How Hydrogen Bonding Amplifies Isomeric Differences in Pyridones toward Strong Changes in Acidity and Tautomerism

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ABSTRACT: Steric hindrance of hydration and hydrogen bond enhancement by localized charges have been identified as key factors for the massive chemical differences between the hydroxypyridine/pyridone isomers in aqueous solution. While all isomers occur mainly in the hydroxypyridine form in the gas phase, they differ by more than 3 orders of magnitude both in their acidity and tautomeric equilibrium constants upon hydration. By monitoring the electronic and solvation structures as a function of the substitution position on the pyridine ring, the amplification of the isomeric differences in aqueous solution has been investigated. Near-edge X-ray absorption fine structure (NEXAFS) measurements at the N K-edge served as the probe of the chemical state. The combination of molecular dynamics simulations, complete active space self-consistent field (CASSCF), and time-dependent density functional theory (TD-DFT) spectral calculations contributes to unraveling the principles of tautomerism and acidity in multiple biochemical systems based on tautomerism.

INTRODUCTION

The proton transfer connecting the enolic hydroxypyridines (HPs) and ketonic pyridones (POs) is prototypical for tautomerism in biological systems.1,2 As seen in Figure 1, both HP and PO occur in three isomeric structures of varying distances between the nitrogen and oxygen functional sites, ranging from ortho (2HP/2PO) to meta (3HP/3PO) to para (4HP/4PO). Due to these geometric differences, the oxygen substituent is conjugated with the pyridine ring in the keto forms of the ortho and para isomers (2PO and 4PO), whereas the meta isomer has no conjugate carbon–oxygen path neither in the 3HP nor the 3PO form.

Independent of these structural variations, the enol form (HP) is favored over the keto tautomer (PO) in the gas phase by all isomers.1 Upon aqueous solvation, massive differences between the equilibration of the ortho and para versus the meta isomers occur with regard to their tautomeric equilibrium, acidity, and photoreactive properties, with significant implications to their functionalities.

In aqueous solution at room temperature, the ortho and para isomers shift equilibrium away from the gas phase 2HP and 4HP forms to the 2PO and 4PO forms, whereas the meta isomer equilibrates as an equal mixture of the 3HP and 3PO tautomers3,4 (see Figure 1). These local configurations in aqueous solution are accompanied by a similar acidity of the ortho (mainly 2PO) and para (mainly 4PO) isomers in contrast to the mixed (3HP/3PO) meta isomer. In addition, the meta isomer in aqueous solution contains with its 3HP form an efficient UV chromophore, whereas the ortho 2PO and para 4PO isomers do not act as such.

Since 3HP/3PO constitutes inter alia the core moiety of vitamin B6,7,8 i.e., increased photosensitivity is caused upon vitamin B6 overdosing in humans, related to the enolic 3HP tautomer in aqueous solution.9

The ortho tautomer is directly related to multiple nucleobases of DNA and RNA,10 where tautomerism and acidity can lead to mutations and diseases, i.e., by altering the hydrogen bond patterns in DNA.11,12 Nevertheless, biological processes such as the removal of damaged nucleobases13 or the versatile catalysis by RNA enzymes14 depend on the acidity and tautomerism of nucleobases. Understanding the principles of the HP/PO protonation states sheds light on the evolution of nucleobases as storage for genetic information in all living organisms15,16 and helps to design active substances for new medical treatments (e.g., viral lethal mutagenesis17).

In this work, it is established, as a function of the protonation state and the position of the O− substituent on the pyridine ring, how charge differences and the steric hindrance of hydrogen bond coordination to the first aqueous solvation shell drive, on a microscopic level, the amplification of the isomeric differences in hydrogen bond coordination to the first aqueous solvation shell drive, on a microscopic level, the amplification of the isomeric differences in
The sample enters the experimental spectroscopic investigation of dissolved organic compounds in deionized water to obtain 0.1 M solutions. The protonation (of Sigma-Aldrich with a minimum purity of 95% and dissolved in deionized water) of the initially small molecular dihedron of the ortho, meta, and para isomers and their fully deprotonated states. They vary in oxygen conjugation, tautomeric equilibrium \( (K_T = [PO]/[HP])^{5,6} \), and acidity \( (pK_a) \). Resonance structures are given in the Supporting Information.

of the initially small molecular differences of the three HP/PO isomers, causing, macroscopically, 3 orders of magnitude variation in tautomeric constants and acidity among the ortho, meta, and para isomers in aqueous solution. The findings are based on the element-specific orbital state populations, chemical sensitivity of near-edge X-ray absorption fine structure (NEXAFS) combined with molecular dynamics (MD) simulations connected to the ab initio X-ray spectrum calculations at the level of complete active space self-consistent field (CASSCF) and time-dependent density functional theory (TD-DFT). In this effort, the ortho, meta, and para pyridone isomers; their fully deprotonated forms, respectively; and the protonated and deprotonated pyridine precursor are investigated in aqueous solution.

**METHODS**

**Experimental Details.** All samples were purchased from Sigma-Aldrich with a minimum purity of 95% and dissolved in deionized water to obtain 0.1 M solutions. The protonation (of pyridine) and deprotonation (of pyridones) were established by a 1.2-fold molar excess of HCl and KOH, respectively.

The transmission NEXAFS endstation facilitates the spectroscopic investigation of dissolved organic compounds in the soft X-ray regime.\(^{15}\) The sample enters the experimental chamber through two nozzles leading to liquid jets, which form a leaf-shaped surface upon collision. The thickness of the liquid sheet can be varied in the region of the penetration depth of soft X-rays so that transmission measurements become feasible. At the same time, radiation-induced sample damage is prevented through continuous sample replenishment. For the present investigation, a flow rate of 2.5–3.0 mL/min was used in combination with 46 μm sized nozzles.

Photonics were provided by Bessy II (Helmholtz-Zentrum Berlin) beamlines UE49-SGM\(^{16}\) for pre-studies and UE52-SGM\(^{17}\) for the quantitative absorption measurements. Bandwidths of 0.16 eV (in the case of 2PO) and 0.11 eV (otherwise) were used. The presented spectra were recorded in 0.05 eV steps with an average acquisition time of 6 s per point.

The extinction coefficient \( (\epsilon) \) was obtained from transmittance \( (T) \) and concentration \( (c) \) according to Beer–Lambert’s law.

\[
\epsilon = -\log_{10} \frac{T}{c \times l}
\]

The sample thickness \( (l) \) was estimated using tabulated values\(^{18}\) of the water transmittance at 396 eV yielding 9.2 ± 0.9 μm for 2PO and 5.6 ± 0.6 μm for the other samples.

Since the first region of the spectrum is predominantly independent of the dissolved sample, it is not only used for the calculation of the thickness but also to remove the background of the water transmittance by a linear fit.

For the energy calibration of the experimental results, the signature of co-dissolved N\(_2\) at 400.84 eV\(^{19,20}\) was used. A fit of these features in the spectrum of the pristine solvent was subtracted from the spectra of deprotonated samples to yield the pure NEXAFS of the investigated substances.

**Computational Details.** All electronic structure calculations were carried out with the Orca package.\(^{21}\) All geometries were optimized at the RI-MP2 level with the aug-cc-pVTZ/C auxiliary basis. The RI-JK correlation was used with the aug-cc-pVTZ/JK fitting basis set.

The MD simulations were performed with the Gromacs package. The OPLS-aa force field has been used for parameterization except for the charges for the intermolecular Coulomb interactions, which were derived by the chelpg\(^{22}\) procedure (based on the relaxed RI-MP2 density). The water molecules were described by the SPC/Fw\(^{23}\) model. Additional details regarding the MD simulations are available in the Supporting Information.

From the MD simulations, minimally solvated models were constructed, including only the hydrogen bonding interactions with the N\(_2\), N=H, O=H, and C=O functional groups. These minimally solvated models were used for the subsequent spectral calculations at the CASSCF and TD-DFT levels of theory.

To model the behavior of the lowest π* resonance across the systematic series, CASSCF calculations were carried out in the gas phase, as well as explicitly solvated. The aug-cc-pVQZ basis set was used for all atoms except for the nitrogen, which was described by the larger aug-cc-pCVQZ. In the solvated cases, the explicit water molecules were described by the smaller cc-pVQZ basis. The bulk-like effects were modeled by the conductor-like polarizable continuum model\(^{24}\) (CPM). The CASSCF calculations used RI-MP2 natural orbitals as a starting point. The π orbitals of the conjugated ring were included in active space, namely, six active electrons in six orbitals, henceforth called CAS(6,6). This setup excludes only the oxygen lone pair orbital, which is usually poorly correlated in CASSCF calculations. Using this active space, a state-averaged (SA) calculation was carried out for the ground state and the lowest valence-excited state. The valence CASSCF solution was then used as a starting point for the core-state calculation. The N(1s) orbital was rotated into the active space leading to CAS(8,7); then, an SA-CASSCF calculation was carried out including only the ground state and...
the lowest core-excited state. This setup is expected to recover the majority of static and dynamic correlations for the \( \pi^* \) transitions.

For Figure 4, the CASSCF transition moments were shifted by \(-0.75\) eV and convoluted with a Voigt profile of \(0.13\) eV lifetime broadening\(^{25}\) and \(0.51\) eV (deprotonated nitrogen) or \(0.80\) eV (protonated nitrogen) experimental broadening to match the experimental results.

TD-DFT calculations were performed to study the remaining resonances of the X-ray absorption spectra. The PBE\(^{26}\) hybrid functional was used in all calculations. Additional details are available in the Supporting Information.

## RESULTS AND DISCUSSION

### General Protonation Shift

Near-edge X-ray absorption fine structure (NEXAFS) at the nitrogen K-edge allows us to distinguish and quantify the protonation state at the nitrogen atom. Both the directly coordinated hydrogen atom and the hydrogen bond toward coordinated water molecules in the first solvation shell are monitored in this approach.

In Figure 2b,\(\text{e}\), the N K-edge NEXAFS of the ortho, meta, and para isomers are presented, as well as their fully deprotonated states in direct comparison to the (pyridine/Py) precursor and its conjugated acid (pyridinium/\(\text{PyH}^+\)). A characteristic blue shift of the lowest unoccupied molecular orbital (LUMO) \(\pi^*\) state is observed for all molecules upon protonation (depicted in Figure 2b,\(\text{e}\)) via the one-electron transition (\(\geq 72\%\)) of the N(1s) electron elevated to the lowest unoccupied \(\pi^*\) orbital.

The photon energy range of \(402-406\) eV monitors excitations into higher unoccupied molecular orbitals, mostly of \(\sigma^*\) character. In the region of \(403.4-403.8\) eV, a shoulder (\(\approx 0.01 \text{ M}^{-1} \text{ µm}^{-1}\)) is observed, which is only present in the protonated state. This feature originates from a transition that is mostly localized at the N–H bond. Consequently, it is missing in the spectra of deprotonated species.

The N K-edge spectra of the pyridones (see Figure 2e) largely resemble the NEXAFS of \(\text{Py(H}^+)\). Only the meta isomer exhibits a second \(\pi^*\) resonance closely below the feature of \(3\PO^-\) due to the presence of the HP tautomer. No indication of the presence of 2HP and 4HP has been found, in agreement with the tautomeric equilibrium constants (see Figure 1).

The shape resonances above \(406\) eV differ both between the pyridone isomers and in comparison with \(\text{Py(H}^+)\). These quasi-bound transitions are known to provide information on interatomic distances.\(^{27}\) The shape resonances are most pronounced in the spectra of \(\text{PyH}^+\) and \(4\PO\), where the nitrogen atom has an equal distance to both neighboring carbon atoms. The position of this feature is in qualitative agreement with the calculated C–N bond distances: \(408.2\) eV for \(4\PO\) (1.351 Å) and \(409.3\) eV for pyridinium (1.342 Å). For \(3\HP/3\PO\) with 1.341–1.346 Å C–N bond distances, the resonance position is intermediate (409.1 eV) and in \(2\PO\) the distances differ so widely (1.356, 1.369 Å) that the shape resonance position cannot be assigned unambiguously.

The shape resonance of the deprotonated species seems to be red-shifted similar to the other features. This effect is not caused by the minor changes of the C–N bond distance upon deprotonation but by a general adjustment of the electronic and solvation structures, as discussed in the following sections.

### Isomer Specific Charge Distribution

Even though all pyridones show a protonation shift similar to that of \(\text{Py(H}^+)\), the exact peak positions differ depending on the position of the \(\text{O}^-\) substituent. This becomes particularly clear from a
comparison of the $\pi^*$ resonances, which also have been used to investigate tautomerism in the past.\textsuperscript{18}

Besides the peak position, the width of the $\pi^*$ resonance depends on the protonation state (of the nitrogen atom in the respective molecule). The broadening (full width at half-maximum (FWHM) + 0.29 eV) for pyridinium/pyridones results mainly from Franck–Condon vibrational excitations\textsuperscript{29} of the N–H bond during the X-ray absorption process, which are absent in the deprotonated species.

The interspecies shift of the $\pi^*$ resonance, first, depends on the electronic density at the (nitrogen) atom, which is excited during the X-ray absorption process.\textsuperscript{30} The better the core charge is screened, the lower is the ionization energy of N(1s) electrons. As far as resonant absorption features are considered, the core electron is excited into a bound state. Since the transition energy depends, inter alia, on the energy of the initial orbital of this electron, excitation and ionization energies are linked by Koopmans’ theorem. Therefore, the NEXAFS peak position contains information on the chemical state of nitrogen in the samples.

2PO$^-$ has the highest charge density at the nitrogen site and the smallest ionization potential, followed by 4PO$^-$ where the charge is more distributed over the whole molecule. In 3PO$^-$, the electron density is shifted to the oxygen, and in Py, the charge at the nitrogen site is even lower, since the molecule is formally neutral. The same argumentation holds for the protonated case, even though the screening capacity is lower for all species (see Figure 1).

Second, the relaxation of the final state due to the Coulomb attraction between the excited electron and the core-hole influences the excitation energy, especially of the lowest NEXAFS feature. As known from benzene,\textsuperscript{31} the higher the contribution of the atomic orbital at the excited atom to the relaxed molecular orbital, the more the red shift of $\pi^*$ resonances. In other words, the $\pi^*$ resonance is lowered, if the lowest unoccupied molecular orbital (LUMO) is centered at the nitrogen atom after its relaxation.

The relaxed $\pi^*$ orbitals are depicted in Figure 2a,c,d,f for the fully deprotonated and protonated Py(H$^+$) and pyridones. It can be seen that in the PyH$^+$ case, the probability of the excited electrons to be close to the carbon atoms in the meta position is nearly zero. Oppositely, in 4PO, the LUMO is distributed over all carbon atoms and has a strong contribution from the oxygen. Consequently, the excitation energy ($h\nu_{\pi^*}$) of PyH$^+$ is expected to be strongly red-shifted, whereas only slight deviations are expected for 4PO. The shift of 3PO and 2PO, due to the described final state effect, is intermediate. If single molecules are compared between their protonation states, it is observed that the deprotonated entity has a LUMO, which is more localized at the nitrogen site. Consequently, the blue shift upon protonation is increased by final state effects.

Taking these relaxation effects into account, it can be understood that the initially expected state order $h\nu_{\pi^*}(2PO^-) < h\nu_{\pi^*}(4PO^-) < h\nu_{\pi^*}(3PO^-) < h\nu_{\pi^*}(Py)$ is disturbed by the creation of the core-hole leading to $h\nu_{\pi^*}(2PO^-) < h\nu_{\pi^*}(PyH^-) < h\nu_{\pi^*}(3PO^-) < h\nu_{\pi^*}(4PO^-)$. In the protonated case, the excitation energy of PyH$^+$ is lowered to beneath that of $h\nu_{\pi^*}(2PO^-)$ due to final state effects. This order of resonance energies is observed both experimentally and in the calculations (see Table 1).

By causing the characteristic $\pi^*$ shifts, the core-hole effect provides detailed information on the isomeric differences in charge distribution. In Py(H$^+$), nitrogen is the most electro-negative element and the excited electron, therefore, remains in close proximity. In the pyridones, nitrogen competes for electron density with oxygen. This effect increases from 3PO$^-$ to 2PO$^-$ to 4PO$^-$\textsuperscript{32}. In 3PO$^-$, the oxygen is not part of the conjugated system and is already in a charged state. In 2PO$^-$, both the oxygen and nitrogen atoms are part of the aromatic structure but pull charge in similar directions. In contrast, the conjugated oxygen in 4PO$^-$ withdraws charge from the nitrogen site, as they lie in opposite sites of the ring.

In short, the constitutinal differences between the pyridone isomers cause considerable differences in the electronic structure and charge distribution. These variations are already present in the gas phase, as the CASSCF calculations show (see Table 1). Nevertheless, without an aqueous environment, the electronic differences are not mirrored by macroscopic properties, i.e., the tautomer equilibrium constants. To gain an insight into the microscopic amplification of the isomeric differences by solvent–solute effects, the solvation structure is analyzed in the following section.

Hydrogen Bonding Interactions. Figure 3 shows how the heteroatoms govern the solvent–solute interactions of Py and the pyridone isomers. In Py, the nitrogen atom accepts on average 0.9 hydrogen bonds (HBS) from water, whereby the bonds are mostly localized out of plane.\textsuperscript{23} If the nitrogen atom is protonated, it naturally cannot accept HBS and,

\begin{table}[h]
\centering
\caption{Experimental and Calculated (CASSCF) N K-Edge $\pi^*$ Resonance Positions in eV}
\begin{tabular}{|l|c|c|c|c|c|c|}
\hline
molecule & deprotonated & protonated & deprotonated & protonated & deprotonated & protonated \\
& calcld gas & calcld aq. & expt & calcld gas & calcld aq. & expt \\
\hline
Py(H$^+$) & 399.68 & 399.78 & 399.03 & 401.78 & 401.21 & 400.52 \\
2PO$^-$ & 399.34 & 399.87 & 399.00 & 402.10 & 401.54 & 400.72 \\
3PO$^-$ & 400.04 & 399.91 & 399.14 & 402.14 & 401.41 & 400.72 \\
4PO$^-$ & 400.25 & 400.09 & 399.37 & 403.15 & 401.67 & 401.07 \\
\hline
\end{tabular}
\end{table}

Figure 3. Competition of hydrogen bond coordination to the first aqueous solvation shell between the nitrogen and oxygen heteroatom as a function of distance between these functional sites. Solvent O and H densities are depicted for isovalues of 0.68 and 0.09, respectively. The additional heteroatom of the pyridones (compared to pyridine) enhances the total coordination number, even though the hydration of the nitrogen site is sterically hindered if the heteroatoms are in close proximity. The nonconjugate O$^-$ substituent in 3PO$^-$ leads to additional coordination.
instead, the NH group donates approximately 0.7 HB to the solvent.

The second heteroatom in PO/HP leads to an increase of solvent density in the first solvation shell compared to Py(H⁺). The solvent structure around the oxygen has the same donutlike shape for all pyridones, and the solvent arrangement at the nitrogen site is similar to that of Py(H⁺). The coordination numbers and hydrogen bonding properties, however, differ drastically between the isomers and their protonation states. The comparison of protonated and deprotonated species demonstrates that higher charges always lead to stronger hydrogen bonding interactions.

Within the deprotonated pyridones, the averaged number of accepted hydrogen bonds at the nitrogen site increases with the distance to the oxygen atom, thus from 2PO⁻ (1.6 HB) to 3PO⁻ (1.9 HB) to 4PO⁻ (2.1 HB). The N···H distance is almost constant at 1.8 Å. These observations reveal that the hydration is sterically hindered in the case of close proximity of the two heteroatoms. This finding is in agreement with the enhanced hydration of the oxygen site in 4PO⁻ compared to that in 2PO⁻. The coordination number at the oxygen site reaches its maximum for 3PO⁻ due to the concentration of charge at the O⁻ substituent, which is not conjugated to the aromatic ring in this isomer. The localized charge enhances hydrogen bond acceptance (3.1 HB) compared to the para (2.9 HB) and the sterically disadvantaged ortho isomers (2.8 HB).

Steric hindrance between the two hydrogen bonding sites and the lack of oxygen conjugation in 3PO are the dominating factors for the hydration of the protonated pyridones as well. 3PO exhibits the highest coordination number both at the oxygen and nitrogen sites, followed by 4PO and 2PO. While hydrogen bond donation by the protonated nitrogen site is similar for all isomers (0.6 HB, 1.9 Å; see the Supporting Information), the number of hydrogen bonds that are accepted by the oxygen atom increase from 2PO (1.6) to 4PO (1.7) to 3PO (2.0) with a common distance of 1.7 Å.

Due to the positive charge of PyH⁺, HBs are more likely (0.7 HB) and contracted (1.8 Å) at the nitrogen site of this molecule. Analogously, HB acceptance by the deprotonated nitrogen atom is reduced in 3HP and Py (0.9 HB, 1.9 Å) compared to that in the anions.

For 3HP, the probability of accepting HBs at the oxygen is reduced to 0.7 due to the bound proton. The hydrogen atom, however, donates approximately 1.5 HBs to water (1.8 Å). The reason for the additionally donated hydrogen bond compared to the N···H case is the more polar O···H bond, as a result of the higher electronegativity of oxygen. The combination of HB donation and acceptance leads to a coordination number at the oxygen site similar to that at 3PO.

The intensity of the solvent–solute interaction as a function of the O⁻ substitution position can be summarized as follows: the steric hindrance of the solvation increases from the para to the meta to the ortho isomer. 3PO⁻ forms additional hydrogen bonds as a result of the nonconjugate C=O⁻ bond. Thereby, the solvation structure underlines the significant differences of 3PO, whose acidity and tautomeric equilibrium differ largely from those of the other isomers.

**Solvent Influence on Acidity and Tautomerism.** Based on the observations of HB formation and the arrangement of water molecules around the pyridones and Py(H⁺), information on the interactions of hydration and the electronic structure can be gained from the NEXAFS. As seen in the CASSCF calculations (see Figure 4b), the π* resonance position can shift by up to 1.48 eV in the presence of a protic solvent like water.

![Figure 4](https://dx.doi.org/10.1021/acs.jpca.0c10873)

As it is known from the sulfur derivative of 2PO (2-thiopyridone), the core-excitation distorts the solvent–solute interactions, which explains the solvation shifts. According to the equivalent core principle, the effect of a core-hole can be approximated by an increase of the nuclear charge by 1; e.g., core-excited nitrogen (N*) largely behaves like oxygen. Consequently, the polarity of the N*···H bond is increased compared to that of N···H, which strengthens the HB in N···H···OH₂. Therefore, the excitation energy is lowered for the protonated species. The magnitude of the red shift is the smallest for 2PO and the largest for 4PO. These differences are mainly caused by the steric hindrance of hydration if NH⁺ and O⁻ are close together, as observed in the MD simulation of the ground state.

The solvation shifts of the deprotonated species are comparatively small, because the solvent–solute interactions are already strong in the ground state. The additional charge in
The excited state destabilizes the hydrogen bonding structure, especially in 2PO⁻, where the LUMO has large contributions at the close-lying N⁶⁻ and O¹⁻ sites.

Taken together, the response of the solvent–solute interaction to the core-excitation leads to a reduction of the protonation shift Δhνπ for all species. This shift correlates (within the errors of the measurement) entirely with the acidity of the keto tautomers. The experimentally observed trend is well reproduced by the CASSCF calculations with explicit solvation. Thereby, the computational findings support the abovementioned reasoning that the smaller Δhνπ is, the stronger the solvent–solute interactions are. Based on the general relation of hydrogen bonding and acidity, it can be stated that the stronger the stabilization of the pyridones by the solvent, the weaker the N–H bond.

It is known from the MD simulations that 3PO has the largest coordination number of all keto forms and that this species also shows a ~0.73 eV calculated π* shift upon solvation (see Figure 4). From this, it becomes apparent why 3PO(⁻) has the lowest Δhνπ* gap of all pyridones. The strong stabilization of 3PO by the solvent is in agreement with the general assumption that the more polar (keto) tautomer is energetically preferable in aqueous solutions. However, this effect is not only mediated by the macroscopic dielectricity of water, as initially assumed in multiple studies, but also by hydrogen bonding interactions. Since these interactions diminish for higher temperatures, it can be expected that the equilibrium shifts to the enol form upon heating as observed for 3HP/3PO.

For 2HP/2PO and 4HP/4PO, the higher polarity of the keto form explains why the tautomeric equilibrium is inverted between the gas phase and aqueous solution. In contrast, 3PO does not seem to obey this rule, as it is the most polar form, but coexists with its enol tautomer in water. In fact, due to missing resonance stabilization of the charges in 3PO, it is severely disadvantaged to the point of not being energetically preferable in an aqueous solution even though it is highly stabilized. This interpretation is in agreement with the rule of Zilberg and Dick that the less stable tautomer receives higher stabilization by the aqueous environment. Additionally, the instability of the zwitterionic 3PO explains the higher acidity of 3HP/3PO compared to the other pyridones.

The acidity difference between 2PO and 4PO can be understood, now, as a result of the higher interaction of 4PO with water. The additional stabilization of 4PO agrees with the inverted order of equilibrium constants between vapor (\( \frac{[4PO]}{[2PO]} > \frac{[2PO]}{[4HP]} \)) and aqueous solution (\( \frac{[4PO]}{[4HP]} > \frac{[2PO]}{[2HP]} \)). Analogous to the 3PO case, the more intense 4PO water interactions stabilize this tautomer but cause its protonation state to be more dependent on the solvent. In other words, the abovementioned finding applies: the higher the stabilization by the solvent, the weaker the N–H bond. Since the solvation, in turn, is dependent on the steric hindrance of the hydration and its enhancement by localized charges, it can be stated that these factors control the acidity and tautomerism of pyridones at the same time.

## CONCLUSIONS

The amplification of isomeric differences of HP/PO by aqueous hydrogen bonding has been investigated using N K-edge NEXAFS spectroscopy accompanied by systematic CASSCF calculations, distinguishing inherent molecular properties and solvation effects: the dominance of the enol form for all HP/PO isomers in the gas phase is mirrored by minor differences in the electronic structure, especially of the ortho and para isomers in the ground state. The core-excited state reveals first differences dependent on the O⁻ substitution position and conjugation to the pyridine ring. In aqueous solution, the keto tautomers of 2HP/2PO and 4HP/4PO have been spectroscopically identified as dominating entities, whereas 3PO coexists with the enol form. This is consistent with previous studies in the ultraviolet range of light.

The HP/PO solvation structure has been investigated in detail by MD simulations. The common theme among all molecules considered in this study is their ability to form strong hydrogen bonds with the solvent, both by donating a hydrogen bond to water or by accepting it, and thus stabilizing the molecule. However, in 2PO, a steric hindrance of the hydration reduces the solvent stabilization. In contrast, 3PO receives the highest stabilization due to its zwitterionic character.

The strong interaction between the electronic structure of the pyridones and the surrounding water molecules shows that the increase of solvent stabilization of the keto tautomer from 2PO to 4PO to 3PO, being expressed by the N(1s) → π* protonation shift, directly correlates with an increase in acidity. It also explains that the tautomeric equilibrium is not only inverted but that the HP form exists as a minuscule fraction for 2HP/2PO and 4HP/4PO upon solvation. 3PO, which does not contain a neutral resonance structure, is energetically so unfavorable that solvent stabilization, even though it is comparatively high, only increases the share of 3PO in aqueous solution to 50%.

The water environment amplifies the tautomeric differences of the pyridone isomers because the intensity of the HB interaction depends both on the localization of charges as a result of the oxygen conjugation and the proximity of the heteroatoms leading to sterical hindrance. Hydrogen bonding not only stabilizes the keto tautomers but also weakens the covalent bond of the proton at the nitrogen site by the same ratio.

These findings clarify the principles of the HP/PO tautomerism and acidity, which have been under investigation for more than 100 years now, and they help to unravel the biological function of nucleobases, vitamin B6, and the pathways of the natural remediation of pyridine.

## ASSOCIATED CONTENT

* Supporting Information*

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcb.0c10873.

Resonance structures of the pyridone isomers; details of the molecular dynamics simulation (including radial distribution functions, chelpg charges, and hydrogen bonding properties); and TD-DFT calculations (PDF)

Optimized geometries of all discussed species in the gas phase and with explicit solvation in Cartesian coordinates (ZIP)

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Combination of AM1 with reaction field theory.

a The coordination numbers are given with respect to the oxygen atoms of water. The underlying radial distribution functions and more details of the MD simulation (including hydrogen bonding analysis) are provided in the Supporting Information.

b Since the π* shift was calculated for the keto tautomers, the pK_a value of the keto tautomers only has been chosen as a measure for the acidity.

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