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Density functional theory study of the corrosion inhibition performance of 6-mercaptapurine and 6-thioguanine expired drugs toward the aluminium (111) surface†

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The potentiality of the 6-mercaptapurine (MP) and 6-thioguanine (TG) expired drugs toward the corrosion inhibition of the aluminium (Al) (111) surface was widely investigated using a series of density functional theory (DFT) calculations. A competition between the anti-corrosive features of the studied drugs in the gas and aqueous phases was conducted on both neutral and protonated forms by means of quantum mechanical descriptors. The results of the electrostatic potential analysis demonstrated the prominent nucleophilic nature of the sulfur and nitrogen atoms over the structures of the examined drugs. The frontier molecular orbital theory findings outlined the higher preferability of TG over MP as a corrosion inhibitor. Upon determining the most beneficial configurations of the MP/TG...Al (111) complexes, first-principles molecular dynamics simulations were executed. Interestingly, the competence of the TG drug in the corrosion inhibition process of Al (111) was more extensive than that of the MP one, which was confirmed by the interaction energy values of -1.79 and -1.64 eV, respectively. Upon obtaining the relaxed complexes, the effect of the presence of water solvent on the adsorption process was studied. These findings provide a foundation for developing green anti-corrosive inhibitors for the aluminium surface.

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1 Introduction

Nowadays, aluminium (Al) and its alloys are used worldwide in versatile industrial applications, such as automotive, aerospace, electronics, and transportation, owing to their mechanical strength and economical manufacturing cost.^{1,2} It is noteworthy that Al and its alloys possess crucial resistance to atmospheric conditions due to a firm adherent metal oxide film.³ Nevertheless, Al-based alloys are susceptible to localized attack in corrosive environments, like media containing chloride ions.⁴⁻⁶

Hydrochloric acid (HCl) is a common solution used for chemical, pickling, and electrochemical etching of Al metal and its alloys.^{7,8} Further, the HCl solution is used to remove dust and rust in the metallurgical industries. Additionally, the surface of Al in an aqueous solution forms a compact and robust passive oxide coating. This film dissolves when submerged in HCl solution due to its amphoteric nature, leading to corrosion issues.⁸⁻¹¹ Failures and losses resulting from the corrosion phenomena lead not only to economic problems but also environmental, security, social, and safety issues. Up to now, the investigation of corrosion inhibition of metallic surfaces has triggered much industrial and commercial interest. Indeed, corrosion could not be eliminated, but it could be only minimized by utilizing several methods comprising anodizing and coatings.^{12,13}

An innovative anti-corrosive approach is the incorporation of organic inhibitors,¹⁴⁻¹⁹ forming a protective barrier by getting adsorbed on the surface of the metals. Organic compounds containing heteroatoms (e.g., N, S, and O) in a conjugated system show high efficiency toward the corrosion inhibition of metals.²⁰⁻²² Notwithstanding, the majority of the above-mentioned compounds are highly expensive and toxic to the environment.²³ In this regard, much scientific interest has been

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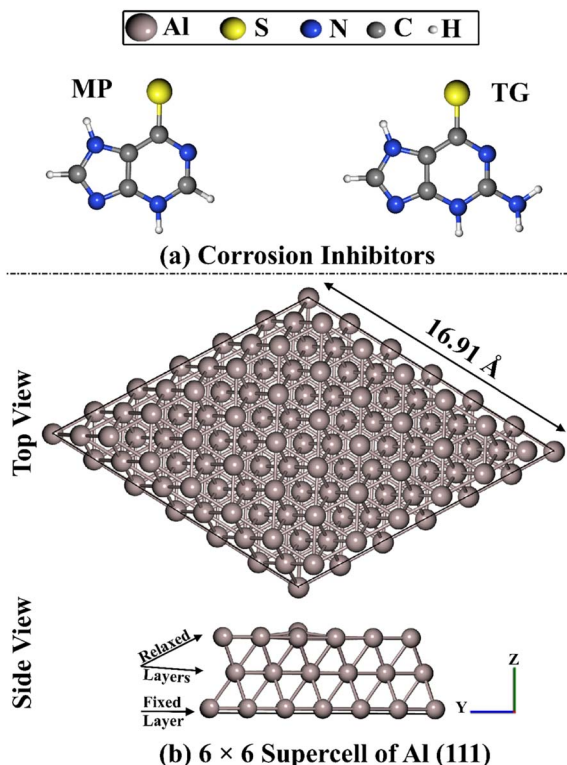


Fig. 1 (a) Illustrative representation for 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs as corrosion inhibitors, and (b) side and top representations of the relaxed 6×6 supercell of aluminum Al (111).

directed toward using pharmaceutical vehicles, more specifically expired drugs, as eco-friendly anti-corrosive inhibitors for metals and their corresponding alloys.^{24–27}

The investigation of drug inhibitory properties has sparked much interest in recent years. A thorough review documented the potential efficiency of drugs as corrosion inhibitors with limited large-scale practical applications due to the high cost of active substances.²⁸ In an attempt to avoid the drawback of the high cost of drugs, corrosion research was accordingly directed to explore the applicability of expired drugs as anti-corrosive inhibitors. It was unveiled that the utilization of expired drugs can solve two more crucial issues, including the reduction of drug disposal costs and limiting environmental pollution with drugs.²⁹

As crucial N-heterocyclic pharmaceutical compounds, purines and their derivatives were deemed to be potential corrosion inhibitors for metals involving steel,^{30–33} aluminium,^{34,35} tin, indium, and their alloys.^{36,37} According to the literature, the 6-mercaptopurine (MP) drug showed promising corrosion inhibition efficiency for copper metal.³⁸ Additionally, the bronze layer was protected by the purine derivative thioguanine (TG).³⁹ The experimental findings revealed that the TG drug exhibited outstanding anodic and cathodic bifunctional corrosion inhibition performance.³⁹ A literature survey unveiled that no solid investigation has been systematically proposed to assess the anti-corrosive efficiency of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs on the Al

(111) surface. In that spirit, the current work is devoted to elucidating the efficacy of the MP and TG drugs as corrosion inhibitors for the Al (111) surface (Fig. 1).

The correlation of the molecular structure with the potentiality of the investigated drugs as corrosion inhibitors was first examined by employing a series of quantum mechanical calculations. Henceforth, molecular dynamics (MD) simulations were conducted for the adsorption process of the MP and TG drugs on the Al (111) surface. Upon obtaining the preferable complexes, the interaction and binding energies were computed. Following that, charge analysis was executed to estimate the role of the charge transfer on the interactions between the investigated inhibitors and the metallic surface. The obtained findings would provide the foundation for utilizing eco-friendly drugs, especially thiopurines, as anti-corrosive inhibitors for the aluminium surface.

2 Computational Methods

2.1. Quantum chemical calculations

DFT computations were carried out for protonated and non-protonated forms of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs in the gas and aqueous phases using the B3LYP/6-311G** level of theory with the help of Gaussian 09 software.⁴⁰ The polarizable continuum model was utilized for the computations in the aqueous phase.⁴¹

Optimization of the geometrical structures of the MP and TG drugs was first performed. Frequency computations were executed to confirm the true minima for their structures. To elucidate the nucleophilic and electrophilic regions over the surface of the chemical systems, electrostatic potential (ESP) analysis was executed on the optimized compounds. Upon earlier recommendations, an electron density envelope with a value of 0.002 au was devoted to extracting the ESP maps.⁴²

To investigate the electronic parameters of the studied thiopurine drugs, FMOs theory was employed. Using FMOs, the electron density distribution of HOMO and LUMO were plotted, and their energies (*i.e.*, E_{HOMO} and E_{LUMO}) were computed. E_{gap} was calculated as the energy difference between E_{LUMO} and E_{HOMO} as follows:

$$E_{\text{gap}} = E_{\text{LUMO}} - E_{\text{HOMO}} \quad (1)$$

Using Koopmans's approximation method, the IP and EA were computed as the negative values of the energies of HOMO and LUMO, respectively, as follows:

$$\text{Koopmans' IP} \approx -E_{\text{HOMO}} \quad (2)$$

$$\text{Koopmans' EA} \approx -E_{\text{LUMO}} \quad (3)$$

With the incorporation of the energy vertical method, the IP and EA were assessed by utilizing the total electronic energy of the neutral molecule (E_{Neutral}) and its corresponding anion ($E_{\text{Anion}}^{\text{Neutral}}$) and cation ($E_{\text{Cation}}^{\text{Neutral}}$), obtained from the geometry of the neutral molecule to keep the external potential constant in the fashion of the following equations:



$$\text{Vertical's IP} = E_{\text{Neutral}}^{\text{Cation}} - E_{\text{Neutral}} \quad (4)$$

$$\text{Vertical's EA} = E_{\text{Neutral}} - E_{\text{Neutral}}^{\text{Anion}} \quad (5)$$

Regarding the energy adiabatic method, the IP and EA were calculated upon obtaining the values of the total electronic energy of the optimized geometry of the neutral molecule (E_{Neutral}), anion (E_{Anion}) and cation (E_{Cation}) based on eqn (6) and (7).

$$\text{Adiabatic's IP} = E_{\text{Cation}} - E_{\text{Neutral}} \quad (6)$$

$$\text{Adiabatic's EA} = E_{\text{Neutral}} - E_{\text{Anion}} \quad (7)$$

Upon calculating the values of IP and EA from the above-mentioned methods, the global descriptors of reactivity, including η , σ , π , X , $\Delta E_{\text{Back-donation}}$, ω , and ΔN , were computed as per eqn (8)–(14).

$$\eta = \frac{\text{IP} - \text{EA}}{2} \quad (8)$$

$$\sigma = \frac{1}{\eta} \quad (9)$$

$$\chi = \frac{\text{IP} + \text{EA}}{2} \quad (10)$$

$$\pi = -\chi \quad (11)$$

$$\Delta E_{\text{Back-donation}} = -\frac{\eta}{4} \quad (12)$$

$$\omega = \frac{\Pi^2}{2\eta} \quad (13)$$

$$\Delta N = \frac{[\phi_{\text{Metal}} - X_{\text{Inhibitor}}]}{2(\eta_{\text{Metal}} + \eta_{\text{Inhibitor}})} \quad (14)$$

The metal work function (ϕ_{Metal}), inhibitor electronegativity ($X_{\text{Inhibitor}}$), and metal hardness (η_{Metal}), and inhibitor hardness (η_{Metal}) were involved in eqn (14). ϕ_{Metal} and η_{Metal} values were 4.26 eV and 0, respectively.^{43–45}

The local reactivity of the investigated drugs was evaluated in terms of the Fukui function indices (*i.e.*, $f_k^+(r)$, $f_k^-(r)$, and $\Delta f(r)$) upon eqn (15)–(17):^{46,47}

$$f_k^+(r) = \rho_k(N+1) - \rho_k(N) \quad (15)$$

$$f_k^-(r) = \rho_k(N) - \rho_k(N-1) \quad (16)$$

$$\Delta f(r) = f_k^+(r) - f_k^-(r) \quad (17)$$

For atom k , the charge values of the neutral ($\rho_k(N)$), anionic ($\rho_k(N+1)$), and cationic ($\rho_k(N-1)$) forms were utilized to evaluate the aforesaid indices.

2.2. Molecular dynamics (MD) simulations

Using the Quantum ESPRESSO 6.4.1 package,^{48,49} molecular dynamics (MD) simulations in the NVT ensemble at 298 K were

performed to investigate the interactions of the **MP** and **TG** drugs with the **Al (111)** surface. According to the experiments and theoretical studies, the **Al (111)** surface has been demonstrated to be the most thermodynamically stable surface.⁵⁰ Therefore, the **Al (111)** surface was designed with three layers of aluminium atoms as a periodic slab, setting a 20 Å layer of vacuum along the z -axis of the **Al (111)** slab. A $6 \times 6 \times 1$ supercell of **Al (111)**, containing 108 atoms, was then constructed (Fig. 1). Afterwards, the **MP** and **TG** drugs were placed on the **Al (111)** surface to assess their ability toward corrosion inhibition (see Fig. 1).

Generalized Gradient Approximation (GGA) within the Perdew–Burke–Ernzerhof (PBE) exchange–correlation functional was utilized.⁵¹ The electron–ion interactions were described *via* the ultrasoft pseudopotential.⁵² The kinetic energy and charge density cutoffs were set to 50 and 500 Ry, respectively. The 10^{-4} eV and 10^{-3} eV Å⁻¹ were set to be the energy and force convergences criteria, respectively. In order to sample the first Brillouin zone, the k -points of the $2 \times 2 \times 1$ grid were utilized for the geometric optimization and charge transfer computations. The Al atoms in the bottom layer were set to be fixed through the relax calculations, while the two top layers of the slab were fully optimized with the inhibitors. For all calculations, the Marzari–Vanderbilt smearing technique was employed.⁵³ Grimme's DFT-D2 correction method was considered to represent the long-range van der Waals interactions.⁵⁴ The MD calculations were performed at a simulation time of 0.5 ps with a time step (dt) of 0.97 fs and steps number (n_{step}) of 1000 steps.

2.3. Interaction and binding energies calculations

Upon the MD simulations, the most stable configurations of the **MP/TG**...**Al (111)** complexes were selected and thoroughly studied. The interaction energies (E_{int}) of the selected complexes were calculated based on the following equation:

$$E_{\text{int}} = E_{\text{MP/TG} \cdots \text{Al (111)}} - (E_{\text{Al (111) in complex}} + E_{\text{MP/TG in complex}}) \quad (18)$$

where $E_{\text{MP/TG} \cdots \text{Al (111)}}$, $E_{\text{Al (111) in complex}}$, and $E_{\text{MP/TG in complex}}$ are the energies of the complex, **Al (111)** surface in complex, and **MP/TG** drugs in complex, respectively. Additionally, the binding energies (E_{bind}) were computed for the investigated complexes as follows:

$$E_{\text{bind}} = E_{\text{MP/TG} \cdots \text{Al (111)}} - (E_{\text{isolated Al (111)}} + E_{\text{isolated MP/TG}}) \quad (19)$$

where $E_{\text{MP/TG} \cdots \text{Al (111)}}$, $E_{\text{isolated Al (111)}}$, and $E_{\text{isolated MP/TG}}$ are the energies of the complex, isolated **Al (111)** surface, and isolated adsorbed drugs, respectively. The Bader charge method^{55,56} was applied to estimate the charge transfer (Q_t) to or from the **Al (111)** surface using eqn (20):

$$Q_t = Q_{\text{combined Al (111)}} - Q_{\text{isolated Al (111)}} \quad (20)$$

where $Q_{\text{combined Al (111)}}$ and $Q_{\text{isolated Al (111)}}$ are the charges of the combined and isolated **Al (111)** surface, respectively. Charge density difference ($\Delta\rho$) maps were depicted using the following equation:



$$\Delta\rho = \rho_{\text{MP/TG}\cdots\text{Al (111)}} - \rho_{\text{Al (111)}} - \rho_{\text{MP/TG}} \quad (21)$$

where $\rho_{\text{MP/TG}\cdots\text{Al (111)}}$, $\rho_{\text{Al (111)}}$, and $\rho_{\text{MP/TG}}$ are the charge densities of the complex, Al (111) surface, and MP/TG drug, respectively. To generate the charge density maps, the VESTA visualization software was used.⁵⁷

For implicit water solvent calculations, the environ code⁵⁸ of Quantum ESPRESSO was utilized with a dielectric constant of 78.3. The solvent effect on the interaction energy of the studied complexes was computed according to the following equation:

$$E_{\text{int}}^{\text{solvent effect}} = E_{\text{int}}^{\text{water}} - E_{\text{int}}^{\text{gas}} \quad (22)$$

where $E_{\text{int}}^{\text{water}}$ and $E_{\text{int}}^{\text{gas}}$ are the interaction energies of the complex in water and gas media, respectively.

3 Results and discussion

3.1. ESP analysis

Electrostatic potential (ESP) analysis is a dependable tool to provide pictorial evidence for either the nucleophilic or electrophilic nature of the chemical systems three-dimensionally.^{59,60} On the optimized structures of the neutral and protonated forms of the MP and TG drugs, molecular electrostatic potential (MEP) maps were portrayed in the gas and aqueous phases using an electron density envelope with a value of 0.002 au (Fig. 2).

Looking at Fig. 2, electron-rich and -deficient centers were identified within the scouted drugs by red- and blue-colored regions, respectively. Evidently, the most negative potentials over the MEP map of the neutral drugs were observed above the sulfur and nitrogen atoms, reflecting their prominent nucleophilic nature. Notably, a negative ESP was found over the hexagonal ring of the drugs under study in the protonated form more than that in the neutral one. The observed nucleophilic character of the studied inhibitors predicted their potential versatility toward interacting with metallic surfaces (*i.e.*, electrophilic surfaces).

3.2. FMOs analysis

The anti-corrosive efficacy of the MP and TG drugs was examined using the frontier molecular orbitals (FMOs) theory. From the FMOs, the distributions of the highest occupied molecular

orbitals (HOMO) and the lowest unoccupied molecular orbitals (LUMO) distributions were plotted for inhibitory compounds under investigation, and are shown in Fig. 3. Energies of the aforesaid molecular orbitals (*i.e.*, E_{HOMO} and E_{LUMO}) along with their gap E_{gap} were evaluated (Table 1).

As can be seen in Fig. 3, the HOMO levels of the two neutral thiopurine drugs in gas and aqueous phases were denoted surrounding the sulfur atom, while HOMO densities of the protonated thiopurine drugs were distributed around the whole molecules. The LUMO distributions were observed over the two rings of thiopurine drugs in the neutral and protonated molecules (Fig. 3).

According to the literature, E_{HOMO} and E_{LUMO} values describe the potentiality of the chemical systems to donate and accept electrons, respectively. From the literature, chemical systems with less negative E_{HOMO} values showed a more prominent tendency toward giving electrons to the electrophiles that have low energy of unoccupied orbitals.^{61,62} It is apparent from Table 1 that the TG drug showed higher values of E_{HOMO} (*i.e.*, less negative) than the MP drug in all of the studied cases, demonstrating its prominent preferability toward donating electrons. Numerically, the E_{HOMO} values of the neutral form of the TG and MP drugs in the gas phase were -5.50 and -5.74 eV, respectively. Comparatively, the neutral form of the investigated thiopurine drugs was denoted with higher E_{HOMO} in the gas phase than in the aqueous phase, and the opposite is true in the protonated form. In both gas and aqueous phases, the studied drugs generally showed higher efficiency in the case of the neutral form than the protonated one. The TG drug showed less negative E_{LUMO} values than the MP drug in the gas and aqueous phases, respectively.

E_{gap} is a descriptive electronic parameter that depends on the reactivity of the inhibitory compound, unveiling its adsorption tendency toward the metallic surface. As E_{gap} decreases (*i.e.*, less positive), the molecular reactivity level is up. In this regard, as listed in Table 1, the TG and MP drugs had lower E_{gap} values in the neutral form compared with the protonated one.

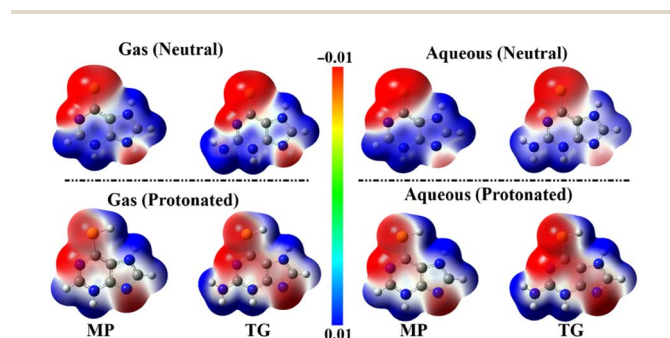


Fig. 2 ESP maps of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs.

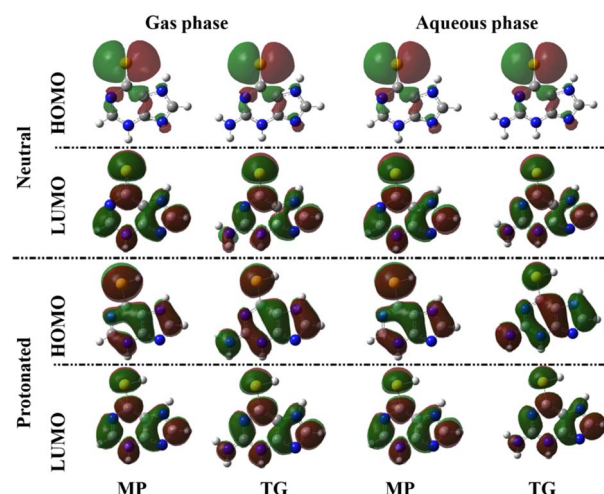


Fig. 3 HOMO and LUMO distributions of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs.



Table 1 E_{HOMO} , E_{LUMO} , and E_{gap} of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs in eV

Compound	E_{HOMO}	E_{LUMO}	E_{gap}
Gas (neutral)			
MP	−5.74	−2.06	3.68
TG	−5.50	−1.69	3.81
Gas (protonated)			
MP	−11.53	−6.99	4.54
TG	−10.96	−6.31	4.64
Aqueous (neutral)			
MP	−6.21	−2.17	4.04
TG	−6.07	−1.82	4.25
Aqueous (protonated)			
MP	−7.72	−3.08	4.64
TG	−7.26	−2.64	4.62

3.3. Global reactivity descriptors

To obtain deeper insight into the reactivity of corrosion inhibitors, a wide scope of global reactivity descriptors, including ionization potential (IP), electron affinity (EA), global hardness (η), chemical potential (π), global softness (σ), electronegativity (χ), total energy change ($\Delta E_{\text{Back-Donation}}$), electrophilicity (ω), and a number of transferred electrons (ΔN), were evaluated by Koopmans, energy vertical and adiabatic methods (Table 2).

By employing Koopmans' theorem, IP and EA can be computed from the E_{HOMO} and E_{LUMO} energies using eqn (2) and (3), respectively. IP is a basic parameter for the reactivity of a molecule. A high IP means high stability and chemical inertness, and the opposite is true for small IP.⁶³ As shown in Table 2, the TG drug showed lower IP and higher EA values than the MP drug. For example, the neutral form of the TG and MP drugs in the gas phase had Koopmans' IP values of 5.50 and 5.74 eV, respectively.

From the view of the hard and soft acid and bases (HSAB) theory, the inhibitory compound with high softness shows superb corrosion efficiency toward the metallic surfaces, and the opposite is true for the hardness property.^{64–66} The inhibitor could be adsorbed onto a metallic surface through the part of a molecule with the lowest hardness and greatest softness.⁶⁷ In both gas and aqueous phases, the neutral form of the studied thiopurine drugs exhibited lower η and higher σ values than the protonated one, outlining their notable reactivity and, accordingly, their favorability in the corrosion inhibition process. Quantitatively, the neutral and protonated forms of the TG drug in the gas phase had Koopmans' η values of 1.90 and 2.32 eV, respectively. Generally, the drugs under study in the gas phase showed lower η and higher σ values compared with the aqueous phase. As numerical evidence, the neutral form of the TG drug in the gas and aqueous phases had Koopmans' η values of 1.90 and 2.13 eV, respectively.

The HSAB theory documented the χ of a molecule as a crucial reactivity property. The χ difference can reveal the electron flow direction between the inhibitors and the surface of

the metal.⁶⁸ It has been reported that the χ value is inversely correlated with the inhibition effectiveness. From Table 2, further suitability of the TG drug over the MP drug toward the corrosion inhibition process was confirmed by the lower positive χ values of the former drug than the posterior one. Obviously, the lower positive χ values were ascribed to the neutral form of the investigated drugs more than the protonated ones, with prominent favorability in the gas phase.

ω allows an assessment of the global electrophilic nature of a chemical system. From the literature, the efficient electrophile and nucleophile are characterized by higher and lower values of ω ,⁶⁹ respectively. The listed ω values in Table 2 show the more eminent nucleophilic nature of the TG drug in all the studied forms and phases compared with the MP drug. Numerically, the neutral forms of the TG and MP drugs in the gas phase exhibited with Koopmans' ω values of 3.39 and 4.13 eV, respectively.

Regarding the ΔN descriptor, a high negative ΔN value indicates the superior preferable inhibition efficiency. For the studied thiopurine drugs, negative ΔN values were obtained, indicating the good inhibition efficiency of the examined drugs.⁷⁰ $\Delta E_{\text{Back-donation}}$ is another characteristic property for the chemical reactivity that was adopted to assess the received charge by each of the interacting species within the corrosion inhibition process. The two thiopurine drugs in all cases had negative values of $\Delta E_{\text{Back-donation}}$, outlining the occurrence of the back donation process, which strengthens the adhesion of the inhibitor to the aluminium surface.

Considering the evaluated global reactivity descriptors using Koopmans, energy vertical and adiabatic methods, the three methods exhibited a nearly similar pattern for the neutral and protonated forms of the studied drugs in the gas and aqueous phases.

3.4. Local reactivity: Fukui functions

Toward identifying the plausible nucleophilic and electrophilic regions, local reactivity of the investigated inhibitory compounds was estimated in terms of Fukui function indices. In this regard, nucleophilic ($f_k^+(r)$) and electrophilic ($f_k^-(r)$) Fukui function indices and their difference ($\Delta f(r)$) were computed, and are summarized in Table 3 for all atoms of the studied thiopurine drugs. Fig. S1† displays the plots of the nucleophilic and electrophilic Fukui function indices iso-surfaces. As shown in Fig. S1,† the sulfur atom was observed with a higher positive purple-coded region than other atoms, reflecting its prominent suitability for the electrophilic attack. It is proven that the most appropriate regions for the nucleophilic and electrophilic attack were, in conjugation, the regions with the highest $f_k^+(r)$ and $f_k^-(r)$ values.^{71,72} High preferability for electrophilic and nucleophilic attacks was perceived for the atoms that possess values more and less than zero, respectively.⁷³ According to the data gathered in Table 3, sulfur and nitrogen heteroatoms were identified as suitable sites for the electrophilic attack by their higher positive $\Delta f(r)$ values compared with other atoms. In general, sulfur atoms of the MP and TG drugs were observed with the highest positive $\Delta f(r)$



Table 2 Global reactivity descriptors of 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs

Compound	Method	IP (eV)	EA (eV)	η (eV)	σ (eV ⁻¹)	π (eV)	χ (eV)	$\Delta E_{\text{Back-donation}}$ (eV)	ω (eV)	ΔN
Gas (neutral)										
MP	Koopmans	5.74	2.06	1.84	0.54	-3.90	3.90	-0.46	4.13	0.10
	Vertical	7.83	0.25	3.79	0.26	-4.04	4.04	-0.95	2.15	0.03
	Adiabatic	7.64	0.39	3.63	0.28	-4.01	4.01	-0.91	2.22	0.03
TG	Koopmans	5.50	1.69	1.90	0.52	-3.59	3.59	-0.48	3.39	0.17
	Vertical	7.57	0.01	3.78	0.26	-3.79	3.79	-0.95	1.90	0.06
	Adiabatic	7.30	0.25	3.53	0.28	-3.78	3.78	-0.88	2.02	0.07
Gas (protonated)										
MP	Koopmans	11.53	6.99	2.27	0.44	-9.26	9.26	-0.57	18.88	-1.10
	Vertical	13.54	5.11	4.22	0.24	-9.32	9.32	-1.05	10.31	-0.60
	Adiabatic	13.42	5.21	4.10	0.24	-9.32	9.32	-1.03	10.57	-0.62
TG	Koopmans	10.96	6.31	2.32	0.43	-8.64	8.64	-0.58	16.06	-0.94
	Vertical	12.82	4.5	4.16	0.24	-8.66	8.66	-1.04	9.01	-0.53
	Adiabatic	12.65	4.67	3.99	0.25	-8.66	8.66	-1.00	9.40	-0.55
Aqueous (neutral)										
MP	Koopmans	6.21	2.17	2.02	0.50	-4.19	4.19	-0.50	4.35	0.02
	Vertical	6.07	2.33	1.87	0.53	-4.20	4.20	-0.47	4.71	0.02
	Adiabatic	5.96	2.45	1.76	0.57	-4.20	4.20	-0.44	5.03	0.02
TG	Koopmans	6.07	1.82	2.13	0.47	-3.94	3.94	-0.53	3.66	0.07
	Vertical	5.93	1.98	1.97	0.51	-3.96	3.96	-0.49	3.96	0.08
	Adiabatic	5.82	2.22	1.80	0.56	-4.02	4.02	-0.45	4.49	0.07
Aqueous (protonated)										
MP	Koopmans	7.72	3.08	2.32	0.43	-5.40	5.40	-0.58	6.28	-0.25
	Vertical	7.67	3.17	2.25	0.44	-5.42	5.42	-0.56	6.52	-0.26
	Adiabatic	7.57	3.28	2.14	0.47	-5.43	5.43	-0.54	6.87	-0.27
TG	Koopmans	7.26	2.64	2.31	0.43	-4.95	4.95	-0.58	5.30	-0.15
	Vertical	7.21	2.75	2.23	0.45	-4.98	4.98	-0.56	5.56	-0.16
	Adiabatic	7.03	3.07	1.98	0.51	-5.05	5.05	-0.50	6.44	-0.20

Table 3 Fukui function indices of S, N, and C atoms of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs

Compound	Atom	Gas Phase						Aqueous Phase					
		Neutral			Protonated			Neutral			Protonated		
		$f_k^+(r)$	$f_k^-(r)$	$\Delta f(r)$	$f_k^+(r)$	$f_k^-(r)$	$\Delta f(r)$	$f_k^+(r)$	$f_k^-(r)$	$\Delta f(r)$	$f_k^+(r)$	$f_k^-(r)$	$\Delta f(r)$
MP	S	-0.368	-0.578	0.209	-0.223	-0.387	0.165	-0.344	-0.724	0.380	-0.196	-0.435	0.239
	N	-0.006	0.017	-0.023	-0.007	-0.018	0.011	-0.010	0.009	-0.019	-0.011	-0.022	0.011
	N	-0.027	-0.026	-0.002	-0.044	-0.033	-0.011	-0.039	-0.017	-0.022	-0.050	-0.029	-0.021
	N	-0.042	-0.044	0.002	-0.042	-0.051	0.009	-0.028	-0.020	-0.008	-0.032	-0.032	0.000
	N	-0.020	0.003	-0.022	-0.035	-0.039	0.004	-0.022	-0.008	-0.014	-0.034	-0.037	0.002
	C	0.003	0.004	-0.002	-0.009	-0.069	0.060	-0.015	-0.007	-0.008	-0.025	-0.083	0.057
	C	-0.065	-0.053	-0.012	-0.073	-0.059	-0.014	-0.083	-0.040	-0.043	-0.095	-0.067	-0.028
	C	-0.037	-0.046	0.009	-0.086	0.026	-0.112	-0.090	-0.044	-0.046	-0.130	-0.006	-0.124
	C	-0.099	-0.059	-0.040	-0.103	-0.068	-0.035	-0.104	-0.038	-0.066	-0.113	-0.066	-0.047
	C	-0.093	-0.054	-0.040	-0.094	-0.051	-0.043	-0.106	-0.038	-0.068	-0.110	-0.048	-0.062
TG	S	-0.355	-0.582	0.227	-0.212	-0.246	0.035	-0.328	-0.742	0.414	-0.190	-0.190	0.000
	N	-0.027	-0.023	-0.003	-0.036	-0.037	0.001	-0.031	-0.013	-0.018	-0.038	-0.037	0.000
	N	-0.011	0.017	-0.028	-0.012	-0.019	0.007	-0.016	0.010	-0.027	-0.017	-0.025	0.009
	N	-0.038	-0.042	0.004	-0.038	-0.050	0.013	-0.025	-0.018	-0.006	-0.028	-0.040	0.012
	N	-0.021	0.003	-0.024	-0.050	-0.035	-0.015	-0.036	-0.008	-0.028	-0.054	-0.036	-0.018
	N	-0.033	-0.036	0.003	-0.041	-0.086	0.045	-0.029	-0.015	-0.014	-0.034	-0.099	0.065
	C	0.007	0.012	-0.005	0.005	-0.093	0.097	-0.008	-0.002	-0.007	-0.012	-0.113	0.102
	C	-0.071	-0.052	-0.019	-0.081	-0.058	-0.022	-0.091	-0.037	-0.054	-0.102	-0.079	-0.023
	C	-0.044	-0.044	0.001	-0.090	0.009	-0.099	-0.097	-0.039	-0.058	-0.136	-0.029	-0.107
	C	-0.061	-0.039	-0.022	-0.058	-0.029	-0.029	-0.073	-0.033	-0.040	-0.078	-0.052	-0.026
	C	-0.098	-0.056	-0.042	-0.103	-0.081	-0.022	-0.108	-0.034	-0.073	-0.114	-0.089	-0.024



values, showing their eminent favorability for electrophilic attack (Table 3). For instance, sulfur atoms of the neutral form of the **MP** and **TG** drugs in the aqueous phase showed the highest $\Delta f(r)$ values of 0.380 and 0.414, respectively.

3.5. Molecular dynamics (MD) simulations

The adsorption behavior of the **MP** and **TG** drugs toward the **Al (111)** surface as effective corrosion inhibitors was unveiled utilizing MD simulations. In this respect, the relationship between the total energy of the **MP**... and **TG**...**Al (111)** complexes and the corresponding simulation time was elucidated, and is plotted in Fig. 4.

Based on the MD simulations, the most stable configurations of the **MP**... and **TG**...**Al (111)** complexes with the highest negative energies were chosen (see Fig. 5). For the most stable configurations, the interaction (E_{int}) and binding (E_{bind}) energies were then computed and are presented in Table 4.

As evident in Fig. 5, S...Al covalent bonds were noticed within the relaxed **MP**... and **TG**...**Al (111)** complexes with lengths of 2.39 and 2.42 Å, respectively. These lengths were smaller than the entire summation of the vdW radii of the S and Al atoms. According to the data in Table 4, negative values of interaction energies (E_{int}) were observed for the **MP**... and **TG**...

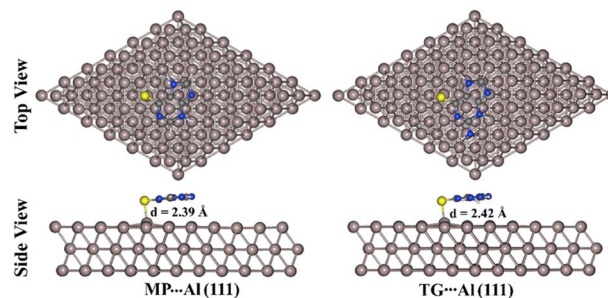


Fig. 5 Top and side representations of the most stable 6-mercaptopurine (**MP**)... and 6-thioguanine (**TG**)... aluminium (**Al (111)**) complexes. The S...Al equilibrium distance (d) is in Å.

Al (111) complexes, indicating that the adsorption processes were carried out. The **TG**...**Al (111)** complex had higher negative interaction energy than the **MP**...**Al (111)** one with values of -1.79 and -1.64 eV, respectively, demonstrating greater efficiency of the corrosion inhibition for the anterior one. The presence of an electron-donating group (*i.e.*, NH_2 group) within the **TG** structure could be the cause of the greater prominence of the molecular inhibitory potency of **TG**.⁷⁴

Considering the structural deformation that occurred during the adsorption process, binding energies (E_{bind}) were estimated (see Table 4). In line with the E_{int} pattern, E_{bind} showed a higher negative value in the case of the **TG**...**Al (111)** complex in comparison with the **MP**...**Al (111)** counterpart. Numerically, the E_{bind} values of the optimized **TG**... and **MP**...**Al (111)** complexes were -1.54 and -1.40 eV, respectively.

Toward a more reliable estimation of the inhibition efficiency, the structures of the most stable configurations of the **MP**... and **TG**...**Al (111)** complexes were reoptimized utilizing five layers of **Al (111)** slab (Fig. S1†). Within the relax calculations, the Al atoms in the last two layers were fixed. At the same time, the three top layers of the slab were fully optimized with the inhibitors. E_{int} and E_{bind} were accordingly computed for the relaxed complexes and are listed in Table S1.† According to Fig. S1,† the relaxed **MP**... and **TG**...**Al (111)** complexes employing five layers of **Al (111)** slab had interatomic distances that were nearly identical to those in the case of using three layers of **Al (111)** slab. For instance, S...Al bond lengths were observed with values of 2.39/2.42 and 2.40/2.42 Å within the relaxed **MP**.../**TG**...**Al (111)** complexes utilizing three and five layers of **Al (111)** slab, respectively.

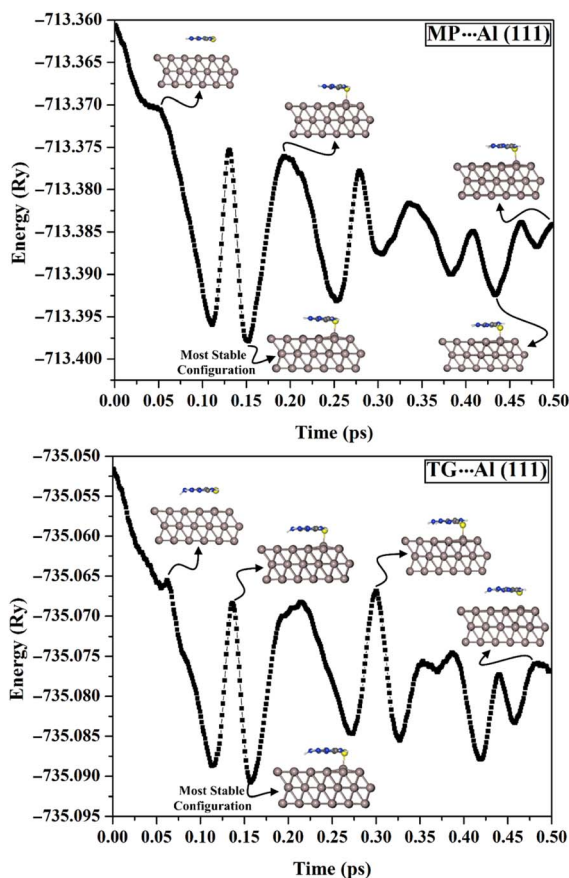


Fig. 4 Correlation between the total energy (in eV) and the molecular dynamics simulation time (in ps) for the 6-mercaptopurine (**MP**)... and 6-thioguanine (**TG**)...aluminium (**Al (111)**) complexes.

Table 4 Interaction (E_{int} , in eV) and binding (E_{bind} , in eV) energies of the most stable 6-mercaptopurine (**MP**)... and 6-thioguanine (**TG**)... aluminium (**Al (111)**) complexes according to the findings of the MD simulations. Interatomic distances (d) are given in Å. Charge transfer (Q_t) values for **Al (111)** surface following the adsorption of the investigated drugs are calculated in e

System ^a	Bond	d (Å)	E_{int} (eV)	E_{bind} (eV)	Q_t^b (e)
MP ... Al (111)	S...Al	2.39	-1.64	-1.40	-0.67
TG ... Al (111)	S...Al	2.42	-1.79	-1.54	-0.52

^a The most stable configurations are depicted in Fig. 5. ^b Q_t was calculated depending on eqn (20).



From the negative E_{int} and E_{bind} values summarized in Table S1,[†] the occurrence of the adsorption process was confirmed. Compared to the adsorption process on the surface of the three layers of the Al (111) slab, five layers of the Al (111) slab showed slightly lower preferability by the energy difference ranging from -0.04 to -0.12 eV. Numerically, the E_{int} values of the adsorbed MP/TG molecule on three and five layers of the Al (111) slab were -1.64 – -1.79 and -1.52 – -1.75 eV, respectively.

3.6. Electronic properties calculations

The Bader charge analysis is an appropriate method to better understand the transfer of the charge between the adsorbent and the substrate.^{55,75} The charge transfer differences (Q_t) of the most stable configurations of the MP and TG drugs on the surface of Al (111) were computed using Bader charge analysis, and the outcomes are summarized in Table 4. Based on the Bader charge findings, the negative values of the Q_t indicate that the charge shifted from the inhibitors to the metallic surface, and the opposite is true for the positive values.

It is clear from Table 4 that the MP and TG drugs transferred electrons to the Al (111) surface *via* the negative Q_t values, indicating that the studied drugs had the potential to interact with the Al (111) surface. Such data consequently ensured the potential anti-corrosive activity of the investigated drugs toward the Al (111) surface. To gain a better understanding of the adsorption process, the maps of the charge density difference ($\Delta\rho$) for the MP... and TG...Al (111) complexes at the most stable configurations were constructed based on eqn (21). The extracted $\Delta\rho$ maps are illustrated in Fig. 6. In the $\Delta\rho$ maps, the sites with increasing and decreasing electron density were detected by colors of yellow and cyan, respectively.

From the data in Table 4, the charge transfer to the Al (111) surface from the investigated drugs was indicated by a negative sign of the Q_t values. The amount of the distributed charge on the complexes was in line with the E_{int} findings, at which the TG...Al (111) complex had an E_{int} value that was larger than that of the MP...Al (111) complex.

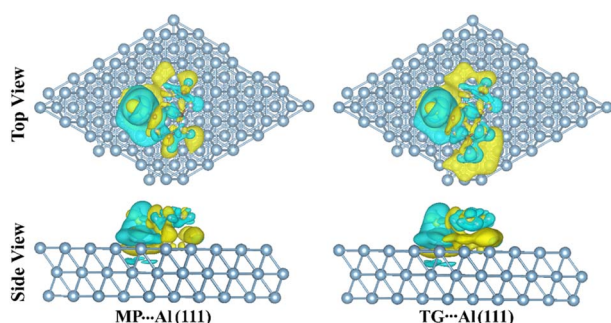


Fig. 6 Top and side views of the charge density difference ($\Delta\rho$) maps of the relaxed structures of the 6-mercaptopurine (MP)... and 6-thioguanine (TG)...aluminium (Al (111)) complexes at the most stable adsorption configurations obtained from the MD simulations. The charge accumulation and depletion were represented by yellow and cyan colors, respectively. The isosurfaces values are in $\text{e} \text{ \AA}^{-3}$.

Looking at Fig. 6, cyan- and yellow-colored regions in the $\Delta\rho$ maps were observed within the MP... and TG...Al (111) complexes, indicating the electron gain and loss upon the adsorption process, respectively. Occurrence of charge accumulation (*i.e.*, yellow-colored regions) confirmed the significant affinity of the studied drugs to interact with the surface of Al (111) as potential corrosion inhibitors.

To better understand the impact of the adsorption process on the electronic properties of the examined drugs, E_{HOMO} , E_{LUMO} , and E_{gap} values were computed. The data for the studied drugs before and after the adsorption on Al (111) surface are listed in Table 5.

From the data in Table 5, notable differences in the E_{HOMO} , E_{LUMO} , and E_{gap} values were observed for the studied drugs before and after the adsorption process. Moreover, the E_{gap} values of MP and TG drugs were changed after their interaction with the Al (111) surface, confirming the occurrence of adsorption processes. For example, the isolated TG had an E_{gap} value of 2.284 eV that changed to -0.341 eV after the adsorption process on the Al (111) surface (Table 5). Interestingly, the E_{HOMO} values of the isolated drugs decreased after the adsorption on the Al (111) surface. For example, the isolated TG drug had an E_{HOMO} value of -4.434 eV that decreased to -4.279 eV after the adsorption process on the Al (111) surface. The latter observation confirmed the electron-donating property of the TG drug, demonstrating that the FMO results were in line with the Bader charge findings (Table 4).

3.7. Solvent effect calculations

To understand the effect of the solvent on the adsorption process within the MP/TG...Al (111) complexes, the interaction energy was evaluated in the presence of a water solvent. Afterwards, the solvent effect ($E_{\text{int}}^{\text{solvent effect}}$) energy was calculated for the most preferable configurations as the difference between the interaction energies of the water and gas media (see the Computational Methods section for details). The obtained ($E_{\text{int}}^{\text{water}}$) and ($E_{\text{int}}^{\text{gas}}$) values are listed in Table 6.

Based on the data in Table 5, the interaction energies of the MP/TG...Al (111) complexes in the water medium showed the

Table 5 E_{HOMO} , E_{LUMO} , and E_{gap} values for 6-mercaptopurine (MP) and 6-thioguanine (TG) before and after the adsorption process on the Al (111) surface

	E_{HOMO} (eV)	E_{LUMO} (eV)	E_{gap} (eV)
Isolated drug			
MP	-4.681	-2.530	2.151
TG	-4.434	-2.150	2.284
Adsorbed drug			
MP	-4.527	-2.648	1.879
TG	-4.279	-2.231	2.048
Complex ^a			
MP...Al (111)	0.810	0.470	-0.339
TG...Al (111)	0.845	0.504	-0.341

^a Based on the most stable configurations (see Fig. 5).



Table 6 Interaction energies of the most stable 6-mercaptopurine (MP)⋯ and 6-thioguanine (TG)⋯ aluminium (Al (111)) complexes in gas ($E_{\text{int}}^{\text{gas}}$, in eV), and water ($E_{\text{int}}^{\text{water}}$, in eV) media

System ^a	$E_{\text{int}}^{\text{gas}}$	$E_{\text{int}}^{\text{water}}$	$E_{\text{int}}^{\text{solvent effect}}$
MP⋯Al (111)	−1.64	−1.23	0.41
TG⋯Al (111)	−1.79	−1.49	0.30

^a Based on the most stable configurations (see Fig. 5).

same trend as those in the gas medium. For example, the E_{int} of the TG⋯Al (111) complex was larger than that for the MP⋯Al (111) complex in water, with values of −1.49 and −1.23 eV, respectively (Table 6). Subsequently, the negative values of E_{int} confirmed the occurrence of the adsorption process in the water medium.

4 Conclusions

Herein, detailed insight into the amplitude of the 6-mercaptopurine (MP) and 6-thioguanine (TG) drugs toward inhibiting the corrosion of the Al (111) surface was established using a series of DFT calculations. The ESP findings first showed the prominent nucleophilic nature of the S and N atoms of the examined drugs. Moreover, the FMO results revealed higher values of E_{HOMO} (i.e., less negative) for the TG drug than the MP drug in all of the studied cases. These higher values demonstrated the greater prominent preferability of the former drug toward donating electrons than the latter one. Generally, the neutral form of the investigated thiopurine drugs had higher E_{HOMO} in the gas phase than in the aqueous phase, and the opposite is true in the protonated form. Using Koopmans, energy vertical and adiabatic methods, the global reactivity descriptors were evaluated, and the higher inhibition efficiency of the TG drug over the MP analog was generally demonstrated. In summary, the obtained values of the QM descriptors indicated that the TG drug showed higher anti-corrosive activity than the MP one. In line with the global reactivity affirmations, the nucleophilic and electrophilic Fukui function indices documented a more noticeable preferability of donating electrons in the case of TG than MP. Consistent with QM descriptors, the interaction energies (E_{int}) confirmed the higher efficiency of the TG drug toward inhibiting the corrosion of the Al (111) surface than the MP one. E_{int} of TG⋯Al (111) and MP⋯Al (111) complexes showed values up to −1.79 and −1.64 eV, respectively. The energetic results of the solvent effect were in line with the interaction energies in the gas medium that demonstrated a more negative interaction energy for the TG⋯Al (111) complex compared with the MP⋯Al (111) complex. Overall, the above-discussed results are vital for using thiopurine-expired drugs as potential corrosion inhibitors, especially for the Al (111) surface.

Author contributions

Conceptualization, Mahmoud A. A. Ibrahim, Tamer Shoeib, and Lamiaa A. Mohamed; methodology, Mahmoud A. A. Ibrahim,

Nayra A. M. Moussa, Amna H. M. Mahmoud, Tamer Shoeib, and Lamiaa A. Mohamed; software, Mahmoud A. A. Ibrahim; formal analysis, Lamiaa A. Mohamed; investigation, Nayra A. M. Moussa, Amna H. M. Mahmoud, and Lamiaa A. Mohamed; resources, Mahmoud A. A. Ibrahim, Shaban R. M. Sayed, and Tamer Shoeib; data curation, Lamiaa A. Mohamed; writing—original draft preparation, Lamiaa A. Mohamed; writing—review and editing, Mahmoud A. A. Ibrahim, Nayra A. M. Moussa, Amna H. M. Mahmoud, Shaban R. M. Sayed, Peter A. Sidhom, Mohamed K. Abd El-Rahman, Tamer Shoeib; visualization, Lamiaa A. Mohamed; supervision, Mahmoud A. A. Ibrahim; project administration, Mahmoud A. A. Ibrahim. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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