

Research Article Free Radical Scavenging Potency of Dihydroxybenzoic Acids

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In order to evaluate the free radical scavenging potency of dihydroxybenzoic acids (DHBAs) the Density Functional Theory (DFT) was used. The M05-2X/6-311++G(d,p) and B3LYP-D2/6-311++G(d,p) theoretical models were applied. Three possible antioxidant mechanisms were examined: hydrogen atom transfer (HAT), single-electron transfer followed by proton transfer (SET-PT), and sequential proton loss electron transfer (SPLET) mechanisms. All of these mechanisms have been studied in nonpolar (benzene and pentylethanoate) and polar solvents (water) using an implicit solvation model (SMD). The following thermodynamic quantities related to these mechanisms were calculated: bond dissociation enthalpy (BDE), ionization potential (IP), and proton affinity (PA). The obtained results indicated the HAT mechanism as the most favourable reaction pathway for antioxidative action of DHBAs in benzene. On the other hand, SPLET is indicated as predominant reaction mechanism in polar solvent. The SET-PT mechanism was not favourable reaction path for antioxidative action in any of the solvents under investigation.

1. Introduction

Oxidative stress plays an important role in the pathogenesis of many diseases [1-5]. To neutralize the damaging effect of free radicals, the organisms often use some external factors including the dietary substances, such as phenolics, which can help in prevention of free radical damage. These substances constitute complex antioxidant defence systems. Phenolic acids are the group of the naturally occurring compounds which, besides one carboxylic acid functionality, have the common structural features as all phenolics: an aromatic ring bearing one or more hydroxyl substituents. There are two types of the phenolic acids: hydroxycinnamic (HCA) and hydroxybenzoic acids (HBA) [6, 7]. They are constituents of almost all vegetables, fruits, and grains. They can be found in the free state, but most commonly they are occurring in plant materials linked through esters, ethers, or as structural components of the cellulose, proteins, and lignins [8–10]. There is some evidence affirming multiple roles and functions of the phenolic acids and indicating these compounds as involved in processes such as the protein synthesis, nutrient uptake, enzyme activity, photosynthesis, and allelopathy [11–13]. There are also a number of the epidemiological investigations [14–17] which confirm a possible connection between the consumption of food containing phenolics and the reduced risk of developing some disorders, including cancer and cardiovascular diseases. However their in vivo role is still unknown.

It has been proposed that the antioxidant activity of HBA depends on the number of hydroxyl groups in a molecule [18] and that it increases following the order: monohydroxy, dihydroxy, and trihydroxy, respectively [19]. Dihydroxybenzoic acids (DHBA) are a subclass of hydroxybenzoic acids possessing two hydroxyl groups whose relative position determines the properties of molecules. There are some experimental evidences which generally confirm good antioxidant activity of DHBAs [20, 21], especially indicating good scavenging potency of 3,4-DHBA and 2,3-DHBA. Protocatechuic acid (3,4-DHBA) is a strong antiradical and antioxidant agent which inhibits the chemical carcinogenesis and show

protection against the hydroperoxide-induced toxicity [22]. Also, a few positive health attributes of 3,4-DHBA such as antibacterial [23], antimutagenic [24], anti-inflammatory, anticoagulatory [25], and antihyperglycemic [26] actions have been reported. Although all these compounds can be found in natural products there are certain findings which implicate the nonenzymatic production of 2,3-DHBA (pyrocatehuic acid), which proceeds upon trapping the hydroxyl radical by salicylic acid [27]. Moreover, pyrocatehuic acid may act as a metabolite of dioxygenases [28].

As mentioned, the antioxidant ability of the phenolic acids is greatly influenced by the number and the relative position of the hydroxyl groups in the ring. It has been also suggested that the proximity of the hydroxyl groups to the acid moiety promotes the hydrogen atom transfer from the phenolic acid (PhO-H) to the radical specie. In the radical scavenging mechanisms the reactive radical specie is inactivated by accepting a hydrogen atom from a hydroxyl group of the phenolic acid. Phenolic acids can scavenge free radicals through three competitive mechanisms, which are generally influenced by the reaction conditions. The suggested mechanisms are hydrogen atom transfer (HAT, (1)), single-electron transfer followed by proton transfer (SET-PT, (2a) and (2b)), and sequential proton loss electron transfer (SPLET, (3a) and (3b)) [29–36]:

$$Ph-OH \longrightarrow Ph-O^{\bullet} + H^{\bullet}$$
(1)

$$Ph-OH \longrightarrow PhOH^{+\bullet} + e^{-}$$
(2a)

$$Ph-OH^{+\bullet} \longrightarrow Ph-O^{\bullet} + H^{+}$$
 (2b)

$$Ph-OH \longrightarrow Ph-O^{-} + H^{+}$$
(3a)

$$Ph-O^{-} \longrightarrow Ph-O^{\bullet} + e^{-}$$
 (3b)

From the presented equations, it is clear that all these mechanisms have the same net result: formation of the corresponding phenoxyl radical which is more stable and less reactive than the free radical specie. The antioxidative mechanisms mentioned above are characterized with the thermodynamic parameters: bond dissociation enthalpy (BDE) related to (1), ionization potential (IP) related to (2a), proton dissociation enthalpy (PDE) related to (2b), proton affinity (PA) related to (3a), and electron transfer enthalpy (ETE) related to (3b).

In this paper, we aimed to provide thermodynamical parameters, related to the antiradical mechanism, that can implicate the antioxidant activity of six DHBAs: 2,3-dihy-droxybenzoic acid (2-pyrocatechuic acid or hypogallic ac-id); 2,4-dihydroxybenzoic acid (β -resorcylic acid); 2,5-dihydroxybenzoic acid (gentisic acid); 2,6-dihydroxybenzoic acid (gamma-resorcylic acid or 2,6-resorcylic acid); 3,4dihydroxybenzoic acid (protocatechuic acid), and 3,5-dihydroxybenzoic acid (α -resorcylic acid) (Figure 1).

2. Methodology Section

All calculations were performed using Gaussian 09 program package [37], and two levels of theory were applied. The



FIGURE 1: Structural formulas of the examined dihydroxybenzoic acids (1, 2, 3, 4, 5, and 6) with atom labeling indicated and their trivial names.

optimized geometries of the investigated dihydroxybenzoic acids and the corresponding radicals, anions, and radical cations were obtained by M05-2X method, in combination with 6-311++G(d,p) basis set [38, 39]. This hybrid functional, initially developed by Truhlar group [38], gives satisfactory results in the thermochemical and kinetic calculations and has been used widely by numerous authors [40–44].

The other applied method is B3LYP-D2, developed by Grimme [45, 46]. This functional can be efficiently coupled with any existing DFT method. It is proved that this method describes the interatomic interactions at short and medium distances (\leq 5 Å) reliably and more accurately than the traditional DFT methods. For including a long-range dispersion contributions to the computed DFT total energy and gradients at the B3LYP level of theory [47, 48], Grimme [46] and Bayach et al. [49] used an atom-atom additive damped empirical potential of the form $-f(R)C_6/R^6$:

$$E_{\rm B3LYP-D2} = E_{\rm B3LYP} - E_{\rm Disp},\tag{4}$$

where C_6 is the dispersion coefficient for the pair of atoms, R is the interatomic distance between atoms, and E_{Disp} is the empirical term. Both M05-2X and B3LYP-D2 methods were chosen, primarily because of being widely used and secondly because these methods describe very well the interatomic interactions at short and medium distances. These methods are more reliable and accurate than traditional density function methods.

Potential energy minima for all the optimized species are verified by the absence of the imaginary frequencies. Influence of water, pentylethanoate, and benzene was estimated using SMD solvation model [50]. The SMD continuum model allows the quantum mechanical approach in studying the interactions of the solvated molecules. The selected solvents enable investigating the behaviour of the molecules in polar and nonpolar environment. The delocalization effects were assessed within the NBO analysis framework [51].

The thermodynamical parameters relevant for the investigated antioxidative mechanisms (BDE, IP, PDE, PA, and ETE) were calculated from total enthalpies of the optimized species using the following equations:

$$BDE = H(Ph-O^{\bullet}) + H(H^{\bullet}) - H(Ph-OH)$$
(5)

$$IP = H (Ph-OH^{\bullet+}) + H (e^{-}) - H (Ph-OH)$$
(6a)



FIGURE 2: The most stable structures of the examined DHBAs.

$$PDE = H(Ph-O^{\bullet}) + H(H^{+}) - H(Ph-OH^{\bullet+})$$
(6b)

$$PA = H(Ph-O^{-}) + H(H^{+}) - H(Ph-OH)$$
(7a)

$$ETE = H (Ph-O^{\bullet}) + H (e^{-}) - H (Ph-O^{-})$$
(7b)

The recommended values of the solvation enthalpies of the protons and electrons were taken from the literature [52]. The reaction enthalpies were calculated at 298.15 K, with the temperature effects not taken into the account.

3. Results and Discussion

The conformational analysis of DHBAs was carried out. For this purpose, different conformations were obtained in the following way. First, two structures of each DHBA were constructed by placing the carboxyl group into two extreme positions (rotation around the C-C bond by 180°). Then, each of these structures was analysed by different orientations of hydrogen atom (antiperiplanar and synperiplanar) of different hydroxyl groups. All the rotamers of DHBAs are presented in Supplementary Material available online at https://doi.org/10.1155/2017/5936239 (Figures S1–S6). The tables shown in Supplementary Material (Tables S1–S6) indicate the energy differences between the investigated rotamers of all the compounds under investigation. The most stable structures are presented in Figure 2.

The structures of the all DHBAs presented in Figure 2 are planar and have at least one internal hydrogen bond (IHB). Namely, compounds which possess the hydroxyl group in position C2 have IHB between hydrogen atom of the

hydroxyl group and carbonyl oxygen of the carboxyl group. Compounds **1** and **4** form one more IHB that contribute to the stability of these molecules. The oxygen atoms of the carbonyl and hydroxyl groups possess the lone electrons located in the antibonding orbitals. These lone pair-antibonding orbital interactions, between the oxygen and the adjacent O-H bonds, are responsible, as revealed by the NBO analysis, for IHB formation. The energy values presented in Tables SI-S6 confirm that the most stable rotamers are those with the highest number of IHBs, especially the ones that include IHBs with carbonyl oxygen. Also it is evident from Figure 2 that compounds **4** and **6** are symmetrical molecules. In further discussion only the species generated from the most stable rotamers will be discussed.

3.1. Radicals and Anions of DHBAs. Investigation of the antioxidant activity of DHBs was conducted to the OH groups. This restriction to the OH groups is supported by the fact that the protons of hydroxyl groups are more acidic than the proton of carboxylic group [53, 54]. The homolytic breaking of the O-H bonds in DHBAs results in formation of the radicals (Figure 3). The stability of the formed radicals, in water, plays the main role in determining the antioxidant activity of the investigated molecules. The obtained values of BDE are given in Tables 1 and 2.

The stability of the formed radicals in water is decreasing in the following order: 3 > 5 > 1 > 4 > 6 > 2, while the order in pentylethanoate and benzene is 5 > 3 > 1 > 6 > 4 > 2. This result implies the homolytic cleavage of the 5-OH bond, in the polar solvent, as the favoured one in compound 3,



FIGURE 3: Spin density distribution in all DHBA's radicals in water.

meaning that the 3-R5 radical is the most stable radical of the investigated compounds. On the other hand, the homolytic cleavage of the 3-OH bond, in nonpolar solvents, is the favoured one in compound 5 (5-R3).

In order to rationalise the differences in BDE and consequently the differences in the reactivity of the individual OH sites the assessment of the spin density distribution was undertaken on the DHBA's radicals. The lower BDE values implicate easier formation of the radicals and more delocalized spin density [55]. The spin density values in water, obtained by the NBO analysis, as well as SOMOs of DHBAs are depicted in Figures 3 and S7. The results show that the radicals formed from compounds 3, 5, and 1 are the most stable. This is consequence of delocalization of their unpaired electrons over oxygen and carbons (O5, C2, C4, and C6 in 3-R5; O3, C2, C4, and C6 in 5-R3; and O3, C2, C4, and C6 in 1-R3). The unpaired electron of 5-R3 can be additionally delocalized only over the one adjacent OH group, 4-OH (Figure 3). Also, the unpaired electrons are well delocalized in radicals obtained by the homolytic cleavage of the O-H bond in other compounds (2, 4, and 6).

The heterolytic cleavage of the O-H bond results in forming the corresponding anions. The obtained PA values of all OH groups (in compounds 1–6), as well as in all solvents under investigation, presented in Tables 1 and 2, follow the sequence 5-A4 > 2-A4 > 1-A2 > 4-A6 > 6-A5 >3-A5. This result implicates the proton transfer from the 4-OH group of the compound 5 as easier than the transfer from the other OH groups. It should be noted that the obtained PA values are significantly lower in water (polar solvent) than the other two, nonpolar, solvents (benzene and pentylethanoate). The reason for this is change of the solvent polarity, which influences the increase of PA values as the solvent polarity increases. This is a consequence of the higher solvation enthalpy of protons.

The natural charge distributions of all anions, formed by heterolytic cleavage of O-H bonds of the DHBAs, are presented in Figure 4. Two, the most stable, anions are obtained by deprotonation of the O-H bonds of the *para* phenolic groups of compounds **2** and **5**. The negative charges, which contribute to the stability of these anions, are delocalized over O4, C1, C3, C5, and C6 atoms in **2**-A4 and over O4, C1, C2, C5, and C6 atoms in **5**-A4. The deprotonated oxygen atom in **5**-A4 forms, with the neighbouring 3OH group, one hydrogen bond, which causes lower PA value and additionally contributes to the stability of this anion.

3.2. Free Radical Scavenging Mechanisms. As mentioned above, three mechanisms of free radical scavenging activity of DHBAs (HAT, SET-PT, and SPLET) were the subject of this study. Reaction enthalpies related to these mechanisms are calculated using (5)–(7b) and by applying two DFT methods. The values of BDE, PA, and IP are used for the estimation of the preferred mechanisms of the antiradical activity of the investigated DHBAs [56–58]. The species needed to complete the thermodynamic calculations are generated from the most stable conformation of the investigated compounds.

If the calculated values obtained by both theoretical methods (Tables 1 and 2) are compared, it is clear that the thermodynamic parameters calculated using the B3LYP-D2

	LET	ETE		376	341		390	412		349	331		378	380		368	386		371	i
Water Benzene Pentylethanoate	SP.	PA		262	273		290	244		298	284		286	264		244	227		269	
	ΡT	PDE		46	35		58	35		71	38		70	51		1	3		26	
	SET	IP	592			622			577			593			611			613		
	HAT	BDE		374	371		415	392		383	350		399	379		347	349		374	
	ET	ETE		370	356		384	406		343	321		371	374		361	380		362	
	SPL	\mathbf{PA}		423	434		452	401		461	444		448	423		401	383		429	
	SET-PT	PDE		120	109		133	105		147	108		145	123		69	70		93	
		IP	673			703			658			674			693			697		
	HAT	BDE		377	366		420	392		389	350		403	381		346	347		375	
	ET	ETE		419	405		442	452		400	386		423	423		411	425		419	
	SPL	PA		132	145		146	129		153	153		149	137		130	119		144	
	SET-PT	PDE		32	25		39	31		49	34		51	39		6	12		27	
		IP	518			550			504			521			532			537		
	HAT	BDE		369	362		408	401		373	358		391	380		361	364		383	
	mpounds	1		2-OH	3-OH		2-OH	4-OH		2-OH	5-OH		2-OH	HO-9		3-OH	4-OH		3-OH	
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TABLE 1: M052X/6-311++G(d,p) calculated parameters of the antioxidant mechanisms for DHBAs in kJ mol⁻¹ in all solvents under investigation.

	.ΕT	ETE		346	314		358	379		320	304		348	350		340	357		341	341
te	SPI	PA		257	278		286	239		293	276		281	261		237	221		263	263
tylethanoa	ΡŢ	PDE		44	34		58	33		71	39		68	50		2	3		27	26
Pen	SET-	IP	559			586			542			562			575			577		
	HAT	BDE		352	341		392	367		362	329		378	360		325	326		353	352
	ET	ETE		344	309		355	377		319	299		345	348		337	355		336	336
	SPL	PA		401	426		431	380		438	420		427	405		378	359		407	406
Benzene	-PT	PDE		102	16		116	87		130	92		126	106		54	54		79	77
	SET	IP	643			670			627			646			660			665		
	HAT	BDE		356	345		397	367		367	330		382	363		325	325		353	352
	ET	ETE		364	345		386	396		346	333		370	370		358	371		365	365
	SPL	PA		135	147		151	131		157	155		148	137		132	120		147	146
Water	-PT	PDE		37	30		45	35		56	40		56	46		16	18		32	31
	SET	IP	462			492			448			461			474			480		
	HAT	BDE		348	341		386	376		352	337		367	356		339	341		361	360
	Compounds	4		2-OH	3-OH	2	2-OH	4-OH		2-OH	5-OH	4	2-OH	HO-9	10	3-OH	4-OH	5	3-OH	5-OH

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FIGURE 4: The natural charge distributions in the most stable anions formed from the investigated DHBAs in water.

functional are mainly lower. Having that in mind, the further discussion will refer to both used methods.

From Tables 1 and 2, it is clear that the IP values for all DHBAs calculated at both theoretical levels, as well as in all solvents investigated, are highly implicating SET-PT as nonoperative mechanism. On the other hand, looking at the obtained results, it is evident that in water and pentylethanoate the PA values are significantly lower than the corresponding BDE values. However, it should be pointed out that PA values are much higher in pentylethanoate than in water which indicates the SPLET mechanism as the most probable reaction pathway in those solvents. Taking into the account BDE and PA values in benzene as classical nonpolar solvent it is evident that BDE values are significantly lower than the corresponding PA values, which proves that the reaction in benzene proceeds via HAT mechanism. Further analysis of the thermodynamic values (Tables 1 and 2) indicates that compounds 5 and 1 have the best antioxidant activity in all solvents under investigation.

The BDE values obtained for the investigated DHBAs are lower than those for phenol (406.4 and 382.8 kJ mol⁻¹), resveratrol (368.8 and 356.2 kJ mol⁻¹), and caffeic acid (359.9 and 342.52 kJ mol⁻¹) [59], obtained with B3LYP method in water and benzene, as solvents. Comparing examined dihydroxybenzoic acids with the other phenolic acids, such as hydroxybenzoic acids and gallic acid [40, 41], it is clear

that dihydroxybenzoic acids show quite good antioxidative properties. Among all the mentioned acids gallic acid shows the best antioxidant activity proving the significance of the number of hydroxyl groups in such activity [41]. Our results are in good accordance with previous results [40, 41, 60].

4. Conclusions

The generally accepted approach based on the thermodynamic parameters (BDE, IP, and PA), related to the HAT, SPLET, and SET-PT mechanisms, was applied to six dihydroxybenzoic acids, their radicals, and the corresponding radical cations and anions. Calculated energy requirements indicate thermodynamically plausible radical scavenging mechanisms. The calculations performed with two theoretical models, B3LYP-D2 and M05-2X, in polar and nonpolar solvents, proved to be in good accordance.

The obtained results indicate HAT as thermodynamically favourable mechanism in benzene and SPLET in water and pentylethanoate. Calculated energy requirements indicated that compounds **5** and **1** have better radical scavenging properties than other DHBAs.

Competing Interests

The authors declare that they have no competing interests.

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