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Regioselective cyclocondensations with thiobarbituric acid: spirocyclic and azocine products, X-ray characterization, and antioxidant evaluation†

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Multicomponent cyclocondensations of 5-amino-3-methyl-1-phenyl-1*H*-pyrazole (AMPZ), thiobarbituric acid, and *p*-formaldehyde under conventional thermal heating or ultrasonic irradiation were studied. Treatment of the reaction mixture in ethanol in an ultrasonic bath for 3 h produced azocine compound **4b**, while the same mixture in ethanol under reflux conditions for 15 h produced spiro compound **4a**. This work encompasses intricate experimental details, X-ray diffraction measurements, and multifaceted computational analyses employing methods such as the density functional theory and Hirshfeld surface analysis. Crystallographic investigations revealed the molecular structure of the compound and clarified its interactions involving hydrogen bonds and weak intermolecular forces. This article describes the synthesis and characterization of a novel spirocyclic compound. The study also evaluated the antioxidant potential *in vitro* using the DPPH and ABTS methods. The results showed that these compounds showed the best free radical scavenging ability, even in very small amounts, and that even at very low concentrations, these compounds showed excellent radical scavenging potential. Surprisingly, these compounds exhibited strong (ABTS⁺) radical scavenging activities, mainly attributed to the HAT mechanism, indicating their potential as therapeutic agents. Facile multipurpose, three-component selective procedures for new spiroheterocycles have been proposed, presenting intriguing perspectives in the field of medicine, particularly in the field of antioxidants. The geometric values of the computationally optimized structure were calculated using the density functional theory in LC-BLYP/6-31(d), aligned with the X-ray diffraction data, reinforcing the precision of our findings.

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1. Introduction

Alkaloids are nitrogenous organic compounds that constitute an important class of secondary metabolites produced by numerous terrestrial and marine organisms (plants, fungi, and bacteria), and are of utmost importance in the defense, proliferation, and reproduction of the species that contain them.^{1,2} These secondary metabolites are usually characterized by many biological activities; therefore, they are of great interest to researchers in organic and pharmacological chemistry. Among alkaloids, the most studied compounds have been characterized as N-heterocycles with five, six, and seven members (indoles, imidazoles, pyridines, azepines, and their respective variants); however, eight-membered alkaloids have not been extensively studied.¹

The structural blocks of 8-membered alkaloids that stand out are azocines and azocanes, which are usually found in fungi such as *Penicillium dimorphosporum*, *Penicillium simplicissimum* and *Laccaria proxima*, as well as in plants like *Campylospermum*

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flavum and some marine sponges, such as the Okinawan sea sponge *Pellina* sp.^{3–7} Azocines and azocanes have been shown to be important biological centers, acting as insecticides,^{7–10} neuroprotective agents,^{11–14} antibacterials,^{3–6,15} antifungals,^{4,6,16} antimalarials,^{17–19} antivirals,²⁰ anticancer^{21,22} and anti-inflammatory.^{23,24}

In contrast, within the group of azocines and azocanes derivatives with bridges (see Fig. 1) have proven to be interesting scaffolds at the pharmacological level; for example, alstomarine B and C molecules have shown significant cytotoxicity against four lines of human osteosarcoma tumor cells (Saos-2, Mg-63, U2-OS, and SOSP-9607).²⁵ Likewise, the Uleine compound inhibits enzymes such as acetylcholinesterase, butyrylcholinesterase, and β -secretase and was patented in 2010 for the prevention and/or treatment of infectious diseases such as HIV.^{26,27} In contrast, FR901483 plays a key role in exerting strong immunosuppressive activity *in vitro* and prolonging skin graft survival time in a mouse allograft model.²⁸ It is still important to highlight that, to date, no studies have reported the antioxidant capacity of bridged azocines, and even less so that they have an antioxidant capacity comparable to that of a reference antioxidant such as ascorbic acid, so this is an interesting area of research to provide a new therapeutic approach for this type of molecule.

Thus, there has been a growing interest in the synthesis of new compounds derived from bridged azocanes and azocines to obtain molecules that are potentially applicable to medicine. However, obtaining these compounds can be challenging for organic chemists because of the difficulty involved in obtaining an eight-membered ring and effectively assembling the bridge.^{29,30} Mohammad *et al.* conducted the synthesis of a new series of dipyrzolo[1,5]diazocine-3,8-diones by the one-pot reaction of 3-aminopyrazolone with substituted benzaldehydes in the presence of catalytic *p*-toluenesulfonic acid.³¹ Also, Vachan *et al.* recently reported the synthesis of 2,6-methanobenzo[d][1,3]diazocines *via* the reaction between primary amine beta-ketoesters and 2-aminoarylaldehydes under reflux for 1 h³² (Scheme 1). To the best of our knowledge, there are no reports on the synthesis of 1,3-diazocines fused with pyrazole derivatives. Furthermore, in this work, a multicomponent synthesis is presented, which, depending on the energy source, can favor the formation of the azocine derivative or a spirone derivative, which together with the pyrazolopyrimidine are two

other types of scaffold widely studied both in terms of reactivity, as well as in a wide range of promising biological properties.^{33–43}

Taking the above into account, in the present work we synthesized, crystallized, and characterized a new nucleus derived from the bridged diazocine system using the experimental techniques XRD, NMR, FT-IR, and HRMS. Likewise, confirmation of the stable crystal structure has been based on quantum chemistry results, such as geometry optimization, Hirshfeld surface analysis, and energy framework calculations of the frontier molecular orbitals. Motivated by the great potential of bridged azocine derivatives, we determined the antioxidant capacity of the compound using DPPH and ABTS assays.

2. Results and discussion

2.1. Chemistry

In the present work, we have synthesized compounds **4a** (62% yield) and **4b** (33% yield) through the implementation of reflux and ultrasound as energy sources, respectively (Scheme 2). The general mechanism proposed for the syntheses of **4a** and **4b** (Scheme 3)⁴⁴ involves the formation of a Knoevenagel product (step A) between thiobarbituric acid (**1**) and paraformaldehyde (**3**). Continuing this mechanism, the formation of an imine (step B) was observed because of the reaction between compound **2** and a molecule of paraformaldehyde (**3**). On the other hand, the imine formed reacts with the Knoevenagel product of step A *via* aza-Diels–Alder (ADA) reaction, generating the intermediate of step C, which due to proton transfer is transformed into compound **4a** (step D). Subsequently, the reaction continues, adding a molecule of ethanol to **4a** (step E), generating an esterification reaction that allows the formation of the intermediate of step F, which attacks a formaldehyde molecule through the nitrogen of the piperidine group, generating the addition of a methanol molecule (step G). Finally, cyclization occurs through dehydration, promoted by the attack of the carbamothioylacetamide nitrogen on the hydroxyl carbon, which leads to the formation of product **4b**.

2.2. FT-IR spectra

Fig. 2 shows the experimental and simulated infrared (IR) spectra obtained using the LC-BLYP/6-31G* method for **4a** and **4b**. On the other hand, in Table 1, the resulting vibrational

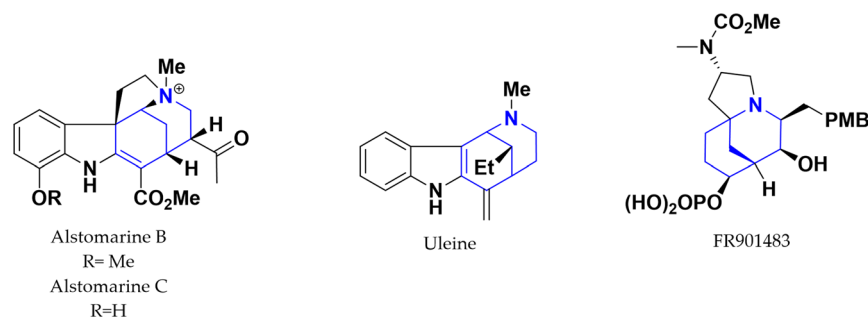
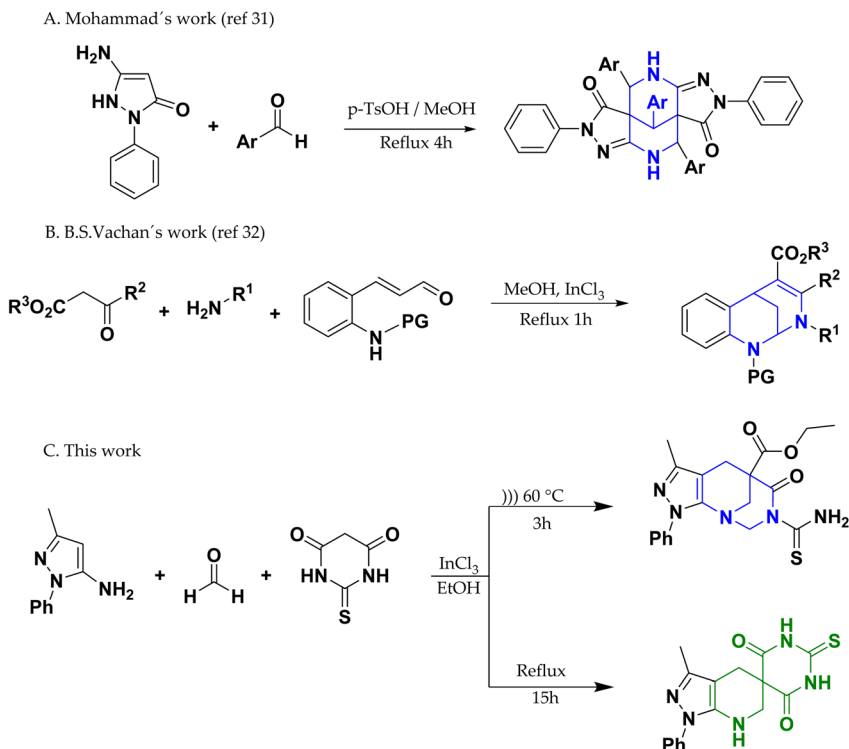


Fig. 1 The derivatives of azocine and azocane have interesting biological properties.



Scheme 1 Synthesis of bridged-azocine.

frequencies for the optimized geometries are shown from highest to lowest, along with the proposed vibrational assignments and IR intensities. Comparisons of the theoretical and experimental IR spectra indicated that the strong vibrations in the experimental spectrum were also strong in the theoretical spectrum.

The IR signals of **4a** and **4b** allowed the identification and characterization of their functional groups (see Table 1). Both present signals for C sp²-H, C sp³-H, and C=C. However, **4b** showed more signals owing to the presence of thioamide and ester groups.

In **4a**, a signal was observed for N-H (stretching) at 3465 cm⁻¹ and one for C=O (stretching) at 1689 cm⁻¹, and there was no signal for C-O. In contrast, **4b** presents two signals for N-H (asymmetric and symmetric stretching) at 3371 and 3234 cm⁻¹, two for C=O (ester and thioamide stretching) at 1724 and 1660 cm⁻¹, and one for C-O (ester) at 1257 cm⁻¹.

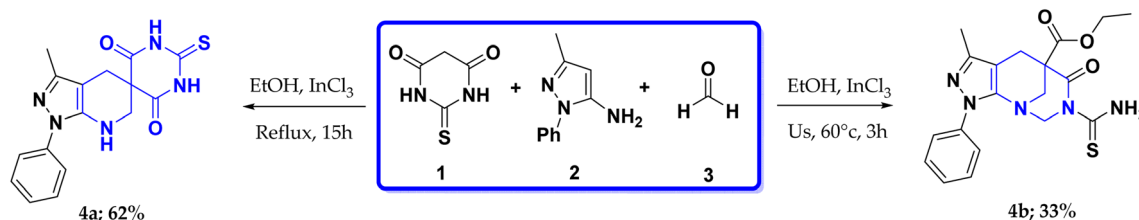
As shown in Table 1, the frequencies calculated using the LC-BLYP/6-31G* method were in good agreement with the

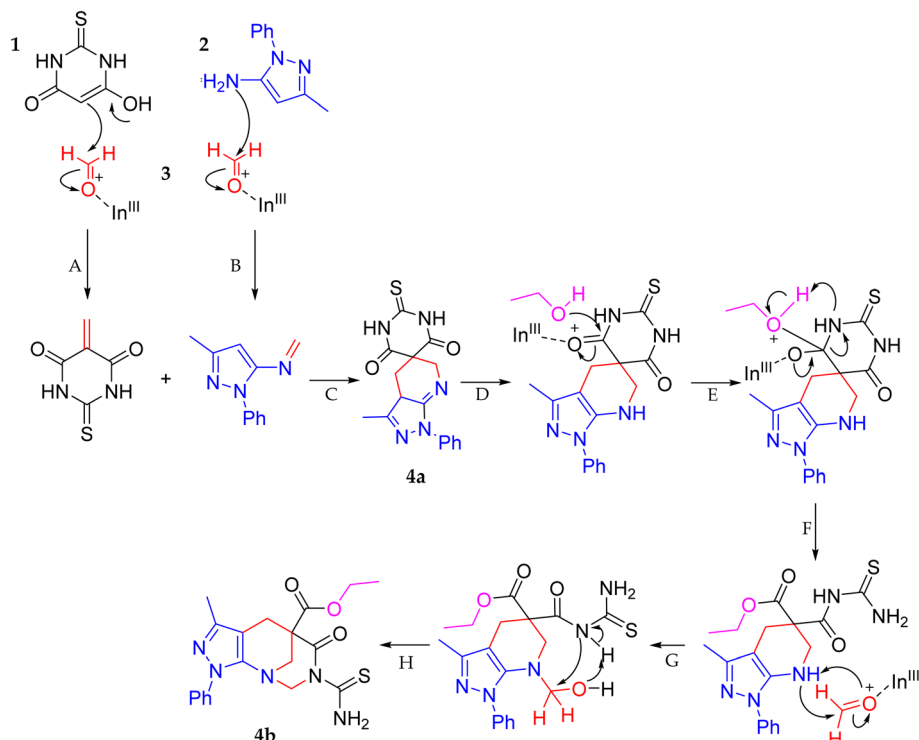
experimentally obtained results. Therefore, this computational method is valuable for identifying important functional groups to characterize the molecules studied here.⁴⁵

2.3. Frontier molecular orbital (FMOs) analysis and molecular reactivity

To obtain information on the reactivity and stability of compounds **4a** and **4b**, descriptors obtained from density functional theory were studied: HOMO and LUMO energies, HOMO-LUMO gap, electron affinity (EA), ionization energy (IE), electronegativity (χ), chemical hardness (η), chemical softness (s), and electrophilicity (w) (see Table 2 and Fig. 3). It was found that for molecule **4a** the values were -784.81, -23.10, 761.74, 83.08, 676.61, 379.85, 296.77, 0.0034 and 243.09, while, for molecule **4b**, the values were -797.83, 88.10, 885.92, 0.12, 756.41, 378.14, 0.0026 and 189.19 kJ mol⁻¹, respectively.

The HOMO energy defines the susceptibility of a compound to donate electrons and undergo electrophilic addition. In

Scheme 2 Conditions and synthesis of spiro-pyrazolopyridine (**4a**) and 1,3-diazocine (**4b**).



Scheme 3 Proposed mechanism for the synthesis of 4a and 4b.

contrast, the LUMO energy provides information about the susceptibility of a compound to accept electrons, undergoes nucleophilic attack, and is related to electronegativity (tendency to attract electron density) and electron affinity (ability to accept

electrons), whereas the HOMO–LUMO gap provides information about chemical reactivity and kinetic stability. A molecule with a high energy gap is associated with low chemical reactivity and high kinetic stability and *vice versa* (see Fig. 3). Finally,

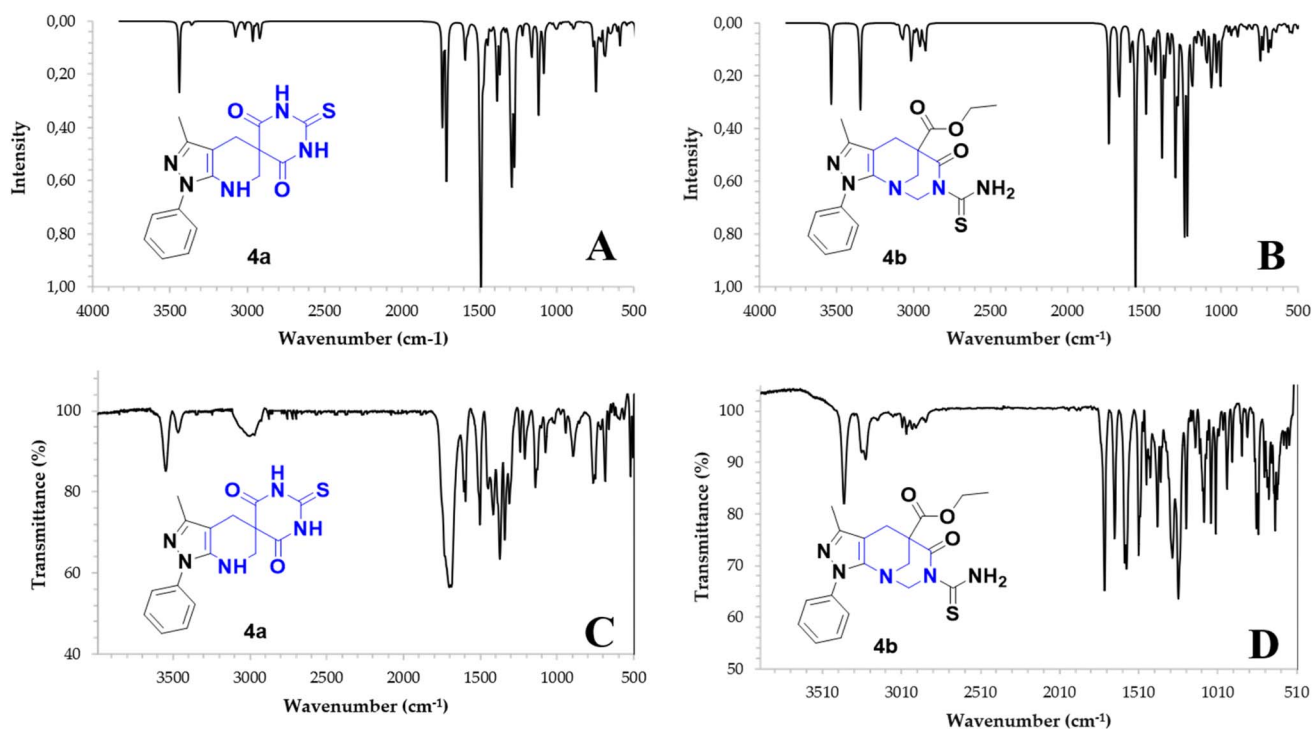


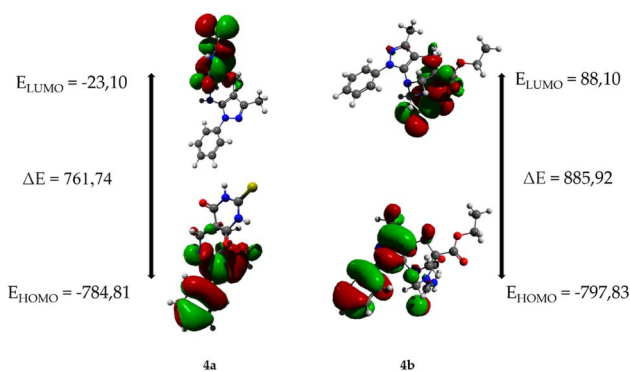
Fig. 2 IR spectra calculated using DFT (B3LYP/6-31G*) (A and B) and experimental (C and D) IR spectra for 4a and 4b.

Table 1 Comparison of calculated and experimental vibration spectra (FT-IR) of compounds **4a** and **4b**

	LC-BLYP/6-31G*	Experimental in this study		
Normal mode	Freq. (cm ⁻¹)	Intensity (km mol ⁻¹)	Freq. (cm ⁻¹)	Approximate assignments
4a				
1	3442	0.27	3465	Symmetric stretching N-H
2	3079	0.06	3074	Symmetric stretching C-H sp ²
3	2967	0.08	2972	Symmetric stretching C-H sp ³
4	2924	0.06	2941	Asymmetric stretching C-H sp ³
5	1714	0.60	1689	Stretch C=O × 2
6	1593	0.14	1597	Stretching C=C
7	1489	1.00	1506	Rocking amide N-H
4b				
1	3537	0.30	3371	Asymmetric stretching N-H ₂
2	3347	0.33	3234	Symmetric stretching N-H ₂
3	3071	0.06	3059	Symmetric stretching C-H sp ²
4	2984	0.14	2974	Symmetric stretching C-H sp ³
5	2924	0.10	2918	Asymmetric stretching C-H sp ³
7	1731	0.46	1724	Stretch C=O (ester)
8	1662	0.27	1660	Stretch C=O (amide)
9	1593	0.15	1583	Stretching C=C
10	1559	1.00	1508	Scissoring N-H ₂
11	1222	0.81	1257	Stretching C-O

Table 2 The electron affinity (EA), ionization energy (IE), electronegativity (χ), chemical hardness (η), softness (s), electrophilicity (w), electron acceptance index (R_a), and electron donation index (R_d) of compounds **4a** and **4b** were determined

	4a	4b
EA	83.08	0.12
IE	676.61	75.41
χ	379.85	378.27
η	296.77	378.14
s	0.0034	0.0026
w	243.09	189.19
R_a	0.49	0.26
R_d	1.47	1.33

**Fig. 3** The frontier molecular orbitals and related energies of **4a** and **4b** are expressed in kJ mol⁻¹.

chemical hardness and softness are related to the polarizability of a molecule; that is, the greater the hardness, the lower the polarizability, and the greater the softness, the greater the polarizability.

Fig. 3 shows the molecular frontier orbitals of compounds **4a** and **4b**. For the **4a** molecule, it is evident that the HOMO orbital shows a high electronic density located on the pyrazole and benzene rings, whereas in the case of the LUMO orbital, the highest probability region is found on the thiobarbituric acid ring, which agrees with the proposed mechanism because it is a zone with the capacity to accept electrons and therefore suffers from nucleophilic attack. On the other hand, for **4b**, the HOMO orbital shows electron density on the thioamide, while the LUMO orbital shows that the highest probability region is located both in the thioamide and the two carbonyl groups present in the molecule.

In general, the **4a** molecule has a smaller HOMO–LUMO gap, lower chemical hardness, greater chemical softness, greater electronegativity, greater electronic affinity, and lower ionization energy than the **4b** molecule, for which the **4a** molecule presents greater reactivity, less kinetic stability, greater polarizability, and greater ability to attract electron density, as well as to accept electrons.

2.4. Biological activity

We used two diverse assays to measure the *in vitro* antioxidant activity: (a) the interaction with the stable free radical DPPH and (b) the interaction with the water-soluble azo compound ABTS⁺, where ascorbic acid was used as a positive control. These radicals are not biologically relevant but are often used as indicator compounds to test the hydrogen transfer capacity, which is

related to antioxidant activity. Antioxidant properties were expressed as EC_{50} values (Table 4).

In the DPPH assay, the EC_{50} values of **4a** and **4b** were $<100 \mu\text{g mL}^{-1}$. Both compounds showed significant ABTS⁺ free radical scavenging activity (EC_{50} : 0.0967 and $1.6820 \mu\text{M}$) compared to the commercially available antioxidants ascorbic acid, quercetin, and Trolox ($EC_{50} > 2 \mu\text{M}$).

The radical scavenging activity of pharmacophore leads depends on electron-donating groups such as hydroxyl (–OH) and –NH groups, independent of their attachment, which can easily abstract free radicals and potentially convert highly reactive free radicals to their non-reactive forms.⁴⁷

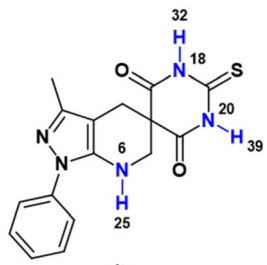
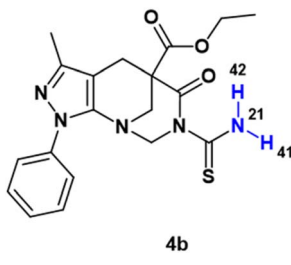
In general, it has been described that antioxidants can act through two mechanisms on radicals, the transfer of a hydrogen atom known as (HAT) and the transfer of a single electron (SET). However, both ABTS and DPPH radicals can be stabilized by hydrogen atom transfer (HAT) or electron transfer (ET) mechanisms, but the reactivity patterns and mechanism are difficult to interpret.⁴⁸

Thus, we calculated IE, EA, w (related to the SET spectra), and the hydrogen bond dissociation energy BDE (related to the HAT mechanism) at the DFT level to determine the possible mechanism of action of the compounds **4a** and **4b**. IE is the minimum energy required to extract an electron from a gaseous atom in its ground state; therefore, the lower the ionization energy of the antioxidant, the easier it is for it to donate an electron and neutralize a radical. On the other hand, EA is the energy released when a gaseous atom in its ground state accepts an electron to form a negative ion. The higher the EA of the antioxidant, the easier it is for it to accept an electron from a radical and neutralize it. W measures the ability of a molecule to attract electrons, thereby allowing it to neutralize them. Finally, BDE is the energy needed to break a hydrogen bond, and the lower it is, the greater is the antioxidant potential of the compound.

In addition, we drew a donor–acceptor map, DAM, as proposed by Martínez.^{49,50} This map, normalized with F as a good electron acceptor and Na as a good electron donor, is a useful graphical indicator. Therefore, this comparison was performed using computational values for the F and Na atoms at the same theoretical level for the molecules studied.

If $R_a = 1$, L is a compound with an electron acceptor efficiency like F. If $R_a > 1$, L is a more effective electron acceptor

Table 3 BDE of compounds **4a** and **4b**

 4a		 4b	
Entry	BDE (kJ mol ^{−1})	Entry	BDE (kJ mol ^{−1})
N ₆ –H ₂₅	228.01	N ₂₁ –H ₄₁	294.80
N ₁₈ –H ₃₂	220.26	N ₂₁ –H ₄₂	269.73
N ₂₀ –H ₃₉	220.17		

than F. Finally, if $R_a < 1$, L is a less effective electron acceptor than F. Similarly, if $R_d = 1$, L is an electron donor with an efficiency similar to that of Na. If $R_d > 1$, then L is a less efficient electron donor than Na. Similarly, if $R_d < 1$, L is a more efficient electron donor than Na is. Plotting R_a and R_d is the correct way to visualize the antioxidant scheme using MAP (Fig. 4).

Compound **4a** has three sp-hybridized –NH protons, which can be transferred to free radicals (e.g., the N-centered hydra- zyl free radical of ABTS⁺) by hydrogen atom transfer (HAT), with BDE of 228.01, 220.26 and 220.17 kJ mol^{−1} for the hydrogens H25, H32 and H39 respectively, while molecule **4b** has a –NH₂ group, with BDE of 294.80 and 269.73 kJ mol^{−1} for the hydrogens H41 and H42, respectively (see Table 3). These results suggest, on the one hand, that the most labile hydrogen in molecule **4a** is 39, while the most labile hydrogen in molecule **4b** is hydrogen 42, so these may be the main responsible for a possible neutralization of the radicals studied by HAT. Likewise, a comparison of the values of the most labile hydrogens in each molecule (H39 for **4a** and H42 for **4b**) shows that molecule **4b** has a higher BDE of 49.56 kJ mol^{−1} with respect to **4a**, which agrees with the experimental antioxidant results obtained, where **4a** presents a greater antioxidant capacity.

In contrast, the IE, EA, and w results indicate that **4a** has a lower ionization potential, greater electronic affinity, and

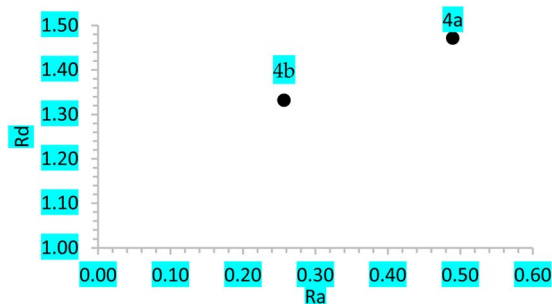


Fig. 4 Donor–acceptor map of the molecules studied in the gas phase.

Table 4 Antioxidant activity results for the **4a** and **4b** compounds, DPPH and ABTS assays

Compound	Antioxidant assay EC_{50}^a (μM)	
	DPPH assay	ABTS ⁺ assay ^a
4a	>292.921	0.0967 ± 0.001
4b	>294.577	1.6820 ± 0.020
Ascorbic acid ^b	8.517 ± 0.2	156.824 ± 3.5
Quercetin ⁴⁶	8.688 ± 0.02	15.487 ± 0.04
Trolox ⁴⁷	—	11.798 ± 0.05

^a Values are expressed as mean $\pm$ SEM of three parallel measurements ($p < 0.05$). ^b Reference compound.

greater electrophilicity than **4b**. Furthermore, compounds **4a** and **4b** presented values of 0.49 and 0.26 for R_a and 1.47 and 1.33 for R_d , respectively. All of the above results allow us to infer that **4a** has greater availability to accept and donate electrons than **4b**, so the latter has less potential as an antioxidant. Likewise, it is important to mention that, in general, the values obtained for BDE compared to those for ionization potential are more than 400 kJ mol⁻¹, which is lower for both molecules, so it is possible that both act mainly by the HAT mechanism.

2.5. Crystallographic studies

The molecular structure of compound **4b** corresponded to ethyl 7-carbamothioyl-3-methyl-6-oxo-1-phenyl-1,2,3,4,7,8-hexahydro-5,9-methanopyrazolo[3,4-*b*]azocine 5(6*H*)-carboxylate. This compound crystallizes in a monoclinic cell system with space group $P2_1/c$ ($Z = 4$). The molecular structure of the compound agrees with spectroscopic characterization and the proposed structure, and both show a centrosymmetric setting with normal bond distances and angles⁵¹ (see Fig. 5). The dihedral angle between the mean planes of the phenyl ring and the 5-membered heterocycle is 19.49(9)°, which is almost coplanar. Additionally, the two fused rings exhibited a fold angle of 91.01(7)°. According to the Cremer & Pople parameters,⁵² the

two fused rings in the title compound have a half-chair conformation, ($Q_T = 0.541(2)/0.558(2)$ Å; $\theta = 5.0(3)/51.3(2)^\circ$; $\phi = 331.8(3)/277.1(3)^\circ$).

All the hydrogen atoms were placed at their calculated positions, assigned fixed isotropic thermal parameters, and constrained to ride on their parent atoms. All geometrical calculations were performed using Platon software.⁵³ A summary of the crystal data, the collection of which is presented in Table 5, and additional crystallographic details are provided in the CIF file. ORTEP views were drawn using OLEX2 software.⁵⁴

Additionally, the crystal packing of the title compound does not present geometrical parameters corresponding to classical hydrogen bonding⁵⁵ and is stabilized by intra- and intermolecular non-conventional hydrogen bond-like interactions, N–H...X (X = O and S) and C–H...O. In the title compound, two intramolecular hydrogen bonds can be described using the graph-set motif $S(6)$. Likewise, intermolecular hydrogen bond interactions generate $R_4^3(13)$ and $R_4^4(22)$ ring motifs along the [100] direction, as a consequence of the 2_1 screw axis. Additionally, the $C_1^1(4)$ and $C_1^1(8)$ graph set motifs (symmetry operations = $+X, 1/2 - Y, -1/2 + Z$, and $1 - X, 1 - Y, 1 - Z$, respectively).⁵⁶ Form extended chains running along the [001] and [100] directions (see Fig. 6 and Table 6).

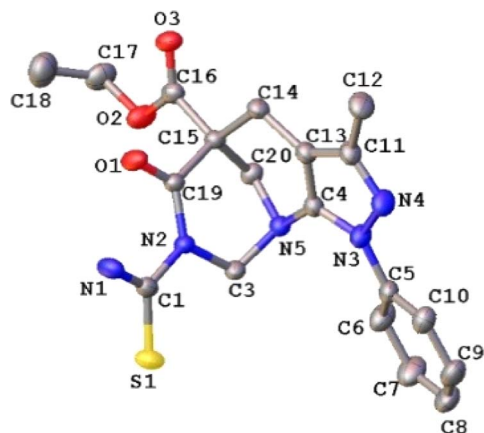


Fig. 5 ORTEP plot of the **4b**. Thermal ellipsoids were drawn at a 30% of probability. The H atoms were omitted for simplicity.

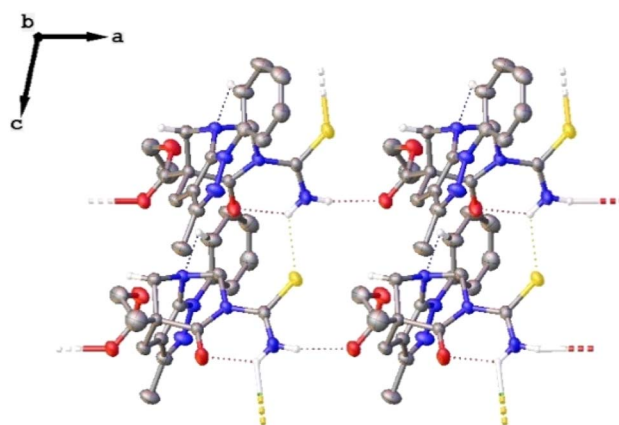


Fig. 6 Crystal packing of **4b**.

Table 5 Crystallographic data and refinement details of the title compound

Empirical formula	C ₂₀ H ₂₂ N ₄ O ₃ S	$F(000)$	840.0
Formula weight	398.47	Crystal size/mm ³	0.19 × 0.17 × 0.02
Temperature/K	297.75	2θ range	7.086 to 56.76
Crystal system	Monoclinic	Index ranges h, k, l	−11/11, −26/27, −14 ≤ l ≤ 14
Space group	$P2_1/c$	Reflections collected	22 466
$a/\text{\AA}$	8.6243(6)	Independent reflections	4675 [$R_{\text{int}} = 0.0372$, $R_{\text{sigma}} = 0.0265$]
$b/\text{\AA}$	20.6113(13)	Comp. q_{max} (%)	99.5
$c/\text{\AA}$	10.7734(7)	Max/min transmission	0.746, 0.669
$\beta/^\circ$	102.261(2)	Data/restraints/parameters	4675/0/255
Volume/Å ³	1871.4(2)	Goodness-of-fit on F^2	1.057
Z	4	Final R indexes [$I \geq 2\sigma(I)$]	$R_1 = 0.0542$, $wR_2 = 0.1572$
ρ_{calc} g cm ^{−3}	1.414	Final R indexes [all data]	$R_1 = 0.0671$, $wR_2 = 0.1689$
μ/mm^{-1}	0.203	CCDC number	2200261

Table 6 Hydrogen bonds in 4b

D	H	A	$d(\text{D-H})/\text{\AA}$	$d(\text{H-A})/\text{\AA}$	$d(\text{D-A})/\text{\AA}$	D-H-A/ $^\circ$
N1	H1A	S1 ^a	0.86	2.87	3.435(2)	125.4
N1	H1A	O1	0.86	1.96	2.590(2)	129.2
N1	H1B	O3 ^b	0.86	2.09	2.939(2)	168.0
C6	H6	N2 ^c	0.93	2.44	3.055(3)	123.2

^a +X, 1/2 - Y, 1/2 + Z. ^b 1 + X, +Y, +Z. ^c +X, 1/2 - Y, -1/2 + Z.

2.6. Hirshfeld surface analysis

To observe other intermolecular contacts across the crystal structure, Hirshfeld surface analysis was performed to complement XRD analysis. The intermolecular interactions were represented by N-H...X (X = O and S). The contacts are shown as red ($d_{\text{norm}} < \text{vdW radii}$), white ($d_{\text{norm}} = \text{vdW radii}$), and blue ($d_{\text{norm}} > \text{vdW radius}$) spots on the d_{norm} surfaces for all compounds. Moreover, there is evidence of another interesting weak contact in the crystal structures of all compounds. The reciprocal contacts, their respective contributions, and all fingerprint plots with d_{norm} (where $d_{\text{norm}} = d_i + d_e$) surfaces for their intermolecular contacts are depicted in Fig. 7. Additionally, for H...H contacts in each compound, does not generate a significant effect on molecular packing in the crystal structure stabilization because their contacts are $d_i + d_e > 2.4 \text{ \AA}$, in other words, these contacts are slightly longer than the sum of the vdW radii for these atoms,⁵⁷ which cannot support the crystal packing in the title compound.

4b shows the same interaction as that described above, where hydrogen-bond-like interactions are observed on the d_{norm} surface. Another type of weak interaction was observed when the Hirshfeld surface was analyzed. For example, aromatic π contact was verified over the title compound (Fig. SI-5†) with a shape-index surface. This was verified using the shape index surface, which allowed us to determine the presence of these weak interactions. The yellow-orange spots show

surface subsidence owing to the proximity of the neighboring moieties, and the blue-green spots show the reciprocal contacts of the moieties that generate the subsidence. The contact is between the ethyl and phenyl fragments in the title compound, with a contribution of approximately 16.8% and a $d_e + d_i$ of $\approx 3.0 \text{ \AA}$, as a clamp pattern in the fingerprint plot.

A C...S chalcogen bond interaction is also observed, and this was verified over the carbonyl and carbamothioyl fragments (Fig. SI-6†) with a contribution of around 0.6% with $d_e + d_i \approx 3.4 \text{ \AA}$, which works as electron donor in a σ -hole noncovalent bond.^{58,59}

Finally, the energy framework⁶⁰ was analyzed to better understand the packing and topology of the crystal structure and supramolecular rearrangement. This method allows the calculation and comparison of different energy components, that is, repulsion (E_{rep}), electrostatic (E_{ele}), dispersion (E_{dis}), polarization (E_{pol}), and total (E_{tot}) energies, based on the anisotropy of the topology of pairwise intermolecular interaction energies (see Fig. SI-7 and Table SI-1†). The thickness of the cylinder radius shows the grade of interactions, is related to the energy magnitude, and provides information regarding the stabilization of the crystal packing.⁶¹ Based on the tube direction, it can be concluded that the formation of the framework is directed by the translational or centrosymmetric elements. However, this rearrangement allowed the formation of weak interactions in the crystal structure.

The results of the calculations revealed that the dispersion interactions exhibited a distorted zigzag ladder-shaped topology in the title compound. The significant difference between E_{ele} and E_{pol} is due to the absence of classical hydrogen bond interactions.

3. Experimental section

3.1. General information

Ultrasound irradiation was performed using a Branson model 1510, 115v, 1.9 L ultrasonic bath, time machine (continuous

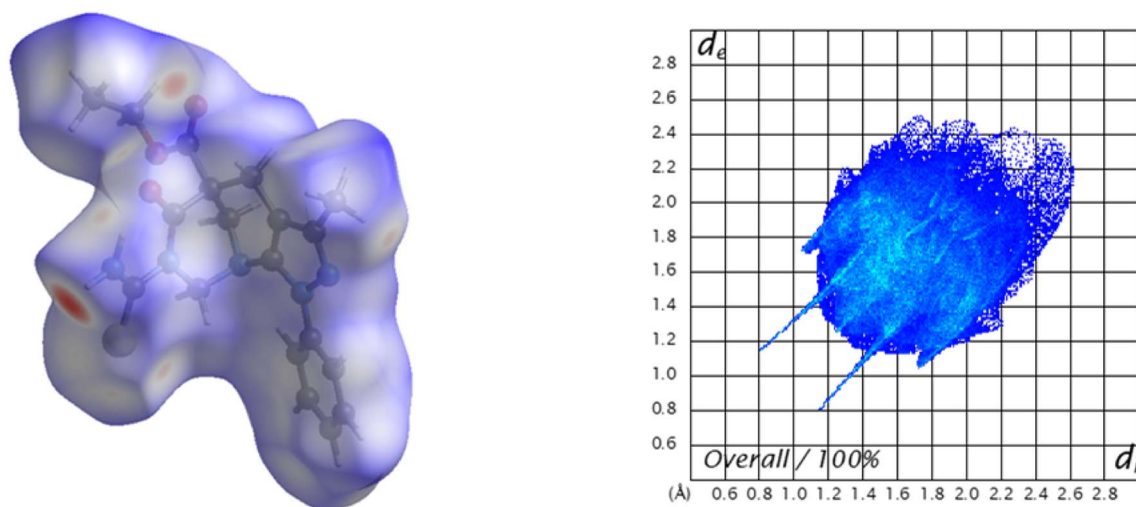


Fig. 7 Overall, the Hirshfeld surface (left) of 4b and the fingerprint plot (right) are shown.

hold for 60 min), and switch heater (47 kHz). ^1H and ^{13}C NMR spectra (400 MHz for protons and 100 MHz for carbon) were recorded on an AM-400 spectrometer (Bruker, Rheinstetten, Germany) using CDCl_3 and $\text{DMSO}-d_6$ as the solvents. Tetramethylsilane (TMS) was used as an internal standard. IR spectra (KBr granules, $500\text{--}4000\text{ cm}^{-1}$) were recorded on a NEXUS 670 FT-IR spectrophotometer (Thermo Nicolet, Madison, WI, USA). ESI-MS and ESI-MS/MS high-resolution mass spectrometry were performed on a high-resolution (Q) hybrid quadrupole time of flight (TOF) mass spectrometer (Waters/Micromass Q-TOF Micro, Manchester, UK) with a constant nebulizer temperature of $100\text{ }^\circ\text{C}$. Melting points (uncorrected) were measured using an IA9100 electrothermal melting point device (Stone, Staff, UK). The progress of the reaction was evaluated by thin-layer chromatography (TLC) using silica gel 60 (Merck, Darmstadt, Germany). All the reagents and solvents were purchased from Merck or Sigma-Aldrich (St. Louis, MO, USA) and used without further purification. The final purification of all the products for the analysis was conducted by column chromatography using silica gel 60 (Merck 7734) with a particle size that fluctuated between 0.063 and 0.200 mm, as eluent phase ether mixtures of petroleum: ethyl acetate in increasing polarity.

3.2. Chemistry

3.2.1 Synthesis of 3-methyl-1-phenyl-2'-thioxo-1,2',3',4,6,7-hexahydro-4'H-spiro[pyrazolo[3,4-b] pyridine-5,5'-pyrimidine]-4',6'(1'H)-dione (4a). A mixture of AMPZ (1.1 mmol), *p*-formaldehyde (excess), thiobarbituric acid (1.1 mmol), InCl_3 (15 mol%), in ethanol is warmed into reflux for 15 h and then treated with brine solution. The biphasic solution was vigorously stirred for 30 min, decanted, and then separated. The collected aqueous phase was extracted using ethyl acetate (EtOAc , $2 \times 10\text{ mL}$). The combined layers were dried over Na_2SO_4 and evaporated under a reduced pressure. The reaction crude was purified by column chromatography using mixtures of petroleum ether: ethyl acetate in increasing polarity as eluant phase to afford the compound target **4a** in 62% yield as a white solid, IR (KBr, cm^{-1}): 3465, 3074, 2972, 2941, 1689, 1597, 1506; ^1H NMR (400 MHz, $\text{DMSO}-d_6$) 2.07 (s, 3H), 2.87 (s, 2H), 3.36 (s, 2H), 5.90 (t, $J = 6.5\text{ Hz}$, 1H), 7.19 (t, $J = 7.3\text{ Hz}$, 1H), 7.40 (t, $J = 8.0\text{ Hz}$, 2H), 7.67 (t, $J = 8.0\text{ Hz}$, 2H), 11.18 (s, 2H); ^{13}C NMR (100 MHz, $\text{DMSO}-d_6$) 12.6 (CH_3), 23.8 (CH_2), 47.8 (C), 51.2 (CH_2), 100.2 (C), 120.5 ($2 \times \text{CH}$), 125.2 (CH), 129.4 ($2 \times \text{CH}$), 139.9 (C), 142.8 (C), 146.0 (C), 150.9 (C), 171.7 ($2 \times \text{C}$); HRMS (ESI, m/z): calculated for $\text{C}_{16}\text{H}_{17}\text{N}_5\text{O}_2\text{S} [\text{M} + 2\text{H}]^+$ 343.1103 found 343.8190.

3.2.2 Synthesis of ethyl 7-carbamothioyl-3-methyl-6-oxo-1-phenyl-1,4,7,8-tetrahydro-5,9-methanopyrazolo[3,4-*d*][1,3]diazocine-5(6*H*)-carboxylate (4b). A mixture of AMPZ (1.1 mmol), *p*-formaldehyde (2.1 mmol), thiobarbituric acid (1.1 mmol), InCl_3 (15 mol%), in ethanol as solvent in an Erlenmeyer flask, was placed in a water bath and sonicated at $60\text{ }^\circ\text{C}$ for 3 h. The reaction mixture was poured into water and extracted with ethyl acetate ($2 \times 10\text{ mL}$). The combined organic layers were dried over Na_2SO_4 . The reaction crude was purified by column chromatography using mixtures of petroleum ether: ethyl acetate in increasing polarity as eluant phase, producing compound (**4b**)

in 33% yield as a green crystal; IR (KBr, cm^{-1}): 3371, 3234, 3059, 2974, 2918, 1724, 1660, 1583, 1508, 1257; ^1H NMR (400 MHz, CDCl_3) 0.99 (t, $J = 7.1\text{ Hz}$, 3H), 2.21 (s, 3H), 3.20 (q, $J = 7.1\text{ Hz}$, 2H), 3.61 (s, 2H) 4.08 (s, 2H), 4.36 (s, 2H), 7.28 (t, $J = 7.5\text{ Hz}$, 1H), 7.40 (t, $J = 7.8\text{ Hz}$, 2H), 7.60 (t, $J = 7.8\text{ Hz}$, 2H); ^{13}C NMR (100 MHz, CDCl_3) 12.1 (CH_3), 14.8 (CH_3), 22.1 (CH_2), 49.5 (CH_2), 64.1 (CH_2), 81.4 (CH_2), 102.6 (C), 124.1 ($2 \times \text{CH}$), 127.2 (CH), 129.1 ($2 \times \text{CH}$), 139.2 (C), 142.4 (C), 146.4 (C); HRMS (ESI, m/z): calculated for $\text{C}_{17}\text{H}_{18}\text{N}_5\text{O}_3\text{S} [\text{M}-\text{C}_2\text{H}_4 + \text{H}]^+$ 372.1125 found 372.9570 (McLafferty rearrangement).

3.3. Biological

3.3.1 Measurement of DPPH radical scavenging activity.

The newly synthesized compounds **4a–b** were evaluated for their $\text{DPPH}^{\cdot+}$ free radical scavenging activity. According to the method previously described and adapted,⁶² a 1 mL aliquot of the tested compound ($10\text{--}100\text{ }\mu\text{g mL}^{-1}$) and the control (2% final DMSO) were taken respectively and mixed with 2 mL of a methanolic $\text{DPPH}^{\cdot+}$ solution (0.02 mg mL^{-1}). The mixture was vigorously stirred and allowed to stand at room temperature for 5 min in the dark. The absorbance of the samples was determined using a spectrophotometer at a wavelength of 517 nm. Free radical scavenging activity was calculated as the percentage of $\text{DPPH}^{\cdot+}$ discoloration using the following equation:

$$\text{Percentage of free radical removal } \text{DPPH}^{\cdot+} = 100 \times (1 - A_E/A_D), \quad (1)$$

where A_E is the absorbance of the solution after the addition of the sample, and A_D is the absorbance of the blank DPPH solution. Ascorbic acid was used as the reference compound, with an EC_{50} value of $1.5\text{ }\mu\text{g mL}^{-1}$.

3.3.2 Measurement of ABTS radical scavenging activity.

The newly synthesized compounds **4a–b** were evaluated for the radical scavenging activity of $\text{ABTS}^{\cdot+}$, according to a published test.⁶³ The stock solution was prepared by mixing equal volumes of 7 mM ABTS and 2.45 mM potassium persulfate, followed by incubation for 12 h at room temperature in the dark to produce a dark-colored solution containing $\text{ABTS}^{\cdot+}$ radicals. The working solution was prepared once it was needed in the assay to avoid oxidation, and 50% ethanol was added to give an initial absorbance of approximately 0.700 (± 0.02) at 732 nm at room temperature. Free radical scavenging activity was evaluated by mixing 300 μL of different compounds ($10\text{--}200\text{ }\mu\text{g mL}^{-1}$ in the respective solvents) with 3.0 mL of $\text{ABTS}^{\cdot+}$ working standard. The decrease in absorbance was measured exactly 1 min after mixing the solutions and the final absorbance was observed after 6 min. Ascorbic acid, with an EC_{50} value of $28\text{ }\mu\text{g mL}^{-1}$, was used as the positive control. The elimination activity was estimated based on the percentage of $\text{ABTS}^{\cdot+}$ radicals eliminated using the following equation:

$$\text{Percentage of free radical removal } \text{ABTS}^{\cdot+} = [(A_0 - A_s)/A_0] \times 100 \quad (2)$$

where A_0 is the absorption of the control and A_s is that of the test compound solution.

3.4. X-ray diffraction measurements

Diffraction data were collected at 297 K on a Bruker D8 Venture diffractometer equipped with a bi-dimensional CMOS Photon 100 detector using graphite monochromated Mo K α ($\lambda = 0.71073$ Å) radiation. The diffraction frames were integrated using the APEX3 package⁶⁴ and corrected for absorption SADABS.⁶⁵ The structure of **4b** was solved by intrinsic phasing⁶⁶ using OLEX2 software⁶⁴ and refined with full-matrix least-squares methods based on F^2 (SHELXL).⁶⁷ Non-hydrogen atoms were refined using anisotropic displacement parameters.

3.5. Computational details

We employed Orca software for optimization and frequency calculations, and the Gaussian 09 software package was used to determine the Wiberg bond indices.^{68,69} The first molecular geometries were generated using the Avogadro molecular editor⁷⁰ to create the corresponding input file. All molecular structures were optimized using LC-BLYP with the 6-31G(d) basis set applied uniformly to all the investigated chemical species (including non-radicals, anions, and cations). The ionization energy (IE) and electron affinity (EA) were calculated adiabatically,⁷¹ where $IE = E_{\text{cation}} - E_{\text{neutral}}$ and $EA = E_{\text{neutral}} - E_{\text{anion}}$. The calculation of bond dissociation energy (BDE) followed the method outlined by Rakiba Rohman *et al.*, resulting in BDE as a product of η and the Wiberg bond index (BIWiberg).⁷²

The determination of electrophilicity involves consideration of electronegativity and hardness. In a system forming N electrons subjected to an external potential $v(r)$ and having a total energy E , the electronegativity is defined as the partial derivative of the energy concerning the number of electrons at a constant potential. In a finite-difference approximation, this corresponds to half the sum of ionization energy (IE) and electron affinity (EA):⁷³

$$x = \left(\frac{\partial E}{\partial N} \right)_{v(r)} \approx \frac{IE + EA}{2}$$

Chemical hardness was calculated according to the definition proposed by Parr and Pearson, involving the differentiation of the chemical potential with respect to the number of electrons at a constant energy potential. The latter can be approximated as half the difference between ionization energy (IE) and electron affinity (EA).⁷⁴ To achieve symmetry with and to ensure electronegativity symmetry, the product is multiplied by half, as highlighted by Pearson:⁷⁵

$$\eta = \left(\frac{\partial^2 E}{\partial^2 N} \right)_{v(r)} \approx \frac{IE - EA}{2}$$

The electrophilicity index is calculated as follows:

$$\omega = \frac{x^2}{2\eta}$$

The ionization energy (IE) and electron affinity (EA) can be used to compute the electron donor (ω^-) and electron acceptor

(ω^+) indices, as suggested by Gázquez and colleagues.⁷⁶ Electron-donating power gauges the tendency to donate charge and is defined by the following equation:

$$\omega^- \approx \frac{(3IE + EA)^2}{16(IE - EA)}$$

The definition of the electron-accepting power or inclination to accept electrons (ω^+) is as follows:

$$\omega^+ \approx \frac{(IE + 3EA)^2}{16(IE - EA)}$$

Reduced ω^- values signify an increased ability to donate charge, whereas elevated ω^+ values indicate a heightened ability to accept charge.

For any given compound L, the electron acceptance index is defined as follows:

$$R_a = \frac{\omega^+ L}{\omega^+ F}$$

Similarly, the electron donation index is defined as follows:

$$R_d = \frac{\omega^- L}{\omega^- Na}$$

The graph has four central regions; the worst antiradical activity was found within the zone with poor donor and acceptor capacities. Two regions with good antiradical behavior correspond to bad acceptors, but good donors, and suitable acceptors, but bad donors. Finally, the best antiradical zone represents both a suitable acceptor and donor (see Fig. 8).

3.6. Hirshfeld surface analysis details

CrystalExplorer 21.3 software⁷⁷ was used to calculate the Hirshfeld⁷⁸ and associated 2D-fingerprint plots⁷⁹ of **4b** using the

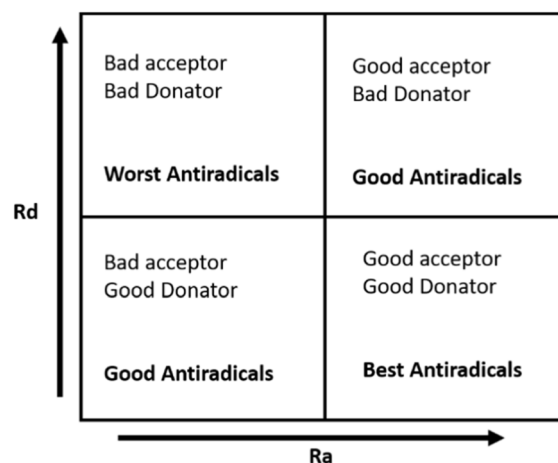


Fig. 8 Donor-acceptor map (DAM) diagram. These four regions were distinguished and described in detail by Martinez *et al.* The dashed lines separating the regions indicate only image clarification.

crystallographic information file (CIF) as input for the analysis. The normalized contact distance d_{norm} , defined in terms of the d_e , d_i , and vdW radii of the atoms, was calculated using eqn (3), where the distance from the Hirshfeld isosurface to the nearest nucleus is external (d_e) or internal (d_i), and vdW is the van der Waals radii of atoms taken from the literature.^{80,81}

$$d_{\text{norm}} = \frac{d_i + r_i^{\text{vdW}}}{r_i^{\text{vdW}}} + \frac{d_e + r_e^{\text{vdW}}}{r_e^{\text{vdW}}} \quad (3)$$

Hirshfeld surface analysis was performed using the 6-31G(d,p) basis set at the B3LYP level of theory over the range of $\pm 0.002 \text{ au}^{78}$ using the TONTO computational package integrated into the CrystalExplorer program.⁸² The bond lengths of the hydrogen atoms involved in the interactions were normalized to standard values obtained from neutron diffraction measurements (C–H = 1.083 Å, N–H = 1.009 Å, and O–H = 0.983 Å).⁸³ The intermolecular forces between pairs of molecules in the crystal packing were calculated at the B3LYP/6-31G(d,p) level of theory in clusters with a 3.8 Å radius around the molecules.^{84,85}

4. Conclusions

In this study, we synthesized a new spiro derivative (**4a**) and a new azocine derivative (**4b**) from a mixture of 5-amino-pyrazole, *p*-formaldehyde, thiobarbituric acid, and InCl_3 using reflux and ultrasound, respectively. Compound **4a** was characterized by means of spectroscopic studies: FT-IR, HRMS, ^1H NMR and ^{13}C NMR; while for compound **4b** the spectroscopic studies were used: FT-IR, HRMS and X-ray crystallography. The X-ray findings showed that **4b** has two fused rings in a half-chair conformation and crystallizes in the monoclinic system with a $P2_1/c$ space group, $Z = 4$, and unit cell parameters $a = 8.624(6) \text{ Å}$, $b = 20.611(7) \text{ Å}$, $c = 10.773(7) \text{ Å}$, $\beta = 102.261(14)^\circ$, and $V = 1871.4(11) \text{ Å}^3$. In general, good agreement was found between all the investigated theoretical properties (structural, electronic, and spectroscopic) and the experimental results. Analysis of the FMOs and chemical reactivity descriptors revealed that **4a** was more reactive and less stable than **4b** was. Hirshfeld surface analysis of the crystal structure indicated that the greatest contribution to stabilization was due to unconventional intermolecular interactions, similar to hydrogen bonds $\text{N-H}\cdots\text{X}$ ($\text{X} = \text{O}$ and S) and $\text{C-H}\cdots\text{O}$. Likewise, compounds **4a** and **4b** presented good antioxidant activities in the ABTS assay, with EC_{50} of 0.033 and $0.571 \mu\text{g mL}^{-1}$ respectively, being better than the positive controls, therefore, in general, both compounds have interesting antioxidant activities that give them good therapeutic potential. Finally, based on the DFT results, it was possible to show that compound **4a** had greater ease in accepting and donating electrons and lower hydrogen bond dissociation energies (BDE) than compound **4b**, which justifies its improved antioxidant capacity. Furthermore, when comparing the ionization energies (IE) of both compounds with their respective BDEs, it is evident that both **4a** and **4b** stabilized the radicals studied *via* the hydrogen atom transfer (HAT) mechanism.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare that they have no conflicts of interest.

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