Dissecting Proton Delocalization in an Enzyme’s Hydrogen Bond Network with Unnatural Amino Acids

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Supporting Information

ABSTRACT: Extended hydrogen bond networks are a common structural motif of enzymes. A recent analysis proposed quantum delocalization of protons as a feature present in the hydrogen bond network spanning a triad of tyrosines (Y16, Y32, and Y57) in the active site of ketosteroid isomerase (KSI), contributing to its unusual acidity and large isotope shift. In this study, we utilized amber suppression to substitute each tyrosine residue with 3-chlorotyrosine to test the delocalization model and the proton affinity balance in the triad. X-ray crystal structures of each variant demonstrated that the structure, notably the O–O distances within the triad, was unaffected by 3-chlorotyrosine substitutions. The changes in the cluster’s acidity and the acidity’s isotope dependence in these variants were assessed via UV–vis spectroscopy and the proton sharing pattern among individual residues with 13C nuclear magnetic resonance. Our data show pKₐ detuning at each triad residue alters the proton delocalization behavior in the H-bond network. The extra stabilization energy necessary for the unusual acidity mainly comes from the strong interactions between Y57 and Y16. This is further enabled by Y32, which maintains the right geometry and matched proton affinity in the triad. This study provides a rich picture of the energetics of the hydrogen bond network in enzymes for further model refinement.

Extended hydrogen bond (H-bond) networks are common features in biological systems, including the active sites of many enzymes. In earlier work mapping the active site electrostatic environment of the enzyme ketosteroid isomerase from Pseudomonas putida (pKSI) by vibrational Stark spectroscopy, it was discovered unexpectedly that the low pKₐ of the oxyanion hole of the D40N mutant (which simulates the intermediate state) was associated with titration of a cluster of tyrosine residues.1 This was established by observing characteristic absorption spectral changes associated with the difference between neutral and ionized tyrosine and detailed 13C nuclear magnetic resonance (NMR) studies exploiting the sensitivity of the chemical shift of the carbon adjacent to the tyrosine hydroxyl group (C₆) to the protonation state of tyrosine.1 A key feature of the tyrosine cluster, known from several high-resolution X-ray structures of KSI, is the short O–O distances among their hydroxyl groups.2 The pKₐ of the tyrosine cluster, involving residues 16, 32, and 57 in the active site (Figure 1), was found to be 6.3 in marked contrast to the pKₐ of tyrosine itself in aqueous solution, which is around 10.1. In addition to this large shift in pKₐ, a further unusually large H–D pKₐ isotope effect was observed. Path-integral molecular dynamics simulations were conducted on this system, leading to a model in which proton delocalization (a nuclear quantum effect) plays a key role to explain both the exceptional stabilization of the negative charge on the ionized tyrosine cluster and this large H–D equilibrium isotope effect.3 The simulations showed that protons delocalize because the zero-point energies of their O–H bonds are comparable to the potential energy barriers for proton transfer.

Figure 1. Proton delocalization in the H-bonding network of the intermediate state of KSI.1 In the intermediate state of apo-KSI, protons delocalize among the three tyrosine residues in the active site, leading to an unusual acidity and a large H–D pKₐ isotope effect of the tyrosine cluster.

An important element that determines the proton delocalization equilibrium in the H-bond network of KSI is the relative proton affinities of the tyrosines in the cluster, which are generally thought to be well-matched. In this study, we are interested in learning how sensitive the collective properties of the cluster are to proton affinity detuning at each site, and whether the interactions between different tyrosine residues contribute equally to proton delocalization in the H-bond network. We do so by selectively modifying the proton affinity of each tyrosine site in turn by using amber suppression to...
incorporate the unnatural amino acid 3-chlorotyrosine (Cl-Y, where Cl is ortho to the hydroxyl group) selectively at positions 16, 32, and 57, denoted KSI-Cl-Y16, KSI-Cl-Y32, and KSI-Cl-Y57, respectively. Because the solution pK_a of Cl-Y is 8.3, its incorporation substantially “detunes” the intrinsic proton affinity balance compared with an all-tyrosine cluster without destroying the H-bond network in the cluster. Depending on the contribution of each tyrosine site in the cluster to proton delocalization, we expect quite different effects on both the cluster’s overall pK_a and the H–D isotope effect on the pK_a and this can be used to further refine the model that describes the H-bond network.

**MATERIALS AND METHODS**

**Synthesis of [13C]3-Chlorotyrosine.** To a 25 mL round-bottom flask containing a suspension of [13C]-L-tyrosine (0.25 g, 1.4 mmol, Cambridge Isotope Laboratories; the ω-carbon is adjacent to the hydroxyl group) in 2 mL of glacial acetic acid was added dropwise SO2Cl2 (152 mg, 1.4 mmol, Cambridge Isotope Laboratories; the remaining 19 naturally occurring L-amino acids (25 μg/mL), using previously published methods.1

The recovered solid was dried under vacuum overnight (220 mg, 70%). The purity of the product was assessed by 1H NMR [300 MHz, D2O (Figure S1)] and 13C NMR [500 MHz, D2O (Figure S2)]; this figure also shows the large chemical shift difference, 9.4 ppm, between the protonated and deprotonated forms of [13C]-3-chlorotyrosine.

**Mutagenesis, Expression, and Purification of KSI-Cl-Y Variants.** QuikChange site-directed mutagenesis (Stratagene) was used to introduce the amber codon TAG at position 16, 32, or 57 in the pKSI gene cloned into the pKK223-3 expression vector.5,6 A His tag was also introduced following a thrombin cleavage site at the C-terminus of the protein to facilitate a cleavage step used to elute KSI from a deoxycholate anion-exchange chromatography followed by His_NTA affinity chromatography (Superdex 75 30/300 GL, GE Healthcare). For [13C]-tyrosine labeling of KSI-Cl-Y variants, protein expression was conducted in M9 minimal medium, augmented with 1 mM MgSO_4, 0.2% glucose, MEM Vitamin Solution (GIBCO), 100 μg/mL 13C-labeled tyrosine (Cambridge Isotope Laboratories), 130 μg/mL 3-chlorotyrosine (Sigma-Aldrich), and the remaining 19 naturally occurring L-amino acids (25 μg/mL), using previously published methods.1

**pK_a Values of D40N-Cl-Y Variants.** A series of buffers were prepared with a pH (where L is H or D) between 4 and 10 as described previously. Samples for titration were prepared by combining 60 μL of 150–200 μM protein (in pure L_O), 150 μL of buffer, and 390 μL of L_O.

UV–visible spectra were recorded for each D40N-Cl-Y variant as a function of pH using a PerkinElmer Lambda 25 spectrophotometer with a 1 cm path length quartz cell.

**X-ray Crystallography.** Crystals of all three D40N-Cl-Y variants were obtained using hanging drop vapor diffusion by mixing 1 μL of 20 mg/mL KSI variant with 1 μL of a reservoir solution [1.1–1.4 M ammonium sulfate, 40 mM potassium phosphate (pH 7.2), and 3–6% 2-propanol]. Blade- or cubic-shaped crystals appeared after incubation for 3 days at 20 °C. Cryoprotection was achieved by directly soaking crystals in 2.0 M sucrose prior to flash-freezing in liquid nitrogen. Single-crystal diffraction data for each D40N-Cl-Y variant were collected at Stanford Synchrotron Radiation Laboratory beamlines BL7-1 and BL11-1 and Advanced Light Source (Lawrence Berkeley National Laboratory) beamline BLS02.

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Table 1. Effects of Chlorotyrosine Substitution on KSI-Catalyzed Isomerization of 5(10)-Estrene-3,17-dione

<table>
<thead>
<tr>
<th></th>
<th>$k_{cat}$ (s$^{-1}$)</th>
<th>$K_M$ (μM)</th>
<th>$k_{cat}/K_M$ ($\times 10^6$ M$^{-1}$s$^{-1}$)</th>
<th>$k_{cat}$ ratio (mutant/WT)</th>
<th>$\Delta \Delta G^\circ$ (kcal mol$^{-1}$)</th>
<th>$k_{cat}/K_M$ ratio (mutant/WT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT$^{19}$</td>
<td>9.0 ± 0.2</td>
<td>16 ± 2</td>
<td>54 ± 7</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>KSI-Cl-Y$^{17}$</td>
<td>1.8 ± 0.1</td>
<td>25 ± 2</td>
<td>7.2 ± 0.7</td>
<td>0.20</td>
<td>0.94</td>
<td>0.13</td>
</tr>
<tr>
<td>KSI-Cl-Y$^{12}$</td>
<td>5.1 ± 0.1</td>
<td>11 ± 2</td>
<td>46 ± 8</td>
<td>0.57</td>
<td>0.34</td>
<td>0.85</td>
</tr>
<tr>
<td>KSI-Cl-Y$^{10}$</td>
<td>1.9 ± 0.1</td>
<td>44 ± 3</td>
<td>4.3 ± 0.4</td>
<td>0.21</td>
<td>0.90</td>
<td>0.080</td>
</tr>
</tbody>
</table>

$^a \Delta \Delta G^\circ = \Delta G^\circ$(mutant) − $\Delta G^\circ$(WT). $\Delta G^\circ = RT\ln k_{cat}$. $T = 293$ K, and $R$ is the gas constant. $^b$Values from ref 19. The addition of the LVP-R C-terminus was found not to affect the catalytic rates.

Data were integrated and scaled using mosflm$^{11}$ and CCP4,$^{12}$ respectively. Data collection and refinement statistics are summarized in Table S2. An initial protein model for each structure was obtained by molecular replacement with Phaser$^{13}$ using the coordinates from the previously published structure of KSI-D40N with bound 3-fluoro-4-nitropheno[Protein Data Bank (PDB) entry 3VGN]. The protein model is chosen to minimize the effect of modeling bias in the structural comparisons (especially the active site residues) between D40N-Cl-Y and the wild type (see Results). Simulated annealing refinement was conducted using a maximum-likelihood likelihood-based target function as implemented in Phenix.$^{14}$ Further refinement was conducted with Phenix, interspersed with manual model building in Coot.$^{33}$ All structural figures were prepared using PyMOL.$^{16}$

$^{13}$C NMR Spectroscopy of D40N-Cl-Y. One-dimensional $^{13}$C NMR spectra were recorded on a Varian UNITYINOVA NMR spectrometer (500 MHz) running VNMR version 6.1C and equipped with a 5 mm PFG switchable probe operating at ambient temperature. Samples contained 0.3–0.5 mM KSI, 40 mM potassium phosphate (pH 7.2), 1 mM EDTA, 2 mM DTT, and 5% D$_2$O as the lock solvent. Spectra were acquired in a 5 mm Shigemi symmetrical microtube for approximately 15000 scans and processed using a 10 Hz line broadening function. Peak positions are reported as the chemical shift value corresponding to the position of maximal intensity for each peak. The $^{13}$C NMR spectra of the ionized and neutral form of 3-chlorotyrosine were recorded in D$_2$O at pD 3 and 4, respectively (Figure S2). All $^{13}$C chemical shifts were referenced to an external standard of sodium 3-trimethylsilylpropionate-2,2,3,3-d$_4$ (0 ppm) under identical solvent and temperature conditions.

Three samples were prepared and measured: D40N-Cl-Y$^{16}$ bearing $^{13}$C-labeled Cl-Y$^{16}$ and unlabeled tyrosines (Y$^{52,74,110}$), D40N-Cl-Y$^{16}$ bearing unlabeled Cl-Y$^{16}$ and $^{13}$C-labeled tyrosines at the other three sites, and D40N-Cl-Y$^{15}$ bearing $^{13}$C$^1$-Cl-Y$^{57}$ and unlabeled tyrosines (Y$^{16,32,110}$). For D40N-Cl-Y$^{16}$, initial measurements were performed on the mixture of two populations from the anion exchange column. The samples were then further purified in an effort to elucidate the chemical shifts corresponding to the first population. The fractional ionization of each residue in the tyrosine triad was calculated using a method similar to that described previously.$^{17}$ The chemical shifts of the neutral and ionized amino acids (Y or Cl-Y) in aqueous solution were used as the reference states for fractional ionization calculations.$^{3}$

RESULTS

Preparation and Kinetic Characterization of KSI-Cl-Y Variants. KSI variants bearing Cl-Y at residue 16, 32, or 57 were expressed and purified and found to have the expected masses by ESI-MS. Tandem MS was further performed on the first population of the KSI variants to verify the incorporation of Cl-Y (Figure S4). The expression yield of KSI-Cl-Y variants is 3–5 mg/L, ~5 times lower than that of wild-type KSI (20–25 mg/L), a decrease typical for amber suppression observed in other systems.$^{18}$ The enzymatic activities of the three KSI-Cl-Y variants were characterized and compared with that of wild-type (WT) KSI (Table 1). The proteins retained most of their activity upon incorporation of Cl-Y, with a modest 1.5–5-fold decrease in $k_{cat}$; $k_{cat}/K_M$ decreased by 8–10-fold for KSI-Cl-Y$^{57}$ and KSI-Cl-Y$^{16}$ and remained the same as that of the WT$^{19}$ for KSI-Cl-Y$^{32}$. The relatively small but systematic changes observed in these conservative mutants suggest structural features very similar to those of the wild type; nonetheless, they provide ideal tests for electric field/catalytic proficiency correlations mapped by vibrational Stark spectroscopy,$^{20}$ and this will be discussed in a future publication (Y. Wu and S. G. Boxer, unpublished data).

X-ray Crystal Structures of D40N-Cl-Y Variants. We determined X-ray crystal structures for the D40N variants bearing Cl-Y at positions 16, 32, and 57 (PDB entries 3D82, 3D83, and 3D81, respectively); the key active site residues in the tyrosine triad are shown in Figure 2, and the data collection statistics can be found in Table S2. The overall structures obtained for these variants were quite similar to those determined previously for P. putida KSI bound to equilin (PDB entry 1OH0),$^{21}$ with average root-mean-square deviation (rmsd) values of 0.24, 0.24, and 0.22 Å for the main-chain atoms of apo D40N-Cl-Y$^{16}$, apo D40N-Cl-Y$^{32}$, and apo D40N-Cl-Y$^{57}$, respectively. Similar results were obtained for comparison with the apo WT structure (PDB entry 3VSY); however, the resolution of that structure (1.5 Å) is worse than that of PDB entry 1OH0 (1.1 Å), so the latter was used for our comparisons. In some cases, coordinate changes of up to 1–1.5 Å are observed in regions distant from the active site, for instance, at the $\alpha$-helices around residue 35. Focusing on the active site residues, especially the O–O distances of the tyrosine triad, we found superposition of Cl-Y-containing KSI with WT KSI structures revealed conformational differences on the same scale as the 0.1–0.2 Å estimated coordinate error for the X-ray crystal structures given their resolutions (Figure 2; see also Figure S5). Because small distances can affect the H-bond interactions in which we are interested, we also used an alternative way to estimate the errors in the O–O distances by refining the distances using Refmac5. The resulting O–O distances differ on average from those refined by Phenix by 0.06, 0.10, and 0.06 Å for apo D40N-Cl-Y$^{16}$, apo D40N-Cl-Y$^{32}$, and apo D40N-Cl-Y$^{57}$, respectively. Therefore, within the coordinate uncertainty at the resolution of the X-ray structures, the incorporation of Cl-Y preserves the relative positioning of the residues participating in the H-bonding network. The highly conservative nature of these mutants, which can be achieved only through unnatural amino acid incorporation, allows us to study subtle aspects of the hydrogen bond network.
as well as the catalytic role of active site electrostatics, in a systematic manner. For D40N-Cl-Y57, two orientations of Cl-Y57 with occupancies of 59 and 41% are observed. Compared to Y57 in the high-resolution WT KSI structure, the plane of the benzene ring is rotated approximately 20° toward either direction in the two conformations of D40N-Cl-Y57 (Figure 2). The O−O distances between Cl-Y57 and Y32 and Y16 are 2.73 and 2.42 Å, respectively, for the major conformation and 2.72 and 2.48 Å, respectively, for the minor conformation (Table 2). Two ordered water molecules within 3 Å of the tyrosine residues are observed in the oxyanion hole (Figure S5A).

For both D40N-Cl-Y32 and D40N-Cl-Y16, two asymmetric units are observed in one unit cell. The two chains of each mutant differ by an overall rmsd value of ∼0.19 Å from each other and have similar O−O distances among the three triad residues (Table 2). The two mutants share more similarities in their active site geometry: Cl-Y32 and Cl-Y16 both adopted a single orientation. Their benzene ring is coplanar to their tyrosine counterparts in KSI (PDB entry 1OH0), and the chlorine is pointing away from the H-bond core of the tyrosine triad. The only difference is that two ordered water molecules are found within 3 Å of the triad residues for D40N-Cl-Y16, while only one was observed for D40N-Cl-Y32 (Figure S5B,C).

**pKₐ Values and Isotope Effect of D40N-Cl-Y Variants.**

We measured the pKₐ and the pKₐ isotope effect of the active site of each D40N-Cl-Y variant using UV absorbance spectroscopy (Figure 3). All measurements were taken in the D40N background as this is the pKSI mutant in which the highly perturbed pKₐ of Tyr57 was originally observed. The D40N variant used here differs from that in previous studies in that it has four extra amino acids at the C-terminus (LVPR)
because of the purification procedure. This variant’s tyrosine cluster has a pK_a of 5.8, which differs from that of the previously studied construct (pK_a 6.3) by 0.5 pK_a unit (Figure S6A). Despite this unexpected difference, the large and unusual isotope effect on the pK_a (Figure S6A) matches that found earlier, and the 13C NMR spectrum of [13C,]tyrosine-labeled protein (Figure S6B) is likewise identical to that observed previously. Combined with the high degree of structural similarity described above (it appears that the active site, in particular the H-bonded Tyr cluster, is unchanged), we suggest that the extra four amino acids at the C-terminus, especially the positively charged arginine, are interacting with the water in a way that results in this difference in absolute pK_a. The C-terminus of KSI ends at the side of the enzyme that has a large opening for water passage. The extra LVPR residues increase the length of the unstructured C-terminus from ~6 to 10 residues. Interestingly, we observed a sharp 13C NMR peak at 159.6 ppm in unlabeled (natural abundance) KSI (Figure S7) that was absent in the previous construct without the LVPR C-terminus. It is likely that this peak originates from the fast rotating arginine side chain at the end of the extended C-terminus (the peak position of Arg’s ζ-carbon in water is at 159.67 ppm.)

The intrinsic pK_a of tyrosine (10.1) and 3-chlorotyrosine (8.2) differ by 1.87 pK_a units (Figure S8), and their H-D isotope effects are both close to 0.5 pK_a unit. The high pH–low pH difference spectra for Tyr and 3-Cl-Tyr (Figure S9A) show that the peak of the difference spectrum red shifts by 7 nm in 3-Cl-Tyr (from 293 to 300 nm), which provides a basis for evaluating the primary site of titration in the three D40N-Cl-Y variants. We discuss each D40N-Cl-Y variant separately.

**D40N-Cl-Y57.** In the case of D40N-Cl-Y57 (Figure S9B), a broad peak centered around 300 nm was observed in the difference spectra with extinction coefficient differences similar to that of Cl-Y in solution, identifying Cl-Y as the primary site being titrated across the pH series. The apparent pK_a of D40N-Cl-Y57 was determined to be 5.1, 0.7 pK_a unit lower than that of D40N (Figure 3A and Table 3). Upon H/D substitution, the pK_a of D40N-Cl-Y57 increased by 0.87 pK_a unit, quite similar to that of D40N (Table 3). The apparent pK_a was determined to be 5.7, 0.1 pK_a unit lower than that of D40N (Figure 3B and Table 3). Upon H/D substitution, the pK_a of D40N-Cl-Y57 increased by 0.87 pK_a unit, quite similar to that of D40N (Table 3).

**D40N-Cl-Y16.** The high pH–low pH absorbance difference spectrum of D40N-Cl-Y16 (Figure S9D) overlapped well with that of tyrosine, similar to that of D40N-Cl-Y32, which indicates the ionization of one tyrosine (mostly Y57) upon pH titration of the mutant. The apparent pK_a of D40N-Cl-Y16 is 4.9, 0.9 pK_a unit lower than that of D40N (Figure 3C and Table 3). Upon H/D substitution, the pK_a of D40N-Cl-Y16 increased by 0.70 pK_a unit, a pK_a isotope effect slightly smaller than that of D40N (Table 3). Results for all variants are summarized in Table 3.

**13C NMR of D40N-Cl-Y Variants.** To further dissect the proton delocalization pattern into the roles of individual residues in the triad, we employed site-specific 13C isotope labeling to look at the fractional ionization of each tyrosine residue. The assignment of peaks to each residue is based on the similarity of their chemical shifts to those of D40N as Cl-Y incorporation is expected to have only local effects confined to the tyrosine cluster (i.e., residue 57, the primary site of ionization, would still have the most downfield chemical shift). For D40N-Cl-Y57, one peak at 163.06 ppm was observed for 13C-labeled Cl-Y57 (Figure 4C). For D40N-Cl-Y16, because only very small amounts of these isotopically modified Cl-Y-modified proteins were made, we initially combined the two populations from ion exchange chromatography. In the spectra with both populations, the peak corresponding to 13C-Y19, a residue at the dimer interface far from the active site, has the same chemical shift as D40N (157.3 ppm (Figure 4A)), two peaks at 157.3 and 158.0 ppm were observed for 13C-labeled Cl-Y (Figure 4B), and two peaks at 158.7 and 159.3 ppm were observed for 13C-Y32 (Figure 4A). For 13C-Y57, two peaks at 164.2 and 164.8 ppm were observed (Figure 4A). Measurements on small samples of the separated population showed a single peak for each of the residues in the tyrosine triad (Figure 4A and Figure 4B, bottom). By comparison, the peaks at 158.0, 158.7, and 164.2 ppm are assigned to Cl-Y16, Y32, and Y57, respectively, of the first population. The small peak at 159.6 ppm (Figure 4B, top) is likely from the same arginine side chain at the end of the C-terminus as that in the unlabeled KSI (Figure S7). The signal-to-noise ratios of the spectra in these cases were sufficient to distinguish this relatively sharp peak from the background signals.

### DISCUSSION

**pK_a Values and Isotope Effects of D40N-Cl-Y Variants.** Proton delocalization within the tyrosine triads of the three KSI-Cl-Y mutants is assessed from the shift in their overall pK_a values and the difference in the pK_a isotope effect upon H/D substitution compared to that of D40N. The earlier modeling and analysis of the origin of the abnormally low pK_a of Y57 in D40N (the pK_a of the primary site associated with the titration) demonstrated that the proton affinities of the three tyrosines in the triad are well-matched to allow strong interactions, and that the phenolic protons formally belonging to Y16 and Y32 on the sides of the cluster could delocalize onto Y57 at the center to stabilize its negative charge (Figure 1). Using this as a starting point, we can make some simple predictions for the effect of Cl-Y incorporation at each site and then compare them with the experimental results.

| Table 3. pK_a Values and pK_a Isotope Effects of D40N-Cl-Y Variants |
|-------------------|-------------------|-------------------|-------------------|
|                   | D40N              | D40N-Cl-Y57       | D40N-Cl-Y16       | D40N-Cl-Y32       |
| pK_a (H)           | 5.76 ± 0.11       | 5.15 ± 0.08       | 5.66 ± 0.08       | 4.86 ± 0.13       |
| pK_a (D)           | 6.65 ± 0.09       | 7.67 ± 0.07       | 6.53 ± 0.08       | 5.56 ± 0.04       |
| pK_a shift relative to D40N  |              | −0.65             | −0.10             | −0.90             |
| H–D pK_a          | 0.87 ± 0.14       | 0.61 ± 0.10       | 0.87 ± 0.11       | 0.70 ± 0.14       |
| isotope effect     | (ΔpK_a)           | (+1.22)           |                     |                   |
| shift in H–D pK_a | −0.26             | −0                   | −0.17             |                   |
| pK_a shift relative to D40N  |              |                      |                   |                   |
| primary site of    | Y^57             | Cl-Y57            | Y^32              | Y^16              |
|      titration     |                     |                     |                   |                   |

*“The pK_a and isotope effect of D40N are used as the reference states for comparison. The partial contribution to the pK_a shift from the loss of proton delocalization (see the text).”*
Addition of chlorine lowers the $pK_a$ of tyrosine by 1.8 $pK_a$ units from a pure inductive effect of the Cl substituent (Figure S8). We focus on two questions: how sensitive are the collective properties of the cluster to proton affinity detuning at each site, and do the interactions between different tyrosine residues contribute equally to proton delocalization? We will use the language developed in ref 3 and the simplified model developed by McKenzie22,23 to describe this qualitatively. McKenzie’s model uses a Morse potential at each site to describe the diabatic states that are then coupled by the H-bond.
interactions to give adiabatic potential curves. This is illustrated in Figure S11. The coupling strength depends sensitively on the O−O distance between the H-bond donor and the acceptor. Because the O−O distances among the hydroxyl groups of the triad residues are conserved within the uncertainty of the coordinates of the X-ray structures upon Cl-Y incorporation (Table 2), for each H-bond pair, we will discuss the changes in the cluster’s overall pKₐ in light of the site proton affinities, even though effects arising from subtle changes in geometry cannot be strictly ruled out. This allows us to directly attribute any change in proton delocalization between a Cl-Y variant and D40N to the proton affinity detuning of a single residue. While all three residues should be considered together, within the framework of this two-site model, we treat the H-bond network such that the interaction between Y³⁷ and Y³₂ or between Y³⁵ and Y¹⁶ stabilizes the system independently. For instance, when residue 16 is modified, because the positions of Y³⁷ and Y³⁵’s diabatic potentials are unperturbed, the energetic benefit from their interaction is not significantly altered from that of the wild type. We then define the in situ proton affinity of Y³⁷ as referring to its diabatic potential after coupling with Y³² (Figure S11A, dashed line), which is lower than the diabatic potential well of Y³⁷ alone, and compare it with the proton affinity of Cl-Y³⁶ to assess the additional stabilization effect from this second pairwise H-bonding interaction.

As shown in Table 3, the tyrosine triads of D40N-Cl-Y mutants exhibit noticeable changes up to 1.2 pKₐ units in their acidities from that of D40N, suggesting that proton affinity detuning effectively alters the proton delocalization behaviors in the H-bond network that provide the extra stabilization energy. Meanwhile, the H−D isotope effects (ΔpKₐ) of the mutants appear to respond less sensitively to the perturbation; the variation among the mutants and D40N is smaller compared to the variation in their overall pKₐ values. However, isotope effects larger than that observed for tyrosine or chlorotyrosine in solution (0.46 ± 0.08) are still present in all mutants, indicating that the tyrosine triad in these mutants preserves some features of the nuclear quantum effect observed in D40N.

In the case of D40N-Cl-Y³⁷, the large intrinsic change in the pKₐ of Cl-Y³⁷ should produce a mismatch in intrinsic proton affinity with the other two sites in the cluster. As a consequence, we expect a net loss of proton delocalization to stabilize the ionized residue 37, counterbalancing the fact that the residue, by itself, is more acidic. Indeed, the observed pKₐ shift of D40N-Cl-Y³⁷ relative to that of D40N (0.65 pKₐ unit) exemplifies the outcome of these two competing effects: the reduction in the level of proton delocalization ostensibly decreased the cluster’s acidity by 1.1 pKₐ units (1.8 − 0.65), corresponding to 1.5 kcal mol⁻¹ of lost stabilization energy at 298 K due to proton affinity mismatch. Moreover, the H−D pKₐ isotope effect of D40N-Cl-Y³⁷ is 0.26 pKₐ unit smaller than that of D40N and in fact is quite similar to that of an isolated chlorotyrosine in solution, consistent with our prediction of a smaller degree of proton delocalization in this variant.

In the case of D40N-Cl-Y³⁵, addition of chlorine lowers the intrinsic proton affinity of residue 32, making it more comparable to the in situ proton affinity of Y³⁷ (the adiabatic potential after coupling with Y¹⁶). We might predict a larger degree of proton sharing between the two to stabilize the ionized tyrosine 37, leading to an effective pKₐ even lower than that of D40N. However, a minimal change in the pKₐ of D40N-Cl-Y³⁵ from that of D40N (0.1 pKₐ unit) was observed along with indistinguishable isotope shifts, indicating that there is virtually no effect on the proton delocalization equilibrium when the intrinsic proton affinity of site 32 is modulated by 1.8 pKₐ units. On the other hand, it was previously found that if the tyrosine at position 32 is substituted with phenylalanine, the pKₐ of the cluster returns to that of a normal tyrosine pKₐ of 9.9. On this basis, we posit that the interaction between Y³² and Y³⁷ is more similar to a normal H-bond, which favors the ionization via electrostatic stabilization (i.e., the dipole moment along Y³²’s O−H bond stabilizes a negative charge on the oxygen atom of Y³⁷). This description is consistent with the X-ray structures, in which the Y³² O atom is consistently found to be more distant from Y³⁷ than Y¹⁶. The longer O−O distance leads to a weaker coupling between the two sites; thus, the energetics of the proton in the adiabatic potential is not significantly lowered. These data suggest that the nuclear quantum effect is smaller for this pairwise interaction, which is consistent with the insensitivity of the tyrosine cluster’s pKₐ and H−D pKₐ isotope effect on the precise affinity matching of Y³² to the other members of the triad. As the degree of proton delocalization across the H-bond in the KSI tyrosine triad strongly depends on the separation between the proton donor and acceptor oxygen atoms, it is not unreasonable to view the role of Y³² as a local modulator: it finely tunes the proton affinity of Y³⁷ relative to that of Y¹⁶ and facilitates their closer approximation in the active site, which the overall protein architecture alone cannot enforce, so that the latter two can afford a larger degree of proton delocalization that is responsible for the unusual acidity of the cluster.

The proposed difference in the role of Y³⁷ and Y¹⁶ in proton delocalization in the triad is further tested by the data we presented for D40N-Cl-Y¹⁶. D40N-Cl-Y¹⁶ is expected to have a degree of proton delocalization between Y³⁷ and Cl-Y¹⁶ larger than that of D40N and a pKₐ lower than that of D40N, because the addition of chlorine brings the intrinsic proton affinity of site 16 closer to the in situ proton affinity of Y³⁷ [the adiabatic potential after coupling with Y³² (Figure S11A)]. Indeed, a large overall pKₐ shift of 0.9 pKₐ unit in the active site upon substitution of Y¹⁶ with Cl-Y is observed (Figure S3 and Table 3); this is a striking observation considering that site 16 is not even the primary site of ionization based on the UV−vis difference spectrum. Moreover, the cluster is rendered even more acidic than when the primary site of ionization (Y³⁷) is chlorinated. This provides compelling evidence that the additional stabilization energy of 1.2 kcal mol⁻¹ is afforded by an increased capacity for the phenolic proton on site 16 to delocalize onto its ionized neighbor. Note that in all three mutants, the measured pKₐ reflects the stabilization of the negative charge at site 57 (Figure S9). In D40N-Cl-Y³⁷, chlorine addition directly affects the stabilization of the negative charge on Cl-Y³⁷. However, in D40N-Cl-Y¹⁶ and D40N-Cl-Y³⁵, the inductive effect of chlorine addition on Cl-Y³⁷ or Cl-Y³⁵ is propagated to Y¹⁶ via the coupling of the two sites’ potentials, which changes the proton delocalization equilibrium and leads to the downward shift of the effective pKₐ. The contrast between the large pKₐ shift of 0.9 pKₐ unit upon substitution of Y¹⁶ with Cl-Y and the small shift upon replacement of Y³² in line with the proposal described above: compared to a normal hydrogen bond, the interaction between Y³⁷ and Y¹⁶ is stronger and more sensitive to the relative proton affinity of the two partners. Their interaction stabilizes the ionized active site by allowing proton delocalization between the two residues and is finely tuned via electrostatic interaction with residue 32.
This larger degree of proton delocalization in the active site of D40N-Cl-Y16 suggests that it should possess the largest H–D pK_a isotope effect; however, the measured isotope shift is instead 0.17 pK_a unit smaller than that of D40N, which appears to be inconsistent with what is expected for its effect on the pK_a (Figure 3C and Table 3). We consider two possible explanations for this observation. First, in the UV−vis measurements, the spectra were fitted to a single titration curve to yield the pK_a of the primary site of ionization in the cluster (site S7). Because a larger degree of proton delocalization between Y57 and Cl-Y16 was observed in D40N-Cl-Y16, it is possible that site 16 also exhibits a noticeable amount of ionization (Figure S9D). In this case, the stabilization energy gained from the larger degree of proton delocalization in the cluster could affect the properties of both residues in the pL range of titration and may only be fully accounted for when the isotope effects of both residues are considered. However, because of the overlap of the basis UV spectra of Cl-Y and Y, it is difficult to deconvolute the spectrum to assess the ionization of Y57 and Cl-Y16 separately at each pL. As we chose to fit the data at the peak position of ionized tyrosine, it is possible that we measured the isotope effect of Y57 that reflects only the partial effect of the proton delocalization in the cluster. Second, an alternative explanation for the apparent small isotope effect is that the energy barrier to H (D) transfer is so low in this system that it is described by a single-well potential. In this scenario, delocalization occurs through a classical mechanism, and quantum effects such as zero-point energy and tunneling are no longer necessary for the protons to delocalize. Indeed, in such a situation, one would expect a smaller pK_a isotope effect (which reports on quantum effects). Interpretation of this curious pK_a isotope effect of D40N-Cl-Y16 is further discussed along with the 13C NMR data in the next section.

**Fractional Ionization of Tyrosine Residues Using 13C NMR.** The changes in the proton delocalization equilibrium in D40N-Cl-Y16 and D40N-Cl-Y57 were clearly reflected from their altered pK_a’s and isotope effects relative to those of the WT. To gain insight into how the ionization is shared among the three residues in the variants’ triads, we determined the fractional ionizations of each residue (Figure 4 and Table S3). The 13C chemical shifts of the ζ-carbons of tyrosine and 3-chlorotyrosine are sensitive to the ionization state of the adjacent hydroxyl group, with a dynamic range of approximately 10 ppm from neutral to fully ionized (Figure S2). Moreover, the proton transfers within the triad are much faster than the NMR time scale, so the observed peak position reflects the population-weighted average of all ionization states present for a given residue. These two observations form the basis for the interpretation of fractional ionization: a larger degree of proton delocalization in the active site of a given variant would cause the fractional ionization of individual residues to be more similar and vice versa. This is an approximate, but residue-specific, method for learning about charge localization.

The chemical shift of 13C-labeled Cl-Y57 in D40N-Cl-Y57 is virtually identical to that of the ionized Cl-Y (Figure 4C), suggesting the (exclusive) localization of negative charge on residue S7. This result agrees well with the proton delocalization model and the conclusion from pK_a titration that addition of chlorine at site S7 causes a larger mismatch of the intrinsic proton affinity among the three residues, largely diminishing the extent of proton delocalization in the active site of the variant.

Unlike D40N-Cl-Y57, the residues in the active site triad of D40N-Cl-Y16 all exhibit 13C chemical shifts between that of the ionized and neutral amino acids, indicating significant sharing of the negative charge among the three. The fractional ionization of each residue is calculated for the first population (Table S3 with details of the calculations). Y57 remains as the major ionized residue in the triad bearing the negative charge >60% of the time, suggesting an appreciable height of the energy barriers for proton transfer among the three residues. When compared with D40N, it shows more extensive sharing between the triad residues: the level of fractional ionization of residue 16 increased by ~6% (27%), while the level of fractional ionization of residue S7 decreased by ~5% (62%). The results indicate an increased degree of proton delocalization in the active site of D40N-Cl-Y16 from that of the wild type, in line with the lower effective pK_a observed. A more quantitative analysis and comprehensive explanation for the one peculiar isotope effect await a more rigorous model following the methods described in ref 3, and this will be presented elsewhere (L. Wang, S. D. Fried, and T. E. Markland, unpublished data).

### CONCLUDING REMARKS

In summary, we have observed a consistent correlation among proton affinity matching, extent of proton delocalization, and acidity enhancement within the tyrosine cluster of ketosteroid isomerase, providing further support for quantum delocalization of protons as a mechanism to perturb the tyrosine pK_a in the active site. The extra stabilization energy mainly comes from the strong interactions between Y57 and Y16, which is further buttressed by Y32 to enforce the right geometry and relative proton affinities. As we modulated the proton affinity balance between the active site residues, KSI-Cl-Y variants showed relatively small but systematic changes in the catalytic rate (k_cat) that will be discussed in greater detail elsewhere (Y. Wu and S. G. Boxer, unpublished data). Natarajan et al. recently described the catalytic consequences of substitution with fluorotyrosines at Tyr16 introduced by protein semisynthesis. Negligible changes in KSI’s catalytic efficiency were observed, however, it has been suggested that the level of pK_a detuning in this case may be too small to significantly perturb the overall energetics of strong H-bonds, leading to the insensitivity. The amber suppression strategy in this study allows both larger and multisite energy perturbations to the active site of KSI while minimizing structural changes. Finally, we note that there are interesting parallels between this system and another well-studied example of the effects of delocalization, namely electron hole delocalization, in the special pair (P') of photosynthetic reaction centers. In that case, delocalization shifts the P/P' reduction potential substantially relative to that of a single bacterio(chlorophyll) molecule, making it easier to oxidize, and this is further modulated by local hydrogen bonds. Thus, in addition to the large diversity of chemical properties among individual amino acids and prosthetic groups, nature also exploits delocalization among multiple copies of amino acids and prosthetic groups to further fine-tune their properties.

### ASSOCIATED CONTENT

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.5b00958.
pK_a, 1H NMR, and 13C NMR of [13C_2]Cl-Y, X-ray crystallography statistics, additional UV–vis and mass spectrometry characterizations of D40N-Cl-Y variants, detailed discussions about the two populations of D40N with supporting pK_a and NMR data, and a qualitative illustration of the Morse potential model (PDF)

[PDF]

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**Notes**
The authors declare no competing financial interest.

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We are very grateful to Professor Jiangyun Wang from the Institute of Biophysics, Chinese Academy of Sciences, for making available plasmids pEVOL-ClYRS and for advice on the chemical synthesis of 3-chlorotyrosine. We thank Drs. Aina Cohen, Clyde Smith, Pete Dunten (SLAC), Marc Allaire (ALS), and Nicolas Levinson for help with X-ray diffraction data collection and analysis and Stephen Lynch for help with the NMR measurements. We thank Dr. Lu Wang and Professor Tom Markland for useful discussions about this work.

**ABBREVIATIONS**
Cl-Y, 3-chlorotyrosine where Cl is ortho to the hydroxyl group; KSI, ketosteroid isomerase from P. putida with additional C-terminal amino acids LVPR; KSI-Cl-Y, KSI with incorporation of Cl-Y in the active site used to study catalytic properties; D40N, KSI with the D40N mutation used to study the pK_a of Cl-Y in the active site used to study catalytic properties; terminal amino acids LVPR; KSI-Cl-Y, KSI with incorporation of Cl-Y in the active site used to study catalytic properties; P. putida ketosteroid isomerase.

**REFERENCES**


Supporting information for:

“Dissecting Proton Delocalization in an Enzyme’s Hydrogen Bond Network with Unnatural Amino Acids”

Yufan Wu, Stephen D. Fried, Steven G. Boxer

Department of Chemistry, Stanford University, Stanford, CA, 94305-5012, United States

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Discussions on the preparation of KSI variants

A. Difference in purification strategies

In this study, a His-tag was engineered to the C-terminal of KSI with a thrombin cleavable linker (Table S1), and the expressed proteins were subsequently purified with Ni-NTA, cleaved and then purified by anion exchange chromatography. Compared to the earlier methods in which the protein was purified using a deoxycholate-sepharose affinity column\textsuperscript{1,2}, two main differences stand out: first, the final protein has four extra residues, LVPR, at the C-terminus; second, the protein remains folded as it was expressed in cell without unfolding and refolding \textit{in vitro}. Careful characterizations on these newly prepared proteins are described below to ensure valid connections between this and the past work.

B. The two populations of KSI and comparison with previous constructs

Two populations of KSI are observed on anion exchange chromatography with close elution time (150 and 170mM NaCl) for all variants prepared, with the first population usually the major peak (Fig. S3). The two populations remained separate upon rerunning, showed identical and correct primary sequence as verified by ESI-MS (Fig. S3) and Tandem MS analysis, and appeared as a single band on a native gel (both are dimers). They both preserve the high binding affinity to the transition state analog equilenin as the previous construct. The \textsuperscript{13}C-NMR chemical shifts of \textsuperscript{13}C\textsubscript{\textgreek{z}}-Tyr show differences < 0.7 ppm for the two populations of D40N-C\textsubscript{I}-Y\textsubscript{16} (Fig. 4A and 4B). This suggests only a subtle difference in the folds of the two populations that are non-interconvertible in the native conditions. Note that only single \textsuperscript{13}C-NMR peaks for each Tyr were observed for the previous KSI constructs\textsuperscript{3}. It is possible that the \textit{in vitro} denaturing and refolding of the protein might explore a different folding energy landscape from that \textit{in vivo} and limit the folding heterogeneity. Because the two populations were observed for all proteins with the LVPR C-terminus prepared this way, including D40N with no Cl-Y substitution, we do not think this has anything to do with the two orientations of Cl-Y observed in D40N-Cl-Y\textsuperscript{57}. In fact, the crystal structure of D40N-Cl-Y\textsuperscript{57} was obtained from a purified protein sample that contained \textgreater 80\% of the first population, while the two Cl-Y
orientations were observed in similar amount (59% vs. 41%, Table 2). We expect that the barrier to rotation (ring-flipping) of Tyr is such that these populations would interconvert at room temperature and not be stably separable by ion-exchange chromatography.

The only noticeable difference between the two populations was observed when the pK\textsubscript{a} of the two populations of D40N were measured (Fig. S10). The first population displayed a conventional single site titration curve with a pK\textsubscript{a} of 5.8, while the second population appeared denatured below pH 5 and the resulting incomplete titration curve gave a pK\textsubscript{a} of 6.9. Further characterization on the first population verified that it exhibits the same isotope effect and \textsuperscript{13}C-NMR shifts as the previous construct, suggesting that its active site has the same properties as what was observed using the earlier purification protocol (Fig. S6, also see main text). Therefore, all data discussed in the main text were collected on the first protein population from the anion exchange chromatography. The only exception is the \textsuperscript{13}C-NMR spectra of D40N-Cl-Y\textsuperscript{16}, where the mixed populations were initially measured, with subsequent separation of the two to elucidate the corresponding peaks of the first population.
## Supplementary Tables

### Table S1 | DNA and primary sequence of pKSI (D40N)

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*a highlighted region corresponds to the extra amino acids from the previous construct. His-tag is eliminated by thrombin digestion in the final protein sample.*
### Table S2 | X-ray data collection and structure refinement statistics

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Table S3 | Fractional ionizations of each tyrosine residue in D40N-Cl-Y^{16}{a}

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<td>%ionized (norm)</td>
<td>$\delta$ [ppm]</td>
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<tr>
<td>Tyr32</td>
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<td>38.0</td>
<td>158.7/159.3</td>
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<td>Tyr57</td>
<td>165.5</td>
<td>92.6</td>
<td>164.2/164.8</td>
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<tr>
<td>total</td>
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</table>

\(a\). Fractional ionizations of tyrosine residues are calculated from the equation:

\[
\%I_{\text{raw}} = \frac{\delta_{\text{apo}} - \delta_{\text{AA}}}{\delta_{\text{AA}}^{\ominus} - \delta_{\text{AA}}},
\]

in which $\delta_{\text{apo}}$ is the chemical shift observed for the active site residue, $\delta_{\text{AA}}^{\ominus}$ and $\delta_{\text{AA}}$ are the chemical shifts of ionized and neutral amino acids (Y or Cl-Y) in Fig. S2 and reference 3. The values were further normalized using a uniform arithmetic correction so that the sum of the fractional ionizations of the three residues equals to unity:

\[
\%I_{\text{norm}} = \%I_{\text{raw}} - \left(\sum\%I_{\text{raw}} - 100\right) / 3
\]

The calculation neglects the effect of local protein environment on the inherent chemical shift of the three residues, assuming that their chemical shifts reflect the changes in the fractional ionization in the same way as the amino acids in the aqueous solutions. We regard the absolute value of $\%$ionizations to be estimates and only attempt to see the trends among different KSI variants.

\(b\). Fractional ionizations are calculated for the first population of D40N-Cl-Y^{16}.
Supplementary Figures

Figure S1 | $^{13}$C decoupled $^1$H NMR of $^{13}$C$_\zeta$-3-chlorotyrosine.

The spectrum demonstrates the purity of the $^{13}$C$_\zeta$-3-chlorotyrosine synthesized. $^{13}$C decoupled $^1$H-NMR of the aromatic protons show identical chemical shifts as those with unlabeled 3-chlorotyrosine.

Figure S2 | $^{13}$C NMR of protonated and deprotonated $^{13}$C$_\zeta$-3-chlorotyrosine.

The spectrum demonstrates the purity of the $^{13}$C$_\zeta$-3-chlorotyrosine synthesized and the large chemical shift difference (9.4 ppm) between its protonated (pH 4, acetic acid) and deprotonated forms (pH 13, sodium hydroxide).
Figure S3 | Elution profile of KSI variants on anion exchange column and the corresponding ESI-MS spectra of the two populations.

Two populations of D40N-CIY\(^{16}\) (with \(^{13}\)C\(_{6}\)-CIY\(^{16}\)) were separated and eluted on an anion exchange column with pH 8, 20 mM Tris buffer, 150 mM and 170 mM NaCl concentrations, respectively. The ESI-MS spectra of the two peaks have the identical fragmentation peaks, giving a molecular weight of 15039±3 Da (theoretical MW=15035.6).
Figure S4 | Tandem-mass data on the first population of KSI variants confirming the incorporation of Cl-Y.

From top to bottom, the peptide fragments contain Cl-Y$^{16}$, Cl-Y$^{32}$ and Cl-Y$^{57}$ are identified, respectively, to confirm the unnatural amino acid incorporation. The position of Cl-Y incorporation is highlighted in red.
Figure S5 | 2Fo-Fc electron density map (contoured at 1.0 $\sigma$) for D40N-Cl-Y variants.

(A) The 1.4-Å structure of D40N-Cl-Y$^{57}$: two ordered water molecules are present in the active site, and further H-bonded to another water molecule. (B) The 1.4-Å structure of D40N-Cl-Y$^{16}$: two ordered water molecules are present in the active site, and further H-bonded to another water molecule. (C) The 1.7-Å structure of D40N-Cl-Y$^{32}$: one ordered water molecule is present in the active site, and further H-bonded to another water molecule.
Figure S6 | The first population of D40N with C-terminal LVPR exhibits same isotope effect and $^{13}$C-NMR of labeled tyrosines as the previous construct (no C-terminal LVPR).

(A) pL titration on D40N (first population with LVPR) yielded a $pK_a$ (H) of 5.8 and $pK_a$ (D) of 6.7. The large isotope effect of $0.9 \pm 0.15$ $pK_a$ unit is similar to the previous construct ($1.1 \pm 0.14$ $pK_a$ unit). (B) $^{13}$C-NMR spectrum for $^{13}$C$\_\zeta$-Tyr-labeled D40N (lower spectrum) shows 4 peaks with chemical shift identical to the previously report values (upper spectrum): 157.4 ppm (Y119), 159.6 ppm (Y32), 160.4 ppm (Y16) and 165.5 ppm (Y57). The upper spectrum is reproduced from Fried, S.D. On the Origins of Catalysis by Ketosteroid Isomerase. Ph.D. Dissertation, Stanford University, Stanford, CA, June 2014. Both pieces of evidence support the preservation of the active site proton delocalization despite the addition of four C-terminal residues.
Figure S7 | $^{13}$C-NMR of unlabeled wild type KSI showing signals from protein backbone and a peak at 159.6 ppm assigned to Arg.

$^{13}$C-NMR of unlabeled wild type KSI (1.6mM) showed signals at 170-180 ppm from the protein backbone carbonyls at natural abundance and a sharp peak at 159.6 ppm. This is likely from the fast rotating arginine side chain at the C-terminus of the protein and is absent in KSI lacking this C-terminal Arg$^{3,4}$. This peak is observed in some spectra of labeled KSI with very good signal-to-noise (c.f. Fig. 4).
Figure S8 | UV-vis titration curves of 3-chlorotyrosine (Cl-Tyr) and tyrosine (Tyr) used to obtain the pKₐ difference due to addition of chlorine.

Titration curves of Cl-Tyr (circles) and Tyr (triangles) in buffered H₂O (red) and D₂O (blue). Fractional ionizations of Cl-Tyr and Tyr are monitored by measuring the change in absorption at 306 nm and 300 nm, respectively. The addition of chlorine is calculated to shift the pKₐ downwards by 1.87 pKₐ unit.
Figure S9 | Superposition of UV-vis absorption difference spectra of D40N-Cl-Y variants with amino acids in aqueous solution.

(A) Difference spectra of aqueous 3-chloro-tyrosine (Cl-Tyr, black) and tyrosine (Tyr, red), calculated from pH 10 minus pH 6, and pH 11.5 and pH 7.5 spectra, respectively. Cl-Tyr exhibits maximum absorption at 300 nm and Tyr at 293 nm. (B) Difference spectrum of D40N-Cl-Y^{57} calculated from pH 9 and pH 4, overlaid with aqueous Cl-Tyr. (C) Difference spectrum of D40N-Cl-Y^{32} calculated from pH 9.5 and pH 4, overlaid with aqueous Tyr. (D) Difference spectrum of D40N-Cl-Y^{16} calculated from pH 9 and pH 4, overlaid with aqueous Tyr.
Figure S10 | Different pKₐ’s were observed for the two populations of D40N.

Two populations of D40N were separated and eluted on an anion exchange column with pH 8, 20 mM Tris buffer, 150 mM and 170 mM NaCl concentrations, respectively. The first population exhibited a pKₐ of 5.8 with a normal shaped single site titration curve. The second population showed disrupted absorptions at pH < 5 (red dots), possibly due to the protein denaturation. The partial titration curve gave a pKₐ of ~7.
Figure S11 | Simplified two-site model for the potential energy curves of the H-bond interaction in the tyrosine triad using the interaction between Tyr\textsuperscript{57} and Cl-Tyr\textsuperscript{16} in D40N-Cl-Y\textsuperscript{16} as an example. ⁵

(A) The diabatic potentials of Cl-Tyr\textsuperscript{16} (purple) and Tyr\textsuperscript{57} (blue, solid line) reflect the intrinsic proton affinity of the two residues. The additional coupling of Tyr\textsuperscript{57} to Tyr\textsuperscript{32} lowers (tunes) the potential of site 57 (blue, dashed line), bringing its in situ proton affinity closer to that of Cl-Tyr\textsuperscript{16}. The red and dark blue lines in each potential energy well indicate the zero point energy (ZPE) of H and D, respectively. The ZPE of H defines the proton affinity of the residue. The values on the r axis are based on the O-O distances found experimentally. The arrow next to the y-axis indicates the direction of lower potential energy. (B) The adiabatic energy potential curve from coupling of the diabatic potentials of the two H-bonded residues. The proton in the H-bonded system possesses lower energy. The energy barrier for proton transfer becomes comparable to the ZPE of H, increasing the degree of proton delocalization to provide extra stabilization energy. The difference between the ZPE of H and D becomes smaller in the shallower adiabatic potential (as compare to the diabatic states), causing a change in the pK\textsubscript{a} isotope effect.
Reference


