Synthesis, crystal structure, Hirshfeld surface investigation and comparative DFT studies of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl] thiazole-4-carboxylate

Muhammad Haroon1,2, Tashfeen Akhtar1*, Muhammad Yousuf3*, Muhammad Nawaz Tahir4, Lubna Rasheed5, Syeda Saniya Zahra6, Ihsan ul Haq6 and Muhammad Ashfaq4,7

Abstract
The ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1), a thiazole ester, was synthesized by refluxing 1-(2-nitrobenzylidene)thiosemicarbazide and ethyl bromopyruvate. The compound is characterized by spectroscopic and single crystal (SC-XRD) techniques. Non-covalent interactions that are responsible for crystal packing are explored by Hirshfeld surface analysis. All theoretical calculations were performed by DFT quantum chemical methods using 6-311G(d,p) and cc-pVTZ basis sets and compared. Theoretical harmonic frequencies of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) were optimized. Confirmation of hydrogen bonding sites was analyzed by molecular electrostatic potential (MEP) and Mulliken population analysis. The vibrational frequencies of characteristic functional groups and chemical shifts were found in good agreement with experimental assignments. Frontier molecular orbital (FMO) revealed relatively small HOMO–LUMO (highest occupied molecular orbital-lowest unoccupied molecular orbital) gap, which speaks off the nearly planar geometry and extended conjugation, as compared to the substituents with no conjugation possible. It has also been observed that –NO2 substituent plays a vital role for this relatively small HOMO–LUMO gap and overall electronic properties when compared with similar thiazole carboxylates (2–6, Table 6). Ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) was also evaluated for its anti-oxidant and anti-microbial activities.

Keywords: Thiazole, DFT calculations, Anti-oxidant, XRD, Spectroscopy

*Correspondence: tashfeenchem@must.edu.pk; nyousuf@postech.ac.kr
1 Department of Chemistry, Mirpur University of Science and Technology (MUST), 10250-Mirpur (AJK), Pakistan
2 Department of Chemistry, Ulsan National Institute of Science and Technology (UNIST), Ulsan, South Korea
Full list of author information is available at the end of the article
Introduction
1,3-Thiazole is an important sulphur and nitrogen containing azole [1]. Binding capability of sulphur and nitrogen in 1,3-thiazole, to induce sigma and pi interactions with receptor moieties, makes it an interesting class of heterocycles in drug designing [2]. It is very significant motif in pharmaceutical chemistry [3–5]. Thiazole derivatives are known for their diverse biological significance like anticancer [6], antiallergic [7], antiviral [8], antibacterial [9], antifungal [10], antitubercular [11, 12] and antiinflammatory [13–15] activities. Besides these, antiparasitic agents viz, nitazoxanide, tenonitrozole and aminitrozole have 1,3-thiazole ring in their structure [16].

Computational chemists use both experimental data and theoretical models in order to determine the structural features like bond lengths, bond angles and torsion angles besides accounting for spectroscopic properties like harmonic frequencies, vibrations and chemical shifts at molecular level. An important tool in the hand of theoretical chemists to compare and analyze the properties like geometry, vibrational frequency and dipole moment [17–24] of the molecules, is Density Functional Theory (DFT). The thiazole ester, ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) was synthesized for the purpose of biological studies. However, it was found interesting to investigate the structural features by comparing theoretical and experimental data and also to see the effect of the substituent i.e. nitro group (–NO₂), on the conjugation leading to difference in the HOMO–LUMO gape with extended conjugation. The analysis of theoretical results revealed that –NO₂ substituent has played a vital role in decreasing the HOMO–LUMO gape, furthermore; it has also exerted its effect on overall electronic properties when compared with compounds 2–6.

Materials and methods
Ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) was synthesized by the reaction of 1-(2-nitrobenzylidene)thiosemicarbazide and ethyl bromopyruvate (Scheme 1). The pure reagent grade chemicals were purchased and used as such. Thiosemicarbazide, 4-nitrobenzaldehyde and ethyl bromopyruvate were purchased form Merck Germany. Pre-coated
Silica 60 HF<sub>254</sub> Aluminum sheets (Merck, Germany) were used to monitor the reaction with thin layer chromatography (TLC). The melting point determination, functional group identification, NMR chemical shifts and single crystal analysis were carried out as reported elsewhere [25]. Dimethyl sulfoxide-deuterated (DMSO-<em>d</em><sub>6</sub>) was used to record the proton and carbon NMR (nuclear magnetic resonance), using 300 and 75 MHz frequency, respectively. High resolution mass spectrometry (HRMS) was carried out on Bruker Micro TOF-ESI (time of flight-electrospray ionization) spectrometer, positive targeted mode.

**Synthesis of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1)**

The desired compound (1) was obtained by refluxing 1-(2-nitrobenzylidene)thiosemicarbazide (0.5 g, 0.00237 mol) and ethyl bromopyruvate (0.3 mL, 0.00238 mol) for 4 h in absolute ethanol (20 mL). The reaction progress assessed by TLC. When the reactants consumed, as shown by TLC, quenching of reaction by adding ice cold water resulted into precipitates, which on filtration and washing with excess of water yielded pure compound [25].

**Table 1** SC-XRD experimental details of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1)

| Crystal data | Chemical formula | C<sub>13</sub>H<sub>12</sub>N<sub>4</sub>O<sub>4</sub>S |
|--------------|-----------------|---------------------------------|
| Temperature (K) | 296 | 320.33 |
| a, b, c (Å) | 14.423 (3), 11.270 (2), 9.2375 (16) | 105.510 (7) |
| V (Å<sup>3</sup>) | 1446.9 (5) | 0.38 x 0.18 x 0.16 |
| Radiation | Mo Kα | 0.38 x 0.18 x 0.16 |
| Absorption correction | Multi-scan (SADABS; Bruker, 2005) | 0.38 x 0.18 x 0.16 |
| No. of measured, independent and observed | 9460, 3409, 1921 | 0.38 x 0.18 x 0.16 |
| R<sub>int</sub> | 0.060 | 0.060 |
| (sin θ/λ)max (Å<sup>-1</sup>) | 0.658 | 0.658 |
| Refinement | | 0.658 |
| R(F<sup>2</sup> > 2τ(F<sup>2</sup>)), wR(F<sup>2</sup>), S | 0.061, 0.166, 1.01 | 0.061, 0.166, 1.01 |
| No. of reflections | 3409 | 3409 |
| No. of parameters | 200 | 200 |
| H-atom treatment | H-atom parameters constrained | H-atom parameters constrained |
| Δρ<sub>max</sub>, Δρ<sub>min</sub> (e Å<sup>-3</sup>) | 0.36, −0.29 | 0.36, −0.29 |

**Fig. 1** The ORTEP diagram drawn at probability level of 50% for (1). H-atoms are displayed by tiny circles of arbitrary radii.
Table 2 Hydrogen-bond geometry (Å, °) for (1)

| D—H···A | D—H  | H···A | D···A | D—H···A |
|---------|------|------|-------|---------|
| N3—H3A···O3i | 0.86 | 2.03 | 2.860 (3) | 161 |
| C5—H5···N2 | 0.93 | 2.44 | 2.764 (4) | 101 |
| C7—H7···O2 | 0.93 | 2.28 | 2.812 (4) | 116 |
| C12—H12A···O3 | 0.97 | 2.30 | 2.679 (5) | 102 |
| C9—H9···N4i | 0.93 | 2.58 | 3.502 (4) | 171 |
| C3—H3···O1ii | 0.93 | 2.55 | 3.432 (4) | 159 |

Symmetry codes: (i) −x, y+1/2, −z+1/2; (ii) −x, y+1/2, −z+1/2; (iii) x+1, y, −z+3/2

$J=7.0\text{ Hz}$, 7.60 (1H, dt, Ar−H, $J=1.5\text{ Hz}$, $J=8.4\text{ Hz}$), 7.78 (1H, t, Ar−H, $J=7.7\text{ Hz}$), 8.02 (2H, m, Ar−H), 7.82 (1H, s, 1,3-thiazole ring C-5), 8.37 (1H, s, −CH=CH=CH−), 12.63 (1H, s, −N−NH−C−); $^{13}$C–NMR: δ (ppm) 14.6 (−CH3), 60.9 (−OCH2−), 120.2 (C-5, 1,3-thiazole ring), 125.2, 128.1, 128.7, 130.4, 137.4, 147.9 (Ar−C), 137.6 (−CH=N−), 143.3 (C-4, 1,3-thiazole ring), 161.7 (C=O), 168.2 (C-2, 1,3-thiazole ring); HRMS: 321.9680 [M+H]$^+$, 343.0700 [M+Na]$^+$. 

Single Crystal X-Rays Diffraction (SC-XRD) studies

The crystallization of ethyl 2-{[2-(2-nitrobenzylidene) hydrazinyl]thiazole-4-carboxylate (1) from tetrahydrofuran (THF) yielded the crystals suitable for diffraction collection using single crystal X-ray diffractions. Table 1 presents the experimental conditions, different parameters and structure refinements. The data reduction and cell refinement carried out using SAINT and data collection by APEX 2 [26]. SHELXS97 [27] and SHELXL97 [28] were used for solving and refining the structure, respectively. PLATON [29], Mercury 3.6 software and ORTEP-3 for Windows [30] were used to draw molecular graphics. The CIF is deposited at Cambridge Crystallographic Data Centre (CCDC=1,951,603).

Fig. 2 Packing diagram showing 2D sheet formed by N–H…O, C–H…N and C–H…O bonding for (1)
Results and discussion
The synthesis of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) was achieved in two steps form thiosemicarbazide via thiosemicarbazone and cyclization with ethyl bromopyruvate. The synthesis was confirmed by spectroanalytical techniques. The spectroscopic data verified the proposed synthesis and the structure was later confirmed by the SC-XRD.

Single crystal X-ray
The SC-XRD inspection showed that ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) (Fig. 1) exists in E-configuration, the benzylidene-hydrazinyl moiety A (C1–C7/N1/N2) and the thiazole ring B (C8/N4/C9/C10/S1) are found to be planar with respective root mean square (r.m.s.) deviations of 0.0287 and 0.0032 Å. The dihedral angle between A/B is 8.41 (9)°. The planarity of the thiazole ring in two most closely related reported crystal structures, one having methyl substituted phenyl ring and second having bromo substituted phenyl ring was obvious by r.m.s. deviation of 0.004 and 0.0072 Å [25]. The nitro group C (N1/O1/O2) and carboxylate group D (C11/O3/O4) are oriented at a dihedral angle of 26.84 (10)° and 4.6 (5)° with the parent groups A and B, respectively. The C-atoms (C12/C13) are at a distance of -0.003 (11) and 1.329 (13) Å, respectively, from the carboxylate group as compared to their distance of 0.034 (16) and 1.440 (21) Å from the carboxylate group in closely related crystal structure having bromo substituted phenyl ring instead of nitro substituted phenyl ring [25]. Intramolecular H-bonding of type C–H…N and C–H…O are responsible for stabilization of molecular configuration. CH of imine functional group is found to engage with O-atom of nitro group (O2) to form S(6) loop through intra C–H…O bonding. Similarly, N-atom of imine functional group is found to engage with CH of phenyl ring to form S(5) loop through C–H…N bonding. Another S(5) is formed by C–H…O bonding, where O-atom is from carboxylate group and CH is from alkyl group (C12/C13). These intramolecular H-bonded loops are displayed in Fig. 1 and specified in Table 2. The molecules are primarily connected to each other in the form of dimers through strong N–H…O and comparatively weak C–H…N bonding to form R21(9) loop as displayed in Fig. 2 and specified in Table 2 [32]. Infinite C7 chain is formed by N3-H3A…O3 bonding whereas C4 chain is formed by C–H…N bonding. These chains extend along b crystallographic axis in a zigzag fashion. The molecules are also interlinked by C–H…O bonding, where para CH of phenyl ring (C1–C6) acts as donor and O-atom is from nitro group. Due to this C–H…O bonding, another zigzag C6 chain is formed along c crystallographic axis. Besides the role of H-bonding in packing of molecules, weak non-covalent interaction named as off-set π–π stacking interaction also play significant role in the crystal packing. Five membered ring of one molecule located

Fig. 3 Graphical view of π–π stacking interaction for (1). Distance shown is measured in Å. H-atoms are omitted for clarity
at asymmetric location is engaged in off-set π–π stacking interaction with the five membered ring of neighboring symmetry related molecule (-x,1-y,-z). Centroid to centroid distance and ring off-set for this particular interaction are found to be 3.92 Å and 1.798 Å, respectively as displayed in Fig. 3.

Hirshfeld surface analysis (HSA)
For further exploration of non-covalent interactions that are accountable for the packing of molecules, Hirshfeld surface inspection is conducted on Crystal Explorer 17.5 [33]. Hirshfeld surface (HS) plotted over d_{norm} can be utilized to represent intermolecular interactions by colour coding [34–36]. The atomic contacts having distance shorter, equal and longer than summation of Van der Waals radii are represented by red, white and blue regions on HS, respectively. Bright red spots on the HS near NH group, carbonyl O-atom, para CH of nitro phenyl ring, one of the O-atom of nitro group indicate that these atoms are involved in H-bonding as displayed in Fig. 5. N-atom and CH of five membered ring are also engaged in H-bonding but no red spot is obvious in Fig. 4 due to graphical limitations. No red spot is found on the HS near imine N-atom indicating that this atom is not engaged in any kind of H-bonding. HS plotted over shape index is utilized to check that whether π–π stacking interaction is present in crystal packing or not. Consecutive triangular regions of red and blue on the HS around the five membered ring indicated that π–π stacking interaction is present in crystal packing as displayed in Fig. 5. As these triangular regions on the HS are not exactly in the center of five membered ring suggesting off-set π–π stacking interaction. In order to find out the percentage contribution of each interatomic contact involved in crystal packing, 2D fingerprint plots are utilized. In these plots, d_i and d_c stands for distance

Fig. 4 HS plotted over d_{norm} in the range — 0.5540 to 1.4207 a.u. for (1)

Fig. 5 HS plotted over shape index in the range — 1 to 1 a.u. for (1)
Fig. 6 2D plots for (1), a for overall interaction, b–f for major contributor in crystal packing.
Fig. 7 a–j 2D plots for minor contributor in crystal packing for (1)
from the HS to nearest atom inside and outside the HS, respectively. 2D fingerprint plot for overall interactions in the crystal packing is displayed in Fig. 6a. The sky blue region in the center of this plot indicates the existence of π–π stacking interaction in crystal packing. The contribution of each individual interatomic contact is calculated including reciprocal contact. It is found that O…H, H…H, C…H, N…H and S…H interatomic contacts are the major contributors in crystal packing with percentage contribution of 29.9%, 28%, 12.4%, 7.4% and 5.5%, respectively as displayed in Fig. 6b–f. C…C, N…C, S…C, O…C, O…N, S…N, N…N, O…O, S…S, and S…O interatomic contacts are found to be minor contributor in crystal packing with percentage contribution of 3.5%, 3.4%, 2.4%, 2.3%, 1.6%, 1.1%, 1%, 0.3%, 0.2% and 0.1%, respectively as displayed in Fig. 7a–j. Furthermore, the interaction of an atom located inside the HS is computed with the atoms of molecules present in the surrounding of HS to have an idea which atom of a molecule is interacting strongly with the atoms of neighboring molecules. H-atoms located inside the HS interact strongly with the atoms of molecules in the neighbouring of HS, percentage contribution of this interaction is found to be 52.4%. The percent (%) contribution of all such interactions are displayed in Fig. 8.

Quantum chemical calculations
Guassian09 package [37–39] and Gauss-View molecular visualization software [40] was used in order to carry out calculation studies on personal computer. DFT (DFT/B3LYP) method [41, 42] with 6-311G(d,p) and cc-pVTZ basis sets were used for geometry optimization.
and further calculations (Fig. 9). The optimized structures obtained from these two methods were compared (Fig. 10) which showed almost same results. Optimized geometry was used in order to ascertain different aspects like vibrational spectra, chemical shifts, quantum chemical and thermodynamic properties of the compound, theoretically [43].

**Vibrational spectra**

The vibrational spectra of the compound were computed theoretically (DFT/B3LYP method with 6-311G(d,p) and cc-pVTZ basis sets). The results obtained by two methods were subjected to scaling factors of 0.966 and 0.966 [44, 45] for 6-311G(d,p) and cc-pVTZ basis sets, respectively. The vibrational modes of the compound are assigned by animation of Guass View software, for both used methods. Total energy distribution (TED) calculations for compound (1) were executed with the scaled quantum mechanical (SQM) program [46, 47]. Several factors were taken into account for matching of the experimental and theoretical vibrational data of the compound (1), such as: animation option of the Gauss View package program, relative intensities of the vibrational bands and previously reported data in the literature [37]. The compound (1) contains 34 atoms and it shows 96 normal vibrational modes having a member of C point group with identity (E) symmetry operation. The conformer having the lowest optimization energy has been selected for calculating vibrational spectra of the molecule. At the end all the experimental results were compared to the theoretical results. Theoretical findings for infrared wavenumbers together with experimental results and TED values of the compound (1) are given in supporting information (Additional file 1: Table S1).

![Fig. 12](image1) Correlation graphics of calculated and experimental chemical shifts of compound (a) $^1$H–NMR (b) $^{13}$C–NMR by B3LYP/6-311G(d,p)

![Fig. 13](image2) Correlation graphics of calculated and experimental chemical shifts of compound (a) $^1$H–NMR (b) $^{13}$C–NMR by B3LYP/cc-pVTZ
In the high wavenumber region bands appeared at around 3493 cm\(^{-1}\) are due to NH vibrations while those appeared at 3273–3036 cm\(^{-1}\) are due to symmetric C–H vibrations. The broadened shape of the band at this region could be attributed to the indication of residue of moisture in the sample and intra and intermolecular interactions. The band appeared at 1685 and 1571 cm\(^{-1}\) are due to C=O and –NO\(_2\) stretchings respectively [48]. The bands at 1298, 1251, 1244, 1160 and 1103 cm\(^{-1}\) are due to C–H stretching. The bands at 1336 and 742 cm\(^{-1}\) are due to C–C stretching [49, 50]. For the sake of better comparison experimental and calculated results were plotted as shown in Fig. 11.

**NMR spectra**

The \(^1\)H- and \(^{13}\)C–NMR spectra of the compound (1) were obtained in deuterated dimethyl sulfoxide and signals calibrated with respect to TMS and assigned the chemical shifts to respective protons and carbons on the basis of the chemical environment. The proton NMR spectrum exhibited a triplet and quartet at 1.28 and 4.24 ppm corresponding to methyl and methylene of ethyl ester moiety in the molecules, respectively. The ortho substitution of the benzene ring was confirmed by the presence of a triplet and a triplet of doublets at 7.60 and 7.78 ppm, respectively. The triplet of doublets indicates one small coupling constant (\(J=1.5\) Hz) corresponding to meta splitting and one large coupling constant (\(J=8.4\) Hz) for the ortho splitting. Two remaining aromatic protons were multiplet at 8.04–7.99 ppm. The singlet at 7.82 ppm, integrating for one proton, was assigned to thiazole ring proton. The benzylidene –CH– proton was observed as singlet at 8.37 ppm. The –NH proton was observed as a downfield singlet at 12.63 ppm.

The carbon NMR spectrum showed thirteen signals corresponding to total number of carbons in the compound. Two signals in aliphatic region at 14.6 and 60.9 ppm were assigned to the ethoxy carbons i.e. –CH\(_3\) and –OCH\(_2\), respectively. The six aromatic carbons of benzene ring were seen between 125.2 to 147.9 ppm. The benzylidene –CH– and thiazole ring C-4 was observed at 137.6 and 143.3, respectively. The carbonyl of ester

### Table 3

| Parameters | B3LYP/6-311G(d, p) | B3LYP/cc-pVTZ |
|------------|-------------------|--------------|
| \(E_{\text{HOMO}}\) | −6.1093 | −6.1148 |
| \(E_{\text{LUMO}}\) | −2.8798 | −2.8763 |
| \(I\) (eV) | 6.1093 | 6.1148 |
| \(A\) (eV) | 2.8798 | 2.8763 |
| \(\chi\) (eV) | 4.4945 | 4.4955 |
| \(\eta\) (eV) | 1.6147 | 1.6192 |
| \(S\) (eV)\(^{-1}\) | 0.3097 | 0.3088 |

### Table 4

| MO(s) | B3LYP/6-311G(d, p) | B3LYP/cc-pVTZ |
|-------|-------------------|--------------|
| \(E(eV)\) | \(\Delta E(eV)\) | \(E(eV)\) | \(\Delta E(eV)\) |
| HOMO | −6.1093 | 3.2295 | −6.1148 | 3.2385 |
| LUMO | −2.8798 | −2.8763 |
| HOMO−1 | 7.6517 | 5.5062 | 7.6517 | 5.4955 |
| LUMO+1 | 2.0425 | 2.0735 |
| HOMO−2 | 1.2770 | 1.3205 |
| LUMO+2 | 1.2770 | 1.3205 |

**Fig. 14** FMOs at B3LYP/cc-pVTZ level
and thiazole C-2 were observed at 161.7 and 168.2 ppm, respectively. The proton and carbon spectra confirmed the proposed structure for the synthesized compound.

GIAO approach was used in order to calculate chemical shifts using B3LYP/6-311G(d,p) and B3LYP/cc-pVTZ, which are then converted to the TMS scale as reported earlier [25] using the formula i.e. δ = Σo – Σ. IEF-PCM (Integral-Equation-Formalism Polarizable Continuum Model) model was employed in order to incorporate the solvent effects [51]. Correlations graphs were produced for comparison of theoretically calculated NMR results to experimental values (Figs. 12 and 13). The correlation co-efficients for B3LYP/6-311G(d,p) (Fig. 12) are 0.9964 for 1H and 0.9977 for 13C NMR, while correlation co-efficients for B3LYP/cc-pVTZ (Fig. 7) 0.9976 for 1H and 0.9982 for 13C NMR were observed which reflects better agreement in experimental and theoretical results. The experimental and theoretically calculated NMR data by both methods has been presented in supporting information (Additional file 1: Tables S2, S3).

### HOMO and LUMO analysis

Frontier molecular orbitals (FMO) have unique importance for calculation of electronic states and stability of a molecule because these orbitals are at frontiers of electron occupation i.e. highest occupied molecular orbital (HOMO) and lowest lying unoccupied molecular orbital (LUMO) [44]. Theoretically calculated HOMO and LUMO are shown in Fig. 8. The compound possessed 83 occupied molecular orbitals. The respective energies for HOMO and LUMO are −6.1093 and −2.8798 eV for B3LYP/6-311G(d,p) levels and −6.1148 and −2.8763 for B3LYP/cc-pVTZ levels, respectively. Ionization potential (I), electron affinity (A) electronegativity (χ), chemical hardness (η) and chemical softness (S) have been obtained from FMO data (Table 3) [52]. Both methods produced almost the same results.

FMO theory assists in the study of photo-electronic properties of the interesting compounds. The energy of HOMO's and LUMO's were computed by DFT method using B3LYP level along with 6-311G(d,p) and cc-pVTZ subset levels. The results are presented in Fig. 8 which shows the electronic contribution in HOMO-2, HOMO-1 and HOMO with LUMO, LUMO+1 and LUMO+2 of the compound. There is the delocalization of HOMO on all atoms of the molecule except ethyl group while concentration of LUMO at all atoms except ester group (Fig. 14). Theoretical data of

| Parameters | \( \lambda_{\text{Exp}} \) (nm) | \( \lambda_{\text{DFT}} \) (nm) | \( E_{\text{DFT}} \) (eV) | \( f \) | MO contribution |
|------------|-----------------|-----------------|-----------------|-----|----------------|
| 6-311G(dp) | 436.53          | 2.8402          | 0.2548          |     | H → L (70%)    |
| 260,350    | 340.35          | 3.6428          | 0.3228          |     | H → L + 2 (66%)|
| 256,338    | 331.35          | 3.7417          | 0.0296          |     | H → L + 1 (44%)|

\( \lambda_{\text{Exp}} \) experimental wavelength; \( \lambda_{\text{DFT}} \) calculated wavelength; E, Excitation energy (ev); MO, molecular orbitals; a, THF, b, DMF, c, DMSO

![Fig. 15 MEP map of compound in gas phase using B3LYP/cc-pVTZ](image-url)
HOMO (H) and LUMO (L) energies has been described in Table 4 which shows almost same values of energies for both methods i.e. 6-311G(d,p) and cc-pVTZ [53].

**UV-Vis analysis**

TDDFT studies (B3LYP method with 6-311G(d,p) and cc-pVTZ basis sets) were carried out in gas phase to get absorption properties. Calculated parameters i.e. wavelength of maximum absorption ($\lambda_{max}$), oscillator strength ($f$), excitation energy ($E_{DFT}$) and orbitals mainly involved for excitation are listed in Table 5 below [54].

The value of computed absorption maxima ($\lambda_{DFT}$) were found to be 436.53, 340.35 and 437.92, 342.88 nm for B3LYP/6-311G(d, p) and B3LYP/ccPVTZ levels respectively with percentage of orbital contribution of [H $\rightarrow$ L (70%) and H $\rightarrow$ L + 2 (66%)]. Calculated wavelength ($\lambda_{DFT}$) 340.35 and 342.88 employing two different basis sets matched nicely with experimental ($\lambda_{Exp}$) value of 350 (DMSO), 338 (DMF) and 360 (THF). It may be concluded that solvent plays its part for the difference between $\lambda_{DFT}$ and $\lambda_{Exp}$ value (Additional file 1: Figure S1).

**Molecular electrostatic potential (MEP)**

The information about electron acceptor and electron-donor regions in a molecule can be easily visualized by MEP. Here the electrostatic potential decreases in the following order red < orange < yellow < green < blue. Molecular electrostatic map of compound is shown in Fig. 15 (B3LYP/cc-pVTZ basis set) where blue and red indicates the areas of strongest attraction and strongest repulsion, respectively. It is concluded that area of strong repulsions i.e. red resides on O1, O2, O3 and N4 atoms while that of strong attractions i.e. blue resides at N2 and N3 atoms, respectively. Therefore, it is assessed that N2 and N3 atom are electron-donor while O1, O2, O3 and N4 atoms are electron-acceptor [55].
Mulliken population analysis

The charge distribution on the molecule was calculated theoretically with the help of B3LYP methods using 6-311G(d, p) and cc-pVTZ basis sets (Fig. 16). It is the charge distribution within the molecule, thereby rendering important information about hydrogen bonding within the molecule [54, 55]. It can also be used to characterize the electronic charge distribution in a molecule and the bonding, antibonding, or nonbonding nature of the molecular orbitals for particular pairs of atoms. Mulliken charges of hydrogen atoms are in positive number while negative for carbons except carbon atoms that are attached to oxygen and nitrogen atoms (due to strong electronegativity of O and N atoms). Oxygen and Nitrogen atoms play the vital role in inter- and intra-molecular hydrogen bonding.

Thermodynamic properties

The vibrational analysis (B3LYP method with 6-311G(d,p) and B3LYP/cc-pVTZ basis sets) was used to calculate the thermodynamical parameters and properties of the compound i.e. entropy ($S^o_m$), heat capacity ($C^o_{p,m}$) and enthalpy ($H^o_m$). Thermodynamical parameters i.e. entropies, standard heat capacities and enthalpies exhibit a direct relation with temperature. Their correlation equations for both B3LYP/6-311G(d,p) (Eqs. 1–3) and B3LYP/cc-pVTZ (Eq. 4–6) levels have been calculated (Fig. 11) and may be used for further studies of the compound and may serve as guideline for similar compounds. From these correlations it is evident that entropy, enthalpy and heat capacity of the compound (1) increases with increase in temperature.

Heat capacity is due to the rotational and vibrational energy that a molecule possesses. At low temperature there is not enough energy in order to excite many rotations and vibrations, so the slope is small. At very high temperature many energy levels are populated, so the rate of increase becomes constant. The same phenomenon has been observed in case of molecule (1) (Fig. 17). As enthalpy is sum of total energies of a molecule, so, it follows the same pattern of slow increase at low temperature but at high temperature it increases steadily due to the dominance of entropy. Entropy of a molecule increases with increase in temperature. The same phenomenon of enthalpy and entropy has also been observed in case of compound (1) also.

$$S^o_m = -8E^{-5}T^2 + 0.3079T + 71.92 \left( R^2 = 0.9911 \right)$$  \tag{1}$$

$$H^o_m = 7E^{-5}T^2 + 0.0363T + 151.81 \left( R^2 = 0.9901 \right)$$ \tag{2}$$

$$C^o_{p,m} = -1E^{-4}T^2 + 0.2593T + 7.5953 \left( R^2 = 0.9917 \right)$$ \tag{3}$$

$$S^o_m = -8E^{-5}T^2 + 0.3081T + 71.98 \left( R^2 = 0.9916 \right)$$ \tag{4}$$

$$H^o_m = 7E^{-5}T^2 + 0.0362T + 151.92 \left( R^2 = 0.9924 \right)$$ \tag{5}$$

$$C^o_{p,m} = -1E^{-4}T^2 + 0.2588T + 7.5854 \left( R^2 = 0.9913 \right)$$ \tag{6}$$

Comparison of HOMO and LUMO analysis of compound with other derivatives

Careful analysis of molecular orbitals of molecules (1–6) reveals that except –CO– and OEt groups, HOMO and LUMO are completely delocalized on all atoms. The

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**Table 6** Comparison of frontier orbital energies, electronegativity, hardness and softness of compounds (1–6)

| Parameter/compound | Present Study | [54] | [25] | [55] | [54] | [55] |
|--------------------|-------------|------|------|------|------|------|
| $E_{HOMO}$         | -6.1148     | -5.8849 | -5.4942 | -5.864 | -5.9649 | -5.989 |
| $E_{LUMO}$         | -2.8763     | -1.9700 | -1.5863 | -1.882 | -2.0568 | -2.170 |
| $\Delta E$        | 3.2385      | 3.9194 | 3.9079 | 3.982 | 3.9081 | 3.819 |
| $I$ (eV)          | 6.1148      | 5.8849 | 5.4942 | 5.864 | 5.9649 | 5.989 |
| $A$ (eV)          | 2.8763      | 1.9700 | 1.5863 | 1.882 | 2.0568 | 2.170 |
| $\chi$ (eV)       | 4.4955      | 3.9274 | 3.5402 | 3.873 | 4.0109 | 4.0795 |
| $\eta$ (eV)       | 1.6192      | 1.9574 | 1.9539 | 1.911 | 1.9541 | 1.9095 |
| $S$ (eV)$^{-1}$   | 0.3088      | 0.2543 | 0.2559 | 0.2511 | 0.2559 | 0.2618 |
| $M$               | -4.4955     | -3.9274 | -3.5402 | -3.873 | -4.0109 | -4.0795 |
energy gap (ΔE) between HOMO and LUMO is summarized in Table 6 below shows lowest energy gap (3.2385 eV) in case of compound (1) having electron withdrawing –NO2 substituent while highest energy gap (3.982 eV) in case of molecule (4) having methyl substituent. The energy gaps of other molecules are in between (3.819 eV–3.9149 eV) although there is a little difference. This data clearly indicates different substituents (electron donor/electron acceptor) has great effect on electronic properties of the molecule (Table 6). Therefore compound (1) possess better capability for electron transfer (lowest ΔE) as compared to other compounds (2–6) [25].

The calculated data of $E_{\text{HOMO}}$ and $E_{\text{LUMO}}$ can be used in order to determine other molecular parameters which are also summarized in Table 6. There is an inverse relationship between energy gap, softness and reactivity of species while a direct relation towards hardness and stability of the molecule. Therefore, compound (4) which possess highest energy gap will resist any change indicating low reactivity while compound (1) will have high reactivity. The calculated energy gap of compound (1) is lowest but ionization potential (6.1148 eV) and electron affinity (2.8763 eV) values of compound (1) are greater than others. The hardness of molecule (1) is lowest (1.6192 eV) while softness is greatest (0.3088 eV) thus confirming the inverse relationship of energy to softness and direct relationship to hardness, respectively. The calculated values of softness (0.2511 eV–0.2618 eV) and hardness (1.9095 eV–1.9574 eV) of other molecules are comparable (Table 6) [58, 59].

The chemical potential ($\mu$) is related to the stability of a molecule and there is a direct relationship. The value of $\mu$ for compound (1) is least (−4.4955 eV) while higher (−3.5402 eV) for compound (3). The order of chemical potential is as under:

$$[1(\mu = -4.4955 \text{ eV})] < [6(\mu = -4.0795 \text{ eV})] < [5(\mu = -4.0109 \text{ eV})]$$
$$< [2(\mu = -3.9274 \text{ eV})] < [4(\mu = -3.873 \text{ eV})] < [3(\mu = -3.5402 \text{ eV})]$$

The data also indicates that −CH$_3$, −Br and −OH substituent has almost comparable effect whereas −NO$_2$ substituent has an immense effect on overall electronic properties of the compound. Based on calculated data of hardness it can be inferred that the compound (2) (1.9574 eV) is the most stable and less reactive compared to compound (6) (1.6192 eV). Although hardness values of compounds (2–6) are almost comparable (Table 6).

### Antioxidant assays [57]

Percent free radical scavenging assay (%FRSA), total antioxidant capacity (TAC) and total reducing power (TRP) studies of compound (1) exhibit 77.52 ± 0.34 percent scavenging, 144.07 ± 0.16 and 62.13 ± 0.84 μgAAE/mg values, respectively. The %FRSA, TAC and TRP results support the use of compound (1) as an antioxidant agent. However, the antifungal and antibacterial studies did not exhibit any significant activity.

### Conclusions

The synthesis of ethyl 2-[2-(2-nitrobenzylidene)hydrazinyl]thiazole-4-carboxylate (1) and its characterization by spectroscopic techniques (UV–Vis, FT-IR, $^1$H- and $^{13}$C-NMR), HRMS and SC-XRD is accomplished. The structure was optimized by computational methods. X-rays analysis revealed that molecular configuration is stabilized by C–H...N and C–H...O bonding whereas crystal packing is stabilized by intermolecular N–H...O, C–H...N, C–H...O bonding and weak off-set π–π stacking interaction. Non-covalent interactions that are responsible for crystal packing are further explored by Hirshfeld surface analysis. Moreover, the experimental and theoretical spectroscopic results were in excellent agreement with each other. The MEP map shows that N2 and N3 atom are electron-donor while O1, O2, O3 and N4 atoms are electron-acceptor. The atoms which are electron rich or electron deficient are responsible for intra- and intermolecular attractive or repulsive interaction. The nearly planar geometry is also reflected by the low HOMO–LUMO gape, due to extended conjugation in the molecule. Theoretical data of compounds (2–6) has also been compared with that of (1) that indicates that −CH$_3$, −Br and −OH substituents has almost comparable effect whereas −NO$_2$ (Compound 1) substituent has great effect on overall electronic properties of the compound. The larger value of hardness than softness reflets greater stability of the compound that is important for technological applications. The compound (1) was also tested for antioxidant (%FRSA, TAC and TRP studies) and antimicrobial (antibacterial and antifungal) potential. It is concluded based on antioxidant results that compound (1) could be used as UV absorber in sunscreen lotions and may protect the skin damage from UV radiations.

### Abbreviations

CCDC: Cambridge Crystallographic Data Centre; DFT: Density Functional Theory; DMSO-d$_6$: Dimethyl Sulfoxide-deuterated; FMO: Frontier Molecular Orbital; FRSA: Free Radical Scavenging Assay; FT-IR: Fourier Transform Infrared; HOMO: Highest Occupied Molecular Orbital; HRMS: Hight Resolution Mass Spectrometry; HS: Hirshfeld Surface; IF-PCM: Integral-Equation-Formalism Polarizable Continuum Model; LUMO: Lowest Unoccupied Molecular Orbital; MEP: Molecular Electrostatic Potential; NMR: Nuclear Magnetic Resonance; ORTEP: Oak Ridge Thermal Ellipsoid Plot; SC-XRD: Single Crystal X-Rays.
Diffraction, TAC; Total Antioxidant Capacity; TDDFT: Time Dependent Density Functional Theory; TED: Total Energy Distribution; TLC: Thin Layer Chromatography; TOF-ESI: Time of Flight-Electrospray Ionization; TRP: Total Reducing Power; UV–Vis: Ultra violet-visible.

**Supplementary Information**

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**Additional file 1.** Synthesis, crystal structure, Hirshfeld surface investigation and comparative DFT studies of ethyl 2-(2-(2-nitrobenzylidene)hydrazinyl)thiazole-4-carboxylate. Table S1. Comparison of the experimental and calculated vibrational wavenumbers (cm⁻¹) of compound 1 in gas phase. Table S2. Experimental and theoretically calculated ¹H NMR. Table S3. Experimental and theoretically calculated ¹³C NMR.

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**Authors’ contributions**

MH has conducted the research, compiled the experimental data and write up of synthetic part. TA Concept idea and supervisor of this research and paper write up. MY and LR performed DFT Quantum chemical calculation and write up of that part. MNT and MA collected XRD single crystal data and Hirshfeld surface analysis and write up. SZ and IH carried out biological studies and write up. All authors read and approved final manuscript.

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**Availability of data and materials**

CCDC (Cambridge Crystallographic Data Center) number 1951603 contains the supplementary crystallographic data for the ethyl 2-(2-(2-nitrobenzylidene)hydrazinyl)thiazole-4-carboxylate (1) This data can be obtained free of charge via http://www.ccdc.cam.ac.uk/conts/retrieving.html, or from the Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: (+44) 1223-336-033; or e-mail: deposit@ccdc.cam.ac.uk. While the computational data is presented in Additional files 1, 2.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

No competing interests.

**Author details**

1Department of Chemistry, Mirpur University of Science and Technology (MUST), 10250-Mirpur (AJK), Pakistan. 2Department of Chemistry, Government Major Muhammad Afzal Khan (Shaheed), Boys Degree College Afzalpur, Mirpur, (Affiliated with Mirpur University of Science and Technology (MUST)), 10250-Mirpur (AJK), Pakistan. 3Department of Chemistry, Ulsan National Institute of Science and Technology (UNIST), Ulsan, South Korea. 4Department of Physics, University of Sargodha, Sargodha, Punjab, Pakistan. 5Department of Chemistry, Division of Science and Technology, University of Education, Township, Lahore, Pakistan. 6Department of Pharmacy, Quaid-I-Azam University, 45320 Islamabad, Pakistan. 7Department of Physics, University of Mannawi, Mannawi, Punjab, Pakistan.

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