Manipulation of a cation- π sandwich expands knowledge of the conformational flexibility of mammalian phenylalanine hydroxylase

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ABSTRACT

Phenylalanine hydroxylase (PAH) is an allosteric enzyme responsible for maintaining phenylalanine (Phe) below neurotoxic levels; its failure results in phenylketonuria. Wild type (WT) PAH equilibrates among long-lived conformations, including resting-state (RS-PAH) and activated (A-PAH), whose equilibrium position depends upon allosteric Phe binding to the A-PAH conformation. The RS-PAH conformation of WT rat PAH (rPAH) contains a stabilizing cation- π sandwich between Phe80, Arg123, and Arg240, which cannot exist in the A-PAH conformation. We report intrinsic protein fluorescence, enzyme kinetics, native PAGE, size exclusion chromatography, limited proteolysis, and ion exchange behavior for F80A, F80D, F80L, and F80R. These data indicate that amino acid substitution at Phe80 affects both RS-PAH and A-PAH conformations so that intermediate, on-pathway conformations are longer lived. For all variants, Phe addition promotes accumulation A-PAH conformation. Kinetic of the characterization of F80A and F80D shows allosteric activation while F80L and F80R are constitutively active. The reaction rates of all Phe80 variants suggest relief of a rate-

determining conformational change present in the wild type protein. Limited proteolysis of WT rPAH in the absence of Phe reveals facile cleavage within a C-terminal 4-helix bundle, reflecting dynamic dissociation of the PAH tetramer in the RS-PAH conformation. Under these conditions, the Phe80 variants show proteolytic hypersensitity in a linker that repositions in the RS-PAH to A-PAH conformational interchange; this cleavage is protected by addition of Phe. We conclude that manipulation of Phe80 perturbs the conformational space sampled by PAH, increasing the population of conformationally dynamic intermediates in the RS-PAH and A-PAH interchange.

Mammalian phenylalanine hydroxylase (PAH, E.C. 1.14.16.1) is a ~453 amino acid, multi-domain, multimeric protein that catalyzes the conversion of phenylalanine (Phe) to tyrosine using tetrahydrobiopterin (BH₄), O₂, and a nonheme iron (1). The role of PAH in human biology is to maintain Phe levels below the threshold for neurotoxicity and sufficient for normal metabolism pigment (e.g. protein, and neurotransmitter biosynthesis). Failure in this

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role results in the most common inborn error of amino acid metabolism, phenylketonuria (PKU) (~1:10,000 births worldwide). Without treatment, PKU has severe neurologic consequences (2). To ensure Phe homeostasis, PAH populates alternate conformations, which include a tetrameric resting state PAH (RS-PAH) conformation and a tetrameric activated PAH (A-PAH) conformation (3). Although their role in PAH allostery remains speculative (e.g. (4)), dimeric conformations are also part of the equilibrium of PAH assemblies.

Figure 1A shows PAH segments that reflect known PAH structures as well as the RS-PAH ⇔ A-PAH interconversion. The sequences of rat and human PAH are compared in Fig S1. The transition between RS-PAH and A-PAH conformations minimally involves a dramatic reorientation of protein domains within each protomer of the multimers (compare Figs 1B and 1C). The reorientation has been proposed to involve the N-terminal 25 - 30% of each protomer (residues 1-117 or residues 1-128; segments 1-4) (4,5). Specific to the RS-PAH conformation is an intra-subunit auto-inhibitory interaction at residue ~ 23 (in segment 2), which partially occludes the enzyme active site (within segment 6), presumably dictating low activity. The A-PAH conformation does not have the autoinhibitory interaction, but contains an intersubunit interface involving residues ~43-77 (within segment 3, a.k.a. the ACT domain), which can be stabilized by allosteric Phe binding through a conformational selection mechanism (4,6). ACT domains serve as ligand sensors in many proteins (7-9). The A-PAH conformation has a more accessible active site, presumably dictating high activity. The Phe-modulated equilibrium among alternate PAH conformations allows PAH to respond to protein intake or catabolism and regulate blood Phe with a more nuanced sensitivity than would be possible for K_M-dependent substrate level control. An individual's basal Phe level can simplistically be thought of as the equilibrium tipping point between the RS-PAH and A-PAH conformations (3). The tipping point for most humans is in the range of $\sim 50 - 120 \mu M$ Phe, while individuals living with PKU can exhibit significantly higher Phe levels (300 μ M – 2.5 mM). PKU is a highly heterogeneous recessive disorder, with more than 1000 disease-associated alleles (10). PKU treatment is further complicated by the fact that the neurologic consequences of a given Phe level can vary considerably among individuals. Treatment strategies under development are diverse (11), many of which would benefit from a deeper understanding of the conformational space available to PAH.

The tetrameric RS-PAH conformation is well described for full-length mammalian PAH (Fig 1B), with the exception of disorder in the Nterminal ~20 amino acids (segment 1), the enzyme active-site lid (within segment 5), and a few C-terminal residues (e.g. PDB: 5DEN, 5FGJ, 6N1K) (5,12,13). Alternate models exist for the A-PAH conformation, some of which are supported by small angle X-ray scattering (SAXS) data, and a crystal structure of allosteric Phe bound to a portion of the protein (residues 34-111, segment 3, the ACT domain, PDB: 5FII) (4,5,12-15). One such SAXS-supported A-PAH conformation model is illustrated (Fig 1C) (5); it dimerized regulatory places the domain considerably farther from the protein's center of mass relative to our original, other previous, and contemporary models (12-15).

There is precedent for single amino acid variants that dramatically alter a protein's multimeric equilibrium and allow crystal structure determination for a physiologically relevant but low-mole fraction assembly in the wild type protein (e.g. (16,17)). Crystal structure determination for the A-PAH conformation has proven elusive and can potentially be revealed using single amino acid substitutions. In the inborn error of metabolism known as ALAD porphyria, such single amino acid variants revealed a structural basis for the disease (18). We have proposed that disease-associated single amino acid variants of PAH can contribute to PKU by shifting the equilibrium between the RS-PAH and A-PAH conformations, thus altering the Phe concentration required to promote activation (i.e. raising the tipping point) (3). To potentially allow A-PAH structure determination, herein we characterize variants designed to destabilize the RS-PAH conformation by disrupting a single intra-subunit interaction. The targeted interaction is a cation- π sandwich, which, depending on the environment, has been estimated to introduce between -45 and +30 kcal/mole to protein with stabilization enhanced stability, by

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neighboring residues that neutralize the repulsion between the two cations (19). Figure 2A illustrates the targeted cation- π interaction between Phe80 in the ACT domain, and Arg123 and Arg420, which are near the N- and C- termini of the catalytic domain (see Fig 1A). In this interaction, charge neutralization is potentially provided by Glu66, with Arg68 likely involved in stabilizing the hydrogen bonding network (12). Substantial domain repositioning in the transition from RS-PAH to A-PAH (see Fig 1C), dictates that this intra-subunit cation- π interaction at Phe80 cannot be present in any of the proposed A-PAH conformations (Fig 2B). Although various models for the A-PAH conformation differ in their details, all contain the ACT domain dimer and all place the Phe80 side chain as available for intra-subunit interactions with the catalytic domain and inter-subunit interactions with C-terminal residues.

Herein we describe the results of Phe80 with alanine, leucine, substituting arginine, and aspartic acid. These substitutions all remove the RS-PAH-specific cation- π sandwich, which might be predicted to destabilize the RS-PAH conformation and favor formation of the A-PAH conformation. Substitution with alanine largely removes side chain interactions. Substitution with leucine retains the hydrophobic side chain character. Substitution with arginine imposes a charge repulsion in the RS-PAH conformation. Substitution with aspartic acid potentially adds an RS-PAH-stabilizing salt bridge. We cannot yet predict how these substitutions impact the A-PAH conformation, where we do not know the molecular interactions of Phe80, as an atomic resolution structure is lacking. We can however predict, based on all A-PAH models, that Phe80 is expected to extend from the inner surface of the ACT-domain dimer towards the rest of the protein (see Fig 1C). Herein we present characterization of the F80A. F80D, F80L and F80R variants as a function of Phe using a variety of tools that probe the conformation of mammalian PAH. These include intrinsic protein fluorescence, enzyme kinetics, size exclusion chromatography (SEC), ion exchange chromatography (IEC), and limited proteolysis. Although the results provide significant insight into PAH conformational dynamics, and point to the power of using multiple complementary approaches, these Phe80 variants have not proven promising toward a crystal structure of PAH in the A-PAH conformation.

The PAH conformational landscape minimally includes the RS-PAH \Leftrightarrow A-PAH equilibrium; any short-lived intermediate conformations/assemblies are largely uncharacterized. However, it has long been realized that PAH activation minimally involves releasing autoinhibition through movement of the N-terminal 24 - 30 amino acids (segment 2, which would by necessity drag along segment 1) (20,21). A recent report on a constitutively active rPAH model of a disease-associated hPAH variant, R68S, suggests the existence of conformations wherein autoinhibition is relieved but ACT domain dimerization is not present, thus uncoupling these phenomena (22). One interpretation of a variety of biochemical and biophysical data is that the R68S variant may be "undocked" such that the entire regulatory domain does not have an intra-subunit interaction with the catalytic domain nor an intersubunit interaction with a neighboring ACT domain, and may sample considerable conformational space. The undocked interpretation is consistent with high thermal instability for R68S (22), studies showing that this amino acid substitution interferes with allosteric Phe binding (23), and in vivo studies indicating protein degradation (24,25). In the following description of the Phe80 variants, we find it useful to consider RS-PAH, A-PAH, as well as putatively undocked intermediate conformations. Two new models of such undocked conformations are illustrated in Fig 1D.

Results

Resolution of conformational isoforms for rPAH and the Phe80 variants

Whether purified using the classic phenyl Sepharose affinity method (26), or cleaved from the N-terminal 6-His-SUMO construct, the resulting SDS-pure rPAH (or Phe80 variants) could be resolved into conformationally distinct isoforms using preparative IEC (4). The IEC fractionation profiles provide novel information on the different multimerization properties of Phe80 variants. As reported previously for wild type rPAH, SDS-pure PAH fractionates on IEC

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into distinct conformations as determined by native PAGE (and confirmed by SEC and/or SEC-SAXS) (12). The order of elution at increasing ionic strength is dimer, a faster migrating tetramer, and a slower migrating tetramer. Nearly all of our published work on the RS-PAH conformation of rPAH has used fractions containing predominantly the faster migrating tetramer (4,6,12). Addition of 1 mM Phe to a sample comprised predominantly of a faster migrating tetramer generally results in loss of dimer (if present), migration to a position similar (on both IEC and native PAGE) to the slower migrating tetramer, and accumulation of higher order multimers (4). Although the similarities/differences structural between different rPAH tetramers observed in the absence of Phe remains unclear. IEC effectively discriminates the behaviors of the Phe80 variants relative to wild type. Below we present native PAGE and analytical IEC as a method to compare preparative IEC fractions from the final purification step of each of the Phe80 variants. A later section presents the use of analytical IEC to explore how the multimeric equilibrium of each of the Phe80 variants responds to the presence of different concentrations of Phe.

IEC behavior of Phe80 variants in the absence of Phe analyzed by native PAGE and SEC.

The preparative IEC behavior of the studied proteins fell into two categories. The first, including rPAH, F80A, and F80D, are resolved to show dimer, and two different tetramer conformations as confirmed by native PAGE (Fig 3A). The tetramers appear less well resolved for the F80A and F80D variants relative to WT rPAH. The second category, comprised of F80L and F80R, show a single tetrameric band with little to no evidence for dimer in the preparative IEC fractions (Fig 3B). All variants show evidence of higher order multimers that elute at the higher ionic strength end of the peak. Western blots (not shown) demonstrate these higher order forms to be PAH. The proportion of higher order multimers was variable among preparations. A remarkable property of WT rPAH is the stable composition of the preparative IEC fractions wherein the multimeric components do not reequilibrate upon lengthy storage at either 4 °C or -80 °C. This implies a high activation energy to

the multimeric interconversion process. The same stability could not be demonstrated for F80A or F80D. The second category of variants comprised of F80L and F80R show an (apparent) single tetrameric conformation in the preparative IEC fractions, consistent with the hypothesis that these variants may be in the A-PAH conformation. It is well established that activated PAH is less prone to tetramer dissociation (e.g. (5)), presumably due to the presence of additional inter-subunit interactions (e.g. the ACT domain dimer interface). To more fully characterize the conformational equilibrium of rPAH and the Phe80 variants in the absence of Phe, analytical IEC was carried out for one or more fractions that had eluted from the preparative IEC columns and analyzed by native PAGE (Fig S2). Taken together with the SEC data in Fig 3, these experiments confirm that the order of elution from the IEC column in the absence of Phe is dimer, faster migrating tetramer, and slower migrating tetramer.

Some of the IEC fractions illustrated in Figs 3A and 3B were also subjected to analytical SEC (see Figs 3C and 3D), where the positions of tetramer and dimer were previously established (4,5). An SEC comparison of WT rPAH and the Phe 80 variants (Fig 3C) shows variations in tetramer size, with F80L being largest, followed by F80R, then F80D, and F80A, which elutes at the same size as WT rPAH. We posit that the average size may reflect contributions from the different conformations, with contributions from the A-PAH or undocked conformations causing apparent size increase. The selected an chromatographic fractions also have different dimeric content. The WT rPAH fraction was specifically chosen to contain dimer. However, for the F80L sample, the high dimer content appears to be a result of the freeze/storage/thaw process (compare with Fig S2C). For rPAH and F80L, Fig 3D shows the effect of 1 mM Phe on the SEC profile. As we had previously demonstrated, addition of 1 mM Phe in the SEC column buffer causes the WT rPAH peak to shift to a slightly larger size (4), consistent with our newest model for the A-PAH conformation (Fig 1C) (5). Addition of Phe also results in a diminished population of WT rPAH dimer and slight accumulation of larger components. For F80L, addition of Phe causes the tetrameric

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component to convert to the same size as WT rPAH. This is consistent with F80L containing a considerable population of undocked conformations, which can be drawn to the A-PAH conformation through interaction with Phe. Consistent with this, addition of Phe also causes the F80L variant to lose nearly all of its dimer content. Notably, in the presence of Phe, F80L gains in population of higher order multimers. We have posited that the appearance of such multimers are Phe-stabilized species wherein PAH tetramers are tethered together by formation of inter-assembly ACT domain dimers (4). Formation of such inter-tetramer interactions could be facilitated by the transient undocking ACT domains.

PAH intrinsic protein fluorescence as a measure of PAH conformational change.

A dramatic change in the PAH intrinsic protein fluorescence upon addition of Phe was one of the first indications that an activated form PAH had а significantly different of conformation relative to its resting state (27). A red-shift of ~ 10 - 15 nm in the tryptophan emission profile is routinely observed for mammalian PAH following addition of 1 mM Phe, which is sufficient to fully activate WT PAH. This change, illustrated in Fig 4A, has been used both as an assay for PAH conformational change and to better understand the structural basis of PAH activation (27-31). Intrinsic fluorescence is a readout of the environment of the three tryptophan residues at positions 120, 187, and 326. Trp120 (noted in Figs 1B-D) is in the linker region (segment 4) between the ACT domain and the catalytic domain; its environment will change in the RS-PAH ⇔ A-PAH conformational transition (3,4). A detailed study using human PAH concluded that Trp120 contributes ~ 61% to the total fluorescence emission changes that are Phe-stabilized, while Trp326 and Trp187 (both in segment 6) are less influential (at ~26% and ~13%, respectively) (29). The WT rPAH intrinsic fluorescence in the absence of Phe reflects tryptophan residues that are largely buried in nonpolar environments. In the presence of Phe, the spectra reflects the tryptophans moving to more polar environment, with at least one becoming largely solvent accessible (fluorescence at 350 nm). In addition to illustrating the intrinsic fluorescence of WT rPAH (+/-1 mM Phe), Fig 4A shows the intrinsic fluorescence of the Phe80 variants in the absence of Phe; there is a remarkable similarity in the spectra of F80L and F80R to that of WT rPAH plus 1 mM Phe. In the absence of Phe, the spectra of F80D and F80A look similar to WT rPAH. Addition of 1 mM Phe (Fig 4B) does not dramatically alter the fluorescence spectra of F80L and F80R, but changes the spectra of F80A and F80D to match that of WT rPAH + Phe.

The intrinsic fluorescence results suggest that F80L and F80R are in an A-PAH conformation while F80A and F80D are in an RS-PAH conformation. We might also conclude that addition of Phe allows each of the Phe80 variants to achieve the A-PAH conformation, as for WT rPAH. Note however that each of these interpretations of the fluorescence data reflects the assumption that the observed red shift reports on a gain of the A-PAH population. If, instead, the red shift reports on a loss of the RS-PAH population, these results are also consistent with the possibility that the Phe80 variants populate conformations intermediate between RS-PAH and A-PAH, where Trp120 is solvent accessible. The suggested conformations have the ACT domain undocked from the intra- and intersubunit interactions characteristic of either the RS-PAH or A-PAH conformations; two modelled examples are illustrated in Fig 1D.

Enzyme Kinetics

For determination of apparent K_M and V_{MAX} values for WT rPAH and Phe80 variants, we capitalize on the sensitivity of PAH activity to pre-incubation with Phe as a proxy for the position of the **RS-PAH** \Leftrightarrow A-PAH conformational equilibrium. For example, the initial velocity of WT rPAH is higher when the enzyme is incubated with Phe prior to starting the assay relative to when the enzyme is not preincubated with Phe. This phenomenon arises from preincubation stabilizing the A-PAH conformation and increasing its population. For the Phe80 variants, kinetic parameters were determined for one fraction off the preparative IEC column, as noted in Fig 3. Table 1 includes the fitted kinetic values; Figure S3 shows the fits of the experimental data. A classic approach to evaluating PAH variants is to quantify the fold

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activation observed when the protein is preincubated with Phe. Here we see the expected ~5-fold activation (reported as comparison of V_{MAX}) for WT rPAH. F80D and F80A show lower fold activation 3.2 and 1.8 respectively. consistent with the notion that these amino acid substitutions destabilize the **RS-PAH** conformation, making it easier to attain the A-PAH conformation upon addition of Phe. F80L and F80R are constitutively active, showing fold activations of ~1. Although this is consistent with F80L and F80R as constitutively in the A-PAH conformation, a closer look at the kinetic parameters for all the Phe80 variants can also be interpreted as supporting constitutively active undocked conformations.

The K_M values for PAH reflect a complex function of Phe binding both to the active site and to the allosteric site at the ACT domain dimer interface. Table 1 shows that the K_M values, like the native PAGE and SEC data, fall into two groups. For WT rPAH, F80D, and F80A, the K_M values are all within experimental error of one another, both without preincubation ($\sim 250 \mu M$) and with preincubation (~155 µM). In contrast, the K_M values for F80L and F80R are significantly lower (~70 µM) and insensitive to the preincubation step. We also note that all of the Phe80 variants show V_{MAX} values larger than wild type (under comparable conditions). One way to compare such values is to calculate a simple V/K ratio. Because the kinetics of PAH is a function of a complex allosteric activation by substrate, the numerical values of the V/K ratio are not mechanistically meaningful. However, Table 1 shows a dramatic variation in V/K suggesting that the rate-limiting step for WT rPAH, which is likely a conformational change, is relieved in the Phe80 variants. The expected conformational flexibility of undocked PAH is consistent with relief of a rate limiting conformational change, perhaps involving breaking the Phe80 cation- π sandwich. Finally, we note that the Hill coefficients for WT rPAH, F80D, and F80A are not significantly different from one another (~2), and likely reflect the formation of two ACT domain dimers per activated tetramer. Our published computational analysis of Phe binding to an isolated ACT domain dimer established the independent (noncooperative) binding of two Phe molecules (6); thus binding of one Phe to each ACT domain dimer of a tetramer will afford full activation.

Analytical IEC as a function of Phe.

We had previously established that addition of 1 mM Phe to rPAH results in a change in an analytical IEC profile using 100 µg protein on a 1 ml HiTrap O column, and that ~50 µM Phe shows a profile containing peaks consistent with both the RS-PAH conformation (no Phe) and the A-PAH conformation (1 mM Phe) (4). Herein we use a slightly different procedure (see Fig 5) to compare and contrast the conformational sensitivity of rPAH and the Phe80 variants to Phe. This study provides additional evidence that all the Phe80 variants are neither fully RS-PAHlike nor fully A-PAH-like in the absence of Phe, notably Fig 5A shows that the IEC chromatographic behavior of all the studied proteins responds to the addition of Phe. Fig 5B shows the progressive change for WT rPAH as Phe is varied from 0 to 1 mM in ~half-log increments. The chromatograms reflect a decrease in the amount of dimer present (first notable at 30 µM Phe), a transition to the A-PAH conformation (the dominant species at 100 µM Phe), and accumulation of higher order multimers (significant by 300 µM Phe). For the F80A and F80D proteins, shown respectively in Fig 5C and 5D, the chromatograms show similar effects, with exception that intermediate the Phe concentrations (e.g. 30 µM) show the mixture of isoforms less well resolved. For WT rPAH, F80A, and F80D, the predominant peak in the absence of Phe elutes at 30.8, 31.0, and 31.2 min, respectively, while the predominant peak at 100 µM Phe elutes at 32.0, 31.8, and 32.0 min respectively (see Fig 5A). In the absence of Phe, F80L and F80R show single peaks, whose elution times are at intermediate values, 31.6 and 31.3 min respectively (see Fig 5A). Addition of Phe alters these elution times to 31.9 and 31.8 min respectively. Notably for both F80L (Fig 5E) and F80R (Fig 5F), this effect is seen at Phe concentrations much lower than their K_M values, which are $\sim 70 \,\mu$ M, and likely reflect Phe binding to the allosteric site on the ACT domain dimer. For F80L, the effect saturates at 10 µM Phe; by 30 µM Phe F80L shows about 50% formation of other assemblies (eluting at 32.8 min). Independent data show that the other assemblies

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are likely higher order multimers. In contrast, for F80R, the change in elution time saturates at 30 μ M Phe and there is little evidence for accumulation of alternate assemblies at 100 μ M Phe. From these results, we feel confident in concluding that F80L and F80R are not fully A-PAH-like in the absence of Phe, in support of the hypothesis that Phe80 side chain interactions are important to both the RS-PAH and A-PAH conformations.

Limited Proteolysis

Limited proteolysis is a classic approach to monitoring conformational change and was first applied to PAH by S. Kaufman and coworkers in the 1980s (32). Herein, we probe the conformations of WT rPAH and the Phe80 variants using trypsin and AspN protease, monitoring limited proteolysis under native conditions at various concentrations of Phe. The results provide the strongest evidence that the Phe80 variants sample conformations are different from either RS-PAH or A-PAH and are highly susceptible to proteolysis in the linker region between the regulatory and catalytic domains (segment 4), as would be predicted for undocked conformations. In addition, proteolysis reveals facile hydrolysis within the C-terminal 4helix bundle when PAH is in the RS-PAH conformation, consistent with transient tetramer dissociation. Proteolytic cleavage sites are included in the sequence alignment illustrated in Fig S1.

Trypsinolysis of WT rPAH and the Phe80 variants is shown in Fig 6, which reveals that the most susceptible site for WT rPAH in the absence of Phe is Lys431. Lys431 is within the C-terminal helix (residues 426-453) and largely inaccessible in the tetrameric RS-PAH crystal structures (e.g. PDB: 5DEN). Proteolysis within 4-helix bundle this C-terminal provides extremelv strong evidence that tetramer dissociation (to dimers or monomers) is facile in the resting state. Under these conditions, proteolysis is relatively slow. Furthermore, native PAGE suggests that cleaved chains are unavailable for tetramer reformation; Fig S4 illustrates that the accumulation of dimer parallels cleavage at Lys431. In contrast, at 1 mM Phe, where WT rPAH is in the A-PAH conformation, Lys113 is the primary cleavage

site, and cleavage is more rapid. Lys113 is within a highly basic region identified to have increased proteolytic susceptibility in Kaufman's original studies (33). In the RS-PAH conformation Lys113, though on the protein surface. is involved in intra-subunit interactions with the carboxyl groups of Asp315 and Asp27, tethering segments 2, 4, and 6 (see Fig 1A). Because the equilibrium of PAH conformations is dynamic, it is not surprising that in the absence of Phe we also observe minor products indicating cleavage at Lys113 as well as products showing cleavage at both Lys113 and Lys431. Likewise, in the presence of Phe, we observe minor products indicating cleavage at Lys431. The N-terminal peptide identified as cleavage at Arg111 for WT rPAH at 1 mM Phe is due to the RDK sequence from 111-113. That Phe-stabilized activation results in rapid cleavage at Lys113 is consistent with an extended structure for segment 4 (see Fig 1A) in the A-PAH conformation (Fig 1C).

In keeping with the intrinsic fluorescence data, all four Phe80 variants show trypsinolysis patterns at 1 mM Phe that are qualitatively indistinguishable from WT rPAH (Fig 6), consistent with the hypothesis that all can reach the A-PAH conformation in the presence of sufficient Phe. However, unlike WT rPAH in the absence of Phe, Lys431 is not a primary cleavage site for any of the Phe80 variants; though it is evident as a minor site in the F80A sample. From this, we might conclude that the F80A variant is most like WT rPAH in its ability to populate the RS-PAH conformation (see also Fig 4A). However, all of the Phe80 variants are cleaved at Lys113 in the absence of Phe, and this cleavage occurs more rapidly relative to WT rPAH in the presence of Phe. This hypersensitivity of cleavage at Lys113 is consistent with the Phe80 variants sampling conformations that are neither RS-PAH nor A-PAH, as would be expected for undocked conformations. In all cases of the Phe80 variants, addition of Phe protects the hypersensitivity of Lys113 to tryptic cleavage. To titrate the sensitivity of the Phe80 variants to protection at Lys113, trypsinolysis was carried out at intermediate Phe concentrations (Fig 6). For F80D, F80A, F80L, and F80R protection becomes apparent at > 100 μ M Phe, ~100 μ M Phe, 30 μ M Phe, and < 30 μ M Phe, respectively. These Phe concentrations are consistent with the

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analytical IEC data (Fig 5) and further supports the hypothesis that all of the Phe80 variants, in the absence of Phe, can sample an open conformation wherein segment 4 (the linker between the ACT domain and the catalytic domain) is hyper-susceptible to trypsinolysis, as might be predicted for undocked conformations. The F80R variant, in the absence of Phe, reveals two additional cleavage sites by the appearance of minor products; these are Lys42 and Lys320. Lys42 is at the edge of a mobile loop that allows the allosteric Phe to bind to the ACT domain dimer interface in the A-PAH conformation (6). In the RS-PAH conformation, its neighbor Glu43 is buried and hydrogen bonded to Tyr204 and His208 of a neighboring subunit. This suggests that F80R contains some population that contains the ACT domain dimer. We cannot comment on the susceptibility of Lys320 at this point.

Digestion with AspN protease (Fig S5) shows SDS gel patterns that are fully consistent with the conclusions drawn from the trypsinolysis studies (Fig 6). Although the AspN protease generated peptides were not identified by mass spectroscopy, side-by-side comparison of the cleavage patterns suggests that WT rPAH without Phe is susceptible to cleavage at Asp425 and Asp435. again suggesting considerably instability in the 4-helix bundle that secures the RS-PAH tetramer. In the presence of 1 mM Phe, AspN protease appears to cleave WT rPAH at Asp112, consistent with the exposed linker region (segment 4 in Fig 1A) expected for the A-PAH conformation. Again, in the absence of Phe, all of the Phe80 variants are more susceptible to cleavage in this linker region, showing hypersensitivity (least so for F80A) that can be protected by addition of Phe. With the exception of F80D, where peptides resulting from AspN protease were not identified, all variants achieve a cleavage pattern expected for the A-PAH conformation at 100 µM Phe. Nevertheless, F80D achieves the expected A-PAH-like cleavage pattern at 1 mM Phe.

The limited proteolysis data uniformly support the hypothesis that substitution at Phe80 destabilizes both the RS-PAH conformation and the A-PAH conformation and promotes population of intermediate states for which the regulatory domain is undocked. Prior limited trypsinolysis studies uniformly focused on

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differential cleavage within a "hinge region" between the regulatory and catalytic domains (residues 111-117) (e.g. (22, 34, 35)).Interestingly, we could not identify a previous report of cleavage within the C-terminal helices (Lys431 in Fig 6; Asp 425 & 435 in Fig S5), despite one study wherein this cleavage is present in the illustrated SDS gels (22). Here we report prominent C-terminal helix cleavage for WT rPAH in the absence of Phe, which is protected in the presence of Phe. This observation is consistent with many reports that Phe stabilizes tetrameric PAH (e.g. (4)). Nevertheless, the observation of proteolysis within the C-terminal helices, which should be a fully protected 4-helix bundle in the tetramer, argues that tetramer dissociation is facile and raises the question as to the physiologic relevance of dimers or monomers in PAH allostery. Our original proposal that activation of PAH was accompanied by ACT domain dimerization also proposed that tetramer dissociation might be part of the allosteric mechanism, consistent with the morpheein model of protein allostery (4). Although the mechanistic significance of monomers or dimers in PAH allostery remains unclear, tetramer dissociation and reassociation can contribute to interallelic complementation seen in PKU patients who are compound heterozygotes (e.g. (36,37)).

Discussion

Herein we report the effects of altering a cation- π sandwich that we had identified as stabilizing the RS-PAH conformation in the first crystal structure of full-length mammalian PAH (WT rPAH, PDB: 5DEN) (12), which remains the highest resolution of all structures determined to date for full length mammalian PAH proteins. Phe80, the aromatic component of this interaction was replaced by alanine, leucine, aspartic acid, and arginine. Notably, all of the Phe80 variants could be isolated in large quantities, stored stably, and used for extensive biochemical characterizations. Working with these proteins did not reveal grossly aberrant behavior, with the exception that F80L showed a higher than normal propensity to form discrete higher order Some multimers. of the reported characterizations, if taken in isolation, could be simplistically interpreted as indicating that F80A and F80D are RS-PAH-like in the absence of Phe,

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while F80L and F80R are A-PAH-like under these conditions. However, the sum of many different types of analyses, most notably limited proteolysis, show that all of the Phe80 variants, to some extent, also sample conformations that are neither RS-PAH-like nor A-PAH-like, but rather more likely reflect intermediates in the allosteric regulation of PAH. Like the reported behavior of human porphobilinogen synthase, this is another example where single amino acid substitutions dramatically alter the conformational space sampled by a medically relevant protein and unmask structural conformers that are, in the wild type protein, too short-lived for extensive characterization (38,39). These variants can also provide а glimpse into the aberrant conformational sampling of disease-associated variants.

Identification of intermediates in the RS-PAH to A-PAH transition

The current work expands our understanding of PAH allostery to include undocked intermediates, simplistically described in Scheme 1.

Scheme 1:	Phe
RS-PAH ⇔ undocked-PA	$AHs \Leftrightarrow A-PAH \Leftrightarrow Phe-stabilized A-PAH$

As per the established conformational selection mechanism (6), first suggested by us in 2013 (4), the population of the A-PAH conformation is increased through stabilization by allosteric Phe binding at the ACT dimer interface (circle in Fig 1C). Interestingly, for WT rPAH, the cation- π sandwich at Phe80 appears to act like a lock on the **RS-PAH** conformation. All tested perturbations lead to an increase in the population of undocked conformers. Possible undocked PAH conformations were recently introduced in a study of the rPAH variant R68S, which is a model for an hPAH disease-associated variant (22). The reported rPAH R68S intrinsic fluorescence, activity, and trypsinolysis results are remarkably similar to what we report herein for the F80L and F80R variants, though the tryptic peptides were not identified and cleavage within the C-terminal helix went unnoted (22). Included in the characterization of R68S was a small angle X-ray scattering study that was evaluated as an ensemble of undocked structures

that allowed flexibility in a linker comprised of residues 110-118. Fitting the data to the undocked conformational ensemble was reported as inferior to an admixture comprised of 33% in the RS-PAH conformation (from crystal structures) and 67% in an A-PAH conformation that held the ACT-domain dimer 8 – 10 Å closer to the tetramer center-of-mass relative to that illustrated in Fig 1C (22). Our ongoing SEC-SAXS studies of rPAH and the Phe80 variants with and without intermediate concentrations of Phe will continue to address undocked models. One such best-fit undocked model is included in Fig 1D (Gupta et al, in preparation); the alternative undocked model in Fig 1D was prepared in conjunction with our ongoing molecular dynamics studies of PAH (Ge, Voelz, et al. in preparation).

Refining A-PAH models

The current work contributes to our building improved models of the A-PAH conformation and provides an improved roadmap towards how to engineer PAH as a structural biology target that is more amenable for study using X-ray crystallography and/or single particle reconstruction using cryo-EM. We initiated this study with a simple model of the PAH quaternary structure equilibrium consisting only of the RS-PAH and A-PAH conformations, positing that destabilizing the former would increase the mole fraction of the latter, thus improving our ability to obtain an A-PAH crystal structure. In designing variants to destabilize the RS-PAH conformation, we focused on disrupting the RS-PAH-specific cation- π interaction. Unexpectedly our data show that the chosen alterations to Phe80 destabilize both RS-PAH and A-PAH conformations increasing the population of intermediate undocked conformations. Thus, we might conclude that Phe80 is important to the stability of both RS-PAH and A-PAH conformations. A maturing model for the A-PAH conformation considers whether disordered segments in the RS-PAH crystal structures become ordered in the A-PAH conformation, and how such disorder ⇔ order transitions might include interactions with Phe80. Two suggested models are described below.

One model proposes that disordered Cterminal basic sequences in RS-PAH become ordered in A-PAH through cation- π interactions

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with Phe80. To date, all full-length PAH crystal structures are in the RS-PAH conformation and all are disordered in the C-termini, where the sequence is QKIK(S) (see Fig S1). Each of the proposed A-PAH models (e.g. those shown in Figs 1C and 2B) include juxtaposition of Phe80, on the inner surface of an ACT domain dimer with the C-termini. In the trans conformation, the predicted cation- π interaction is inter-subunit; in the cis conformation, it is intra-subunit (5). Arguing against this model is the A-PAH-like character of F80R, where only 10 µM Phe appears to be required to reach the A-PAH conformation. If Phe80 in the A-PAH conformation is interacting with the basic Cterminal residues, F80R should introduce charge repulsion and thus not be A-PAH-like.

An alternative model considers a disorder to order transition within the N-terminal ~20 residues (segment 1 of Fig 1A). This segment might move into the space between the ACT domain dimers and the rest of the tetramer. As shown in Fig 1C, this is a fairly large space with an 8 - 10Å distance between the surfaces of the catalytic domains and the ACT domains. The Nterminal sequence of rPAH is AAVVLENGVLSRKLS, where the residues conserved with human PAH are in boldface. Such a relocation of the N-terminal residues could potentially be stabilized by hydrophobic interactions with Phe80. The proposed model would have residues ~1-15 threaded into the space between the ACT domain dimer and the rest of the tetramer with residues ~18-32 as a potentially disordered linker. We note that Ser16 (italic) is а phosphorylation site and phosphorylation is known to reduce the Phe concentration required for full activation (40,41). It is possible that phosphorylation of Ser16 might facilitate the relocation of segment 1 through interaction with Asp112. In support of the notion that the positioning of segment 1 is important to the A-PAH conformation is the reported failure of Δ 1-26rPAH to yield an A-PAH crystal structure (13).

Conclusion

The current study provides a deeper understanding of the dynamic behavior of mammalian PAH and the profound effect of Phe80 variants on the conformational equilibrium

among RS-PAH, A-PAH and undocked intermediate structures. If such undocked structures are relatively more common among disease-associated human PAH variants, this would contribute to the off-cited generalization that disease-associated proteins are highly susceptible to degradation (e.g. (25)). Nevertheless, Phe80 variants are not among the ~1000 different documented PKU-associated alleles (42). The sum of studies presented herein argues that Phe80 plays a pivotal role in locking PAH into the RS-PAH conformation, potentially providing the rate-determining step in PAH activation. It has long been known that PAH activation is relatively slow and dependent upon the order of addition of reaction components in vitro (4,43). Our original interpretation of the unusual kinetic properties of PAH included intermediate dimeric (or monomeric) structures in the pathway between tetrameric RS-PAH and A-PAH conformations, in part to account for the slow timescale of the transition, though this remains only a hypothesis (4). The limited proteolysis studies of WT rPAH in the RS-PAH conformation provide strong evidence that tetramer dissociation is facile, consistent with our proposal that multimerization could be a significant part of PAH regulation. As of yet, there has been no systematic inspection of the dynamic oligomerization properties of PKUassociated variants of PAH. Similar to the case of the Phe80 variants reported here, single-residue variants of the wild-type sequence of other proteins, such as a PBGS, have served to successfully predict the behavior and structures of disease-associated and single-residue variants (17, 18, 39).

Experimental procedures *Protein production and purification methods*

Proteins were overexpressed heterologously in an *E. coli* system using using a cleavable N-terminal His-SUMO affinity tag as previously described (5,44). WT rPAH was also expressed and purified using the Shiman method as previously described (4,5,12,26). The plasmids encoding the variants were produced by Genewiz. The resultant PAH contains no non-PAH residues at either end of the full-length PAH protein. The classic two-step purification on Ni-Sepharose using a Tris-HCl buffer as previously

described (5) yielded nearly homogeneous PAH per SDS PAGE. The final purification step uses IEC fractionation (either a 1-mL or 5-mL HiTrap Q column) in 30 mM Tris-HCl, pH 7.4, 15% glycerol, and a 30 column volume linear gradient to 400 mM KCl. IEC fractionation of SDS-pure rPAH is established to separate quaternary structure isoforms (4,12). Two-mL fractions were collected, analyzed by SDS and native PAGE, flash frozen and stored at -80 °C. The preparative IEC column on Phe80 variants ($\sim 15 - 50$ mg per 1 mL column run) yielded 5 - 7 two-ml fractions at ~0.5 - ~6 mg/ml giving a final overall yield of \sim 7 mg purified protein/g cells, for each variant . In some instances, where the HiTrap O column had been heavily loaded (e.g. >~25 mg for the 1mL column), fractions were pooled, diluted with elution buffer to a final salt concentration of <20mM KCl, and the column was re-run in an attempt to isolate fractions containing only one quaternary structure isoform. PAH is routinely stored (either at 4 °C or -80 °C) as the preparative Hi-Trap Q fractions; once brought to room temperature, PAH is generally allowed to equilibrate at RT for ~1 h prior to experimental observations.

Analytical ion exchange chromatography

For each of the PAH proteins, one or more fractions was chosen from the preparative IEC column and used for an analytical IEC investigation of the effect of Phe on the separable conformations. For these studies, 100 µg of protein is diluted to 5 ml of 30 mM Tris-HCl, pH 7.4, 20 mM KCl, the sample is pumped onto a 1 ml HiTrap Q column (0.5 ml/min), followed by a 40 min wash of the same buffer containing a set concentration of Phe (0, 10, 30, 100, 300, and 1000 µM). Elution used a 30 ml linear gradient to the same column buffer at 400 mM KCl. Blank gradients (one or more) were performed at each Phe concentration prior to each protein injection.

Intrinsic PAH fluorescence methods

Intrinsic fluorescence of PAH at a concentration of 0.5μ M subunit is measured at 25 °C by exciting the protein solution with a light of wavelength 295 nm and collecting the emission spectrum between 305 and 400 nm. Intrinsic fluorescence measurements were obtained for each of the Phe80 variants and for

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WT rPAH in the absence and presence of 1 mM Phe.

PAH enzyme kinetics

PAH activity was determined at 25 °C by continuously monitoring tyrosine fluorescence intensity after initiating the assay by addition of PAH, which was at a concentration between 0.1 and 0.4 mg/mL (i.e. ~ 2-8 μ M subunit). Prior to addition to the assay mixture, the protein was preincubated with or without Phe, at the assay concentration, for 1 minute at room temperature. The continuous fluorescence-based assay is based on that of Gersting et al. (45) with modifications introduced by Bublitz (46). Tyrosine fluorescence measurements use an excitation wavelength of 275 nm and an emission wavelength of 305 nm on a PTI Fluorescence System spectrophotometer equipped with an USHIO Xenon Short Arc Lamp and FelixTM for Windows Software. The standard reaction is performed in a final volume of 2 mL containing 20 mM bis-tris propane (pH 7.3), Phe (at concentrations between 10 µM and 1 mM), 10 µM ammonium iron (II) sulfate, 40 µg/mL catalase (800 - 3200 units), 75 μ M BH₄ and 75 µM DTT. Initial velocity reflects the first 50 sec. The initial velocity data were fitted to a 3parameter Hill equation using the SigmaPlot v. 10.0 (Systat Software, San Jose, CA). Calibration curves of standard tyrosine concentrations (10, 20, 30, 40 µM in the assay solution) were obtained on the day of the activity analysis.

Native proteolytic digestion

WT rPAH and the Phe80 variants were subjected to trypsinolysis at 25 °C, at 1 mg/ml and various concentrations of Phe (0, 30 µM, 100 µM, 1 mM), 30mM Tris-HCl pH 7.4, and 150 mM KCl using a PAH:trypsin ratio of 200:1. The digestions were monitored for 1 hour at RT, taking samples at 1, 15, 30, 45, and 60 min; stopping the reaction by addition of 100 µM PMSF. AspN protease digestions used a 1:250 protease:PAH ratio and were monitored at various concentrations of Phe (0, 100 µM, 1 mM). AspN digestion was stopped by exposure to 100 °C for 5 min. Proteolytic digestion was monitored by SDS PAGE using 12.5% PhastGels. Peptides were identified in select digests by separation using BioRad gels and submission to the Wistar

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Proteomics Facility. For WT rPAH, without Phe, proteolysis was also monitored using native PAGE and quantified using Image-J software (47).

Analytical SEC

SEC analyses were performed using a calibrated Superdex 200 10/300 GL column (GE

Healthcare) in 30 mM Tris-HCl, pH 7.4, 150 mM KCl, without or with 1mM Phe at a flow rate of 0.5 ml/min at RT. Samples were from individual preparative IEC fractions, containing 15% glycerol, flash frozen, stored at -80 °C, defrosted at 25 °C, and allowed to sit for an hour. Samples were adjusted to a protein concentration of ~ 1.5 mg/ml when injected onto the pre-equilibrated column.

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FOOTNOTES

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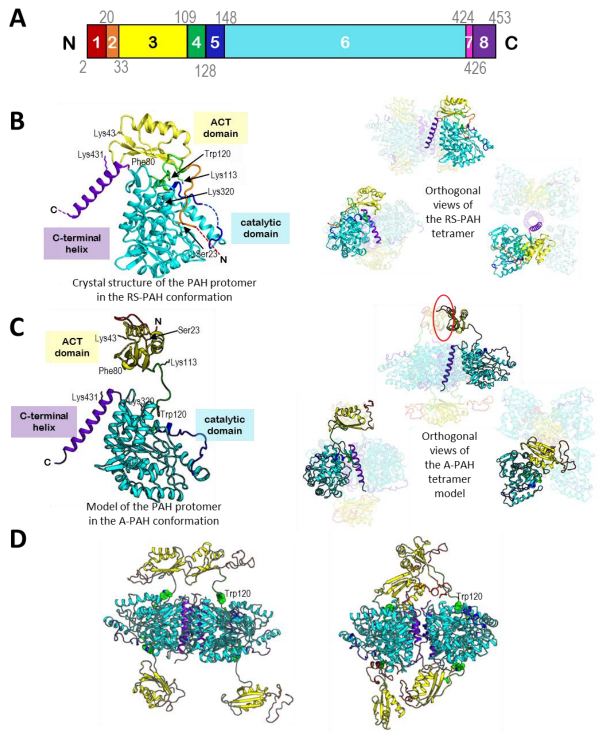
The abbreviations used are: IEC, ion exchange chromatography; Phe, phenylalanine; PAH, phenylalanine hydroxylase; rPAH, rat PAH; PKU, phenylketonuria; PAGE, polyacrylamide gel electrophoresis; SDS, sodium dodecyl sulfate; A-PAH, activated PAH; RS-PAH, resting-state PAH; BH₄, tetrahydrobiopterin; SEC, size exclusion chromatography; SAXS, small angle X-ray scattering; PDB, protein data bank; WT, wild type.

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	Км (μМ)	V _{MAX} (nmol Tyr/min/mg)	Hill coefficient	Fold Activation	V/K
WT rPAH (-Phe)	203 ± 45	465 ± 46	2.2 ± 0.8		2.2
F80D (-Phe)	327 ± 107	$1,210 \pm 230$	1.6 ± 0.5		3.7
F80A (-Phe)	229 ± 18	$3,470 \pm 141$	1.8 ± 0.2		15.1
F80L (-Phe)	59 ± 3	$7,600 \pm 147$	1.3 ± 0.1		129.0
F80R (-Phe)	78 ± 8	$4,100 \pm 151$	1.6 ± 0.3		52.6
WT rPAH (+Phe)	158 ± 58	2,560 ± 337	2.6 ± 1.7	5.5	16.2
F80D (+Phe)	133 ± 33 142 ± 12	$3,890 \pm 164$	3.2 ± 0.7	3.5	27.4
F80A (+Phe)	180 ± 15	6,310 ± 208	2.2 ± 0.2	1.8	35.1
F80L (+Phe)	65 ± 6	$8,\!130\pm282$	1.2 ± 0.1	1.1	125.0
F80R (+Phe)	79 ± 7	3,880 ± 142	1.4 ± 0.2	0.9	49.1

Table 1: Kinetic Parameters for WT rPAH and Phe80 variant in the absence or presence

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Two example tetramer models of undocked PAH

Figure 1. *The structures of phenylalanine hydroxylase (PAH).* (A) The domain structure of mammalian PAH is comprised of eight segments. Segment 1 (red) is disordered in RS-PAH crystal structures. Segment 2 (orange) contains the autoinhibitory interaction specific to the RS-PAH conformation. Segments 1 + 2 have previously been called the autoinhibitory region. Segment 3 (yellow) is the ACT domain. Segments 1 - 3 are commonly called the regulatory domain. Segment 4 (green) is a linker region, some or all of which

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repositions during the RS-PAH ⇔ A-PAH conformational interchange. Segment 5 (blue) contains the active site lid (residues $\sim 136 - 142$ are disordered in crystal structures of the RS-PAH conformation). Segments 5 & 6 (blue & cyan) comprise the catalytic domain. Segment 7 (pink) is a hinge whose conformation differs among the four chains in all documented RS-PAH tetramer structures (5,12,13). Segment 8 (purple) is a C-terminal helix, which has been seen as both bent and straight, and which forms a 4-helix bundle; it does not contain the classic pattern of leucine residues found in other aromatic amino acid hydroxylases (48). The last three to six C-terminal amino acids are disordered in all crystal structures of the RS-PAH conformation, with chain-to-chain variation within each structure. (B) The X-ray crystal structure of WT rPAH in the tetrameric RS-PAH conformation (PDB: 5DEN (12)) is shown. One subunit, colored as part A, is enlarged on the left. Three orthogonal illustrations of the tetramer are shown on the right with three subunits faded. In this conformation, the ACT domain of each subunit is docked to the catalytic domain of a neighboring subunit. On the left image, we show in sticks the side chains of residues Phe80, Trp120, Lys113, Lys431, Lys43, and Lys320, which are discussed in this paper. (C) An A-PAH model is shown in the "trans" conformation, as previously described (5); one subunit is enlarged on the left with the same residues depicted. Three orthogonal illustration of the tetramer are shown on the right with three subunits faded. In this conformation, the ACT domain of one subunit is docked to the ACT domain of another subunit (red circle). This is where the allosteric Phe binds (14). The RS-PAH and A-PAH conformations differ in the position of the regulatory domain relative to the rest of the protein, but both conformations contain ACT domains that are docked. Only the A-PAH conformation includes dimerized ACT domains, docked to each other across the diagonal of the tetramer. All proposed A-PAH conformations have uncertainty in the position of segments 1 + 2. (D) Two models of "undocked" tetrameric PAH conformations are illustrated. The model on the left was prepared using the program SASSIE (49) and is part of a detailed SEC-SAXS analysis of the Phe80 variants (Gupta et al, in preparation). The model on the right is the result of a molecular dynamics investigation of tetrameric PAH (Ge, et al, in preparation), expanding methods we previously applied to a Phe-bound ACT domain dimer (6).

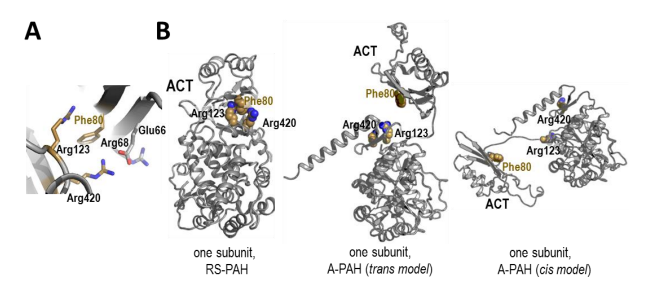


Figure 2: *The juxtaposition of Phe80, Arg123, and Arg420 of one PAH protomer in alternate conformations.* (A) The cation- π sandwich first identified in the rPAH crystal structure (12), with potential charge neutralization provided by Glu66 and involvement of Arg68. (B) Alternate positioning of residues Phe80, Arg123, and Arg420 (shown in space filling) in different PAH conformations. The RS-PAH crystal structure is left and shows the cation- π sandwich. The commonly assumed A-PAH trans conformation is in the center, while the recently proposed A-PAH cis conformation is right (5). The A-PAH conformations cannot include cation- π interactions between Phe80, Arg120, and Arg123.

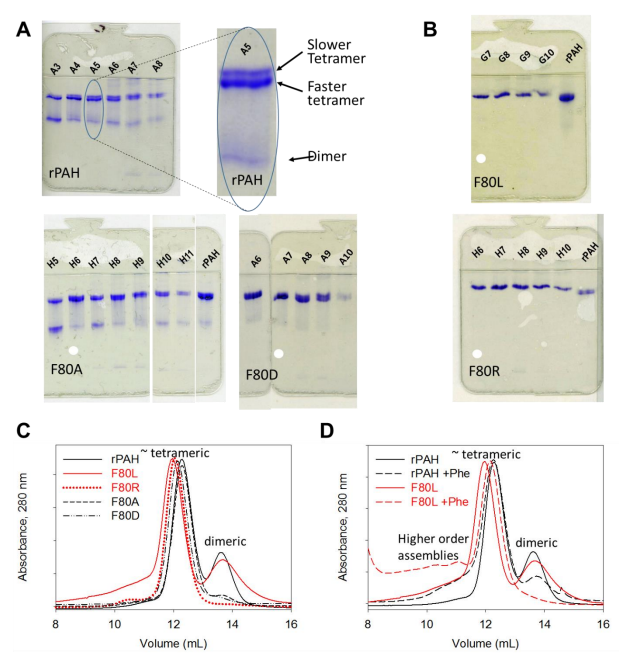


Figure 3. Individual fractions from the preparative IEC purification step were analyzed using 12.5% native *PhastGels and analytical SEC*. The white dots indicate fractions used for determination of enzyme kinetic parameters. (**A**) WT rPAH, F80A, and F80 D show multiple quaternary structure isoforms. WT rPAH fractionates into a dimeric and two tetrameric components. Shown right is a 3.5x enlargement of fraction A5. F80A and F80D behave similarly, though the tetramer bands are less sharply resolved. (**B**) Preparative IEC fractions F80L and F80R show only one tetramer band and no significant dimer. (**C**) Analytical SEC on preparative IEC fractions of WT rPAH and all the Phe80 variants following storage at -80. (**D**) WT rPAH and F80L chromatograms are compared in the presence (dashed lines) and absence (solid lines) of 1 mM Phe in the column buffer.

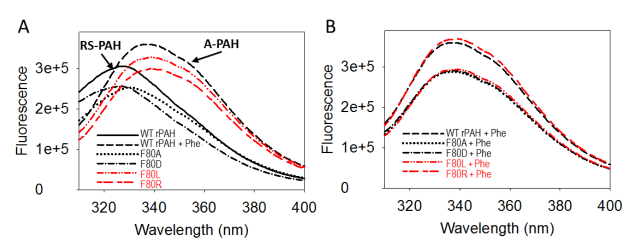


Figure 4. *Comparative intrinsic fluorescence of rPAH and the Phe80 variants.* (A) Emission spectra are shown for rPAH and the Phe80 variants in the absence of Phe. The dashed black line shows WT rPAH in the presence of 1 mM Phe. (B) Emission spectra are shown for rPAH and the Phe80 variants in the presence of 1 mM Phe.

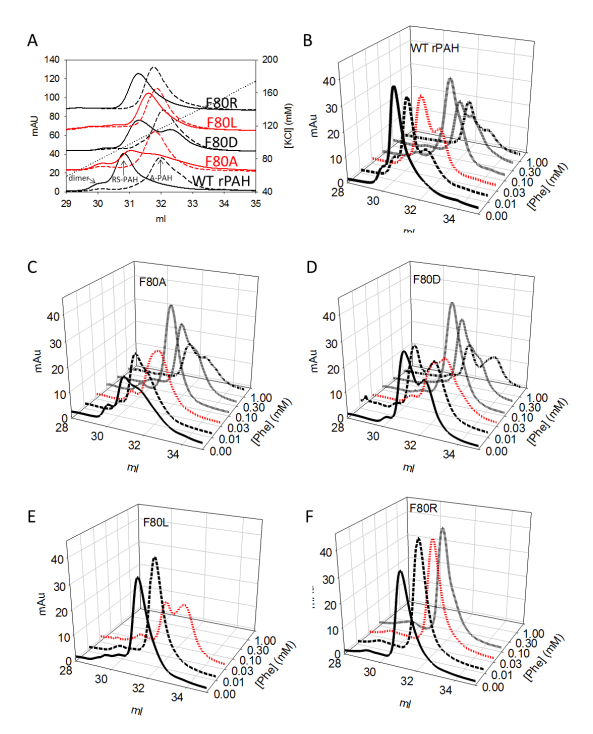


Figure 5: Analytical IEC on rPAH and Phe80 variants as a function of [Phe]. For all separations, 100 μ g of protein was diluted into 5 ml start buffer (no Phe) and applied to a 1 ml HiTrap Q column that had been pre-equilibrated with the set concentration of Phe (10 min). The column was washed with is starting buffer for 40 min and protein isoforms were eluted with a linear gradient to 400 mM KCl at the set Phe concentrations that corresponds to the predominant peak in the A-PAH conformation. For WT rPAH, F80A, and F80D, this is 100 μ M Phe; for F80L it is 10 μ M Phe; for F80R it is 30 μ M Phe. Parts B-F illustrate the progression of chromatographic changes for each variant as the Phe concentration is increased. To facilitate comparison, the chromatogram at 30 μ M Phe is shown in red.

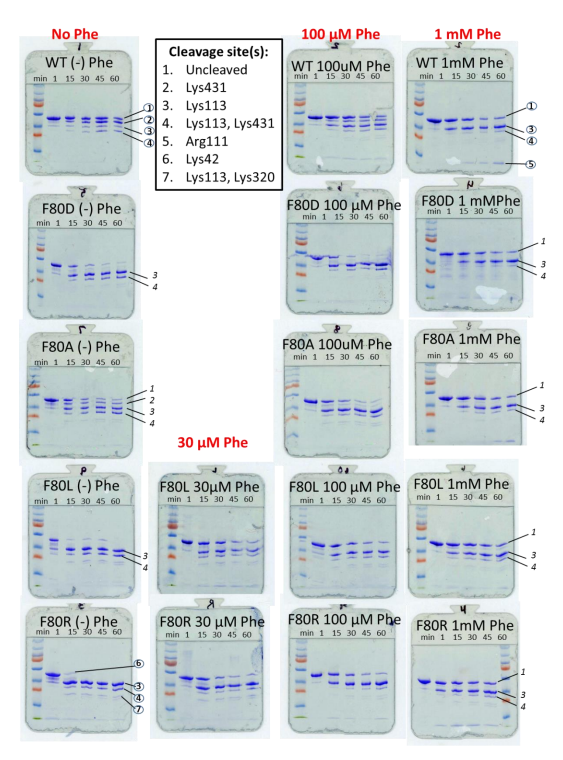


Figure 6: *PhastGel analysis of tryptic digests as a function of time and [Phe].* SDS PAGE analysis of a tryptic digest time course for WT rPAH and the Phe80 variants. Circled numbered bands for WT rPAH no Phe, WT rPAH + 1 mM Phe, and F80R no Phe were identified by submission to the Wistar Proteomics Facility using standard mini-gels. For all other bands numbered in italics, identification is by inference. A similar study using AspN protease is shown in Fig S3. The annotated PAH sequence is in Fig S1.