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New 1*H*-benzimidazole-2-yl hydrazones with combined antiparasitic and antioxidant activity†

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Parasitic infections, caused mainly by the species *Trichinella spiralis* (*T. spiralis*), are widespread around the world and lead to morbidity and mortality in the population. Meanwhile, some studies have showed that these parasites induce oxidative stress in the infected host. With the aim of developing a class of compounds combining anthelmintic with antioxidant properties, a series of new benzimidazolyl-2-hydrazones **5a-l**, bearing hydroxyl- and methoxy-groups, were synthesized. The anthelmintic activity on encapsulated *T. spiralis* was studied *in vitro* thus indicating that all hydrazones were more active than the clinically used anthelmintic drugs albendazole and ivermectin. **5b** and **5d** killed the total parasitic larvae (100% effectiveness) after 24 hours incubation period at 37 °C in both concentrations (50 and 100 µg ml⁻¹). The antioxidant activity of the target compounds was elucidated *in vitro* against stable free radicals DPPH and ABTS as well as iron induced oxidative damage in model systems containing biologically relevant molecules lecithin and deoxyribose. The two 2,3- and 3,4-dihydroxy hydrazones **5b** and **5d** were the most effective radical scavengers in all studied systems. DFT calculations were applied to calculate the reaction enthalpies in polar and nonpolar medium and estimate the preferred mechanism of antioxidant activity. The relative radical scavenging ability of compounds **5a-l** showed a good correlation to the experimentally observed trends. It was found that the studied compounds are capable to react with various free radicals – ·OCH₃, ·OOH and ·OOCH₃, through several possible reaction pathways – HAT in nonpolar medium, SPLET in polar medium and RAF in both media.

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1. Introduction

Trichinellosis is a natural and synanthropic focal zoonanthroposis, which is caused by parasites of the genus *Trichinella*.¹ Human trichinellosis is usually associated with the species *Trichinella spiralis* (*T. spiralis*) which has a cosmopolitan distribution, but is most often found in the areas with temperate climate where swine are raised.²⁻⁴

The complex interactions between the *Trichinella* parasites and the host lead to different clinical manifestations of the disease, which can be grouped into two phases related to the life

cycle of the parasite – gastrointestinal (enteral) phase⁵ and muscular (parenteral) phase.⁶ Conventional treatment of the human trichinellosis is complex and includes the use of the benzimidazole anthelmintic drugs albendazole or mebendazole (Fig. 1) in combination with antipyretics, anti-inflammatory drugs and vitamins to control the temperature and intense inflammatory reactions accompanying the clinical picture of the disease.⁷

The tissue damages and immune system dysfunction which arise during trichinellosis as well as toxic manifestation caused from the massive destruction of parasites during the treatment with anthelmintic drugs can be in many cases conditioned by the oxidative stress state that accompanies this infection.⁸ Studies have shown that changes in the levels of some antioxidant enzymes such as glutathione peroxidase (GPx), superoxide dismutase (SOD), glutathione-S-transferase (GST) and catalase (CAT) in the host's muscle tissues during trichinellosis were observed.⁹⁻¹¹ Therefore, antioxidants are expected to help protecting the hosts against these injurious factors. Gabrashanska *et al.*⁸ examined the effect of dietary selenium in the form of organic compound Sel-Plex on the oxidative and antioxidant status, mortality, and body weight in rats experimentally infected with the nemataode *T. spiralis*. Their results confirmed

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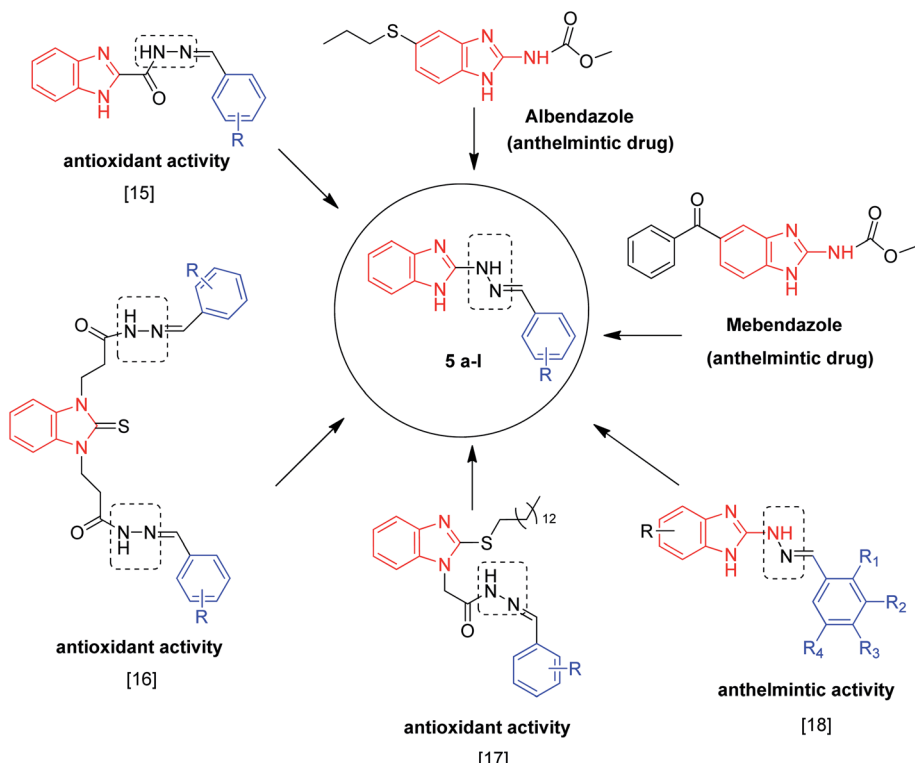


Fig. 1 Designed prototype scaffold based on antioxidant and anthelmintic active pharmacophores.

that selenium at low doses contributes to antioxidant response and improvement of the antioxidant status in host has been observed. Analogously, resveratrol, orally applied at a dose of 20 mg kg^{-1} once daily for two weeks, significantly improved the redox status of the small intestine and muscles during the intestinal and muscular phases of *T. spiralis* infection in mice.¹²

In view of the above, we present here the design and synthesis of new benzimidazole derivatives as potential anti-trichinella compounds that combine anthelmintic and antioxidant actions in one molecule. Thus, in addition to their ability to kill parasites, these biologically active substances may lead to an improvement of the antioxidant status in the host.

The benzimidazoles are a large chemical family widely used in veterinary and human medicine to treat helminthic infections¹³ and an appropriate structural platform for antioxidant discovery efforts.¹⁴ In examining the recent literature, certain fragments were observed in the structure of benzimidazole-based antioxidants – a benzimidazole core, phenolic hydroxyl and/or methoxyl groups and hydrazone moiety (Fig. 1).^{15–17}

On the other hand, in our previous paper,¹⁸ we reported the synthesis of a series of benzimidazolyl-2-hydrzones of fluoro-, hydroxy- and methoxy-substituted benzaldehydes and 1,3-benzodioxole-5-carbaldehyde (Fig. 1). The benzimidazoles showed potent anthelmintic activity against isolated muscle larvae of *T. spiralis* in an *in vitro* experiment as well as moderate antiproliferative activity against MCF-7 breast cancer cells.¹⁸ The antiparasitic and anticancer effects were correlated to the ability of the compounds to interfere with tubulin polymerization, as demonstrated by *in vitro* testing. As a next step in our

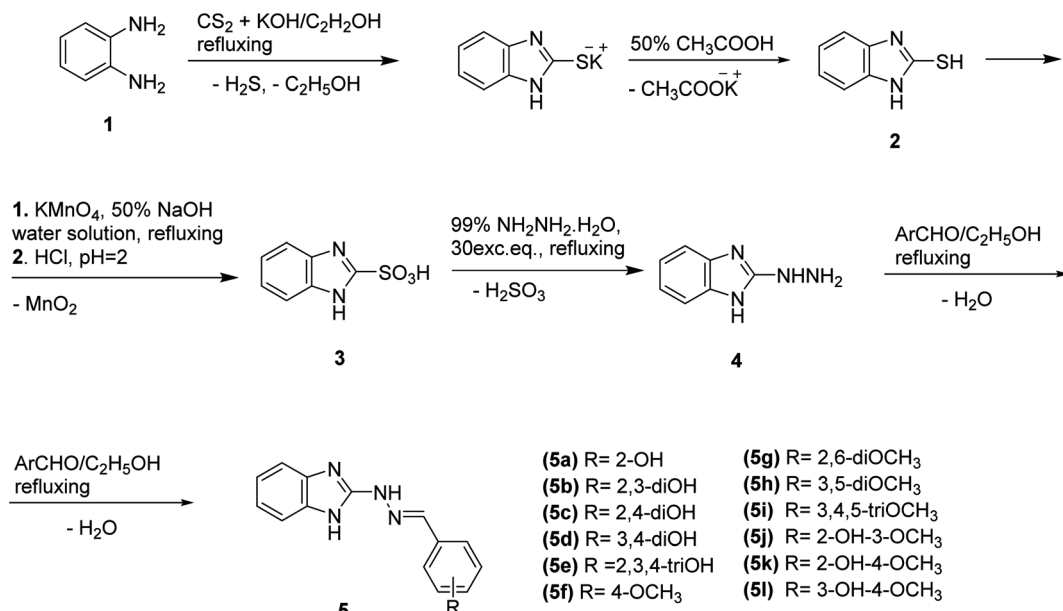
research¹⁸ and on the basis of the above mentioned facts, the present study reports the synthesis of a new series of 1*H*-benzimidazol-2-yl hydrazones containing hydroxyphenyl and methoxyphenyl moieties, their anthelmintic activity as well as antioxidant and radical scavenging properties determined using *in vitro* assays – DPPH, ABTS, and iron induced oxidative damage in model systems containing lecithin and deoxyribose. In order to rationalize the observed effects, a detailed computational study of the radical scavenging properties using density functional theory (DFT) was carried out. Thermodynamic parameters related to the main antioxidant mechanisms were computed. Feasibility of the reactions with $\cdot\text{OCH}_3$, $\cdot\text{OOH}$ and $\cdot\text{OOCH}_3$ were estimated by thermodynamic calculations as well. The polarity of the medium was accounted in the study by incorporation of water and benzene solvent.

2. Results and discussion

2.1. Chemistry

According to the literature benzimidazole-2-thiols could be synthesized thorough various methods such as reaction of benzene-1,2-diamine with carbon disulfide and potassium hydroxide in the presence of ethyl alcohol and water; with potassium dithiocarbamate (potassium ethyl xanthate),¹⁹ reaction of benzene-1,2-diamine with thiophosgene in chloroform²⁰ as well as refluxing solution with thiocyanate at 120–130 °C.²¹ In this study the new benzimidazole-2-yl hydrazones 5a–I were obtained according to the following reaction scheme (Scheme 1).





Scheme 1 Synthesis of $1H$ -benzimidazol-2-yl hydrazone derivatives 5a-l.

The synthesis of the $1H$ -benzimidazol-2-yl-thiol 2 was carried out by refluxing ethanol–water solution of potassium hydroxide, carbon disulphide and *o*-phenylenediamine according to the procedure described by us earlier.²² $1H$ -Benzimidazol-2-yl-sulfonic acid 3 was obtained by oxidation of $1H$ -benzimidazol-2-thiol in a 50% sodium hydroxide solution of potassium permanganate for 1 h. The filtrate was acidified with hydrochloric acid to pH = 1 and the resulting precipitate of the sulfonic acid was filtered off and washed with water.²³ The $1H$ -benzimidazol-2-yl-sulfonic acid 3 was converted in hydrazine 4 by refluxing it for 3 hours in excess of 99% hydrazine hydrate according to the described protocol.²⁴

The target compounds 5a-l were synthesized through reaction of condensation between $1H$ -benzimidazol-2-yl-hydrazine 4 and various hydroxyl- and methoxy-benzaldehydes in molar ratio 1 : 1 and using ethanol, 99%, as a solvent.

Identification of the $1H$ -benzimidazol-2-yl hydrazone was achieved by spectroscopic techniques FT-IR, ^1H NMR and ^{13}C NMR. The spectral data are given in the Experimental section.

The formation of the azomethine bond was identified by disappearance of the strong doublet for N–H stretching vibration, characteristic for the hydrazine 4, and appearance of a bands for the C=N stretching in the region 1600–1620 cm^{-1} of the IR spectra of the $1H$ -benzimidazol-2-yl hydrazone products. The N–H bonds in the benzimidazolyl fragment and hydrazone moiety gave rise to strong peaks varying within the region 3370–3160 cm^{-1} . The C–O stretching vibrations of the methoxyl groups, were found as intense broad bands around 1265 and 1135 cm^{-1} . In the IR spectra of the compounds containing hydroxyl groups, another band for C–O stretching was observed around 1200 cm^{-1} , while the bands of the corresponding O–H stretching vibrations appeared from 3332 to 3468 cm^{-1} .

In the ^1H -NMR spectra of 5a-l the signals for the protons of the azomethine group were found at 7.9–8.0 ppm as singlets. The benzimidazole protons produced multiplet signals within the interval 6.9–7.3 ppm in DMSO solvent, while the phenyl protons resonated in the interval 6.9–7.8 ppm. The chemical shift values for the protons for the labile NH and OH protons varied in a broader range – 11.26–11.55 and 9.87–10.95 ppm.

Generation of nitrosamine impurities during the synthetic process, degradation or biotransformation is considered a severe toxicity issue due to the mutagenic and carcinogenic properties of nitrosamines.^{25–30} In the past few years, nitrosamine impurities (above the maximum acceptable limit that is in the range from 26.5 to 96 ng per day) were detected in various drugs and lead even to recalls from the market of several sartans used as angiotensin II receptor blockers, the diuretic drug hydrochlorothiazide, the anti-ulcer agents ranitidine and nizatidine, the antidiabetic drug metformin, and other medicines.^{25,26,31}

The key step in the synthesis of the drug valsartan is the formation of the tetrazole ring by the reaction of nitrile intermediate with tributyltin azide. Sodium nitrite is being used to expel remaining sodium azide reagent and under acidic condition, the nitrite ion could form nitrous acid, which could react with products from the degradation process of the solvent dimethylformamide (DMF) used in the route of synthesis.^{28,32} The tetrazole ring formation in the drug losartan involved using the solvent DMF and NaN_3 , which were precursors for obtaining of the nitrosamines. The synthetic pathways of irbesartan^{33,34} and candesartan³⁵ also contained azide–nitrile coupling to form tetrazole ring, which were risk factors for nitrosamine contamination. Ranitidine, that is a H_2 -blocker for control of the acidity in the stomach, has been reported as the most reactive precursor of *N*-Nitrosodimethylamine (NDMA).^{26,36} Metformin in reaction of oxidation could form NDMA under

high-pressure oxidative conditions or heat.³⁷ As well as, dimethylamine is the main precursor in the synthesis of ranitidine and metformin, thus obtaining NDMA impurities.³⁸ Last year FDA has found new nitrosamine impurities in the life-saving drugs rifampin and rifapentine (1-methyl-4-nitrosopiperazine (MNP) and 1-cyclopentyl-4-nitrosopiperazine (CPNP)) but they were not recalled from the market due to their importance.³⁹

The nitrosamine derivatives could be obtained in presence of primary, secondary, tertiary amines or quaternary ammonium salts along with nitrosating agents such as sodium nitrite, nitrous acid (HNO_2) or nitrite (NO_2) under acidic conditions. Some carbamates, *N*-alkyl amides, cyanamides, guanidines, amidines, hydroxylamines, hydrazines, hydrazones, and hydrazides might form nitrosamine impurities.^{26,40} In presence of O_2 could be formed other nitrosating agents – nitrosyl chloride (NOCl), nitrogen oxides (N_2O_3 , N_2O_4), nitrosonium tetrafluoroborate (NOBF_4)⁴⁰ and nitric oxide (NO)⁴¹ that could produce nitrosamines.²⁶ In addition, decomposition of solvents used in the route of synthesis of the drugs, such as dimethylformamide (DMF), dimethylacetamide (DMAc) or diethylacetamide (DEA), could lead to producing the mutagenic nitrosamines impurities.^{27,28}

In this context, it should be mentioned that the synthetic route suggested by us does not include the use of nitric acid, nitrite or solvents containing dialkylamino fragment and the structure of **5a-l** does not comprise tetrazole ring, dimethyl or piperazine fragments. In effort to assess the potential formation of nitrosamine impurities in the synthetic route, compound **5a** was studied by UV and ^{15}N -NMR spectroscopy. The UV spectra of nitrosamines in ethanol are characterized by two absorption bands – one of high intensity around 230 nm (similar to the band characterizing a N-NO_2 group), and another one of lower intensity within the region 345–374 nm (attributed to resonance structures close to the N=N band).⁴² However hydrazone **5a** showed strong own absorption around 230 nm in ethanol which hampers the detection of potential nitrosamine impurities

(Fig. S6, *cf.* ESI†). In the ^{15}N -NMR spectrum of **5a** in DMSO-d_6 one signal was registered at 320 ppm downfield of ammonia, taken as 0.0 ppm (Fig. S7, *cf.* ESI†) corresponding to the N-atom from the azomethine bond, and no other signals were found that might be attributed to nitrosamines. Reported chemical shifts for nitrosamine compounds are above 500 ppm on the ammonia scale.⁴³ Other studies reported for simple symmetrically-substituted nitrosamines NMR shifts from –107.2 to –146.7 ppm for $\text{N1}(-\text{N=O})$ and from +154.6 to +175.9 ppm for $\text{N2}(=\text{O})$, referenced to $^{15}\text{NO}_3\text{Na}$.⁴⁴ In unsymmetrically substituted nitrosamines N1 resonates respectively from –103.7 to –153.6 ppm, while N2 – from +157.0 to +176.9 ppm.⁴⁴

The nitrosation and formation of the nitrosamines could be also derived endogenous in the stomach. The conversion of L-arginine to citrulline in poly-morphonuclear leukocytes, is catalyzed by nitric oxide synthases (NOS) that lead to generation of nitric oxide (NO) which could react with oxygen thus producing nitrosating agents, such as N_2O_3 and N_2O_4 . Under certain conditions, these nitrosating agents could react with secondary amines to form nitrosamines.⁴⁵ On the other hand, it was shown that simultaneous presence of antioxidants such as ascorbate is able to block the intragastric nitrosation of drugs.⁴⁶

The low risk of nitrosamine formation from benzimidazole fragment is supported by the data available for the benzimidazole-containing drug albendazole whose principal route for primary metabolism is the rapid oxidation of its sulphide group to sulphonide, followed by further oxidation to sulphone, and deacetylation of the carbamate group to an amine.⁴⁶

2.2. Anthelmintic activity

The larvicidal effects of the newly synthesized compounds **5a-l** on encapsulated *T. spiralis* ML were studied *in vitro*. Biological assay results, shown in Table 1, indicate that all benzimidazoles

Table 1 *In vitro* activity of compounds **5a-l** against *T. spiralis* ML after 24 h and 48 h of exposure^a

Comp.	Concentrations 50 (100) $\mu\text{g ml}^{-1}$ in μM	Efficacy ^b (%)		Efficacy ^b (%)	
		After 24 h		After 48 h	
		50 $\mu\text{g ml}^{-1}$	100 $\mu\text{g ml}^{-1}$	50 $\mu\text{g ml}^{-1}$	100 $\mu\text{g ml}^{-1}$
5a	0.198 (0.396)	72.45	90.05	80.07	95.44
5b	0.186 (0.372)	100.00	100.00	100.00	100.00
5c	0.186 (0.372)	95.03	95.78	98.05	98.87
5d	0.186 (0.372)	100.00	100.00	100.00	100.00
5e	0.176 (0.352)	90.03	100.00	94.87	100.00
5f	0.188 (0.376)	5.74	15.34	24.60	40.25
5g	0.169 (0.337)	95.45	95.67	98.58	100.00
5h	0.169 (0.337)	10.33	95.20	28.17	97.98
5i	0.153 (0.306)	90.46	95.00	94.95	98.06
5j	0.177 (0.354)	40.15	100.00	61.47	100.00
5k	0.177 (0.354)	90.09	95.87	95.00	98.79
5l	0.177 (0.354)	85.43	100.00	91.03	100.00
Albendazole	0.188 (0.377)	10.68	10.80	14.63	15.61
Ivermectin	0.057 (0.114)	45.39	49.10	62.17	78.83

^a Control – 100 parasites. ^b $p < 0.05$.



in the tested concentrations were more active than the clinically used anthelmintic drugs albendazole and ivermectin.

The evaluation of the anthelmintic properties of the benzimidazole derivatives indicated that the hydrazones bearing only hydroxyl groups on the phenyl ring (compounds **5a–5e**) showed remarkable activity. **5a** with a single hydroxyl group in 2-position of the phenyl ring exhibited 90% larvicidal effect at concentration $100 \mu\text{g ml}^{-1}$ after 24 h. The introduction of a second hydroxyl group at 4-position of hydrazone **5a** to give compound **5c** produced 5% enhancement in activity. The shift of OH group from 4- to 3-position to give hydrazone **5b** led to further increase in activity (100% efficacy at a concentration of $50 \mu\text{g ml}^{-1}$). In accordance with these and our earlier results,¹⁸ it can be assumed that the presence of a OH group in the 3-position of the phenyl moiety of the molecule accounts for a larger share of the anti-*Trichinella spiralis* activity. The benzimidazolyl-2-hydrazone of 2,3-dihydroxybenzaldehyde (compound **5b**) and of 3,4-dihydroxybenzaldehyde (compound **5d**), respectively, killed the total parasitic larvae (100% effectiveness after 24 hour incubation period at 37°C) in

concentrations of $50 \mu\text{g ml}^{-1}$. A comparison of the biological effects revealed that the presence of two or three hydroxyl groups in the phenyl moiety led to benzimidazole derivatives **5b**, **5c**, **5d** and **5e** endowed with a much better anthelmintic activity against *T. spiralis* as compared with that of the naturally occurring stilbenoid polyphenol – resveratrol, which in study *in vitro* has showed a lethal effect on muscle *Trichinella* larvae only upon exposure to very higher concentrations (440 and $880 \mu\text{M}$) for 72 h.⁴⁷ On the other hand, there are numerous reports of natural phenolic compounds (including flavonoids) with anthelmintic effects against parasitic nematodes other than *T. spiralis*^{48–50} and their anthelmintic activity increases with the number of hydroxyl groups.⁵⁰

However, the presence of a hydroxyl group at 3-position in combination with a 4-methoxyl moiety (compound **5l**) led to slight reduction in activity (85% larvicidal effect after 24 h in concentration of $50 \mu\text{g ml}^{-1}$) as compared to hydrazone **5d**. Methoxy-analogues (**5f** and **5h**) showed less pronounced activities against the parasites, comparable to that of the anthelmintic albendazole, after 24 h of exposure in $50 \mu\text{g ml}^{-1}$

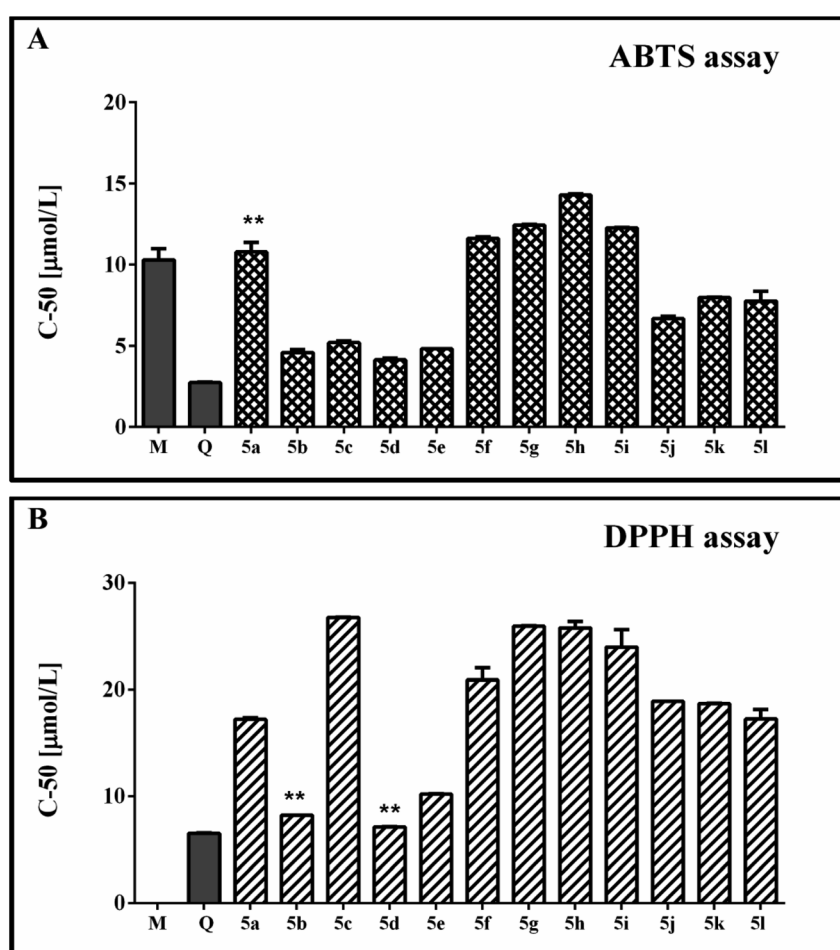


Fig. 2 Anti-radical properties of compounds **5a–l** in spectrophotometric model systems containing stable free radicals. Results have been presented as C-50 values calculated on the base of the concentration–RSA% relationship. (A) Data from the ABTS cation radical containing model systems; (B) data obtained using the DPPH assay. Melatonin (M) and quercetin (Q) have been used as reference compounds. Data are expressed as the mean \pm SD. Groups were compared by one-way ANOVA with Bonferroni post hoc test. The results were considered statistically significant when $p < 0.05$. ** ($p > 0.05$) compared with Melatonin; ● ($p > 0.05$) compared with quercetin.

concentration. The larvicidal effects of the tested compounds (excluding **5f**, **5h**, **5j** and **5l**) were dose-independent *i.e.* the viability of *T. spiralis* larvae was almost unchanged with 2-fold increase in the dose (100 µg ml⁻¹, Table 1).

Based on the *in vitro* activity of hydrazones **5a–l** on *T. spiralis* ML as well as that of other recently investigated 1*H*-benzimidazolyl-2-hydrazone¹⁸ it was outlined that the combination of benzimidazole and hydrazone pharmacophores in addition with hydroxyl and/or methoxyl groups on the arylidene moiety favorably modulates the anti-*Trichinella spiralis* activity. Indeed, the studied benzimidazole hydrazones showed remarkable larvicidal effect superior to the benzimidazole-containing derivatives earlier synthesized by us.^{22,51–53} Thus, compounds **5a–l** deserve more detailed pharmacological and toxicological investigations for the development of potential new anti-*Trichinella spiralis* agents.

2.3. Experimental estimation of the radical scavenging activity

The observed radical scavenging effects of the new 1*H*-benzimidazole-2-yl hydrazones against stable free radicals ABTS and DPPH are illustrated in Fig. 2A and B. The performed experiments denoted that the absorbance values of the working solutions of both radicals decreased as the sample concentrations of the tested compounds increased. The curves for the relationship between the concentration and the radical scavenging activity (RSA), obtained on the basis of these results, were with well-expressed linear dependences ($R^2 < 0.95$). They were used to estimate the C-50 values of the compounds and to compare their scavenging activity. Two known antioxidants – quercetin and melatonin, were used as reference compounds in the experiments in order to judge better on the relative radical scavenging properties of the 1*H*-benzimidazole-2-yl hydrazones **5a–l**. Melatonin contains an indolyl heterocycle, structurally close to the benzimidazole one, while quercetin is a flavonoid compound including a catechol fragment which is also a structural unit relevant the hydrazones studied by us.

The data obtained from the ABTS model system denoted C-50 values of the studied 1*H*-benzimidazole-2-yl hydrazones close or lower than that of the reference melatonin. The hydrazones bearing only methoxy groups in the phenyl ring (mono-, di- and tri-methoxy) had higher C-50 values compared to the references, suggesting necessity of higher amount of these compounds compared to melatonin and quercetin in order to achieve 50% RSA. Within this group of compounds, **5h** was the least active – with the highest observed C-50 value – 14.30 µmol l⁻¹. **5f**, **5g** and **5i** had statically identical C-50 values. They were lower compared to **5h** suggesting better scavenging activity. **5f** had an equivalent C-50 value with the mono-hydroxyphenyl hydrazone **5a**.

The presence of both methoxy and hydroxy groups in the tested 1*H*-benzimidazole-2-yl hydrazones induced significant decrease in the C-50 value compared to **5f** and **5a**. The C-50 values of this subgroup comprising **5j–l** were lower than 8 µmol l⁻¹. They demonstrated necessity of lower concentration of the active substance to induced 50% decrease of the RSA%

compared to melatonin, but did not reach the C-50 concentration of quercetin.

The di- and trihydroxy substituted 1*H*-benzimidazole-2-yl hydrazones **5b–e** demonstrated the best RSA within the whole series. They had very similar and in some cases even identical C-50 values ranging from 4.15 to 5.20 µmol l⁻¹. The observed effect was much higher compared to melatonin, but still did not reach the quercetin's effectiveness.

In the DPPH model system, the reference melatonin did not exert capability to decrease the absorbance value of the radical solution. As observed in the ABTS system, among the group of the di- and trihydroxy substituted benzimidazoles were the most potent scavengers. The C-50 value of **5e** was slightly higher than quercetin (with a statistically significant difference). **5b** and **5d** denoted identical concentration as the mentioned reference. The hydrazone with the mono-hydroxyphenyl moiety **5a** and the hydrazones with mixed methoxy/hydroxy substitution *i.e.* **5j–l** showed equivalent C-50 values. The compounds with di- and trimethoxyphenyl moiety, similarly to the ABTS system, exerted the highest C-50 values together with the dihydroxy substituted benzimidazoles **5c**.

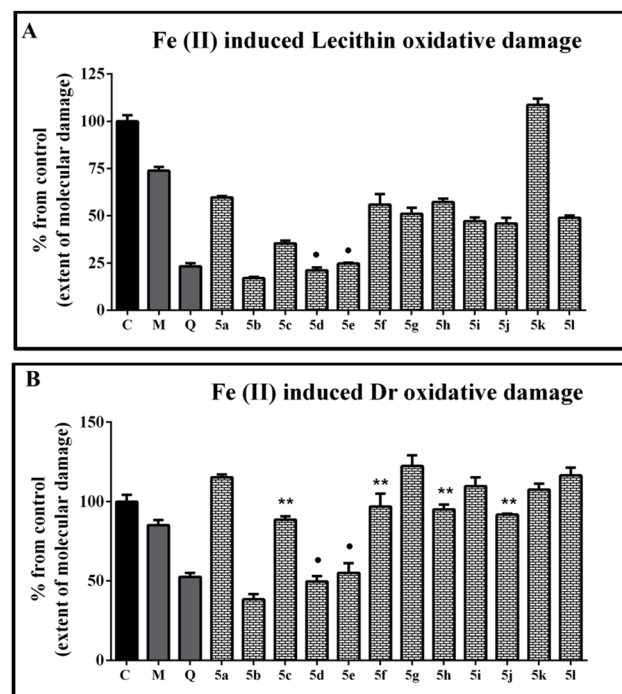


Fig. 3 Protection effects of compounds **5a–l** on iron induced oxidative damage of biologically relevant molecules: (A) extent of molecular oxidative damage observed in lecithin containing model system (1 mg ml⁻¹) in the presence of compounds **5a–l** (90 µM); (B) extent of molecular oxidative damage observed in deoxyribose containing model system (0.5 mmol l⁻¹) in the presence of compounds **5a–l** (80 µM). Melatonin (M) and quercetin (Q) at the same concentration have been used as reference compounds in both systems. Data were compared by one-way ANOVA with Bonferroni post hoc test. Results are presented as "% of molecular damage" means ± SD. The results were considered statistically significant when $p < 0.05$. ** ($p > 0.05$) compared with melatonin; ● ($p > 0.05$) compared with quercetin.



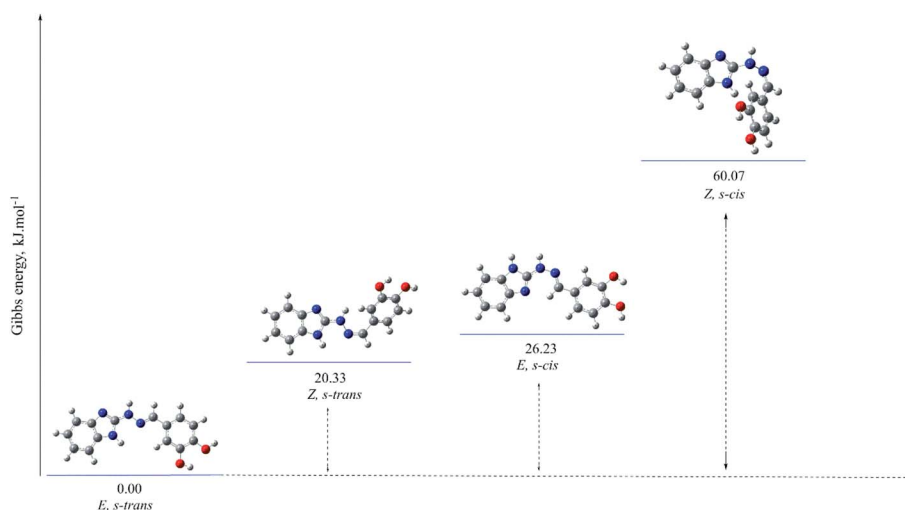


Fig. 4 Optimized molecular structure and relative Gibbs energies (in kJ mol^{-1}) of the possible isomers of compound **5d** in amino tautomeric form, obtained at B3LYP/6-311++G(d,p) level of theory in gas phase.

The effect of the tested new *1H*-benzimidazole-2-yl hydrazones on iron induced lecithin oxidative damage is presented in Fig. 3A. In the samples containing the reference compounds and all hydrazones, except for **5k**, administrated at concentration 90 μM , the absorbance values at 532 nm were lower compared to the controls. This indicates decrease of the extent of molecular damage and well expressed protection effect against ferrous induced peroxidation. The estimated extent of molecular damage in the presence of quercetin was around 20%. In the samples containing mono-, di- and trimethoxyphenyl substituted hydrazones the observed extent of molecular damage was higher than 45%. The effect of the mono-methoxy hydrazone **5f** was statistically identical to those of the dimethoxy substituted derivatives **5g** and **5h**. The modulation effect of **5f** was also comparable to the results for the hydrazone with mono hydroxyphenyl moiety **5a**. The trimethoxy substituted hydrazone **5i** denoted more substantial protection compared to **5f** but it was still equivalent to the one of **5g** as well to that of **5j** and **5l**. The di- and trihydroxy substituted hydrazones **5b**, **5d** and **5e** showed excellent protective effect – for **5d** and **5e** equal to the effect of quercetin.

The ability of the compounds to protect the deoxyribose molecules from iron induced molecular degradation is shown in Fig. 3B. In this system the tested benzimidazoles displayed both prooxidant and antioxidant effects depending on the molecular structure. The extent of the observed prooxidant effect was negligible to modest. The mono-, di- and trimethoxy substituted hydrazones either increased the absorbance values compared to the controls or did not exert any modulation effect. More substantial deoxyribose degradation and enhancement of the induced by iron molecular damage has been observed in the samples containing **5a**, **5g** and **5l**. For all of these samples the estimated parameter was over 115%.

As in the lecithin containing system, the derivatives possessing hydroxyl groups in vicinal positions in the phenyl ring were better molecular protectors under the conditions of

ferrous iron induced peroxidation. Compounds **5d** and **5e** had an effect equivalent to that of the reference quercetin and denoted extent of molecular damage close to 50%. The strongest protection effect was observed with the samples containing **5b** which exerted extent of molecular damage lower than 40% – considerably better protection effect compared to both reference compounds.

Comparing the structures and the exerted modulation effects of the studied compounds it is inferred that the presence of a more than one hydroxyl group in the molecular structure of the derivatives is a crucial factor affecting their capability to protect the biologically important molecules in both chosen model systems. This relationship follows the trends found in the anthelmintic study.

The performed assays denoted a higher radical scavenging ability of the hydroxyl substituted *1H*-benzimidazole-2-yl hydrazones than other earlier studied benzimidazole derivatives as for example benzimidazoles containing *N*-methyl-1,3,4-thiadiazol-2-amine and 4-methyl-2*H*-1,2,4-triazole-3(4*H*)-thione moieties,⁵⁴ 2-amino- and 5-aryl-1,3,4-oxadiazole moieties⁵⁵ and

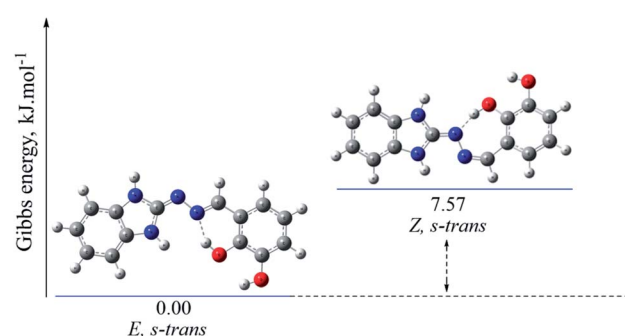


Fig. 5 Optimized molecular structure and relative Gibbs energies (in kJ mol^{-1}) of the possible isomers of compound **5b** in imino tautomeric form, obtained at B3LYP/6-311++G(d,p) level of theory in gas phase.



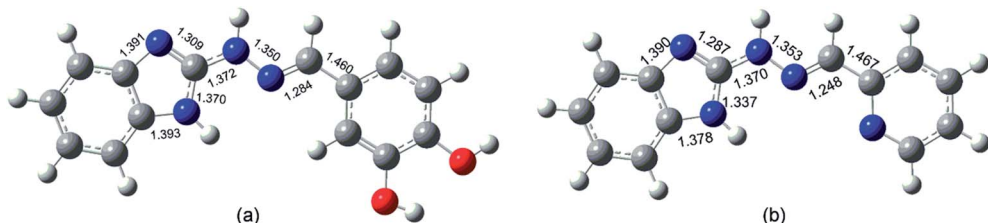


Fig. 6 General view of the molecule of compound 5d (a) and benzimidazolyl-2-hydrazone of pyridine-2-carbaldehyde (b,⁵⁸) with selected bond lengths.

N,N-disubstituted benzimidazole-2-thione hydrazone derivatives.¹⁶ On the other side, taking into account the comparison to the reference compounds used in the experiments, it could be concluded that the observed effects of the hydroxyl substituted 1*H*-benzimidazole-2-yl hydrazones are close to those of catechin and flavonol compounds⁵⁶ and superior to melatonin derivatives.⁵⁷

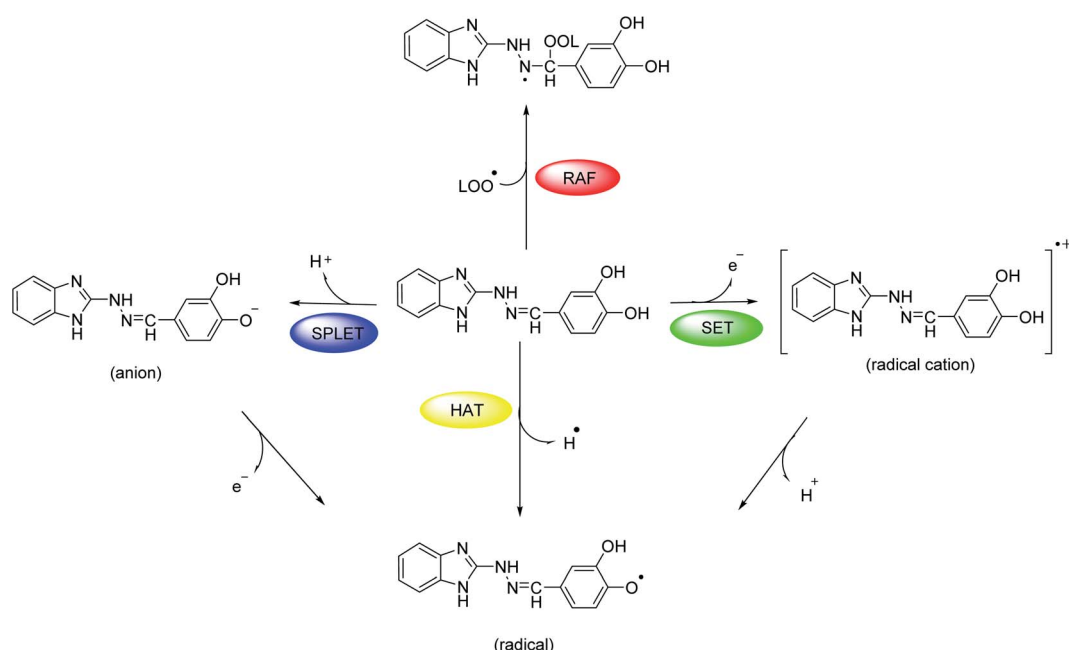
2.4. Molecular structure optimization

In order to rationalize the observed effects, the molecular structure and the radical scavenging properties of compounds 5a–l were characterized by DFT methods. In consideration of the fact that the studied compounds can exist in two tautomeric forms – amino and imino (Fig. 4), their geometries were optimized fully using (U)B3LYP density functional theory (DFT) method and 6-311++G** basis set in gas phase and solvents: water and benzene. Solvent effects were included in the optimizations by using the Polarizable Continuum Model (PCM). Single-point (SP) energy calculations were performed additionally at the M062X/6-311+G** level of theory, on the optimized B3LYP geometries. The calculated geometries included also Z

and *E* configurations of the double C=N (azomethine) bond, *cis* and *s-trans* configuration of the N–N bond, and different orientations of the substituents in the phenyl ring for both tautomeric forms (Fig. 4). The calculated Gibbs free energies *G* of the possible isomers of all studied compounds in amino and imino form are listed in the ESI.†

Summarizing the computations, the imino form strongly prevails in all compounds containing hydroxyl group in *ortho* position of the phenyl ring, otherwise the amino form is the favourable one. The energy difference between them is about 2.06–12.88 kJ mol^{−1} (cf. ESI†).

For compound 5d containing two hydroxyl groups at positions 3 and 4 in the phenyl ring and showing the most potent radical scavenging effect towards free stable radicals ABTS and DPPH, the amino : imino ratio is: 81.66 : 18.34 (gas phase); 91.97 : 8.03 (water); 96.18 : 3.82 (benzene) according to the B3LYP calculations. The results obtained with SP M06-2X functional indicate even greater prevalence of the amino form in all kind of media: 90.60 : 9.40 (gas phase); 96.87 : 3.13 (water); 98.37 : 1.63 (benzene) respectively.



Scheme 2 Probable mechanisms of radical-scavenging activity.



On the other hands, for compound **5b** containing also two hydroxyl groups, but at positions 2 and 3 in the phenyl ring and showing the highest protection effect towards lecithin and deoxyribose, the amino : imino ratio is 3.48 : 96.52 (gas phase); 9.98 : 90.02 (water); 4.60 : 95.40 (benzene) according to the B3LYP calculations.

The optimized molecular structures of the most stable isomers of compounds **5b** (in imino tautomeric form) and **5d** (in amino tautomeric form) are presented for illustration in Fig. 4 and 5. The computational results showed that *E* configuration of the double azomethine bond is more stable for both amino or imino form, as well as *s-trans* – for the arrangement around the N–N bond. The molecular geometry of the preferred amino and imino isomers is essentially flat.

The gas-phase energy differences between the different isomers of the compounds with preferred amino forms are within the following ranges: *E*, *s-trans* → *E*, *s-cis* (ΔG 16.90–47.29 kJ mol⁻¹) → *Z*, *s-trans* (ΔG 16.91–34.60 kJ mol⁻¹) (cf. ESI†).

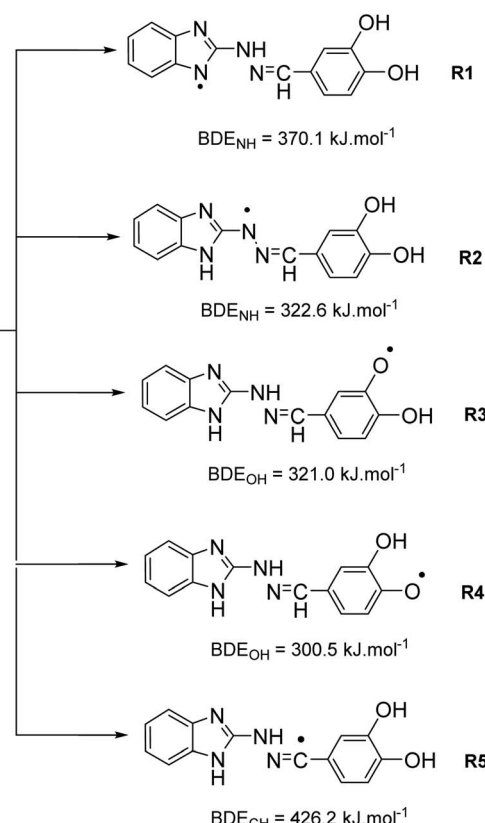
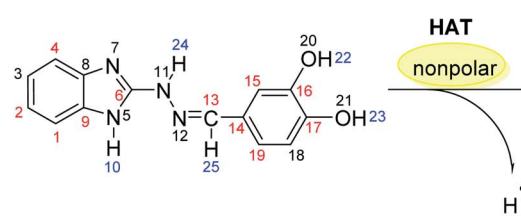
The theoretically estimated geometry of the amino *E*, *s-trans* forms is in a significant correlation with earlier reported X-ray experimental data.⁵⁸ The calculated bond lengths of compound **5d** are close to the values of the benzimidazolyl-2-hydrazone of pyridine-2-carbaldehyde examined by Pervova *et al.* (Fig. 6).⁵⁸ The *E* configuration of the double C=N bond of the target hydrazones is compatible with the reported structures of hydrazones containing benzothiazole⁵⁹ as well as benzimidazole fragment.⁵⁸

The gas-phase energy differences between the different isomers of the compounds with preferred imino forms vary as follows: *E*, *s-trans* → *E*, *s-cis* (ΔG 0.88–35.34 kJ mol⁻¹) → *Z*, *s-trans* (ΔG 4.17–28.34 kJ mol⁻¹) (cf. ESI†).

2.5. Computational study of the radical scavenging activity

At molecular level, the studied hydrazones **5a–l** can exert their radical scavenging activity through several probable mechanisms (Scheme 2). The first one of them comprises direct hydrogen atom transfer from the antioxidant to the free radical (HAT mechanism). Other probable mechanisms are the two-steps reaction pathways where the electron and proton are transferred in separate steps from the antioxidant to the free radical: single electron transfer followed by proton transfer (SET-PT mechanism) and sequential proton loss electron transfer (SPLET mechanism). A fourth possible way for deactivation of the free radicals is radical adduct formation (RAF mechanism) where the antioxidant covalently binds to the free radical.

In the same way, the molecular structure of the studied 1*H*-benzimidazole-2-yl hydrazones suggest several possible sites for hydrogen atom transfer, deprotonation and radical adduct formation. For instance, compound **5d** might cleave the bonds N5–H10, N11–H24, O20–H22, O21–H23 and C13–H25 (for numbering cf. Scheme 3) in order to transfer a hydrogen atom or a proton as shown in blue in Scheme 3. Radical attack for adduct formation is also possible at several reaction sites: C1,



Scheme 3 Possible sites for hydrogen atom transfer (in blue) and radical adduct formation (in red) for compound **5d** along with the radicals formed by HAT and the respective bond dissociation enthalpies, calculated at B3LYP/6-311++G(d,p) level of theory in benzene.



C2, C4, C6 *etc.* (shown in red in Scheme 3). The reactivity for HAT of the above mentioned sites in compound **5d** was estimated by calculating the respective bond dissociation enthalpies (BDE) resulting in the formation of radicals **R1–R5** (Scheme 3). The most reactive site is denoted by the lowest enthalpy. It was previously demonstrated that DFT/B3LYP calculations are a good choice for estimation of the reaction enthalpies as they can provide a fairly accurate agreement between the calculated values and the available experimental data as well as describe reliably the structure–activity relationship for antioxidants with diverse chemical structure at reasonable computational time.^{60–64} On the other hand, the use of at least double-zeta basis set, including polarization and diffuse functions, is recommended,⁶⁵ therefore the B3LYP//6-311++G(d,p) combination was applied in the present calculations. The results obtained at this computational level allow a useful comparison with earlier studied antioxidants.

The calculations in gas phase and benzene medium outlined the hydroxyl group at *para* position in the phenyl ring as the most active one (BDE in benzene 300.5 kJ mol⁻¹; BDE in gas phase 293.2 kJ mol⁻¹), followed by the hydroxyl group at *meta* position in the phenyl ring (BDE in benzene 321.0 kJ mol⁻¹; BDE in gas phase 313.4 kJ mol⁻¹) and the amino group in the hydrazine chain (BDE in benzene 322.6 kJ mol⁻¹; BDE in gas phase 316.5 kJ mol⁻¹). Moreover, the calculated BDE values are close to the calculated BDE values of known phenolic antioxidants such as α -tocopherol (BDE in gas phase 297 kJ mol⁻¹, calculated at the same level of theory⁶⁰) and quercetin (with lowest O–H BDE value estimated to 305 kJ mol⁻¹ at the same level of theory⁶⁶).

The amino group in the benzimidazolyl fragment and the C–H bond from the azomethine group of compound **5d** are characterized by a much lower reactivity – with BDE values in

benzene 370.1 kJ mol⁻¹ (BDE in gas phase 366.7 kJ mol⁻¹) and 426.2 kJ mol⁻¹ (BDE in gas phase 418.7 kJ mol⁻¹), thus not expected to contribute significantly to the radical scavenging properties of the compound. For comparison, the BDE of the indolyl N–H bond in melatonin and the methylene group next to the indolyl fragment were also calculated. The first one is 355 kJ mol⁻¹, whereas the latter – 428 kJ mol⁻¹, respectively.

The radical scavenging ability of the antioxidants is related to the stability of the formed radicals, which on turn is affected by the delocalization of the unpaired electron over the conjugated system.⁶⁷ In order to estimate the stability of **R4**, **R3** and **R2** radicals of **5d**, the Natural Bond Orbital (NBO) spin density distribution was examined and compared (Fig. 7).

Radical **R4** showed the highest delocalization of spin density – only 0.176 of the unpaired electron are found on O21, while the greater part of it is spread over the phenyl ring and the hydrazone chain. In radical **R3** the spin density is delocalized only over the phenyl ring with 0.231 e⁻ localized on O20, while in radical **R2** it is limited within the hydrazone chain and part of the benzimidazole ring, with 0.311 e⁻ localized on N11. These data indicate **R4** as the most stable radical, followed by **R3** and **R2**. The order of stability of radicals **R4**, **R3** and **R2**, correspond to the order of reactivity at the different reaction sites established by the calculated BDE values. Such correlation between the two parameters has been previously discussed for other phenolic antioxidants.^{68–71}

Further the reaction enthalpies for HAT, SET-PT and SPLET mechanisms of all hydrazones **5a–l** were computed in gas phase and water. The ability to deactivate free radical *via* HAT mechanism was described by the BDE values. The ionization potentials (IP) were used to describe the ability of the compounds to transfer a single electron *via* SET-PT mechanism, while the proton affinities (PA) were used to estimate the ability to

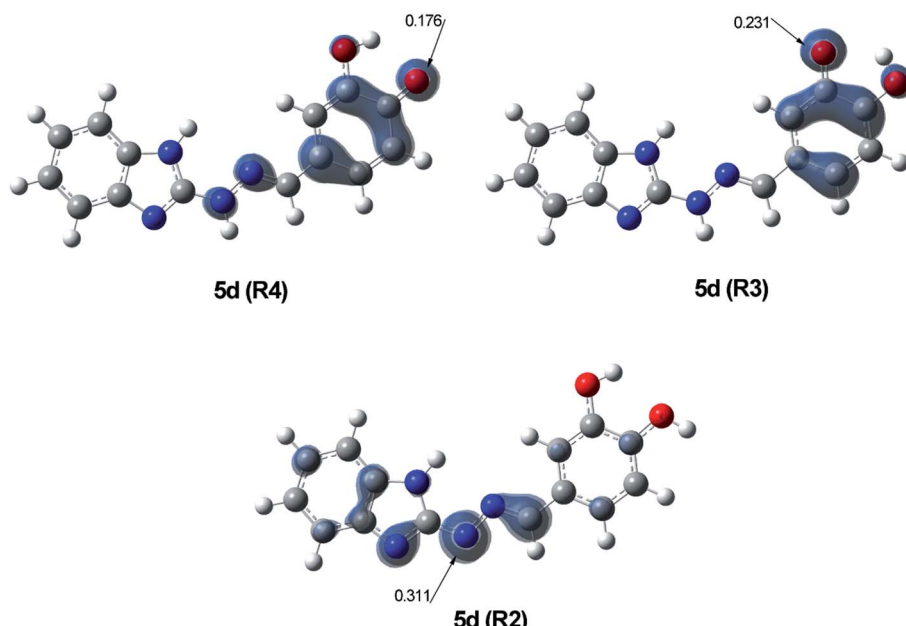


Fig. 7 Visualisation of spin density of the three most stable radicals **R4**, **R3** and **R2** of **5d**, calculated at B3LYP/6-311++G(d,p) level of theory in benzene.



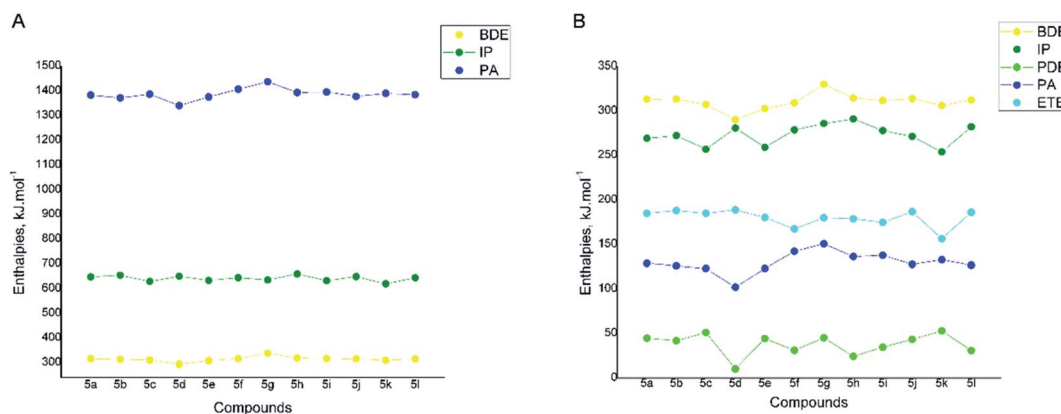


Fig. 8 Reaction enthalpies for compounds **5a-l** in gas phase (A) and water (B): dissociation enthalpy (BDE), ionization potential (IP), proton dissociation enthalpy (PDE), proton affinity (PA), and electron transfer enthalpy (ETE), calculated at B3LYP/6-311++G** level of theory.

transfer a proton *via* SPLET. Based on the comparison of the BDE values to IPs and PAs, it is possible to judge which mechanism is expected to govern the antioxidant action of the studied compounds. The lower the reaction enthalpy, the more easily would be followed the particular reaction pathway. On the other hand, the comparison of the reaction enthalpies of the different compounds might clarify the relationship between the molecular structure and the experimentally determined radical scavenging activity.

The reaction enthalpies for each compound were modelled at all possible reaction sites. Fig. 8A presents the lowest BDEs, IPs and PAs found for all compounds in gas phase. As it could be seen the IP and PA values lay far above the respective BDE values for all studied compounds (Fig. 8A), pointing out that the SET-PT and SPLET mechanism require significantly more energy than HAT and therefore are not likely to occur in nonpolar phase.

It was established that for compounds **5d** and **5l**, bearing hydroxyl groups in 3 and 4 position of the phenyl ring, the lowest BDE corresponds to the cleavage of one of the O–H bonds and are 293.21 and 314.49 kJ mol⁻¹, respectively. For compounds **5f-i**, where only methoxyl groups are present in the phenyl ring, the lowest BDEs are connected to the cleavage of the N5–H10 bond from the benzimidazole fragment. Correspondingly they are higher than the O–H BDE values of **5c-e** and range from 315.24 to 373.81 kJ mol⁻¹. For all derivatives containing hydroxyl groups in 2-position of the phenyl ring the lowest BDE are connected to the cleavage of the N5–H10 bond similarly to **5f-i**. It should be noted that the compounds bearing vicinal hydroxyl groups *i.e.* **5b-e** possess the lowest BDEs within the whole studied series and the 3,4-dihydroxyl derivative **5d** is outlined as the most active one with BDE of 293.21 kJ mol⁻¹. Based on that, it is expected to be the most active *via* HAT mechanism in nonpolar medium.

In water the BDE values of compounds **5a-l** are slightly lowered in comparison to the gas phase ones (they vary in the range 291.75–330.01 kJ mol⁻¹), but follow the same trend (Fig. 8B).

On the other hand, the calculated O–H and N–H proton affinities of **5a-l** in water are much lower than the respective BDEs showing that in polar medium the first step (deprotonation) of SPLET mechanism is favored over HAT. The PAs are connected to the same functional groups as in the HAT process. PAs are between 108.30 and 150.47 kJ mol⁻¹ – the lowest values were found again for the compounds bearing vicinal hydroxyl groups **5b-e**, and particularly for the 3,4-dihydroxyl derivative **5d**. The second step (electron transfer) of the SPLET mechanism, described by the electron transfer enthalpy (ETE), also requires lower energy than the homolytic cleavage of the O–H or N–H bond for all studied compounds.

The ionization potentials of **5a-l** in water are lower than the BDEs, but still higher than the PA values which indicate that the SET-PT mechanism is less probable than the SPLET in polar medium.

Summarizing the data in polar medium, it could be concluded that in this case SPLET is regarded as the most probable mechanism of antioxidant action and the derivatives with vicinal hydroxyl groups are expected again to show the highest activity. These results are in good agreement with the experimentally determined radical scavenging effects in DPPH, lecithin and deoxyribose assays, where the hydroxyl substituted hydrazones demonstrated the most potent radical scavenging effect. The presence of a phenyl fragment with vicinal hydroxyl groups has been associated to improved radical scavenging properties due to its conversion to *ortho*-quinone intermediates.^{72,73}

The ability of the studied compounds to deactivate different free radicals was modeled by transition states involving attack by $\cdot\text{OCH}_3$, $\cdot\text{OOH}$ and $\cdot\text{OOCH}_3$, at all reactive sites identified in the preceding computational study. The calculations were carried out with compound **5d** at SP M06-2X/6-311++G** level of theory. M06-2X/6-311++G** was selected for the computations taking into account the reliability of this method for transition state studies.^{69,74,75} The activation energies (ΔG^\ddagger) for all studied transition states in gas phase and solvent (water and benzene) for all three attacking radicals are presented in Table 2.



Table 2 Gibbs free energies of activation (ΔG^\ddagger) for the formation of transition states (TSs) of compound **5d** with various free radicals corresponding to HAT mechanism, at 25 °C, in kJ mol⁻¹

TSs	'OCH ₃			'OOH			'OOCH ₃		
	Gas phase	Water	Benzene	Gas phase	Water	Benzene	Gas phase	Water	Benzene
TS H10	67.64	69.23	77.44	90.83	81.24	90.59	95.07	86.47	97.32
TS H24	33.15	89.83	—	64.81	77.91	74.32	74.46	82.93	84.91
TS H22	—	18.08	—	75.71	81.95	86.09	80.93	81.31	90.32
TS H23	—	18.30	—	68.97	83.31	74.56	71.73	83.13	67.47
TS H25	71.01	83.94	79.93	115.15	125.92	122.74	119.82	132.74	128.83

The most stable transition states **TS H24**, **TS H22** and **TS H23** correspond to the formation of the most stable radicals **R2**, **R3** and **R4** for all attacking species. Unfortunately, we could not

locate transition states between **5d** and **'OCH₃** for **TS H24** in benzene and **TS H22** and **TS H23** in gas phase and benzene. The scans made (5d)-O···H···O-CH₃ for **TS H22**'s and **TS H23**'s and

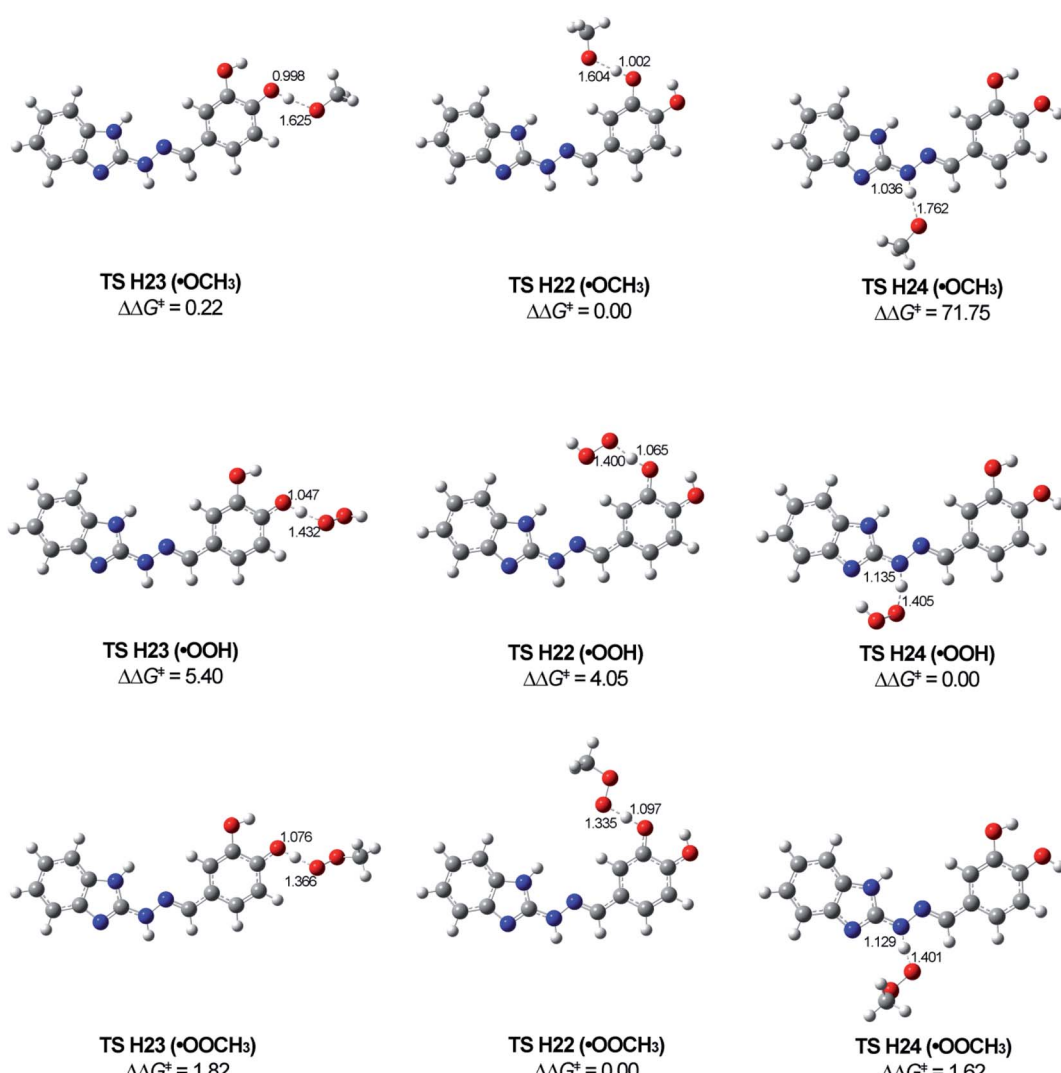


Fig. 9 Optimized structures of the TSs for the most stable radicals R2, R3 and R4 corresponding to the HAT mechanism. $\Delta\Delta G^\ddagger$ shows the difference in the activation energies between the TSs for every one of the studied radicals. Calculations are done on SP M06-2X/6-311++G** method in solvent water.



Table 3 Gibbs free energies of activation (ΔG^\ddagger) for the formation of transition states (TSs) of compound **5d** with various free radicals corresponding to RAF mechanism, at 25 °C, in kJ mol⁻¹

TSs	ΔG^\ddagger [(U)M06-2X/6-311++G**//(U)B3LYP/6-311++G**]			'OOH			'OOCH ₃		
	Gas phase	Water	Benzene	Gas phase	Water	Benzene	Gas phase	Water	Benzene
TS C1	68.15	74.93	75.89	102.16	104.87	107.80	115.53	119.58	122.07
TS C2	70.23	72.73	74.54	101.96	100.35	107.29	115.26	114.57	119.53
TS C4	72.23	71.69	75.85	102.99	100.31	107.88	117.04	116.04	121.04
TS C6	83.64	92.37	91.53	101.62	109.66	109.84	107.90	118.34	116.40
TS C9	73.18	78.53	80.13	107.77	107.56	112.57	121.37	121.36	126.40
TS C13	52.87	60.20	59.55	73.13	82.25	80.70	80.72	91.13	88.93
TS C14	82.30	83.24	87.81	113.28	114.10	117.98	125.51	127.23	130.16
TS C15	70.08	70.10	76.12	102.25	101.91	107.36	112.21	113.95	117.81
TS C16	70.29	72.34	82.25	107.55	101.44	108.84	117.23	115.83	121.77
TS C17	47.72	61.16	57.31	83.81	91.66	91.33	95.10	104.29	102.26
TS C19	65.02	70.42	71.00	94.11	97.89	98.95	106.31	112.12	112.56

(**5d**)-N···H···O-CH₃ for **TS H24** do not indicate maximum in the energies corresponding to transition state (*cf.* ESI, Fig. S1†). Having in mind the activation energies for **TS H22** ('OCH₃) and **TS H23** ('OCH₃) in water (around 18 kJ mol⁻¹ only) and the activation energies of the TS's for the other attacking radicals (values in gas phase and benzene lower than in water), it seems that the not detected TS's should have lower than 18 kJ mol⁻¹ activation energy.

The optimized structures of the most stable TS's (Fig. 9) show similarity in regard to the bond distances (in Å) for the breaking of O···H and N···H and forming new H···O bonds with the different free radicals. The distances in **TS H24**'s are longer compared to the other two transition states (**TS H23** and **TS H22**) for the same attacking free radical.

According to our theoretical study, the radical scavenging can go through RAF mechanism as well. The calculations

showed that the most stable transition states (TS) between the studied radicals and **5d** are at C13 position (Table 3).

The optimized geometries of the possible transition states involved in the RAF mechanism are shown in Fig. 10. The most relevant parameter is the C···O distance corresponding to the forming new bond and it was found to be around 2 Å for all calculated TS's.

The values of $\Delta G_{\text{reaction}}$ in Table 4 for this mechanism showed that concerning the 'OCH₃, the reaction can proceed at C13 and C17 position in **5d** in water and benzene; when the radical is 'OOH, the reaction still will be possible at C13 (water and benzene), while when the reacting species is 'OOCH₃, the reaction will follow this mechanism with difficulty.

Hypothetically, the RAF mechanism can occur at N12 (Scheme 3) as well, but any attempt to locate the corresponding products and transition states brought us failure.

When the reacting species is 'OOH both mechanisms (HAT and RAF) can be followed: HAT – at H23, H22 and H24, and RAF – at C13 position. The most stable TSs for HAT and RAF mechanism have similar values of ΔG^\ddagger (HAT: H24–77.91 (water) and 74.32 (benzene) kJ mol⁻¹ and RAF: C13–82.25 (water) and 80.70 (benzene) kJ mol⁻¹).

The 'OOCH₃ could be deactivated only by HAT mechanism. The calculations showed a lower reactivity of the radical compared with the other two studied radicals. According to the $\Delta G_{\text{reaction}}$, the reactions are slightly exothermic and in the same time the ΔG^\ddagger of the most stable TSs are higher than the ΔG^\ddagger of the other two radicals at the respective positions.

All gathered computational data demonstrated that the newly synthesized 1*H*-benzimidazole-2-yl hydrazones possess very versatile radical scavenging properties – several reaction sites characterized by relatively low reaction enthalpies and possibility to act simultaneously through several possible reaction pathways. It was also found that the predicted activities of compounds **5a–l** are in good correlation with the experimentally observed antiradical activity.

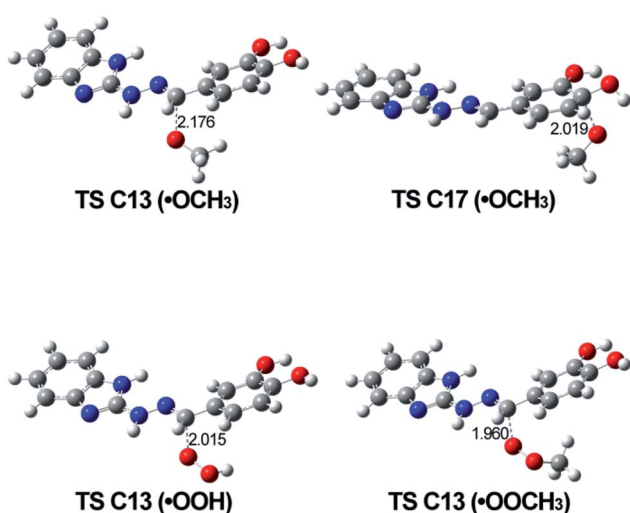


Fig. 10 Optimized structures of the most probable TSs corresponding to the RAF mechanism for every one of the studied radicals. Calculations are done in solvent water.



Table 4 Gibbs free energies of reaction ($\Delta G_{\text{reaction}}$) corresponding to hydrogen atom transfer (HAT) and radical adducts formation (RAF) of compound **5d** with various free radicals, at 25 °C, in kJ mol⁻¹, with respect to the isolated reactants^a

Reaction site	$\Delta G_{\text{reaction}}((\text{U})\text{M06-2X}/6-311++\text{G}^{**}/(\text{U})\text{B3LYP}/6-311++\text{G}^{**})$								
	'OCH ₃			'OOH			'OOCH ₃		
	Gas phase	Water	Benzene	Gas phase	Water	Benzene	Gas phase	Water	Benzene
HAT									
H10	-36.36	-46.88	-37.65	34.50	23.15	32.96	40.00	30.75	39.62
H24	-84.27	-87.42	-82.32	-13.41	-17.39	-11.71	-7.91	-9.79	-5.05
H22	-84.32	-84.46	-83.50	-13.46	-14.43	-12.89	-7.96	-6.83	-6.23
H23	-99.63	-99.29	-98.46	-28.77	-29.26	-27.85	-23.27	-21.66	-21.20
H25	3.36	2.21	5.44	74.22	72.24	76.05	79.71	79.84	82.70
RAF									
C13	-72.93	-59.30	-63.33	-12.69	2.89	-1.05	-1.74	14.23	7.63
C17	-33.50	-20.52	-24.00	33.22	43.65	41.08	43.15	54.25	51.38
C19	-7.15	3.53	0.05	49.64	57.48	56.23	61.73	69.46	67.33
C6	-7.44	6.19	2.97	46.14	59.22	55.08	59.45	73.11	69.60
C15	-0.32	6.64	6.26	55.90	61.67	62.62	66.13	71.97	72.00
C1	5.92	16.26	13.97	63.08	70.87	70.68	76.25	84.87	83.93
C2	12.27	20.53	19.17	67.01	70.94	72.33	80.20	85.44	87.28
C4	3.91	10.93	11.13	61.61	63.63	66.99	72.86	75.68	79.21
C9	32.02	42.70	40.60	91.60	96.73	98.32	106.09	110.98	112.45
C14	31.47	37.22	37.58	87.01	92.49	91.45	98.65	103.71	106.64
C16	4.70	8.28	11.17	68.99	69.92	73.65	80.22	81.57	86.95

^a Having in mind the values of $\Delta G_{\text{reaction}}$ and ΔG^{\ddagger} for the TSs of HAT and RAF mechanisms, the reaction between $'\text{OCH}_3$ and **5d** will go mainly through HAT mechanism at H23, H22 and H24 position.

3. Conclusions

To conclude, several 1*H*-benzimidazole-2-yl hydrazones as potent multi target drugs, combining antioxidant and anthelmintic activity, were synthesized using *o*-phenylenediamine as precursor and variety of methoxy- and hydroxyl-substituted benzaldehydes. The newly synthesized 1*H*-benzimidazole-2-yl hydrazones exhibited higher anthelmintic activity against isolated *Trichinella spiralis* muscle larvae *in vitro* than the clinically used anthelmintic drugs albendazole and ivermectin in the tested concentrations. The 1*H*-benzimidazole-2-yl hydrazones containing more than one hydroxyl groups in the phenyl moiety (compounds **5b–e**) showed a potent anthelmintic activity – above 90% larvicidal effect at concentration 50 µg ml⁻¹ after 24 h, while the methoxy derivatives showed less pronounced activities against the parasites. Compounds **5b** and **5d** were the most active ones with 100% effectiveness after the 24 hour incubation period at 37 °C in concentrations of 50 µg ml⁻¹ and 100 µg ml⁻¹.

The *in vitro* evaluation of the antioxidant properties of the new 1*H*-benzimidazole-2-yl hydrazones based on the stable free radicals ABTS and DPPH as well as ferrous iron induced oxidative molecular damage in lecithin and deoxyribose, revealed that the di- and trihydroxy substituted 1*H*-benzimidazole-2-yl hydrazones **5b–e** were the most potent scavengers within the whole series. The strongest protection effect was observed with compounds **5d** and **5e** – denoting extent of molecular damage close or lower than 50%, respectively, and compound **5b** – having similar or even ameliorated

protection effect compared to the reference quercetin. In this way, it was established that the presence of more than one hydroxyl group especially in the case when occupying positions next to each other in the phenyl moiety of the studied 1*H*-benzimidazole-2-yl hydrazones is beneficial for the manifestation of both anthelmintic and radical scavenging activity.

The molecular geometry of the synthesized hydrazones was studied by computational methods, suggesting that the *E, s-trans* isomer is the favorable one. Regarding the possible tautomeric forms (amino and imino forms), it was found that the imino form is strongly preferred in all compounds containing hydroxyl group in *ortho* position of the phenyl ring, while the amino form is the favourable one for all others. In order to clarify the radical-scavenging activity of the 1*H*-benzimidazole-2-yl hydrazones, the reaction enthalpies for HAT, SET-PT and SPLET mechanisms of all compounds **5a–l** were computed in gas phase, benzene and water at B3LYP/6-311++G^{**} level of theory. In nonpolar medium, it could be concluded that the most probable mechanism is HAT. However, in polar medium due to the lower proton affinities of the compounds than the BDEs, SPLET mechanism is favored over HAT. Based on the gained computational data the derivatives possessing vicinal hydroxyl groups are expected to show the highest activity which could explain the results obtained from the *in vitro* assays. It was also shown that the studied compounds are capable to react with various free radical through several possible reaction pathways, including RAF, with C13 position being the most active for radical adduct formation.



4. Experimental

4.1. Reagents and materials

The *o*-phenylenediamine, 98% and the benzaldehydes (2-hydroxybenzaldehyde, 97%, 2,3-dihydroxybenzaldehyde, 98%, 2,4-dihydroxybenzaldehyde, 98%, 3,4-dihydroxybenzaldehyde, 98%, 2,3,4-trihydroxybenzaldehyde, 98%, 4-methoxybenzaldehyde, 98%, 2,6-dimethoxybenzaldehyde, 98%, 3,5-dimethoxybenzaldehyde, 98%, 3,4,5-trimethoxybenzaldehyde, 98%, 2-hydroxy-3-methoxybenzaldehyde, 99%, 2-hydroxy-4-methoxybenzaldehyde, 99%, 3-hydroxy-4-methoxybenzaldehyde, 99%) used for the preparation of the target benzimidazolyl hydrazones were obtained from Alfa-Aesar (United Kingdom) and Sigma Aldrich.

The progress of the reaction was monitored by thin layer chromatography (TLC) on standard silica gel pre-coated plates (60 F254, 0.25 mm, Merck KGaA, Darmstadt, Germany) and eluted by benzene–methanol (3 : 1, v/v).

The melting points (mp) were measured by Büchi B-540 instrument (Büchi Labortechnik AG, Flawil, Switzerland) and were uncorrected. The IR spectra of the synthesized compounds in solid state were recorded on a Bruker Tensor 27 FT spectrometer in ATR (attenuated total reflectance) mode with a diamond crystal accessory. The spectra were referenced to air as a background by accumulating 64 scans, at a resolution of 2 cm⁻¹. ¹H and ¹³C NMR spectra were measured on a Bruker Avance II⁺ 600 MHz NMR instrument in deuterated dimethyl sulfoxide-d₆ (DMSO-d₆) as a solvent, at room temperature. Chemical shifts (δ) are reported in parts per million (ppm), coupling constants (J) – in Hz, and splitting patterns are denoted as follows: s (singlet), d (doublet), t (triplet), m (multiplet).

4.2. Synthesis

4.2.1. Synthesis of 1*H*-benzimidazol-2-yl-thiol 2. To a solution of potassium hydroxide (0.09 mol) in ethanol (95 ml) and carbon disulphide (0.09 mol) was added in portions *o*-phenylenediamine (0.08 mol) and water (15 ml). The reaction mixture has been refluxed for 3 h. Charcoal was added cautiously and the solution has been refluxed 10 more minutes followed by filtration and removing of the charcoal. The filtrate was heated to 60–70 °C and 95 ml warm water and 50% acetic acid (15 ml) was added by good stirring. After cooling the solution in refrigerator for 3 h the product the crystallization was completed. The 1*H*-benzimidazol-2-yl-thiol 2 was prepared as described in previous papers.^{18,22,23}

Yield: 75%; R_f = 0.65 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3153 ($\nu_{\text{N-H}}$); 3114 ($\nu_{\text{Ar-H}}$); 2559 ($\nu_{\text{S-H}}$); 1623, 1511, 1466 ($\nu_{\text{C-C-Ar}}$); 1178 ($\nu_{\text{C-S}}$); 743 (δ_{Ar}).

4.2.2. Synthesis of 1*H*-benzimidazol-2-yl-sulfonic acid 3. To a boiling solution of 1*H*-benzimidazol-2-yl-thiol 2 (0.05 mol) in water (40 ml) was added 50% sodium hydroxide (17 ml) followed by addition of potassium permanganate (0.11 mol) in 275 ml water in small portions by stirring. After the complete addition of potassium permanganate the reaction solution was refluxed for 45 min more. The formed manganese dioxide was filtered off and hydrochloric acid was added to the filtrate to reach pH = 2, by cooling. The obtained product of the reaction

was filtered off and washed with water. The reaction was conducted as described in previous papers.^{18,22,23}

Yield: 60%; R_f = 0.20 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3080, 3059, 3014 ($\nu_{\text{Ar-H}}$), 1631 ($\delta_{\text{N-H}}$), 1619 ($\nu_{\text{C=N}}$), 1192 ($\nu_{\text{Sao}}^{\text{as}}$), 1057 ($\nu_{\text{Sao}}^{\text{as}}$), 653 ($\nu_{\text{S-O}}$).

4.2.3. Synthesis of 1*H*-benzimidazole-2-yl-hydrazine 4. 1*H*-Benzimidazole-2-yl-hydrazine 4 was prepared by refluxing a solution of 1*H*-benzimidazolyl-2-sulfonic acid 3 (0.0176 mol) and hydrazine hydrate in excesses (99%, 0.53 mol, 26 ml) for 3 h according to the previously described procedure.²⁴ After cooling the reaction mixture using an ice bath, the product crystallized, it was filtered off and washed with cold water.

Yield: 76%; mp. 221.8–223.9 °C; R_f = 0.29 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3314, 3273 ($\nu_{\text{N-H}}$), 3091, 2993 ($\nu_{\text{Ar-H}}$), 1664 ($\delta_{\text{N-H}}$), 1620 ($\nu_{\text{C=N}}$), 1587, 1556 ($\nu_{\text{C-C-Ar}}$).

4.2.4. General procedure for preparation of 2-[2-benzylidenehydrazinyl]-1*H*-benzimidazole 5a–l. To a solution of 1*H*-benzimidazol-2-yl-hydrazine 4 (0.002 mol) in absolute ethanol (99%, 5 ml) was added the respective aromatic hydroxyl- and methoxy-aldehyde (0.002 mol). The reaction mixture was refluxed for 3–4 h while the reaction was monitored using TLC (benzene/methanol = 3 : 1). The solid product was filtered off and recrystallized from ethanol.

4.2.4.1. 2-(2-Hydroxybenzylidene)-1-(1*H*-benzimidazol-2-yl)hydrazine (5a). The mixture was refluxed for 3 hours. Yield: 71%; mp. 275.1–277.7 °C; R_f = 0.69 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3351 ($\nu_{\text{O-H}}$), 3285 ($\nu_{\text{N-H}}$), 3063, 3046, 3019 (ν_{Charom}), 1627 (δ_{NH}), 1608 ($\nu_{\text{C=N}}$), 1245 ($\nu_{\text{Ar-O}}$); 727 ($\gamma_{\text{C-H}}$); ¹H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.50 (bs, 1H, NH), 11.30 (bs, 1H, NH), 10.19 (s, 1H, OH), 8.31 (s, 1H, N=CH), 7.79–7.77 (d, J = 7.42 Hz, 1H, Ar), 7.21–7.17 (m, 3H, Ar-bz, Ar), 6.98–6.92 (m, 2H, Ar), 6.89–6.86 (m, 2H, Ar-bz); ¹³C NMR (150 MHz, DMSO-d₆) δ (ppm): 156.28, 130.37, 127.73, 121.07, 119.85, 119.69, 116.40, 109.85.

4.2.4.2. 2-(2,3-Dihydroxybenzylidene)-1-(1*H*-benzimidazol-2-yl)hydrazine (5b). The mixture was refluxed for 3 hours. Yield: 70%; mp. 298.0–300.4 °C; R_f = 0.25 (benzene : methanol = 4 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3403 ($\nu_{\text{O-H}}$), 3382 ($\nu_{\text{N-H}}$), 3060 (ν_{Charom}), 1628 (δ_{NH}), 1614 ($\nu_{\text{C=N}}$), 1262, 1212 ($\nu_{\text{Ar-O}}$); 720, 710 ($\gamma_{\text{C-H}}$); ¹H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.50 (bs, 1H, NH), 9.33 (s, 1H, OH), 8.30 (s, 1H, N=CH), 7.21–7.18 (m, 3H, Ar-bz; Ar), 6.95 (s, 2H, Ar-bz), 6.78–6.76 (dd, J = 7.75 Hz, 1.52 Hz, 1H, Ar), 6.70–6.67 (t, J = 15.54, 7.72 Hz, 1H, Ar); ¹³C NMR (150 MHz, DMSO-d₆) δ (ppm): 145.92, 145.01, 121.50, 119.42, 118.31, 116.15.

4.2.4.3. 2-(2,4-Dihydroxybenzylidene)-1-(1*H*-benzimidazol-2-yl)hydrazine (5c). The mixture was refluxed for 4 hours. Yield: 40%; mp. 284.5–286.8 °C; R_f = 0.66 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3401 ($\nu_{\text{O-H}}$), 3338 ($\nu_{\text{N-H}}$), 3056 (ν_{Charom}), 1626 (δ_{NH}), 1610 ($\nu_{\text{C=N}}$), 1219, 1219, 1261 ($\nu_{\text{Ar-O}}$), 732 ($\gamma_{\text{C-H}}$); ¹H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.39 (s, 1H, NH), 10.95 (s, 1H, NH), 10.19 (s, 1H, OH), 9.70 (s, 1H, OH), 8.19 (s, 1H, N=CH), 7.52–7.51 (d, J = 8.98 Hz, 1H, Ar), 7.18–7.15 (m, 2H, Ar), 6.95–6.89 (m, 2H, Ar-bz), 6.32–6.30 (m, 2H, Ar); ¹³C NMR (150 MHz, DMSO-d₆) δ (ppm): 159.94, 157.95, 129.38, 112.63, 107.92, 102.81.

4.2.4.4. 2-(3,4-Dihydroxybenzylidene)-1-(1*H*-benzimidazol-2-yl)hydrazine (5d). The mixture was refluxed for 4 hours and 30



minutes. Yield: 50%; mp. 215.1–217.3 °C; R_f = 0.20 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3407 ($\nu_{\text{O-H}}$), 3257 ($\nu_{\text{N-H}}$), 3057 (ν_{CHarom}), 1635 (δ_{NH}), 1606 ($\nu_{\text{C=N}}$), 1268 ($\nu_{\text{Ar-O}}$), 744, 751 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.46 (bs, 1H, NH), 9.28 (bs, 2H, OH), 7.85 (s, 1H, N=CH), 7.24 (d, J = 2.09 Hz, 1H, Ar), 7.23–7.21 (d, J = 8.37 Hz, 2H, Ar-bz), 7.02–7.01 (dd, J = 8.16 Hz, 2.02 Hz, 1H, Ar), 6.93 (s, 2H, Ar-bz), 6.78–6.76 (d, J = 8.10 Hz, 1H, Ar); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 154.00, 147.22, 145.98, 141.88, 127.13, 119.46, 115.87, 113.78.

4.2.4.5. 2-(2,3,4-Trihydroxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5e). The mixture was refluxed for 4 hours. Yield: 56%; mp. 248.5–250.5 °C; R_f = 0.27 (benzene : methanol = 4 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3468, 3332 ($\nu_{\text{O-H}}$), 3252 ($\nu_{\text{N-H}}$), 3062 (ν_{CHarom}), 1631 (δ_{NH}), 1612 ($\nu_{\text{C=N}}$), 1212, 1266 ($\nu_{\text{Ar-O}}$), 730 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.46 (s, 1H, NH), 9.87 (bs, 1H, OH), 9.42 (bs, 1H, OH), 8.14 (s, 1H, N=CH), 7.17–7.16 (m, 2H, Ar-bz), 6.96–6.92 (m, 3H, Ar-bz, Ar), 6.37–6.36 (d, J = 8.47 Hz, 1H, Ar); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 153.76, 147.92, 146.55, 133.13, 119.30, 113.14, 107.87.

4.2.4.6. 2-(4-Methoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5f). The mixture was refluxed for 3 hours. Yield: 63%; mp. 241.7–243.8 °C; R_f = 0.46 (benzene : methanol = 4 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3388 ($\nu_{\text{N-H}}$), 3060, 3030 (ν_{CHarom}), 2927 ($\nu_{\text{CH}_3}^{\text{as}}$), 2835 ($\nu_{\text{CH}_3}^{\text{s}}$), 1651 (δ_{NH}), 1609 ($\nu_{\text{C=N}}$), 1458 ($\delta_{\text{CH}_3}^{\text{as}}$), 1378 ($\delta_{\text{CH}_3}^{\text{s}}$), 1244, 1136 ($\nu_{\text{C-O-C}}$), 728, 714 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.43 (bs, 2H, NH), 7.97 (s, 1H, N=CH), 7.76–7.73 (m, 2H, Ar), 7.24–7.20 (m, 2H, Ar), 7.00–6.96 (m, 2H, Ar-bz), 6.97–6.92 (m, 2H, Ar-bz), 3.80 (s, 3H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 160.38, 153.98, 140.99, 128.37, 114.55, 55.70.

4.2.4.7. 2-(2,6-Dimethoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5g). The mixture was refluxed for 3 hours. Yield: 75%; mp. 227.5–229.7 °C; R_f = 0.72 (benzene : methanol = 4 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3321 ($\nu_{\text{N-H}}$), 3105, 3054, 3019 (ν_{CHarom}), 2946 ($\nu_{\text{CH}_3}^{\text{as}}$), 2847 ($\nu_{\text{CH}_3}^{\text{s}}$), 1640 (δ_{NH}), 1605 ($\nu_{\text{C=N}}$), 1459 ($\delta_{\text{CH}_3}^{\text{as}}$), 1384 ($\delta_{\text{CH}_3}^{\text{s}}$), 1253, 1086 ($\nu_{\text{C-O-C}}$), 743, 727 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.26 (bs, 1H, NH), 11.15 (bs, 1H, NH), 8.16 (s, 1H, N=CH), 7.32–7.28 (t, J = 16.75 Hz, 8.38 Hz, 1H, Ar), 7.23–7.20 (dd, J = 7.71 Hz, 2.85 Hz, 2H, Ar-bz), 6.97–6.94 (m, 1H, Ar-bz), 6.91–6.87 (m, 1H, Ar-bz), 6.72–6.69 (d, J = 8.44 Hz, 2H, Ar), 3.81 (s, 6H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 158.80, 153.64, 136.83, 133.76, 130.76, 120.91, 119.33, 115.62, 111.98, 110.14, 104.66, 56.35.

4.2.4.8. 2-(3,5-Dimethoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5h). The mixture was refluxed for 3 hours. Yield: 66%; mp. 251.7–253.8 °C; R_f = 0.73 (benzene : methanol = 4 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3389 ($\nu_{\text{N-H}}$), 3060 (ν_{CHarom}), 2960 ($\nu_{\text{CH}_3}^{\text{as}}$), 2838 ($\nu_{\text{CH}_3}^{\text{s}}$), 1647 (δ_{NH}), 1638 ($\nu_{\text{C=N}}$), 1458 ($\delta_{\text{CH}_3}^{\text{as}}$), 1357 ($\delta_{\text{CH}_3}^{\text{s}}$), 1266, 1127 ($\nu_{\text{C-O-C}}$), 733 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.56 (bs, 1H, NH), 11.48 (bs, 1H, NH), 7.94 (s, 1H, N=CH), 7.27–7.22 (m, 2H, Ar-bz), 6.99–6.92 (m, 4H, Ar-bz, Ar), 6.50–6.49 (m, 1H, Ar), 3.80 (s, 6H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 161.11, 153.83, 141.01, 137.56, 133.62, 121.03, 119.73, 115.52, 109.81, 104.92, 101.30, 55.81.

4.2.4.9. 2-(3,4,5-Trimethoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5i). The mixture was refluxed for 3 hours. Yield: 60%; mp. 216.1–218.5 °C; R_f = 0.4 (benzene : methanol = 4 : 1,

v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3380 ($\nu_{\text{N-H}}$), 3060, 3005 (ν_{CHarom}), 2940 ($\nu_{\text{CH}_3}^{\text{as}}$), 2834 ($\nu_{\text{CH}_3}^{\text{s}}$), 1655 (δ_{NH} , $\nu_{\text{C=N}}$), 1612 ($\nu_{\text{C=N}}$), 1462 ($\delta_{\text{CH}_3}^{\text{as}}$), 1359 ($\delta_{\text{CH}_3}^{\text{s}}$), 1228, 1121 ($\nu_{\text{C-O-C}}$), 728 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.55 (bs, 2H, NH), 7.95 (s, 1H, N=CH), 7.27–7.25 (m, 2H, Ar-bz), 7.11 (s, 2H, Ar), 6.99–6.95 (m, 2H, Ar-bz), 3.86 (s, 6H, OCH₃), 3.69 (s, 3H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 153.76, 141.38, 138.76, 131.08, 104.35, 60.58, 56.52.

4.2.4.10. 2-(2-Hydroxy-3-methoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5j). The mixture was refluxed for 2 hours and 30 minutes. Yield: 74%; mp. 252.7–254.4 °C; R_f = 0.74 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3363 ($\nu_{\text{O-H}}$), 3305 ($\nu_{\text{N-H}}$), 3061, 2997 (ν_{CHarom}), 2965 ($\nu_{\text{CH}_3}^{\text{as}}$), 2836 ($\nu_{\text{CH}_3}^{\text{s}}$), 1626 (δ_{NH}), 1606 ($\nu_{\text{C=N}}$), 1479 ($\delta_{\text{CH}_3}^{\text{as}}$), 1369 ($\delta_{\text{CH}_3}^{\text{s}}$), 1262 ($\nu_{\text{Ar-O}}$); 1240, 1065 ($\nu_{\text{C-O-C}}$), 738, 718 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.52 (bs, 1H, NH), 8.32 (s, 1H, N=CH), 7.41–7.40 (d, J = 7.70 Hz, 1H, Ar), 7.20–7.19 (m, 2H, Ar-bz), 6.95–6.93 (m, 3H, Ar-bz, Ar), 6.83–6.81 (t, J = 15.78 Hz, 7.85 Hz, 1H, Ar), 3.82 (s, 3H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 153.92, 148.32, 145.82, 121.45, 119.34, 112.55, 56.32.

4.2.4.11. 2-(2-Hydroxy-4-methoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5k). The mixture was refluxed for 4 hours. Yield: 53%; mp. 251.0–253.1 °C; R_f = 0.76 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3402 ($\nu_{\text{O-H}}$), 3236 ($\nu_{\text{N-H}}$), 3068, 3054 (ν_{CHarom}), 2963 ($\nu_{\text{CH}_3}^{\text{as}}$), 2838 ($\nu_{\text{CH}_3}^{\text{s}}$), 1656 (δ_{NH}), 1625 ($\nu_{\text{C=N}}$), 1460 ($\delta_{\text{CH}_3}^{\text{as}}$), 1378 ($\delta_{\text{CH}_3}^{\text{s}}$), 1267 ($\nu_{\text{Ar-O}}$); 1222, 1027 ($\nu_{\text{C-O-C}}$), 730, 713 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.44 (bs, 1H, NH), 11.15 (bs, 1H, NH), 10.38 (bs, 1H, OH), 8.23 (s, 1H, N=CH), 7.64–7.63 (d, J = 8.49 Hz, 1H, Ar), 7.17 (s, 2H, Ar-bz), 6.94–6.93 (d, J = 14.77 Hz, 2H, Ar-bz), 6.49–6.48 (d, J = 8.66 Hz, 1H, Ar), 6.45 (s, 1H, Ar), 3.79 (s, 3H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 161.42, 157.88, 153.94, 129.27, 121.04, 114.06, 106.54, 101.34, 55.62.

4.2.4.12. 2-(3-Hydroxy-4-methoxybenzylidene)-1-(1H-benzimidazol-2-yl)hydrazine (5l). The mixture was refluxed for 3 hours. Yield: 55%; mp. 267.1–269.5 °C; R_f = 0.5 (benzene : methanol = 3 : 1, v/v); IR ($\nu_{\text{max}}/\text{cm}^{-1}$): 3382 ($\nu_{\text{O-H}}$), 3265 ($\nu_{\text{N-H}}$), 3058, 3007 (ν_{CHarom}), 2954 ($\nu_{\text{CH}_3}^{\text{as}}$), 2834 ($\nu_{\text{CH}_3}^{\text{s}}$), 1629 (δ_{NH}), 1602 ($\nu_{\text{C=N}}$), 1461 ($\delta_{\text{CH}_3}^{\text{as}}$), 1375 ($\delta_{\text{CH}_3}^{\text{s}}$), 1274 ($\nu_{\text{Ar-O}}$); 1233, 1094 ($\nu_{\text{C-O-C}}$), 733 ($\gamma_{\text{C-H}}$); ^1H NMR (600 MHz, DMSO-d₆) δ (ppm): 11.51 (bs, 1H, NH), 7.89 (s, 1H, N=CH), 7.33–7.32 (d, J = 1.93 Hz, 1H, Ar), 7.22 (s, 2H, Ar-bz), 7.11–7.09 (dd, J = 8.24 Hz, 1.99 Hz, 1H, Ar), 6.96–6.92 (m, 3H, Ar-bz, Ar), 3.80 (s, 3H, OCH₃); ^{13}C NMR (150 MHz, DMSO-d₆) δ (ppm): 153.95, 149.19, 147.05, 141.53, 133.73, 128.54, 120.95, 119.45, 115.50, 113.13, 112.15, 109.79, 56.10.

4.3. *In vitro* anthelmintic assay

Benzimidazole derivatives **5a–l** were tested for anthelmintic activity against *T. spiralis* muscle larvae (ML) provided from The National Centre for Infectious and Parasitic Diseases, Department of Parasitology and Tropical medicine, Sofia, Bulgaria.

In brief, freshly decapsulated larvae of the parasite (100 specimens for 1 ml physiological solution) were exposed to 50 and 100 $\mu\text{g ml}^{-1}$ concentrations of experimental compounds dissolved in dimethyl sulfoxide (DMSO), and were incubated at



37 °C in "humid" chamber with a thermostat. Albendazole and ivermectin were used in this test as standard anthelmintics. The microscopic observations for vitality of *T. spiralis* ML were carried out after 24 h and 48 h, respectively. The efficacy (%) of newly synthesized 1*H*-benzimidazol-2-yl hydrazones on the viability of the parasites was calculated as the average value of three experiments as previously described.¹⁸

$$\text{Efficacy\%} = (\text{the number of dead parasite larvae/total parasite larvae}) \times 100\%$$

4.4. Estimation of the radical-scavenging activity and protective effect

4.4.1. Stable free radicals containing model systems. Two model systems containing alternative biologically irrelevant indicators for scavenging activity respectively ABTS^{•+} (2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonate radical cation)) or DPPH[•] (2,2-diphenyl-1-picrylhydrazyl stable radical) have been chosen. Both assays are operationally simple, relatively cheap and fast. They are based on the spectrophotometric estimation of the absorbance change of the chromophore preformed solutions. On the base of the observed decrease of the absorbance value of the ABTS or DPPH working solution, after interaction with the new designed compounds have been made conclusions concerning the capability of the potential antioxidants to eliminate both radicals. The obtained data help to make suggestions concerning the capability of the tested hydrazones to participate in reactions *via* hydrogen atom transfer (HAT), single-electron transfer (SET-PT) and sequential proton loss electron transfer (SPLET). Before each experiment fresh stock and working solution of each radical have been prepared. Both methods comprise the preparation of the two following types of measurement compositions having identical final concentration of the used in the system radical: (1) controls where the tested hydrazones have been omitted and (2) samples containing the investigated compounds and the used in the system stable free radical.

ABTS assay – the experiments were performed as described by Re *et al.*⁷⁶ As first step ABTS^{•+} stock solution was prepared by mixing 14 mM ABTS and potassium persulfate 2.45 mM (final concentration). The resulting mixture was allowed to stay in the dark for 12–16 h (until a primary solution of the radical with stable absorbance has been obtained). The suspension was then diluted until obtaining final working solution with absorbance 0.70 ± 0.01 units at 734 nm. The reading of the absorbance values has been done exactly 1 hour after mixing the ABTS radical working solution with different concentrations of the tested compounds.

DPPH assay – the experimental procedure was performed according to Grupy *et al.*⁷⁷ a standard purple colored working solution with absorbance of 1 at 517 nm (characteristic band for DPPH[•]) was prepared in ethanol. Two milliliters of the DPPH radical solution were allowed to react with different concentrations of the tested hydrazones for 1 hour. After the incubation the relative decrease in the absorbance at 517 nm was registered and the radical scavenging activity was calculated.

On the base of the obtained data has been estimated the radical scavenging activity (RSA %) calculated in the following way:

$$\text{RSA\%} = ((A_{\text{control}} - A_{\text{sample}})/A_{\text{control}}) \times 100$$

where A_{control} and A_{sample} represent the corresponding absorbance values for the control and the sample measurement composition using both methods. The first parameter helped to estimate the absorbance of the characteristics signal of the stable free radical and the second its relative decrease due to the ongoing free radical processes. The determined RSA% for the individual sample along with the number of the performed measurements, were used for statistical estimation of the relative differences between the scavenging activities of the tested compounds.

4.4.2. Model systems based on ferrous iron induced oxidative molecular damage. The thiobarbituric acid reactive-substances (TBA-RS) assay has been used to estimate the modulation effect of the compounds on the peroxidation of biologically important molecules. The experiments were performed according to Asakawa and Matsushita and are based on reaction between the secondary end product of the oxidation process with two molecules of thiobarbituric acid (TBA).⁷⁸ Two alternative variations of the methods containing different oxidizable substrates (lecithin or deoxyribose) have been applied. The induced oxidative damage was initiated by FeCl₂ (0.1 mmol l⁻¹). Two types of measurement compositions have been prepared: (i) controls containing the biologically important molecules and the agent used to induce oxidative damage and (ii) samples containing also the tested hydrazones. All samples have been prepared in phosphate buffer (pH 7.4) with equivalent concentrations of the substrate (lecithin (1 mg ml⁻¹) or deoxyribose (0.5 mmol l⁻¹)). The samples have been incubated at 37 °C for 30 min. After the incubation, 2.5 ml of 0.5% thiobarbituric acid and 0.5 ml of 2.8% trichloroacetic acid were added. Another incubation at 100 °C in a water bath for 20 min was performed, following by centrifugation at 3000 rpm for 20 min. The absorbance of the generated pinkish-red chromophore was estimated at 532 nm (characteristic band). The results were presented as "extent of molecular damage" calculated as percentage of the treated only with ferrous iron control samples.

4.5. Computational methods

Full geometry optimizations have been performed with the Gaussian 09 software package⁷⁹ employing density functional theory (DFT) with the hybrid functional B3LYP⁸⁰ and the 6-311++G** basis set.^{81,82} Solvent effects in water and benzene were included in the optimizations by using the Polarizable Continuum Model (PCM).⁸³ Harmonic vibrational frequencies have been calculated for all located stationary structures to verify whether they are minima or transition states. Zero-point energies and thermal corrections have been taken from unscaled vibrational frequencies. Single-point energy calculations in gas phase and single-point PCM energy calculations, in water and benzene, were performed at the M06-2X/6-311++G**⁸⁴



levels of theory. All energies are in kJ mol^{-1} and were calculated at 25 °C. All bond lengths are reported in Å.

Dissociation enthalpy (BDE), ionization potential (IP), proton dissociation enthalpy (PDE), proton affinity (PA), and electron transfer enthalpy (ETE) of the most stable conformers will be calculated according eqn (1)–(5).⁶⁰

$$\text{BDE} = \text{H}(\text{A}^\cdot) + \text{H}(\text{H}^\cdot) - \text{H}(\text{A}-\text{H}) \quad (1)$$

$$\text{IP} = \text{H}(\text{A}^{+\cdot}) + \text{H}(\text{e}^-) - \text{H}(\text{A}-\text{H}) \quad (2)$$

$$\text{PDE} = \text{H}(\text{A}^\cdot) + \text{H}(\text{H}^+) - \text{H}(\text{A}^{+\cdot}) \quad (3)$$

$$\text{PA} = \text{H}(\text{A}^-) + \text{H}(\text{H}^+) - \text{H}(\text{A}-\text{H}) \quad (4)$$

$$\text{ETE} = \text{H}(\text{A}^\cdot) + \text{H}(\text{e}^-) - \text{H}(\text{A}^-) \quad (5)$$

The enthalpy of hydrogen atom, $\text{H}(\text{H})$, for each solvent are obtained by the same method and basis set. All reaction enthalpies will be calculated for 25 °C. The enthalpies of proton $\text{H}(\text{H}^+)$, and electron, $\text{H}(\text{e}^-)$, in gas phase: 6.197 kJ mol^{-1} and 3.145 kJ mol^{-1} , were applied in accordance with previously estimated values.⁶⁰ The respective enthalpies of hydrated proton, $\text{H}(\text{H}^+)$, and hydrated electron, $\text{H}(\text{e}^-)$, were used also as reported in the literature:⁶⁰ $\text{H}(\text{H}^+) = -1083.5 \text{ kJ mol}^{-1}$ and $\text{H}(\text{e}^-) = -232.8 \text{ kJ mol}^{-1}$.

Author contributions

M. A., D. Y. and K. A. conceived and designed the study, M. A. performed the chemical synthesis, N. A. performed radical-scavenging activity experiments, D. V. and G. P.-D. performed parasitological experiments, M. A., M. G. and D. Y. performed theoretical calculations. All the authors contributed to the paper writing.

Conflicts of interest

The authors declare that there is no conflict of interests related to this study.

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