# Joint neutron crystallographic and NMR solution studies of Tyr residue ionization and hydrogen bonding: Implications for enzyme-mediated proton transfer

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Human carbonic anhydrase II (HCA II) uses a Zn-bound OH<sup>-</sup>/H<sub>2</sub>O mechanism to catalyze the reversible hydration of CO<sub>2</sub>. This catalysis also involves a separate proton transfer step, mediated by an ordered solvent network coordinated by hydrophilic residues. One of these residues, Tyr7, was previously shown to be deprotonated in the neutron crystal structure at pH 10. This observation indicated that Tyr7 has a perturbed pK<sub>a</sub> compared with free tyrosine. To further probe the pK<sub>a</sub> of this residue, NMR spectroscopic measurements of [<sup>13</sup>C]Tyr-labeled holo HCA II (with active-site Zn present) were preformed to titrate all Tyr residues between pH 5.4-11.0. In addition, neutron studies of apo HCA II (with Zn removed from the active site) at pH 7.5 and holo HCA II at pH 6 were conducted. This detailed interrogation of tyrosines in HCA II by NMR and neutron crystallography revealed a significantly lowered pK<sub>a</sub> of Tyr7 and how pH and Tyr proximity to Zn affect hydrogenbonding interactions.

proton transfer | neutron crystallography | perturbed pKa | solution NMR

Carbonic anhydrases (CAs) are ubiquitous enzymes that cat-alyze the interconversion of  $CO_2$  and  $HCO_3^-$ . In humans, CAs are involved in a great variety of physiological processes, from respiration, ureagenesis, gluconeogenesis, and cerebrospinal fluid production to general acid/base homeostasis (1). The most abundant and well-characterized human CA isoform is human carbonic anhydrase II (HCA II), which displays a bellshaped pH dependence on catalytic activity, with a maximal  $k_{cat}$ of  $10^{6}$  per second at pH 7.2. In the hydration direction, the first step of catalysis is a Zn-OH<sup>-</sup>-mediated nucleophilic attack on  $CO_2$  to form  $HCO_3^-$  and one H<sup>+</sup>, leaving one water molecule bound to the zinc. The second, rate-limiting step occurs via proton transfer (PT) that removes one H<sup>+</sup> from the metal-bound water to regenerate the catalytic Zn-OH<sup>-</sup> (2). This PT step is thought to adopt classic Grotthuss-type hopping and it is mediated by a well-ordered network of hydrogen-bonded (H-bonded) water molecules that spans the ~8-Å distance between the Znwater and proton-shuttling residue, His64 (Figs. 1-3). In previous neutron structures, the Zn-bound solvent has been observed to be an H<sub>2</sub>O (or exchanged D<sub>2</sub>O) and this may imply that H<sub>2</sub>O is the favored species over OH<sup>-</sup> in the absence of substrate (3). In chemical crystallography extended X-ray absorption fine structure can be used to determine Zn:Zn-solvent distances as a way to discriminate between H<sub>2</sub>O or OH<sup>-</sup>. For Zn-OH<sup>-</sup> the distance should be  $\sim 1.80$  Å and for Zn-H<sub>2</sub>O it should be closer to 2.0 Å (4). Despite the availability of ultra-high-resolution X-ray crystal structures of holo HCA II (no substrate present) the range of Zn:Zn-solvent distances observed has led to inconclusive results. Also, within the coordinate error it is unfeasible to determine the Zn-bound solvent species using distance with any certainty (5, 6).

His64 is thought to be one component in the delivery of excess H<sup>+</sup> to the bulk solvent, resetting the active site for another round of catalysis (7–9). The His64Ala mutant retains 5–10% wild-type PT activity. This observation in combination with computational analysis supports the existence of alternate PT pathways (5, 8). The O atoms of the water network are observed intact in X-ray crystal structures determined from pH 5-10 without positional change (10). The water molecules are coordinated by a number of hydrophilic active-site residues, including Tyr7, Asn62, Asn67, Thr199, and Thr200 (Figs. 1 and 3). Numerous mutagenesis, computational, structural, and kinetic studies have shown the importance of these hydrophilic residues in (i) organizing and coordinating the active-site water network and (ii) maintaining an appropriate electrostatic environment for optimal PT (11-13). A previous study focused on the role of hydrophilic residues Tyr7, Asn62, and Asn67 in the active site. These residues were mutated to hydrophobic residues Phe7, Leu62, and Leu67. The mutants with Leu at positions 62 and 67 displayed slower PT kinetics than wild type but the Tyr7Phe mutant was observed to have significantly enhanced PT activity (approximately sevenfold over wild type) in the bircarbonate dehydration reaction. This

#### Significance

Proton transfer is a fundamental mechanism at the core of many enzyme-catalyzed reactions. It is also exquisitely sensitive to a number of factors, including pH, electrostatics, proper active-site geometry, and chemistry. Carbonic anhydrase has evolved a fast and efficient way to conduct protons through a combination of hydrophilic amino acid side chains that coordinate a highly ordered H-bonded water network. This study uses a powerful approach, combining NMR solution studies with neutron protein crystallography, to determine the effect of pH and divalent cations on key residues involved in proton transfer in human carbonic anhydrase. The results have broad implications for our understanding of proton transfer and how subtle changes in ionization and H-bonding interactions can modulate enzyme catalysis.

The authors declare no conflict of interest.

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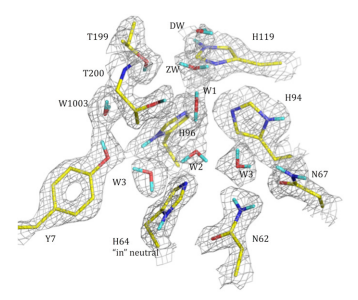
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Data deposition: Crystallography, atomic coordinates, and structure factors have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 4q49 and 4y0j).

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**Fig. 1.** The pH 7.5 *apo* HCA II active site determined by neutron crystallographic studies. The  $2F_o - F_c$  nuclear density map is shown in gray mesh (contour 1.3 $\sigma$ ). Exchanged Ds are in cyan; unexchanged H atoms are not shown for clarity. Active-site residues and solvent molecules are labeled.

unexpected behavior was ascribed to two effects: the loss of Tyr at position 7 seemed to lower the  $pK_a$  of His64 from 7 to ~6, and the solvent structure between the Zn-bound solvent and His64 was disrupted to form an unbranched network. It was believed that this and His64's having a more acidic  $pK_a$  led to the Tyr7Phe mutant's having improved PT kinetics compared with wild type (11).

Neutron protein crystallography provides insight into enzyme mechanisms. It is the only technique that can directly, and at high resolution, visualize H and its isotope deuterium, D. Unlike X-rays, neutrons are scattered very well by H/D atoms, similarly to the other common elements found in proteins, such as C, N, and O. This allows detailed analysis of water coordination, H bonding, ligand binding, and the protonation state of certain amino acids such as His and Lys (14, 15). This information is crucial to understanding how H bonds and water mediate not only enzyme catalysis but also ligand binding. Previously, the neutron crystal structure of HCA II crystallized from a pH 10 buffer showed an incomplete H-bonded water network in the active site and a  $D_2O$  molecule bound to the Zn (3). The structure also revealed that a key active-site residue, Tyr7, was deprotonated (Fig. 3). Interestingly, all of the other seven tyrosines in HCA II, including solvent-exposed surface Tyr residues that were expected to be sensitive to bulk solution pH changes, were observed to be neutral in the crystal structure (3). Although deprotonation of Tyr7 was not totally unexpected, because tyrosine in solution and in random coil polypeptides has a pKa of  $\sim 10$ , this observation indicated that the pK<sub>a</sub> of this residue is significantly lowered (16, 17). In enzyme systems where tyrosines are involved with general acid-base catalysis, site-specific pKa measurements with NMR spectroscopy have revealed that some tyrosines exhibit pK<sub>a</sub>s as low as 6 (18). Other examples of enzymes with catalytic tyrosines that have perturbed pKas include β-lactamase, ketosteroid isomerase, and UDP-galactose 4-epimerase (19-21). It is an important, although less understood, phenomenon that enzymes are able to fine-tune local electrostatics to support a particular function (22).

Subsequent to the characterization of the high-pH neutron structure of HCA II, an additional neutron crystal structure was determined for crystals grown at pH 7.8. Here, as expected, Tyr7 was neutral and involved in a bifurcated H bond (Fig. 3). To

date, the catalytic role for Tyr7 in HCA II has not been proposed, although it is unusual for a noncatalytic tyrosine to have a perturbed  $pK_a$ . Of course, the extent of the perturbation is difficult to gauge using neutron crystal structures alone because they provide only static snapshots of the residue in question at a few selected pH values. Moreover, estimating the pH inside a crystal is challenging; we are limited to measuring the pH of the crystallization solution, which gives only an approximation of the crystal's inner pH. Furthermore, because at pH close to  $pK_a$  residues are partially protonated, it is possible that one of the charged forms selectively crystallizes, shifting the observed equilibrium.

There are eight Tyr residues in HCA II with residue numbers 7, 40, 51, 88, 114, 128, 191, and 194. Tyr7 is located on the edge of the active site and is exposed to solvent. Tyr40, 114, and 128 sit on the surface and are also solvent-exposed, whereas Tyr51, 88, and 191 have their side chains buried and participate in H bonding with other parts of the protein. In this study, we probed the protonation state of all tyrosine residues in HCA II and measured their  $pK_a$  in solution. We combined these studies with analysis of neutron crystallographic structures to shed light on the effect of pH on important active-site residues (such as Tyr7 and His64) and the biophysical effect of the divalent cation (Zn) in the active site.

To address these questions, we determined the neutron structure of apo HCA II (with active-site Zn removed) crystallized at pH 7.5 to detect any influence of the divalent cation on the nearby residues and those farther away from the active site. Another neutron crystal structure of holo HCA II (with active-site Zn present) crystallized at pH 6.0 was determined to observe pH effects on all Tyr residues and the proton shuttle, His64. [13C]Tyr-labeled CA was prepared for NMR analysis and the spectra were measured from pH 5.4-11.0. It was possible to assign all eight Tyr in HCA II, providing the opportunity to interrogate their individual behaviors in solution and to compare the results to the crystal structures. To our knowledge, this is the first report of the use of neutron crystallography and NMR spectroscopy together to map tyrosine protonation states as a function of pH and the divalent cation. The observation of a perturbed pKa for Tyr at physiological pH has implications for our understanding of the PT mechanism and the overall enzyme-catalyzed reaction trajectory.

## **Results and Discussion**

Apo HCA II at pH 7.5. The neutron structure of apo HCA II was determined to 1.8-Å resolution with a refined  $R_{cryst}/R_{free}$  of 20.3/21.7. Detailed analyses of  $F_o$  –  $F_c$  and  $2F_o$  –  $F_c$  nuclear maps show that the active-site water network has the same H bonding and coordination as *holo* HCA II crystallized at pH 7.8 (Fig. 1: PDB ID code 3tmj). In the absence of an active-site metal, there is still a complete H-bonded network of active-site water molecules stretching to the bulk solvent. Also, as expected, Asn62, Asn67, Thr199, and Thr200 remain unchanged (Fig. 1). Although the overall water pattern and most of the active-site residues seem to be similar to holo structures, there are notable differences in the catalytically significant residues. The proton shuttling residue, His64, is neutral and well-ordered in the "inward" conformation. This is in contrast to other studies in which His64 was observed in two conformations, "inward" and "outward," relative to the activesite opening (Fig. 1) (8, 23, 24). In the holo pH 7.8 neutron structure, there are three discernable states of His64 that differ in charge and imidazole ring orientation (PDB ID code 3tmj).

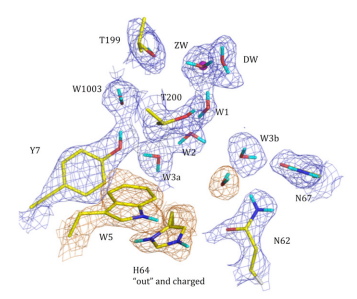
**Holo HCA II pH 6.** Because the protein does not crystallize below pH 7, it was challenging to obtain the low-pH crystal form of HCA II. To obtain crystals at this low pH, we subjected them to vapor acidification, which may have led to lower-quality neutron data observed. The neutron structure of *holo* HCA II crystallized at pH 6 was determined to 2.0-Å resolution and the final model

was refined to  $R_{cryst}/R_{free}$  26.3/29.1. Despite the lower quality of this dataset with regard to resolution and completeness, we could observe several readily apparent structural features.

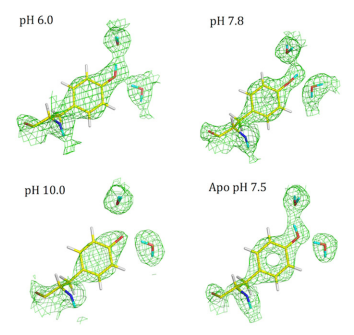
Overall, the active-site residues seem to be very similar to the *apo* structure with some subtle changes in the water network. Most notably, the deep water (DW) has moved compared with its position in the substrate-free pH 7.8 and 10 neutron crystal structures (Fig. 2). A report of the *holo* HCA II in a complex with CO<sub>2</sub> substrate also identified this water molecule in a shifted position (25). In the low-pH neutron structure we analyzed, the DW oxygen (O) atom is shifted by about 1.6 Å, similar to the O atom of the DW in the HCA II:CO<sub>2</sub> complex structure.

The most interesting structural feature is His64 in the "outward" conformation while protonated and  $\pi$  stacking with Trp5 (Fig. 2; see close-up view in Fig. S1). This outward position has been observed in numerous X-ray crystal structures and His64 was speculated to be protonated in this position (24). Our lowpH neutron crystal structure offers the first direct evidence to our knowledge of the protonation of His64 in the "outward" conformation. Also, we observed an additional water molecule occupying the former "inward" His64 position with its O atom acting as an H-bond acceptor (HBA) from an imidazole H. The two main features of the low-pH HCA II neutron structure, namely DW moving outward and His64 in the "outward" conformation, are the same as those seen in the high-resolution X-ray crystal structure of the HCA II:CO<sub>2</sub> complex structure (25).

**Tyrosine Residues.** Further investigation of all eight tyrosine residues in HCA II reveals additional features (Figs. S2–S4 and Table 1). Out of the eight, Tyr7 is the only residue that is observed in three distinct states: neutral [HBA and H-bond donor (HBD) to water molecules W3a and W1003], neutral (bifurcated HBD to two water molecules), and deprotonated/charged (HBA) (Fig. 3). As expected, we observed the other seven tyrosines always as neutral (Table 1). Four tyrosine residues—Tyr40, Tyr114, Tyr128, and Tyr191—were unchanged in H-bonding interactions over the pH range studied by crystallography. These four tyrosines were also insensitive to the presence or absence of the divalent cation Zn in the active site (Table 1). The remaining three Tyr



**Fig. 2.** The pH 6 *holo* HCA II active site determined by neutron crystallographic studies. The  $2F_o - F_c$  nuclear density map is shown in blue mesh (contour 1.3 $\sigma$ ). Exchanged Ds are in cyan; unexchanged H atoms are not shown for clarity. Active-site residues and solvent molecules are labeled. The nuclear density for Trp5 and His64 is shown in orange.



**Fig. 3.** Protonation state and H-bonding interactions of Tyr7 in HCA II as a function of pH and in the absence of Zn. The  $2F_o - F_c$  nuclear density map is shown in green mesh (contour  $1.2\sigma$ ). Exchanged D atoms are in cyan; unexchanged H atoms are shown in white.

residues—Tyr51, Tyr88, and Tyr194—underwent subtle changes in their H-bonding patterns between the *holo* and *apo* forms to compensate for the absence of metal and to maintain H-bonding interactions (Figs. S2–S4). Table 1 summarizes the Tyr residues and their interactions from all four neutron structures of HCA II.

The position of the D atom shared between Tyr and His residues is challenging to assign with absolute confidence based on crystallographic data alone. However, we know from the NMR data that all Tyr residues other than Tyr7 are neutral in the pH range studied. In these cases, we also calculated a series of omit maps to help guide the placement of the D atom, while taking the surrounding H bonds into account. Interestingly, Tyr51, 88, and 194 side chains are buried and interact with other residues and water. Also, these changing tyrosine residues are all between 7 and 17 Å away from the zinc. They seem to be insensitive to pH but may be sensitive to the absence of metal. That these effects operate within a certain distance from the zinc site seems to indicate possible long-range electrostatic effects operating in HCA II. An alternate explanation is that the changes in Tyr H bonding could also be due to a series of local effects that are propagated over a distance, culminating in local changes.

NMR Spectroscopy of Tyr in HCA II. Although neutron crystal structures give explicit information about Tyr residues and their charged states, they provide discrete snapshots and do not indicate protonation changes over a broad pH range and associated pK<sub>a</sub> values. NMR spectra measured on [<sup>13</sup>C]Tyr-labeled HCA II from pH 5.4-11.0 fill in these gaps and provide details not obtainable from static crystal structures alone. Owing to specific <sup>13</sup>C[Tyr labeling, only the Tyr residues were observed and analyzed. The assignments of Cy carbon frequencies from HNCACB, HSQC, and CG(CB)HB spectra are shown in Fig. S5. The assignments of C<sup>ζ</sup> carbon frequencies are shown in Fig. S6. Fig. 4 shows the chemical shift changes for all Tyr residues between pH 5.4 and 11.0. The integrity of the protein was followed using  ${}^{1}\text{H}{}^{-15}\text{N}$ heteronuclear single quantum coherence (HSQC) experiments, which showed well-resolved and dispersed spectra in the pH range from 5.4 to 11.0. When the pH was raised to 11.47, the HSQC

Table 1.	Tyrosine in HCA II interactions, H bonds, and distance to active-site Zn
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Tyr no. (distance from Zn, Å)	pH 6 <i>holo</i> (this work, 4y0j)	pH 7.8 <i>holo</i> (PDB ID code 3tmj)	pH 10 <i>holo</i> (PDB ID code 3kkx)	pH 7.5 <i>apo</i> (this work, 4q49)
7 (7)	Neutral (HBD and HBA to 2 waters)	Neutral (bifurcated HBD to 2 waters)	Deprotonated/charged (HBA to water)	Neutral (HBD and HBA to 2 waters)
40 (28)	Neutral (HBD to water)	Unchanged	Unchanged	Unchanged
51 (14)	Neutral (*HBA to protonated His122)	Neutral (HBA to protonated His122)	Neutral (HBA to protonated His122)	Neutral (HBD to neutral His122)
88 (17)	Neutral (HBD to –CO Asp139, HBA to 2 waters)	Neutral (HBD to –CO of Asp139, HBA to 2 waters)	Neutral (HBD to water)	Neutral (HBD and HBA to 2 waters)
114 (18)	Neutral (HBD and HBA to 2 waters)	Unchanged	Unchanged	Unchanged
128 (21)	Neutral (HBD to Asp139)	Unchanged	Unchanged	Unchanged
191 (20)	Neutral (HBD to –CO of Lys45)	Unchanged	Unchanged	Unchanged
194 (10)	Neutral (HBA to protonated His107, HBD to Ser29)	Neutral (HBA to protonated His107, HBD to Ser29)	Neutral (HBA to protonated His107, HBD to Ser29)	Neutral (HBD to neutral His107, HBA to Ser29)

spectrum showed the collapse of resonances, indicating protein unfolding. No data at higher pH could therefore be obtained. Tyr7 (black squares in Fig. 4) is the only Tyr that changes at close to neutral pH, consistent with observations of the neutron crystal structures (Table 1 and Fig. 3) (3). Close to pH 10, two additional Tyr residues, Tyr40 and 114, start to deprotonate. Tyr51, 128, and 194 begin shifting when the pH is greater than 10.5. This is not surprising because this pH is close to the expected pK<sub>a</sub> for free Tyr. Fig. 5 shows the chemical shift dependence of Tyr7 with a calculated pK<sub>a</sub> of 7.1.

Although accurate  $pK_a$  values for Tyr40 and Tyr114 could not be determined because the chemical shift values for these residues do not level off at the highest pH accessible, the fit of the initial part of the titration curves yields lower bounds for these  $pK_a$  values. These are 10.25 for Tyr40 and 10.45 for Tyr114. Both of these residues are solvent-exposed; thus, it is not surprising that their  $pK_a$  values are similar to those in model peptides. Given the initial shift change for Tyr51 above pH 10.5, we estimated its  $pK_a$  to be in the 11.5–12.0 range, whereas for Tyr88, 128, 191, and 194 the estimated the  $pK_a$  values will be greater than 12. This wide range of  $pK_as$ , from ~7–12 as measured for residues that are either buried or solvent-exposed indicates modulation by local environments and H bonding and how they affect the ionization of residues.

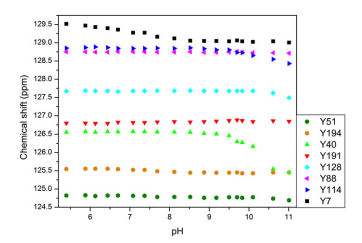


Fig. 4. Chemical shift changes of  $C\gamma$  resonances for all tyrosine residues in HCA II as a function of pH.

An interesting observation from the NMR data is that the total chemical shift changes for Tyr7 during the titrations seem to be very small. As determined previously, chemical shift changes upon deprotonation for <sup>13</sup>C $\gamma$  and <sup>13</sup>C $\zeta$  resonances in peptides are -6.2 and +10.4 ppm, respectively (26). In photoactive yellow protein, chemical shift changes for tyrosine <sup>13</sup>C $\gamma$  were determined to be between -3.8 and -4.8 ppm (27), while Baturin et al. observed changes for tyrosine <sup>13</sup>C $\zeta$  resonances in *B. circulans* xylanase between +9.0 and +9.5 ppm (19). In this study, we observed a total chemical shift change for Tyr7 <sup>13</sup>C $\gamma$  of -0.5 ppm, with an apparent pK<sub>a</sub> of 7.1, which is very low for a tyrosine hydroxyl group (Fig. 5). The data for <sup>13</sup>C $\zeta$  of Tyr7 showed a total chemical shift change of +0.3 ppm, with similar apparent pK<sub>a</sub> (Fig. S7*B*).

One could argue that the small overall chemical shift changes for Tyr7 carbons reflect electrostatic changes in the neighborhood of Tyr7 rather than actual deprotonation. Tyr7 is in fact close (<5Å) to His64 with a pK<sub>a</sub> of ~7.2 and deprotonation of this histidine could change the electrostatics around Tyr7 and lead to small chemical shift changes of Tyr7 resonances. The evidence contradicts this argument, however. First, it is the Tyr7 C $\zeta$  carbon that is closer to His64 rather than the C $\gamma$  carbon, yet the C $\gamma$  carbon experiences larger chemical shift changes. Additionally, as evidenced by the neutron crystal structures, at pH 6 Tyr7 is fully protonated, whereas at pH 10.0 it is deprotonated. Because the NMR titrations extended over a pH range from 5.4 to 11.0, the chemical shift changes of Tyr7 carbon resonances should then reflect the deprotonation of this residue.

The reason for the decreased chemical shift sensitivity of Tyr7 carbon resonances remains ambiguous. One possibility is the unique participation of Tyr7 in the proton-conducting water network extending from the Zn atom to His64. These water molecules are well ordered and seem to be conserved in all crystal structures discussed here. Tyr7 forms relatively short and strong H bonds with water molecules from this ordered network. At low pH, the O–H bond of Tyr7 is polarized by participating in the H bond as a donor. At high pH, the electron density is withdrawn from the charged O by participating in the H bond as an acceptor. If these H bonds are very stable, as suggested by the conservation of the water network and the short O...O distances, the overall change in electron density on the tyrosine ring between the neutral and charged state may be small, thus leading to small overall chemical shift changes.

Tyr7 plays a coordinating role in the proton-conducting water network, but it was previously surmised that it is mostly involved in maintaining an appropriate electrostatic environment to support

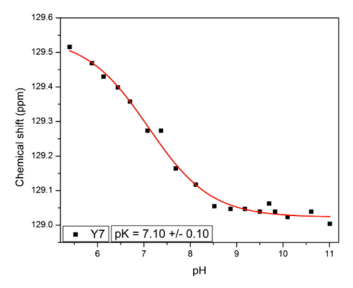


Fig. 5. Chemical shift changes of  $C\gamma$  resonance for Tyr7 in HCA II as a function of pH. The fit gives the apparent  $pK_a$  of  ${\sim}7.1.$ 

PT (11–13). The pH of normal blood is 7.35–7.50, which is slightly above the pK<sub>a</sub> for Tyr7, suggesting that roughly half the population would be deprotonated (28). Given this  $pK_a$  that is within the physiological pH range, there are profound implications for the role of Tyr7 in PT under normal conditions. When protonated at physiological pH, Tyr7 acts as a bifurcated HBD to W1003 and W3a in the water network (Figs. 1, 3, and 4). This "normal" configuration supports the H bond-connected water network and is seemingly ready to transport the excess proton from the Znbound solvent to the proton-shuttling residue, His64. However, Tyr7 was observed in the neutron crystal structure to be deprotonated at high pH, causing a rearrangement of the water molecules into a configuration where W3a is an HBD to Tyr7. Such a configuration would not support PT through the water molecules alone, and the proton moving from the Zn-bound water may be captured by the phenolate of Tyr7 before it reaches His64. In this way, Tyr7 can operate as a proton trap in wild-type HCA II, a possibility that would support the kinetic measurements of the Tyr7Phe HCA II mutant that indicated faster kinetics than in wild type (11). In the next step Tyr7 can easily donate its H to the adjacent water that in turn passes it along to His64, or His64 can occupy a rotamer that favors direct transfer from Tyr7. This direct transfer scenario would necessitate the rearrangement of two or three water molecules as well as the rotation of the His64 side chain along  $\chi^2$ , which should be energetically unfavorable.

The pK<sub>a</sub> of the proton donor:acceptor groups (ZW, His64) in the HCA II active site are closely matched at ~7 and our data indicates that Tyr7 also belongs to this group. For efficient PT the pK<sub>a</sub> values of the proton donor: acceptor groups need to closely match. We propose that involvement of Tyr7 in the H-bonding network and proximity to His64 results in lowering the pK<sub>a</sub> of Tyr7 by ~3 pH units. For His64 to switch between "in" and "out" conformations with minimal energetic cost, Tyr7 should participate in rearranging of the H-bonded network. This is possible if Tyr7 pK<sub>a</sub> is close to that of His64. This suggests several scenarios for the PT pathway in wild-type HCA II. All involve proton movement from the ZW to His64 through (i)W1-W2, (ii) W1-W2-Tyr7 and W3a, or (iii) W1-W2-Tyr7-His64. We believe that Tyr7 is also a catalytic residue. Pathways ii and iii are longer and/or may involve significant solvent rearrangement that can be expected to be kinetically and energetically less favorable. Tyr7 as a PT partner directly with His64 or a proton trap further supports the observation that the Tyr7Phe

HCA II mutant has even higher PT activity than does the wildtype HCA II (11). From the Tyr7Phe kinetic results it was shown that Tyr7 has a significant role in modulating the  $pK_a$  of His64 at near-neutral values and that removing Tyr7 causes the His64 to become more acidic, lowering His64  $pK_a$  to ~6. This makes His64 a better proton donor in the bicarbonate dehydration reaction, causing an increase in PT activity in the Tyr7Phe mutant (11–13). These earlier measurements support our proposal that Tyr7 is a catalytic residue involved in stabilizing the PT network, thus enhancing PT.

The presence of a strongly electronegative Tyr in a finely balanced electrostatic environment will have a major impact on PT, electrostatics, and the overall energetics of PT. Tyrosinate could also modulate alternative, less-efficient PT routes out of the active site under certain nonideal pH conditions, such as during blood alkalosis. It is also possible that Tyr7 can act as a proton hole when it is deprotonated. The mechanistic determinants of a proton hole in this environment require that OH<sup>-</sup> is generated first by protonation of the acceptor with subsequent transfer of the OH<sup>-</sup> to the donor via the active-site water network (29). The explicit protonation information on active-site residues determined by static and dynamic (crystallographic and NMR spectroscopy) methods supports mechanisms involving the effects of electrostatics on proton holes and catalysis in HCA II (22, 29). Such a mechanism can operate in conjunction with the classic Grotthuss mechanism, or it can represent an alternate route under certain active-site conditions (30).

#### Conclusion

To our knowledge, this is the first example of using NMR spectroscopy and neutron crystallography complementarily to gain insight into the electrostatics of a protein active site and how such electrostatics affect the charged states and H bonding of amino acid residues. In HCA II, the behavior of Tyr7 has implications for how PT can proceed under different conditions, especially at physiological pH. Similar effects may operate in other enzyme systems that use PT during catalysis. This is a relatively unexplored area and the results are useful for computational biology where experimental knowledge of the positions of H atoms is important. The results show that the protonation states of His and Tyr residues are sensitive to both pH and the presence of metal cofactor. Although this observation seems intuitive, this is the first study to our knowledge that explores these effects in a systematic way. These factors change how His and Tyr residues H-bond to water and other amino acids and we have shown that it is possible to study these directly and at high resolution. It is noteworthy that the combination of neutron crystallography and NMR spectroscopy allows for detailed interpretation of experimental data, creating a powerful combined approach.

## **Experimental Procedures**

Crystal Preparation for Neutron Crystallography. Several batches of wild-type HCA II (~400 mg total) were prepared and the expression and purification protocols are described in detail in Supporting Information. Both apo and holo HCA II were crystallized at room temperature for the neutron experiments by mixing HCA II in a 1:1 ratio with 1.15 M sodium citrate and 100 mM Tris (pH 8.5) in a 500-µL drop. The drops were allowed to undergo vapor diffusion against a 25-mL reservoir consisting of 1.4 M sodium citrate and 100 mM Tris at pH 8.5. To produce the low-pH form, crystals were grown at pH 8.5 using the same protocol for the apo, but before mounting the pH in the drop was lowered to  $\sim$ pH 6 by adding 5 mL glacial acetic acid directly to the reservoir. The sandwich box was closed and the acetic acid vapor acidified the drop. After  $\sim 10$  min the pH in the drop was checked with a microprobe pH meter and it was measured to be below pH 6. Suitable HCA II crystals were selected and mounted in thick-walled guartz capillaries with a liquid plug of perdeuterated reservoir solution. The crystals were left for ~4 wk to undergo vapor H/D exchange before neutron data collection.

**Room-Temperature X-Ray and Neutron Data Collection, Structure Determination, and Refinement.** X-ray and neutron data collection details, the dataset, and refinement statistics can be found in *Supporting Information* and Table S1. Room-temperature X-ray data were collected on an in-house Rigaku HighFlux home source using a standard approach. Room-temperature neutron data to 1.8-Å resolution were collected from a H/D-exchanged *apo* HCA II crystal (~2 mm<sup>3</sup>). These data were collected at the monochromatic neutron diffractometer BIODIFF located at the FRM-II reactor in Garching, Germany. Time-of-flight Laue neutron data to 2.0-Å resolution were collected from an H/D-exchanged, low-pH *holo* HCA II crystal (~2 mm<sup>3</sup>) at the Protein Crystallography Station (PCS) at Los Alamos Neutron Science Center (LANSCE). Experimental data and model coordinates of the *apo* and low pH structures have been deposited with the Protein Data Bank (ID codes 4q49 and 4y0), respectively).

**Isotope-Labeled HCA II for NMR.** Several <sup>13</sup>C isotopically Tyr-labeled versions of wild-type HCA II were prepared for pH titrations to ultimately determine the pK<sub>a</sub> values of the tyrosine residues. Details on expression and purification can be found in *Supporting Information*. The isotopically labeled Tyr used for these experiments was either [U-<sup>13</sup>C, <sup>15</sup>N]tyrosine (CNLM-439-0.5; Cambridge Isotopes) or [<sup>13</sup>Cζ]tyrosine synthesized in-house.

NMR Spectroscopy Measurements and Assignments. All experiments were recorded on 1.0–1.3 mM samples of labeled CA on a Bruker AVANCE III 700 MHz instrument equipped with a TCI cryoprobe and processed using TOP-SPIN 3.2 software (Bruker Biospin). Several spectra at various pH were recorded on labeled HCA II to determine pH-dependent chemical shift

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changes for Tyr signals (Fig. 4). Specific details on the measurements and assignments can be found in *Supporting Information*. Specific assignments of Tyr residues were obtained from a combination of the following 2D and 3D NMR experiments: <sup>1</sup>H-<sup>15</sup>N HSQC, HNCACB, HBCB(CGCD)HD, HBCB(CGCDCE) HE, and aromatic HMBC. Amide proton and nitrogen assignments previously obtained by Venters et al. (31) were used as a starting point. Assignments of <sup>13</sup>C<sub>Y</sub> resonances were obtained from a combination of HNCACB, HSQC, and CG(CB)HB experiments.

**pK**<sub>a</sub> **Determination from NMR Data.** The chemical shifts of all assigned resonances were plotted as a function of pH using Origin 7 software. The data were then fitted to a sigmoidal Boltzmann function using a nonlinear least squares algorithm in Origin 7 and the midpoint of the transition was reported as the pK<sub>a</sub> value. The fits were performed for both C $\zeta$  and C $\gamma$  titrations. The chemical shift error was estimated as 0.01–0.02 ppm based on the resolution of the spectra and repeated experiments.

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