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Experimental and computational investigation of 3,5-di-*tert*-butyl-2-(((3-((2-morpholinoethyl)(pyridin-2-ylmethyl) amino) propyl)imino) methyl)phenol and related reduced form as an inhibitor for C-steel

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HIGHLIGHTS

GRAPHICAL ABSTRACT

- Two inhibitors containing morpholine tags have been synthesized and characterized.
- The corrosion studies show an excellent corrosion inhibitor of two ligands.
- Scanning electron microscopy evaluations confirm the protection of carbon steel.
- The adsorption mechanisms preferred the Langmuir adsorption isotherm.
- Molecular modeling provides more information into the inhibition performance.

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ABSTRACT

The effect of new compounds containing morpholine moiety including: 3,5-di-tert-butyl-2-(((3-((2-morpholinoethyl)(pyridine-2-ylmethyl)amino)propyl)imino)methyl)phenol (L^1) and its reduced form, 3,5-di-tert-butyl-2-(((3-((2-morpholinoethyl)(pyridine-2-ylmethyl)amino)propyl)amino) methyl)phenol (L^2) on the corrosion inhibiting of C-steel in 1.0 M HCl and 3.5 wt% NaCl was tested by experimental and computational techniques. The concentration of inhibitor varied in the range of 0.2–2.0 mM. The corrosion behavior of C-steel was investigated using electrochemical techniques, including open circuit potential (OCP), potentiodynamic polarization (Tafel), electrochemical impedance spectroscopy (EIS) and scanning electron microscopy (SEM) inhibitor-free and inhibitor containing solutions. From the Tafel diagram, the i_{corr} of the C-steel electrode in the 1.0 M hydrochloric acid solution represents the highest value among the other inhibitor-containing solutions. The i_{corr} for C-steel in L^1 containing solution decreases gradually from 1.43×10^{-4} to 0.91×10^{-4} A/cm², with the concentration of L^1 increasing from 0.2 to 2.0 mM. Along with the increase of the L² concentration from 0.2 to 2.0 mM, the i_{corr} is reduced by about 41%. ElS studies demonstrated that the capacitance of the double layer increased by the addition of inhibitor-containing acidic solution. SEM images confirmed the corrosion results in which

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Received 19 August 2021; Received in revised form 21 March 2022; Accepted 16 April 2022 Available online 19 April 2022 0254-0584/© 2022 Elsevier B.V. All rights reserved. inhibitor-containing solution protects the C-steel surface against the attack of species. The results showed that in all solutions containing equivalent concentrations of inhibitors, the reduced form of L^1 had a better inhibition efficiency than the L^1 inhibitor. Inhibitors, L^1 and its reduced form exhibit Langmuir adsorption isotherm, in which monolayer adsorption of inhibitors is carried out on the C-steel electrode. The properties of the studied molecules to be corrosion inhibitors were examined with the Gaussian software program. The calculations of the molecules were made in different methods and basis sets.

1. Introduction

A broad range of organic inhibitors applied in the corrosion domain is expensive [1]. Electron pairs and negative ions are transferred from the inhibitors to the metal d orbitals, resulting in coordination complexes with specific geometries [1]. It is well-known that the coupling between an aldehyde and amine can be readily carried out and that the produced imine compound undergoes hydrolysis under acidic conditions [2]. These compounds have attracted significant and increasing interest in various aspects [3]. There are many studies, as well as some of ours on the synthesis of various Schiff base ligands containing piperazine and morpholine moieties [4–7]. Also, there are some reports on the application of imines for metal corrosion prevention [8–11]. There is a desire to use iron alloys in the industry due to the low cost and high strength in different applications.

Meanwhile, carbon-containing iron alloys (i.e., steel) play a significant role in various industrial sections, including constructions of building and railway, gas and oil industry, etc. [12]. Since C-steel alloys are used in common aqueous environments, the corrosion behavior of these types of alloys seems to be very important for the management of corrosion costs [13]. There are two main strategies to improve the corrosion rate and elongate the life of C-steel in harsh acidic media, including the use of highly resistant alloys in corrosion media and the addition of organic inhibitor compounds in the acidic solutions [14]. The availability and low cost of organic inhibitors result in extensive usage in acidic solutions [15–17]. Recently, the potential of corrosion protection of inhibitors containing imines [18], azines [19], sulphonamides [20], thiols [21], etc. has been investigated in the literature. Generally, organic inhibitors contain O, N, P and S heteroatoms which show inhibiting behavior in acidic media [12,22]. The protection mechanism of inhibitors can be explained by the adsorption of organic atoms on the steel surface and then filling the surface of the particles and removing corrosive ions from the C-steel surface [15]. Although there are many common inhibitors in the industries, numerous efforts are being made to introduce new groups of organic inhibitors to increase the inhibition efficiency and minimize the corrosion costs [23]. Therefore, there is a serious need to research this field and introduce new materials to achieve the mentioned aims.

Barmatov and Hughes [24] stated that most of the inhibitors used in sulfuric acid and hydrochloric acid solutions have insufficient hydrolytic stability at low pH conditions. Only a few studies have been performed to investigate the corrosion resistance of reduced forms of inhibitors [15,



Scheme 1. Synthesis of L^1 and L^2 .

24-26].

Here, the main goal is to investigate the behavior of L^1 and its reduced form as an inhibitor on C-steel corrosion in a 1.0 M hydrochloric acid solution by using electrochemical impedance measurements (EIS). Theoretical calculations are a better guide than experimental procedures. With the quantum chemical parameters found as a consequence of theoretical calculations, the inhibitory properties of molecules are expected to be predicted. This process is faster and more accessible [27]. The calculations of the molecules were made in different methods and basis sets and afterward, the comparison of the inhibition properties of the inhibitor molecules with the obtained parameters was made.

2. Experimental

N1-(2-morpholinoethyl)-N1-((pyridine-2-yl)methyl)propane-1,3diamine and 3,5-di-*tert*-butyl-2-hydroxy benzaldehyde were synthesized according to literature methods [6,27,28]. Pyridine-2-carbaldehyde and 2-aminoethylmorpholine were purchased from Sigma Aldrich Company. IR spectra were collected on a BIO-RAD FTS-40A spectrophotometer (4000-400 cm⁻¹). Nuclear NMR spectra were recorded on a Bruker 500 spectrometer operating at 500.06 MHz, respectively. Mass spectra were measured on a Bruker micro TOFQ. Standard microanalysis for all complexes was carried out using a CHNS/O elemental analyzer (model 2400, Perkin-Elmer).

2.1. Synthesis

2.1.1. Synthesis of 3,5-di-tert-butyl-2-(((3-((2-morpholinoethyl)(pyridin-2-ylmethyl) amino)propyl)imino)methyl)phenol (L^1)

An ethanolic solution of N1-(2-morpholinoethyl)-N1-((pyridine-2-yl) methyl)propane-1,3-diamine (0.5 mmol, 0.139 g) was added dropwise to the stirring solution of 3,5-di-tert-butyl-2-hydroxy benzaldehyde (0.5 mmol, 0.117 g) in ethanol (50 mL). The mixture was refluxed for 12 h. A brown oil was obtained that was filtered off, washed with cold ethanol and dried in vacuo. Yield: (85%). Anal. Calc. for C₃₀H₄₆N₄O: C, 75.27; H, 9.69; N, 11.70. Found: C, 75.00; H, 9.83; N, 11.22%. IR (ATR, cm⁻¹): 3424, 1633 v(C=N), 1590, 1470 v(C=C). EI-MS (*m*/*z*): 494.710, Found: 494.500 [L]⁺. ¹H NMR (CDCl₃, ppm) $\delta = 1.32$ (s, 9H, H-r); 1.43 (s, 9H, H-w); 1.88 (p, 2H, H-f); 2.40 (t, 4H, H-b); 2.49 (t, 2H, H-e); 2.69 (t, 4H, H-c, H-d); 3.61 (t, 2H, H-g); 3.68 (t, 4H, H-a); 3.79 (s, 2H, H-i); 7.06 (d, 1H, H-k); 7.10 (t, 1H, H-m); 7.47 (t, 1H, H-s); 7.49 (s, 1H, H-p); 7.61 (t, 1H, H-l); 8.31 (s, 1H, H-h); 8.51 (d, 1H, H-n); 13.81 (s, 1H, H-y). ¹³C NMR (CDCl₃, ppm) δ = 29.28 (c-r); 29.43 (c-s); 31.32 (c-v); 31.52 (c-w); 28.46 (c-f); 51.47 (c-c); 52.08 (c-d); 54.08 (c-b); 56.99 (c-e); 57.16 (c-g); 60.94 (c-i); 66.91 (c-a); 117.80 (c-m); 121.85 (c-o); 122.83 (c-p); 125.67 (c-k); 126.68 (c-t); 127.84 (c-u); 131.85 (c-l); 136.33 (c-q); 148.91 (c-n); 157.80 (c-x); 160.14 (c-h); 165.96 (c-j) (Scheme 1).

2.1.2. Synthesis of 3,5-di-tert-butyl-2-(((3-((2-morpholinoethyl)(pyridin-2-ylmethyl)amino)propyl)amino) methyl)phenol (L^2)

To an ethanlic solution (50 mL) of 2-((3-(2-morpholinoethylamino)-N3-((pyridine-2-yl)methyl) propylimino) methyl)pyridine (0.5 mmol, 0.239 g) was added slowly sodium borohydride (0.25 mmol, 0.009 g). The mixture was refluxed for 12 h. A brown oil was obtained that was filtered off, washed with cold ethanol and dried in vacuo. Yield: (85%). Anal. Calc. for C30H48N4O: C, 74.95; H, 10.06; N, 11.65. Found: C, 75.03; H, 9.89; N, 11.92%. ⁻¹ IR (ATR, cm): 3232 υ(N–H)_{str}, 1675 υ (N-H)_{bend}, 1591 v(C=N), 1479 v(C=C). EI-MS (m/z): 496.73, Found: 497.50 [L+1]⁺. ¹H NMR (CDCl₃, ppm) δ = 1.33 (s, 9H, H-s); 1.43 (s, 9H, H-r); 1.75 (p, 2H, H-f); 2.38 (t, 4H, H-b); 2.43 (t, 4H, H-c); 2.49 (t, 2H, Hd); 2.62 (t, 2H, H-e); 2.75 (t, 2H, H-g); 3.63 (t, 4H, H-a); 3.75 (s, 2H, Hh); 3.93 (s, 2H, H-i); 6.79 (d, 1H, H-k); 6.88 (t, 2H, H-m); 7.12 (s, 1H, Hp); 7.24 (t, 1H, H-l); 7.29 (s, 1H, H-t); 8.45 (d, 1H, H-n). ¹³C NMR (CDCl₃, ppm) δ = 29.67 (c-r); 29.72 (c-s); 30.06 (c-f); 31.72 (c-v); 31.80 (c-w); 34.13 (c-g); 34.90 (c-h); 45.96 (c-c); 47.19 (c-d); 51.10 (c-h); 54.03 (c-b); 56.84 (c-e); 60.73 (c-i); 66.82 (c-a); 122.02 (c-m); 122.98 (c-

Table 1

Chemical co	nposition	(in wt. %)	of the base	metal.		
Flements	C	Si	Ni	Mn	p	

Elements	C	51	INI	IVIII	P	CI
wt. %	0.049	0.010	0.003	0.221	0.013	0.001

o); 136.37 (c-k); 148.88 (c-t); 148.96 (c-u); 153.41 (c-n); 154.75 (c-j); 157.96 (c-l); 153.41 (c-n); 159.70(c-x) (Scheme 1).

2.2. Immersion time measurements

The weight loss measurements were performed using coupons with a dimension of $1 \times 1 \times 0.1$ cm³ cuts from the C-steel. The electrodes were polished with emery papers, followed by rinsing with acetone and ethanol and dried in hot air and kept in a desiccator. The C-steel samples were immersed in 50 ml of 1.0 M HCl solution (inhibitor-free) and different concentrations of inhibitors (0–2 mM) for 12 h. These experiments were done at a temperature of 22 °C. After every 1 h, the samples were taken out and rinsed several times with deionized water and acetone. Next, they were dried in an oven and then weighed. The experiments were repeated at least twice under similar conditions, and the average result was recorded.

2.3. Calculation method

Calculations were made using the Gaussian software program [29] to compare the theoretical activities of inhibitor molecules. Inhibitor molecules were optimized by these calculations. Many quantum chemical parameters were obtained in calculations using these optimized structures. For the optimization of molecules, calculations were made on the B3lyp, HF, and M062X method 6-31++g(d, p) basis set. Many quantum chemical parameters were calculated as a result of the calculations made on these basis sets. Many quantum chemical parameters have been calculated from these calculations. These are EHOMO (Highest Occupied Molecular Orbital), ELUMO (Lowest Unoccupied Molecular Orbital), ΔE (HOMO-LUMO energy gap), electronegativity (χ), chemical potential (μ), chemical hardness (η), electrophilicity (ω), nucleophilicity (ε), global softness (σ) and proton affinity (PA) [30,31].

2.4. Solutions, electrodes and electrochemical experiments

Electrochemical measurement was carried out with a conventional three-electrode-electrochemical cell using a Potentiostat/galvanostat set (Iviumstat compact 20250 H) controlled by IviumSoft electrochemistry software and a GSTAT101 N Potentiostat (Metrohm Autolab) controlled by Nova software (Version 2.1.4). The C-steel was used as the working electrode, and its chemical composition is shown in Table 1. An Ag/AgCl electrode (Metrohm) filled with 3.5 M KCl (217 mV vs. SHE at 22 °C) and a platinum electrode was employed as a reference and counter electrodes, respectively. Before electrochemical testing, C-steel was mounted by epoxy cold resin to have a surface area of 1 cm² exposed to the corrosive solution. The electrodes were ground wet to 1500 grit SiC followed by rinsing with acetone and ethanol and dried in hot air. 1.0 M hydrochloric acid and 3.5 wt % NaCl solutions were used as electrolyte for all electrochemical experiments. The electrochemical experiments were carried out under non-stirred, ambient pressure, and a controlled temperature of 22 °C. Two new substances, including L^1 and L^2 forms, were used to compare the inhibitors' results. The concentration of inhibitors was varied within the range of 0.2–2.0 mM.

To ensure the accuracy of analyses, experiments were started only after stabilizing the open circuit potential (OCP) within ± 5 mV [32]. For the Tafel tests, the current values were recorded in the range ± 0.70 V vs. OCP at a constant scan rate of 10 mV/s and converted to current density by figuring the surface area. Electrochemical impedance spectroscopy (EIS) tests were conducted in 100 ml of a 1.0 M HCl solution with and without inhibitors at OCP. The frequency range was 10^{-1} – 10^{5} Hz with



Fig. 1. The OCP value of C-steel in 1.0 M HCl at different concentrations of (a) L¹ and (b) L².

peak-to-peak amplitude of 10 mV. Each electrochemical experiment was repeated at least twice to ensure the reproducibility of results.

2.5. Surface characterization

The C-steel surface of the sample was characterized by an FEI Quanta 450 scanning electron microscope (SEM), which was equipped with energy-dispersive X-ray spectroscopy (EDS). The C-Steel was cut into 1 \times 1 \times 1 cm³, and then the surface of the samples was polished on emery paper of 400, 800, 1000 and 1500 grades. The polished surface was washed twice with distilled water and ultrapure acetone, respectively. Subsequently, the specimens were immersed in 100 ml of 1.0 M HCl solution with and without L¹ and L² inhibitors for 24 h. Finally, the samples were rinsed with distilled water and acetone to remove any adhered particles.

3. Result and discussion

3.1. Characterization of L^1 and L^2

A new Schiff base ligand (L^1) was prepared through the condensation reaction of *N1-(2-morpholinoethyl)-N1-((pyridine-2-yl)methyl)propane-1,3-diamine* and *3,5-di-tert-butyl-2-hydroxy benzaldehyde*, with a molar ratio of 1:1, for 6 h in ethanol. The prepared ligand and related reduced form were characterized by microanalysis and some spectroscopic techniques.

3.1.1. FT-IR and mass spectra

The FT-IR spectrum of the L¹ showed a sharp band at 1633 cm⁻¹ related to the stretching vibration frequency of the imine group, indicating the condensation of the precursors to produce the Schiff base ligand. The mass spectrum of the L¹ showed the molecular ion peak at m/z = 494.71 amu, which is consistent with the proposed molecular formula. The L² was prepared by an in-situ reduction of the Schiff-base ligand and characterized by microanalysis, IR, EI-MS and ¹H and ¹³C NMR spectroscopy. The IR spectrum of L² showed bands at 1591, 1570 and 1479 cm⁻¹, associated with the v(C=N) and v(C=C) vibrations from the pyridine ring. The v(N-H) bending and stretching bands appeared at 1675 and 3232 cm⁻¹, respectively. The mass spectrum of L² showed the molecular ion peak at m/z = 497.50 seen to be consistent with the proposed molecular formula.

3.1.2. NMR spectral studies

The ¹H and ¹³C NMR spectra of ligands (L^1 and L^2) were recorded in CDCl₃. The peaks obtained were consistent with the structures of the synthesized Schiff base and its reduced compound. For both compounds, the aromatic protons appeared as multiples between 7.06 and 8.51 ppm. The imino proton, for L^1 , was observed at 8.31 ppm, while in the reduced form, the singlet at δ 3.75, due to $-CH_2$ - group, confirmed the

formation of the L^2 . The ¹³C NMR spectrum showed the signals due to the methyl carbon at 51.10 ppm, and the peaks which appeared in the range 122.02–159.70 ppm reflected the aromatic carbons. The peak due to the imino carbon in L^1 appeared at 160.14 ppm.

3.2. Electrochemical measurements in 1.0 M HCl

3.2.1. OCP measurements

The OCP magnitude for samples can be precisely a sign of their electrochemical activity. The OCP evolution of the C-steel in 1.0 M HCl solution in the absence and presence of different concentrations of L¹ and L² is shown in Fig. 1. After immersion of the C-steel electrode in inhibitor-free solution (1.0 M HCl), the OCP was almost unchanged during its measurement for 900 s. The OCP of the C-steel generally reached a steady-state condition in the first 200 s after being immersed in test solutions. More evident in Fig. 1a, the C-steel OCP in the L^{1} containing solutions showed a higher value than the L¹-free solution. The reaching OCP value for C-steel in the L¹-containing solution occurs in the early stage of immersion in 1.0 M HCl (c.a. 200 s), as is clear from Fig. 1a. During the first 200 s, the OCP value varied around 6 mV for the C-steel in the L¹ solution. After that OCP value fluctuated in a small range (less than 5 mV), which can be assigned to the competitive dynamic process by desorption and adsorption of the inhibitor on the Csteel surface [33,34]. The OCP value gradually increases from -446 to -422 mV vs. Ag/AgCl by increasing the L¹ concentration in the range of 0.2–2.0 mM. The positive shift of the OCP can be attributed to the terms of the adsorption of inhibitor molecules on the C-steel layer [35]. It is supposed that the anodic reactions like iron oxidation in the presence of inhibitors can retard under OCP conditions [36]. These results confirm that a higher concentration of L¹ promotes the inhibition efficiency of samples in a 1.0 M HCl solution.

While the OCP variation with time follows from Fig. 1b that the addition of inhibitor L² to the 1.0 M hydrochloric acid solution generates new characteristics in the OCP curve. The OCP value for C-steel in three different concentrations of inhibitor L² (e.g. 0.2, 1.0 and 2.0 mM) initially decreases from c.a. -420 mV Ag/AgCl to a minimum value of -440 mV vs. Ag/AgCl for 200 s and then stabilizes. These findings could be explained by the inhibition of the cathodic reaction on the C-steel surface for the first 200 s after electrode immersion into 1.0 M HCl solution. It is evident that the slow increase in OCP value at a concentration of 1.0 mM and 2.0 mM of L² inhibitor may be attributed to the anodic reaction being inhibited after 200 s [37]. Similar to the L¹-containing solution, the OCP of the C-steel in the L²-containing solutions show a positive shift compared with the inhibitor-free solution. This positive shift in the value of the OCP was discovered to enhance with an increase in the concentration of L². The maximum change in OCP was found to be 26 mV at a concentration of 2.0 mM for inhibitor L^2 . The OCP of the C-steel in the L²-containing solutions is in the potential range of -445 to -437 mV vs. Ag/AgCl, which shows a narrow potential range



Fig. 2. Tafel diagram for C-steel in 1.0 M HCl solution in the absence and various concentration inhibitors of L^1 and L^2 .

compared to the L¹-containing solutions. Like the result of the L¹, a 1.0 M HCl solution containing L² has a higher OCP compared to L²-free solution. The increase in OCP values by the addition of L¹ and L² inhibitors as compared to inhibitor-free demonstrates the importance of inhibition performance, which helps to protect the C-steel surface from the corrosion process.

3.2.2. Potentiodynamic polarization

Tafel diagram, as presented in Fig. 2, can be used to determine electrochemical parameters, including the corrosion current density, i_{corr} , and corrosion potential of C-steel (E_{corr}). Additionally, the cathodic coefficients (β_c) and the apparent anodic coefficients (β_a) are calculated by the aid of the slopes of the Tafel diagram and Tafel equations, as follows:

$$\log i_{ox} = \log i + \frac{\alpha_a n F E_{corr}}{2.3 R T}$$
(1)

$$\log i_{red} = \log i - \frac{\alpha_c n F E_{corr}}{2.3 RT}$$
⁽²⁾

where i_{red} and i_{ox} are the reduction and oxidation current densities, while *n*, *F*, *T*, and *R* demonstrate the number of transferred electrons, faradaic constant, absolute temperature (K), and ideal gas constant, respectively. The Tafel curves of the C-steel in the absence and presence of inhibitors revealed similar polarization diagrams, implying no considerable difference in the polarization result between the inhibitorfree and inhibitor-containing solutions.

The i_{corr} for C-steel is precisely the evidence of its corrosion damage. As a result, an electrode with a higher i_{corr} will corrode faster than a lower i_{corr} at fixed E_{corr} [38,39]. As shown in Table 2, the i_{corr} of the C-steel electrode in 1.0 M HCl solution is 2.33×10^{-4} A/cm² representing the highest value among the other inhibitor-containing

solutions. The i_{corr} for C-steel in the L¹-containing solution decreases gradually from 1.43×10^{-4} to 0.91×10^{-4} A/cm² by raising the concentration of L¹ from 0.2 to 2.0 mM. Table 2 shows that the L¹ has lower i_{corr} compared to L² at all concentrations, which could be due to the polarized functional groups of C=O and the NH₂ with better adsorbing capability on the steel surface compared to the CH₂-NH group of lower polarity in L^2 . It is seen that the addition of the reduced form of L^1 in the 1.0 M HCl solution provided a superior result with the best resistance to corrosion as compared to the L¹-free and containing solutions. Along with the increase of the L^2 concentration from 0.2 to 2.0 mM, the i_{corr} is reduced by about 41%. A similar conclusion can be drawn by using the results of corrosion rate, in which the C-steel showed a higher corrosion resistance in the L²-containing solution in comparison to the inhibitor-free and L¹-containing solutions. By Assessing Table 2, resistance polarization (R_P) of the C-steel in the free-inhibitor solution is the lowest one (108.73 Ω cm²) compared with the C-steel in inhibitor-containing solutions. It is again confirmed the lowest corrosion resistance among the other samples. The more concentration of inhibitors L¹ and L² leads to the high values of R_p. In addition, the inhibitor L^2 shows higher R_p than that L^1 in the equal concentration, indicating the higher corrosion performance C-steel in the L²-containing solution.

From Table 2, at all solutions (except inhibitor-free solution), β_a is greater than β_c . If β_a is greater than β_c , the double layer formed on the electrode surface shows the p-type semiconductor behavior [40]. Therefore, the double layer formed on the C-steel surface in inhibitor-containing solutions acts as p-type semiconductor behavior. Generally, both apparent charge transfer coefficients decrease by increasing the concentration of inhibitors. In addition, it is reported that a reduced form of the inhibitor has higher stability hydrolytically in hydrochloric acid media because the reduced amine does not endure the hydrolytic process [15,24]. In this study, the imine bond (-C=N-) is converted to an amine bond (-C-NH) by the reduction process.

By assessing the obtained data, the change in E_{corr} between the Csteel electrode in the presence inhibitor with respect to the blank 1.0 M HCl was less than 20 mV for all concentrations. Previously published stated if the difference in E_{corr} was less than 85 mV, the inhibition mechanism could be recognized as a mixed-type, showing that the process rate of anodic inhibition was nearly equal to the cathodic reactions [41,42]. However, based on the OCP and Tafel polarization measurements for the C-steel in the absence and presence of inhibitors, it is believed that L¹ and L² act as mixed a mixed type system with anodic inhibition predominant.

3.2.3. EIS study

The simulated and measured Nyquist impedance spectra of C-steel at the OCP in the absence and presence of inhibitors, L^1 and L^2 , are shown in Fig. 3a and b. Only one time constant can be seen in the testing frequency range for the samples in 1.0 M HCl with and without inhibitors, which is generally attributed to a process that mainly corresponds to a charge transfer phenomenon on the electrode surface. The Nyquist diagram obtained in different concentrations of L^1 and L^2 is attributed to the capacitive impedance of a double–layer with a charge transfer resistance on the surface of the C-steel electrode. It is worth mentioning that the shape of Nyquist diagrams is a depressed semi-circle and accommodates the non-ideal behavior of a capacitance. This phenomenon

Table 2

Tafel diagram data for C-steel in 1.0 M HCl solution in the absence and various concentration inhibitors of L¹ and L².

•							
Sample	Conc. (mM)	E _{corr.} (mV)	i _{corr.} (A/cm ²)	β_a (V/dec)	$-\beta_c$ (V/dec)	$-\beta_a/\beta_c$	$R_P (\Omega.cm^2)$
1.0 M HCl	0	-448.80	$2.33 imes10^{-4}$	0.111	0.123	0.90	108.73
L^1	0.2	-449.75	1.43×10^{-4}	0.130	0.097	1.34	168.68
	1.0	-445.07	1.24×10^{-4}	0.140	0.096	1.45	199.42
	2.0	-446.37	0.91×10^{-4}	0.130	0.113	1.15	288.46
L^2	0.2	-464.11	1.25×10^{-4}	0.160	0.106	1.51	221.48
	1.0	-464.07	$0.88 imes10^{-4}$	0.149	0.087	1.71	271.03
	2.0	-464.73	$0.74 imes10^{-4}$	0.139	0.093	1.49	326.95



Fig. 3. Simulated and measured Nyquist impedance spectra of C-steel at OCP in 1.0 M HCl at different concentrations of (a) L¹ and (b) L², and (c) an equivalent circuit on the electrode surface.

Table 3					
EIS parameters	for an	equivalent	circuit	of Fig.	3c

Inhibitor type	Concentration (mM)	R _{sol.}	$R_{ct} (\Omega cm^2)$	Q _{dl}		χ^2
		$(\Omega \text{ cm}^2)$		$Y_0 (\Omega s^{-n})$	n	
1.0 M HCl	0.0	2.41	126.06	$1.31 imes 10^{-4}$	0.89	0.013
L ¹	0.2	2.41	585.60	1.35×10^{-4}	0.75	0.001
	1.0	2.00	868.21	1.18×10^{-4}	0.75	0.009
	2.0	2.98	1098.00	$1.07 imes10^{-4}$	0.76	0.006
L ²	0.2	7.19	912.20	$1.08 imes10^{-4}$	0.78	0.004
	1.0	4.94	1136.10	$1.01 imes 10^{-4}$	0.77	0.008
	2.0	6.73	1379.40	1.00×10^{-4}	0.75	0.002

illustrates using constant phase element (CPE), which includes two electrochemical parameters; Y_0 (the admittance of CPE) and n (empirical constant). The value below 1.0 arises from the roughness of the C-steel surface and inhomogeneity owing to the inhibitors adsorption together with the corrosion products [8].

In addition, an equivalent electrical circuit is depicted in Fig. 3c and its electrochemical parameters are shown in Table 3. An equivalent electrical circuit designed to fit EIS diagrams, contains a resistance of 1.0 M HCl solution ($R_{sol.}$) in series with Q_{dl}/R_{ct} . As already known, charge transfer resistance is a characteristic quantity for corrosion resistance; the higher the charge transfer resistance, the higher the corrosion resistance. From charge transfer resistance values given in Table 3, the resistance obtained in the presence of L^1 was quite higher than that in the absence of L^1 (126.06 Ω cm²). Also, the trend for the inhibition efficiency for L^1 follows the order of 2.0 mM > 1.0 mM > 0.2 mM. The charge transfer resistance for C-steel in the L¹-containing solution increases significantly from 585.6 to 1098.0 Ω cm² by promoting the concentration of L¹ from 0.2 to 2.0 mM. It is apparent in Table 3 that with increasing the L^2 concentration, an increment in the value of R_{ct} is observed (912.20–1379.40 Ω cm²). As a result, the EIS data for inhibitors confirms that the L²-containing solutions have much better corrosion resistance of the C-steel than the L¹ compound.

The obtained results in EIS measurements are in good accordance with the Tafel experiments. Compared to EIS and Tafel data, the polarization resistance of C-steel in inhibitor-free solution demonstrates the minimum value. While, the R_{ct} and R_P values for inhibitor L¹ and L² rise meaningfully in the range of 0.2–2.0 mM in 1.0 M HCl solution, confirming the improvement of the inhibition performance at higher concentration. The charge transfer resistance observed for L¹ and L² was

similar to that observed in Tafel experiments, so as to C-steel electrode in the L²-containing solution demonstrates higher R_{ct} than that L¹ in the equal concentration. It is clear that both EIS and Tafel experiment confirm that L² inhibitor has higher corrosion resistance at a maximum concentration (2.0 mM), thanks to its higher R_p and R_{ct} makes its use when the environment is too corrosive and aggressive. A similar view has been reported previously that the higher concentration of inhibitors causes the higher charge transfer resistance for metal electrodes [38,43, 44].

Generally, an increase in inhibitor concentration leads to an increment in the thickness and surface coverage inhibitor in the steel surface, which effectively adsorbed on the electrode surface [39]. On the other hand, the inhibitor molecules replace water ions on the C-steel surface [45]. Based on the FTIR results, the other reason for this phenomenon can be their inherent feature in that N-containing functional groups play as electron donors to the electrode surface [16]. The existence of hetero elements and aromatic rings such as nitrogen and sulfur on the structure of amino acids causes a weighty increment in inhibition efficiency [46].

The lower capacitance of the double-layer can be evidence of the higher efficiency of the inhibitor performance. The value of the double-layer capacitance is calculated by the following equation [47]:

$$C_{dl} = \left(\frac{Y_0}{R_{c-1}^{d-1}}\right)^{\frac{1}{n}}$$
(3)

where C_{dl} is capacitive of double layer, Y_0 represents the value of CPE, R_{ct} is the charge transfer resistance, and n is the phase exponent which is always in the range of 0–1. As the following equation, C_{dl} is conversely related to the thickness of the double layer, which serves as a barrier layer for protection from corrosion.

The capacitance of double layer and inhibition efficiency values of C-steel in 1.0 M HCl solution containing different concentrations of inhibitors.

Inhibitor type	Inhibitor-free	L^1			L^2			
	1.0 M HCl	0.2 mM	1.0 mM	2.0 mM	0.2 mM	1.0 mM	2.0 mM	
C _{dl} (μF cm ⁻²) IE (%)	79.0 -	57.94 78.47	55.21 85.48	54.41 88.51	56.18 86.18	52.17 88.90	51.67 90.86	



Fig. 4. Tafel diagram data for C-steel in 3.5% NaCl solution in the absence and presence of 2.0 mM inhibitors of L^1 and L^2 .

$$C_{dl} = \frac{\varepsilon \varepsilon_0 A}{d} \tag{4}$$

where d is the thickness of the electrical double layer, ε_0 is vacuum permittivity, ε is dielectric constant, and A is the electrode area in the electrolyte. It is clear from the data in Table 3 that an increase in L¹ concentration leads to a gradual decrease in the value of double-layer capacitance (57.94–54.41 $\mu F~cm^{-2}\mbox{)},$ which implies a reduction in dielectric constant or/and increase in the double layer thickness [8]. This phenomenon reflects that the L¹ inhibitors' presence led to the decrement in the capacitance of the double-layer due to the replacement of H₂O molecules by inhibitor ions at the surface of the C-steel electrode. As a result of the inhibitors adsorption on the electrode surface, the effective value of the area (A), which can intensify the corrosion rate, decreases, and the efficiency of surface protection is increased. The adsorption of L¹ molecules on the electrode surface acts as a barrier for ions and charge transfers between metal and electrolyte, thus enhancing the protecting surface from corrosion. These findings confirm the data obtained in EIS analysis, in which higher charge transfer resistance led to higher protection efficiency while the C_{dl} for C-steel in the L²-containing solutions decreases from 56.18 to 51.67 $\mu F\,cm^{-2}$ with increasing the inhibitor concentration, which demonstrates lower C_{dl} values than the L¹-containing solutions. Therefore, C-steel shows high corrosion resistance in the L^2 -containing solutions compared to the L^1 .

Table 4 also shows the inhibition efficiency (IE) calculated from charge transfer resistance values obtained in EIS diagrams shown in Fig. 3 as follows:



Fig. 5. Nyquist impedance spectra and equivalent circuit of C-steel electrode at OCP in 3.5% NaCl solution in the absence and presence of 2.0 mM inhibitors of L^1 and L^2 .

$$IE(\%) = \frac{R_{ct} - R_{ct}^0}{R_{ct}} \times 100$$
(5)

where R_{ct} and R_{ct}^0 present the charge transfer resistance of C-steel with and without inhibitors, respectively. The IE increased by adding more L¹-inhibitor with a maximum IE of 88.51% at 2.0 mM concentration. As shown in Table 4, there is a gradual increment in the inhibition efficiency of the L¹-containing solution from 78.47% to 88.51%, which represents only a 10.04% improvement in corrosion protection by increasing inhibitor concentration. As shown in Table 4, the IE obtained in the presence of L² is far greater than the values obtained in the L¹containing solution. Again, these results suggested that the L¹ is hydrolyzed by HCl into its precursor molecules, C=O and the NH₂ with higher polarity has better absorbance onto the steel surface which increases the C_{dl} value and is slightly higher than the L² with lower polarity and adsorbing capability.

3.3. Electrochemical measurements in 3.5 wt % NaCl

Fig. 4 shows Tafel diagram data for C-steel in 3.5% NaCl solution in the absence and presence of 2.0 mM inhibitors of L^1 and L^2 . The corrosion parameters including i_{corr} , E_{corr} , β_a and β_c and R_p were measured from the Tafel curves shown in Table 5. The E_{corr} of C-steel in the free-inhibitor solution (3.5% NaCl) was -871.03 mV vs. Ag/AgCl

Table 5

Tafel diagram data for C-steel in 3.5 wt % NaCl in the absence and presence of 2.0 mM inhibitors of L^1 and L^2 .

0		· · · · · · · · · · · · · · · · · · ·				
Sample	E _{corr.} (mV)	i _{corr.} (A/cm ²)	β_a (V/dec)	$-\beta_c$ (V/dec)	$-\beta_a/\beta_c$	$R_P (\Omega.cm^2)$
3.5% NaCl	-871.03	3.89×10^{-4}	0.017	0.375	0.046	18.45
2.0 mM L ¹	-910.15	$1.78 imes10^{-4}$	0.053	0.630	0.084	119.14
2.0 mM L ²	-935.34	$1.25 imes 10^{-4}$	0.039	0.539	0.072	126.70

EIS parameters for an equivalent circuit of Fig. 5

Sample	$R_s (\Omega.cm^2)$	$R_{ct} (\Omega.cm^2)$	Q _{dl}		W (Ω .s ^{-0.5})	χ^2
			$Y_0 (\Omega.s^{-n})$	n		
3.5% NaCl	9.8	26.3	0.099	0.47	1.9	0.025
2.0 mM L ¹	16.1	43.2	0.140	0.45	1.4	0.023
2.0 mM L ²	17.2	51.5	0.110	0.38	1.1	0.015

which is more positive than C-steel in containing-inhibitor solutions. Like the inhibition mechanism of L¹ and L² in 1.0 M HCl, the mixed-type mechanism controls the inhibiting process on the C-steel surface in 3.5% NaCl solution, due to the E_{corr} difference being lower than 85 mV [41, 42]. It is fair to say that the inhibition mechanism in C-steel is similar in both 1.0 M HCl and 3.5% NaCl solutions. From Table 5, the icorr of C-steel in 3.5% NaCl was 3.89×10^{-4} , the highest one compared with 2.0 mM L^1 and L^2 -containing. As C-steel showed the highest corrosion resistance in 1.0 M HCl solution, the same trend can be seen for C-steel in NaCl solution, in which the i_{corr} for inhibitor L² illustrated by 67.9% and 29.8% reduction in corrosion rate compared with the absence inhibitor and 2.0 mM L¹. Moreover, a big difference between the potentiodynamic curve of C-steel in 1.0 M HCl and 3.5%NaCl is the formation of a passive region in sodium chloride solution. At the range -750 to -450 mV vs. Ag/AgCl, in the anodic branch, corrosion rate directly enters the passive region by increasing the applied potential, indicating a hydroxide iron formation on the C-steel surface in contact with the solution [48]. Again, resistance polarization (R_P) of the C-steel in the free-inhibitor solution showed the lowest one (18.45 Ω cm²) compared with the C-steel in inhibitor-containing solutions. In contrast to the results of the 1.0 M HCl solution, at all solutions, β_a is lower than β_c .

Fig. 5 shows the Nyquist impedance spectra and equivalent circuit of C-steel electrode at OCP in 3.5% NaCl solution in the absence and presence of 2.0 mM inhibitors of L^1 and L^2 . As shown in Fig. 5, all Nyquist shows a depressed circle at high frequency and a straight line at low-frequency range, indicating a completely different corrosion mechanism compared with C-steel in 1.0 M HCl. The first semi-circle is generally related to the charge transfer phenomenon. The second part corresponds to the diffusion process through the iron hydroxide layer (passive film) on the surface [49].

Moreover, the Nyquist plots were simulated by an equivalent circuit as shown in Fig. 5 and their electrochemical were demonstrated in Table 6. The circuit consists of a solution resistance (R_s) in series with a parallel capacitive loop charge transfer resistance and capacitance (R_{cl} /



Fig. 6. Corrosion rate of C-steel in the absence and presence of inhibitor L^1 and L^2 solution measured by weight loss experiments.

 Q_{cl}) and a parallel Warburg element. From R_{ct} values given in Table 6, C-steel showed the lowest value (26.3 Ω cm²). On the other hand, the trend for the inhibition efficiency for C-steel follows the order of 2.0 mM L^2 > 2.0 mM L^1 > inhibitor-free. Therefore, maximum inhibition efficiency can be found in the L²-containing solution for C-steel. Similar to EIS results in HCl, it can be said that the L²-containing solutions have much better corrosion resistance for the C-steel than that of the L¹ compounds. The EIS results are in good agreement with the Tafel data in 3.5% NaCl solution. The Warburg element in the equivalent circuit is related to the diffusion process through passive films, which acts as a barrier in mass



Fig. 7. (a) SEM image of the as-received C-steel surface, and SEM image of C-steel taken after immersion in (b) 1.0 M HCl (inhibitor-free), (c) 2 mM inhibitor L^1 and (d) 2 mM inhibitor L^2 .



Fig. 8. EDS analysis of the as-received C-steel before dipping in the electrolyte, and after immersion in 1.0 M HCl (inhibitor-free), 2.0 mM inhibitor L^1 and (d) 2.0 mM inhibitor L^2 .

transfer. A lower value of Warburg indicates a barrier effect of the passive film slowing down the diffusion of corrosive ions [50]. The Warburg values increased with decreasing inhibition efficiency due to the greater aggressive corrosion agent diffusing through the passive film. Such an increase can be observed for C-steel in free-inhibitor 3.5% NaCl solution.

3.4. Immersion time measurements

The weight loss measurement was used to determine the corrosion rate of C-steel in 1.0 M HCl solution in the absence and presence of inhibitors L^1 and L^2 (as depicted in Fig. 6). It is a way to confirm the obtained data from electrochemical measurements on a time scale as well as the stability of inhibitors. The corrosion rate for C-steel in inhibitorfree solution shows the highest value compared with samples in inhibitor-containing solutions. The obtained data from the immersion test for the sample in 1.0 M HCl is well accordance with our finding in Tafel and EIS measurements, indicating a minimum corrosion resistance for the inhibitor-free solution. As illustrated in Fig. 6, the addition of both inhibitors L¹ and L² cause a dramatic decrease in the corrosion rate of C-steel in 1.0 M HCl solution, indicating the improvement in corrosion behavior even a long time. In addition, the corrosion rate decreases gradually from 0.27 to 0.16 mg cm⁻² h^{-1} for inhibitor L^1 and from 0.22 to 0.13 mg cm⁻² h^{-1} for inhibitor L^2 . Similar results have been reported by previous studies [43,51]. It is fair to say that all corrosion rate data have confirmed the EIS and Tafel polarization experiments at different concentrations of inhibitors.

3.5. Surface characterization

SEM images in Fig. 7 show the corrosion inhibition efficiency of the inhibitors L¹ and L² on the C-steel surfaces exposed in 1.0 M HCl. Fig. 7a shows the as-received C-steel before dipping in the electrolyte, which depicts a smooth and clear surface. In an inhibitor-free solution, the surface severely deteriorated at the end of the 24 -hour immersion in 1.0 M HCl, which was confirmed by irregular and rough surface (Fig. 7b). As the inhibitors L¹ and L² add to the 1.0 M HCl solution, the corrosion degree of the C-steel surface is decreased which is shown in Fig. 7c and d. This confirmed that the inhibitor-containing solution protects the C-steel surface against corrosive species. It is apparent that the reduced form of L¹ in the 1.0 M HCl solution, confirming a relatively smoother and cleaner surface.



Fig. 9. Langmuir adsorption isotherms involving L¹ and its reduced form (L²).

Fig. 8 shows the EDS profile analyses of as-received C-steel before dipping in the electrolyte, the sample immersed in 1.0 M HCl, 2.0 mM L¹ and the L²-containing solutions. The EDS profile obtained on the C-steel surface illustrates that the main peak is iron and carbon, representing the steel composition. The surface of the C-steel immersed in the 1.0 M HCl shows one more peak, which corresponds to oxygen. The additional peak owing to oxygen indicates the formation of the steel corrosion product on the surface. The EDS spectrum of C-steel exposed in the 1.0 HCl solutions in the presence of 2.0 mM L¹ and L² do not demonstrate the oxygen peak, implying the protective layer of inhibitors on the C-steel surface. The results confirmed that the addition of L¹ and L² in 1.0 M HCl effectively hinders the corrosion of the C-steel.

3.6. Adsorption mechanism

The adsorption of inhibitors on the surface of the C-steel electrode is the vital step in an inhibition mechanism [17]. The main reason for higher corrosion protection in the presence of inhibitors is the adsorption of this material on the C-steel surface. Based on the previous studies [15,39], the Langmuir adsorption model is the best model to explain the adsorption process of the inhibitors of L^1 and L^2 on the surface of C-steel in 1.0 M HCl. The Langmuir adsorption isotherm, which is based on a simple kinetic model for monolayer adsorption on the surface, is presented as follows:

$$\frac{C_{inh}}{\theta} = \frac{1}{K_{ads}} + C_{inh} \tag{6}$$

where K_{ads} the adsorption equilibrium constant, C_{inh} is the concentration of inhibitor, and θ is the fraction of surface coverage by the inhibitor molecule, which is calculated by the following equation [38]:

$$\theta = 1 - \frac{C_{dl}^i}{C_{dl}^b} \tag{7}$$

where C_{dl}^i is the capacitive response of the carbon steel electrode resulting in the inhibitor-containing and C_{dl}^b is capacitive of the double layer in the free-inhibitor solution. Langmuir relationship, a plot of C_{inh}/θ against C_{inh} should yield a straight line via the regression coefficient, R^2 (shown in Fig. 9). The R^2 values give a good agreement between the EIS and Langmuir data for L^1 ($R^2 = 0.9996$) and L^2 ($R^2 = 0.9999$) inhibitors. The near-unity slope for L^1 and L^2 -containing solutions (i.e., slope = 1.12 and 1.05, respectively) supports the suitability of the

The calculated quantum chemical parameters of inhibitors.

	E _{HOMO}	E _{LUMO}	Ι	А	ΔΕ	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3LY	/P/6-31+g LE	VEL											
1	-5,4274	-0,8675	5,4274	0,8675	4,5598	2,2799	0,4386	3,1474	-3,1474	2,1725	0,4603	4,7422	-41884,5606
2	-5,2581	-0,5018	5,2581	0,5018	4,7563	2,3782	0,4205	2,8799	-2,8799	1,7438	0,5735	4,0596	-41917,1468
HF/6	5-31+g LEVE	L											
1	-8,0642	2,5987	8,0642	-2,5987	10,6629	5,3314	0,1876	2,7327	-2,7327	0,7004	1,4278	2,4128	-41605,3465
2	-8,2315	3,5144	8,2315	-3,5144	11,7459	5,8729	0,1703	2,3586	-2,3586	0,4736	2,1115	4,1761	-41636,1458
M06	2X/6-31+g L	EVEL											
1	-7,0269	0,1039	7,0269	-0,1039	7,1308	3,5654	0,2805	3,4615	-3,4615	1,6803	0,5951	4,4003	-41866,4346
2	-6,9373	0,5382	6,9373	-0,5382	7,4756	3,7378	0,2675	3,1995	-3,1995	1,3694	0,7302	3,8441	-41899,0131

Table 8

The calculated quantum chemical parameters of inhibitors.

	E _{HOMO}	E _{LUMO}	I	А	ΔΕ	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3LY	/P/6-31++g L	EVEL											
1	-5,7193	-1,1875	5,7193	1,1875	4,5318	2,2659	0,4413	3,4534	-3,4534	2,6316	0,3800	4,9934	-41886,1091
2	-5,5803	-0,9121	5,5803	0,9121	4,6681	2,3341	0,4284	3,2462	-3,2462	2,2574	0,4430	4,0488	-41918,7373
HF/	5-31++g LEV	EL											
1	-8,6508	0,8934	8,6508	-0,8934	9,5442	4,7721	0,2096	3,8787	-3,8787	1,5763	0,6344	2,8098	-41605,9430
2	-8,3850	0,8906	8,3850	-0,8906	9,2756	4,6378	0,2156	3,7472	-3,7472	1,5138	0,6606	4,1414	-41637,1374
M06	2X/6-31++g	LEVEL											
1	-7,2459	-0,2634	7,2459	0,2634	6,9825	3,4912	0,2864	3,7547	-3,7547	2,0190	0,4953	4,5038	-41867,6736
2	-7,1863	-0,4338	7,1863	0,4338	6,7526	3,3763	0,2962	3,8100	-3,8100	2,1498	0,4652	3,7428	-41900,3243

Table 9

The calculated quantum chemical parameters of inhibitors.

	E _{HOMO}	E _{LUMO}	Ι	А	ΔE	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3L	/P/6-31++g (d	1,p) LEVEL											
1	-5,9373	-1,1712	5,9373	1,1712	4,7661	2,3831	0,4196	3,5542	-3,5542	2,6505	0,3773	4,4151	-41899,6892
2	-5,8804	-0,8955	5,8804	0,8955	4,9849	2,4924	0,4012	3,3880	-3,3880	2,3026	0,4343	3,7854	-41932,2514
HF/	5–31++g (d,p) LEVEL											
1	-8,0848	0,9802	8,0848	-0,9802	9,0650	4,5325	0,2206	3,5523	-3,5523	1,3921	0,7184	4,3616	-41625,4290
2	-8,1970	0,8985	8,1970	-0,8985	9,0955	4,5477	0,2199	3,6492	-3,6492	1,4641	0,6830	3,7140	-41656,8335
M06	2X/6-31++g	(d,p) LEVEL											
1	-7,2369	-0,2465	7,2369	0,2465	6,9904	3,4952	0,2861	3,7417	-3,7417	2,0028	0,4993	3,9070	-41880,3065
2	-7,3063	-0,4240	7,3063	0,4240	6,8824	3,4412	0,2906	3,8651	-3,8651	2,1707	0,4607	3,5402	$-41912,\!8041$

Table 10

The calculated quantum chemical parameters of inhibitors.

	E _{HOMO}	E _{LUMO}	Ι	Α	ΔE	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3L	YP/6-31+g LEV	/EL											
1	-6,9420	-4,8219	6,9420	4,8219	2,1201	1,0600	0,9434	5,8819	-5,8819	16,3189	0,0613	33,5474	-41893,5312
2	-7,0315	-4,4850	7,0315	4,4850	2,5465	1,2732	0,7854	5,7582	-5,7582	13,0210	0,0768	23,9338	-41925,5171
HF/	6–31+g LEVEL												
1	-9,4147	-0,0038	9,4147	0,0038	9,4109	4,7054	0,2125	4,7092	-4,7092	2,3565	0,4244	29,0210	-41614,2315
2	-11,0906	0,1951	11,0906	-0,1951	11,2857	5,6429	0,1772	5,4478	-5,4478	2,6297	0,3803	13,4799	-41646,8833
M06	2X/6-31+g LE	EVEL											
1	-8,3044	-3,5122	8,3044	3,5122	4,7922	2,3961	0,4173	5,9083	-5,9083	7,2843	0,1373	33,7350	-41875,1812
2	-8,7749	-3,3696	8,7749	3,3696	5,4053	2,7027	0,3700	6,0723	-6,0723	6,8215	0,1466	22,7482	-41907,2417

Table 11

The calculated quantum chemical parameters of inhibitors.

	E _{HOMO}	E _{LUMO}	Ι	Α	ΔΕ	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3LYP/6-31++g LEVEL													
1	-7,2070	-5,3196	7,2070	5,3196	1,8874	0,9437	1,0597	6,2633	-6,2633	20,7846	0,0481	33,3897	-41894,9229
2	-7,3558	-5,1041	7,3558	5,1041	2,2518	1,1259	0,8882	6,2300	-6,2300	17,2365	0,0580	23,3850	-41926,9453
HF/6–31++g LEVEL													
1	-9,4049	-2,4265	9,4049	2,4265	6,9784	3,4892	0,2866	5,9157	-5,9157	5,0148	0,1994	33,2438	-41614,6495
2	-11,2218	-1,8956	11,2218	1,8956	9,3262	4,6631	0,2144	6,5587	-6,5587	4,6124	0,2168	13,2422	-41647,7705
M062X/6-31++g LEVEL													
1	-8,5096	-4,4679	8,5096	4,4679	4,0417	2,0209	0,4948	6,4887	-6,4887	10,4172	0,0960	33,7687	-41876,4072
2	-9,4705	-3,4986	9,4705	3,4986	5,9719	2,9859	0,3349	6,4845	-6,4845	7,0412	0,1420	12,1952	-41910,7910

The calculated quantum chemical parameters of inhibitors.

	-		-										
	E _{HOMO}	E _{LUMO}	I	А	ΔE	η	Σ	Х	Рİ	ω	ε	dipol	Energy
B3LYP/6-31++g (d,p) LEVEL													
1	-7,1898	-5,0959	7,1898	5,0959	2,0939	1,0470	0,9551	6,1429	-6,1429	18,0211	0,0555	31,9798	-41908,3685
2	-7,3558	-5,1041	7,3558	5,1041	2,2518	1,1259	0,8882	6,2300	-6,2300	17,2365	0,0580	23,3850	-41926,9453
HF/6-31++g (d,p) LEVEL													
1	-9,3009	-2,4561	9,3009	2,4561	6,8448	3,4224	0,2922	5,8785	-5,8785	5,0486	0,1981	32,8781	-41634,2688
2	-10,5687	-2,3620	10,5687	2,3620	8,2067	4,1034	0,2437	6,4653	-6,4653	5,0934	0,1963	22,4671	-41665,0021
M062X/6-31++g (d,p) LEVEL													
1	-8,4892	-4,2970	8,4892	4,2970	4,1922	2,0961	0,4771	6,3931	-6,3931	9,7494	0,1026	32,1212	-41888,8771
2	-9,5129	-2,5783	9,5129	2,5783	6,9346	3,4673	0,2884	6,0456	-6,0456	5,2706	0,1897	11,2549	-41930,2612



Fig. 10. Representations of optimized structures, HOMO, LUMO, and ESP shapes of inhibitor molecules.

Langmuir model for obtained results in the present study.

The value of K_{ads} can be calculated by considering the width of the original value of the C_{inh}/θ against C_{inh} diagrams. The K_{ads} obtained for L^1 and L^2 inhibitors are 49.26 and 102.04 (mM)⁻¹, respectively. The Langmuir K_{ads} value can be interpreted as the adsorption/desorption equilibrium for each inhibitor on the surface of the C-steel. In other words, it describes the fraction of the electrode surface, which is covered by inhibitor molecules. The higher the adsorption equilibrium constant, the more corrosion protection there appears to be.

3.7. Theoretical studies

Today, the use of theoretical approaches to compare inhibitor molecules has become widespread. However, in the calculations made by theoretical methods, many quantum chemical parameters about inhibitor molecules are obtained. These quantum chemical parameters enable us to gain information faster than experimental methods to explain the inhibitory properties of inhibitor molecules. Among these quantum chemical parameters, the two most important parameters are HOMO and LUMO. The HOMO parameter of the molecule shows the ability of inhibitor molecules to donate electrons from the highest energy-filled orbitals to metal atoms. On the other hand, the LUMO parameter of molecules indicates the ability of molecules to receive electrons from metal atoms to the lowest energy empty molecular orbitals. These two parameters either create a chemical interaction by giving electrons from metal atoms to inhibitor molecules for the molecules to hold onto the metal surface, or make a chemical interaction by donating electrons to metal atoms from inhibitor molecules. The numerical values of all calculated parameters with various basis sets are given in Tables 7–12.

The calculations of the molecules were made in different methods and basis sets. Calculations of 6-31+g, 6-31++g, and 6-31++g(d,p)basis sets were made in this method and basis sets B3LYP, HF, M062X method. In these calculations, many quantum chemical parameters of the molecules are calculated. After the heteroatoms in the molecule were examined, the nitrogen atom with the most negative charge was protonated. In experimental processes, the acid molecule in solution causes the protonation of this most negatively charged heteroatom. For this reason, calculations of the protonated states of the molecules were made. As a result of the calculations, it was seen that the inhibitory activity of L² was higher than that of L¹.

Although many parameters are obtained in the calculations, there is a formal demonstration of the two parameters. The first picture in Fig. 10 shows the optimized structures of the inhibitor molecules. In the second and third pictures, there are HOMO and LUMO representations of the inhibitor molecules. The last picture shows which atoms in inhibitor molecules have greater electron density [52].

HOMO and LUMO parameters of molecules are calculated with theoretical calculations. With these calculations, it is possible to comment on the electron exchanges and inhibitory activities of molecules. However, in ESP shapes of molecules, the red colored regions of the molecules are electron-rich regions. The blue colored regions are electron-poor regions. The electron-rich regions of the molecules allow the molecules to show inhibitory activity by donating electrons. However, the electron-poor regions of the molecules allow the molecules to accept electrons and show inhibitory activity. The ESP results of the molecules with the calculations made the nitrogen atom in the center of the L¹ molecule and the oxygen atoms at the end of the molecule are shown in red. These regions are electron-rich regions. On the other hand, it is seen that the same atoms in the L² molecule are colored red. In general, the blue colored regions are on the methyl groups attached to the benzene ring. Since the methyl group is not able to accept electrons, it is believed that the inhibitory mechanism will proceed through the red colored regions. Since the methyl group is not able to accept electrons, it is thought that the inhibitory mechanism will proceed through the red colored regions. This situation shows heteroatoms in molecules, that is, atoms such as oxygen, nitrogen, and sulfur, other than carbon atoms in organic and inorganic compounds. They show inhibitory activity by donating lone electron pairs on these atoms.

4. Conclusion

The corrosion behavior of C-steel in 1.0 M HCl with and without inhibitors was investigated using electrochemical techniques. The main conclusions of the present study are summarized below:

- 1. From Tafel's results, the highest value of *i*_{corr} corresponded to the inhibitor-free solution, suggesting a higher corrosion rate than other inhibitor-containing solutions.
- 2. The i_{corr} for C-steel in the L¹-containing solution decreases gradually by raising the concentration of L¹ from 0.2 to 2.0 mM. It is seen that the addition of a reduced form of L¹ in the 1.0 M HCl solution provided a superior result with the best resistance to corrosion as compared to the L¹-free and containing solutions.
- 3. The addition of L^1 and L^2 inhibitors to the 1.0 M HCl electrolyte decreases the corrosion rate of steel in the electrolyte. The compound L^2 was found to show a better inhibitor characteristic than L^1 due to the higher stability in HCl solution.

- 4. The result showed that the inhibition efficiency of L^2 enhances with increasing the concentration and presented the highest value at a concentration of 2.0 mM (e.g., 90.86%). The same trend can be found in the L¹-containing solution, in which the corrosion rate of the C-steel tends to decrease by an increment of L¹ and presented a minimum rate at 2.0 mM in the range of 0.1–2.0 mM.
- 5. SEM images supported the protection of inhibitors against the corrosion of the C-steel surface in 1.0 M HCl solution.
- Both adsorption of L¹ and its reduced form on the surface of the Csteel followed the Langmuir adsorption isotherm.
- 7. As a result of the theoretical calculations, many quantum chemical parameters have been obtained. Each parameter obtained provides important information about the inhibitory properties of inhibitor molecules. The numerical values of the parameters obtained indicate that the L^2 inhibitor molecule is better than the other.

CRediT authorship contribution statement

Saeid Karimi: Data curation, Methodology, Writing – review & editing. Majid Rezaeivala: Conceptualization, Writing – review & editing. Koray Sayin: Software, Investigation, Validation. Burak Tuzun: Software.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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