


 Cite this: *RSC Adv.*, 2020, **10**, 2037

Synthesis and amide \leftrightarrow imidic prototropic tautomerization in thiophene-2-carbohydrazide: XRD, DFT/HSA-computation, DNA-docking, TG and isoconversional kinetics *via* FWO and KAS models†

 Nabil Al-Zaqri, ^a Tamer Khatib, ^b Ali Alsalme, ^a Fahad A. Alharthi, ^a Abdelkader Zarrouk^c and Ismail Warad ^{*de}

Thiophene-2-carbohydrazide as a novel small-molecule amide tautomer has been synthesized with an acceptable yield under microwave radiation (MW) conditions. The amide \leftrightarrow imidic thiophene-2-carbohydrazide prototropic tautomerization *via* single proton intramigration was computed using the DFT B3LYP/6-311G(d,p) level of theory. The *endo*-isomer amide structure of thiophene-2-carbohydrazide was proved by XRD and is considered to be the kinetically favored isomer. The DFT-structure parameters were compared to their corresponding XRD-experimental parameters. Several H-bond interactions were detected in the crystal lattice experimentally using the XRD-packing model then correlated to MEP and HSA calculations. The manual and calculated electronic parameters such as, frontier molecular orbital energies, excitation energy, absorption, dipole moment, DOS, GRD quantum parameters and TD-SCF/B3LYP were DFT computed. The thiophene-2-carbohydrazide isomers together with their prototropic (E)/(Z)-thiophene-2-carbohydrazonic acid tautomers were docked against 1BN DNA. FWO and KAS isoconversional kinetic methods were applied, and the thermal behavior and estimated $E_a - \alpha$ relations were determined.

Received 29th November 2019
 Accepted 24th December 2019

DOI: 10.1039/c9ra09831c
rsc.li/rsc-advances

1. Introduction

Methods to prepare hydrazides/hydrazide-derivatives of $-\text{C}(=\text{O})-\text{NH}-\text{N}^{\text{H}}-$ as general framework and active intermediate compounds have been recently reported for the synthesis and development of several types of N-N donor ligands.¹⁻³ Thiophene-2-carbohydrazide as a N-S-O-chelate small molecule ligand and its metal ion complexes have some wide applications, mainly catalytic, biological and industrial.²⁻⁴ The presence of oxygen, nitrogen and sulfur as electron-donor atoms in the backbone of thiophene-2-carbohydrazide exhibit broad types of hydrogen bonds which enhance the structural therapeutic effects and variety of its pharmaceutical activities

like: antitumor, antifungal and anti-HIV.⁵⁻⁸ Moreover, the carbohydrazide small molecule function as an essential pharmacophore in many therapeutically valuable materials, for example, the antiviral, antibacterial and antitubercular biological activities were dramatically increased by inserting the thiophene-carbohydrazide in the structure of big molecules.⁵⁻¹² Therefore, a huge number of heterocyclic carbohydrazide derivatives have been made available.⁶⁻¹⁵

The prototropic tautomers are compounds that can be interconverted from molecule to another *via* single proton migration from group to the close neighbor atom.¹⁶ It's known that there is no pure separating line between such tautomers and isomers: tautomer is obviously isomer that transform with a comparatively depressed activation energy < 20 kcal mol⁻¹.¹⁷⁻²⁴

The most well-known single-proton prototropic tautomerization reactions are illustrated in Scheme 1a-d, such reactions may play a vital role in biological suits.¹⁸ For instance, the hydrogen-bonds which connected bases pairs like G-C and G:G-T together (Scheme 1) to build the DNA is a remarkable example emphasizes the importance of such proposition.¹⁷ In mother nature, tautomerization in nitrogenous bases likely cases genetic mutation *via* H-migration, therefore, in organic molecules prototropic tautomerization toward more stable or (and) active forms play essential role in structural shape selectivity (SSS).¹⁵⁻¹⁹

^aDepartment of Chemistry, College of Science, King Saud University, P. O. Box 2455, Riyadh 11451, Saudi Arabia

^bDepartment of Energy Engineering and Environment, An-Najah National University, Nablus 97300, Palestine

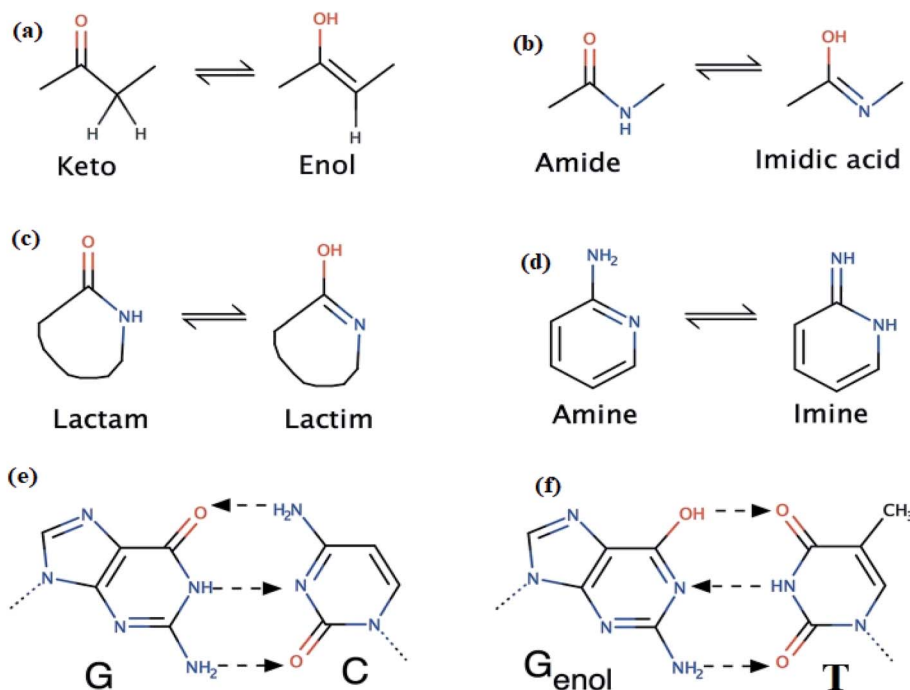
^cLaboratory of Materials, Nanotechnology and Environment, Faculty of Science, Mohammed V University, 4Av. Ibn Battuta, B. P. 1014, Rabat, Morocco

^dDepartment of Chemistry and Earth Sciences, Qatar University, PO Box 2713, Doha, Qatar. E-mail: ismail.warad@qu.edu.qa

^eDepartment of Chemistry, Science College, An-Najah National University, P. O. Box 7, Nablus, Palestine

† CCDC 936463. For crystallographic data in CIF or other electronic format see DOI: 10.1039/c9ra09831c





Scheme 1 (a-d) Prototropic tautomerization reactions and G-C and G_enol-T DNA-bases pairs (e and f).

Thiophene-2-carbohydrazide can be pointed as an excellent model simple-small molecule, for example, Balachandran and coworkers have geometrically optimized thiophene-2-carbohydrazide without having its XRD-crystal structure as trustable structural reference.¹⁰ Moreover, theoretical amide \rightleftharpoons imidic tautomerization of thiophene-2-carbohydrazide to thiophene-2-carbohydrazonic acid *via* single proton intra-migration has been never studied before; such tautomerization process was DFT-computed in this study, the transition states energies and structures were estimated by QST2 method of calculation. Additionally, the XRD-structure of thiophene-2-carbohydrazide confirmed the *endo*-isomer existence of molecule as kinetic favored isomer has been matched to the DFT-calculated parameters. Applying of KAS and FWO isoconversional kinetic methods resulting the multistep reaction decomposition. To establish the intermolecular forces in crystal lattice, MPE and HSA computed result were compared to experimental XRD-packing result. The TD-SCF/DFT/B3LYP calculation, FMO, DOS and GRD quantum parameters were matched with its relative's experimental electronic one. The thiophene-2-carbohydrazide *endo* and *exo* isomers reflected no DNA-docking effect, meanwhile its (*E*)/(*Z*)-thiophene-2-carbohydrazonic acid isomers showed cisplatin like DNA-binding mode.

2. Materials and methods

2.1. XRD

A colorless block single-crystal of thiophene-2-carbohydrazide with the dimension of $0.71 \times 0.12 \times 0.08$ mm was used. The structure was solved using *SHELXL* and *SHELXS* programs, respectively (Table 1).²⁵

2.2. Computations

The DFT-computations were performed at the DFT/B3LYP level of theory in gaseous phase using Gaussian09 software.²⁶ AUTO-DOCK version 4.5 was used for docking, CRYSTAL EXPLORER 3.1 program was used for HSA analysis.²⁷

2.3. Kinetic FWO and KAS methods

In FWO and KAS methods, the correlation between the heating rate β_i and temperature $T_{\alpha i}$ is given by eqn (1) (ref. 28) and (2), respectively.²⁹

Table 1 Crystallographic refined parameters of desired molecule structure

Empirical formula	C ₅ H ₆ N ₂ OS
CCDC	936463
Temperature	293(2) K
Formula weight	284.36
Wavelength	0.71073 Å
Crystal system, space group	Monoclinic, P2 ₁ /c
Volume	639.55(5) Å ³
Unit cell dimensions	$a = 6.06900(4)$ Å $b = 8.50100(2)$ Å $c = 12.51800(3)$ Å $\beta = 98.048(4)$ °
Crystal size	0.71 × 0.12 × 0.08 mm ³
Z	4
Absorption coefficient	0.42 mm ⁻¹
F_{000}	296
Reflections collected/unique	1131/[$R(\text{int}) = 0.021$]
Refinement method	Full-matrix least-squares on F ²
Final R indices [$I > 2\sigma(I)$]	$R_1 = 0.038$, $wR_2 = 0.094$,
Largest diff. peak and hole	0.21, -0.22 e Å ⁻³



$$\ln(\beta_i) = -1.052(E_\alpha/RT_{\alpha i}) + \text{constant} \quad (1)$$

$$\ln(\beta_i/T_{\alpha i}^2) = -E_\alpha/RT_{\alpha i} + \text{constant} \quad (2)$$

2.4. Synthesis of thiophene-carbohydrazide

Methyl thiophene-2-carboxylate (1.0 mmol) and hydrazine monohydrated (5.0 mmol) was dissolved in 40 ml of methanol in 250 ml round bottom flask containing 5 small pieces of boiling cheeps, to a void solvent evaporation during radiation an effective reflux system with two considers was setup, the reaction mixture in the bottom flask only was subjected to MW radiation for 5 min, the white precipitate was collected then filtered off after direct cooling of the reaction in ice bath, yield: 86.7%. The mp = 136–138 °C, EI-MS *m/z*: found 143.2 [M⁺ + 1], cal.: 142.3 [M⁺]. CHN-EA for C₅H₆N₂OS, obs. C, 42.24; H, 4.25; N, 19.70%, cal.: C 42.12; H 4.11; N 19.58%.

3. Result and discussion

3.1. Synthesis

A one-pot eco-synthesis method has been developed for the preparation of thiophene-2-carbohydrazide simple molecule using water-methanol as solvents and cheap kitchen 1500 W normal microwave as source of radiation. In the eco-synthesis one-pot time process was performed by subjecting a combination of hydrazine monohydrated with methyl thiophene-2-carboxylate to 5 min microwave period radiation, as seen in Scheme 2.

The well-known thiophene-2-carbohydrazide compound was prepared in an excellent yield, short time reaction and without side products, moreover; the excess hydrazine monohydrated

was allowed to be evaporated under the heat of MW radiation. The formation of the thiophene-2-carbohydrazide compound was proved by XRD.

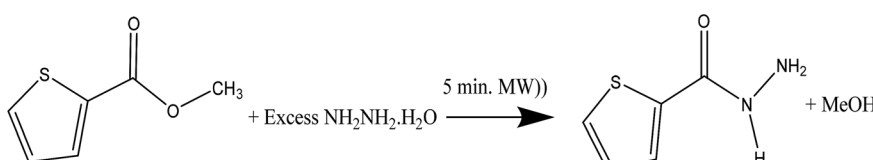
3.2. XRD analysis

Novel conformal thiophene-2-carbohydrazide *endo* isomer (H of amide *trans* to O of carbonyl, *exo*) has been proven by XRD single crystal measurement, as seen in Scheme 3. The rotation around C_{sp²}-N_{sp³} single bond exactly at 180° led to the less stable conformational *exo* isomer (H of amide *cis* to O of carbonyl, *endo*) was not detected by XRD yet.

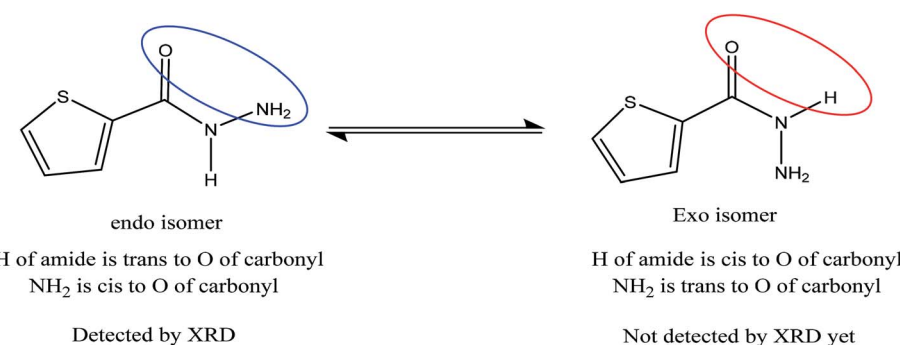
The details crystal structure parameters of the *endo* thiophene-2-carbohydrazide isomer were illustrated in Table 2, moreover, Fig. 1 showed the numbering molecular structure scheme together with its packing unit cell. The thiophene-2-carbohydrazide, C₅H₆N₂OS, crystallized as a monoclinic//P2₁/c space-group, four molecules were crystalline in the unit cell (Z = 4), as seen in Fig. 1b. The structure is look like planer around the carbonyl and the thiophene centers, the carbonyl oxygen atom is in *cis* to the S of the thiophene with dihedral angle = 4.7° and *trans* to the H of the amide with dihedral angle = 170.9°, such seen raised up the polarity of the compound to more than 3.00 D and enhance its metal ions chelate effect.

3.3. Structure optimizations

The *endo* isomer structure of thiophene-2-carbohydrazide which was confirmed by XRD-single crystal analysis was optimized at B3LYP/6-311G(d,p) theory. All the structural parameters like bond lengths and angles were compared to the XRD/exp. collected data. An excellent matching between calculated and theoretical (excepting N-H bonds) was illustrated in Fig. 2, the correlation between the calculated/experimental bond lengths is 0.9783 (Fig. 2a and b). Similarly, the correlation



Scheme 2 Synthesis of thiophene-2-carbohydrazide.

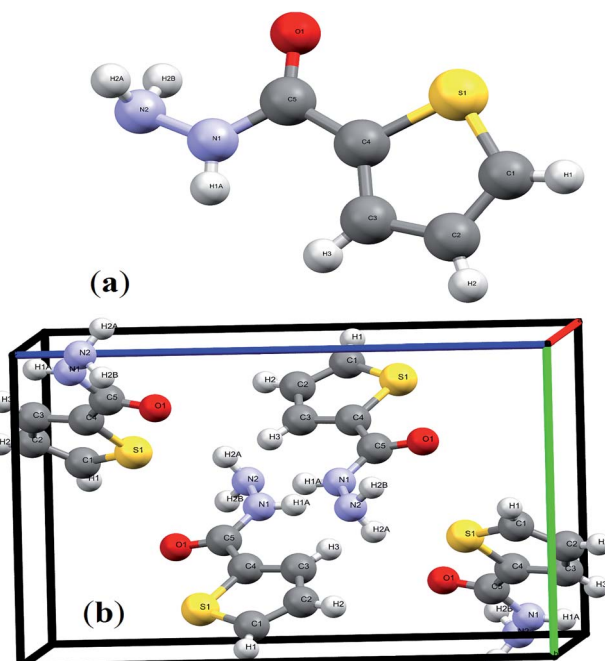


Scheme 3 *Endo/exo*-thiophene-2-carbohydrazide isomerization.



Table 2 XRD-exp. bond lengths (Å) and angles (°) compared to the DFT-calculated

Bond no.	Bonds	Exp. XRD	DFT	Angles no.	Angles (°)	Exp. XRD	DFT
1	S1 C4	1.713	1.7431	1	C4 S1 C1	91.4	91.25
2	S1 C1	1.698	1.7276	2	O1 C5 N1	122.4	121.76
3	C5 O1	1.234	1.2287	3	O1 C5 C4	122.3	122.57
4	C5 N1	1.327	1.378	4	N1 C5 C4	115.3	115.68
5	C5 C4	1.474	1.4789	5	C5 N1 N2	122	119.62
6	N1 N2	1.411	1.4141	6	C5 N1 H1A	122	118.72
7	N1 H1A	0.84	1.0111	7	N2 N1 H1A	113	113.2
8	C4 C3	1.376	1.3773	8	S1 C4 C5	118.1	117.79
10	C3 C2	1.408	1.4222	9	S1 C4 C3	111.4	111.15
11	N2 H2A	0.88	1.0207	10	C5 C4 C3	130.5	131.04
12	N2 H2B	0.89	1.023	11	C4 C3 C2	111.9	112.99
14	C2 C1	1.347	1.3716	12	N1 N2 H2A	107	106.93
				13	N1 N2 H2B	107	106.48
				14	H2A N2 H2B	104	105.03
				15	C3 C2 C1	112.7	112.31
				16	S1 C1 C2	112.6	112.29

Fig. 1 (a) Endo-isomer structure and (b) packing unit cell with $Z = 4$ in thiophene-2-carbohydrazide.

between the calculated/experimental angles is 0.9857 (Fig. 2c and d).

3.4. Crystal interactions, HSA, 2D fingerprint (FP) and MEP investigation

In the crystal lattice, several interactions shorter than 3 \AA were detected experimentally, since four heteroatoms 2N, 1O and 1S in addition to three polar H atoms cited in the backbone of ligand, therefore, four shorter head-to-tail H-bonds as $-\text{HN}-\text{H}\cdots\text{O} = 2.138\text{ \AA}$ and $-\text{N}-\text{H}\cdots\text{NH}_3 = 2.163\text{ \AA}$ strongly dimerized molecules with its neighbors as illustrated in Fig. 3a. Moreover,

two longer H-bonds were detected as $\text{C}=\text{O}\cdots\text{H} = 2.702\text{ \AA}$ and $-(\text{CH}_2)_2-\text{S}\cdots\text{HNH}^- = 2.993\text{ \AA}$ (Fig. 3b).

In HSA, three big red spots were detected close to the O and N atoms on the d_{norm} surface supporting their roles in formation of several types of H-bonds (Fig. 4a). Moreover, each molecule is connected with surrounding molecules *via* three types of H-bonds (Fig. 4b), which built a 3D network around the center molecule.^{2,30-35} The shortest H-bonds are identified as $\text{C}=\text{O}\cdots\text{H}-\text{N}$ type with 2.138 \AA length, followed by two $-\text{N}-\text{H}\cdots\text{N}$ types with 2.336 and 2.546 \AA which is consistent with the XRD result. No true $\text{C}-\text{H}\cdots\text{S}$ thiophene H-bond was detected by the Hirshfeld surface; meanwhile, XRD analysis reflected such bond as H-bond with 2.993 \AA bond lengths.

The MEP/B3LYP reflecting electrophilic/nucleophilic atoms in the molecule as indicated by colors [red (highest) > orange > yellow > green > blue (lowest)]; the electrostatic potential ratios are illustrated not only by colors also by different in lines computing intensity, as seen in Fig. 4c and d. The MEP proved the presence of both e-poor/e-rich positions in the ligand, for example, the line intensity and the red color reflected the O atom of the carbonyl as strong nucleophile site, on the other hand, the blue color of amide H atom showed a strong electrophilic.^{30,31} Since both red and blue colors detected by MEP in the same molecule this proved probability of $\text{H}\cdots\text{O}$ hydrogen bonds formation. The result is consistent with XRD experimental and computed HSA result of having such $\text{NH}\cdots\text{O}=\text{C}$ H-bonds.

The 2D-fingerprint plots over the HSA computed surface molecule reflected the presence of inter-contacts as: $\text{H}\cdots\text{H}$ (28.8%) > $\text{H}\cdots\text{O}$ (9.1%) > $\text{H}\cdots\text{S}$ (6.9%) > $\text{H}\cdots\text{C}$ (5.9%) > $\text{H}\cdots\text{N}$ (4.0%) with $\text{H}\cdots$ overall connections (54.7%), as depicted in Fig. 4e.

3.5. Amide \leftrightarrow imidic tautomerization

In prototropic tautomerization the proton atom migrated from functional group to the neighbor one casing H-bond shift *via* pseudo four, five or six (favor) membered ring transition state.²⁻⁸



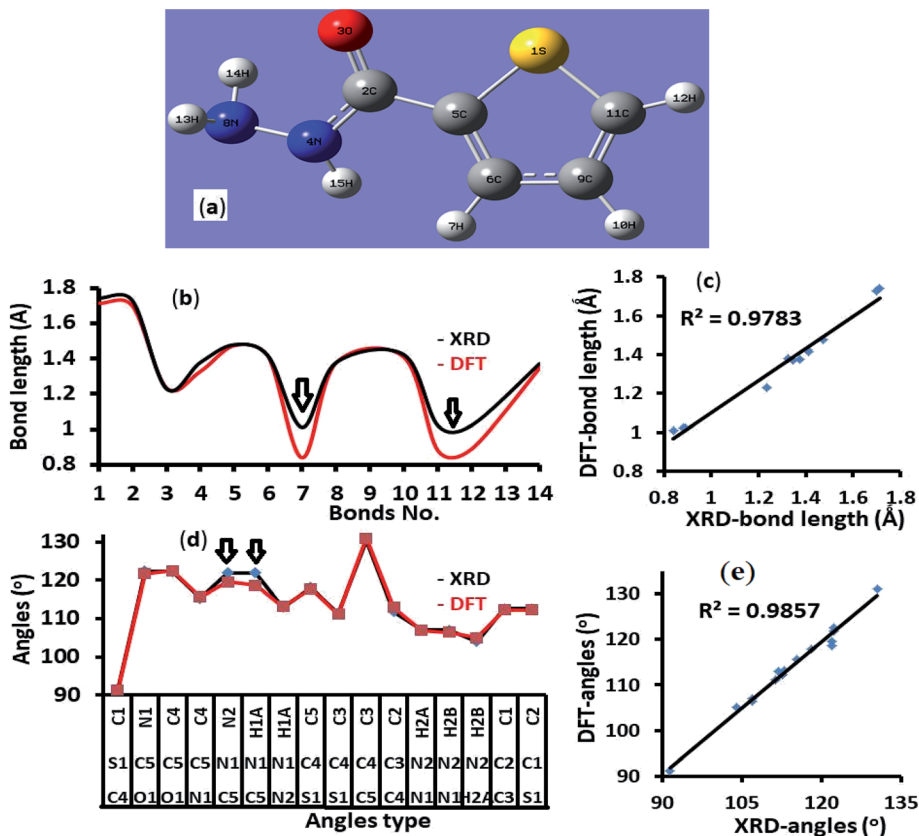


Fig. 2 (a) B3LYP/6-311G(d) structure, (b) XRD/DFT bond lengths histogram, (c) XRD/DFT bond lengths graphical correlation, (d) XRD/DFT angles histogram, and (e) XRD/DFT angles graphical correlation.

This tautomeric behavior can place between enol \rightarrow keto, lactim \rightarrow lactam, imine \rightarrow amine, and amide \rightarrow imidic functionality resulted in single sigma bond migration followed mostly by double bond shift.¹⁵⁻¹⁹

All the XRD and DFT-B3LYP/6-311G(d)-calculations supported the *endo*-thiophene-2-carbohydrazide as favor amide isomer (Fig. 1). Therefore, *endo*-isomer considered to be the most stable tautomeric-isomer (kinetic favored isomer) refer to zero-point-reference-energy with global total DFT energy = -777.0233007 a.u.

The gaseous DFT calculation showed that two steps are needed to tautomerize for **A** \rightarrow **C** (Scheme 4), when crossing amide \rightarrow imidic C=O oxygen to OH parallel to N in NH to C=N bonds functional groups performed prototropic transformed.^{15,16} The first reaction-path involves the conversion of **A** \rightarrow **B**, the H of amine should be in the same direction of carbonyl O atom, for that, **A** performed simple flip-rotation around $C_{sp^2}-N_{sp^3}$ with O-C-N-H dihedral angle = 0.43° to produce the **B** isomer. The transition state (**T.S₁**) of this step is very simple, a perpendicular $C_{sp^2}-N_{sp^3}$ rotation around C-N single bond to make the O-C-N-H dihedral angle around $\sim 90^\circ$ was detected, as seen in Scheme 4.

The second reaction-path involves the conversion of **B** to **C** (*E* or *Z* isomer as final product) *via* single proton-transfer, the proton is intra-migrated from N to O atoms establishing a pseudo-four-membered-ring O···C···N···H as transition state

T.S₂ (Fig. 5). The QST2-computed method of the **T.S₂** demonstrated the H-transfer inside a group containing a C=O moiety in a direct neighborhood of the N-H group, at this level Hammond's postulate was applied since the proton lost its N-attractive correlation and be trapped by O forming O-H new bond, therefore the **T.S₂** structure should be closed to the product shape. This seen is consistent with **T.S₂** computed structure which generated with O1···H1 and N1···H1 distances 1.31 and 1.37 Å, respectively.

The energy profiles in gaseous-phase calculation of amide \rightleftharpoons imidic prototropic tautomerism *via* the single-proton transfer mode is illustrated in Fig. 5. The XRD and DFT-computation revealed the amide as favored isomer over imidic tautomer one libeled with zero-point-energy. The ΔE tautomerization energy was estimated for each step as the energy differences between the tautomer and its transition state. The $E_{T.S_1}$ calculated to 64.14 kJ mol⁻¹, the energy difference between **A** and **B** tautomers were found to be 20.90 kJ mol⁻¹. The $E_{T.S_2}$ calculated to 188.72 kJ mol⁻¹, the energy difference between **B** and **C(Z)** tautomer was found to be 41.45 kJ mol⁻¹, the energy difference between **B** and **C(E)** tautomer was found to be 61.44 kJ mol⁻¹, the energy difference between **C(E)** and **C(Z)** was found to be 19.29 kJ mol⁻¹. In conclusion, the energy calculations in gaseous-phase supported the amide \rightleftharpoons imidic prototropic tautomerism possibility under mild condition.

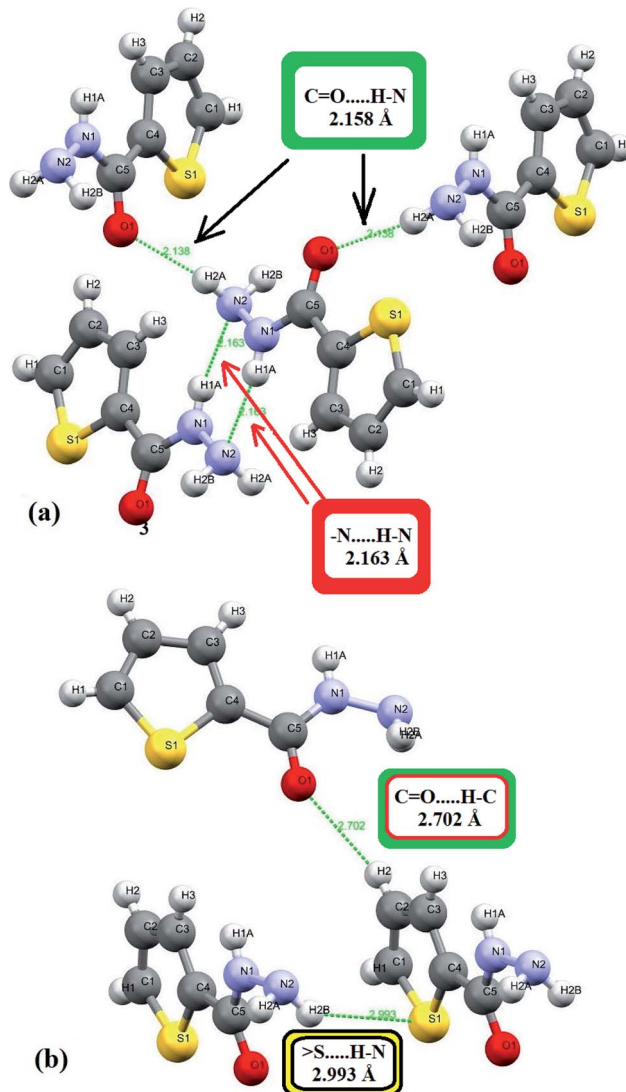


Fig. 3 H-bonds interactions types and lengths: (a) short ($\text{N}-\text{H}\cdots\text{O}$ and $-\text{N}-\text{H}\cdots\text{N}$) and (b) long ($\text{C}=\text{O}\cdots\text{H}$ and $\text{S}\cdots\text{H}-\text{N}$) interactions.

In this process, the amide tautomer of thiophene-2-carbohydrazide was detected by XRD as *endo*-isomer where it classified as kinetic favored isomer, the DFT-calculation supported the probability of amide \rightleftharpoons imidic prototropic tautomerization, since the energy required is not so high it can be easily provided from surrounding temperature or solvents. Therefore, imidic acid isomer form in this can be isolated under suitable conditions, this fact was seen in similar work, the imidic acid tautomer in methanol solvent reaction was isolated by XRD as predominant product.²⁸

3.6. HOMO-LUMO, DOS, exp. electronic transfer, TD-SCF-B3LYP, Mulliken/NPA and GRD

$\text{HOMO} \rightarrow \text{LUMO}$ and $\text{HOMO-1} \rightarrow \text{LUMO+1}$ shapes and energy diagram reflected the electron donation capacity in the ultra-violet region with $\Delta E_{\text{HOMO/LUMO}} = 5.301 \text{ eV}$ (233.88 nm) and

$\Delta E_{\text{HOMO-1/LUMO+1}} = 7.646 \text{ eV}$ (162.15 nm) as illustrated in Fig. 6a.

Experimental-spectral analysis of the thiophene-2-carbohydrazide was carried out in methanol. One broad peak $\lambda_{\text{max}} = 250 \text{ nm}$ was recorded, which was attributed to $\pi \rightarrow \pi^*$ e-transition. No other peaks in the visible region were detected (Fig. 6b). The computed TD-SCF/DFT-B3LYP/6-311G(d,p) in the same solvent exhibited same electronic behavior, a mean broad band at $\lambda_{\text{max}} = 245.8 \text{ nm}$ corresponding mainly to HOMO to LUMO (87%) e-transition (Fig. 6c). The other minor e-transfer together with the molecular contribution ratios were illustrated in Table 3.

In general, the molecular orbital energy levels together with their energy gap (5.301 eV) are strongly agreed with energy gap generated from DOS (5.434 eV) as well as the UV-experimental result as seen in Fig. 6d, the small bathochromic shift ($\sim 5 \text{ nm}$) detected between exp./DFT can be resonated to solute-solvent interactions behavior.^{17,30-35}

NPA and Mulliken charges play a critical role in theoretical quantum-charge distribution; it revealed information about electrophilic and nucleophilic sites on molecule.^{17,30} B3LYP/6-31G(d) Mulliken population and NPA charge analysis of the desired compound were illustrated in Fig. 6e. In general, the Mulliken showed lower atomic-charges compared to NPA, the NPA and Mulliken proved the 2N, O and all carbons atoms except C2 as nucleophilic atoms. The electrophilic sites are S, C2 carbonyl carbon and all the hydrogen atoms, the highest electrophilic hydrogen were sited to the amide proton (H15). The NPA and Mulliken charge result is strongly consistent with XRD-packing, MEP and HSA results.

The global reactivity descriptors (GRD) like: hardness (η), the chemical potential (μ), electrophilicity (ω), softness (σ) and electronegativity (χ) of the molecule were elaborated by using the following equations:

$$I: \text{ionization potential} = -E_{\text{HOMO}} \quad (3)$$

$$A: \text{electron affinity} = -E_{\text{LUMO}} \quad (4)$$

$$\Delta E_{\text{gap}}: \text{the energy gap (eV)} = E_{\text{LUMO}} - E_{\text{HOMO}} \quad (5)$$

$$\chi: \text{absolute electronegativity} = \frac{I + A}{2} \quad (6)$$

$$\eta: \text{global hardness} = \frac{I - A}{2} \quad (7)$$

$$\sigma: \text{global softness} = \frac{1}{\eta} \quad (8)$$

$$\mu: \text{chemical potential} = -\chi \quad (9)$$

$$\omega: \text{electrophilicity} = \mu^2/2\eta \quad (10)$$

The GRD data values were collected in Table 4.

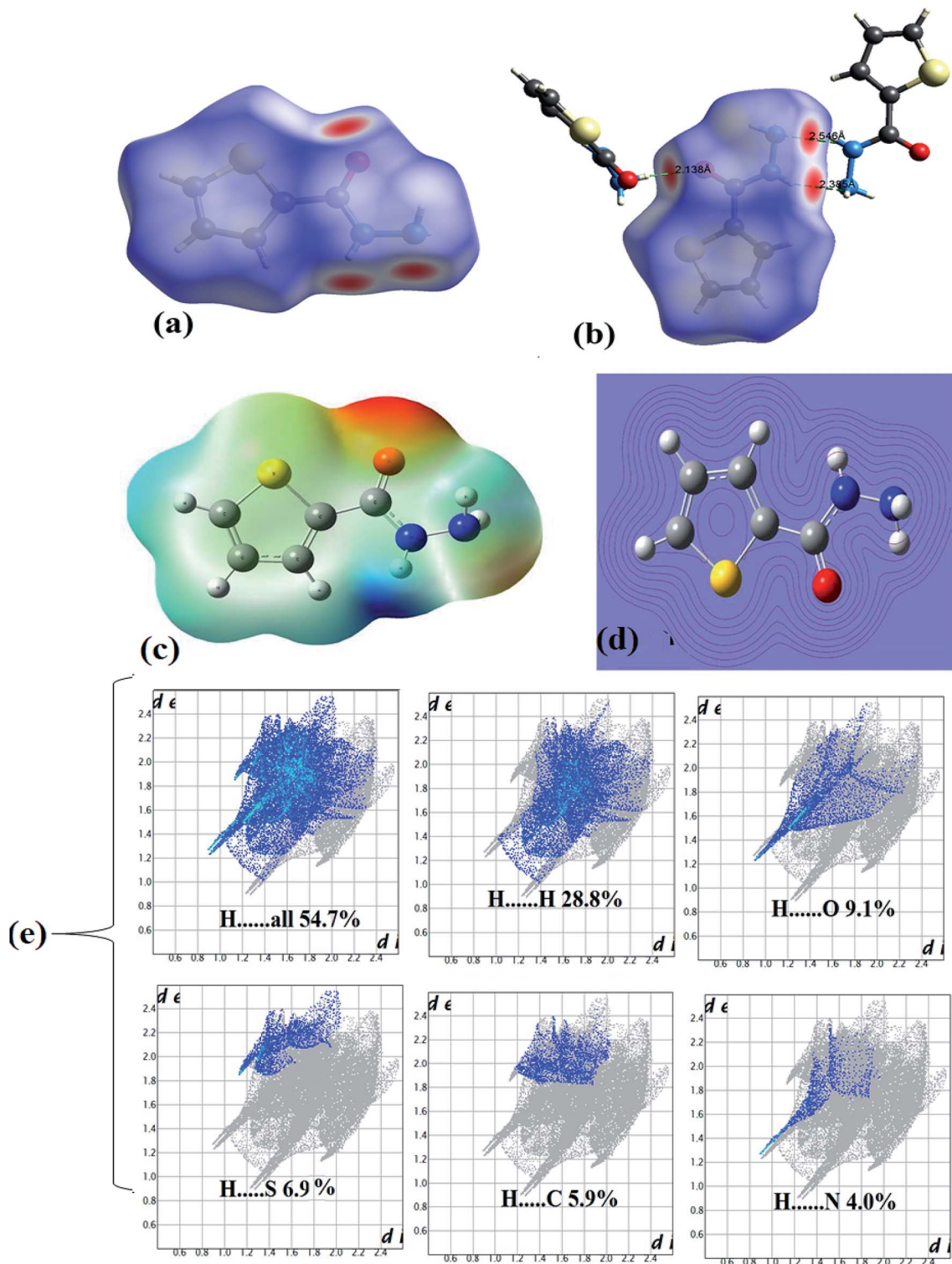
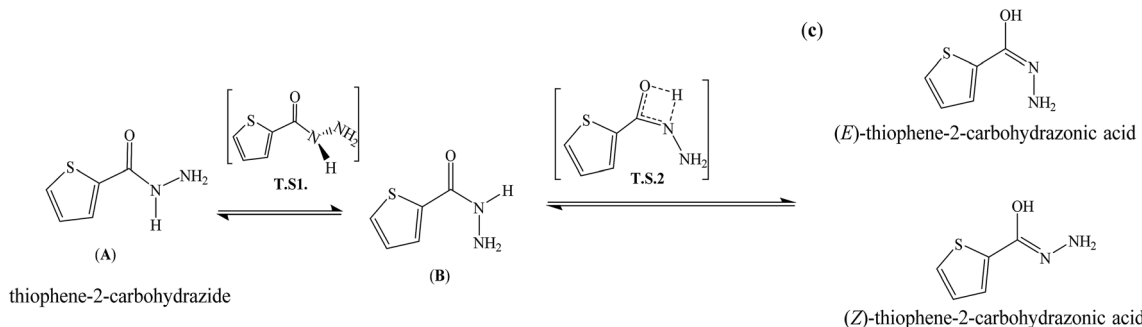
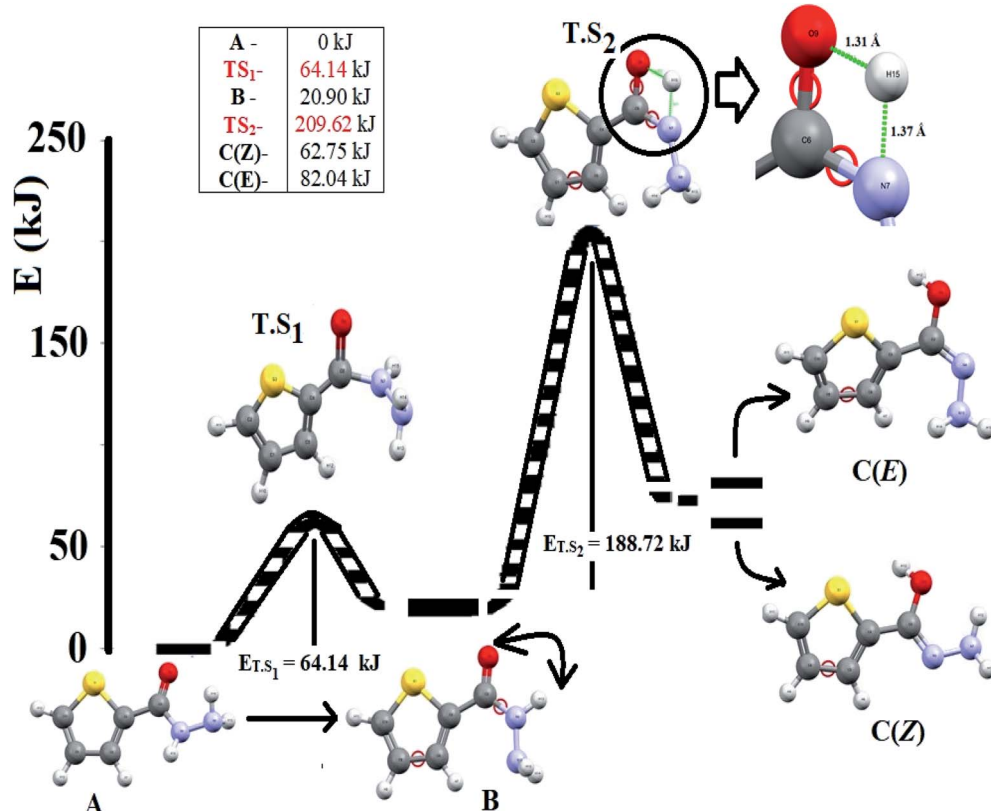


Fig. 4 (a) d_{norm} , (b) H-bonds interactions, (c) MEP, (d) contour and (e) FP inside···outside maps.

3.7. Molecular docking

Model small-molecule, *endo-exo* thiophene-2-carbohydrazide and its (*E*)/(*Z*)-thiophene-2-carbohydrazonic acid tautomers were theoretically molecular docked under the same level of theory using DNA (PDB ID: 1BNA) available data.

Interestingly, *endo/exo* thiophene-2-carbohydrazide isomers (see Scheme 3) reflected no docking effect with 1BNA, meanwhile the (*E*)/(*Z*)-thiophene-2-carbohydrazonic acid resemble an excellent docking by crosslinking both chains of DNA double helix like the cisplatin binding mode.

Scheme 4 Amide \rightleftharpoons imidic prototropic tautomerization reaction.Fig. 5 Structures and energies profiles of amide \rightleftharpoons imidic prototropic tautomerization.

Several hydrogen bonds between DNA:(*E/Z*)drug were detected meanwhile, no non-covalent $\pi-\pi$ stacking interactions were observed (Fig. 7a). The greater binding affinity of *E*-isomer reflected close contact with the surface of molecule through minor groove intercalation mode (Fig. 7a). Three short hydrogen bonds: DNA DG16:H22 \cdots O with 1.98 Å, DNA DG10:H21 \cdots N with 1.86 Å and DNA DA17:O4 \cdots H with 2.084 Å were detected (Fig. 7b and c). Usually docking data are good when the Root Mean Square Deviation (RMSD) value is close or less than 2 Å.³⁶

In the *Z*-thiophene-2-carbohydrazonic acid tautomer, small structural conformation with \sim 20 kJ mol $^{-1}$ isomerization energy difference compared to *E*-isomer,

a dramatically changes in the DNA binding behavior was detected. Due to the structural appropriate of *Z*-isomer it moved deeper between DNA double helix, therefore deep groove intercalation mode with new binding position was observed as in Fig. 7d and e. Moreover, four short hydrogen bonds like: DNA-DG10:H21 \cdots S with 2.154 Å (new bond), DNA-DG16:H22 \cdots N with 2.159 Å, DNA-DG10:H3 \cdots O with 1.747 Å and DNA-DG10:O4 \cdots H with 2.205 Å (Fig. 7f). Since *Z*-isomer energetically and structurally more favored over *E* one, therefore, it can be claimed that (*Z*)-thiophene-2-carbohydrazonic acid is better DNA-binder; this result is consistent with structural shape selectivity method.³⁶



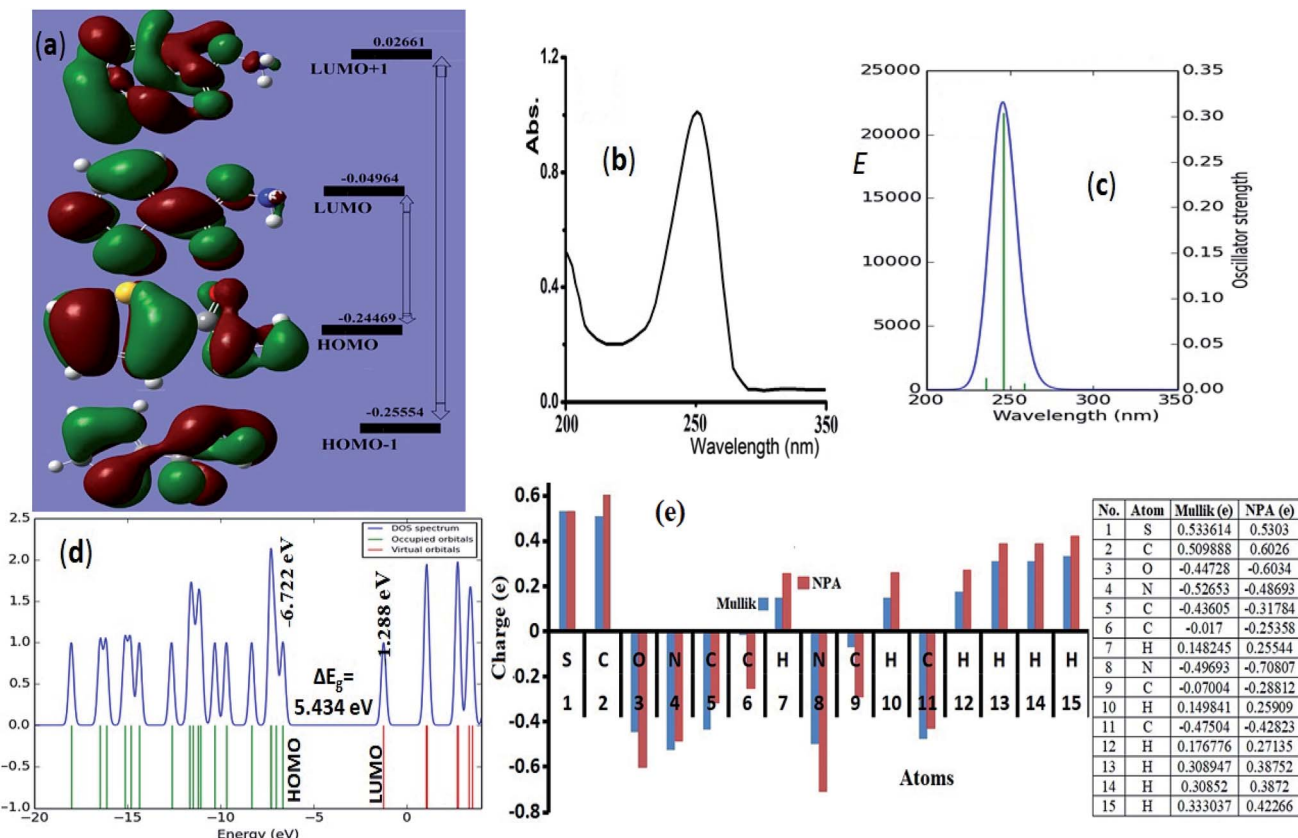


Fig. 6 (a) HOMO/LUMO FMO, exp. (b) and calculated electronic spectra in methanol (c), (d) DOS and (e) NPA (—) and Mulliken (—) atomic charge-distribution.

Table 3 TD-SCF electronic parameters

No.	Wavelength (nm)	Osc. strength	Major contribs	Minor contribs
1	258.65	0.0072	H-3 → LUMO (91%)	H-1 → LUMO (3%), HOMO → LUMO (4%)
2	245.85	0.3041	HOMO → LUMO (87%)	H-3 → LUMO (4%), H-2 → LUMO (5%)
3	235.76	0.0127	H-1 → LUMO (92%)	H-3 → LUMO (2%)

3.8. Thermal and isoconversional kinetics analysis *via* FWO and KSA

The TG/DTG-curves in the temperature range 0 to 400 °C under various heating rates in an open atmosphere revealed the desired compound with good thermal stability, it reflected an acceptable stability up to $T_o = 110$ °C under heat rate = 5 °C min⁻¹, increased to 155 °C by increasing the heat rate to reach up to 20 °C min⁻¹. In all the trials 1–4, the compound decayed from 100% weight to ~zero residue weight *via* one broad step with T_{off} 250–270 °C and full thermal decomposition (Fig. 8a and b).

The TG-DTG-thermograms not served only for thermal analysis of the desired ligand, but also to study its isoconversional kinetic-thermal decomposition. Accordingly, the FWO/KAS isoconversional kinetics models performed TG-analysis at four various heating rates: 20, 15, 10 and 5 °C min⁻¹ to record enough data for E_a 's activations. The T_{DTG} max was shifted to

Table 4 Calculated quantum chemical parameters of free ligand

GRD	Value
Global total energy	E_T 777.0233007 a.u.
Low unoccupied molecular orbital	LUMO -1.3516 eV
High occupied molecular orbital	HOMO -6.6524 eV
Energy gap	ΔE_{gap} 5.3008 eV
Electron affinity	A 1.3516 eV
Ionization potential	I 6.6524 eV
Global hardness	η 2.6504 eV
Global softness	σ 0.3773 eV
Chemical potential	μ -4.0020 eV
Absolute electronegativity	χ 4.0020 eV
Electrophilicity	ω 1.5099 eV
Dipole moment	μ 2.9936 D



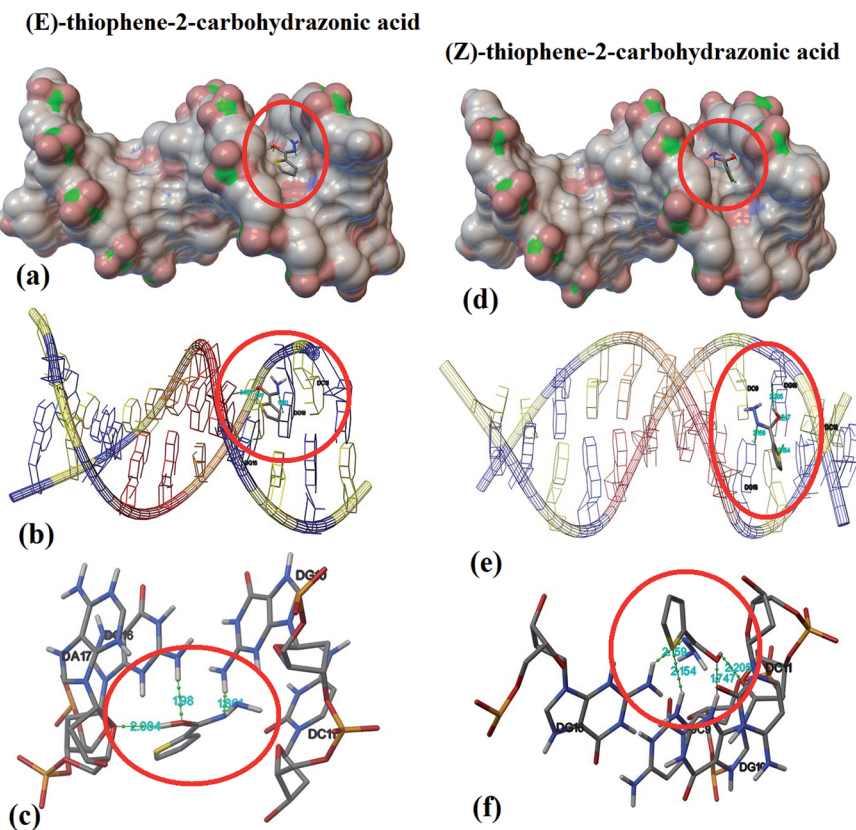


Fig. 7 DNA binding position for *E* (a) and for *Z*-isomer (d), nuclides-binding types for *E* (b) and for *Z*-isomer (e), and DNA:drug H-bonds interactions for *E* (c) and for *Z*-isomer (f).

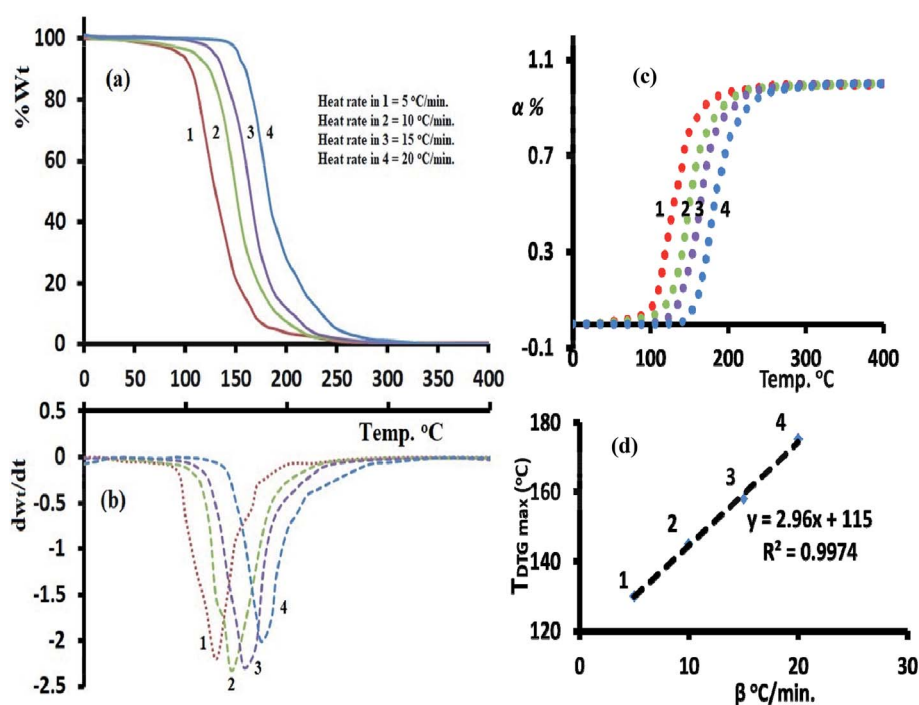


Fig. 8 (a) TG, (b) DTG, (c) conversions and (d) $\beta-T$ curves at four different heating rates.



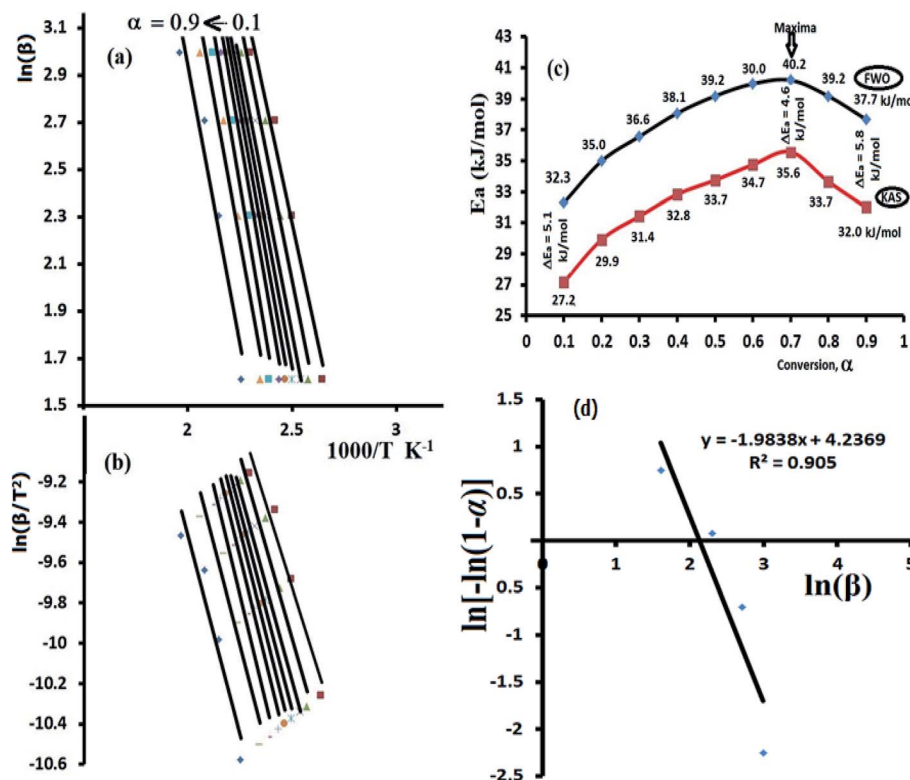


Fig. 9 E_a 's calculations via FWO (a), KAS (b), $\alpha-E_a$'s relation (c) and Avrami's plot (d).

high temperature by rising up β 's as seen Fig. 8b. A perfect linear relation with $R^2 = 0.997$ between β 's and the T_{DTG} max values were recorded as seen Fig. 8d.

Both FWO and KAS as model-free isoconversional methods assumed that the conversion% (α) dependence directly on the temperature values, therefore the activation energy (E_a) of the thermal decomposition reaction can be estimated *via* plotting $\ln(\beta)$ or $\ln(\beta/T^2)$ versus $1000/T$ using FWO Fig. 9a and KAS model Fig. 9b respectively.

The activation energies were changed by rising α from 0.1 to 0.9 (Fig. 9c), at $\alpha = 0.1$ FWO model revealed E_a with 32.3 kJ mol^{-1} , while KAS gave it 27.2 kJ mol^{-1} . Both models reached up to the maxima at $\alpha = 0.7$ with 40.2 kJ mol^{-1} (FWO) and 35.6 kJ mol^{-1} (KSA) then decreased back with the extent of conversion to 37.7 kJ mol^{-1} at $\alpha = 0.9$ (FWO) and 32.0 kJ mol^{-1} by KSA method. In general, a high degree of matching between the two models was recorded with about $\Delta E_a \sim 5 \text{ kJ mol}^{-1}$ in all the trials. Moreover, the significant variation in E_a values indicated a complex kinetics stimulated decomposition process, the non-linear relationship between E_a and α provided collected *via* both models indicated that the ligand is prospective to be decomposed *via* more than one steps reaction ($n > 1$) and not through one broad step as thermal analysis reflected.³⁷⁻³⁹ Therefore, Avrami kinetic equation³⁹ was applied at fixed suitable temperature $160 \text{ }^\circ\text{C}$ and in $\alpha = 0.1-0.9$ range to support the complexity (multi-step) of thiophene-2-carbohydrazide thermal degradation reaction as in eqn (11).

$$\ln[-\ln(1-\alpha)] = \log \chi(T) - n \ln \beta \quad (11)$$

Plotting of $\ln[-\ln(1-\alpha)]$ *vs.* $\log \beta$ at $160 \text{ }^\circ\text{C}$ reflected a straight line with slope = -1.98 (Fig. 9d), therefore, the kinetic order n of the process is ~ 2 (second order reaction), Avrami theory plot supported the non-simplicity decomposition of thiophene-2-carbohydrazide as well as the FWO/KAS models did.

4. Conclusion

The small-molecule *endo*-thiophene-2-carbohydrazide isomer was confirmed by XRD-crystal structure (CCDC 936463); the DFT/XRD-structure parameters reflected a semi-unity graphical correlations. An excellent matching in the hydrogen bonds computed by (HSA and MEP) was recorded compared to the experimental XRD-packing result. Both nucleophilic and electrophilic sites on the molecule surface were detected by Mulliken and NPA population charge analysis. A high degree of matching between the theoretical HOMO/LUMO, DOS, GRD and TD-SCF/DFT/B3LYP compared to the experimental spectral results.

The DFT-computation of amide \rightleftharpoons imidic prototropic tautomerization through the QST2 method provided the possibility of such process *via* two steps reaction, simple flip-rotation with T.S_1 and single proton intramigration with T.S_2 (pseudo-four-membered-heterocyclic ring) formation.

The dependence of E_a 's on α was confirmed by applying both FWO and KAS isoconversional thermal kinetic methods, the kinetic decomposition process is likely to be multistep reaction,



Avrami's kinetic plot supported the second order decomposition reaction.

(E)/(Z)-thiophene-2-carbohydrazonic acid isomers reflected an excellent DNA-docking with cisplatin binding mode, moreover, Z-isomer is consider to be more active since more shorter H-bonds in [drug:DNA] complex were detected, meanwhile, no docking effect was detected in both *endo* and *exo*-thiophene-2-carbohydrazide isomers.

Conflicts of interest

The authors declare that they have no conflicts of interest.

Acknowledgements

The authors extend their appreciation to the Deanship of Scientific Research at King Saud University for funding this work through research group no (RG-1440-141).

References

- Y. L. Niu and T. Li, *Anal. Chim. Acta*, 2019, **1049**, 196–212.
- G. Kumar, V. S. Krishna, D. Sriram and S. M. Jachak, *Eur. J. Med. Chem.*, 2018, **156**, 871–884.
- W. Pan, J. Liu and X. Wang, *Tetrahedron*, 2018, **17**, 1468–1475.
- L. K. Kummar, M. S. Butler, E. Furlong, R. Blundell and A. Robertson, *Bioorg. Med. Chem.*, 2018, **26**, 5408–5419.
- S. Tolosa, N. Mora-Diez, A. Hidalgo and J. A. Sanson, *RSC Adv.*, 2014, **4**, 44757–44768.
- M. Boobalan, S. Ramalingam, M. Amaladasan, D. Tamilvendan, G. Venkatesa Prabhu and M. Bououdina, *J. Mol. Struct.*, 2014, **1072**, 153–172.
- B. Parrino, S. Ullo, A. Attanzio, S. Cascioferro and P. Diana, *Eur. J. Med. Chem.*, 2018, **158**, 236–246.
- Z. Wojnarowska, P. Włodarczyk, K. Kaminski, K. Grzybowska, L. Hawelek and M. Paluch, *J. Chem. Phys.*, 2010, **133**, 94507–94515.
- Z. Wojnarowska, K. Grzybowska, L. Hawelek, M. Dulski, R. Wrzalik, I. Gruszka and M. Paluch Drug, *Mol. Pharmaceutics*, 2013, **10**, 3612–3627.
- V. Balachandran, A. Janaki and A. Nataraj, *Spectrochim. Acta, Part A*, 2014, **118**, 321–330.
- E. Mateo-Martí and C. M. Pradier, *Spectrochim. Acta, Part A*, 2013, **109**, 247–252.
- R. Arora, U. Issar and R. Kakkar, *Comput. Theor. Chem.*, 2017, **1105**, 18–26.
- A. Rajavel, A. Aditya Prasad and T. Jeyakumar, *J. Mol. Struct.*, 2017, **1130**, 138–149.
- A. Yıldırım, M. Yıldırım and Ç. Kastas, *J. Mol. Struct.*, 2017, **1127**, 275–282.
- A. K. Srivastava, A. Kumar, N. Misra, P. S. Manjula, B. K. Sarojini and B. Narayana, *J. Mol. Struct.*, 2016, **1107**, 137–144.
- B. Bax, C. Chung and C. Edge, *Acta Crystallogr., Sect. D: Struct. Biol.*, 2017, **73**, 131–140.
- M. R. Aouad, M. Messali, N. Rezki, N. Al-Zaqri and I. Warad, *J. Mol. Liq.*, 2018, **264**, 621–630.
- P. Gilli, V. Bertolasi, L. Pretto, A. Lyčka and G. Gilli, *J. Am. Chem. Soc.*, 2002, **124**, 13554–13567.
- B. Bandyopadhyay and P. Biswas, *RSC Adv.*, 2015, **5**, 34588–34593.
- H. Tavakol, *J. Mol. Struct.: THEOCHEM*, 2010, **956**, 97–102.
- H. Tavakol, *Int. J. Quantum Chem.*, 2012, **112**, 1215–1224.
- H. Tavakol, *Struct. Chem.*, 2011, **22**, 1165–1177.
- H. Tavakol, *Int. J. Quantum Chem.*, 2011, **111**, 3717–3724.
- H. Tavakol, T. Hadadi and H. Roohi, *J. Mol. Struct.*, 2012, **53**, 649–658.
- G. M. Sheldrick, *Acta Crystallogr.*, 2008, **64**, 112–122.
- M. J. Frisch, G. W. Trucks and H. B. Schlegel, *et al.*, *Gaussian 09*, Gaussian Inc., Wallingford CT, 2009.
- S. K. Wolff, D. J. Grimwood, J. J. McKinnon, D. Jayatilaka, and M. A. Spackman, *Crystal Explorer 2.1*, University of Western Australia, Perth, 2007.
- H. E. Kissinger, *Anal. Chem.*, 1957, **29**, 1702–1706.
- T. Ozawa, *Bull. Chem. Soc. Jpn.*, 1965, **38**, 1881–1886.
- A. Barakat, M. Islam, A. Al-Majid, H. A. Ghabbour, S. Atef, A. Zarrouk and I. Warad, *J. Theor. Comput. Chem.*, 2018, **17**, 1–23.
- M. R. Aouad, M. Messali, N. Rezki, M. A. Said, D. Lentz, L. Zubaydi and I. Warad, *J. Mol. Struct.*, 2019, **1180**, 455–465.
- A. Titi, T. Shiga, H. Oshio, R. Touzani, B. Hammouti, M. Mouslim and I. Warad, *J. Mol. Struct.*, 2020, **1199**, 126995–127003.
- I. Warad, F. F. Awwadi, B. Abd Al-Ghani, A. Sawafta, N. Shivalingegowda, N. K. Lokanath, M. S. Mubarak, T. Ben Hadda, A. Zarrouk, F. Al-Rimawi, A. B. Odeh and S. A. Barghouthi, *Ultrason. Sonochem.*, 2018, **48**, 1–9.
- I. Warad, Y. Al-Demeri, M. Al-Nuri, S. Shahwan, M. Abdoh, S. Naveen, N. K. Lokanath, M. S. Mubarak, T. B. Hadda and Y. N. Mabkhot, *J. Mol. Struct.*, 2017, **1142**, 217–227.
- F. A. Saleem, S. Musameh, A. Sawafta, P. Brandão, C. J. Tavares, S. Ferdov, A. Barakat, A. Al Ali, M. Al-Noaimi and I. Warad, *Arabian J. Chem.*, 2017, **10**, 845–855.
- M. V. P. Ravi and S. P. Tewari, *J. Phys. Chem. A*, 2013, **117**, 10162–10169.
- J. Shi, Y. Lou, K. Zhou and D. Pan, *Spectrochim. Acta, Part A*, 2018, **204**, 209–216.
- I. Warad, S. Musameh, A. Sawafta, P. Brandão, C. J. Tavares, A. Zarrouk, S. Amereih, A. Al Ali and R. Shariah, *Ultrason. Sonochem.*, 2019, **52**, 428–436.
- N. Koga and J. Sestak, *J. Therm. Anal.*, 1991, **37**, 1103–1114.

