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Ethyl ester/acyl hydrazide-based aromatic sulfonamides: facile synthesis, structural characterization, electrochemical measurements and theoretical studies as effective corrosion inhibitors for mild steel in 1.0 M HCl†

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In this research paper, aromatic sulfonamide-derived ethyl ester (p-TSAE) and its acyl hydrazide (p-TSAH) were directly synthesized, characterized, and employed for the first time as prospective anticorrosive agents to protect mild steel in 1.0 M HCl conditions. The corrosion efficiency was probed by electrochemical methods including polarization, impedance, and frequency modulation measurements. Optimal efficiencies of 94% and 92% were detected for the hydrazide and ester, respectively, revealing excellent corrosion inhibition. Moreover, both the hydrazide and ester molecules combat the cathodic and anodic reactions correspondingly in a mixed-type manner. The electron transfer (ET) at the inhibitor/ metal interface was evaluated using DFT at the B3LYP/6-31g(d,p) level. Natural bond orbital analysis (NBO) and frontier molecular orbital analysis (FMO) calculations showed superior capabilities of the synthesized inhibitors to easily reallocate charge into the metal surface. However, the hydrazide molecules showed slightly better inhibition efficiency than the ester due to the strong interaction between the lone pairs of the nitrogen atoms and the d-orbitals of the metal. The chemical hardness of the hydrazide and ester are 2.507 and 2.511 eV, respectively, in good accordance with the recorded electrochemical inhibition efficiencies for both molecules. Good and straightforward correlations between the experiments and calculations are obtained. PAPER

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1. Introduction

Organic molecules are widely used as effective corrosion inhibitors for mild steel in aqueous media as they are rich in adsorption centers, i.e. hetero atoms (O, N and S), and a small amount of the inhibitor can provide a high inhibition efficiency, which makes it cost effective.¹⁻³ The rate of both cathodic and anodic corrosion reactions at the metal/solution interface can be suppressed to a high extent by the mutual interactions between the metal surface and the organic film, which are governed by an adsorption mechanism.4,5 While the rate of corrosion can be probed by electrochemical methods such as electrochemical impedance spectroscopy, electrochemical

frequency modulation and potentiodynamic polarization, the metal–inhibitor interactions can be probed by theoretical calculations.^{6,7}

Aromatic/heterocyclic sulfonamides are a broad collection of privileged organic sulfur motifs that have received a lot of interest in the pharmaceutical industry as drugs and potential therapeutic candidates.⁸ In 2020, Mondal and Malakar documented in their review article the advanced protocols for the chemical synthesis of sulfonamides and their important synthetic and biological implementations.⁹ More recently, Iakovenko et al. (2022) described a novel approach for the synthesis of N-alkyl heteroaryl sulfonamides in good yields via the direct oxidative coupling of readily available heterocyclic thiols and primary amines under mild conditions.¹⁰ This landmark work has been extended to transform the prior products to the corresponding N,N-dialkyl heteroaryl sulfonamides through a microwave-assisted Fukuyama–Mitsunobu alkylation reaction.¹⁰ In 2021, Mkrtchyan and Iaroshenko reported a successful mechanochemical strategy to construct diverse aromatic sulfonamides by the Pd-mediated one-pot coupling transformation of $K_2S_2O_5$ (a source of SO_2) and

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Table 1 Comparison of our outcomes with previously reported studies on the employment of hydrazides as anti-corrosive agents

 a Inhibition efficiencies (%) are tabulated according to the electrochemical impedance spectroscopy (EIS) measurements.

amines with aryl bromides or aromatic carboxylic acids as arylation reagents.¹¹

Abd El-Lateef's (2020) study showed that a new salicylidene isatin hydrazone sodium sulfonate inhibits the corrosion of carbon steel (CS) in 1.0 M HCl by up to 87.8%, and when merged with nickel (Ni^{2+}) cations, the inhibition ability was increased up to 99.2%.¹² In 2021, Verma and his colleagues¹³ reported the synthesis and characterization of three N-hydroxybenzothioamide analogues that exhibited inhibition efficiencies of over 90% at concentrations of 300 ppm. In March 2021, Abd El-Lateef et al.'s published article¹⁴ described the construction of a new hydrazide–hydrazone and its metal complexes with $Zn(\pi)$ and $ZrO(\pi)$, derived from nicotinic acid hydrazide and salicylaldehyde-5-sodium sulfonate salt. Chelatebased ZrO showed the best anti-corrosive potency with an efficacy reaching 97.4% at a dose of 5 \times 10⁻⁴ mol L⁻¹. In fact, several hydrazides and hydrazones have attracted a high level of attention in corrosion studies recently, as shown in Table 1.¹⁵–²² This table shows a comparison of recent reports and currently investigated compounds as corrosion inhibitors for mild steel in 1.0 M HCl, of which our investigated compounds show significant and promising efficiencies compared to the other published compounds.

A combination of the hydrazide functional group and sulfurcontaining compounds such as sulfonamides for application as corrosion inhibitors is rare in the literature, indicating a gap that needs to be filled. According to the best of our information, there is one research article, published in 2020 by Laggoun and co-authors, who described the protection impact of *p*-toluenesulfonyl hydrazide (p -TSH) on the copper surface in a 0.5 M HCl medium.²³ In a promising result, p-TSH successfully blocked the corrosion sites on the copper surface with an approximate inhibition efficiency of 90% at 5 mM. Consequently, this finding encouraged us to synthesize ethyl 4-(4-methylphenylsulfonamido)benzoate (p-TSAE, 8) and 4-(4-methylphenyl-sulfonamido)benzohydrazide (p-TSAH, 9), and utilize them for the first time ever as potential anticorrosive agents to protect mild steel under acidic conditions due to 1.0 M HCl. We used DFT calculations with the B3LYP method and 631g(d,p) basis set to obtain molecular reactivity parameters on both the molecular and atomic levels. New and promising findings were obtained relating the molecular aspects of the synthesized inhibitors to mutual inhibitor–metal interactions.

2. Experimental and computational details

2.1 Materials and instrumental techniques

For the chemical synthesis, starting materials and reagents were obtained from commercial sources, particularly Acros Organics™ (Geel, Belgium), and used without further purification unless otherwise indicated. Solvents were obtained from Thermo Fisher Scientific™ and employed without additional purification. All reactions were carried out in oven-dried glassware. The purity of the synthesized compounds was investigated by TLC and performed on Merck precoated silica gel 60 $F₂₅₄$ aluminum sheets with a solvent mixture of DCM-MeOH (99–1) as the eluent system. Spots were visualized under UV illumination at 254 nm. Chemical synthesis procedures were not optimized. Compound names are derived from ChemOffice Suite 2021 v21.0.0.28 and are not necessarily identical to those according to IUPAC nomenclature. Schemes were produced by the same version of ChemOffice. IR and MS data were plotted by OriginPro 2021. NMR spectra were delineated via MestReNova v12.0.2.

Melting points were determined on a SMP50 Digital APP (Bibby Scientific, Staffordshire, UK) 120/230 V, in open capillaries and are uncorrected. The mid-IR spectra (KBr, v/cm^{-1}) were recorded on a CARY 630 FT-IR spectrophotometer (Agilent, Santa Clara, CA, USA). NMR spectra $(^1H$ NMR and ^{13}C NMR) were recorded on a Bruker EX-500 MHz ($^1\rm H$: 500 MHz, $^{13}\rm C$: 125 MHz) spectrometer (Bruker, USA) at room temperature. Tetramethylsilane (TMS) was used for internal calibration (¹H NMR and 13 C NMR: 0.00 ppm). Chemical shifts were reported in parts

Scheme 1 Previous organic transformations and our approach for the chemical synthesis of 8 and 9. Reagents and conditions: (a) 1 (16.5 g 0.1 mol, 1.0 equiv.), 2 (19.1 g, 0.1 mol, 1.0 equiv.), DCM, TEA (10.1 g, 0.1 mol, 14.4 mL, 1.0 equiv.), stirring, 8 h, 95% yield; (b) 8 (16 g, 0.05 mol, 1.0 equiv.), NH₂NH₂·H₂O 100% (10 mL, 0.2 mol, 4.0 equiv.), pyridine (3.95 g, 4.0 mL, 0.05 mol, 1.0 equiv.), EtOH, reflux, 10 h, 91% yield.

per million (ppm) on the δ scale and relative to residual solvent peaks (DMSO- d_6 : ¹H: 2.50 ppm, ¹³C: 39.5 ppm). Coupling constants (J) are reported in Hz with the following abbreviations used to indicate splitting: $s = singlet$, $d = doublet$, $t = triplet$, q $=$ quartet, $m =$ multiplet, $br =$ broad signal. Electron ionization mass spectra (EI-MS) were obtained using a GC MS-QP 2010 plus mass spectrometer (Shimadzu, Kyoto, Japan) at an electron voltage of 70 eV, with a time of 10 min, scan speed of 1000, and m/z range of 50-500.

2.2 Synthesis and characterization measurements

2.2.1 Chemical synthesis of ethyl 4-(4-methylphenylsulfonamido)benzoate (p-TSAE, 8, CAS: 739-33-3). 24,25 For the large-scale synthesis under a nitrogen atmosphere, into a 500 mL, two-necked, oven dried flat-bottomed flask that was equipped with a magnetic stir bar, ethyl 4-aminobenzoate (6, benzocaine, CAS: 94-09-7, 16.5 g, 0.1 mol, 1.0 equiv.) was dissolved in dichloromethane (DCM, 50 mL) containing triethylamine (TEA, 10.1 g, 0.1 mol, 14.4 mL, 1.0 equiv.) (Scheme 1). A solution of 4-toluenesulfonyl chloride (TsCl, 5, CAS: 98-59-9, 19.1 g, 0.1 mol, 1.0 equiv.) in DCM (50 mL) was added portionwise over 30 min and the reaction mixture was stirred at room temperature for 8 h. The precipitated solid was isolated *via* vacuum filtration and washed with DCM (3×20 mL), then air dried. The obtained ester 8 was employed in the next step without additional purification.

yield: 95%; mp 205 °C (lit. mp 202–206 °C (ref. 24)); mid FT-IR (KBr, v/cm⁻¹) 3216 (N-H stretching), 3062, 2985 (aromatic C-H stretching), 2942, 2904 (aliphatic C–H stretching), 1693 (ester $C=O$ stretching), 1604 ($C=C$ stretching), 1511 (N-H bending),

1477 (aliphatic C-H bending), 1338 (asym. $O = S = O$ stretching), 1288 (aromatic amine C–N stretching), 1234 (aromatic ester C–O stretching), 1158 (sym. O=S=O stretching); 1 H NMR (500 MHz, DMSO- d_6 , ppm) δ 10.81 (s, 1H, NH), 7.80–7.76 (m, 2H, ArCH), 7.71–7.67 (m, 2H, ArCH), 7.28 (d, $J = 8.0$ Hz, 2H, ArCH), 7.22– 7.19 (m, 2H, ArCH), 4.17 (q, $J = 7.1$ Hz, 2H, ester CH₂), 2.24 (s, 3H, ArCH₃), 1.19 (t, $J = 7.1$ Hz, 3H, ester CH₃). ¹³C NMR (125 MHz, DMSO, ppm) δ 165.66 (ester C=O), 144.11 (quat-C-CH₃), 142.99 (quat-C-NH), 137.02 (quat-C-SO₂), 131.05 (ArCH), 130.25 (ArCH), 127.30 (ArCH), 125.21 (quat-C-COOEt), 118.63 (ArCH), 118.57 (ArCH), 60.92 (ester CH₂), 21.36 (ArCH₃), 14.58 (ester CH₃); EI-MS for C₁₆H₁₇NO₄S [M]⁺ m/z (%): 319 (30), 274, 226, 182, 155, 92, 91 (100), 65.

2.2.2 Chemical synthesis of 4-(4-methylphenylsulfonamido)benzohydrazide (p-TSAH, 9).²⁶ An ovendried 500 mL, one-necked, flat-bottomed flask, equipped with a magnetic stirring bar, was charged with carboxylate ester 8 (16 g, 0.05 mol, 1.0 equiv.) in EtOH (150 mL) containing pyridine (3.95 g, 4.0 mL, 0.05 mol, 1.0 equiv.). Next, the reaction mixture was heated at 60 °C for 10 min to activate the ester group. After that, hydrazine hydrate 100% (10 mL, 0.2 mol, 4.0) equiv.) was added and the reaction mixture was heated under reflux conditions in an oil bath for 10 h. A colorless solid was formed while hot and the reaction solution was then cooled to room temperature. The precipitated solid was collected by vacuum filtration and washed with hot EtOH (3×20 mL), then air dried.

yield: 91%; mp 272–274 °C (lit. mp 270–272 °C (ref. 26)); mid FT-IR (KBr, v/cm^{-1}) 3321, 3279 (NH₂ stretching), 3146 (N–H stretching), 3048, 2940 (aromatic C–H stretching), 2875 (aliphatic C-H stretching), 1649 (hydrazide C=O stretching), 1607 (C=C

stretching), 1579, 1536 (N–H bending), 1476 (aliphatic C–H bending), 1335 (asym. $O = S = O$ stretching), 1292 (aromatic amine C-N stretching), 1152 (sym. O=S=O stretching); 1 H NMR (500 MHz, DMSO- d_6 , ppm) δ 10.51 (s, 1H, NH), 9.63-9.59 (m, 1H, NH), 7.67 (dd, $J = 8.2$, 2.6 Hz, 4H, ArCH), 7.31 (d, $J = 8.0$ Hz, 2H, ArCH), 7.12 (d, $J = 8.5$ Hz, 2H, ArCH), 4.45-4.38 (m, 2H, NH₂), 2.27 (s, 3H, ArC H_3); ¹³C NMR (126 MHz, DMSO, ppm) δ 165.93 (hydrazide $C=O$), 144.09 (quat-C-CH₃), 140.99 (quat-C-NH), 137.01 (quat-C-SO₂), 130.35 (quat-C-CONHNH₂), 128.85 (ArCH), 128.75 (ArCH), 127.28 (ArCH), 118.82 (ArCH), 21.48 (ArCH₃); EI-MS for $C_{14}H_{15}N_3O_3S$ $[M + 2]^+$ m/z (%): 307 (7), 295, 241, 196, 162, 108, 92, 91 (100), 63.

2.3 Electrochemical measurements

A three-electrode cell containing a fresh 100 ml aerated solution of 1.00 M HCl with/without different concentrations of the synthesized inhibitors was first left for 1 h to obtain a steady state open circuit potential. After the solution had reached the open circuit potential (E_{OCP}) , three consecutive electrochemical techniques, electrochemical frequency modulation (EFM), electrochemical impedance spectroscopy (EIS) and potentiodynamic polarization (PDP) were conducted at a temperature of 303.15 K. The first cell electrode was a mild steel working electrode (WE), with a total exposed surface area of 1 cm^2 and elemental composition of 97.929% Fe, 0.330% C, 0.030% Mo, 0.836% Mn, 0.043% Al, 0.389% Si, 0.157% Cr, 0.033% S, 0.075% Ni, 0.014% Co, 0.149% Cu, and 0.015% W. The second electrode was a graphite counter electrode (CE), and the third was the saturated calomel electrode (SCE) as a reference electrode. The investigated inhibitor concentrations were 1.00 \times 10 $^{-5}$, 5.00 \times 10^{-5} , 1.00×10^{-4} , 5.00×10^{-4} and 1.00×10^{-3} M. EFM was achieved by applying 2 Hz and 5 Hz frequencies, with a base frequency of 0.1 Hz, and a perturbation signal of 10 mV, for 16 cycles of repeated sinusoidal waveform. The highest frequency should be at least twice the lower one, and slow enough to eliminate the double layer charging contribution to the current response. EIS measurements were carried out at the OCP in a frequency range from 1.0×10^5 to 1.0×10^{-1} Hz using 10 mV as the AC sinusoidal perturbation. PDP tests were carried out from −500 mV to 500 mV against the OCP with a sweeping rate of 0.5 mV s^{-1} . Prior to each experiment, the WE was grounded with Si–C paper (300–800 grit), washed with fresh double distilled water, degreased with acetone, and dried. A Gamry REF 3000 potentiostat/galvanostat/Zra instrument was used for the measurements and the Echem Analyst 7.8.2 program for analysis, fitting and graphing of the data.

2.4 Surface examination by SEM and XPS

Scanning electron microscopy (SEM) and X-ray photoelectron spectroscopy (XPS) were used to examine the appearance of the mild steel surface, and therefore the element properties after immersion in the corrosive solution for 24 h with and without the addition of p -TSAE and p -TSAH molecules. SEM tests were done on an SEM, JOEL, JSM-T20, Japan instrument with an acceleration voltage of 30.00 kV and a magnification of 1000

times. XPS was carried out on a Scienta Omicron Nanotechnology instrument, Germany.

2.5 Theoretical models

The geometries of the synthesized molecules in three dimensions were built using the Gaussview 6 (ref. 27) cartesian coordinates builder and were optimized following the convergence criteria of the Gaussian 09 software package.²⁸ The theoretical procedure included Becke-style 3-parameter density functional theory using the Lee–Yang–Parr correlation functional, B3LYP, which was of reasonable computational cost for the specified molecules.²⁹–³¹ The convergence was reached at a maximum force of 0.1 \times 10⁻⁵ and displacement of 1.6 \times 10⁻⁵, while the thresholds for energy and displacement were 45×10^{-5} and 180 \times 10⁻⁵, respectively. The molecular orbitals were represented by the Gaussview 6 MO editor to accurately approximate the exact orbital densities. The frontier molecular orbital surfaces were adjusted to an isovalue of 0.02 and density of 0.0004 and presented as mesh surfaces using the molecular editor of Gaussview 6.²⁷ The natural bond orbital surfaces were obtained using NBO version 3.1 (ref. 32) incorporated in the Gaussian 09 package. The NBO surfaces were extracted using the same edits as for the frontier molecular orbitals to accurately compare them. The parameters accounting for global reactivity were calculated using the HOMO and LUMO eigenvalues obtained from the DFT calculation.³³ The structures were then used in a single point calculation to obtain the molecule energies, charges and populations using Hirshfeld surface analysis, in which there was a neutral molecule, singly-charged cation and singly-charged anion, in order to perform local reactivity analysis using Fukui functions. The closed shell model in Gaussian 09 forces each electron pair, in neutral molecules, into a single spatial orbital, while switching to the open shell model, to treat electrons spin up and spin down, in charged molecules. Paper

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Monte Carlo (MC) simulation was used to investigate the interactions of p -TSAE and p -TSAH with the mild steel surface. The adsorption locator module of the Materials Studio 2017 software was utilized to examine how these molecules interacted with the Fe (110) surface.³⁴ The COMPASS force field was used in all system adsorption computations. To verify that a sufficient depth was obtained, a crystal of Fe (110), with an edge of 30 Å, was built. This crystal was subsequently magnified to form a supercell (11 \times 11). The optimal structures of **p-TSAE** and p -TSAH were investigated in order to find the most stable adsorption configurations on the Fe (110) surface in the neutral and protonated forms in the gas phase and in the presence of water and HCl molecules.

3. Results and discussion

3.1 Synthesis and characterization of inhibitors

3.1.1 Synthetic route of the ester (p-TSAE, 8) and hydrazide (p-TSAH, 9). As shown in Scheme 1, our target sulfonamide ethyl ester $(p\text{-TSAE}, 8)$ has been previously synthesized via modern organic transformations that were mediated by Pd catalysis according to Shekhar et al.'s paper²⁴ and Cao et al.'s

Fig. 1 FT-IR spectra of the ester 8 (p-TSAE) and the acyl hydrazide 9 (p-TSAH).

Fig. 2 $^{-1}$ H NMR spectra (500 MHz, DMSO- d_6) of the ester 8 (p-TSAE) and the acyl hydrazide 9 (p-TSAH).

work.²⁵ The corresponding methyl ester was also recently synthesized via a two-step conventional method that was documented in 2020 by Hamdani group's,²⁶ who further converted it to our target hydrazide (p-TSAH, 9) with 78% yield.²⁶ In our large scalable approach that depends on the Hinsberg reaction, commercially available starting material 4-toluenesulfonyl chloride (tosyl chloride, 5) was directly coupled with ethyl 4-aminobenzoate (6) in dichloromethane (DCM) containing triethylamine $(Et₃N)$ to afford the desired sulfonamide ethyl ester $(p\text{-TSAE}, 8)$ in excellent yield (95%). Then, the pyridinecatalyzed hydrazinolysis of 8 with 100% hydrazine hydrate was carried out in refluxing ethanol to generate the corresponding acyl hydrazide derivative (p-TSAE, 8) in high yield (91%). The target ester 8 and its acyl hydrazide 9 were characterized by physical properties, FT-IR, NMR, and mass spectrometry analysis. Paper

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3.1.2 FT-IR spectral analysis. The mid-infrared spectrum of 8, recorded in KBr, shows a strong band at 3216 cm⁻¹ corresponding to the N-H stretching vibration, 26 which proves the sulfonamide formation. The sp^2 -(aromatic stretching) and sp^3 -CH (aliphatic stretching) valence vibrations are assigned to 3062-2904 cm^{-1} . Additionally, strong bands in the 1693 and 1604 cm⁻¹ regions can be attributed to C=O and C=C stretching vibrations. The $O=$ S $=$ O stretching bands are observed at 1338 and 1158 cm^{-1} (ref. 26) (Fig. 1). The solid

mode IR spectrum of acyl hydrazide 9 shows new frequencies, one of which is a slightly split medium band at 3321 and 3279 cm⁻¹ due to NH₂ vibration, and the other a medium band at 3146 cm−¹ , arising from N–H stretching. Moving to the double bond region, we detect two medium bands at 1649 and 1607 cm−¹ , which are probably attributable to the acyl hydrazide $C=O$ and aromatic $C=C$ moieties. Upon comparison with ester 8, $O=$ S $=$ O shows two similar bands with slight differences at 1335 and 1152 cm⁻¹ (ref. 26) (Fig. 1).

3.1.3 NMR spectroscopic analysis. The 1 H NMR spectrum of sulfonamide ethyl ester 8 (Fig. 2), recorded in DMSO- d_6 inspected from left to right first reveals a singlet ¹H resonance at δ 10.81 ppm, which is attributable to the NH proton,²⁴ proving the formation of a sulfonamide linkage. In the aromatic region, eight ¹H chemical shifts are generated from δ 7.78 to 7.20 ppm, which correspond to the typical spin AA′ XX′ pattern of two 1,4 disubstituted phenyl rings. Moving to the upfield region, we find three proton resonances. Two among them are quartets (q) one at δ 4.17 ppm, and a triplet (t) signal at δ 1.19 ppm, which clearly belongs to the ester group $(J = 7.1 \text{ Hz})$. The remaining singlet signal at δ 2.24 ppm with the highest intensity must belong to aromatic methyl protons.²⁴ In the clear carbon NMR analysis of 8 (Fig. 3), a ¹³C resonance at δ 165.66 ppm can be directly assigned to the carbonyl ester group.^{24,25} The nonprotonated and CH aromatic carbon signals ranging from

Fig. 3 13 CNMR spectra (125 MHz, DMSO) of the ester 8 (p-TSAE) and the acyl hydrazide 9 (p-TSAH).

 δ 144.11 to 118.57 ppm are also shown. In the aliphatic region, methylene ester and its methyl carbon show resonances at δ 60.92 and 14.58 ppm, respectively, whereas the aromatic methyl carbon is seen as expected at δ 21.36 ppm.^{24,25}

As shown in the $1D^{-1}H$ NMR spectrum of acyl hydrazide derivative 9 (Fig. 2), the sulfonamide NH is slightly shielded at δ 10.51 ppm and a new peak corresponding to the hydrazide group –CONHNH₂ is present at δ 9.61 ppm, confirming the successful hydrazinolysis step.²⁶ The typical AA'XX' pattern of two 1,4-disubstituted benzene rings is observed between δ 7.67 and 7.12 ppm. Prior to the aliphatic region, the $NH₂$ group is observed as a broad singlet at δ 4.42 ppm.²⁶ The absence of a peak corresponding to an ethyl ester group is indicative of the complete conversion of 8 to 9 via the hydrazinolysis reaction. As expected, the aromatic methyl group is shown as a singlet signal with the highest intensity at δ 2.27 ppm. Additionally, the wellanalyzed 13 C NMR spectrum of 9 (Fig. 3), reported for the first time, confirms the absence of the ethyl ester group, and shows the carbonyl chemical shift at δ 165.93 ppm, whilst the methyl of the tosyl fragment is shown at δ 21.48 ppm.

3.1.4 EI mass spectrometry. In the EI mass spectrum of 8, the tallest basic peak at $m/z = 91$ (100%)²⁵ is assigned to the formation of the p-tolyl fragment ion. Another peak with a relative intensity of 20% is generated at $m/z = 155$ due to the formation of the tosyl ion. Additionally, in the mass stick diagram of 8, the line generated by the heaviest ion flowing via the machine at $m/z = 319 \,[\mathrm{M}]^+$ (30%) is related to the parent ion

(Fig. 4). Concerning the fragmentation pattern of methylsulfonamide ester 8 (Scheme 2), radical cation A at $m/z = 319$ undergoes homolysis via sigma bond cleavage (σ -cleavage) of the carbon–oxygen (C–O) bond giving rise to the fragment ion B at $m/z = 274$ after the removal of an ethoxy radical (C₂H₅O⁻) of 45 Da. The radical migrates from the alpha carbon to the carbonyl oxygen radical that has a strong tendency for electron pairing, which acts as a driving force of this fragmentation type (radical site-initiated fragmentation). Upon the further fragmentation of B by heterolytic dissociation of the S-N bond,³⁵ a common fragment ion C of aromatic sulfonamides at $m/z =$ 155 is formed.36,37 The inductive effect of the positive nitrogen results in the fragmentation step (charge site-initiated cleavage). In addition, the extrusion of SO_2 (loss of 64 Da)³⁷ through the direct heterolytic cleavage of the C–S bond affords the most common *p*-tolyl fragment ion D_1 at $m/z = 91$ with the highest abundance (100%, base peak). Further skeletal rearrangement leads to the conversion of D_1 to D_2 and D_3 ions, which affords the last fragment ion **E** at $m/z = 65$ due to the expulsion of acetylene gas of 26 Da. RSC Advances

8 14.1.11 to 118.57 ppm are also shown in the aliphatic region, (Fig. 1). Concerning the figherein and model

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Mass analysis supported the chemical structure of 9 (nonreported) with a molecular ion peak at $m/z = 307$ [M + 2]⁺ (5%) and the p-tolyl fragment ion as the same base peak as that of 8 at $m/z = 91$ (100%) (Fig. 4). Ionization occurs by protonation of 9 to form the molecular ion peak at $m/z = 307$. Protonated hydrazide $[M + 2H]^{2+}$ is fragmented to afford the corresponding amide **F** at $m/z = 290$ by elimination of NH₃ (loss of 17 Da).³⁸

Fig. 4 EI-MS spectrum of the ester 8 (p-TSAE) and the acyl hydrazide 9 (p-TSAH)

Additional fragmentation of F via inductive cleavage of the C–N bond yields the corresponding acylium ion **B** at $m/z = 274$ after the loss of NH_3 of 17 Da.³⁹ Additionally, F can undergo decarbonylation (loss of CO, 28 Da) to produce amine G at $m/z = 262$. Both B and G may further dissociate to exhibit the prospective phenyl cation H at $m/z = 246$. Fragmentation of H after the neutral loss of a diacetylene gas molecule of 50 Da resulted in the formation of azirine ion I_1 at $m/z = 196$. Additionally, H can be fragmented by the loss of HCN of 27 Da to produce another possible ion I_2 at $m/z = 219$. Both fragment cations I_1 and I_2 can further disassociate to form the known ion C at $m/z = 155$,^{36,37} which upon protonation converts to cation J. The rearrangement of J drives the elimination of SO and the formation of an ion at $m/z = 109$,³⁶ which dehydrates to generate the base peak at $m/z = 91$ (Scheme 3).³⁷

3.2 Electrochemical measurements

3.2.1 Open circuit potential (OCP). The corrosion potential, simply the potential difference between the working and reference electrode, was probed versus time to obtain a steady

Scheme 3 Proposed fragmentation pathways of methyl-sulfonamide hydrazide 9.

Fig. 5 OCP–time curves for mild steel in 1.0 M HCl solution with and without different concentrations of p -TSAH and p -TSAE compounds at 30 ° \mathcal{C}

state for the investigated systems prior to the following electrochemical experiments. At a steady state, the rate of the corrosion reactions is assumed to be constant. A one-hour experiment was conducted at a temperature of 303.15 K for this purpose. The resultant potential vs. time curves for the mild steel/1.0 M HCl/ $(p$ -TSAH or p -TSAE) systems are presented in Fig. 5. It can be inferred from the curves that the potential through all the solutions of p -TSAH or p -TSAE reached minimal fluctuations $(\pm 5 \text{ mV or less})$ almost at the same time. While measuring the corrosion potential is an easy task for a potentiostat, it guarantees the accuracy of the following EFM, PDP and EIS measurements. Moreover, upon adding concentrations of p-TSAH or p-TSAE inhibitors to the blank solutions, a positive shift in the potential can be noticed, implying the adsorption of these molecules to the mild steel surface and blocking of the surface corroded active sites.⁴⁰

3.2.2 Electrochemical frequency modulations. After the OCP curves of the tested solutions had achieved stability, EFM was immediately carried out to evaluate the corrosion inhibition behavior of the p -TSAH and p -TSAE molecules for the mild steel surface in a 1.0 M HCl solution. The EFM technique has several advanced characteristics, including being a nondestructive and reliable corrosion testing technique.⁴¹ Fig. 6 shows the intermodulation spectra for mild steel immersed in the corrosive medium in the absence and presence of p-TSAH molecules at a temperature of 303.15 K. These spectra show two intense and characteristic peaks at 0.2 and 0.5 Hz related to the selected excitation frequencies, besides other low intensity peaks related to the harmonic sum, differences, and multiples of the selected two excitation frequencies.⁴² The current responses at the two selected excitation frequencies determine

the corrosion rate, and therefore the inhibition efficiencies. The extracted electrochemical parameters from the EFM measurements are shown in Table 2. The reliability of the obtained data was checked through the values of the causality factors. It is shown in Table 2 that the empirical values of CF-2 and CF-3 are close to their theoretical values (2 and 3).⁴³ The EFM spectra for the inhibited solutions are observed in more negative current areas than the corrosive 1 M HCl solution, so the presence of p-TSAH and p -TSAE molecules decreases the current flow between the steel surface and the reference electrode. The highest value of i_{corr} being in the inhibitor free solution with respect to the solutions containing p -TSAH or p -TSAE molecules indicated that corrosive ions such as Cl− are more activated and a high corrosion rate occurred. However, after the addition of p -TSAH or p-TSAE molecules, the corrosion current densities are reduced, and the steel surface protection is enhanced. This is related to the adsorption of p -TSAH or p -TSAE via the organic hetero atoms (S, N and O) and the aromatic moieties, blocking the corrosion active sites and slowing down the current flow.⁴⁴ The values of the corrosion current are 2791 μ A cm⁻² for the blank 1 M HCl solution, 1453 µA cm⁻² for the **p-TSAE** compound and 1287 μA cm⁻² for the **p-TSAH** compound at the lowest concentration of 1.00 \times 10⁻⁵ M, so the corrosion of steel is decreased by the addition of p -TSAH or p -TSAE molecules. The successive addition of the inhibitor molecules gradually decreases the current density. As an example (for the p -TSAH molecule), i_{corr} is 599.5 µA cm⁻² at 5.00 × 10⁻⁵ M, 463.6 µA cm⁻² at 1.00 × 10⁻⁴ M, 303.5 µA cm⁻² at 5.00 × 10⁻⁴ M and 158.6 μA cm⁻² at 1.00 × 10⁻³ M. This situation is observed also for the corrosion rate, as the highest value was observed for the 1.0 M HCl solution (1275.00 mpy), and this was greatly reduced

Fig. 6 Intermodulation spectra for steel in 1.0 M HCl in the absence and presence of different concentrations of the p-TSAH compound at 30 °C.

Table 2 Electrochemical kinetic parameters^a obtained by the EFM technique for steel in the absence and presence of various concentrations of p-TSAE and p-TSAH inhibitors in 1.0 M HCl at 30 °C

Inhibitor name	Conc. (M)	$i_{\rm corr}$ (µA cm ⁻²)	$\beta_{\rm a}$ (mV dec ⁻¹)	β_c (mV dec ⁻¹)	$CF-2$	$CF-3$	k (mpy)	θ	$\eta_{\rm EFM}$ %
Blank		2791	100.4	113.1	1.763	3.155	1275.00		
p -TSAE	1.00×10^{-5}	1453	99.04	186	2.013	2.853	664.10	0.4794	47.94
	5.00×10^{-5}	745.4	102.5	131.1	1.989	2.992	340.60	0.7329	73.29
	1.00×10^{-4}	534.8	116.3	121.3	2.204	3.076	244.40	0.8084	80.84
	5.00×10^{-4}	380.6	116.5	124	1.992	2.887	173.90	0.8636	86.36
	1.00×10^{-3}	209	94.66	114.1	1.862	3.131	95.50	0.9251	92.51
p -TSAH	1.00×10^{-5}	1287	81.73	133.6	1.967	3.424	587.90	0.5389	53.89
	5.00×10^{-5}	599.5	87.95	120.8	1.986	3.061	273.90	0.7852	78.52
	1.00×10^{-4}	463.6	117.6	120.9	1.982	3.037	211.80	0.8339	83.39
	5.00×10^{-4}	303.5	111.4	115.1	2.074	3.099	138.70	0.8913	89.13
	1.00×10^{-3}	158.6	84.18	90.8	1.729	2.975	72.46	0.9432	94.32

 a $E_{\rm corr}$ is the corrosion potential; $i_{\rm corr}$ is the corrosion current density; β_a and β_c are Tafel constants for the anode and cathode; k is the corrosion rate; θ is the surface coverage; η_{EFM} is the inhibition efficiency.

to 664.10 mpy and 587.90 mpy by the addition of the lowest dose $(1.00 \times 10^{-5} \text{ M})$ of the studied molecules. The values of β_a and β_c are impacted by the addition of **p-TSAH** or **p-TSAE** molecules as evidence of the mixed inhibition performance of these two molecules by the adsorption on both cathodic and anodic active sites. The protective ability of p -TSAH and p -TSAE were estimated from i_{corr} as per the following:⁴⁵

$$
\eta = \theta \times 100 = \frac{\left(i_{\text{corr}}^0 - i_{\text{corr}}\right)}{i_{\text{corr}}^0} \times 100\tag{1}
$$

 i_{corr}^0 = corrosion current density of the blank solution, i_{corr} = corrosion current density of any desired concentration of p-TSAH or **p-TSAE** molecules.

The results confirmed that the high protecting ability of the p-TSAH and p-TSAE molecules approached 94.32% (p-TSAH) and 92.51% (p -TSAE) at 1.00 × 10⁻³ M. The slight difference in inhibition efficiency between the p -TSAH molecule and the p -TSAE molecule might be due to the presence of extra nitrogen atoms in the molecular skeleton of the former. This facilitates its adsorption to enhance the protection.

3.2.3 Potentiodynamic polarization measurements. Tafel plots, from the potentiodynamic polarization method, for the corrosion of mild steel in 1.0 HCl solution (blank, with p-TSAH and with p -TSAE), are presented in Fig. 7. It can be observed through the plots that the polarized curves were affected when the inhibitors were added to the blank solution. A noticeable decrease in the rate of both anodic and cathodic reactions could be inferred from the comparison of the blank ν s. inhibitor (ν - TSAH and p-TSAE) curves. However, the anodic reactions were altered more noticeably by the presence of p -TSAH and p -TSAE molecules than the cathodic ones. The latter are in the close vicinity of the curves of the 1.0 M HCl blank solution. Overall, a reduction in the corrosion reactions in both the anodic and cathodic directions could be observed, nevertheless the anodic reactions (signifying mild steel dissolution) were more retarded.⁴⁶ The parallel polarization curves support the idea that the addition of p -TSAH or p -TSAE does not alter the pathway of the corrosion reaction.

The results derived from the Tafel plots, by fitting and extrapolation of the log current versus potential curves back to their intersections, in addition to the inhibition efficiency and surface coverage, are shown in Table 3. The data reveal that the Tafel constants $(\beta_a$ and $\beta_c)$ move to lower values when **p-TSAH** and p -TSAE samples are used in the corrosion systems. This suggests that p -TSAH and p -TSAE molecules contribute to the electrochemical reactions (both cathodic and anodic) through an adsorption mechanism promoting an intact metallic surface.⁴⁷ Accordingly, the addition of p -TSAH or p -TSAE molecules significantly decreases the corrosion current i_{corr} ; the minimum obtained value is 580 µA cm⁻² in the case of **p-TSAH** and 710 µA cm⁻² in the case of **p-TSAE** at 1.00 \times 10⁻³ M, Table 3. Correspondingly, E_{corr} shifted to more positive values in the presence of p-TSAH or p-TSAE molecules while the difference, *i.e.* ($\Delta E_{\text{corr}} = |E_{\text{corr}}(\text{blank}) - E_{\text{corr}}(\text{inhibitor})|$), is less than 85 mV in most solutions. Therefore, a mixed type inhibition behavior of p -TSAH and p -TSAE molecules could be suggested.⁴⁸ RSC Advances

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Fig. 7 Potentiodynamic polarization curves for the corrosion of steel in 1.0 M HCl in the absence and presence of different concentrations of p -TSAH and p-TSAE compounds at 30 °C.

Table 3 Electrochemical parameters^a for steel dissolution in 1.0 M HCl solution containing different concentrations of the p-TSAE and p-TSAH inhibitors obtained from polarization measurements at 30 °C

Inhibitor name	Conc. (M)	$E_{\rm corr}$ vs. SCE (mV)	$i_{\rm corr}$ (µA cm ⁻²)	$\beta_{\rm a}$ (mV dec ⁻¹)	β_c (mV dec ⁻¹)	$k \text{ (mpy)}$	$\Delta E_{\rm corr}$ (mV)	θ	$\eta_{\rm PDP}$ %
Blank		-523	8624	305	333	4137			
p -TSAE	1.00×10^{-5}	-471	4050	607.8	572.6	1852	52	0.530	53.02
	$5.00\,\times\,10^{-5}$	-432	2730	526	429.2	1248	91	0.683	68.33
	1.00×10^{-4}	-454	1670	234.9	249.6	762.6	69	0.806	80.63
	5.00×10^{-4}	-466	1170	295.1	302.6	536	57	0.864	86.43
	1.00×10^{-3}	-437	710	183.8	209.4	324.5	86	0.918	91.76
p -TSAH	1.00×10^{-5}	-471	3740	579.1	599.6	1709	52	0.566	56.61
	5.00×10^{-5}	-465	2550	243.2	243.5	1164	58	0.704	70.42
	1.00×10^{-4}	-467	1410	219.4	232.1	646	56	0.836	83.64
	5.00×10^{-4} 1.00×10^{-3}	-450	843	195.3	221.9	385.1	73	0.902	90.22
a $E_{\rm corr}$ is the corrosion potential; $i_{\rm corr}$ is the corrosion current density; β_a and β_c are Tafel constants for the anode and cathode; k is the corrosion rate; θ is the surface coverage; $\eta_{\rm p}$ is the inhibition efficiency.									
			Furthermore, the tendency for passivation to take place by p - TSAH and p -TSAE is a function of i_{corr} according to eqn (1), as		by the successive addition of p -TSAH or p -TSAE molecules because of their adsorption on the metallic surface and the				
described in section 3.2.2.			According to the data presented in Table 3, the protection abilities, or η_{PDP} , of p-TSAH and p-TSAE from the PDP measurements reached maxima of 93.27% (p-TSAH) and 91.76% (p-TSAE) at 1.00 \times 10 ⁻³ M. The protection ability gradually improved upon the successive addition of either p- TSAH or p-TSAE molecules as extra molecules were adsorbed,		formation of a layer that protects it from corrosion. Adsorption considerations will be discussed in detail in a separate section. The equivalent circuit proposed for the studied systems is a physical model which demands that the model's components are attributed to a physical process in the electrochemical corrosion reaction. Accordingly, the proposed circuit model in Fig. 8 was built up using three components: two resistors, one				
			and more of the mild steel surface was covered. The adsorbed organic layers reduced the interaction between the metallic		accounting for the solution resistance (R_s) and the other accounting for the charge transfer resistance (R_{ct}) in addition to				
			surface and corrosive mineral acid. ⁴⁹ Therefore, the presence of		a component with a constant phase. In general, the impedance				
			several hetero atoms (S, N and O) in addition to donor-acceptor		of a constant phase element CPE (Z_{CPE}) can be determined				
			interactions from the aromatic conjugation π -systems to the		using the following equation: ⁵¹				
increased the protection ability.			empty d-shell of the mild steel facilitated the adsorption and			$Z_{\rm CPE} = [Q(j\omega)^n]^{-1}$			

3.2.4 Electrochemical impedance spectroscopy. To highlight the mechanism of the electrochemical process that takes place at the mild steel surface, EIS measurements were performed. EIS outcomes for the mild steel/1.0 M HCl/ $(p$ -TSAH or p-TSAE) systems are presented in Fig. 8, which includes Nyquist and Bode plots and the corresponding equivalent circuit model. Furthermore, corresponding EIS fitting data are collected in Table 4. Both the Nyquist and Bode plots are characterized by a single time constant that is related to the corrosion as a charge transfer process taking place at the metallic surface.⁵⁰ The semicircles obtained for the Nyquist plots are imperfect capacitive loops, at the high frequency regions, which could be attributed to metallic surface heterogeneity and roughness caused by exposure to the aggressive mineral acid. In addition, it can be seen in Fig. 8 that the semi-circle is single and depressed in the case of both the blank and the inhibited solutions. Capacitive semicircles corresponding to charge-transfer and adsorption processes are often depressed, therefore, it can be concluded that the charge transfer pathway controls both the corrosion and the inhibition reactions, while the used inhibitors control these reactions through adsorption to the metal surface. Additionally, the diameter of the semi-circle is gradually increased

$$
Z_{\rm CPE} = [Q(j\omega)^n]^{-1} \tag{2}
$$

Q = proportionality factor; ω = angular frequency; $j = -1$, and n = variation parameter ($-1 \le n \le +1$). If $n = 0$, CPE = pure resistor; $n = -1$, CPE = inductor; $n = +1$, CPE = pure capacitor.

While a double-layer capacitor is expected to proceed ideally, EIS experiments often do not behave ideally. In such a case, the double-layer capacitor in a real cell acts like a constant phase element (CPE).⁵²

The capacitance of the double layer could then be determined by the following relationship:

$$
C_{\rm dl} = [QR_{\rm ct}^{1-n}]^{1/n} \tag{3}
$$

The C_{dl} reduction (Table 4) in the presence of **p-TSAH** or **p**-TSAE could be attributed to a decreased local dielectric constant and increased double layer thickness due to the replacement of H2O molecules adsorbed on the metallic surface by the organic inhibitors through the active sites.⁵³ The protecting ability is related to the charge transfer resistance R_{ct} by the relationship:

$$
\eta_z = \theta \times 100 = \frac{(R_{\rm ct} - R_{\rm ct}^{\circ})}{R_{\rm ct}} \times 100
$$
 (4)

Fig. 8 (a) Nyquist and (b) Bode modulus, phase angle plots (the dots show experimental results and the lines show fitted data), and (c) the equivalent circuit model used to fit the EIS data for steel in 1.0 M HCl in the absence and presence of different concentrations of p-TSAH and p-TSAE compounds at 30 °C.

When the capacitance goes down, in contrast, the impedance goes up when the connection is in series. This is a consequence of the inverse relationship between the capacitance and impedance. It can be seen in Table 4 that R_{ct} gradually increases upon the addition of p -TSAH or p -TSAE inhibitors to reach maxima of 109.8 Ω cm² (p-TSAH) and 72.26 Ω cm² (for p-TSAE)

Table 4 EIS parameters^a for the corrosion of steel in 1.0 M HCl in the absence and presence of different concentrations of p-TSAE and p-TSAH inhibitors at 30 °C

Inhibitor	Conc. (M)	$R_{\rm s}$ $(R_{\rm u})$ $(\Omega$ cm ²)	$R_{\rm ct}$ $(R_{\rm p})$ $(\Omega$ cm ²)	Q $(\mu \Omega^{-1} s^n cm^{-2})$	\boldsymbol{n}	C_{dl} $(\mu$ F cm ⁻²)	chi squared	S	α°	θ	η_z (%)
Blank		1.082	6.884	478.50	0.8836	225.363	2.25×10^{-2}	-0.365	-42.07		
p -TSAE	1.00×10^{-5}	1.612	17.15	366.70	0.8980	267.822	6.60 \times 10 $^{-3}$	-0.469	-49.87	0.5986	59.86
	5.00×10^{-5}	1.688	22.94	154.90	0.8780	97.423	6.83 \times 10 $^{-3}$	-0.557	-51.01	0.6999	69.99
	1.00×10^{-4}	1.628	30.30	240.30	0.7819	115.712	7.86×10^{-3}	-0.515	-47.67	0.7728	77.28
	5.00×10^{-4}	2.169	43.48	75.59	0.8570	42.753	3.34×10^{-3}	-0.657	-54.69	0.8417	84.17
	1.00×10^{-3}	1.264	72.26	123.00	0.7149	18.702	1.48×10^{-3}	-0.580	-50.23	0.9047	90.47
p -TSAH	1.00×10^{-5}	1.602	18.65	201.30	0.8670	121.667	6.78 \times 10 $^{-3}$	-0.420	-48.79	0.6309	63.09
	5.00×10^{-5}	1.582	24.6	218.20	0.8389	124.431	6.95×10^{-3}	-0.429	-50.02	0.7202	72.02
	1.00×10^{-4}	1.996	34.33	109.00	0.8303	55.692	5.04×10 $^{-3}$	-0.517	-50.63	0.7995	79.95
	5.00×10^{-4}	1.257	54.29	479.00	0.7459	138.167	1.15×10^{-3}	-0.502	-50.18	0.8732	87.32
	1.00×10^{-3}	1.116	109.8	377.50	0.8446	210.160	7.65×10^{-4}	-0.746	-65.77	0.9373	93.73

 $a_{\rm Rs}$ = solution resistance, $R_{\rm ct}$ = charge transfer resistance, Q, n = variation parameters, $C_{\rm d}$ = double layer capacitance, θ = surface coverage, η_z = inhibition efficiency.

at 1.00 \times 10⁻³ M. The corresponding protection abilities at the same concentration are 93.73% for **p-TSAH** and 90.47% for **p**-TSAE. Overall, the EIS electrochemical outcomes support both the PDP and EFM results, which in turn promote the ability of both p-TSAH and p-TSAE molecules as candidates for effective corrosion inhibition.

Despite the limitless informative outcomes of the Nyquist Plots, they have one major shortcoming in that they cannot give any information about the frequency perturbation within the system under study. Another common representation of EIS data is the Bode plot, in which the frequency is plotted against both the absolute values of the impedance and the phase-shi (Fig. 8b). This representation provides much information about the interfacial organic layer, p -TSAH or p -TSAE molecules, adsorbed between the metallic surface and the corrosive mineral acid, and contains a single time constant that corresponds to the electrical double layer at the interface. For the blank solution curve in Fig. 8, the phase angle diverges from the standard value for a pure capacitor, which is −90°. This could a result of frequency scattering due to the surface

inhomogeneity caused by acid attack. On the other hand, the addition of either p -TSAH or p -TSAE molecules shifts the angles to more negative values away from those of the blank solution as an indication of enhanced capacitive behavior and that the surface becomes smoother upon the formation of the adsorbed organic layer. As shown in Fig. 8 and Table 4, the maximum angle deviation is −65.77° for p-TSAH and −50.23° for p-TSAE at 1.00×10^{-3} M. Moreover, the slopes at the middle frequencies shift much closer to the standard value of a pure capacitor of -1 (ref. 54) in the presence of both p -TSAH and p -TSAE molecules (Table 4).

3.3 Adsorption considerations

Adsorption isotherms are widely used to investigate the adsorption and protection mechanisms at the mild steel/1.0 M HCl interface. Several traditional adsorption isotherm equations were utilized to fit the results obtained from the EFM measurement of p -TSAH and p -TSAE molecules.¹ The fitting data for the six adsorption models are shown in Fig. 9 and Table

Fig. 9 The different adsorption models for p-TSAH and p-TSAE compounds on the mild steel surface in 1.0 M HCl at 30 °C.

Table 5 Adsorption isotherm models of the inhibitors with values of R^2 , slopes, intercepts, $K_{\rm ads}$, and $\Delta G_{\rm ads}$ obtained by using data from EFM measurements^a

	Adsorption isotherm model Linear form equation	Inhibitor	Slope	Intercept	\mathbb{R}^2	K_{ads} , M^{-1}	ΔG_{ads} , kJ mol ⁻¹
Freundlich	$\log \theta = \log K + 1/n \log C$	p -TSAE	0.13066	0.38343	0.85418	2.4178	-12.34
		p -TSAH	0.11061	0.32809	0.84755	2.1286	-12.02
Langmuir	$\frac{c}{\theta} = \frac{1}{K} + C$	p -TSAE	1.07480	0.00002	0.99906	56463	-37.68
		p -TSAH	1.05515	0.00001	0.99937	69 636	-38.21
Frumkin	$\log \frac{\theta}{(1-\theta) C} = \log K + 2a\theta$	p -TSAE	-1.99780	6.02670	0.77986	1.0634×10^{6}	-45.08
		p -TSAH	-2.15565	6.33479	0.77446	2.1617×10^{6}	-46.86
Temkin	$\theta = -\frac{1}{2a} \ln C - \frac{1}{2a} \ln K$	p -TSAE	10.10181	-16.72345	0.90502	0.1910	-5.95
		p -TSAH	11.07539	-17.87078	0.89339	0.1992	-6.05
Flory-Huggins	$\log\left(\frac{\theta}{c}\right) = \log K + n \log(1-\theta)$	p -TSAE	2.16061	5.33216	0.95973	2.1486×10^{5}	-41.05
		p -TSAH	2.06039	5.46038	0.96296	2.8865×10^5	-41.79
Kinetic-thermodynamic	$\log\left(\frac{\theta}{1-\theta}\right) = \log K + y \log c$	p -TSAE p -TSAH	0.51945 0.52835	2.62065 2.76446	0.96524 0.96795	417.4933 581.3738	-25.32 -26.15
	a R^2 = regression correlation coefficient, K = binding constant, θ = surface coverage, c = concentration. 5. The quality of the fitting is measured by R^2 values that are					adsorption, which describes electron sharing from the active	
	close to unity. According to the R^2 values, the quality of the fitting is in the order Langmuir > kinetic-thermodynamic >					sites of the organic materials (lone pairs of hetero atoms, π electrons and aromatic moieties) to the empty orbitals of the	
	Flory-Huggins > Temkin > Freundlich > Frumkin. Therefore,					metal. The third category is for values between -20 kJ mol ⁻¹	
	the Langmuir model was found to be the most appropriate for					and -40 kJ mol ⁻¹ , considered as mixed type. ⁵⁸ Our case	
	the chosen EFM results. This isotherm considers that the					matches with the third category as mixed type, but it is greatly	
	metallic adsorbent surface is uniform (the active adsorption					shifted towards -40 kJ mol ⁻¹ , so the priority is to chemical	
	sites are equivalent), there is no interaction between the		adsorption.				
	adsorbate (inhibitor) molecules, and the adsorption process						
	occurs by the exchange of single adsorbed water molecules by						
	single inhibitors. ⁵⁵ This isotherm is expressed as: ⁵⁶		3.4 Surface analysis				
	$\frac{c}{\theta} = \frac{1}{K_{ads}} + C$	(5)				3.4.1 SEM analysis. SEM analysis was performed to confirm	

$$
\frac{c}{\theta} = \frac{1}{K_{\text{ads}}} + C \tag{5}
$$

The adsorptive binding constant (K_{ads}) is the inverse of the intercept of a c/θ vs. C plot. The K_{ads} values are found to be 69 636 M⁻¹ for the **p-TSAH** molecule and 56 463 M⁻¹ for the **p-TSAE** molecule. These large values of K_{ads} are a good indicator of the strength of the adsorption process, with p-TSAH having priority over p-TSAE molecules.

The adsorption free energy is estimated from the relationship:⁵⁷

$$
\Delta G_{\rm ads} = -RT \ln 55.5 K_{\rm ads} \tag{6}
$$

$$
(R = 8.314
$$
 J mol⁻¹ K⁻¹, $T = 303.15$ K, [H₂O] = 55.5 mol L⁻¹)

 $\Delta G_{\rm ads}$ values were found to be −38.21 kJ mol⁻¹ for the p-**TSAH** molecule and -37.68 kJ mol⁻¹ for the **p-TSAE** molecule. The adsorption occurred in a spontaneous way as indicated from the negative values. ΔG_{ads} values can be grouped into three categories. The first is for values that are more than [−]20 kJ mol−¹ , considered as physical adsorption, which describes the electrostatic attraction between the metallic surface and the organic materials. The second category is for values less than -40 kJ mol⁻¹, considered as chemical

3.4 Surface analysis

3.4.1 SEM analysis. SEM analysis was performed to confirm the adsorption of p -TSAE and p -TSAH on the surface of the mild steel specimens. SEM pictures of the morphology of the mild steel samples after 24 hours in the acid solution with and without 1 mM p -TSAE and p -TSAH are shown in Fig. 10. In the blank sample (mild steel/1.0 M HCl solution), the surface was very rough, badly corroded and destroyed. The addition of p-TSAE and **p-TSAH** significantly reduced the extent of the corrosion and the surface became smoother and less damaged. This suggests the formation of an organic protective layer that can effectively resist corrosion.⁵⁹

3.4.2 XPS analysis. The XPS technique was used to investigate the formation of an adsorbed organic layer on the steel surface. The analysis was done on mild steel in 1.0 M HCl containing 1.0 \times 10⁻³ M of **p-TSAH** after immersion for 24 h and the data is shown in Fig. 11. The surface survey proved the presence of C, N, S and O with atomic percentages of 52.16%, 2.63%, 0.9% and 33.44%, respectively. The deconvoluted spectrum of C1s shows three bands corresponding to $C-C/C=C/C$ – H aromatic (285.18 eV), C–O/C=O (286.97 eV) and C–N, C–N⁺ (288.72 eV). The N1s spectrum shows one peak at 400.03 eV, corresponding to N–C/N–H. The deconvoluted spectrum of O1s shows three bands at 530.28, 531.79 and 532.74 eV, corresponding to several iron oxides such as $Fe₂O₃/Fe₃O₄/FeOOH$ and organic species such as $C-O/C=O/C-OH$. The S2p spectrum shows two peaks at 168.26 and 169.87 eV, corresponding to $S-Fe/S=O$.

Fig. 10 SEM images of the steel surface after 24 hours in the absence and presence of 1.0×10^{-3} M of p-TSAH and p-TSAE compounds at 30 °C.

3.5 Molecular reactivity

3.5.1 Global reactivity. Molecular reactivity is an old expression that was used for many decades in organic synthesis before it was used in corrosion studies. An early use of this expression was to describe the reactivity of an organic molecule as a function of its ability to share electrons from its HOMO to the LUMO of another organic molecule.⁶⁰ The same concept is then applied to the ability of organic inhibitors to donate their

electrons to metal d-orbitals, so that corrosion reactions are suppressed.⁶¹ The reactivity requires a low ionization potential and narrow gap between the HOMO and LUMO so that electrons can be easily reallocated into the metal surface.⁶² p-TSAH and p -TSAE are different only in one functional group, an ester group (with one terminal oxygen) versus a hydrazine group (with two terminal nitrogen atoms), which suggests close global reactivities. However, the hydrazine should be slightly more reactive due to the ease of reallocation of the two lone pairs of the two nitrogen atoms. This is shown clearly in the calculated values of the energy gap⁶³ as shown in Table 6. The calculated LUMO–HOMO energy difference (ΔE) for **p-TSAH** and **p-TSAE** are 5.015 and 5.022 eV, respectively. These values were calculated by subtracting the eigenvalue of the highest occupied molecular orbital (HOMO) from the eigenvalue of the lowest unoccupied molecule orbital (LUMO), both obtained directly from DFT results. IPs (ionization potentials) of 6.228 and 6.440 eV for p -TSAH and p -TSAE, respectively (Table 6), indicate a slightly greater tendency of p -TSAH to donate its HOMO electrons to the metal surface and therefore suppress surface corrosion somewhat more effectively. In addition, both p-TSAH and **p-TSAE** show low chemical hardness of 2.511 and 2.507 eV, respectively, which should be reflected in an effortlessness interaction between inhibitor molecules and mild steel. It is worth mentioning that the low hardness is due to the low energy gap - in molecules with filled HOMOs, the electronegativity separates the energy gap into two parts, and their combination gives the chemical hardness.

The tendency of electrons to flow between the inhibitor and the metal surface can be estimated using the following equation:⁶⁴

$$
\Delta N = \frac{(\chi_{\text{metal}} - \chi_{\text{inh.}})}{2(\eta_{\text{metal}} + \eta_{\text{inh.}})}
$$
(7)

where the absolute value of ΔN refers to the direction of the electron flow from an inhibitor to the vacant metal orbitals and thereby is a dimensionless quantity (Table 6). The calculated ΔN values for p -TSAH and p -TSAE are 1.152 and 1.109, respectively, indicating a greater ability of p -TSAH to donate electrons to the metal than p-TSAE.

A qualitative description of both the molecular and atomic reactivity can be explored further by investigating both the frontier molecular orbital and natural bond orbital densities. Fig. 12 shows the FMOs for p -TSAH and p -TSAE, specifically the HOMO and LUMO for each molecule. It shows that the HOMO of the p-TSAE molecule is delocalized mainly over the phenyl ring (attached to the ester group), and the nitrogen and oxygen atoms of the ester group, and less so over the other phenyl ring. It is worth noting the missing contribution of the sulfur atom to the HOMO of p -TSAE, while having three lone pairs with low electronegativity. This could be explained in that it is surrounded by two oxygen atoms and one nitrogen atom with higher electronegativity (N/O) (3.04/3.44) than sulfur (2.58). A fruitful synthesis of organic inhibitors then demands a delocalized distribution of hetero atoms across the molecule so that they don't cancel the donating ability of each other. The same surface density distribution stands for the HOMO of the **p-TSAH**

Table 6 Parameters of global reactivity for p -TSAH and p -TSAE from B3LYP/6-31g(d,p) calculations

^a The units used are eV except for ΔN which is a dimensionless quantity. ^b EHOMO and ELUMO were taken from DFT computation.

Fig. 12 Frontier molecular orbital densities for p-TSAH (left) and p-TSAE (right); LUMO (upper), HOMO (middle), and HOMO−1 (bottom)

molecule in Fig. 12, with a lack of electron density over the nitrogen atoms of the hydrazine group, which, instead, contributes to HOMO−1. Overall, the FMO analysis shows the priority of electron donation to the phenyl ring attached to either the ester or hydrazine group with contributions from the oxygen of the carbonyl group and the nitrogen atom at the linkage of the two aromatic rings being similar.

To evaluate the priority of individual atoms/groups involved in the HOMO of each molecule in sharing their electrons with the metal d-orbitals, the NBOs were analyzed.⁶⁵ Tables 7 and 8 show the orbital hybridization schemes for the proposed sites of interactions ordered by the priority of the electron donating ability along with the corresponding eigenvalues in eV. In addition, atom numbering is provided in Fig. 13. The corresponding surface densities are provided in Tables S1 and S2 in the ESI† in the same order. According to Table 7, hybridizations of **p-TSAH** are in the following order: LP (1) C_{20} > LP (2) O_{31} > BD (2) C₁₇-C₂₁ > BD (2) C₁₈-C₁₉ > BD (2) C₁-C₆ > BD (2) C₂-C₃ > BD (2) C₄-C₅ > LP (1) N₃₂ > LP (2) O₁₃ > LP (3) O₁₃ > LP (3) O₁₄ > LP (2) O_{14} > LP (1) N₁₅ > LP (1) N₃₄ > BD (2) C₂₆-O₃₁, which corresponds

to the orbitals from HOMO to HOMO−13 (Table S1†). This order corresponds to the abilities of the hybridizations to reallocate their electron density into d-orbitals of the metal; i.e. LP (1) C_{20} is at the top of the natural bond orbitals (HOMO), and hence the easiest donation takes place from it, followed by LP (2) O₃₁ (HOMO−1), *etc.* **p-TSAE** shows similar donating behavior according to the sequence: LP (1) (C_{20}) > LP (2) (O_{31}) > BD (2) $C_{17}-C_{21}$ > BD (2) $C_{18}-C_{19}$ > BD (2) $C_{1}-C_{6}$ > BD (2) $C_{2}-C_{3}$ > BD (2) C_4-C_5 > LP (3) (O_{13}) > LP (3) (O_{14}) > LP (2) (O_{13}) > LP (2) (O_{14}) > LP (1) (N_{15}) > LP (2) (O_{32}) > BD (2) C_{26} -O₃₁. These hybridizations correspond to the orbitals from HOMO to HOMO−13 (Table S2†). Therefore, in both molecules most of the donating ability goes to the phenyl ring as well as the carbonyl group attached to it, followed by the other phenyl group and the oxygen atoms of the sulfonyl group. The oxygen and nitrogen atoms of the ester and hydrazine groups, respectively, show the lowest donating ability.

3.5.2 Local reactivity. A molecular electrostatic potential (MEP) map divides a molecule into low potential regions and high potential regions corresponding to electron-rich and

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			Table 7 p -TSAH NBOs at expected inhibitor-metal interactions ordered according to their energies (highest to lowest)				
Bond	Occupancy	Energy	NBO	$s \%$ (atom 1)	p % (atom 1)	$\,$ % $\,$ (atom 2)	p % (atom 2)
$LP(1) C_{20}$	1.057	-0.104	P	$\bf{0}$	99.99		
LP (2) O_{31}	1.865	-0.237	p	0	99.78		
BD (2) $C_{17}-C_{21}$	1.669	-0.246	0.7256 $p^{1.00}$ + 0.6881 $p^{1.00}$	0.02	99.93	$\bf{0}$	99.95
BD (2) $C_{18}-C_{19}$	1.682	-0.251	0.7329 $p^{1.00}$ + 0.6803 $p^{1.00}$	0.01	99.95	$\mathbf{0}$	99.95
$BD(2) C_1-C_6$	1.627	-0.255	0.6920 $p^{1.00}$ + 0.7219 $p^{1.00}$	0.01	99.95	$\mathbf{0}$	99.96
$BD(2)C_2-C_3$	1.658	-0.259	0.7189 $p^{1.00}$ + 0.6951 $p^{1.00}$	$\overline{0}$	99.96	$\mathbf{0}$	99.95
$BD(2) C_4-C_5$	1.689	-0.273	0.7500 $p^{1.00}$ + 0.6614 $p^{1.00}$	0.04	99.95	$\bf{0}$	99.95
LP (1) N_{32}	1.758	-0.278	$sp^{21.54}$	4.44	95.54	$\overline{}$	
LP (2) O_{13}	1.816	-0.287	p	0.07	99.63		
$LP(3) O_{13}$	1.783	-0.287	P	0.08	99.61		
LP (3) O_{14}	1.772	-0.287	p	0.07	99.62		
LP (2) O_{14}	1.82	-0.289	p	0.06	99.64		
LP (1) N ₁₅	1.833	-0.314	${\rm sp}^{8.95}$	10.04	89.91	$\overline{}$	$\overline{}$
						$\overline{}$	$\overline{}$
	1.962 1.981	-0.34 -0.405	${\rm sp}^{3.75}$ 0.5520 sp ^{40.39} + 0.8339 sp ^{30.29}	21.02 2.41	78.91 97.4	3.12	
LP (1) N_{34} BD (2) C_{26} -O ₃₁			Table 8 p -TSAE NBOs at expected inhibitor-metal interactions ordered according to their energies (highest to lowest)	$s \%$	$p \%$	$s \%$	$p \%$
Type	Occupancy	Energy	NBO	(atom 1)	(atom 1)	(atom 2)	96.57 (atom 2)
$LP(1) C_{20}$	1.065	-0.113	$\mathbf{p}^{1.00}$	0	100		
LP (2) O_{31}	1.849	-0.253	$p^{0.99}$	0.01	99.74		
	1.671	-0.255	0.7243 p + 0.6895 p	0	99.94	0	
BD (2) $C_{17}-C_{21}$ BD (2) C_{18} - C_{19}	1.682	-0.258	0.7326 p + 0.6807 p	0	99.95	$\mathbf{0}$	
BD (2) $C_1 - C_6$	1.625	-0.259	$0.6919 p + 0.7220 p$	0.01	99.95	$\mathbf{0}$	
$BD(2)C_2-C_3$	1.657	-0.263	0.7190 p + 0.6950 p	$\boldsymbol{0}$	99.96	0	
	1.688	-0.278	0.7506 p + 0.6608 p	0	99.95	$\mathbf{0}$	
BD (2) $C_4 - C_5$ LP (3) O_{13}	1.782	-0.291		0.08	99.61		
$LP(3) O_{14}$	1.771	-0.291	$p^{0.99}$	0.06	99.63		
	1.815	-0.292	$p^{0.99}$ $p^{0.99}$	0.07	99.63		
$LP(2) O_{13}$ $LP(2) O_{14}$	1.819	-0.293		0.07	99.63	$\overline{}$	
	1.829	-0.318	$p^{0.99}$	9.65	90.31		
LP (1) N ₁₅ LP (2) O_{32}	1.796	-0.318	$sp^{9.36}$ $p^{1.00}$	$\bf{0}$	99.91		99.95 99.95 99.96 99.95 99.95

Table 8 p-TSAE NBOs at expected inhibitor-metal interactions ordered according to their energies (highest to lowest)

electron-poor regions, labeled red and blue, respectively (Fig. 13). The electrophile (in this case the metal) tends to interact with the inhibitor through the red sites where it can find electrons. The map shown in Fig. 13 shows that p -TSAE has two extreme electron-rich regions (red regions) and one extreme electron-poor region (blue region) while p -TSAH has two electron-rich against two electron-poor sites. It is worth noting that the number of electrophile and nucleophile sites doesn't necessarily reflect the overall reactivity of the molecule, however it is an indication of the locations where the metal can interact more effectively. In both p -TSAH and p -TSAE, the electron-rich sites are those surrounding oxygen atoms while the electronpoor sites are those surrounding nitrogen atoms, presumably due to the withdrawing effect of both oxygen atoms and phenyl rings on the lone pairs of the nitrogen atoms. On the other hand, the green colored sites of the π -system of the phenyl rings indicates moderate electron density. While the NBOs suggested that, on the basis of energy, the phenyl rings respond before the

oxygen atoms in inhibitor-metal interactions, the MEP map reveals that oxygen atoms, electron rich sites that come later in the NBO energy scale, provide a greater amount of electron density. Nevertheless, condensed Fukui functions can be used to quantitatively determine the reactivity locations within a molecule on an atom-by-atom basis using the following formulae:⁶⁶

$$
f_{k}^{-} = [q_{k}(N) - q_{k}(N-1)] \tag{8}
$$

$$
f_{k}^{+} = [q_{k}(N+1) - q_{k}(N)] \tag{9}
$$

where $q_k(N)$, $q_k(N+1)$, and $q_k(N-1)$ are the net charges on the atom k in neutral, cationic and anionic species, respectively. Atoms with higher values of $f_{\mathbf{k}}^-$ are subject to electrophilic attack, while atoms with $f_{\rm k}^{\rm t}$ are subject to nucleophilic attack. 60,67 The calculated Fukui indices for the p-TSAH and p-TSAE molecules are listed in Table 9.

Fig. 13 Optimized structures (upper panel) and molecular electrostatic potential (MEP) map (lower panel) of p-TSAH (right panel) and p-TSAE (left panel).

The electrophilic Fukui indices f^- show that when a metal surface approaches *p***-TSAE**, the S₁₀ atom $f^-=0.123$) has the largest electronic density contribution to the d-orbitals of the

Table 9 Condensed Fukui functions^a for local reactivities in p -TSAH and p-TSAE molecules

p -TSAH			p -TSAE		
Atoms	f^*	f^-	Atoms	f^*	f^-
C_1	0.035	0.057	S_{10}	0.019	0.123
O_{31}	0.076	0.056	O_{31}	0.049	0.057
S_{10}	0.022	0.047	C_{26}	0.013	0.055
O_{13}	0.04	0.043	C_1	0.039	0.051
C_{26}	0.019	0.043	C_{20}	0.079	0.045
C_{20}	0.075	0.042	C_{16}	0.047	0.042
C_4	0.012	0.037	C_{21}	0.036	0.034
C_5	0.021	0.037	O_{13}	0.043	0.034
C_6	0.022	0.036	C_{19}	0.038	0.033
C_{16}	0.05	0.036	C_6	0.024	0.029
C_3	0.006	0.035	C_{18}	0.047	0.028
C_{21}	0.034	0.034	C_4	0.015	0.027
O_{14}	0.028	0.033	C ₅	0.023	0.024
C_{19}	0.038	0.033	O_{14}	0.032	0.024
C_2	0.024	0.028	C_{17}	0.052	0.024
C_{18}	0.044	0.027	C_3	0.008	0.022
C_{17}	0.051	0.025	C_2	0.028	0.021
N_{32}	0.021	0.023	O_{32}	0.014	0.014
N_{15}	0.083	0.018	C_7	0.013	0.007
C_7	0.012	0.017	C_{33}	0.008	0.001
N_{34}	0.013	0.008	C_{34}	0.006	-0.003
			N_{15}	0.095	-0.022

 a Calculated using Hirshfeld charges at B3LYP/6-31G(d,p).

metal, however, on the basis of energy, because S_{10} orbitals come late within the NBOs, it should have a low probability of approaching the metal surface. On the other hand, phenyl rings show high electrophilic contributions to the metal surface, in agreement with the order of the NBOs, which confirms their ability to adhere to the metallic surface. The atoms of the p -TSAH molecule have close electrophilic Fukui indices to those of the atoms of p -TSAE molecule, however the nitrogen atoms of the hydrazine group of the p -TSAH molecule have the lowest electrophilic indices (N_{32} and N_{34} are 0.023 and 0.008, respectively) which confirms the electrochemical results in which p -TSAH has better corrosion inhibition than p-TSAE.

3.6 Monte Carlo simulations

MC simulations were used to identify the inhibitor molecule (p-TSAH and p-TSAE) attractions to the Fe surface as well as to offer a clear understanding of the adsorption mechanism. As a result, Fig. 14 shows that the most stable adsorption configuration for the protonated inhibitors is approximately flat. This flat and parallel orientation achieves the highest proper adsorption formations for p -TSAH and p -TSAE molecules on the Fe interface in the acidic solution, implying an enhancement in the adsorption and maximum surface coverage.⁶⁸ As a result, we may conclude that p -TSAH and p -TSAE molecules prefer to interact with the Fe surface even after protonation through several functional groups, heteroatoms, and π bonds. Table 10 contains the adsorption and binding energy parameters. The results demonstrate that the examined p -TSAH and p -TSAE molecules exhibit substantial negative adsorption energy,

Fig. 14 Monte Carlo simulations for the most favorable modes of adsorption obtained for p-TSAE-H⁺ and p-TSAH-H⁺ on the Fe (110) surface, side and top view.

Table 10 The outputs and descriptors calculated by the Monte Carlo simulations for the adsorption of p -TSAH and p -TSAE on Fe (110) (in kcal mol−¹)

Phase	Inhibitor	Total energy (kcal mol^{-1}	Adsorption energy $(kcal mol-1)$	Rigid adsorption energy (kcal mol^{-1}	Deformation energy $(kcal mol-1)$	(dE_{ads}/dNi) (kcal mol^{-1}	Binding energy $(kcal mol-1)$	IE^a (%)
Gas phase	p -TSAE	-111.363	-550.819	-161.848	-388.971	-550.819	550.819	92.51
	p -TSAH	-173.753	-622.299	-177.272	-445.026	-622.299	622.299	94.32
	p -TSAE- H^+	-145.089	-734.915	-175.178	-559.736	-734.915	734.915	92.51
	p-TSAH- H^+	-205.944	-752.517	-190.172	-562.344	-752.517	752.517	94.32
Aqueous	p -TSAE	-5966.133	-6565.058	-6156.983	-408.075	-478.562	478.562	92.51
phase	p-TSAH	-5975.518	-6583.533	-6113.947	-469.586	-517.565	517.565	94.32
	p -TSAE- H^+	-5939.723	-6695.038	-6105.430	-589.608	-668.719	668.719	92.51
	p-TSAH- H^*	-6094.487	-6806.550	-6225.349	-581.200	-866.154	866.154	94.32

 a Inhibition efficiency values obtained from EFM measurements.

which supports the enhanced and spontaneous inhibitory effect of these compounds on the mild steel surface. These findings suggest that the adsorbed H2O and HCl molecules on the Fe

(110) surface can be gradually substituted by inhibitor molecules, resulting in the creation of a protective film between the metallic surface and the aqueous corrosive solution.⁶⁹

Fig. 15 Pictorial representation of the adsorption of p -TSAE and p -TSAH on the steel surface in 1.0 M HCl.

3.7 The inhibition mechanism

The adsorption process on the metal interface is the key factor in determining the inhibition capacity of the organic molecules. This adsorption process includes the formation of a protective barrier layer over the metal surface, which shields it from aggressive components.⁷⁰ Fig. 15 shows the adsorption modes of p-TSAH and p-TSAE in order to inhibit the mild steel corrosion in a hydrochloric acid solution. As indicated from the Langmuir isotherm model and the values of ΔG , our investigated inhibitors are mixed type between the physical and chemical adsorption modes, and the chemical mode predominates. This is in addition to the parallel adsorption configuration through multiple active sites deduced from the computational calculations. As a result, the electrons of the aromatic moieties, lone pairs of electrons on the hetero atoms (S, N and O) and π electrons (C=O and S=O), can exchange their electrons with the vacant "d" orbitals of Fe as chemical adsorption modes to limit further corrosion of the metal exposed to the corrosive solution.⁷¹ In the hydrochloric acid solution, the p -TSAH and p -TSAE molecules can be easily protonated. Thus, the electrostatic interaction between the positively charged protonated inhibitor species and the negatively charged Cl− ions (originally bonded to the steel surface) generates the physisorption mode. Furthermore, the extra electrons on the steel surface can be transferred back to the empty π^* regions of the inhibitor molecules (retro-donation).⁷²

4. Conclusions

(1) The facile synthesis of two corrosion inhibitors has been achieved in a large, scalable approach based on the Hinsberg reaction, forming commercially available chemicals that can be applied in industrial applications.

(2) The investigated synthesized compounds p -TSAH and p -TSAE were highly efficient inhibitors of corrosion, while p-TSAH had a better inhibition efficiency than p -TSAE in an acid medium.

(3) The adsorption of p -TSAH and p -TSAE on steel was tested against several isotherms and found to be in line with the Langmuir model.

(4) The adsorption of the compounds is of mixed type and controlled by the charge transfer mechanism.

(5) The design of organic inhibitors requires a delocalized distribution of hetero atoms across the molecule to achieve the optimal donating ability of these atoms.

 (6) The active adsorption regions were confirmed using NBO and MC analysis.

Conflicts of interest

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

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