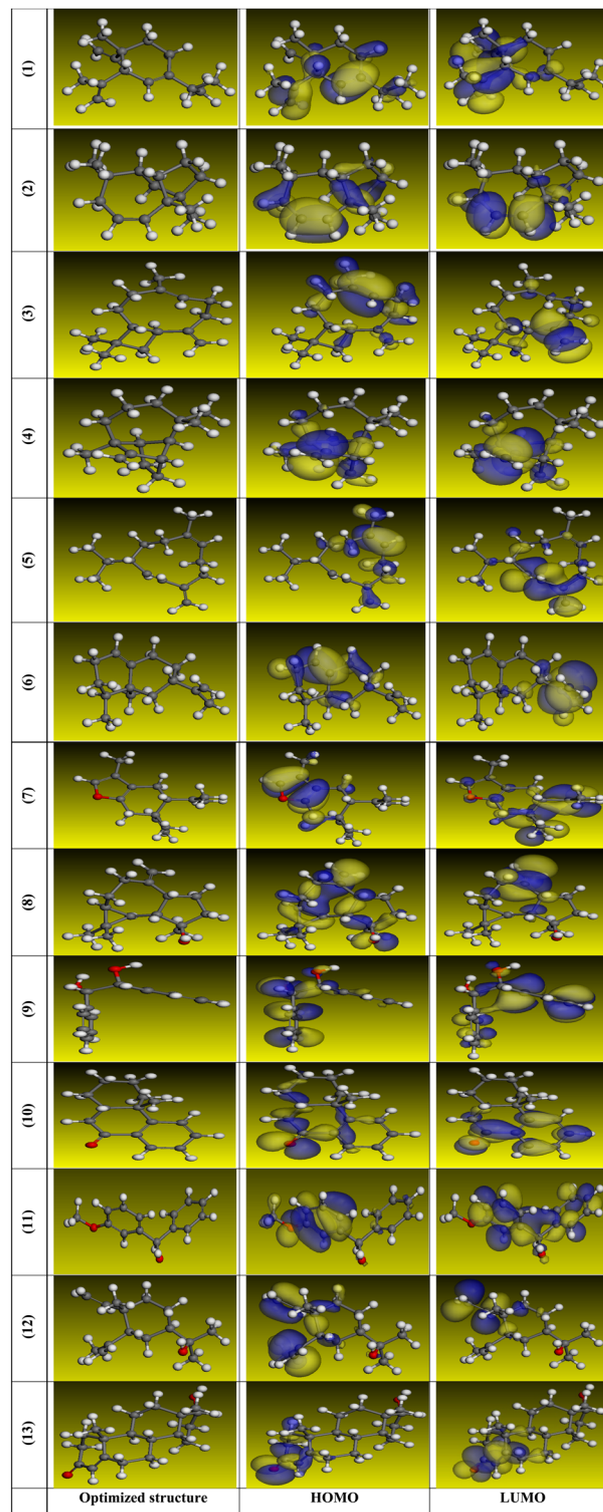


Fig. 20 Optimized structures, LUMO and HOMO for the 13 neutral MRE molecules.

coatings that protect the SS substrate from corrosion induced by HCl solutions. Fig. 21, S7 and S8, in conjunction with Tables 11, S3 and S4, illustrate the adsorption dynamics of MRE molecules in both vacuum and acidic (neutral and protonated)



Table 11 Parameters computed by MD simulation for MRE molecules

Parameter (kcal mol <sup>-1</sup> )	Compound				
	(1)	(2)	(3)	(4)	(5)
Total energy	-5156.390	-5.112	-4257.168	-6938.518	-6874.990
Adsorption energy	-5130.234	-5092.202	-4244.306	-6917.882	-6875.450
Rigid adsorption energy	-5317.082	-5271.337	-4.426	-7110.053	-7057.550
Deformation energy	186.850	179.130	181.600	192.170	182.100
dE <sub>ads</sub> /dN <sub>i</sub>	-110.960	-55.440	-105.160	-94.230	-111.740
dE <sub>ads</sub> /dN <sub>i</sub> for water	-9.490	-7.710	-10.530	-8.200	-9.720

Parameter (kcal mol <sup>-1</sup> )	Compound				
	(6)	(7)	(8)	(9)	(10)
Total energy	-6866.785	-6890.549	-6852.649	-6749.567	-6900.022
Adsorption energy	-6842.330	-6874.151	-6901.110	-6775.957	-6893.540
Rigid adsorption energy	-7031.989	-7069.073	-7097.571	-6964.140	-7090.820
Deformation energy	189.660	194.920	196.460	188.180	197.280
dE <sub>ad</sub> /dN <sub>i</sub>	-1.670	-116.970	-93.800	-114.170	-122.220
dE <sub>ads</sub> /dN <sub>i</sub> for water	-9.820	-7.700	-13.770	-9.340	-10.500

Parameter (kcal mol <sup>-1</sup> )	Compound		
	(11)	(12)	(13)
Total energy	-6907.988	-6954.281	-6971.699
Adsorption energy	-6.928.218	-6902.147	-6902.404
Rigid adsorption energy	-7135.060	-7098.410	-7095.140
Deformation energy	206.840	196.260	192.730
dE <sub>ad</sub> /dN <sub>i</sub>	-129.220	-103.620	-121.600
dE <sub>ads</sub> /dN <sub>i</sub> for water	-8.620	-10.840	-11.890

environments. The adsorption energies in the context of an aqueous solution surpass those in a vacuum environment, indicating a more effective adsorption of myrrh molecules onto the SS surface. To quantify the energy associated with the adsorption of MRE onto the SS(110) interface, the following equations may be employed:<sup>97,98</sup>

$$E_{\text{binding}} = -E_{\text{interact}} \quad (15)$$

$$E_{\text{interact}} = E_{\text{tot}} - (E_{\text{sub}} + E_{\text{inh}}) \quad (16)$$

$E_{\text{sub}}$  = energy contributions from both water molecules and the substrate.

Predominantly, regions exhibiting negative electrostatic potential are represented in red, indicating areas of significant electrical activity and electrophilicity. The majority of heteroatoms (O, N) and conjugated double bonds can be observed in regions of high electron density. Functional groups containing oxygen serve as indicators of zones that facilitate electrophilic attacks through negative potential. Fig. 22, S9 and S10 illustrate the nucleophilic attack on the H atoms (blue) of the 13 molecules of MRE according to MESP investigations.

It is worth noting that the active sites in MRE structures are regions that show an increased propensity to interact with the SS substrate. Fig. 23 delineates a comparable behavioral trend,

wherein the total negative charge (TNC) escalates concomitantly with the protonation of the MRE molecules.<sup>90</sup> An augmentation in the structural interaction among protonated entities is observable through the electrostatic attraction that exists between the protonated components and the Cl<sup>-1</sup> that have been adsorbed onto the SS surface.<sup>53</sup> Organic components of myrrh extract, inclusive of heteroatoms, have the ability to undergo protonation in aqueous HCl, thus producing positively charged entities. They can then engage in interactions with the Cl<sup>-1</sup>, which diffuses abundantly across the metal surface. Fig. 24 and S11 delineate the atomic structures of the MRE components under examination, highlighting elevated values of Fukui functions ( $f_k^+$  and  $f_k^-$ ) in both protonated and non-protonated scenarios, serving as supplementary indicators (FI). The findings indicate that each molecular entity possesses distinctive atoms pertinent to both electrophilic and nucleophilic frameworks. The LUMO, representing the molecular segment with the most significant  $f_k^+$ , exhibits augmented reactivity towards donor reagents and is consequently predisposed to nucleophilic assault. The atom within the target molecule that exhibits the highest  $f_k^-$  values, correlated with HOMO, facilitates reactivity directed towards an acceptor species, culminating in electrophilic engagement. The presence of protons (H<sup>+</sup>) obstructing the atomic centres in the protonated configuration results in a diminution of  $f^-$  values for the



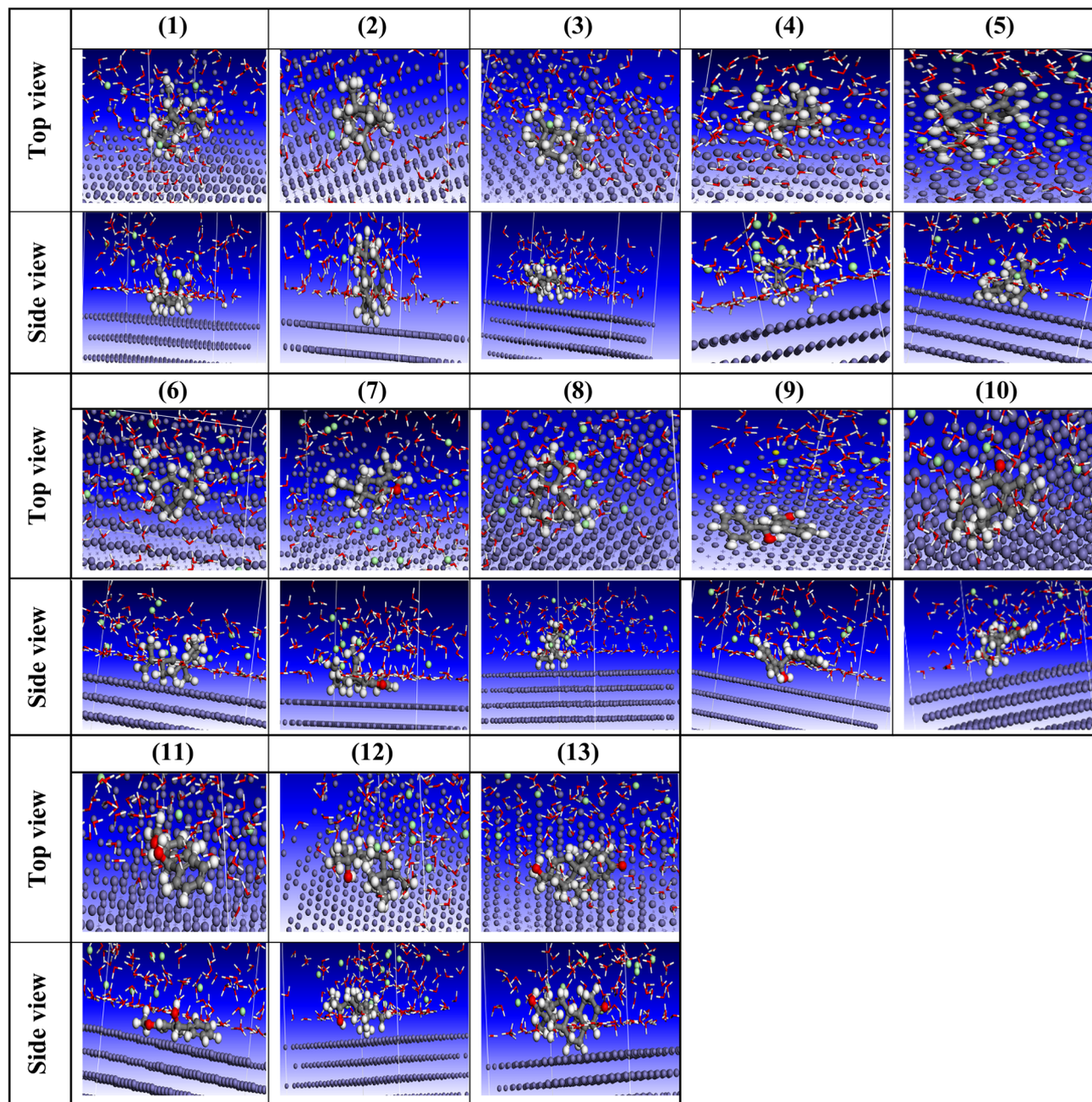


Fig. 21 Top and side views of the most stable adsorption configuration of the 13 molecules of MRE computed using MD simulation on Fe(110) substrate.

corresponding atoms, whereas the  $f^+$  values for the entities experience an upsurge owing to the atoms' enhanced capacity for electron acceptance (Fig. 24 and S11). Moreover, altered atoms within analogous components manifest divergent  $f^+$  and  $f^-$  values, thereby signifying that the MRE exhibits a dual effect.

### 3.3 The inhibition mechanism

The efficiency of a corrosion inhibitor is determined by its adsorption capacity and its bond strength to the surface of the material. Myrrh extract contains 13 compounds (Table 2) with

unique structures that undoubtedly make it a promising green corrosion inhibitor. Statistical calculations (MD and DFT) verified that the inhibitory properties of the MRE originate from the electron-dense active sites, such as the  $\pi$ -bond of the benzene ring, OH group, and C=O. These sites enable MRE species to form bonds with the SS surface and protect it. Functional groups in organic molecules are most likely to accept a proton in aqueous acidic solutions and be protonated.<sup>98</sup> Based on the MRE structure, the SS surface is expected to be significantly protected either by: (i) Electrostatic interactions occur between



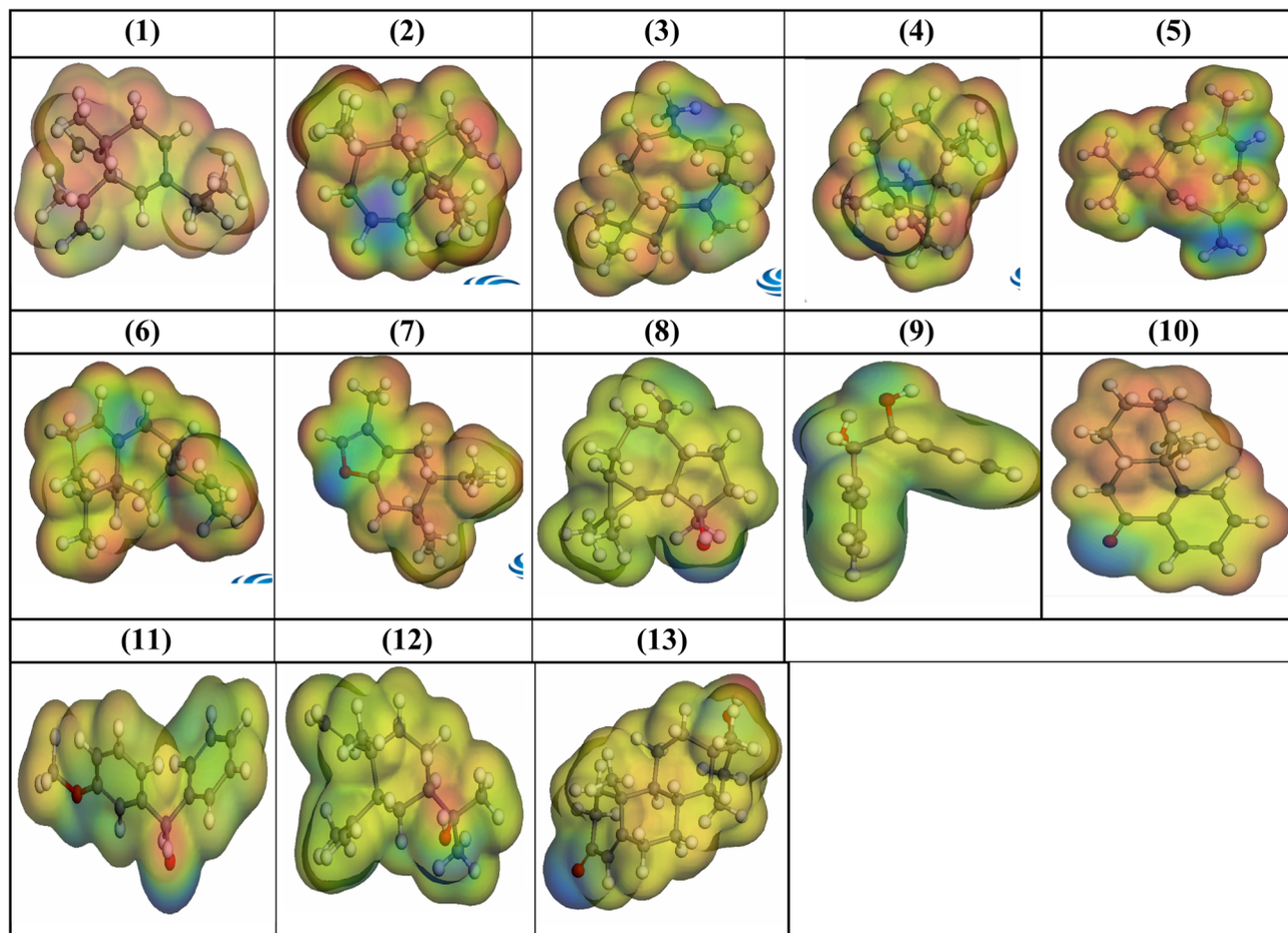


Fig. 22 Illustrations showing the nucleophilic attack on the H atoms (blue) of the 13 molecules of MRE.

chloride ions covering the SS surface and protonated MRE molecules (physisorption).<sup>99</sup> (ii) A protective layer is formed on the SS surface by the conjugated bonds of the aromatic rings and the unshared  $\pi$ -electrons of the inhibitor.<sup>26</sup> (iii) The presence of heteroatoms such as O that donate their lone pairs of electrons to the vacant d-orbital of Fe also forms a strong protective layer on its surface (chemisorption).<sup>53</sup>

Another interesting factor in this study that had a strong influence on the inhibition capacity was temperature. The results of this factor were surprising during the WL measurements shown in Table 2. The IE% value increased drastically from 77.6 to 93.6% with the temperature increasing from 25 to 55 °C, respectively, confirming the dominance of chemical adsorption at elevated temperatures. These results were also supported by the calculated  $\Delta G_{\text{ads}}$  value of  $-41.78 \text{ kJ mol}^{-1}$  at 55 °C (Table 3), indicating the formation of stable chemical bonds between the adsorbed MRE molecules and the SS surface.

An important issue that has not yet been discussed is the stability of MRE components in an acidic environment at high temperatures. Based on experimental results, there is no indication that the inhibitor molecules decompose under the

reaction conditions here. The significant increase in the % IE value to 93.6% with increasing temperature to 55 °C, confirms the formation of stable chemical bond (chemisorption). In fact, the stability of an organic molecule is linked to its chemical bonds and structure. If organic molecules decompose, chemical bonds break, and new chemical species may form. XPS can identify the presence of these new species by detecting new peaks in the spectrum or changes in the relative peak intensities of the elements present, which was not the case with myrrh extract. Further support for the stability of MRE molecules under reaction conditions is that most of its components (Table 1), such as the phenanthrene derivative (49.41%), include stable aromatic rings and functional groups (ketones, ethers).

Strictly speaking, this is a superior property of this inhibitor over others in industrial applications requiring high-temperature processes, such as: cleaning and polishing steel equipment, which is typically performed at temperatures between 50 and 60 °C, and oil well acidification, where the oil may be at the bottom of the well at high temperatures. In air coolers that operate at temperatures higher than ambient



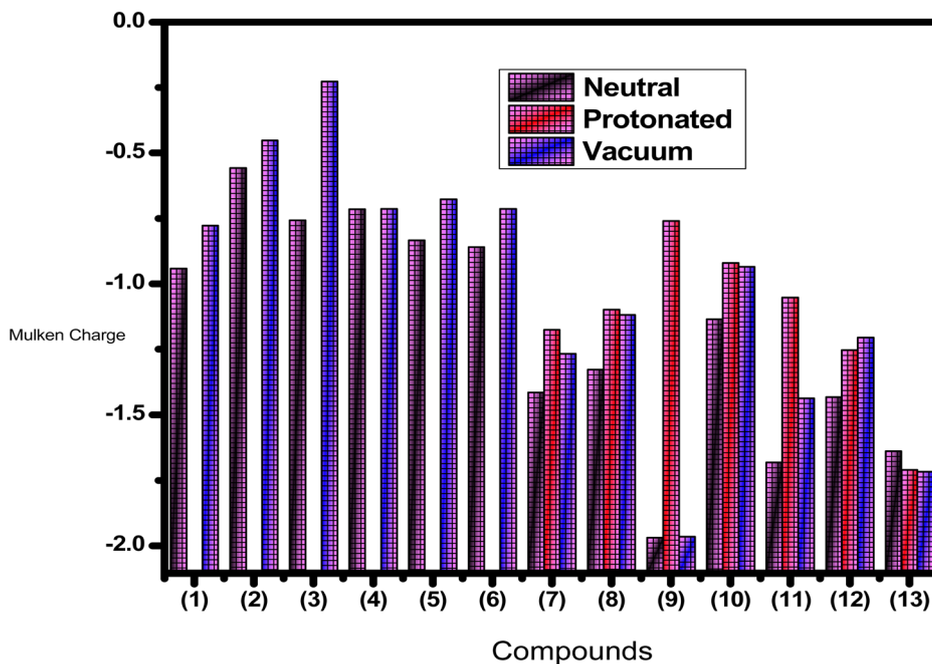
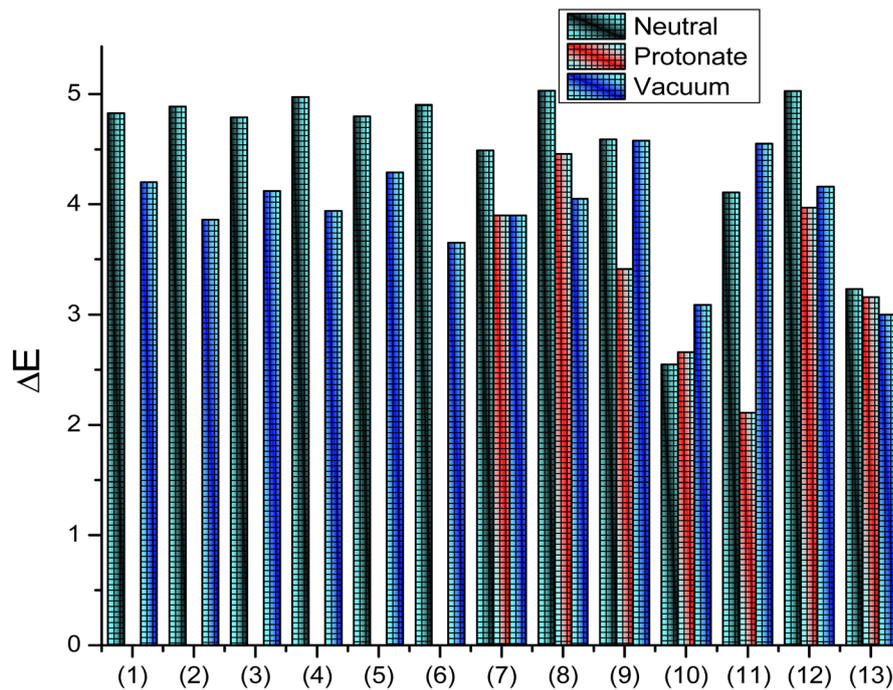


Fig. 23 Mulliken charges (MAC) and  $\Delta E$  values for the 13 MRE molecules.

temperatures as well. Another advantage of this green inhibitor is its low cost compared to other inhibitors, as the percentage of myrrh plant extract reaches 15%, making it commercially and economically feasible and highly effective for the industrial

sector. Strictly speaking, green inhibitors like MRE represent a more sustainable option because they are less harmful than their traditional counterparts.



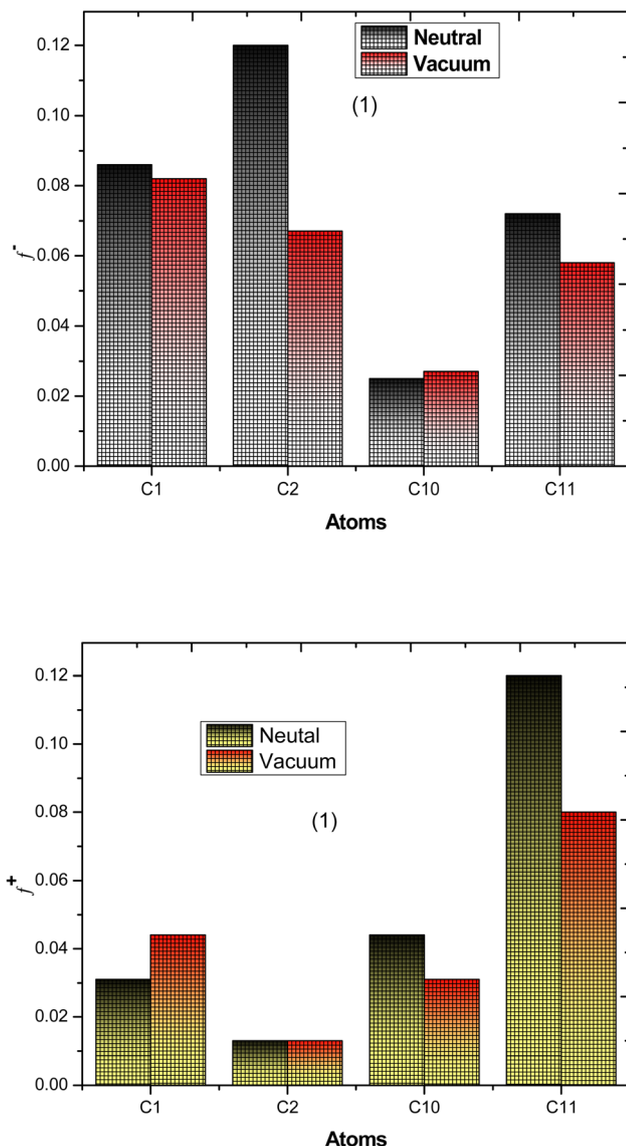


Fig. 24 Atomic structures of the MRE component (1), highlighting elevated values of Fukui functions ( $f_k^-$  and  $f_k^+$ ).

## 4 Conclusions

Herein, green MRE extract was investigated as a potential corrosion inhibitor for SS in 1.0 M HCl solution at different temperatures using multifaceted experimental-computational methods. With the increase of MRE concentration and temperature, the inhibition efficiency increased dramatically. The Langmuir and Frumkin adsorption isotherms' models govern the myrrh extract's adsorption on the SS surface. The MRE functioned as a mixed-type inhibitor. As seen by XPS and AFM, a smooth and uniform protective layer from MRE particles was constructed on the SS surface. The thermodynamic and kinetic parameters were calculated and analyzed. The computational techniques provided more insights into the mechanism of inhibition of the myrrh extract. All experimental results of different techniques were in an excellent agreement with each

other and with the employed computational methods. Green inhibitors, like MRE, represent a more sustainable option because they are less harmful than traditional inhibitors. The results of this study may have an impact on the rational development of a strong, effective, environmentally friendly, and inexpensive inhibitor for protecting metals in harsh acidic environments, fulfilling environmental, industrial, and economic aspirations.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The datasets generated and analyzed during this study are available from the corresponding author upon reasonable request.

Supplementary information is available. See DOI: <https://doi.org/10.1039/d5ra07138k>.

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