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# Synthesis and inhibition corrosion effect of two thiazole derivatives for carbon steel in

### 1□M HCl

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#### Abstract

Inhibition of C38 carbon steel corrosion by Phenyl-3-phenyl-2(3H)-thiazolethione (TO1) and 4-methyl-2-(methylthio)-3- phenylthiazolethiac (ST1) in 1 M HCl was investigated by weight loss and electrochemical methods. All of the data obtained reveal that the two compounds act as good inhibitors in this media. At optimized concentration TO1 and ST1 showed the highest inhibition efficiency of 98.8 % (2.10<sup>-4</sup>M) and 93.86 % (10<sup>-3</sup>M) respectively. Polarization curves show that inhibitor molecules act as mixed type inhibitors. The impedance study showed that an increase in the concentration of the two inhibitors is accombanied by an increase in polarization resistance and a decrease in double layer capacitance. The Langmuir isotherm very well describes the adsorption of inhibitors to the surface of the corroding metal and the thermodynamic parameters showed that the adsorption of the two compounds was strong and chemical nature. X-ray photoelectron spectroscopy (XPS) confirms and describes the absorption of inhibitors under investigation on the metal surface.

**Keywords:** Corrosion inhibition; C38 carbon steel; HCl; thiazole derivatives, XPS

### 1. Introduction

Carbon steel is one of the most widely used metals in the automotive industry, machinery, construction structures, among many other uses. This material has been extensively studied as a model for corrosion inhibition. This is due to the fact that carbon steel corrodes very easily in different environments, but if we talk about an acidic medium which is very corresive, the corrosion of carbon steel increases considerably [1-4]. In general, the inhibitors are organic compounds in whose structures are oxygen, sulfur, nitrogen and phosphorous atoms and unsaturated bonds together with aromatic rings. Thiazole-type compounds have been proven to be potent corrosion inhibitors. This is due to their strong absorption susceptibility to metal surfaces in aggressive media via their polar groups and their potential for complexing with the metal surface. In addition, they exhibit different agricultural, industrial and biological properties such as anticancer, antiviral, antibacterial antifungal, etc. [5-18].

All of these properties have manifested their intr-toxic character. They also possess abundant pie-electrons and unshared electron pairs on the nitrogen atom, which can interact with any metal's d-orbitals to provide a projective film1 [13-14]. Thus in recent years research on derivative thiazole inhibitors has been a major topic. Researchers have made many efforts to examine new thaz le derivatives in the hope of improving its inhibition effectiveness. A common route for enhancing a given heterocycle's corrosion inhibition efficiency is to alter its structure with different moieties or functional groups [15-18].

This work is focused at synthesizing two thiazolic derivatives: 4-methyl-3-phenyl-2(3H)-thiazolethione (**TO1**) and 4-methyl-2-(methylthio)-3- phenylthiazol-3-ium (**ST1**) and study their inhibition efficiency as inhibitors for C38 carbon steel in 1M HCl solution, using electrochemical methods, weight loss measurement and XPS surface study.

### 2. Experimental Section

### 2.1 Materials

# 2.1.1. Composition of C38 Sample.

The pieces of steel sheet (used as working electrode and for weight loss experiments) were prepared from C38 carbon steel, where the composition and method of preparation was given in our already published work [14].

### 2.1.2. Solutions.

The concentration of the used inhibitors ranged from 10<sup>-6</sup>M to 10<sup>-3</sup>M in 1 M ICN orepared from 37% HCl in distilled water).

# 2.1.3. Synthesis of corrosion inhibitors

## a) 4-methyl-3-phenyl-2(3H)-thiazolethione (**TO1**)

Aniline (18.5 mL, 0.2 mol) was added dropwise on a mixture of  $CS_2$  (14.5 mL, 0.2 mol) and 24 mL of NH<sub>4</sub>OH, stirred at 0°C. After stirring for 2 hour at room temperature, the mixture was filtered and washed with  $Et_2O$  (3 x 25 mL) to give the yellow dithiocarbamate (30 g, 80%). 1-Chloro-2-propanone (9 mL, 0.1mol, 1 eq) was added to a suspension of (18.6 g, 0.1mol) of the dithiocarbamate call in et lanol (20 mL). After stirring for 2 hours at r.t., 0.5 mL of HCl (36%) were added and me mixture was refluxed for 3 hour. After filtration, the solid was washed with vater (3 x 5 mL) and dried to give (TO1) (15.13 g, 73%, mp = 146°C). Detailed reaction mechanism is presented in figure 1a and NMR spectra of TO1 is presented it figure 1b.

$$NH_2 + S = C = S$$
 $NH_2 + S = C = S$ 
 $NH_4 \cdot OH$ 
 $NH_5 = S \cdot OH$ 
 $NH_4 \cdot OH$ 

Figure 1a. Synthesis of 4-methyl-3-phenyl-2(3H)-thiazolethione (TO1

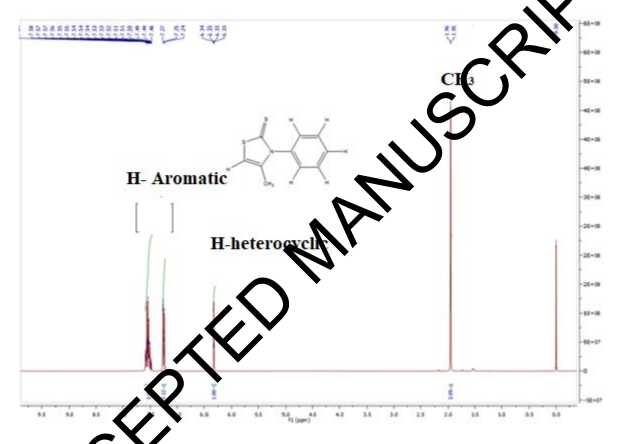


Figure 16. NMR spectra of 4-methyl-3-phenyl-2(3H)-thiazolethione (TO1)

# b) Vmethyl-2-(methylthio)-3- phenylthiazol-3-ium (ST1)

The method adapted to the synthesis of this thiazolium salt (ST1), provides the condensation of thiazolin-2-thione, with the alkyl iodine in acetonitrile with magnetic stirring at room temperature according to the alkylation method of thiazol-2-thione described by literature [20-21] as shown in figure 1c and NMR spectra of ST1 is presented in figure1d.

In a 200 ml monocolumn flask, 100 mg (10 mmol, 2 g, 1 eq) of thiazolin-2-thione are solubilized in acetonitrile (50 ml). Then 6.84 ml (11 eq) of methyl iodide is added using a syringe. The solution is stirred magnetically for 24 h at room temperature. The product is obtained by sinter filtration and dried (85%, mp = 220°C).

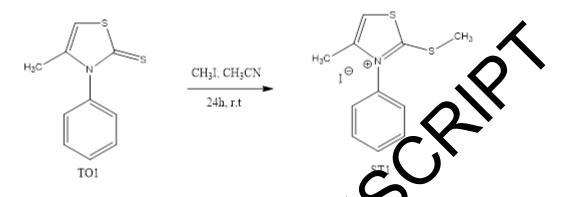


Figure 1c. Synthesis thiazolium iodides via thiazol. \_\_-thione.

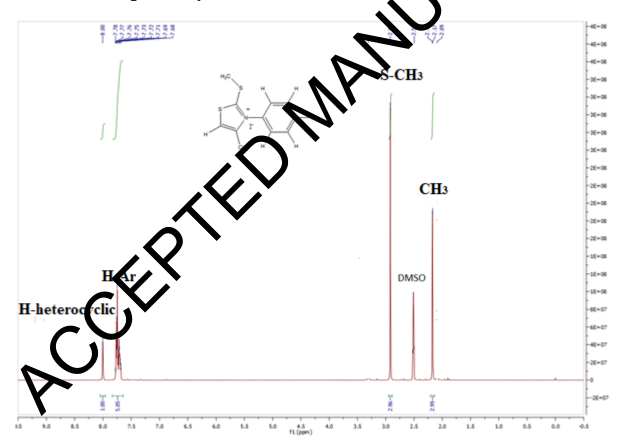


Figure 1d. NMR spectra of 4-methyl-2-(methylthio)-3- phenylthiazol-3-ium (ST1)

### 2.2. Methods

For weight loss studies the dimension of C38 steel is  $1.5~\rm cm \times 1.5~\rm cm \times 0.2~\rm cm$ . The measurements were carried out at  $30^{\circ}\rm C$  for 1 h of immersion into 100 mL of 1 M hydrochloric acid in the absence and the presence of various concentrations of inhibitors.

After immersion, the samples were washed, degreased with acetone and weighed. All the manipulations were repeated three times in order to have reproducibility. Equations (1), (2) and (3) give the procedure for calculating the corrosion rate (C.R), the inhibitary afficiency (IE%) and the recovery rate ( ) [4]:

$$-$$
 (1)

where  $\Delta W$  = weight loss (mg), S = area of specimen (cm<sup>2</sup>), and = posure time (hours).

$$(2)$$

(3)

where C.R and C.R' are the corrosion rate with and without inhibitor, repectively.

Electrochemical measurements were carried out at a temperature of 30 °C in a Pyrex glass thermostatted cell with three electrodes (carbon steel C38 as working electrode with an area of 1 cm<sup>2</sup>, platinum as rounter-electrode (1 cm<sup>2</sup>) and a saturated calomel electrode as reference electrode). The solution is subjected to stirring and azote gas bubbling. The Solartron instruments SI 1287 potentiostat assisted by a personal computer via a GPIB interface and the CorrWare 2.80 software were used to perform the electrochemical tests and collect the experimental data. The steel sample was allowed to corrode freely and its open circuit potential (OCP) for 1 hour (sufficient time to have the stable potential regime). The last OCP value recorded corresponds to the corrosion potential (E<sub>corr</sub>) of the working electrode.

The inhibition efficiency (IE %) was calculated using the following equation:

where <sub>0</sub> and <sub>ihn</sub> are corrosion current without and with inhibitor, respectively.

AC impedance measurements were performed with the same electrochemical system using ZPlot 2.80 and ZView 2.80 software to fit the spectra obtained. The system was excited with alternating current with frequency range from  $10^5$  Hz to  $10^{-2}$  Hz and a peak-to-peak amounted of 10 mV with 10 points per decade. All EIS diagrams were registered at  $E_{corr}$  potential. The values of charge transfer resistance were used to calculate the inhibition efficiency (EI<sub>Rt</sub> (%)) as previously described [4].

The detailed description of the method X-ray photoelectron spect oscop, (XPS) as well as the treatment of the results is given in our published paper [4].

### 3. Results and Discussion

### 3.1. Weight loss measurements

The table 1 gives the values of the corrosion rate and inhibition efficiency of **TO1** and **ST1** in 1M HCl. An overview on this table, we clearly notice that the both inhibitors are effective against corrosion, even at low concentration and that their efficacies are related to the inhibitor concentrations.

Maximum IE% of each compound was achieved at  $2 \cdot 10^{-4}$  M for **TO1** and  $10^{-3}$  M for **ST1**. The use of lower concentration for the **TO1** inhibitor can be explained in terms of non-binding pairs in the molecule. While **TO1** has two unbonded electron pairs, the **ST1** molecule has only one pair. This fact indicates a greater adsorption and attraction to the metal of the **TO1** inhibitor.

Table 1. Weight loss results of C38 carbon steel in presence of TO1 and ST1 in 1M HCl

Conc.	Corrosion rate	IE
( <b>M</b> )	$(mg.cm^2.h^{-1})$	(%)

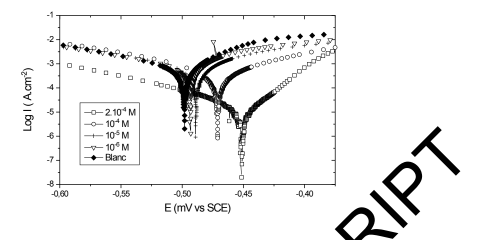
	D1 1	1.00	_
	Blank	1.00	
	10 <sup>-6</sup>	0.61	39
	10 <sup>-5</sup>	0.42	58
TO1	$10^{-4}$	0.22	78
	$2.10^{-4}$	0.07	93
	10 <sup>-6</sup>	0.48	52
ST1	10 <sup>-5</sup>	0.30	70
	$10^{-4}$	0.24	76
	5.10 <sup>-4</sup>	0.13	
	10 <sup>-3</sup>	0.06	94
			<del>)                                    </del>

Moreover, we note from the same table that the S(X) is more efficient compared to the TO1 at the concentration lower than  $10^{-4}$ M. This is probably due to the strong adsorption at low concentration of the compound TO1 compared to ST1.

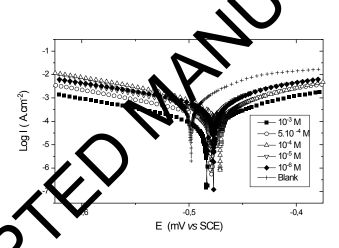
# 3.2. Electrochemical experiment

# 3.2.1. Polarization curves

Under the same conditions as the gravimetric measurements (t = 1h and T = 30  $^{\circ}$  C), the anodic and catholic polarization curves in the absence and in the presence of both inhibitors at different concentrations were obtained (figure 2a and 2b). The scan rate of the electrochemical tests is 0.5 mV/s [22-23].



**Figure 2a.** Potentiodynamic polarization curves for C38 steel in 1 M ACI containing different concentrations of **TO1.** 



**Figure 2b.** Potentio synamic polarization curves for C38 steel in 1 M HCl containing different concentrations of **S11**.

In both inhibitors, a decrease in the anodic and cathodic currents is observed. The values extracted from these figures such as corrosion potential, Tafel slopes, corrosion current and inhibitor efficiency are shown in table 2.

From this table, we can draw the following conclusions:

 $\bullet$  The presence of the two inhibitors moves the  $E_{corr}$  values towards anodic values.

- As reported elsewhere, when the values of corrosion potential in the presence of inhibitor is greater than ±85 mV compared to the value in the absence of inhibitor, the inhibitor is considered to be either anodic or cathodic. In this study, the maximum displacement in corrosion potential observed is ±32 mV suggests that the two inhibitors behaves as mixed type of inhibitor [4].
- The values of b<sub>c</sub> change with increasing concentration of both inhibitors. There results indicate that the kinetic of the hydrogen evolution reaction is influenced by the presence of **TO1** and **ST1**. Such behavior has been largely observed and discussed in the literature [24-25].
- The increase in the concentration of inhibitors has the onsequence of increasing their inhibition efficiency.

**Table 2.** Electrochemical potentiodynamic polarization parameters for C38 steel in 1 M HCl containing **TO1** and **ST1**.

	Conc.	-E <sub>corr</sub>	<b>-b</b> <sub>c</sub>	$\mathbf{I_{corr}}$	$\mathbf{EI}_{\mathbf{Icorr}}$
	(Mol/L)	(nV s SCE)	(mV/dec)	$(mA/cm^2)$	(%)
	Panc	-498	138	1115	
	10-6	-494	137	796	28.60
TOI	10-5	-488	106	500	55.15
	10 <sup>-4</sup>	-470	52	253	77.30
ر ا	$2.10^{-4}$	-466	73	34	96.95
	10-6	-477	135	529	52.55
	10 <sup>-5</sup>	-491	96	348	68.78
ST1	10 <sup>-4</sup>	-473	65	305	72.64
	5.10 <sup>-4</sup>	-486	85	160	85.65
	$10^{-3}$	-487	93	70	93.72

# 3.2.2. The electrochemical impedance spectroscopy (EIS)

Nyquist Impedance diagrams obtained at  $E_{corr}$  for C38 steel in HCl 1M in the absence and in the presence of **TO1** and **ST1** are given in figure 3 (a and b).

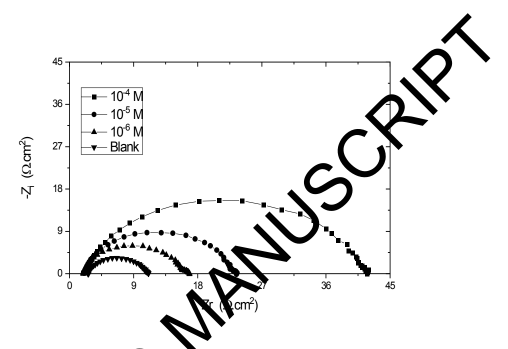


Figure 3a. Nyquist plot for C38 stee and M HCl with and without TO1 at 30°C.

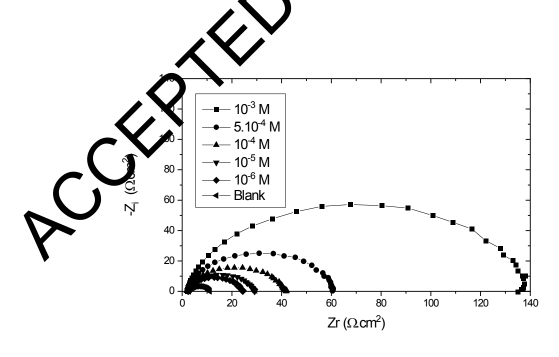


Figure 3b. Nyquist plot for C38 steel in 1 M HCl solution with and without ST1 at 30°C.

The diagrams are not perfect semicircles and have been attributed to frequency dispersion [26]. This approximately semicircular of impedance diagrams shows that the corrosion phenomenon is controlled by a charge transfer process [6, 27-29]. The increasing of the diameter with increasing of the concentration of the inhibitor indicates the good inhibition effect of our inhibitor [6]. The modeling of these diagrams by the ZView software led us to propose the following equivalent electrical circuit: an electrolyte resistor in series with a charge transfer resistor which in parallel with a CPE (figure 4).

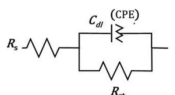


Figure 4. Electrical equivalent chronic

The obtained results are given in Table 3 (the charge transfer resistance  $R_t$ , n values, the CPE (Q) values, the double layer capacitance  $C_{dl}$  and inhibition efficiency). Ddetailed method of calculating  $C_{dl}$  values from CPE values is identical to that given in our previous work [4].

It can be seen that the presence of the two inhibitors affects the values of  $R_t$  towards the high values and the values of C in the opposite direction. On the otherhand, in all the case of TO1 and **ST1**, n reaches approximately the same value of 0.80. This result can be interpreted as an indication of the degree of heterogeneity of the metal surface, corresponding to a small depression of the double layer capacitance semicircle.

These result confirmed the obtained results by the above methods.

**Table 3.** Impedance parameters for the corrosion of the system C38 steel / 1 M HCl in the presence of **TO1** and **ST1**.

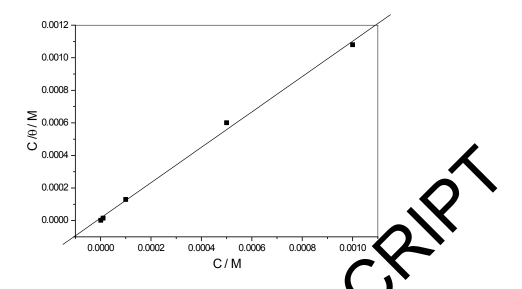
Conc.	$\mathbf{R}_{\mathbf{t}}$	Q	n	$C_{dl}$	$\mathbf{EI}_{\mathbf{Rt}}$
(Mol/L)	$(\Omega.cm^2)$	$(\Omega^{-1}.cm^{-2}.s^n)$		(µF cm <sup>-2</sup> )	(%)

	Blank	7.77	9.81x10 <sup>-4</sup>	0.8	290	
	10 <sup>-6</sup>	13.57	$7.05 \times 10^{-4}$	0.79	205	42.74
	10 <sup>-5</sup>	20.02	4.84x10 <sup>-4</sup>	0.78	131	61.18
TO1	10 <sup>-4</sup>	36.4	3.24x10 <sup>-4</sup>	0.80	107	78.65
	2.10 <sup>-4</sup>	252.8	6.7x10 <sup>-5</sup>	0.79	23	98.80
	10 <sup>-6</sup>	20.69	5.14x10 <sup>-4</sup>	0.81	177	2.44
	10 <sup>-5</sup>	25.08	4.30x10 <sup>-4</sup>	0.78	120	69.01
ST1	10 <sup>-4</sup>	36.4	3.24x10 <sup>-4</sup>	0.80	10	78.65
	5.10 <sup>-4</sup>	55.55	2.35x10 <sup>-4</sup>	0.82	91	86.01
	10 <sup>-3</sup>	126.7	1.08x10 <sup>-4</sup>	0.00	37	93.86
			•			

# 3.3. Adsorption isotherm

The Langmuir adsorption isotherm is plotted in the figure 4 (example for ST1) using the parameters from weight loss measurements and the following equation ( $C_{inh}/\theta=1/K+C_{inh}$ ) with  $C_{inh}$  is the inhibitor concentration, the  $\theta$  is the degree of surface coverage and K is the adsorption constant [30-37].

The plot of C/ $\theta$  vs. C (Fig. 5) yields a straight line with correlation coefficient close to 1. The Values of K was found to be  $1.42 \times 10^5 \, \text{M}^{-1}$  and  $0.71 \times 10^5 \, \text{M}^{-1}$  for **TO1** and **ST1** respectively. The strong adsorption of the two inhibitors on C38 steel surface is reflected by high value of adsorption equilibrium constant.



**Figure 5.** Curve fitting of the corrosion data of C38 stellar the presence of **ST1** to Langmuir isotherm

The relation between the standard free enthalpy,  $\Delta G_{N_0}$ , and the adsorption constant (K) is given by the relation:

$$K = 55.5 \text{ Mp} \Delta G_{ads}/RT$$

The  $\Delta G_{ads}$  was calculated as -40.0 M/mol and -38.05 kJ/mol for **ST1** and **TO1** respectively.

The spontaneous adsorption on the two molecules on the metal surface is clearly identified by the large negative values of the standard absorption enthalpy.

According to the interactive, values of  $\Delta G_{ads}$  around of -40 KJ.mol<sup>-1</sup> or less are attributed to the formation of strong covalent bonds between the surface of the metal and the free electrons existing on the neteroatoms of the inhibitory molecules therefore chemisorption [27, 32-34]. In our study, the calculated values of  $\Delta G_{ads}$  indicate the chemisorptive nature of the adsorption of inhibitors in 1M HCl solution [35-36].

The use of thiazole derivatives as corrosion inhibitors has been discussed in several publications [37-42]. Table 4 shows the protection efficiency and  $\Delta G_{ads}$  obtained of some thiazole derivatives exploited as corrosion inhibitors of steel in a 1 M HCl medium. Such comparison shows that the values of the standard free enthaply ( $\Delta G_{ads}$ ) obtained in our study

and those of the literature are in good agreement where we clearly notice that the majority of thiazole derivatives are strongly adsorbed with a mixed adsorption (physisorption with a tendency to chemisorption).

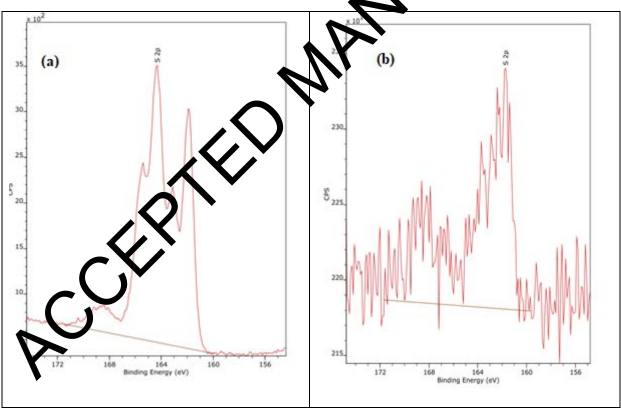
Table 4. Corrosion protection parameter for thiazole derivatives in 1M HCl

Thiazole derivative	Optimum Concentration	Metal	Protection efficiency	-ΔG <sub>ads</sub> (kJ/A ol)
4-methyl-3-phenyl-2(3H)-	$2x10^{-4}M$	C38	98.80%	40.0
thiazolethione [this work]		steel	•	
thiazol: 4-methyl-2-(methylthio)-3-	10 <sup>-3</sup> M	C38	93.86%	38.05
phenylthiazol-3-ium [this work]		steel		<b>\</b> '
3-((4-amino-2-methylpyrimidin-5-	40 ppm	Carbon	91. %	48
yl)methyl)-5-(2-hydroxyethyl)-4-		steel		
methylthiazol-3-ium chloride [ 37]		C		
4-(pyridin-4-yl)thiazol-2-amine [38]	2x10 <sup>-4</sup> M	M\d steel	96.06%	36.33
2-(2'-hydroxylphenyl)benzothiazole [39]	50 ppm	Mile teel	92.1%	34.31
2-(2',5'-	50 pp.	Mild	51.6%	31.87
dihydroxyphenyl)benzothiazole [39]		steel		
(4-benzothiazole-2-yl-phenyl)-	50 p.m.	Mild	96.8%	40.80
dimethyl-amine [39]	M.	steel		
2-(n-hexylamino)-4-(3 -N,N-	0 <sup>-3</sup> M	Carbon	99%	34.09
dimethylamino-propyl)amino6		steel		
(benzothiazol-2-yl)thio-1,3.5-s-				
triazine [40]				
and 2-(n-octylamino)-4 3 -N,N	10 <sup>-3</sup> M	Carbon	99.3%	37.92
dimethylaminopropyl)amh o-6-		steel		
(benzothiazol-2-y )thib-1,3,5-s-				
triaz ne [4.1]	7			
Mercaptobenzothiazole [40]	$10^{-3}$ M	Carbon	57.3%	31.72
	7	steel		
4-(1-hexad cyl-1 H-benzo[d]imidazol-	$10^{-3}$ M	Mild	94.5%	38.2
2-yr)miazole [41]	3	steel		
1,3 diheyadecyl-2-(thiazol-4-yl)-	10 <sup>-3</sup> M	Mild	95.3%	41.6
Ha enzo[d]imidazol-3-iumbromide		steel		
[41]	5			
2-(2 Hydroxyphenyl)	7.10 <sup>-5</sup> M	Mild	96.8%	-37
benzothiazole [42]		steel		

# 3.4 Analysis of the organic film formed by XPS photoelectron spectroscopy

For a better understanding of the inhibition mechanism, we performed photoelectron spectroscopy (XPS) surface analyzes on the organic inhibitor alone and on the organic film adsorbed on the steel surface. In this case we study only the results of **TO1** compound.

The characterization by XPS is carried out on the pure **TO1** pressed in the form of a pellet  $(13x2x2 \text{ mm}^3)$  and on the steel disc immersed for 24 hours in 1M HCl in the presence of the optimal concentration of **TO1**  $(2 \times 10^{-4} \text{ M})$ . The latter is removed from the corrosive medium, rinsed with distilled water and degreased in ethanol under ultrasound and finally fried. The presence of TO1 on the steel surface was detected based on the signal characteristic of the sulfur atom S2p (Fig. 6b). For comparative purposes, the XPS spectrum of ture TO1 (Fig. 6a) were also obtained.



**Figure 6.** XPS spectrum of S2p for: (a) pure TO1 and (b) the surface of the steel exposed to the solution  $(2 \times 10^{-4} \text{M} \text{ TO1} + 1 \text{M} \text{ HCl})$  for 24 h at 30 ° C.

For the surface exposed to hydrochloric acid containing TO1, the large signal from sulfur observed in the binding energy range of 161 to 172 eV, can be attributed to several

components with different binding energies. We observed that the S2p peaks can be deconvolved into several components formed of three doublets. The first deconvolved peak in 162.0 and 163.3 eV is attributed to the oxidation states of sulfur. The doublet at 164.6 and 165.9 eV, which is characteristic of the thione function and thiazole (structure -S-), shows the presence of **TO1** on the steel surface [43] and a doublet at 168.5 and 169.8 eV is due to the presence of sulphates,  $SO_4^{2-}$  which is due to the oxidation of  $S^{2-}$  [44-45].

The surface of the steel covered by chemisorbed **TO1** molecules (Fig. 7) reveals the existence of a binding energy component to 711 eV (Fe2p 3/2) and at 724 eV (Fe 2p 1/2) characteristic of Fe<sup>3+</sup>, showing the oxidation of the steel surface [46-47]. The same figure shows à little Fe2p peak at 706.6 eV characteristic of Fe° [48-49]. The increase in the Fe2p signal confirms the adsorption and the film formation on the steel surface [50.51].

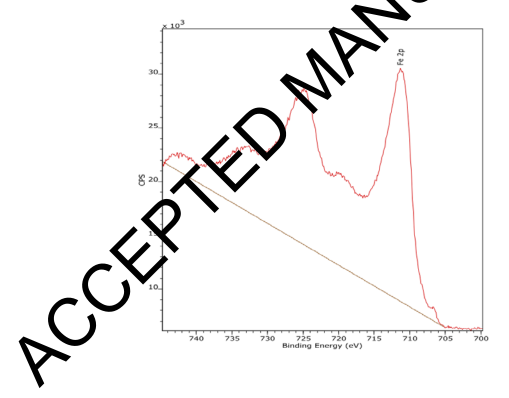


Fig. 7. Spectre Fe2p pour la surface de l'acier oxydée exposée à une solution ( $2\times10^{-4}$  M TO1 + HCl 1M) pendant 24 h à  $30^{\circ}$ C.

N 1s spectrum of metal treated with TO1 in 1 M HCl shows one peak located at around 399 - 402 eV (Fig. 8). This peak can be partly associated to =N- structure in the thiazole ring [52].

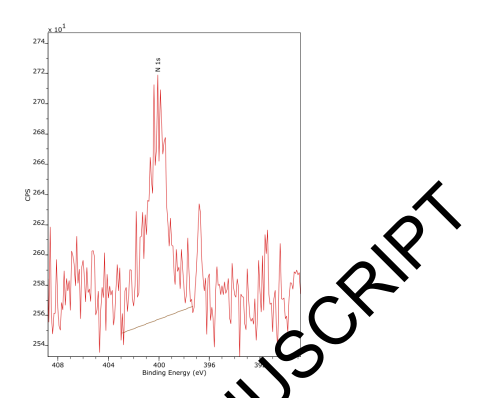


Fig. 8. XPS profile of N 1s for TON treated steel.

### 3.4. Conclusion

The results of inhibition of two compour and ST1) were studied in 1 M HCl for corrosion of C38 steel by differ methods (weight loss and the electrochemical good inhibitors with better efficiency for TO1. The measurements). **TO1** and **ST**1 ere classified as cathodic type inhibitors. The above compounds under inves ation ed by EIS study where the charge transfer resistance increases and conclusions were confirm decreases as the concentration of the inhibitor increases. Adsorption double layer ca on the C38 steel surface obeyed to the Langmuir isotherm. An XPS Infirm that the adsorption of the two molecules was done through nitrogen and ar atoms.

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