Solvent influence on intramolecular interactions and aromaticity in meta and para nitroanilines

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Abstract

Theoretical density functional theory (B3LYP/6-31G**) was used to study the intra- and intermolecular interactions of nitrobenzene, aniline, and meta and para nitroaniline in various solvation models. The studied molecules were solvated by one or two water molecules in the presence of continuum solvation (the PCM model) or without it. Finally, the studied molecules were surrounded by a cluster of water molecules. For comparison, calculations were also made for separated molecules. Geometries, energies, hydrogen bonding between solutes and solvent molecules, atomic charges, and aromaticity were examined. The analysis was based on the Atoms in Molecules methodology and the Harmonic Oscillator Model of Aromaticity (HOMA) index. As a result, an extensive description of the solvation of nitro and amino groups and the effect of solvation on mutual interactions between these groups in meta and para nitroanilines is provided. It was found that in general, the PCM description of the hydration effect on the electronic structure of the studied systems (substituents) is consistent with the approach taking into account all individual interactions (cluster model).

Keywords Substituent effect · Benzene derivatives · Theoretical calculations · Water environment · Charge of the substituent active region

Introduction

Intramolecular interactions in para and meta nitroanilines belong to the category of substituent effects (SE) in widely studied systems of a general structure $X-R-Y$, where $Y$ is a fixed functional group (reaction site), $X$ is a variable substituent, and $R$ is a transmitter. At the beginning, this kind of systems has been subject of the application of the Hammett-type approaches [1–5] with original substituent constants $\sigma$ [6] or one of their modifications [7] as explanatory descriptors.

Introduction of quantum chemistry-based descriptors of substituent electron donating/accepting properties opened a new chapter in this field of investigations. Introduced by Taft et al. [8], energies of homodesmotic reactions as

\[
\Delta E = \{E(X-R-Y) + E(Y-R)\} - \{E(X-R) + E(Y)\}
\]

allowed to quantify the substituent effect. When $\Delta E > 0$, then the interactions between $X$ and $Y$ in $X-R-Y$ system are stabilizing. The $\Delta E$ term was later named as substituent effect stabilization energy (SESE) [9] and revealed well correlations with substituent constants [9–11].

Almost 30 years later, Sadlej-Sosnowska presented a new descriptor of the substituent effect based on quantum chemistry: cSAR(X) (abbreviation from charge of the substituent active region) [12, 13] defined as a sum of atomic charges of the substituent $X$ and the substituted carbon atom, $C_{ipso}$

\[
cSAR(X) = q(X) + q(C_{ipso})
\]

Unlike atomic charges at substituents $X$, cSAR(X) nicely correlated with substituent constants in series of monosubstituted benzenes. Moreover in para substituted derivatives of nitrosobenzene and N,N-dimethylaniline, cSAR(X) described changes in geometries of amine and nitroso groups even slightly better than the Hammett substituent constants [14]. For the same reaction series, it was shown that the charge transferred from

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substituents to the NO groups, expressed by the difference cSAR (NO) – cSAR(X), well correlates with the Hammett substituent constants, whereas this difference for charges at X and NO completely failed [15]. It is important to stress that cSAR(X) values estimated by various atomic charge assessments lead to well-correlated regression lines, whereas for charges at substituents, these relations failed [16].

From the beginning of application Hammett-like relationships, it was well known that sensitivity of the substituent effect expressed by the so-called reaction constants \(\rho\) is strongly dependent on the environment in which the physicochemical processes are carried out [3]. The acid-base equilibria of substituted benzoic acids are very sensitive to solvent: The reaction constant (slope) for the Hammett relation is for water equal 1.00 (by definition), but for dimethyl sulfoxide is 2.48 [17], whereas for the gas phase measurements is 5.6 [18], and for wider review, see [19]. Recently it was shown [20] that the substituent effect in olefinic (cyclohexa-1,3-diene) and aromatic (benzene) derivatives exhibits a substantially greater sensitivity in water represented by polarizable continuum model (PCM model) [21] than in the gas phase (GP). For saturated 2,2,2-bicyclooctane derivatives, the effect of solvent was significantly weaker [20].

The purpose of this report is to confront results obtained in the gas phase and polarizable continuum model with the approach based on a model that takes into account chemical interactions of solute (aniline, nitrobenzene, and meta and para nitroaniline) molecules with explicit water molecule(s). The solvation of studied molecules was simulated by surrounding them by the 50 water molecules and full geometry optimization of the whole system at the DFT level, see the “Methods” section for details. This model (abbreviated here as \(\text{H}_2\text{O}_{\text{sim}}\)) was successfully used for the investigation of aromaticity of the benzene molecule in water [22]. For better understanding what happens when the single molecule is fully solvated by discrete solvent molecules, we added also some more models representing intermediate stages between the gas phase and the full solvation. They are the first model in which each substituent (–NO, or –NH2) is solvated by one water molecule (the \(\text{H}_2\text{O}\) model) and in the second model where the just mentioned model, the PCM environment is added (the \(\text{H}_2\text{O}+\text{PCM}\) model). In cases of meta nitroaniline and para nitroaniline, models with solvated only one functional group, NH2 (\(\text{H}_2\text{O}_{\text{amino}}\)) or NO2 (\(\text{H}_2\text{O}_{\text{nitro}}\)), are also included. Corresponding models with continuum solvation added (\(\text{H}_2\text{O}_{\text{amino}}+\text{PCM}\) and \(\text{H}_2\text{O}_{\text{nitro}}+\text{PCM}\)) are studied too. The electronic properties of substituents in this study are described by the cSAR model estimated by the use of Bader [23] methods for the atomic charge assessments. The aniline, nitrobenzene, meta nitroaniline, and para nitroaniline molecules were chosen since nitro and amino groups are well known as very strong electron-attracting [24] and donating [25] substituents, respectively. Thus, intramolecular interactions between them have been the subject of numerous studies [26–30]. Hence, they may be an excellent probe for studying the environmental impact on their mutual interactions. Another aspect of the effect of solvation is its influence on the aromaticity of benzene ring in the studied systems which will be studied by the use of aromaticity index HOMA [31, 32].

Methods

Full geometry optimizations for studied molecules and their different water clusters were performed using the Gaussian’03 package [33]. A standard optimization procedure implemented in the Gaussian package was used in all cases. The nature of optimized structures was checked by frequency calculations. No one imaginary frequency was detected. Thus, all calculated structures are minima. Three different minima were optimized for water clusters (\(\text{H}_2\text{O}_{\text{sim}}\), model) with each studied compound. The B3LYP [34] DFT functional and 6-31G basis set [35] enriched with a polarization function [36] for all the atoms (6-31G**) were used. Complexation energies of one or two water molecules to studied compounds were calculated for the gas phase according to the supramolecular approach (energy of complexation reaction, i.e., the energy difference of the product and the reactants) and corrected by the basis set superposition error values estimated by the counterpoise method [37].

Atoms in Molecules (AIM) [23] charges were calculated by means of the AIMAll software [38], which was also used to detect hydrogen bonds between a solute and water molecule(s).

Harmonic Oscillator Model of Aromaticity (HOMA) is an aromaticity index based on the geometry of studied systems (bond lengths of the ring under investigation) [31, 32]. It is calculated using the HOMA = 1 − [\(\alpha/N\) (\(R_{\text{opt}} − R_{\text{ref}}\))^2] formula, where \(R_{\text{opt}}\) and \(R_{\text{ref}}\) are optimal bond lengths and bond lengths in the real system, respectively. \(N\) is the number of bonds in the studied system and empirical factor \(\alpha\) sets the HOMA value equal to 0 for the Kekule structure of benzene (reference non-aromatic system) and 1 for the real benzene structure (reference aromatic system).

Results and discussion

Series of three subsequent figures present visualization of studied compounds: aniline, nitrobenzene, and meta and para nitroanilines in their complexes with water in various interactions with one or two water molecule(s), and these complexes in PCM model and finally when solute molecules interact with a cluster of 50 water molecules, see Figs. 1, 2, and 3. The lengths of the observed H-bonds are shown in Table 1. In addition, in the same table, energies connected with the...
complexation of one or two water molecules in the gas phase are presented.

To begin with, let us compare the strength of hydrogen bonds formed by the studied molecules with water in the gas phase. The amino group can act as both a proton donor and a proton acceptor. In the case of aniline, both H-bonds are formed in the gas phase: (i) between water oxygen and hydrogen of the amino group, denoted $\text{H}_2\text{O}(\text{H} \cdots \text{O})$, and (ii) between hydrogen of a water molecule and the amino’s nitrogen, denoted $\text{H}_2\text{O}(\text{N} \cdots \text{H})$. The first H-bond is shorter than the second (2.054 Å and 2.067 Å, respectively), but the second is slightly stronger (by 2.12 kJ/mol). The presence of a nitro group in the molecule significantly strengthens the $\text{H}_2\text{O}(\text{H} \cdots \text{O})$ H-bond ($d_{\text{H} \cdots \text{O}} = 1.975$ and 2.008 Å, with energies $-23.57$ and $-20.22$ kJ/mol, respectively for para and meta nitroaniline) and eliminates the second hydrogen bond ($\text{H}_2\text{O}(\text{N} \cdots \text{H})$). The nitro group acts only as a proton acceptor, but both oxygen atoms can participate in hydrogen bonding. This is observed in the gas phase for nitrobenzene or nitroaniline complexes with one molecule of water. The addition of an amino group to nitrobenzene does not cause such significant changes in characteristics of the H-bonds between the nitro group and water molecule as in the case of the abovementioned analogous amino group interactions, see Table 1. In addition, the simultaneous complexation of both groups by water molecules in para and meta nitroanilines enhances intermolecular interactions (by 3.64 and 2.55 kJ/mol, respectively); this strengthening is also documented by H-bond shortening. This means that intermolecular hydrogen bonds also cause changes in intramolecular interactions, i.e., the electronic structure of the remaining part of nitroaniline (solute).

Figure 1 shows also nitrobenzene and aniline molecules interacting with a water molecule in the PCM environment. As mentioned above, there is only one binding mode of a single water molecule to the nitro group of nitrobenzene. In the gas phase, the water connects to nitrobenzene using its both hydrogen atoms, see Fig. 1. It is quite curious that this seemingly logical and symmetrical structure changes in conditions of continuous solvation simulated by the PCM model, where only one hydrogen bond is formed. The reason why only one hydrogen bond is formed in the presence of solvation (here with the PCM solvation, but the same is observed in case of discrete solvation in the $\text{H}_2\text{O}_{\text{solv}}$ model) is that hydrogen bonds between water and the nitro group in the gas phase are relatively weak. The single hydrogen bond in the asymmetrical structure observed for solvated nitrobenzene is stronger (hydrogen bond in asymmetrical structure is shorter by about 0.2 Å, see Table 1). At the same time, the second hydrogen of the coordinated water molecule can interact with the environment.

On the contrary to the nitrobenzene-water complex, there are two modes of water interactions with the amino group of the aniline molecule. Water can interact through one of its hydrogens (the N...H hydrogen bond is formed) or by its oxygen atom (in this case, the H ...O hydrogen bond appears). The results of performed calculations reveal that the energetic preference of these H-bonds depends on the environment. In the case of the aniline-water complex without the PCM environment, the structure with the H ...O bond has energy lower by 3.84 kJ/mol. On the other hand, the introduction of the PCM environment changes this so that the complex with the N ...H hydrogen bond becomes more stable by 5.61 kJ/mol.

The nitro group types of interaction in meta and para nitroanilines follow rules observed in the case of nitrobenzene, see Fig. 2. The nitro group without continuum solvation interact with water by two hydrogen bonds between oxygen atoms of the nitro group and hydrogen atoms from water. This changes after using the PCM model, and the nitro group is interacting with water through one hydrogen bond only for both nitroaniline molecules.

The behavior of the solvated amino group in meta and para nitroanilines is more complicated. Not always two binding modes of the amino group with a water molecule are possible. In the case of both compounds, meta nitroaniline and para nitroaniline, geometries started with suggested N ...H hydrogen bond easily transform during optimization procedure to complexes with the H ...O hydrogen bond, if only additional continuum solvation is not present. Thus, such structures are not
present in Fig. 2 and in all the following tables. In structures in which we can compare relative stability between complexes with N–H and H–O motifs (i.e., $\text{H}_2\text{O}_{\text{amino}}+\text{PCM}(\text{H}–\text{O})$ vs. $\text{H}_2\text{O}_{\text{amino}}+\text{PCM}(\text{N}–\text{H})$, and $\text{H}_2\text{O}+\text{PCM}(\text{H}–\text{O})$ vs. $\text{H}_2\text{O}+\text{PCM}(\text{N}–\text{H})$), energies of structures containing the H–O bond are lower by about 11 and 2 kJ/mol, for para and meta derivatives respectively. All these data suggest, as expected, that the N–H hydrogen bond is more difficult to form than the H–O one.

Typical modes of studied compound interactions with solvent molecules in water clusters are presented in Fig. 3. The nitrobenzene molecule interacts with the solvent sometimes through one, sometimes through two oxygen(s), which formed always two hydrogen bonds. But always a particular
water molecule interacts with the nitro group by one hydrogen bond. Structure in which one of the water molecules in cluster forms two hydrogen bonds with the NO₂ group (the structure preferred in the gas phase) is not observed.

The amino group of aniline interacts strongly with at least three water molecules. Both possible types of hydrogen bonds (N⁻H and H⁻O) are observed. Sometimes a hydrogen atom of the amino group takes part in two hydrogen bonds, and then aniline interacts directly with four water molecules (see the structure of aniline-b, Fig. 3). Thus, the amino group interacts with water in a more differentiated way than the nitro ones. This is the reason why aniline is slightly soluble in water while nitrobenzene is considered water insoluble compound.

Different modes of solvations are observed also in cases of para and meta nitroanilines. The nitro group directly interacts always with two water molecules but in two modes: either by two H⁻O bonds between each of oxygen atoms in the NO₂ group and two hydrogen atoms of different water molecules (see structures p-nitroaniline-b and m-nitroaniline-b, Fig. 3), or by two H⁻O from different water molecules to the same oxygen atom of the nitro group (structures p-nitroaniline-a and p-nitroaniline-c, Fig. 3). Sometimes, both these types of interactions are present, see the structure m-nitroaniline-a, where the meta nitroaniline molecule interacts with three water molecules.

The nitro group of the meta nitroaniline is hydrated stronger than this group in the para nitroaniline (see Table 1 and
In the para nitroaniline, always two water molecules strongly interact with the solute molecule. In the case of meta derivative, three or two water molecules are bound to the nitro group, and the structure with three water molecules is more frequent. Also an amino group of meta nitroaniline is hydrated by more water molecules than the para derivative. Always three water molecules form hydrogen bonds with the amino group of meta nitroaniline (two N–H and one H–O hydrogen bonds) while the amino group of para nitroaniline is involved in strong interactions with three (the same type of interactions as in case of the meta nitroaniline) or two water molecules. In the case of the p-nitroaniline-c structure, the nitrogen atom of the amino group has not formed any hydrogen bond (as it was shown above, the N–H hydrogen bond is not as easy to form.

Table 1 Lengths of hydrogen bonds (HB, in Å) and BSSE corrected water complexation energies (in kJ/mol) in complexes of the studied derivatives with water; in parentheses, the type of hydrogen bond (O–H or N–H)

| Aniline                      | HB with amino group | HB with nitro group | Complexation energies |
|------------------------------|---------------------|---------------------|-----------------------|
| H2O(N–H)                    | 2.009               | 2.283               | −14.79                |
| H2O(N–H)+PCM                | 2.017               | 2.024               | −17.14                |
| H2O(N–H)+PCM+PCM            | 2.071               | 2.147               | −20.77                |

| Nitrobenzene                 |                     |                     |                       |
|------------------------------|---------------------|---------------------|-----------------------|
| H2O(N–H)                     | 1.975               | 2.011               | −23.57                |
| H2O(N–H)+PCM                 | 2.033               | 2.114               | −26.94                |
| H2O(N–H)+PCM+PCM             | 2.077               | 2.162               | −30.29                |

| Para nitroaniline            |                     |                     |                       |
|------------------------------|---------------------|---------------------|-----------------------|
| H2O(N–H)                     | 1.993               | 2.034               | −20.22                |
| H2O(N–H)+PCM                 | 2.051               | 2.127               | −23.57                |
| H2O(N–H)+PCM+PCM             | 2.095               | 2.173               | −26.94                |

| Meta nitroaniline            |                     |                     |                       |
|------------------------------|---------------------|---------------------|-----------------------|
| H2O(N–H)                     | 1.984               | 2.025               | −20.22                |
| H2O(N–H)+PCM                 | 2.043               | 2.121               | −23.57                |
| H2O(N–H)+PCM+PCM             | 2.088               | 2.166               | −26.94                |
as the H–O one). These all observations suggest that para nitroaniline effectively interacts with a lower number of water molecules than its meta counterpart. This effect is probably responsible for relative water solubility of these compounds—meta nitroaniline (0.090 g/100 g H₂O, 25 °C) is slightly more soluble than para nitroaniline (0.057 g/100 g H₂O, 25 °C) [39].

The main topic of this paper is to study the influence of solvent (water molecules) on electronic structure and the substituent effects in studied compounds by means of the analysis provided in the cSAR approach. As it was presented above, studied molecules can be solvated in different ways. With no doubts, this can affect effective cSAR values evaluated in our calculations. This may be some inconvenience but anyway, performed calculations for many different water clusters surrounding studied compounds allow us to understand better the nature of interactions and their impact on the electronic structure of functional groups as well as the interactions between them. However, such calculations are time and computer resources consuming. Thus, we must base our analysis on a limited number of cases.

The obtained cSAR (NH₂), cSAR (NO₂), and charges at both groups q (NH₂) and q (NO₂) are gathered in Tables 2 and 3. To facilitate a quantitative view, the above data are expressed also as percentage quantities in relation to the values estimated in the gas phase for NO₂ and NH₂ groups (which in all cases correspond to 100%).

Changes of cSAR (NH₂) and cSAR (NO₂) values in studied molecules depend significantly on the hydration and the differences observed are due to the type of interactions of NH₂ group with water molecules and the applied model of hydration. To show this, let us consider first results for aniline and nitrobenzene.

| Table 2 | Charges (Q) of amino and nitro groups as well as their cSAR values in the gas phase and in different types of solvations for aniline and nitrobenzene. Relative percentage changes compared with the gas phase are given in brackets |
|---------|-------------------------------------------------|
|         | Aniline                                         | Nitrobenzene                                   |
|         | Q (NH₂)  | cSAR (NH₂)  | Q (NO₂) | cSAR (NO₂) |
| GP      | −0.3648  | 0.0494 (100.00%) | −0.5350 | −0.3013 (100%) |
| PCM     | −0.3426  | 0.0636 (128.74%) | −0.5988 | −0.3582 (118.88%) |
| H₂O     | −0.3540  | 0.0832 (168.42%) | −0.5666 | −0.3238 (107.47%) |
| H₂O(H⁻O) | −0.3406  | 0.0270 (54.66%) | −0.5350 | −0.3013 (100%) |
| H₂O(N⁺H) | −0.3118  | 0.0552 (111.74%) | −0.3359 | −0.3428 (100%) |
| H₂O+PCM  | −0.3438  | 0.0772 (156.28%) | −0.3359 | −0.3428 (100%) |
| H₂O+PCM(H⁻O) | −0.3364  | 0.0123 (24.90%) | −0.3359 | −0.3428 (100%) |
| H₂O+PCM(N⁺H) | −0.3320  | 0.0569 (115.18%) | −0.3359 | −0.3428 (100%) |

For NH–O (water) interactions, an increase of cSAR (NH₂) is very significant—by 68%, whereas for N⁺-H (water), a great decrease, by 45%, is observed (Table 2). This is fully understandable: In the first case, the hydrogen bond NH–O results in drawing back proton from the NH bond towards oxygen atom in water molecule leading to an increase of a negative charge at the nitrogen atom and hence on the whole NH₂ group. The opposite is in another case: the H-bond in the N⁺-H (water) system where hydrogen atom of water molecule attracts pi-electron pair of the amino group leading to a decrease of a charge at the whole group and hence decreasing cSAR (NH₂) values. The application of the PCM approach leads also to significant changes. Direct comparison of cSAR (NH₂) values in the gas phase and in PCM results in an increase of 29%. When the PCM model is applied to interactions NH–O (water) and N⁺-H (water), then PCM computation leads to a significant increase of cSAR (NH₂) by 56% and only by 12%, respectively. It is well known that NH in the amino group is a better proton-donating system than the lone pair in this group, a proton-accepting moiety [40, 41]. Thus, the PCM model, which describes the medium by means of its dielectric constant, supports the interactions with a stronger charge separation i.e., stronger H-bonding, whereas the weaker does decrease less significantly. In the case of interactions with many water molecules, N⁺-H interactions are even strengthened: The decrease of cSAR (NH₂) values are lower than in the gas phase by 54% and 75% which may be related to the additional interactions with neighboring water molecules stabilizing the proton transfer process.

In the case of nitrobenzene, it is shown that cSAR (NO₂) values are much less dependent on interactions with water molecules and on the choice of the hydration model applied. In all cases, an increase of cSAR (NO₂) values is observed. The greatest changes are observed for PCM applied to the
It is well known that resonance effect (or intramolecular charge transfer effect) is much stronger for para substituted species than for the meta ones [42]. In order to facilitate description of changes in cSAR values for all cases considered, apart from original values of cSAR(X) (X = NH₂ or NO₂), their relative changes are given in a percentage scale, taking as 100% the values for cSAR(X) for the individual molecules in the gas phase.

The first observation is that cSAR(NH₂) values for both para and meta nitroaniline molecules are dramatically higher than for the mono substituted species. Even the gas phase data are considerably greater for amino group in para and meta nitroaniline than for individual aniline by 124.9 and 101.2%, respectively. In contrast, significantly weaker is the effect revealed for the nitro group. When cSAR(NO₂) in nitrobenzene alone, by 19%, and for NO₂ interacting with a water molecule, by 21%, whereas for nitrobenzene interacting with a water molecule in the gas phase, the change is only 7%.

Commenting these two cases, it may be said that the stronger H-bond type interactions the higher changes in cSAR(X) and the greater influence of dielectric constant of the medium, as observed via the PCM approach. Both cases of nitroanilines reveal one more interesting problem: How far different intramolecular electron interactions between the nitro group and the amino group differentiate the influence of hydration on cSAR values of both the nitro and amino groups. It is well known that resonance effect (or intramolecular charge transfer effect) is much stronger for para substituted species

### Table 3

Charges (Q) of amino and nitro groups as well as their cSAR values in gas phase and in different types of solvations for meta and para nitrobenzenes. The percentages in brackets refer to GP values of para or meta nitroanilines and to GP values of aniline or nitrobenzene, respectively.

| Para nitroaniline | Q (NH₂) | Q (NO₂) | cSAR (NH₂) | cSAR (NO₂) |
|-------------------|---------|---------|------------|------------|
| GP                | −0.3617 | −0.5976 | 0.1111 (100.00%, 224.90%) | −0.3356 (100.00%, 111.38%) |
| PCM               | −0.3324 | −0.7198 | 0.1828 (164.54%, 370.04%) | −0.4368 (130.15%, 144.97%) |
| H₂Oₐmino(H⁻O)     | −0.3562 | −0.6247 | 0.1518 (136.63%, 307.29%) | −0.3569 (106.35%, 118.45%) |
| H₂Oₜₜₜ(H⁻N)      | Transform during optimization to the H₂Oₐ₢₢₢(H⁻O) structure |
| H₂Oₜₜₜ        | −0.3601 | −0.6425 | 0.1293 (116.38%, 261.74%) | −0.3640 (108.46%, 120.81%) |
| H₂O(H⁻O)         | −0.3562 | −0.6754 | 0.1734 (156.08%, 351.01%) | −0.3887 (115.82%, 129.01%) |
| H₂Oₜₜₜ(N⁻H)     | Transform during optimization to the H₂Oₜₜₜ(H⁻O) structure |
| H₂Oₜₜₜ+PCM(H⁻O)  | −0.3353 | −0.7508 | 0.2095 (188.57%, 424.09%) | −0.4613 (137.46%, 153.10%) |
| H₂Oₜₜₜ+PCM(N⁻H)  | −0.3002 | −0.6757 | 0.1432 (128.89%, 289.88%) | −0.4038 (120.32%, 134.02%) |
| H₂Oₜₜₜ+PCM       | −0.3295 | −0.7603 | 0.2061 (185.51%, 417.21%) | −0.4564 (136.00%, 151.48%) |
| H₂O+PCM(H⁻O)     | −0.3277 | −0.7888 | 0.2278 (205.04%, 461.13%) | −0.4789 (142.70%, 158.94%) |
| H₂O+PCM(N⁻H)     | −0.2993 | −0.7090 | 0.1585 (142.66%, 320.85%) | −0.4191 (124.88%, 139.10%) |
| H₂Oₜₜₜ           | −0.3180 | −0.8442 | 0.2209 (198.83%, 447.17) | −0.5012 (149.34%, 166.35%) |
| H₂Oₜₜₜ           | −0.2811 | −0.7419 | 0.1752 (157.70%, 354.66) | −0.4702 (140.11%, 156.06%) |
| Mean H₂Oₜₜₜ value| −0.3018 | −0.7922 | 0.1954 (175.88%, 395.55%) | −0.4901 (146.04%, 162.66%) |

| Meta nitroaniline | Q (NH₂) | Q (NO₂) | cSAR (NH₂) | cSAR (NO₂) |
|-------------------|---------|---------|------------|------------|
| GP                | −0.3488 | −0.5417 | 0.0994 (100.00%, 201.21%) | −0.3043 (100.00%, 101.00%) |
| PCM               | −0.3213 | −0.6081 | 0.1340 (134.82%, 271.26%) | −0.3672 (120.67%, 121.87%) |
| H₂Oₐmino(H⁻O)     | −0.3409 | −0.5513 | 0.1343 (135.11%, 271.86%) | −0.3163 (103.94%, 104.98%) |
| H₂Oₜₜₜ(N⁻H)      | Transform during optimization to the H₂Oₜₜₜ(H⁻O) structure |
| H₂Oₜₜₜ            | −0.3458 | −0.5739 | 0.1123 (112.98%, 227.33%) | −0.3275 (107.62%, 108.70%) |
| H₂O(H⁻O)         | −0.3383 | −0.5854 | 0.1474 (148.29%, 298.38%) | −0.3406 (111.93%, 113.04%) |
| H₂O(N⁻H)         | Transform during optimization to the H₂O(H⁻O) structure |
| H₂Oₜₜₜ+PCM(H⁻O)  | −0.3248 | −0.6153 | 0.1503 (151.21%, 304.25%) | −0.3765 (123.73%, 124.96%) |
| H₂Oₜₜₜ+PCM(N⁻H)  | −0.2910 | −0.5986 | 0.1187 (119.42%, 240.28%) | −0.3548 (116.60%, 117.76%) |
| H₂Oₜₜₜ+PCM       | −0.3190 | −0.6278 | 0.1431 (143.96%, 289.68%) | −0.3748 (123.17%, 124.39%) |
| H₂O+PCM(H⁻O)     | −0.3229 | −0.6358 | 0.1594 (160.36%, 322.67%) | −0.3849 (126.49%, 127.75%) |
| H₂O+PCM(N⁻H)     | −0.2882 | −0.6181 | 0.1271 (127.87%, 257.29%) | −0.3627 (119.19%, 120.38%) |
| H₂Oₜₜₜ            | −0.3060 | −0.7066 | 0.1513 (152.21%, 306.28%) | −0.4250 (139.66%, 141.06%) |
| H₂Oₜₜₜ            | −0.2992 | −0.6606 | 0.0909 (91.45%, 184.01%) | −0.3709 (121.89%, 123.10%) |
| Mean H₂Oₜₜₜ value| −0.2992 | −0.6812 | 0.1254 (126.16%, 253.85%) | −0.3942 (129.54%, 130.83%) |
is compared with the values of para and meta nitroanilines, the increase is only by 11.4 and 1.0%, respectively. The same tendencies are observed when the mean values of the overall effect of variously realized hydration are concerned ($H_2O_{\text{solv}}$ model). The averaged cSAR (NH$_2$) value for these cases for aniline complexes, compared with aniline gas phase data, gives an increase of approximately 15% and for para and meta nitroaniline values higher by 295% and 154%, respectively. These changes are very significant but in a relative agreement with those observed for aniline and nitroanilines without intermolecular interactions, which caused only a minor increase.

In the case of cSAR (NO$_2$), the changes due to hydration are again decisively much weaker in comparison with those estimated for cSAR (NH$_2$). The appropriate cSAR (NO$_2$) values expressed in percentage scale in relation to the gas phase in nitrobenzene are as follows: in nitrobenzene, an increase by 31%, while in para and meta nitroanilines, an increase by about 63% and 31%, respectively. Tables 2 and 3 present all details of relative percentage values, whereas some summaries of the above discussion are shown in Table 4.

Table 5 shows explicitly how various interactions with water molecules influence the intramolecular charge transfer measured by values of the Charge Flow Index (CFI) [43] between amino and nitro groups ($\text{CFI} = \text{cSAR (NH}_2\text{)} − \text{cSAR (NO}_2\text{)}$) estimated by AIM atomic charge assessment. The obtained CFI data (Table 5) reveals that for para nitroaniline and its H-bonded complexes, CFI values are always higher than for the meta ones. The last right column of Table 5 shows the increase in CFI values in para isomers in comparison with the meta ones, presented in a percentage scale. Taking into account individual interactions, the differences are between 15 and 30%, indicating numerically stronger charge transfer for para nitroanilines than for the meta ones. Usually, PCM approach reveals greater changes of CFI in both cases, inside the series of meta or para isomers as well as in the relation between para isomer and the meta one.

When the CFI values for meta derivative are plotted against the para ones, then the slope is 0.71, with correlation coefficient 0.93. Once again, the transmission of charge via resonance effect for meta substituted nitroaniline is significantly weaker than for the para substituted one.

The CFI index informs about the transfer of electrons between the substituents through the ring. One more information about this effect can be found by analyzing changes in aromaticity of the ring in studied systems. Table 6 presents these data.

In the gas phase, aromaticity of the ring, estimated by the HOMA index [31, 32], for nitrobenzene is equal to 0.988, whereas for aniline, it is 0.966. In PCM, these values are 0.985 and 0.965, respectively. Because the substituent effect of amino and nitro groups is similar, in their moduli of substituent constant values (−0.66 and 0.78 for the Hammett $\sigma_p$), respectively) [44], it allows concluding that benzene ring is...
more sensitive for accepting pi-electrons than for withdrawing them. Neither aniline nor nitrobenzene does not demonstrate any substantial influence of solvation, independently of its mode; for aniline, the HOMA values are between 0.948 and 0.979 (incidentally 0.915 for one case of H₂Osolv). For nitrobenzene between 0.972 and 0.988. The case of meta nitroaniline does not differ from these data, with the range between 0.946 and 0.978. The case of para nitroaniline is significantly different, and the range of HOMA index is greater, between 0.848 and 0.945 (incidentally 0.840 and 0.841 for two cases of H₂Osolv). This may be attributed to strong NH···OH₂ interactions (see Table 1), which in the para position of NH₂ group in nitroaniline causes pi-electron pair at the amino group looser bound and easier to be involved in intramolecular charge transfer. In consequence, a quinoid-like structure is induced associated with a partial decrease of the aromatic character of the ring [45].

| Table 5 | Charge Flow Index, CFI, values of meta and para nitroanilines in different environments |
|---------|------------------------------------------------------------------------------------------|
|         | Para nitroaniline | Meta nitroaniline | Percentage (para vs. meta) |
|         | CFI | Percentage of GP | CFI | Percentage of GP |                          |
| GP      | 0.4467 | 100.0  | 0.4037 | 100.0  | 111.5                   |
| PCM     | 0.6196 | 138.7  | 0.5012 | 124.7  | 123.6                   |
| H₂Oₐmino (H⁻O) | 0.5087 | 112.9  | 0.4506 | 111.6  | 112.9                   |
| H₂Oₐmino (N⁻H) | Transform during optimization to the H₂Oₐmino (H⁻O) structure |
| H₂Oₐntro | 0.4933 | 110.4  | 0.4398 | 109.9  | 112.2                   |
| H₂O (H⁻O) | 0.5621 | 125.8  | 0.4880 | 120.9  | 115.2                   |
| H₂O (N⁻H) | Transform during optimization to the H₂O (H⁻O) structure |
| H₂Oₐmino+PCM (H⁻O) | 0.6708 | 150.2  | 0.5268 | 130.5  | 127.3                   |
| H₂Oₐmino+PCM (N⁻H) | 0.5470 | 122.5  | 0.4735 | 117.3  | 115.6                   |
| H₂Oₐntro+PCM | 0.6625 | 148.3  | 0.5179 | 128.3  | 127.9                   |
| H₂O+PCM (H⁻O) | 0.7067 | 152.2  | 0.5443 | 134.2  | 129.8                   |
| H₂O+PCM (N⁻H) | 0.5776 | 129.3  | 0.4898 | 121.3  | 117.9                   |
| H₂Oₛolv | 0.7221 | 166.7  | 0.5140 | 123.3  | 140.5                   |
|         | 0.6454 | 144.5  | 0.4777 | 118.3  | 135.1                   |
|         | 0.6888 | 154.2  | 0.5209 | 129.0  | 132.2                   |
| Mean H₂Oₛolv value | 0.6854 | 154.1  | 0.5042 | 124.9  | 135.9                   |

| Table 6 | HOMA values for aniline, nitrobenzene, and meta and para nitroanilines in various modes of solvation |
|---------|--------------------------------------------------------------------------------------------------|
|         | Aniline | Nitrobenzene | Para nitroaniline | Meta nitroaniline |
| GP      | 0.966   | 0.988       | 0.945            | 0.968            |
| PCM     | 0.956   | 0.985       | 0.901            | 0.959            |
| H₂Oₐmino (H⁻O) | 0.956 | –           | 0.922            | 0.957            |
| H₂Oₐmino (N⁻H) | 0.973 | –           | –               | –               |
| H₂Oₐntro | –       | 0.987       | 0.933            | 0.966            |
| H₂O (H⁻O) | –       | –           | 0.904            | 0.955            |
| H₂O (N⁻H) | –       | –           | –               | –               |
| H₂Oₐmino+PCM (H⁻O) | 0.948 | –           | 0.869            | 0.948            |
| H₂Oₐmino+PCM (N⁻H) | 0.968 | –           | 0.941            | 0.972            |
| H₂Oₐntro+PCM | –       | 0.983       | 0.879            | 0.958            |
| H₂O+PCM (H⁻O) | –       | –           | 0.848            | 0.946            |
| H₂O+PCM (N⁻H) | –       | –           | 0.929            | 0.971            |
| H₂Oₛolv | 0.982   | 0.967       | 0.841            | 0.967            |
|         | 0.915   | 0.978       | 0.905            | 0.978            |
|         | 0.979   | 0.972       | 0.840            | 0.972            |
| Mean H₂Oₛolv value | 0.959 | 0.972       | 0.862            | 0.972            |
Conclusions

A quantum chemical study, carried out for variously solvated molecules of aniline, nitrobenzene, and meta and para nitroaniline, enabled us to make several conclusions. It is well known that both NO₂ and NH₂ groups are able to interact with water molecules. However, the type and number of hydrogen bonds formed are different in the gas and solvated phases. It was found that, in general, the PCM description of the hydration effect on the electronic structure of studied systems (substituents) is in line with the approach taking into account all individual interactions (cluster model). It was detected that interactions of the amino group with water lead to much larger changes in its electronic structure than those observed for the nitro group. Obtained data indicate that the intramolecular charge transfer from the amino group to the nitro group is significantly stronger for the para nitroaniline than for the meta nitroaniline. Moreover, it is much more affected by interactions with water molecules. A comparison of the CFI values for variously hydrated NH₂ and NO₂ groups reveals the changes in the range between 15 and 30%. Direct correlation between the CFI values and meta and para nitroanilines reveals a slope of 0.71. Such a result again indicates much weaker charge transfer in meta substituted species. It is documented that the effect of solvation on the ring aromaticity for nitrobenzene, aniline, and meta nitroaniline is also very weak. Oppositely to that, this effect is significantly greater for para nitroaniline. This is due to the NH bond involved in H-bonding with a water molecule and hence, lone pair at nitrogen may be involved in charge transfer, leading to a quinoid-like structure with a low aromatic character.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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