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ANALYSIS OF ACID-BASE PROPERTIES OF SOME p-SUBSTITUTED AROMATIC HYDRAZONES IN AQUEOUS PERCHLORIC ACID BY SPECTROPHOTOMETRIC AND SEMIEMPIRICAL METHODS

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The acid-base properties of some p-substituted aromatic hydrazones were examined in aqueous perchloric acid medium by spectrophotometric methods. The changes in the UV spectra occurred due to the protonation reaction of the investigated hydrazones. The possible site where protonation may take place was discussed using the values of the total energy, binding energy, heat of formation and proton affinity, calculated according to the semiempirical methods AM1 and PM3. Furthermore, the stability of the protonated forms, as well as the stability of the isomers (E and E) was discussed. Electronic absorption studies were utilized to determine the dissociation constants of the protonated form ($pK_{\rm BH}^+$), numerically and graphically, at different ionic strengths (0.1, 0.25 and 0.5 mol/dm³), as well as the thermodynamic dissociation constants. The obtained values ranged between 3.08 and 3.47. The influence of the solvent on the appearance of the spectra was compensated by the use of the method of characteristic vector analysis (CVA).

Keywords: UV spectroscopy; protonation; thermodynamic dissociation constants; *p*-substituted aromatic hydrazones; AM1 and PM3 semiempirical methods

АНАЛИЗА НА КИСЕЛИНСКО-БАЗНИТЕ СВОЈСТВА НА НЕКОИ p-СУПСТИТУИРАНИ АРОМАТИЧНИ ХИДРАЗОНИ ВО ВОДНИ РАСТВОРИ ОД ПЕРХЛОРНА КИСЕЛИНА СО СПЕКТРОФОТОМЕТРИСКИ И СЕМИЕМПИРИСКИ МЕТОДИ

Киселинско-базните својства на некои p-супституирани ароматични хидразони се испитани во водни раствори од перхлорна киселина со спектрофотометриски методи. Како резултат на реакцијата на протонирање се забележани промени во UV-спектрите. Местото каде протонирањето може да се одвива е дискутирано со користење на вредностите на вкупната енергија, сврзувачката енергија, топлината на формирање и протонскиот афинитет, пресметани со примена на семиемпириските методи AM1 и PM3. Исто така е дискутирана стабилноста на протонираните форми, како и стабилноста на изомерите (E и Z). Електронските апсорпциони спектри се искористени за определување на константите на дисоцијација на протонираната форма ($pK_{\rm BH}^+$), нумерички и графички, при различна јонска сила (0,1, 0,25 и 0,5 mol/dm³), како и термодинамичките константи на дисоцијација.

Добиените вредности се движат во границите од 3,08 до 3,47. Влијанието на растворувачот на изгледот на спектрите е компензирано со примена на методот на карактеристична векторска анализа (CVA).

Клучни зборови: UV-спектроскопија; протонирање; термодинамички константи на дисоцијација; *р*-супституирани ароматични хидразони; семиемпириските методи AM1 и PM3

1. INTRODUCTION

Hydrazones are considered one of the most important classes of organic compounds. Their derivatives constitute an important class of organic compounds showing diverse biological activity such as: anticonvulsant, antidepressant, analgesic, antiinflammatory, antiplatelet, antimicrobial, antimalarial, antitumoral, antiviral, vasodilator and antitubeculosis [1–6]. This activity of hydrazones and their derivatives is a result of having an azometine proton (-NH-N=CH-). As a result of the presence of this proton, hydrazones find wide applications in many areas. They can also be used in agriculture as herbicides, insecticides and plant growth stimulants [7]. Furthermore, they are used in synthetic organic chemistry for the preparation of other compounds, and in analytical chemistry for the identification of carbonyl compounds [8, 9]. The hydrazones also find applications as spectrophotometric reagents, because they can act as multidentate ligands with transition metals [10].

Acidity measurements of organic compounds, like those investigated here, are significant in studies that involve the formation and investigation of biological activities. An understanding of the acidity or basicity of organic compounds is fundamental to molecular design. The reaction mechanisms are also very important for understanding the chemical and biological processes that may take place at the azometine site [11]. It is well-known that acid-base properties affect the toxicity and pharmaceutical characteristics of organic acids and bases [12]; the biological activity of hydrazones depends on the ionic forms in which they exist in solution [13].

Moreover, knowledge of the ionization constant values of ionizable groups is important for understanding of many areas of chemistry. It should be pointed out that the values of these constants are important, especially for elucidating the

reaction mechanism, especially those involving proton transfers. There are several different methods, such as potentiometric, chromatographic, electrophoretic and spectrophotometric methods, that can be used for the determination of the ionization constants of compounds that behave as weak bases in acidic media. It is known that spectrophotometric methods are highly sensitive and suitable for studying chemical equilibria in solution [14].

The proton-acceptor and proton-donor capacities of drug molecules are significant factors in the functioning of biological systems. With knowledge of the ionization constants, one can calculate the ratio of the molecules with different protonation sites at various pH values in certain compartments of the body. It is also important to know the average protonation state of the molecule when designing in vitro chemical reactions and analytical procedures [15]. So far, no precise data have been available in the literature referring to the sites where the molecule loses and accepts protons. However, the recent development of a calculation method in quantum chemistry has facilitated a new approach to the analysis of acid-base properties of biologically active compounds [16].

In view of the above observations, this work focused on following the behavior of some p-substituted aromatic hydrazones in acidic media (pH range from 1 to 7), as well as on the determination of the ionization constants of their protonated forms (p $K_{\rm BH}^+$). Moreover, the thermodynamic ionization constants were evaluated from the p $K_{\rm BH}^+$ values determined at different ionic strengths. Additionally, applying the semiempirical methods AM1 and PM3, the proton affinities, site of protonation and stability of the protonated forms and that of the isomers (E and E) were determined.

The *p*-substituted aromatic hydrazones that were the subject of this study have the general structural formula shown in Table 1.

Table 1

The structure and nomenclature of the investigated hydrazones

	R NH—N=CH—OCH3	Substituents
Compound	IUPAC name	
H_1	N-benzaldehyde-p-metoxybenzoilhydrazone	–H
H_2	N- p -methylbenzaldehyde- p -metoxybenzoilhydrazone	$-CH_3$
H_3	N-p-metoxybenzaldehyde-p-metoxybenzoilhydrazone	$-OCH_3$
H_4	N-p-chlorobenzaldehyde-p-metoxybenzoilhydrazone	–Cl
H ₅	$\emph{N-p}$ -hydroxybenzaldehyde- \emph{p} -metoxybenzoilhydrazone	-ОН

These hydrazones were synthesized in our laboratory and their structures were confirmed by several methods, including UV, IR, ¹H NMR, ¹³C NMR and elemental analysis [17].

2. EXPERIMENTAL

2.1. Solutions and measurements

Stock solutions were prepared by dissolving an accurate weight of the compound in the appropriate volume of 98% ethanol. The concentration of the investigated hydrazones in these solutions was about $1\cdot 10^{-3}$ mol/dm³. The stock solutions were stable over a long period of time, under ordinary conditions. The test solutions of individual hydrazones were prepared with redistilled water; the concentration of hydrazone was approximately $3\cdot 10^{-5}$ mol/dm³. In order to obtain more precise data, a three series of test solutions were prepared, and $pK_{\rm BH}^+$ values were calculated as the average value of these data.

In all test solutions, sodium perchlorate (1 mol/dm³) was added to adjust the ionic strength (0.1, 0.25 and 0.5 mol/dm³), while the pH was varied in the range between 1 and 7 by adding a suitable amount of perchloric acid (0.5 mol/dm³). After each pH adjustment, the UV spectra were recorded. Simultaneously, the blanks were prepared with the same composition as the working solutions, without the investigated hydrazone.

The purity of the investigated hydrazones was confirmed by elemental analysis, as well as constant measurement of melting points. The perchloric acid and sodium perchlorate were of analytical grade p.a.

Absorbance measurements were carried out using a quartz cell with a 1 cm path length in a Varian Cary 50 UV/Vis spectrophotometer (the maximum scan rate is 24 000 nm/min and resolution is 1.5 nm), immediately after preparation of the solutions, at room temperature.

All absorption spectra were digitized at five data points per nanometer in the wavelength region between 190 and 400 nm. The pH measurements were carried out using a digital pH meter with a combined glass electrode (1 < pH < 13), calibrated with two buffer solutions at pH 4 and pH 7.

2.2. Calculations

 pK_{BH}^{+} values can be calculated from the absorbance value which corresponds to the absorption maximum; more than one absorbance value

can be used. Our previous studies demonstrated that the obtained results are more accurate when four absorbance values are used [18]. Furthermore, the pK_{BH}^{+} values calculated from the absorbance values at four selected wavelengths around the absorption maximum in neutral media are statistically identical with those in acid media. It has been suggested that both maxima can be used for the selection of wavelengths for further calculations [18]. Taking into consideration these results, the calculations of the pK_{BH}^{+} values were performed from the absorbance values from experimental and reconstructed spectra at four selected wavelengths around the absorption maximum that appears in neutral media: 306 nm, 310 nm, 314 nm and 318 nm. To determine the molar absorption coefficients, the absorbance values of the neutral (pH 6.7) and protonated (pH 2.1) form were measured. For this purpose, the UV spectra were recorded at three different concentrations $(2.40 \cdot 10^{-5}, 3.00 \cdot 10^{-5})$ and 3.60 10⁻⁵ mol/dm³) of the investigated hydrazones in solution at ionic strengths 0.1 mol/dm³, 0.25 mol/dm³ and 0.5 mol/dm³. Furthermore, the ionization ratio $I(I = c(BH^+)/c(B))$, i.e. the ratio between the concentration of the protonated (BH⁺) and neutral (B) form of the hydrazones, was determined. The calculations were performed using the molar absorption coefficient values and the spectrophotometric data obtained at the selected wavelengths, in accordance with Beer's law. In order to minimize the influence of random errors, an overdetermined system of four equations (for absorbance values) with two unknown parameters (concentration of the protonated and unprotonated form) was set up. The pK_{BH}^{+} values were calculated using the Equation (1) presented below:

$$pK_{BH}^{+} = n \cdot pH + \log I, \qquad (1)$$

where pK_{BH}^{+} is the dissociation constant of the protonated form, I is the ionization ratio and n is the number of protons.

The influence of the solvent can be eliminated by employing various methods such as principal component analysis (PCA) [19], the target-testing method of factor analysis [20] and characteristic vector analysis (CVA) [21]. One of the most commonly used is CVA, which has been tested and proven to be applicable in the analysis of spectroscopic data, to investigate problems dealing with protonation [22].

Additionally, the p $K_{\rm BH}^+$ values of the investigated hydrazones were determined graphically [23], as the intercept of the dependence of $\log I$ on pH. When $c({\rm BH}^+)=c({\rm B})$, $\log I=0$, the graphically

determined pK_{BH}^+ value is equal to the pH value of the solution. Finally, with extrapolation of the curve $pK_{BH}^+ = f(\sqrt{\mu})$ to zero ionic strength, the thermodynamic pK_{BH}^+ values were evaluated, as the intercept [24].

The semiempirical calculations (heat of formation, proton affinities, total energy, binding energy and Gibbs energies of formation) were performed using semiempirical the AM1 (Austin Model 1) and PM3 (Parametric Method 3) methods [25, 26]. Preliminary optimization was performed by the molecular mechanics MM⁺ with application of the Polak-Ribiere algorithm and a gradient of 0.1 kcal/(A·mol). The same methods were applied to optimize the geometry of the analyzed hydrazones and to determine the stability of the possible isomers.

All the results were obtained using the computer programs Excel, Grams Version 4.10, and HyperChem Version 8.

3. RESULTS AND DISCUSSION

3.1. The absorption spectra and characterization of electron transitions

The UV spectra of N-benzaldehyde-p-meto-xybenzoilhydrazone (\mathbf{H}_1), N-p-methylbenzaldehyde-p-metoxybenzoilhydrazone (\mathbf{H}_2), N-p-metoxybenzaldehyde-p-metoxybenzoilhydrazone (\mathbf{H}_3), N-p-chlorobenzaldehyde-p-metoxybenzoilhydrazone (\mathbf{H}_4) and N-p-hydroxybenzaldehyde-p-metoxybenzoilhydrazone (\mathbf{H}_5) were obtained in aqueous perchloric acid medium (pH from 2 to 6) at room temperature. These spectra were recorded in the wavelength region from 190 to 400 nm, as shown for hydrazone \mathbf{H}_3 (Figure 1). Predictably, similar spectra were obtained for the other investigated hydrazones.

As can be seen from the UV spectra (see Figure 1), it is clear that in the investigated wavelength region, two absorption bands exist in the spectra of the hydrazone $\mathbf{H_3}$, in the pH range from 2 to 6. The first absorption band that appears in the neutral region, at a wavelength around 195 nm ($\varepsilon = 307632 \text{ cm}^{-1} \text{ mol}^{-1} \text{ dm}^3$), is a result of $\pi \rightarrow \pi^*$ electron transitions. Moreover, the second absorption band, found at around 314 nm ($\varepsilon = 344548 \text{ cm}^{-1} \text{ mol}^{-1} \text{ dm}^3$), is due to low energy $n \rightarrow \pi^*$ electron transition of the azometine group, involving the lone pair on the azometine nitrogen atom, and characterizes the neutral form of the compound. This absorption band was interesting for our further investigations.

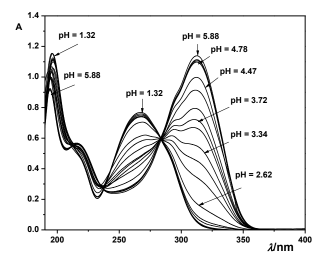


Fig. 1. The experimental UV spectra for *N-p*-meto-xybenzaldehyde-*p*-metoxybenzoilhydrazone ($c = 3.21 \cdot 10^{-5}$ mol/dm³) in perchloric acid media (pH from 1.3 to 5.8) at an ionic strength of 0.1 mol/dm³ (NaClO₄)

Upon increasing the acidity of the solution until pH 4.78, there were no changes in the position and intensity of this absorption band. On the other hand, when the pH was lower than 4.47, the intensity of the absorption band decreased (Figure 1), while when the pH of the solution was below 3.53, despite no change in intensity except intensity, a hypsochromic shift of the absorption band of about 20 nm was observed. In more acidic media (pH 2.44), the absorption band reached the final position (266 nm) and there were no further changes in its intensity. However, as can be seen in Figure 1, acidification of the investigated solutions resulted in an approximate 50 nm hypsochromic shift in the absorption band.

The described changes in the experimental spectra suggested that the reaction of protonation of the investigated hydrazones probably occurred in acidic media. Namely, it is known that this absorption band which is a result of $n \rightarrow \pi^*$ transitions despaired in acidic media when the protonation process of compounds that contain an azometine group take place, i.e. transitions where a proton is added to the azometine group are hindered. The absorption band which appears in acidic media (pH 2.4) at lower wavelengths (around 266 nm) is probably a result of the $\pi \rightarrow \pi^*$ electron transitions of the protonated form of hydrazones. Furthermore, changes were noted in the intensity and the position of the first absorption band. However, as mentioned before, this absorption band was not interesting for our further investigations.

In addition, in Figure 1 it can clearly be seen that isosbestic points can be expected in the expe-

rimental spectra of the hydrazone **H**₃. In order to determine their exact positions, the experimental spectra were reconstructed using the CVA method. The obtained spectra after reconstruction are shown in Figure 2.

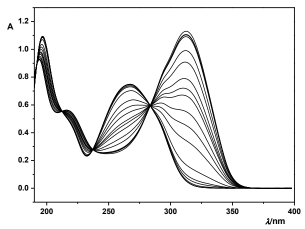


Fig. 2. The reconstructed UV spectra for *N*-*p*-metoxybenzaldehyde-*p*-metoxybenzoilhydrazone $(c = 3.21 \cdot 10^{-5} \text{ mol/dm}^3)$ at an ionic strength of 0.1 mol/dm³ (NaClO₄)

From the reconstructed spectra (see Figure 2), it can be seen that for the hydrazone \mathbf{H}_3 , the isosbestic points around 214, 238 and 286 nm are clearly visible. The position and the intensity of the absorption bands do not change after reconstruction. The existence of the isosbestic points indicates that a reaction is taking place in the studied solutions and there are at least two molecular species present capable of absorbing electromagnetic radiation.

Based on the changes in the UV spectra for all investigated hydrazones, we concluded that the reaction that occurs in acidic media is the protonation of hydrazones. This conclusion was additionally supported by the variation of the absorbance at 314 nm after changing the acidity of the solution. These curves (A = f(pH)) for all investigated hydrazones have a sigmoidal "S" shape (see Figure 3).

The curves shown in Figure 3 have only one step, which is an indication that only one reaction takes place. Its initial part is almost horizontal until a pH of about 5.0. After that, the curves rise until a pH of 2.5 and then the absorbance values remain practically constant. The changes in the sigmoidal curve of the hydrazone $\mathbf{H_3}$ (see Figure 3) showed that formation of the protonated form of this hydrazone occurred when the pH of the solution was below 2.5. The identical situation for the other investigated hydrazones was observed. Using the

"S"-shaped curve, the pH range of protonation of each investigated hydrazone (steep part), and the pH values at which only the neutral or the protonated form (horizontal part) exist, can be determined (see Figure 3).

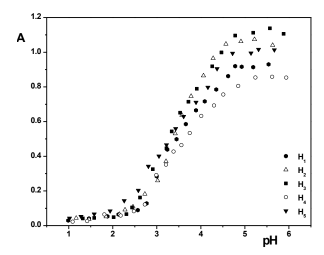


Fig. 3. The absorbance values at $\lambda = 314$ nm, for hydrazones $\mathbf{H_1}$ – $\mathbf{H_5}$ as a function of pH in aqueous perchloric acid media

For the protonation equilibrium of a base, the following equation can be written:

$$B + H_3O^+ = BH^+ + H_2O$$
 (2)

In order to make it applicable to a given reaction only, the exact position of the site at which protonation takes place must be determined. To achieve that, the proton affinities of possible protonation sites and some physicochemical parameters (total energy, binding energy and heat of formation) we calculated, using the semiempirical methods AM1 and PM3.

3.2. Physicochemical parameters and proton affinities

Possible sites of protonation of a hydrazone molecule are the azometine nitrogen atoms *i.e.* the sp² (imino) and sp³ (amino) hybridized nitrogen atoms, which can exist in acidic solution as cations. The electronic properties of the investigated hydrazones and their stabilities were determined with full geometry optimization by the AM1 and PM3 semiempirical methods. The obtained results for total energy (E_{tot}), binding energy (E_{bind}) and heat of formation ($\Delta H^{\text{o}}_{\text{f}}$) are given in Tables 2 (AM1) and 3 (PM3).

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•	meters of nitrogen cations $(N-sp^2$ and $N-sp^3)$ lated with the AM1 method

	Form	$^*E_{ m tot}$	$E_{\rm bind}$	ΔH_f^{o}	Charge
H ₁	$N-sp^2$ $N-sp^3$	-73165.90 -73155.40	-3517.13 -3506.66	172.86 183.33	-0.043 -0.306
\mathbf{H}_2	$N-sp^2$	-76760.94	-3800.45	164.45	-0.043
	$N-sp^3$	-76752.70	-3792.25	172.83	-0.306
\mathbf{H}_3	$N-sp^2$	-84141.20	-3891.12	133.52	-0.045
	$N-sp^3$	-84133.80	-3883.71	140.94	-0.306
H_4	$N-sp^2$	-81468.50	-3499.27	167.61	-0.047
	$N-sp^3$	-81458.10	-3488.86	178.02	-0.305
H ₅	$N-sp^2$ $N-sp^3$	-80559.80 -80550.00	-3621.52 -3611.65	128.04 137.90	-0.045 -0.306

^{*}All energy parameter values are in kcal/mol

Table 3

Physicochemical parameters of nitrogen cations calculated with the PM3 method

	Form	$^*E_{ m tot}$	$E_{ m bind}$	$\Delta H^{\rm o}_{\rm f}$	Charge
\mathbf{H}_{1}	$N-sp^2$ $N-sp^3$	-67133.70 -67126.40	-3520.92 -3513.64	169.07 176.35	-0.116 -0.009
\mathbf{H}_2	$N-sp^2$ $N-sp^3$	-70586.80 -70580.50	-3806.01 -3799.65	159.07 165.44	-0.117 -0.008
H_3	$N-sp^2$ $N-sp^3$	-77348.10 -77342.00	-3894.74 -3888.72	129.90 135.93	-0.116 -0.011
H_4	$N-sp^2$ $N-sp^3$	-74083.30 -74076.50	-3503.30 -3496.52	163.58 170.36	-0.120 -0.008
H ₅	$N-sp^2$ $N-sp^3$	-73911.60 -73904.10	-3626.33 -3618.83	123.22 130.72	-0.116 -0.011

^{*}All energy parameter values are in kcal/mol

It was assumed that the most probable structure is the one for which the calculated values are the lowest. From the results obtained using the semiempirical AM1 method, it was seen that the sp^2 hybridized nitrogen cation had lower energies ($E_{\rm tot}$, $E_{\rm bind}$ and $\Delta H^{\rm o}_{\rm f}$) than the sp³ hybridized one (see Table 2). These results were confirmed with the semiempirical method PM3 (see Table 3).

On the other hand, the lowest energies for both the $N-sp^2$ and $N-sp^3$ protonated forms were obtained for hydrazone $\mathbf{H_3}$, while the highest energies were observed for hydrazone $\mathbf{H_1}$. This suggests that the protonated form of hydrazone $\mathbf{H_3}$ was more stable in solution than those of the other hydrazones. It must be pointed out that all values of the analyzed energies for the investigated hydrazones were close (Tables 2 and 3). However, according to these calculations, we could conclude that the protonation of the imino hybridized nitrogen atom (sp²) was most probable.

Furthermore, the conclusion regarding the protonation site can be tested by comparison of the partial charges calculated with the semiempirical methods. These values evaluated with the AM1 semiempirical method (Table 2) indicate protonation of the sp³ hybridized nitrogen atom, while those evaluated using PM3 (Table 3) suggest the opposite situation, *i.e.* protonation of the sp² hybridized nitrogen atom. To resolve this situation, the proton affinity values of both nitrogen atoms (sp² and sp³ hybridized) should be calculated. The calculations were performed using the heat of formation (ΔH_f) and Gibbs energy of formation (ΔG_f), according to the Equations (3) and (4):

$$PA = \Delta G_f(B) + \Delta G_f(H^+) - \Delta G_f(BH^+)$$
 (3)

$$PA = \Delta H^{o}_{f}(B) + \Delta H^{o}_{f}(H^{+}) - \Delta H^{o}_{f}(BH^{+})$$
 (4)

where PA is the proton affinity, $\Delta H^{0}_{f}(B)/\Delta G_{f}(B)$ is the heat/Gibbs energy of formation for the mole-

cule, $\Delta H^{\circ}_{f}(BH^{+})/\Delta G_{f}(BH^{+})$ is the heat/Gibbs energy of formation for the cation and $\Delta H^{\circ}_{f}(H^{+})/\Delta G_{f}(H^{+})$ is the heat/Gibbs energy for the proton formation and has the value of 367.2 /362.57 kcal/mol [27, 28].

The results from both semiempirical methods (AM1 and PM3), provided using Equation (3), are presented in Table 4.

Table 4 $\Delta H_f^o of \ neutral \ (B) \ and \ protonated \ (BH^+) \ forms \ (sp^2 \ and \ sp^3 \ hybridized \ nitrogen \ atoms)$ and PA of hydrazones H_I – H_5

		Neutra	al form	sp^2 protonated form			sp ³ protonated form		
Compound		$^*\Delta G_f(\mathrm{B})$	$\Delta S(B)$	$\Delta G_f(\mathrm{BH}^+)$	$\Delta S(BH^+)$	$PA(G_f)$	$\Delta G_f(\mathrm{BH}^+)$	$\Delta S(BH^+)$	$PA(G_f)$
_	$\mathbf{H_1}$	-16.26	0.1258	131.65	0.1383	214.65	143.36	0.1341	202.94
	\mathbf{H}_{2}	-25.64	0.1308	120.25	0.1489	216.67	127.80	0.1511	209.12
AM1	H_3	59.06	0.1396	88.46	0.1512	333.16	96.21	0.1501	325.41
<.	H_4	-25.24	0.1326	124.45	0.1448	212.87	136.68	0.1387	200.64
	H_5	62.82	0.1312	84.83	0.1450	340.55	96.98	0.1373	328.40
	H_1	-25.45	0.1381	210.24	0.1417	210.24	135.19	0.1381	201.89
~	H_2	-34.94	0.1381	113.45	0.1531	214.17	122.02	0.1457	205.60
PM3	H_3	68.05	0.1513	83.56	0.1555	347.05	90.22	0.1534	340.39
Η.	H_4	-33.93	0.1441	119.32	0.1485	209.31	126.70	0.1465	201.93
	H_5	72.58	0.1427	79.33	0.1473	355.81	88.58	0.1414	346.56

 $^{^*\}Delta G = \Delta H - T\Delta S, \Delta G_f; PA - kcal/mol; \Delta S - kcal/mol \cdot K$

The obtained results (see Table 4) show that the $\rm sp^2$ hybridized nitrogen atom has higher proton affinity values compared to those of the $\rm sp^3$ hybridized one. This indicates that the protonation reaction takes place at the nitrogen atom which is $\rm sp^2$ hybridized, *i.e.* the imino nitrogen atom. This conclusion is in accordance with the results of the protonation site of hydrazones known from the literature [16, 29, 30]. Furthermore, the hydrazones $\rm H_5$

and \mathbf{H}_3 have greater proton affinity compared to the other investigated hydrazones, which have similar values for proton affinity. The lowest proton affinity was 212.87/209.32 kcal/mol (AM1/PM3) for hydrazone \mathbf{H}_4 . According to the calculated values of proton affinities and data known from the literature, the chemical reaction describing the protonation of hydrazones in acidic media can be presented as follows:

The similarity of the PA values of both nitrogen atoms suggests that the influence of the substituents present in the investigated hydrazone molecules on the protonation site is not significant.

Furthermore, we did some investigations related to the stability of the syn(Z) and anti(E) stereo isomers of the neutral and protonated forms of hydrazone molecules. As is well-known, the syn(Z) and anti(E) stereo isomers are possible for the amide hydrazone structures of the investigated compounds. Applying the above mentioned semiempirical methods, the geometries of the E and E isomers were optimized. The stability of the isomers and their E spotonated forms was deter-

mined using the values of E_{tot} (in kcal/mol) and ΔH^{0}_{f} (in kcal/mol). The PA values (in kcal/mol) for both isomers were calculated using the heat of formation values, according to Equation (4). The obtained results for the neutral and sp² protonated forms are given in Tables 5 (AM1) and 6 (PM3).

The obtained results indicate that the E isomer is more stable (it has a lower value of E_{tot} ; see Tables 5 and 6) than the Z isomer in neutral media, for all the investigated hydrazones. The situation is different in acidic media, when the cation is formed as a result of protonation of the sp² hybridized nitrogen atom. In this case, the stability of the Z isomer is greater compared to the E isomer.

This can be explained by the PA values of both isomers. As seen in Tables 5 and 6, the Z isomers of all hydrazones possess greater proton affinity than the E isomers. Hence, despite the fact that the stability of E isomers is higher than that of the E forms, the cation formation ability of E isomers is

higher than that of E isomers. The obtained results suggest that the Z isomer is more easily protonated than the E isomer, which is more stable in neutral solutions. The obtained results suggest that quantum chemical calculations can be used to determinate the protonation sites and stability of the isomers.

Table 5 $E_{tot} \Delta H_f^o \text{ and PA of the Z and E isomers calculated by the AM1 semiempirical method}$

AN	/ 11	Neutral	form	sp^2 protonated form			
Comp	ound	$E_{ m tot}$	$\Delta H^{o}_{f}(\mathbf{B})$	$E_{ m tot}$	$\Delta H^{o}_{f}(BH^{+})$	PA	
\mathbf{H}_{1}	E Z	-73003.34 -73000.88	20.48 22.95	-73160.64 -73163.94	178.09 174.79	209.59 215.36	
\mathbf{H}_2	$E \ Z$	-76599.40 -76595.60	16.27 15.08	-76754.10 -76758.8	171.51 166.81	211.96 215.47	
\mathbf{H}_3	$rac{E}{Z}$	-83980.70 -83975.50	-13.84 -15.64	-84133.4 -84139.40	141.32 135.32	212.04 216.24	
$\mathbf{H_4}$	$E \ Z$	-81307.60 -81304.90	13.57 16.30	-81462.90 -81466.30	173.18 169.76	207.59 213.74	
H_5	$E \ Z$	-80397.30 -80394.80	-24.31 -21.85	-80554.10 -80557.80	133.75 130.04	209.14 215.31	

Table 6 $E_{tob} \Delta H_f \text{ and PA of Z and E isomers calculated by the PM3 semiempirical method}$

PM3		Neutral	form	sp^2 protonated form			
Comp	ound	$E_{ m tot}$	$\Delta H^{o}_{f}(\mathbf{B})$	$E_{ m tot}$	$\Delta H^{o}_{f}(BH^{+})$	PA	
\mathbf{H}_{1}	E Z	-66934.94 -66933.50	14.26 15.74	-67127.06 -67129.30	175.71 173.47	205.75 209.47	
\mathbf{H}_2	$rac{E}{Z}$	-70387.40 -70386.20	4.96 6.10	-70579.20 -70583.00	166.73 162.87	205.43 210.43	
H_3	$E \ Z$	-77147.70 -77147.10	-23.34 -22.74	-77339.70 -77344.6	138.24 133.39	205.62 211.07	
\mathbf{H}_4	$rac{E}{Z}$	-73885.30 -73884.00	8.03 9.29	-74075.90 -74079.3	170.96 167.58	204.27 208.91	
H ₅	$E \ Z$	-73711.90 -73711.20	-30.64 -29.90	-73903.70 -73908.10	131.12 126.77	205.44 210.53	

3.3. Experimental pK_{BH}^+ values

As indicated earlier, pK_{BH}^+ values are important to understand reaction mechanisms which involve proton transfers. The variation of the absorbance value that occurred when the pH of the solution was increased made it possible to determine the values of pK_{BH}^+ . The calculations were performed according to the procedure mentioned in the experimental part of this work. The statistical data (standard deviation (SD), relative standard deviation (RSD) and coefficient of determination (R^2)) were calculated, as well. The pH range of

protonation, ionization ratio (log*I*) and pK_{BH}^+ values calculated from the absorbance data at an ionic strength of 0.5 mol/dm³ are given in Table 7, while those calculated from the data at an ionic strength of 0.25 and 0.5 mol/dm³ are not shown.

The corresponding average pK_{BH}^{+} values for all investigated hydrazones obtained with the spectrophotometric method and their confidence intervals at the 95% confidence level are presented in Tables 8 and 9. The presented values are the average of the measurements of three series of solutions; the number of data used for calculations is 10.

Table 7 pH, logI and pK_{BH}^+ values of hydrazones H1-H5 (experimental spectra), $\mu=0.5$ mol/dm³

10		H_1			H_2			H_3		
n	pН	$\log I$	$pK_{\rm BH}^{+}$	pН	$\log I$	$pK_{\rm BH}^{+}$	pН	$\log I$	$pK_{\rm BH}^{+}$	
1	4.7	-1.3084	3.3816	4.4	-0.8746	3.5054	4.4	-0.9419	3.5181	
2	4.4	-1.1234	3.3166	4.2	-0.7197	3.4903	4.2	-0.6525	3.5975	
3	4.2	-0.7957	3.4043	3.8	-0.3131	3.5369	3.9	-0.3804	3.5296	
4	4.0	-0.6523	3.3577	3.6	-0.1721	3.4979	3.7	-0.1634	3.5666	
5	3.7	-0.3943	3.3657	3.5	0.0114	3.4914	3.5	-0.0128	3.5272	
6	3.5	-0.2112	3.3288	3.3	0.1672	3.4672	3.3	0.2022	3.5422	
7	3.4	-0.0337	3.3463	3.2	0.3292	3.4892	3.1	3.1 0.3991 3		
8	3.1	0.2685	3.3785	2.8	0.6886	3.5386	2.9	0.6130	3.533	
9	2.8	0.4787	3.3487	2.5	0.9481	3.5081	2.7	0.8661	3.5561	
10	2.7	0.7161	3.4161	2.4	1.1307	3.5207	2.5	1.1255	3.5955	
n		H_4			H_5					
n ·	pН	$\log I$	$pK_{\rm BH}^{+}$	pН	$\log I$	$pK_{\rm BH}^{+}$	_			
1	4.6	-1.3981	3.2519	4.4	-1.2525	3.1675	_			
2	4.4	-1.0968	3.2932	4.2	-1.0019	3.1981				
3	4.1	-0.8839	3.2761	3.9	-0.7937	3.1263				
4	3.8	-0.6328	3.2472	3.6	-0.5699	3.0901				
5	3.6	-0.4038	3.2662	3.5	-0.2517	3.2183				
6	3.5	-0.3076	3.2024	3.3	0.0114	3.2714	n = numh	er of data used	l for	
7	3.3	-0.1208	3.2192	3.1	0.1208	3.2108		lations	. 101	
8	3.1	0.1138	3.2238	2.8	0.2735	3.1235		nization ratio		
9	2.8	0.4282	3.2582	2.6	0.6012	3.2212		dissociation co	nstants	
10	2.6	0.7508	3.3408	2.3	0.8128	3.1228	_	of the protonat		

Table 8 $pK_{BH}^{+} \ values \ (numerically \ and \ graphically), \ thermodynamic \ pK_{BH}^{+} \ values \ and \ statistical \ data \ (SD, RSD \ and \ R^{2}) \ of \ hydrazones \ \boldsymbol{H_{1}} - \boldsymbol{H_{5}} \ (experimental \ spectra)$

	μ [mol/dm ³]	$\mathcal{E}_{ ext{BH}}$	$\mathcal{E}_{\mathrm{BH}}^{}^+}$	1 p K_{BH}^{+}	2 p K_{BH}^{+}	1SD	1RSD	${}^{1}R^{2}$
	0.1	299768	27285	3.31±0.02*	3.32	0.03	0.91	0.999
$\mathbf{H_1}$	0.25	300371	27728	3.33 ± 0.03	3.35	0.04	1.36	0.997
	0.5	300837	28079	3.36 ± 0.01	3.39	0.03	0.94	0.998
	Thermo	dynamic p $K_{ m BH}$	+	3.26	3.26			
	0.1	302959	27772	3.43±0.03	3.46	0.05	1.49	0.997
\mathbf{H}_2	0.25	303523	28161	3.46 ± 0.04	3.49	0.06	1.72	0.996
1112								
	0.5	303892	28431	3.50 ± 0.01	3.53	0.02	0.64	0.999
	Thermo	dynamic p $K_{ m BH}$	+	3.37	3.41			
	0.1	369997	23088	3.50 ± 0.04	3.52	0.07	2.11	0.994
H_3	0.25	370304	23333	3.52 ± 0.01	3.54	0.01	0.42	0.999
	0.5	369489	23389	3.55 ± 0.02	3.57	0.03	0.83	0.998
	Thermo	dynamic p <i>K</i> _{BH}	+	3.46	3.48			
	0.1	223812	31037	3.23±0.02	3.25	0.04	1.22	0.998
H_4	0.25	223870	31591	3.24 ± 0.02	3.26	0.03	0.96	0.998
	0.5	224163	31163	3.26 ± 0.02	3.29	0.04	1.23	0.998
	Thermo	dynamic p $K_{ m BH}$	+	3.21	3.24			
	0.1	423830	29887	3.13±0.03	3.12	0.05	1.52	0.997
H_5	0.25	424423	30506	3.15 ± 0.04	3.14	0.07	2.22	0.995
	0.5	424812	30950	3.17 ± 0.03	3.18	0.06	1.82	0.996
	Thermo	dynamic p $K_{ m BH}$	+	3.10	3.08			

 $^{{}^1}pK_{BH}^+$ calculated from over determinated system of equations; ${}^2pK_{BH}^+$ graphically; 1SD – standard deviation; 1RSD – relative standard deviation; ${}^1R^2$ – coefficient of determination; ε – molar absorption coefficients ([dm³ mol^1 cm^1]) of neutral (ε_{BH}) and protonated (ε_{BH}^+) form; * confidence interval at the 95% confidence level

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Table 9
pK_{BH}^{+} values (numerically and graphically), thermodynamic pK_{BH}^{+} values and statistical data
(SD, RSD and R^2) of hydrazone $H_1 - H_5$ (reconstructed spectra)

	μ [mol/dm ³]	$\mathcal{E}_{\mathrm{BH}}$	$\mathcal{E}_{\mathrm{BH}}^{}^+}$	1 p $K_{\mathrm{BH}}^{^{+}}$	2 p K_{BH}^{+}	¹ SD	¹ RSD	${}^{1}R^{2}$
	0.1	299783	30189	3.27±0.03*	3.24	0.05	1.59	0.997
$\mathbf{H_1}$	0.25	300195	30518	3.31 ± 0.02	3.27	0.04	1.32	0.997
	0.5	300697	30721	3.36 ± 0.04	3.33	0.07	2.06	0994
	Thermody	ynamic pK _{BH} +		3.20	3.16			
	0.1	302268	27378	3.41±0.02	3.42	0.03	0.95	0.998
H_2	0.25	302917	27960	3.43 ± 0.01	3.45	0.02	0.85	0.999
	0.5	303343	28262	3.46 ± 0.02	3.49	0.03	0.83	0.999
	Thermody	ynamic pK _{BH} +		3.37	3.36			
	0.1	370064	23091	3.50 ± 0.03	3.52	0.05	1.37	0.997
H_3	0.25	370488	23341	3.52 ± 0.01	3.53	0.02	0.68	0.999
	0.5	370946	23826	3.54 ± 0.02	3.55	0.03	0.94	0.998
	Thermod	ynamic pK _{BH} +		3.47	3.51			
	0.1	222757	30358	3.22±0.03	3.25	0.04	1.38	0.997
H_4	0.25	222611	31170	3.23 ± 0.03	3.26	0.04	1.34	0.997
	0.5	222372	30438	3.24 ± 0.04	3.28	0.06	1.78	0.996
	Thermody	ynamic pK _{BH} +		3.21	3.23			
	0.1	426187	32306	3.12±0.02	3.11	0.04	1.28	0.998
H_5	0.25	426707	32690	3.14 ± 0.04	3.13	0.07	2.32	0.994
	0.5	427297	33149	3.17 ± 0.04	3.16	0.06	1.99	0.995
	Thermod	ynamic pK _{BH} +		3.08	3.07			

 $^1pK_{BH}^+$ calculated from over determinated system of equations; $^2pK_{BH}^+$ - graphically; 1SD - standard deviation; 1RSD - relative standard deviation; $^1R^2$ - coefficient of determination; ε - molar absorption coefficients ([dm³ mol¬ cm¬]) of neutral (ε_{BH}) and protonated (ε_{BH}^+) form; * confidence interval at the 95% confidence level

From the results presented in Tables 8 and 9, it can be seen that the pK_{BH}^+ values of all investigated hydrazones were quite similar. A small difference (lowest pK_{BH}^+ values) was noticed for hydrazone \mathbf{H}_5 , which exhibited the strongest basic properties compared to the other investigated hydrazones. This may be expected by the influence of the hydroxyl group in its molecule.

The similarity between pK_{BH}^{+} values was observed by comparison of the calculated pK_{BH}^{+} values obtained from experimental data and from the reconstructed spectra. This result was expected because, as was already mentioned, in the experimental spectra, clear isosbestic points appear. This suggests that the pK_{BH}^{+} values were not significantly influenced by the solvent and the use of CVA in aqueous solutions is completely unnecessary.

The ionic strength of the solution also influenced the pK_{BH}^+ values. Hence, the obtained pK_{BH}^+ values at an ionic strength of 0.5 mol/dm³ were greater than those calculated at the ionic strengths of 0.1 and 0.25 mol/dm³.

The graphically obtained pK_{BH}^+ values are presented in Tables 8 and 9. For illustration, the method of the graphical determination of the pK_{BH}^+ value of the hydrazone \mathbf{H}_3 is shown in Figure 4.

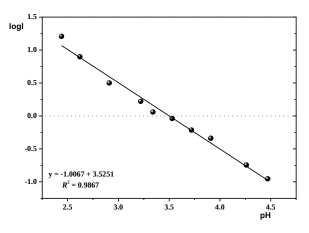


Fig. 4. Dependence of the ionization ratio (log*I*) for *N*-p-metoxybenzaldehyde-*p*-metoxybenzoilhydrazone ($c = 3.21 \cdot 10^{-5} \text{ mol/dm}^3$) on the pH of the solution, experimental spectra

As it can be seen in Figure 4, the correlation between $\log I$ and pH values is excellent with a regression coefficient $R^2 \approx 1$. The slope of the correlation line is about unity. Also, there are no significant differences between the p $K_{\rm BH}^+$ values obtained numerically and graphically (see Tables 8 and 9).

As mentioned before, the thermodynamic pK_{BH}^{+} values (Tables 8 and 9, indicated in bold)

were evaluated at zero ionic strength. The dependence of pK_{BH}^+ (at ionic strengths of 0.1, 0.25 and 0.5 mol/dm³) on the $\sqrt{\mu}$ of the hydrazone \mathbf{H}_3 obtained from the experimental spectra are shown in Figure 5.

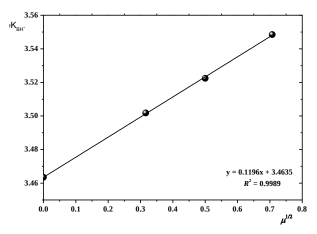


Fig. 5. Dependence of pK_{BH}^+ values on $\sqrt{\mu}$ for N-p-metoxybenzaldehyde-p-metoxybenzoilhydrazone, experimental spectra

The similarity between the thermodynamic pK_{BH}^+ values obtained from the experimental and reconstructed spectra is one more confirmation that the solvent has no important influence on the appearance of the spectra. The thermodynamic dissociation constants were also similar for all the investigated hydrazones and they were lower compared to the pK_{BH}^+ values at the ionic strength of 0.1, 0.25 and 0.5 mol/dm³. The calculated pK_{BH}^+ values of the investigated hydrazones were similar to those obtained for a similar class of compounds [31, 32].

4. CONCLUSION

The acid-base properties of five p-substituted aromatic hydrazones were described using UV spectroscopy. The spectrophotometric data demonstrated that the reaction of protonation of the investigated hydrazones occurred in acidic media. The dissociation constant values of the protonated form were performed using the absorbance values selected at four wavelengths. The thermodynamic pK_{BH}^+ values were evaluated at zero ionic strength from the pK_{BH}^+ values determined at different ionic strengths. The calculated statistical data show that this method gives very precise values. The semiempirical methods AM1 and PM3 were employed in order to calculate some of the physicochemical parameters (E_{tot} , E_{bind} , ΔH^o_f and

 $\Delta G_{\rm f}$) and proton affinities of both nitrogen (sp² and sp³ hybridized) protonated forms. The obtained results indicate that the imino (sp² hybridized) nitrogen form is favored. Although the stability of the *E* isomer is greater than that of the *Z* isomer, the protonation of the *Z* isomer is favored in acidic media. The results demonstrate that semiempirical calculations can be used successfully to determine the site of protonation, the stability of protonated forms and the stability of isomers.

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