Optimization of the Influencing Variables on the Corrosion Property of Steel Alloy 4130 in 3.5 wt.% NaCl Solution

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In this work, the aqueous Lawsonia inermis extract (LI) is investigated as an economic and green deterioration inhibitive formula for steel alloy 4130 in 3.5 wt.% NaCl solutions. The water-based extraction process is considered as one of the cheapest techniques for preparation of active ingredients of natural products. These ingredients play an important role in corrosion mitigation of steel alloy 4130 in saline media. This extract was subjected to three different parameters: inhibitor concentration, rotation speed, and temperature in 3.5 wt.% NaCl solutions. The electrochemical techniques are used to perceive the corrosion behavior, and the obtained results were dedicated to theoretical explorations to assess the features of corrosion inhibition and the adsorption over the steel substrate in 3.5 wt.% NaCl solutions. Affording to the electrochemical techniques of LI showed very promising results against corrosion depending on the inhibitor concentrations. The inhibition efficiency of LI was additionally appraised at three diverse temperatures, and the results disclosed that the inhibition efficiency is decreased. Additionally, the theoretical aspects illuminated that the main active ingredients of LI have a proclivity to coagulate on the steel substrates allowing these areas to paradigm a protecting layer on the steel surfaces. This behavior is in provision of investigational results. Statistical studies were used to examine the consequence of chief constraints (i.e., inhibitor concentration, temperature, and rotation speed) on the inhibition efficiency and the rate of corrosion of steel alloy 4130. The inhibitive effect of LI in contradiction of the corrosion of steel alloy 4130 surfaces is considered by resources of DFT/6-31G(d) calculations. The quantum chemical parameters interrelated to the inhibition efficiency are considered.

1. Introduction

Currently, 3.5 wt.% NaCl solutions are commonly utilized in various investigations due to the broad-spectrum existence in industrial zones [1]. It is well known that this medium is considered as an unpleasant medium which strictly deteriorates the metals and alloys, generating growth of more stable layers by direct attack with the surrounding constituents [2, 3]. Consequently, the unfavorable deterioration procedure conveys approximately severe cost through the energy loss and failure of metallic structures. Throughout current years, emerging approaches for shielding of metals in contradiction of decomposition have been a significant area of exploration. Generally, corrosion inhibitors are the greatest common and economic technique for monitoring the corrosion of metals against aggressive chloride media [4–7]. It has been verified that the organic compounds including atoms such as N, O, P, and S in extra exist in aromatic rings and binary or tripartite bonds. All these factors can make the organic molecule to act as efficient corrosion inhibitors. Unfortunately, utmost of these inhibitors display poisonous action in contradiction to the environment [8–12]. The green chemistry has been speedily developed to diminish the antagonistic possessions...
accompanying with the toxic organic inhibitors. Natural products are commonly used to get the alternative source for the toxic compounds. Diverse fragments of natural products have been earlier laboring and used for monitoring the deterioration of metallic substrate. Based on the source and the mode of action, the safe and green chemicals are obtainable in the formulae of biopolymeric forms or in the form of ionic liquids [13]. The Pedalium murex L extract is used to protect the steel surface in pickling electrolyte and conveyed an efficacy of about 89% [14]. With a concentration of about 1000 ppm, it shows an inhibition efficiency of about 93% for protection of mild steel in acidic solution [15]. In addition, the optimum concentration of Salvia officinalis alcoholic extract was applied to protect of the carbon steel in an acidic solution with inhibition efficiency of about 87% [16]. The leaf extract of Cinnamomum verum is used to protect mild steel in acidic environment with an inhibition efficiency of about 93.1% [17]. The inhibition efficiency of 5 g/l from Akeek apple seed extract was investigated for controlling the decomposition of the steel in a medium of HCl and experimentally reported the inhibition efficiency of about 86.9% [18]. The Achyranthes aspera extract was used to protect mild steel in an acidic medium with 90% inhibition efficiency [19]. The root of Glycyrrhiza glabra is applied as green corrosion inhibitor with 99% inhibition efficiency after immersion of steel coupon for 72 h in 3.5 wt.% NaCl solution. In case of 800 ppm from leaves of Glycyrrhiza glabra, the inhibition efficiency is decreased to 88% after immersion of mild steel in acidic medium for 24 h [20]. In case of 300 ppm of Musa paradisiaca extract, the inhibition efficiency is reported to be 82% in acidic solution [21]. The peel extract of Musa paradisiaca was reported to be 89% in acidic medium after 24 h immersion [22]. At the present time, several factors play an important role in assortment of a corrosion inhibitor for the application like efficiency, cost effective, availability, and ecological impact must be considered. Additionally, water solubility of corrosion inhibitors is a significant representative in their abstraction procedure. In case of abstraction of the safe and green corrosion monitoring compounds with partially soluble in water, an extra solvent should be used but not recommended because it is not safe and executes undesirable ecological possessions. Water extract is very an important part from environmentally friendly aspects since the procedure depends on water which is more cost-effective if equated with other solvents. Corrosion is one of the main factors for the degradation of an engineering structure in marine environments, occurring due to chemical or electrochemical reactions between the marine environment and metals. Numerous curricula of organic composites are extensively applied for controlling the metallic corrosion in acidic atmospheres [23–27]. Investigational resources are convenient in explaining the mode of action of inhibition process; nonetheless, these techniques are frequently none economic and wasting time. Continuing computer-based techniques have been used for controlling the use of hypothetical understanding in the study of corrosion monitoring. Accordingly, several techniques such as quantum chemistry and molecular modeling have been achieved to associate the inhibiting performance to the molecule’s property of LI [28, 29]. Application of hypothetical parameter shows two chief compensations: (i) the mixtures with their several parts of substituents can be categorized by their molecular structures; (ii) the planned mode of action of interface can be attributed for the reactivity of chemical compounds [30–33]. By bearing in mind these aspects, the present study was planned to use the aqueous Lawsonia inermis extract (LI), which is cost-effective, for monitoring the decomposition of mild steel alloy 4130 in 3.5 wt.% NaCl solutions. Lawsonia inermis extract comprises organically active mixtures with biological and medical values and potential corrosion inhibitor for metallic substrates. The electrochemical techniques were used to calculate the inhibition efficiency of the inhibitor. The morphology after exposure in 3.5 wt.% NaCl solutions of steel alloy 4130 surfaces was analyzed without and with gradual concentrations of green inhibitor. The optimum conditions among the variable parameters (i.e., concentration of inhibitor, rotation speed, and temperature) which effect on the corrosion rates and inhibition efficiencies were tested by utilization of investigational arithmetical enterprise (Box–Behnken). Additionally, the theoretical assessments were studied for the adsorption landscapes of the investigated compound over the steel alloy 4130 substrate and their reactive sites. Remarkable found that the oxygen containing compound, which promotes more inhibiting activity, is among diverse ingredients of LI.

2. Materials and Methods

2.1. Materials

2.1.1. Steel Alloy 4130. Inhibition possessions of Lawsonia inermis extract (LI) were inspected on the steel alloy 4130 electrode (1 cm × 1 cm) with composition as mentioned in Table 1. The electrodes were graceful using SiC papers (several grades). The refined sample was washed away with bidistilled water and formerly dried out. The destructive 3.5 wt.% NaCl solutions protected by LI at several concentrations were equipped by NaCl (Merck Co.) and bidistilled water.

2.1.2. Lawsonia inermis Extract (LI) Preparation. Water has been commonly employed as an appropriate solvent for abstraction of inhibitor from the dried Lawsonia inermis powder. Water is used a proper solvent for the two main reasons: (i) make the extraction process cost-effective and environmentally acceptable; (ii) only the water-soluble components can be extracted from Lawsonia inermis during the water extraction process. These complexes do not demonstrate good controlling rate in destructive media. 100 g/100 ml distilled water of Lawsonia inermis powder was poured into 1000 ml beaker, boiled for 30 min, and then filtered, and the filtrate is concentrated to 100 ml. The resulting clear solution was then used for preparation of the examination electrolytes in electrochemical studies. The major existing compound in LI is 2-hydroxy-1,4-naphthoquinone (HNQ; lawsone). Lawsone is the standard natural
dye delimited at 1.0–1.4% in the leaves of LI (Henna) [35]. Figure 1 represents the chemical structures of lawsone that is the main component in the aqueous extract of LI [36].

2.2. Methods

2.2.1. Electrochemical Techniques. A GILL AC expedient fortified with gill analyst software was working for EIS tests. The value of frequency was between 0.01 and 10,000 Hz. Also, the Z view software package was castoff for the examination of impedance results. Electrochemical assessments were performed in a common three electrodes including working (steel alloy 4130), counter (platinum), and reference electrode (Ag/AgCl).

2.2.2. Statistical Analysis. For studying the corrosion rate and inhibition efficiency of steel alloy 4130 using of LI as inhibitor, Box–Behnken experimental design [37,38] for the variables (concentration of inhibitor, rotation speed, and temperature) is shown in Table 2. The response variables were developed by designs of the rejoinder superficial outlines and the greatest analytical replicas. The Box–Behnken enterprise can accept the subsequent prototypical formula [39]:

\[
E(y) = \beta_0 + \sum_{i=1}^{3} \beta_i X_i + \sum_{i=1}^{3} \sum_{j=1}^{3} \beta_{ij} X_i X_j,
\]

where \(y\) is the rejoinder adjustable estimation, \(X_i\) are the autonomous parameters (inhibitor load, rotation speed, and temperature) that are recognized for each experimental run, and \(\beta_0, \beta_i, \) and \(\beta_{ij}\) are the worsening limits. Software package, Design-Expert 6.1, Stat-Ease, Inc., Minneapolis, USA, was used for regression examination of investigational statistics and to design rejoinder superficial. The statistical parameters were subjected to approximate the variance analysis (ANOVA). The amount of appropriate investigational consequences to the polynomial prototypical calculation was articulated to determine the coefficient, \(R^2\).

2.2.3. Theory and Computational Details. The DFT method was used to optimize the molecular structure of the investigated compound. The optimization of the total structure together with the vibrational analysis is implementing by revenue of Gaussian 03 package [40]. The molecule was assembled by means of the Gauss View 3.0 instigated in Gaussian 03 package. The adsorption centers of the inhibitor molecule were predicted using frontier molecular orbitals (HOMO and LUMO). Intended for the modest transmission of electrons, the adsorption must happen at the portion of the fragment anywhere the gentleness, \(\sigma\), has the maximum rate. Rendering to Koopman’s theory, the \(E_{\text{HOMO}}\) and \(E_{\text{LUMO}}\) of the corrosion inhibiter formula are connected to the potential of ionization (I) and the affinity of electron (A), correspondingly [41]. The extraquantum parameters of the inhibitor formula that bounce appreciated data about the responsive performance are ionization potential (IP), electron affinity (EA), electronegativity (\(\chi\)), chemical potential (\(\mu\)), hardness (\(\eta\)), and softness (\(\sigma\)). These parameters were designed by the subsequent formula:

\[
I = -E_{\text{HOMO}},
\]

\[
A = -E_{\text{LUMO}},
\]

\[
\mu = -\chi,
\]

\[
\mu = \frac{E_{\text{HOMO}} + E_{\text{LUMO}}}{2},
\]

\[
\eta = \frac{E_{\text{LUMO}} - E_{\text{HOMO}}}{2}.
\]

The softness (\(\sigma\)) is the converse of the universal stiffness and calculated as follows:

\[
\sigma = \frac{1}{\eta}.
\]

The optimization of lawsone compound displayed in Figure 1 was theoretically achieved applying density functional theory (DFT) to measure their electronic possessions. To recognize these atoms, the possible protonated site including O and C atoms with binary bonds (Figure 1) was primary protonated. These protonated molecules were calculated using DFT performance. Subsequently, the subsequent protonated fragments with augmented arrangements and the ethics of protonation affinity (PA) and basicity (B) were examined [42–45].

<table>
<thead>
<tr>
<th>Element</th>
<th>C</th>
<th>P</th>
<th>S</th>
<th>Mn</th>
<th>Si</th>
<th>Mo</th>
<th>Cr</th>
<th>Fe</th>
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<tr>
<td>%</td>
<td>0.28–0.33</td>
<td>0.035</td>
<td>0.04</td>
<td>0.4–0.6</td>
<td>0.2–0.35</td>
<td>0.15–0.25</td>
<td>0.8–1.1</td>
<td>Rem</td>
</tr>
</tbody>
</table>

*Equivalent carbon content 0.664 [34].

*Table 1: Chemical analysis of steel alloy 4130 used in the present study.*
3. Results and Discussion

3.1. Electrochemical Measurements

3.1.1. Open Circuit Potential. Open circuit potential (OCP) magnitudes were examined for the steel alloy 4130 substrate during immersion in 3.5 wt.% NaCl media with diverse LI loads, and the outcomes are offered in Figure 2. Rendering to Figure 2, the OCP of the steel alloy 4130 sampling after immersion in 3.5 wt.% NaCl solutions rapidly declined and touched a steady rate after 200s. The disbanding of oxide layer [46, 47] and the adsorption particles on the substrate of steel alloy 4130 are the foremost explanations for the detected OCP weakening [48–50]. It is well-known that by growing the period of immersion and the LI concentration the OCP values turn out to be more negative, approving the role of foremost cathodic inhibition of the LI. It is also originated that the most negative values of OCP were present in solution injected by 10% of LI, demonstrating the adsorption of LI composites on the steel surface.

3.1.2. Electrochemical Impedance Spectroscopy. Investigation of electrochemical impedance spectroscopy (EIS) was finished to examine the LI inhibition ability on the steel alloy 4130 electrode in 3.5 wt.% NaCl solutions. Nyquist diagrams for steel alloy 4130 electrodes engrossed in the saline solutions that are threatened by diverse LI concentrations that are assumed in Figure 3 authorize a modest capacitive circele for all trials, closefitting that the leading governing response at the boundary of steel surface and 3.5 wt.% NaCl solutions is charge transmission. The loop of capacitance with abundant superior width can be detected in the Nyquist plots of the electrolytes comprising 10% LI compared to the uninhibited solution [51]. In the occurrence of LI inhibitor, a great capacitive loop gives the impression in the electrolyte of low-slung frequency assortment. Figure 4 illustrates the operation of a modest prototypical R(Q) as exemplified in Figure 3 and the corresponding analogous circuit. The impedance conspiracies were form fitting, and the limits obtained are exposed in Table 3. Solitary remark of one-time continual in the Nyquist designs of the reserved trial announces that the constant period which associated with the double layer is determined at a small range of the frequency and protected layer at growth frequency variety. Consequently, in the used model, $R_s$, $R_t$, and $CPE_t$ are correspondingly revealing the solution resistance, total resistance ($R_t$ + inhibitor film resistance ($R_f$)), and the total constant phase element. The efficiency ($\eta$) principles of LI in 3.5 wt.% M NaCl media and diverse loads of LI.

Table 2: Corrosion rate and inhibition efficiency of steel alloy 4130 in 3.5 wt.% NaCl using LI as an inhibitor.

<table>
<thead>
<tr>
<th>Run number</th>
<th>Conc. of inhibitor (%)</th>
<th>Rotation speed (rpm)</th>
<th>Temperature (°C)</th>
<th>Thrust pressure (Pa)</th>
<th>Total time (s)</th>
<th>Corrosion rate (mL/Y)</th>
<th>Inhibition efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5</td>
<td>150</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>23.9</td>
<td>50.51</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>150</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>5.2</td>
<td>89.23</td>
</tr>
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<td>3</td>
<td>5</td>
<td>250</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>28.7</td>
<td>49.47</td>
</tr>
<tr>
<td>4</td>
<td>10</td>
<td>250</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>6.7</td>
<td>88.20</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>200</td>
<td>25</td>
<td>101325</td>
<td>1200</td>
<td>14.9</td>
<td>64.18</td>
</tr>
<tr>
<td>6</td>
<td>10</td>
<td>200</td>
<td>25</td>
<td>101325</td>
<td>1200</td>
<td>4.1</td>
<td>90.14</td>
</tr>
<tr>
<td>7</td>
<td>5</td>
<td>200</td>
<td>75</td>
<td>101325</td>
<td>1200</td>
<td>39.3</td>
<td>42.54</td>
</tr>
<tr>
<td>8</td>
<td>10</td>
<td>200</td>
<td>75</td>
<td>101325</td>
<td>1200</td>
<td>15.3</td>
<td>77.63</td>
</tr>
<tr>
<td>9</td>
<td>7.5</td>
<td>150</td>
<td>25</td>
<td>101325</td>
<td>1200</td>
<td>4.97</td>
<td>87.72</td>
</tr>
<tr>
<td>10</td>
<td>7.5</td>
<td>250</td>
<td>25</td>
<td>101325</td>
<td>1200</td>
<td>5.89</td>
<td>86.71</td>
</tr>
<tr>
<td>11</td>
<td>7.5</td>
<td>150</td>
<td>75</td>
<td>101325</td>
<td>1200</td>
<td>12.51</td>
<td>69.12</td>
</tr>
<tr>
<td>12</td>
<td>7.5</td>
<td>250</td>
<td>75</td>
<td>101325</td>
<td>1200</td>
<td>14.43</td>
<td>65.31</td>
</tr>
<tr>
<td>13</td>
<td>7.5</td>
<td>200</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>9.66</td>
<td>76.77</td>
</tr>
<tr>
<td>14</td>
<td>7.5</td>
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<td>50</td>
<td>101325</td>
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<td>15</td>
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<td>76.77</td>
</tr>
<tr>
<td>16</td>
<td>7.5</td>
<td>200</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>9.66</td>
<td>76.77</td>
</tr>
<tr>
<td>17</td>
<td>7.5</td>
<td>200</td>
<td>50</td>
<td>101325</td>
<td>1200</td>
<td>9.66</td>
<td>76.77</td>
</tr>
</tbody>
</table>

Blank corrosion rate: 150 rpm = 40.5 mpy; 200 rpm = 41.6 mpy; and 250 rpm = 44.3 mpy.

4. Journal of Chemistry
where $R_i$ and $R_0$ are the total resistance in the occurrence and nonattendance of LI, correspondingly.

The whole resistance is the totality of solution resistance and charge transfer resistance [52–54]. The calculated ethics of efficacy are tabularized in Table 3. According to Table 3, with growth in the LI concentration, the efficiency and $R_t$ values expressively enlarged. The supreme productivity was got for the protected solution by 10% of aqueous extract of LI. The better $R_t$ morals theme to the growth of shielding coat development on the surface of steel alloy 4130 [55]. Concluding the subsequent equation [56], the total capacitance ($C_t$) parameter, representative of the reduction or growth of the resident dielectric constant, thinness of the double layer, and the adsorption of inhibitor layer [57], was considered and conveyed in Table 3:

$$\epsilon_t = \frac{n}{1/n} \times \left( \frac{R_p}{R_s + R_t} \right)^{1-n/n}. \quad (9)$$

The compact $C_t$ that principles as an outcome addition of inhibitor load can be ascribed to the decrease in resident dielectric width and the argument of electrolyte fragments with inhibitor composites at the boundary of steel alloy 4130/3.5 wt.% NaCl [58].

### 3.1.3. Potentiodynamic Polarization Test Results.

Potentiodynamic polarization designs of the steel alloy 4130 trials uncovered to the 3.5 wt.% NaCl solutions together with LI are exemplified in Figure 5. Numerous electrochemical constraints such as cathodic Tafel slopes ($\beta_c$), corrosion potential ($E_{corr}$), and corrosion current density ($i_{corr}$) are gotten from the extrapolation process of Tafel plot, and the consequences are conserved in Table 4. It is expressed that, in the anodic designs of the protected steel alloy 4130 examples, no visible anodic Tafel region is undescribed and deliberated as the anodic concepts of the Tafel slope ($\beta_c$). It is clearly noticed that, as in Figure 5, the current density of both anodic and cathodic twigs significantly condensed in the occurrence of LI extracts [59]. Nevertheless, the extra effect of LI on the current density of cathodic branch, since it reduced and the move of $E_{corr}$ to extra negative magnitudes, proves that the LI represented as a cathodic inhibitor rather than an anodic one. Observing the equivalent cathodic

![Figure 4: Electrochemical equivalent circuit simulated for steel alloy 4130 sample engrossed in 3.5 wt.% NaCl electrolyte and diverse loads of LI.](image)

![Figure 3: Nyquist plots for the steel alloy 4130 sample engrossed in 3.5 wt.% NaCl electrolyte and diverse loads of LI.](image)

![Figure 5: Potentiodynamic polarization designs of steel alloy 4130 sample engrossed in 3.5 wt.% NaCl solution and diverse loads of LI.](image)

### Table 3: The electrochemical impedance parameters of steel alloy 4130 sample occupied in 3.5 wt.% NaCl solution with diverse loads of LI.

<table>
<thead>
<tr>
<th>Medium, %</th>
<th>$R_s$ ($\Omega \text{cm}^2$)</th>
<th>$R_p$ ($\Omega \text{cm}^2$)</th>
<th>$R_{p1}$ ($\Omega \text{cm}^2$)</th>
<th>$R_{p2}$ ($\Omega \text{cm}^2$)</th>
<th>L/H</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank</td>
<td>0.593</td>
<td>5.2x10⁻³</td>
<td>30.60</td>
<td>2.587</td>
<td>7.188</td>
</tr>
<tr>
<td>1</td>
<td>0.642</td>
<td>5.1x10⁻³</td>
<td>3.791</td>
<td>1.806</td>
<td>7.562</td>
</tr>
<tr>
<td>3</td>
<td>1.150</td>
<td>6.8x10⁻³</td>
<td>8.395</td>
<td>3.467</td>
<td>7.972</td>
</tr>
<tr>
<td>5</td>
<td>1.663</td>
<td>7.7x10⁻³</td>
<td>14.006</td>
<td>5.405</td>
<td>14.926</td>
</tr>
<tr>
<td>7.5</td>
<td>2.387</td>
<td>7.2x10⁻³</td>
<td>19.321</td>
<td>6.961</td>
<td>17.877</td>
</tr>
<tr>
<td>10</td>
<td>3.682</td>
<td>8.4x10⁻³</td>
<td>35.609</td>
<td>11.99</td>
<td>26.748</td>
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</table>
75°C, the corrosion rate decreases with increasing inhibitor concentration and temperature increases from 25 to 75°C decreases the corrosion rate from 6.75 to 8.66%, indicating no effect. By increasing inhibitor concentration to 7.5, 10%, the corrosion rate decreases. Also, increasing the temperature from 50 to 75°C increases the corrosion rate from 25 to 75°C decreases. Inhibiting the cathodic reaction deprived of altering the hydrogen development mechanism [60].

The provided data in Table 4 establish that the corrosion current density and corrosion potential constraints dropped in contradiction of an upsurge in LI concentration. This remark authenticates the actual inhibitive achievement of the LI on the drop of steel alloy 4130 corrosion mostly over the cathodic reaction conquest. The effect of LI on the anodic outlet figure clearly validates the fragments of inhibitor which interrelated through the energetic anodic spots and altered the steel alloy 4130 corrosion mechanism. The cations of steel produced throughout the anodic disbanding of steel alloy 4130 can vary developments with the organic particles of LI. The empty orbital of steel cations can receive the free lone pair of electrons of oxygen heteroatoms, which exist in the structure of inhibitor. Accordingly, the LI particles settle on the anodic spots and guard the steel from unembellished disbanding. At small loads, the coverage surface is imperfect, hence diminution of $i_{\text{corr}}$ with an upsurge in the inhibitor load, foremost to the upsurge in superficial coverage of the corrosion inhibitor [61].

### 3.2. Statistical Analysis

#### 3.2.1. Optimization of the Experimental Conditions

To optimize the three individual factors (inhibitor concentration, rotation speed, and temperature), there were a total of 17 experimental runs in the Box–Behnken design (Table 2).

#### 3.2.2. Corrosion Rate

Figure 6 shows the contour plot of the consequence of inhibitor concentration and rotation speed on the corrosion rate of steel alloy 4130 at different temperatures 25, 50, and 75°C. It is clear that, at 25°C, the maximum corrosion rate was 18.67 mL/Y. Also, increasing the temperature from 50 to 75°C increases the maximum corrosion rate to 27.58 and 38.68 mL/Y, respectively.

Figure 9 shows the 3D plot of corrosion rate of steel alloy 4130 as a function of inhibitor concentration and rotation speed at different temperatures 25, 50, and 75°C. It shows that, at 25°C, the maximum corrosion rate was 18.67 mL/Y. Also, increasing the temperature from 50 to 75°C increases the maximum corrosion rate to 27.58 and 38.68 mL/Y, respectively.

Figure 10 shows the 3D plot of the consequence of temperature and inhibitor load on the deterioration rate of steel alloy 4130 at different rotation speeds 150, 200, and 250 rpm. It shows that, at 150 rpm, the maximum corrosion rate was 34.13 mL/Y. Also, increasing the rotation speed from 200 to 250 shows insignificant increase in the maximum corrosion rate to 37.59 and 38.57 mL/Y, respectively [62].

Figure 11 shows the 3D plot of corrosion rate of steel alloy 4130 as a function of rotation speed and temperature at different inhibitor concentration 5, 7.5, and 10%. It shows that, at 5 %, the maximum corrosion rate was 32.71 mL/Y. Also, increasing the inhibitor concentration from 7.5 to 10% decreases the maximum corrosion rate to 13.67 and 10.23 mL/Y, respectively.

All the investigational data, composed of the 3D cubic, as publicized in Figure 12, exposed that the corrosion rate which fluctuated from 4.49 to 38.57 mL/Y can be produced. The lowest corrosion rate, 4.49 mL/Y, can be obtained at high concentration of inhibitor and small stages of temperature and rotation speed. On the contrary, the highest corrosion rate can be achieved at low concentration of inhibitor and high levels of temperature and rotation speed. However, increasing the concentration of inhibitor leads to decrease in the corrosion rate which is desirable. This is attributed to the LI inhibitor, which is well known as inhibiting agent, since it is stable at <50°C and 150 rpm. Figure 13 displays the normal

#### Table 4: Tafel constraints obtained from polarization curves from Tafel extrapolation technique of steel alloy 4130 trial immersed in 3.5 wt.% NaCl medium and diverse loads of LI.

<table>
<thead>
<tr>
<th>Medium (%)</th>
<th>$B_0$ (mV dec$^{-1}$)</th>
<th>$E_{\text{corr}}$ (mV)</th>
<th>$j_{\text{corr}}$ ($\mu Acm^{-2}$)</th>
<th>$B_n$ (mV dec$^{-1}$)</th>
<th>$R_p$ (Ωcm$^2$)</th>
<th>$R_{\text{corr}}$ (mmole$^{-1}$)</th>
<th>IE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank</td>
<td>$-102.96$</td>
<td>$-614$</td>
<td>$3145$</td>
<td>$224.86$</td>
<td>$154.3$</td>
<td>$44.4$</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>$-127.17$</td>
<td>$-605$</td>
<td>$3033$</td>
<td>$89.18$</td>
<td>$5.57$</td>
<td>$32.6$</td>
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<td>$76.45$</td>
<td>$4.77$</td>
<td>$4.8$</td>
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shapes through the minor slope variation specifies that LI inhibited the cathodic reaction deprived of altering the hydrogen development mechanism [60].

### Table 4: Tafel constraints obtained from polarization curves from Tafel extrapolation technique of steel alloy 4130 trial immersed in 3.5 wt.% NaCl medium and diverse loads of LI.
Figure 6: Contour plot of corrosion rate as a purpose of inhibitor load and rotational speed at the temperatures (a) 75°C, (b) 50°C, and (c) 25°C (mL/Y).

Figure 7: Continued.
plot of residuals in which almost all the residuals fall on a straight line, reflecting that errors were expected to distribute normally, namely, the model exhibits a good stability to predict the response [62, 63].

Figure 7: Contour plot of corrosion rate as a purpose of inhibitor load and temperature at the rotational speeds (a) 250 rpm, (b) 200 rpm, and (c) 150 rpm (mL/Y).

Figure 8: Contour plot of corrosion rate as a purpose of rotational speed and temperature at inhibitor loads (a) 10%, (b) 7.5%, and (c) 5% (mL/Y).
Figure 9: 3D plot of corrosion rate as a purpose of inhibitor load and rotation speed at temperatures (a) 75°C, (b) 50°C, and (c) 25°C.

Figure 10: Continued.
Figure 10: 3D plot of corrosion rate as a purpose of inhibitor load and temperature at rotation speeds (a) 250 rpm, (b) 200 rpm, and (c) 150 rpm.

Figure 11: 3D plot of corrosion rate as a purpose of rotational speed and temperature at the inhibitor loads (a) 10%, (b) 7.5%, and (c) 5%.
Final equation in terms of actual factors is indicated as follows:

\[
\text{Corrosion rate (mL/Y)} = +46.40000 \\
(i) \text{Concentration of inhibitor (\%)} = -18.31300 \\
(ii) \text{Rotational speed (rpm)} = +0.26115 \\
(iii) \text{Temperature (°C)} = +0.44920
\]

3.3. Inhibition Efficiency. Figure 14 shows the contour plot of the consequence of inhibitor concentration and temperature on the inhibition efficiency of steel alloy 4130 at different concentrations of inhibitor 5, 7.5, and 10%. It is clear that, at 5%, the inhibition efficiency was increased from 57.20 to 90.28% with increasing inhibitor concentration from 5 to 10%. Also, by increasing temperature from 25 to 75°C, the inhibition efficiency has no effect. The inhibition efficiency has no effect by increasing rotation speed to 200 and/or 250 rpm.

Figure 15 shows the contour plot of the consequence of inhibitor concentration and rotation speed on the inhibition efficiency of steel alloy 4130 at different concentrations of inhibitor 5, 7.5, and 10%. It is clear that, at 5%, the inhibition efficiency has no effect with increasing rotation speed from 150 to 250 rpm. Also, increasing inhibitor concentration from 5 to 10% increases the inhibition efficiency from 50.90 to 86.03%. By increasing inhibitor concentration to 7.5 and/or 10%, the inhibition efficiency increases.

Figure 16 shows the contour plot of the consequence of rotation speed and temperature on the inhibition efficiency of steel alloy 4130 at different concentrations of inhibitor 5, 7.5, and 10%. It shows that, at 150rpm, the maximum inhibition efficiency was 41.91%. Also, increasing the temperature from 250 to 75°C decreases the inhibition efficiency to 39.75 and 38.78%, correspondingly.

Figure 17 shows the 3D plot of the inhibition efficiency of steel alloy 4130 as a purpose of inhibitor load and rotation speed at different temperatures 25, 50, and 75°C. It shows that, at 25°C, the maximum inhibition efficiency was 62.85%. Also, increasing the temperature from 50 to 75°C decreases the inhibition efficiency to 51.13 and 38.78%, correspondingly.

Figure 18 shows the 3D plot of the inhibition efficiency of steel alloy 4130 as a purpose of inhibitor load and temperature at diverse rotation speeds 150, 200, and 250 rpm. It shows that, at 150 rpm, the maximum inhibition efficiency was 41.91%. Also, increasing the rotation speed from 200 to 250 rpm shows insignificant increase in the inhibition efficiency to 39.75 and 38.78%, respectively.

Figure 19 shows the 3D plot of the inhibition efficiency of steel alloy 4130 as a purpose of rotation speed and temperature at diverse inhibitor concentration 5, 7.5, and 10%. It shows that, at 5%, the inhibition efficiency was 44.99%. Also, increasing the temperature from 7.5 and 10% increases the maximum inhibition efficiency to 71.45 and 81.90%, correspondingly.

All the investigational data, composed at the 3D cubic as given in Figure 20, exposed that the inhibition efficiency which fluctuated from 38.78 to 93.67 (%) can be produced. The lowest inhibition efficiency, 38.78%, can be obtained at low concentrations of inhibitor and high levels of temperature and rotation speed. On the contrary, the highest inhibition efficiency can be realized at a great concentration of inhibitor and small stages of temperature and rotation speed [64]. However, growing the load of inhibitor leads to increase in the inhibition efficiency, which is desirable. This is attributed to the L1 inhibitor, which is well known as the inhibiting agent, since it stable at <50°C and 150 rpm. Figure 21 displays the normal plot of residuals in which almost all the residuals fall on a straight line, reflecting that errors were expected to distribute normally, namely, the model exhibits a good stability to predict the response [65].

Final equation in terms of actual factors indicated the following:

\[
\text{Inhibition efficiency (%)} = -8.27500
\]
Figures 14 and 15: Contour plots of the inhibition efficiency as a function of inhibitor load and rotational speed at different temperatures. (a) 75°C, (b) 50°C, and (c) 25°C.
### 3.3.1. DFT Results

To examine the consequence of operational constraints on the inhibition efficacy of the inhibitor, the quantum chemical calculations have been achieved. The adsorption mode of action on the metallic outward is

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**Concentration of inhibitor (%)** = +24.30700

**Rotational speed (rpm)** = −0.083375

**Temperature (°C)** = −0.50985

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**Figure 15:** Contour plot of inhibition efficiency as a purpose of inhibitor load and temperature at the rotational speeds (a) 250 rpm, (b) 200 rpm, and (c) 150 rpm.

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**Figure 16:** Contour plot of inhibition efficiency as a purpose of rotational speed and temperature at the inhibitor loads (a) 10%, (b) 7.5%, and (c) 5%.
The geometrical and electronic structure of lawsone is studied. The optimization of their bond angles, dihedral angles, and bond lengths. Figure 22 represents the calculated optimized molecular structure with the minimum energy. The calculated quantum chemical parameters can be used for the inhibition efficiency of inhibitor. Table 5 summarizes the calculated quantum chemical constraints as the energy of the lowermost empty molecular orbital (ELUMO), energy of the maximum engaged molecular orbital (EHOMO), the separation energy (ELUMO-EHOMO), and ∆E, representing the meaning of activity, dipole moment (D), chemical potential (µ), electronegativity (χ), softness (σ), and hardness (η). These results indicate that the geometrical structure of lawsone is planner, as shown in Figure 22. Rendering to the frontier molecular orbital theory (FMO), the chemical activity is a purpose of the collaboration amongst HOMO and LUMO planes of the acting types [66]. The EHOMO designates the capability of the fragment to provide electrons to a suitable receptor through unfilled molecular orbitals, and ELUMO designates its aptitude to receive electrons. The higher is the value of EHOMO of the inhibitor, the superior is its ease of donation electrons to the unoccupied d-orbital of metallic superficial, and the superior is its inhibition efficacy. The lower is the rate of ELUMO, and more is the ability of the fragment to receive electrons (Figure 23). The HOMO-LUMO energy gap (∆E) method, which is a significant steadiness guide, is practical to progress hypothetical replicas for explanation of the assembly and conformational fences in numerous molecular structures. The minor is the worth of ∆E, with extra possibility that the complex has inhibition efficacy [67, 68]. From Table 5, the molecule has the smallest HOMO-LUMO gap (∆E), and hence, the molecule has more inclination to adsorb on the metallic superficial. The dipole moment (D), the primary plagiaristic of the energy regarding a practical electric arena, is castoff to deliberate and vindicate the construction of the molecule [69]. In the literature, there is a deficiency of arrangement on the relationship amongst µ and inhibition effectiveness [70]. On the contrary, the hypothetical outcomes indicate that there is no noteworthy connection around amongst µ and inhibition efficacy, referring to Figure 17: 3D plot of inhibition efficiency as a function of inhibitor concentration and rotation speed at temperatures (a) 75°C, (b) 50°C, and (c) 25°C.
Table 5. The absolute softness ($\sigma$) and hardness ($\eta$) are significant possessions that quantify individually the constancy and activity of the structural formula. A firm fragment has a great energy hole, and a lenient fragment has a minor energy hole. The lenient fragments are supplementary responsive than the firm ones since they could simply suggest electrons to a receptor. Intended for the humblest transmission of electrons, adsorption could happen at the portion of the molecule where $\sigma$, which is a resident possession, has the maximum rate [71]. In a deterioration arrangement, the inhibitor performs by way of a Lewis base although the metallic surface performs as a Lewis acid, correspondingly. Consequently, it is determined that the inhibitor by means of the uppermost $\sigma$ rate has the maximum capability inhibition effectiveness (Table 5), which is in a decent arrangement with our investigational data. Moreover, our results show that the inhibitor has the lowest $\omega$, and $\chi$ (Table 5) which lead to growth of its contribution capability to the metallic superficial and consequently ornamental its inhibition efficacy. Based on the above results, we can simply conclude that the calculated quantum chemical parameters are confirming that the inhibitor has high inhibition efficacy which decides fit through the investigational explanations. The molecular electrostatic potentials are very supportive in the negative areas that can be observed as nucleophilic cores although areas through positive electrostatic potential are potential electrophilic spots. Additionally, the electrostatic potential types the polarization of the observable electron compactness. Figure 24 shows that the biphenyl moiety and the oxygen atom of hydroxyl group have negative electrostatic potential which resources that these spots are the energetic cores for the requisite to the metallic superficial. The electronic-structure DFT investigates and demonstrated the interactions between the giver and receiver of responsive spots of the corrosion inhibitive particles. We remarkably noticed that the major watersoluble ingredient for the LI, the O-containing lawsone
Figure 19: 3D plot of inhibition efficiency as a purpose of rotational speed and temperature at the inhibitor loads (a) 10%, (b) 7.5%, and (c) 5%.

Figure 20: 3D plot of inhibition efficiency as a function of inhibitor concentration, rotation speed, and temperature variables.

Figure 21: Normal plot of error distribution for inhibition efficiency.
A compound which includes a perfect formula, can provide a good corrosion inhibiting effect due to the presence of more than one polar oxygenated center. The potentiality of inhibition was computationally confirmed by DFT theory through the protonation attraction foremost to its probable physisorption and projected adsorption energy.

4. Conclusions

The characteristics of corrosion inhibition of green inhibitor based on Lawsonia inermis extract (LI) in 3.5 wt.% NaCl solutions on steel alloy 4130 surfaces were estimated by collective investigational and computational approaches. Examinations by EIS designated that, by cumulating the load of LI from 1 to 10%, the protection efficacy pointedly amplified and touched 89.19%. The LI performed as a diverse-type corrosion inhibitor and measured the degree of
corrosion of equally anodic and cathodic responses. The water-soluble components of LI can interact with steel alloy 4130 superficial over distribution of their lone pair of electrons through empty d-orbital of steel alloy 4130 molecules.

Data Availability
The data used to support the findings of this study are included within the article.

Conflicts of Interest
The authors declare that there are no conflicts of interest.

Authors’ Contributions
All authors contributed equally to this work.

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