







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Molecular basis of regio- and stereo-specificity in biosynthesis of bacterial heterodimeric diketopiperazines

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Bacterial heterodimeric tryptophan-containing diketopiperazines (HTDKPs) are a growing family of bioactive natural products. They are challenging to prepare by chemical routes due to the polycyclic and densely functionalized backbone. Through functional characterization and investigation, we herein identify a family of three related HTDKP-forming cytochrome P450s (NasbB, Nas_{S1868} and Nas_{F5053}) and reveal four critical residues (Qln65, Ala86, Ser284 and Val288) that control their regio- and stereo-selectivity to generate diverse dimeric DKP frameworks. Engineering these residues can alter the specificities of the enzymes to produce diverse frameworks. Determining the crystal structures (1.70–1.47 Å) of Nas_{F5053} (ligand-free and substrate-bound Nas_{F5053} and its Q65I-A86G and S284A-V288A mutants) and molecular dynamics simulation finally elucidate the specificity-conferring mechanism of these residues. Our results provide a clear molecular and mechanistic basis into this family of HTDKP-forming P450s, laying a solid foundation for rapid access to the molecular diversity of HTDKP frameworks through rational engineering of the P450s.

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Natural products derived from tryptophan-containing diketopiperazine (TDKP) comprise a large class of secondary metabolites¹⁻³. Among them, heterodimeric tryptophan-containing diketopiperazines (HTDKPs) are particularly attractive for their unique structural architecture and fascinating bioactive properties, ranging from anticancer, antiplasmodial, anti-HIV, and neuroprotective activities¹⁻⁶. TDKPs are primarily produced by fungal systems, in which two pyrroloindoline units are predominantly fused together by a C3-C3' bond^{3,7,8}. Bacterially-sourced HTDKPs are much rarer, but their structural architectures are more versatile^{4-6,9-11}. Based on their connectivities and stereochemistry, the dimeric DKP frameworks can be classified into five different types (Fig. 1): (I) C3-C7', 2R-3S (e.g., naseazepine B or NAS-B^{4,9}); (II) C3-C7', 2S-3R (e.g., NAS-3⁶); (III) C3-C6', 2S-3R (e.g., naseazepine C or NAS-C⁵); (IV) C3-C6', 2R-3S (e.g., iso-NAS-B¹¹); and (V) N1-C7' (e.g., aspergilazine A or Asp-A¹⁰).

The regio-specificity and stereo-specificity in the densely functionalized frameworks, especially at the quaternary stereocenter at the C3 position, renders chemical synthesis of bacterial HTDKPs very challenging^{9,12-14}. To develop efficient biocatalytic approaches, we recently investigated the biosynthesis of naseazepine C (NAS-C) and identified a key diketopiperazine (DKP) forming P450 enzyme (NascB)⁶. NascB catalyzes a radical cascade reaction to form intramolecular and intermolecular carbon-carbon bonds with both regio-specificity and stereo-specificity, which is very efficient in constructing the HTDKP frameworks and has been used to create 30 type I-IV NAS analogs employing different DKP substrates⁶. Very recently, Li and coworkers further identified two other P450 enzymes, AspB and NasB, which are responsible for the predominant formation of aspergilazine A (ASP-A) and NAS-B, respectively¹¹ (Supplementary Table 1). Unusually, HTDKP-forming P450s have relaxed regio-specificity and stereo-specificity and can generate

products with different frameworks, e.g., AspB is able to convert cyclo-L-Trp-L-Pro (cW_L-P_L) into NAS-C (type III) and iso-NAS-B

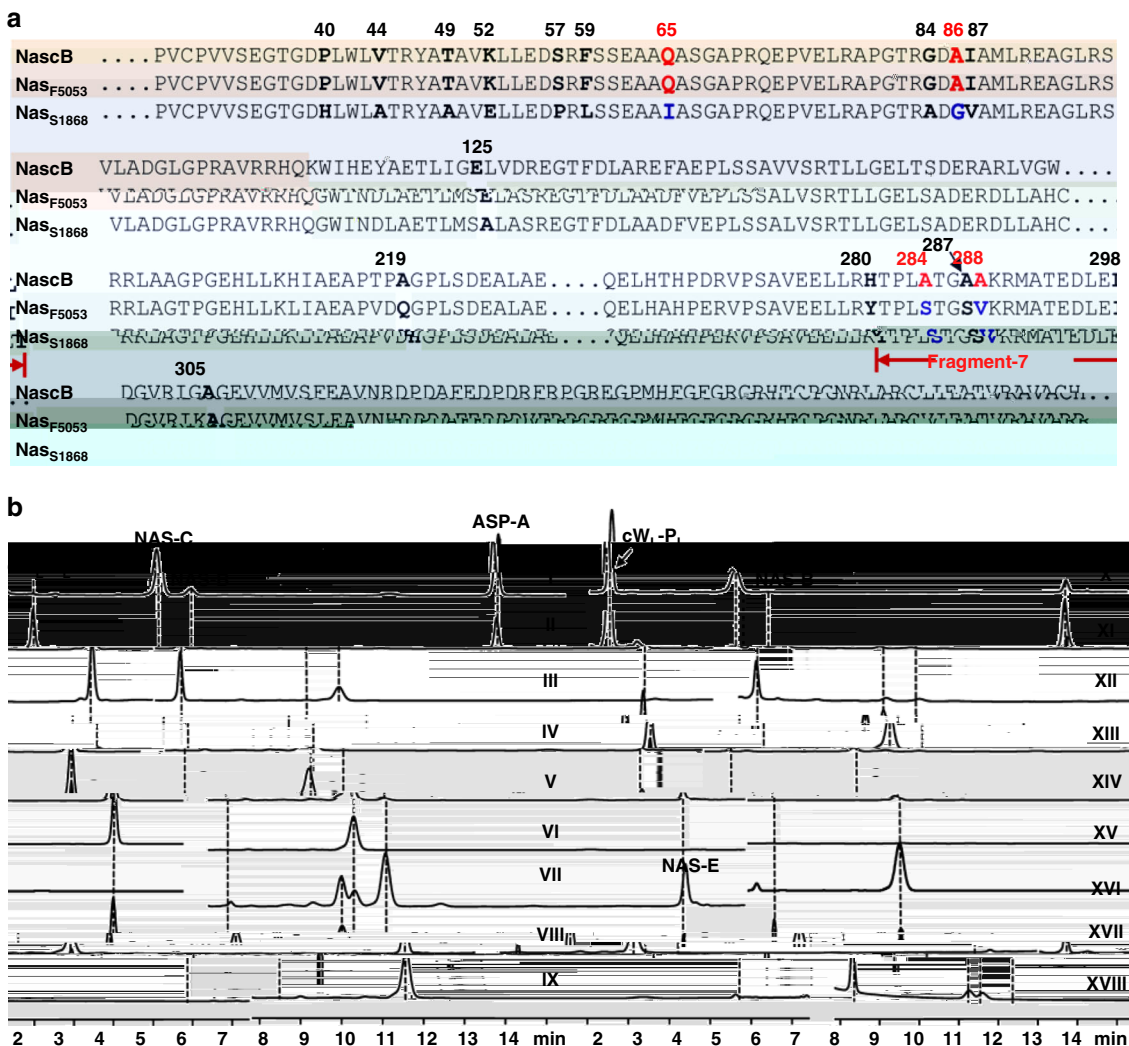


Fig. 2 Deciphering and engineering the regio-specificity and stereo-specificity of HTDKP forming P45Os. **a** The sequence alignments of NascB, Nas_{F5053}, and Nas_{S1868}. The identical N-terminal and C-terminal parts of NascB/Nas_{F5053} and Nas_{F5053}/Nas_{S1868} are shaded in orange and light green, respectively. Residues less important are not shown and indicated by dashed lines. The four critical residues are bolded and highlighted by colors. **b** In vitro characterization of P45Os and their mutants using cW_L-P_L as substrate. (I) Nas_{F5053}; (II) Nas_{S1868}; (III) NascB; (IV) Nas_{F5053}-Q65I; (V) Nas_{F5053}-A86G; (VI) Nas_{F5053}-Q65I-A86G; (VII) NascB; (VIII) NascB-Q65I; (IX) NascB-A86G; (X) NascB-Q65I-A86G; (XI) NascB-S1868fragment-7; (XII) NascB-Q65I-A86G-A284S-A288V; (XIII) Nas_{F5053}-S284A; (XIV) Nas_{F5053}-V288A; (XV) Nas_{F5053}-S284A-V288A; (XVI) Nas_{F5053}-A86K-V288P; (XVII) NAS-E synthetic standard; (XVIII) Nas_{F5053}-Q65P-A86W-S284C.

recycling system (NADP, glucose and glucose dehydrogenase), NascB was confirmed to efficiently dimerize the cW_L-P_L into NAS-B (Fig. 2b, trace III; NMR and HRMS data see Supplementary Fig. 1, Supplementary Fig. 2 and Supplementary Table 2). In the meantime, Li et al identified a homologous P450 (NasB, 96% identity to NascB) from *Streptomyces* NRRL S-1868, also generating NAS-B¹¹.

As the P450 enzymes with highly similar sequences can produce different products, we were interested in establishing the relationship between enzyme sequences and the corresponding products. Although NascB and NascB share 68% sequence identity, it is difficult to extract the key residues responsible for the difference in product formation. In order to identify more P450s which could potentially generate some other kinds of C3-aryl pyrroloindolines, we used simple sequence searches to find genes homologous to *nascB* or *nasbB*. We identified two previously uncharacterized P450 proteins: Nas_{F5053} and Nas_{S1868} from the *Streptomyces* strain sp. NRRL F-5053 and sp. NRRL S-1868, respectively. Soluble recombinant Nas_{F5053} (CYP

nomenclature: CYP1190B1) could be expressed in *E. coli* BL21 (DE3), while soluble Nas_{S1868} (CYP nomenclature: CYP1190B1) could only be expressed in *Mycobacterium smegmatis* MC² 155.

Using an in vitro assay employing the electron-transport system (Fd and FdR) from spinach, Nas_{S1868} can convert cW_L-P_L into Asp-A (Fig. 2b, trace II; NMR and HRMS data see Supplementary Fig. 2, Supplementary Fig. 3 and Supplementary Table 3); a similar observation was also made by Li et al.¹¹). Further, the in vitro assay confirmed that Nas_{F5053} could produce NAS-C (47.4%), Asp-A (44.4%), as well as a minor product NAS-B (8.2%) (Fig. 2b, trace I). This catalysis profile is well correlated with the sequence alignments of Nas_{F5053}, NascB, Nas_{S1868} and NascB. Based on the sequence alignments, Nas_{F5053} can be viewed as a chimeric form of NascB and Nas_{S1868}. The first 112 residues of Nas_{F5053} are exactly the same as NascB (Fig. 2a). Except for mismatches in residues 125, 219, and 305, the C-terminal 273 residues of Nas_{S1868} and Nas_{F5053} are identical (Fig. 2a). This simple protein chimera strongly implies that the N-terminal portion of Nas_{F5053} contains residues that are involved in the

generation of NAS-C, while the C-terminal part of Nas_{F5053} harbors residues that are involved in the generation of ASP-A. Therefore, Nas_{F5053}, NascB, and Nas_{S1868} provide a suitable portfolio of P450 to reveal the relationships between enzyme sequences and the corresponding products.

Identifying the keys residues which determine the regio-configuration and stereo-configuration. Next, to identify the roles of critical residues in regulating and controlling the product profiles of the P450 enzymes, a series of protein variants was generated with mutations in the N-terminal and C-terminal portions of Nas_{F5053} and NascB, and the resultant mutant proteins were tested by enzyme assays. In the N-terminal part of Nas_{F5053}, we converted the following residues to their corresponding amino-acids in Nas_{S1868}: V44A-T49A-K52E (triple mutant), and P40H, Q65I, G84A, A86G, and I87V (point mutants). The in vitro enzyme assays of the Nas_{F5053} mutants (P40H could not be tested as it was insoluble) using cW_L-P_L showed that the simultaneous mutation of V44A, T49A, and K52E and single mutation of G84A or I87V imposed a slight change to the ratio of NAS-C and ASP-A (Supplementary Fig. 4, trace I-IV). Similarly, none of the eight point-mutations (P40H, V44A, T49A, K52E, S57P, F59L, G84A, and I87V) in the N-terminal portion of NascB were able to engineer NascB to produce ASP-A (Supplementary Fig. 4, trace V-XII). These eight positions were thus not investigated further. On the other hand, the single mutations Q65I and A86G in Nas_{F5053} dramatically reduced the production of NAS-C (Fig. 2b, trace IV, V). Furthermore, the double mutation A86G-Q65I almost abolished the production of NAS-C, leaving ASP-A as the only detectable product (Fig. 2b, trace VI). These results clearly indicate that Q65 and A86 are the two crucial residues in Nas_{F5053} that direct the enzyme to produce NAS-C.

In the following steps, the same mutations (Q65I and A86G) were introduced into NascB. The Q65I mutation in NascB also impacted on the production of NAS-C; it decreased the production of NAS-C and increased the production of ASP-A from none to reach the NAS-C to ASP-A ratio of 5:7 (Fig. 2b, trace VIII). The single mutation A86G in NascB had a negligible effect on the production of ASP-A, whereas it neutralized the effect of Q65I in the A86G-Q65I double mutant, which is opposite to the synergistic effect observed in Nas_{F5053} (Fig. 2b, trace IX, X). The contrasting effect of Q65 and A86 in NascB, as compared to Nas_{F5053}, prompts us to hypothesize that more residues in the C-terminal portion of NascB contribute to regulating the production of NAS-C and ASP-A. Because the C-terminal portion of NascB exhibits significant sequence differences, compared to Nas_{S1868}, the C-terminal part of NascB was divided into eight fragments (Supplementary Fig. 5). Each of the eight fragments was then replaced by the corresponding fragment in Nas_{S1868}, with Q65I and A86G mutations already in place. Each of eight NascB mutants was purified and enzyme assays revealed that the seventh fragment (fragment-7, carrying five mutations: H280Y, A284S, A287S, A288V, and L298I) almost abolished the production of NAS-C and generated ASP-A as the sole product (Fig. 2b, trace XI and Supplementary Fig. 6).

Point mutants were made to identify the effect of every single mutation in fragment-7 on the product profile. All five single point mutations produced more NAS-C than ASP-A (Supplementary Fig. 7, trace I-V), indicating more than one-substitution in fragment-7 is required to make ASP-A as the predominant product. We therefore restored, one by one, each of five-point mutations in the fragment-7 to its wild-type amino-acid of NascB, and observed that S284A or V288A counteracted the overall effect of five-point mutations in fragment-7 most

(Supplementary Fig. 7, trace VI-X). The results suggest that S284 and V288 are the two critical residues for the generation of ASP-A, while A284 and A288 are essential for the generation of NAS-C. Finally, the combination of Q65I-A86G-A284S-A288V in the NascB quadruple mutant was confirmed to make ASP-A as the major product (Fig. 2b, trace XII). In the case of Nas_{F5053}, the single mutations S284A or V288A significantly reduced the production of ASP-A, while the production of NAS-C was unaffected; and the double mutation S284A-V288A almost completely abolished the production of ASP-A (Fig. 2b, trace XIII-XV), hence unequivocally confirming the crucial role for these residues in determining the selective production of ASP-A and NAS-C. In addition to impact on the production of ASP-A

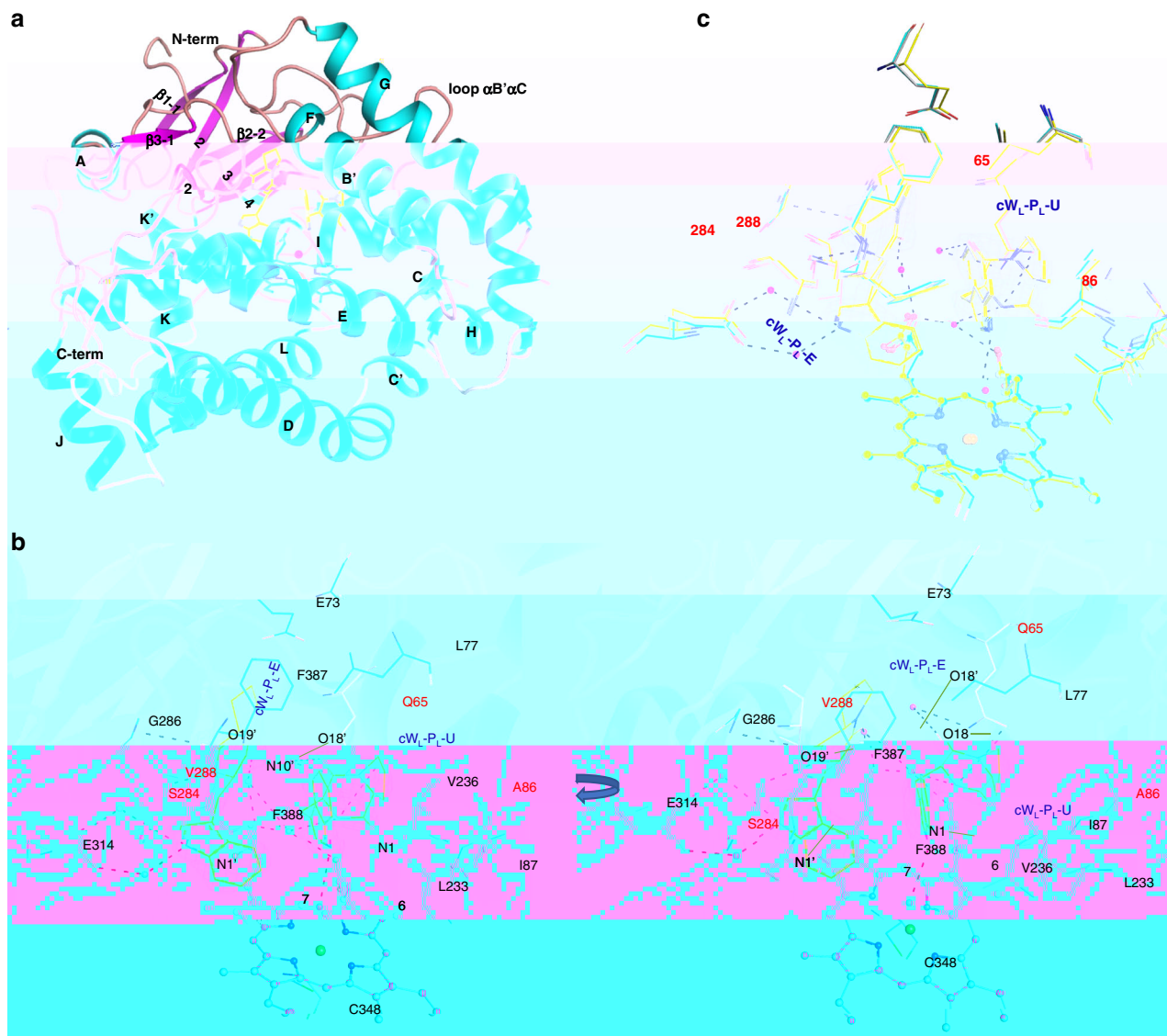


Fig. 3 Crystal structures of Nas_{F5053} and its mutants. **a** Cartoon representation of the structure of Nas_{F5053} bound to cW_L-P_L. Elements of secondary structure and the N/C-termini are labeled; α -helices are shown in cyan and β -strands in magenta. The iron in the heme is shown as a brown sphere and water molecules are displayed as magenta spheres. Other parts of the heme and cW_L-P_L are displayed as green and yellow sticks, respectively. **b** A representation of the active site of Nas_{F5053} in complex with cW_L-P_L-E and cW_L-P_L-U shown as yellow sticks. Oxygen and nitrogen atoms are shown in red and blue, respectively. The heme is displayed in cyan ball-and-stick representation, with the iron presented as a brown sphere. Nas_{F5053} residues are colored in cyan. Left: "side" view of the active site; right, "top" view. Probable H-bonds between Nas_{F5053}, cW_L-P_L, heme propionate and water molecules (magenta spheres) are indicated as dotted lines. Four critical residues (Q65, A86, S284, and V288) are highlighted in red and shown as orange sticks. **c** Superposition of the active sites of Nas_{F5053} (cyan sticks), Nas_{F5053}-Q65I-A86G (gray sticks), and Nas_{F5053}-S84A-V288A (yellow sticks) bound to cW_L-P_L-E and cW_L-P_L-U. The locations of four critical residues (65, 86, 284, and 288) are highlighted. The three complex structures are nearly identical, except for the mutated and adjacent residues. The cW_L-P_L-E and cW_L-P_L-U substrates in the Nas_{F5053} complex structure are surrounded by Fo-Fc electron density omit map, which is calculated after 20 cycles of refinement in the absence of the ligands and contoured at 2.0 σ level (blue mesh).

heme iron is coordinated by the axial ligand Cys348 in helix L. At the distal side of the heme, the iron is coordinated by a water molecule (Supplementary Fig. 11), consistent with the EPR data for NascB⁶ and CYP121¹⁵ that water is coordinated predominantly to low-spin Fe (III).

Comparison between the substrate-free and substrate-bound Nas_{F5053} structures reveals binding of substrates only invokes minimal conformational changes, with a root-mean-square deviation (RMSD) of 0.362 Å (for 388 Ca atoms) between the two forms (Supplementary Fig. 12). Instead, substrate binding is associated with rearrangements of some of the residues lining the substrate-binding cavity. Upon substrate binding, the side-chains

of both D85 and E73 rotate along the Ca-C β axis away from the binding site, to accommodate the substrates. Q65 undergoes a 2.0 Å shift (measured on the C δ atom) toward the binding site, to interact with one of the substrate molecules (cW_L-P_L-U; see below). Notably, Q65, D85, and E73 all reside in the long α B'- α C loop.

In the substrate-bound Nas_{F5053} structure, two cW_L-P_L molecules (cW_L-P_L-E and cW_L-P_L-U; E and U indicate extended and U-shaped, respectively) are present in the binding site, with full occupancy. cW_L-P_L-E adopts an extended conformation and forms multiple contacts with the heme group, loop β 3-1- β 3-2, loop α B'- α C, and loop α K- β 1-4. F387 and L77 form hydrophobic

interactions with the proline portion of extended cW_L-P_L-E . Formation of hydrogen bonds is observed between S284 and the backbone amide nitrogen of G286, with N10' and O19' of the substrate, respectively. O18' and N1' are indirectly in contact with 7-propionate of the heme and E314, respectively, mediated via hydrogen bonding with water molecules. The hydrophobic side of V288 also protrudes towards 7-propionate of the heme and indole ring in cW_L-P_L-E (Fig. 3b).

On the other side, cW_L-P_L-U is mainly in contact with the heme, αI , $\alpha B'$ and long loop $\alpha B'-\alpha C$, including a T-shaped stacking interaction network between the F388 side-chain and the indole rings of both cW_L-P_L-E and cW_L-P_L-U . The DKP ring of cW_L-P_L-U is further restrained by the side-chain of Q65 and extensively stabilized by secondary interactions with water, 6-propionate of the heme, N10, O19, O18, the side-chain amide of Q65 and the backbone NH of A86. Multiple hydrophobic interactions are also observed with residues lining the binding site, including V236, L233, I87, and Q65. These interactions therefore force cW_L-P_L-U into a U-shaped folded conformation, bringing the indole and prolyl entities into close proximity (Fig. 3b). Notably, the folded conformation of cW_L-P_L-U brings its C2 and N10 into close contact (3.2 Å distance), making the intramolecular cyclization between W_L and P_L in cW_L-P_L-U possible. Importantly, the indole ring of cW_L-P_L-U is positioned perpendicular to the heme group plane, with N1 forming a hydrogen bond with the heme-ligating water molecule (Fig. 3b), consistent with the initial step of N-deprotonation reaction by P450 compound I⁶. The indole rings from the two substrate molecules also form a T-shaped stacking interaction with each other. Hence, the complex structure between Nas_{F5053} and its substrate reveals a sophisticated orchestrated enzymatic environment where the heme, two identical substrates but different conformations, and the residues lining the substrate-binding cavity, are intimately interwoven.

However, our wild-type Nas_{F5053} crystal structures in complex with cW_L-P_L could not explain how Nas_{F5053} produced two different products: NAS-C and ASP-A. To explain product selectivity, we therefore determined two more cW_L-P_L substrate-bound crystal structures, of the mutants Nas_{F5053} -Q65I-A86G (PDB ID 6VZA) and Nas_{F5053} -S284A-V288A (PDB ID 6VZB). Comparisons among the three substrate-bound structures showed that all substrate-interacting residues, the heme, and the two substrates superimpose well (RMSD in this region between any two structures <0.26 Å; Fig. 3c), except for the mutated residues and the adjacent residues such as K289 and I87. These identical crystal structures indicates a common starting conformation for the reactions. To characterize Nas_{F5053} -catalyzed reactions further, we performed UV-Vis spectroscopic analysis and molecular dynamics (MD) simulations, to delineate the mechanism of regio-, stereo-selectivity and product profile regulation in Nas_{F5053} and its re-engineered variants.

Spectroscopic characterization of cW_L-P_L binding to Nas_{F5053} .

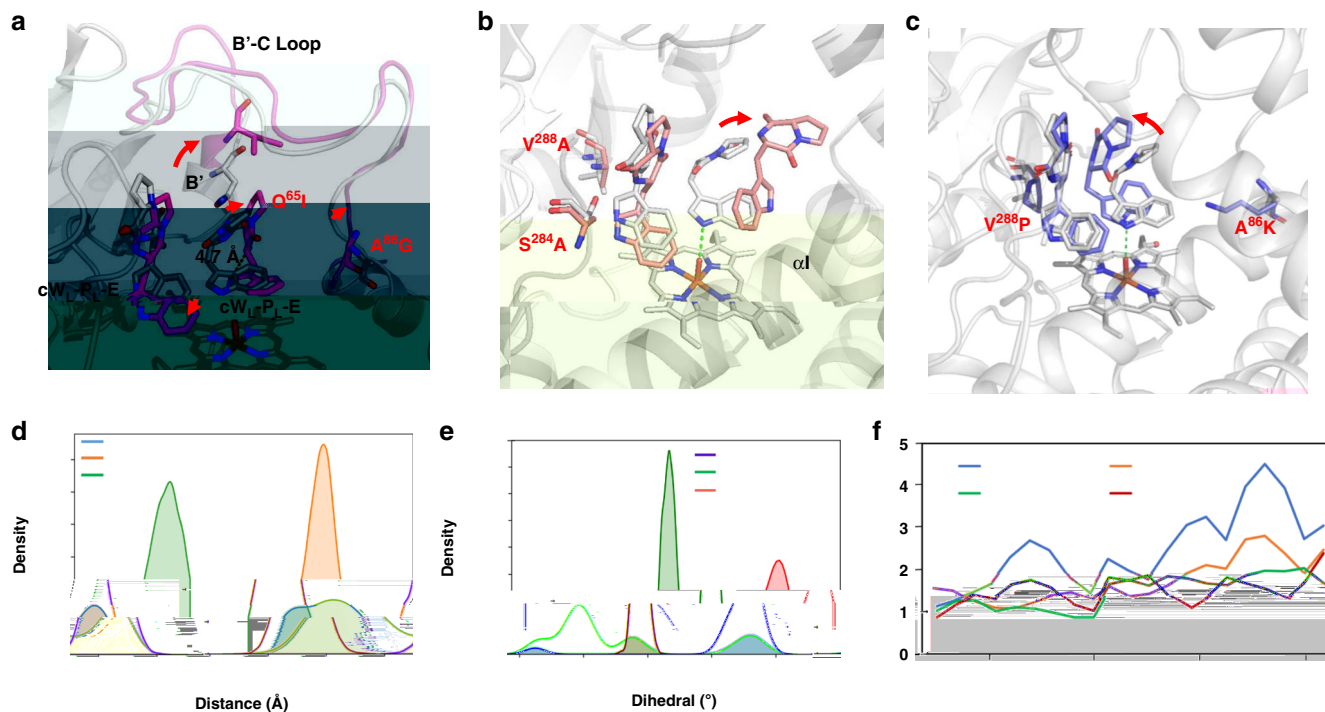
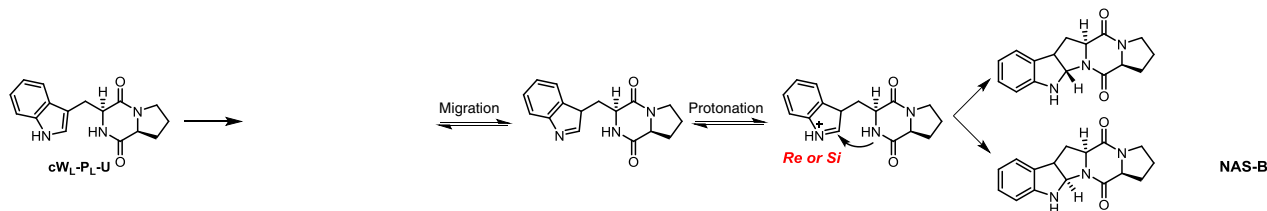
We measure UV-Vis absorption and difference spectra to probe the interaction in solution between cW_L-P_L and each of three enzyme variants, i.e., Nas_{F5053} , Nas_{F5053} -Q65I-A86G, and Nas_{F5053} -S284A-V288A. Binding of cW_L-P_L to Nas_{F5053} and its double mutants are all shifting a major Soret band from 418 nm to 387 nm, associating with the transition of the heme iron from the low spin (LS) to high spin (HS) state¹⁶. This transition, however, is not complete because a small but significant fraction of LS signal still remains even in the saturating cW_L-P_L concentration (Supplementary Fig. 13).

Then the difference spectra are used to calculate the spectral variations with OriginPro software. The plotting of the spectral

variation as a function of cW_L-P_L concentration is fitting to a rectangular hyperbola curve, yielding a binding constant of $11.6 \pm 2.1 \mu M$ for the interaction between cW_L-P_L and wild-type Nas_{F5053} , $25.6 \pm 1.0 \mu M$ for cW_L-P_L with Nas_{F5053} -Q65I-A86G and $4.81 \pm 0.26 \mu M$ for cW_L-P_L with Nas_{F5053} -S284A-V288A (Supplementary Fig. 13). Data fitting to a rectangular hyperbolic shape also models the case of CYP121 with single substrate¹⁵, suggesting that two cW_L-P_L substrates with Nas_{F5053} lack cooperativity for binding and catalysis. This assertion is further supported by a two-ligands complex structure where cW_L-P_L occupies one site and cW_L-P_L occupies the other site, which is reported in a published on-line research paper¹⁷ when we are revising our manuscript.

Molecular dynamics analysis. To characterize Nas_{F5053} -catalyzed reactions further, we performed molecular dynamics (MD) simulations with Amber (Supplementary Fig. 14), to delineate the mechanism of regio-selectivity, stereo-selectivity, and product profile regulation in Nas_{F5053} and its re-engineered variants. MD simulations were performed particularly to analyze the conformational changes associated with the proposed cW_L-P_L-U radical (Int1, Fig. 4) at the compound II stage⁶. The Q65-A86 and S284-V288 patches orchestrate the regio-specificities and stereo-specificities by distinct mechanisms. The Q65-A86 patch is involved in regulating the motion of the long loop $\alpha B'-\alpha C$, where Q65 and A86 reside at its two ends. The conformation of the $\alpha B'-\alpha C$ loop influences the conformation of the cW_L-P_L-U radical. Based on the MD of native Nas_{F5053} , the cW_L-P_L-U radical rotates anticlockwise along the axis of N1-Fe (IV)-OH, until two indole rings of the two substrates are almost in a plane. The Q65I and A86G mutations result in a shift of the $\alpha B'-\alpha C$ loop away from cW_L-P_L-U radical (Fig. 4a). The consequent relaxation of the restraints on Int1 unfolds Int1 (the distance between N10 and C2 is approximately 4.7 Å in most distance distributions; Fig. 4d). The results exclude the intramolecular cyclization of the cW_L-P_L-U radical to form a pyrroloindoline, without affecting the formation of ASP-A; this observation is consistent with our data that Nas_{F5053} -Q65I-A86G exclusively produce Asp-A.

The mutations of S284-V288 regulate regio-selectivity and stereo-selectivity by adjusting the relative positions of the two substrates and their conformations. Given that S284 and V288 contribute to lining the binding pocket for cW_L-P_L-E , mutations to less bulky Ala residues create space for cW_L-P_L-E to move towards the heme. This movement disturbs the interactions with the cW_L-P_L-U radical, and in turn pushes away the DKP and propyl rings of cW_L-P_L-U towards αI (orange sticks in Fig. 4b). This movement rigidifies the DKP ring of the cW_L-P_L-U radical as evidenced by decreased root-mean-square fluctuations (RMSFs) (Fig. 4f). According to MD, a positive C2-C3-C12-N10 dihedral angle between positions N10 above C2 (i.e., N10 attacks the Re face of the indole ring), generating an intermediate that leads to NAS-B, while a negative dihedral angles leaves N10 beneath C2 (i.e., N10 attacks the Si face of the indole ring), producing a different intermediate that leads to NAS-C. In the native form, there is a ~4–5 times higher probability for this dihedral angle to be negative (leading to NAS-C) than positive (leading to NAS-B), which is consistent with the experimental data that Nas_{F5053} produces more NAS-C than NAS-B. The dihedral angle can only be negative in the S284-V288 mutant, echoing that this double mutant can catalyze the formation of only NAS-C (Fig. 4e). MD also shows that the probability of the C2-N10 distance in the wild-type protein being >4 Å or <4 Å is almost equal, which means that native Nas_{F5053} could catalyze the formation of the products either requiring or escaping intramolecular cyclization. In the S284-V288 mutant, however, this



distance is fixed between 3.0 and 3.5 Å, making intramolecular cyclization inevitable (Fig. 4d).

The re-positioning and conformational changes of the substrates can also be achieved by the combined mutations in both the Q65-A86 and S284-V288 sites, such as the A86K-V288P double mutant. Opposite to its wild-type form, the A86K-V288P mutant produces NAS-B as the major product and NAS-C as the minor product. The long side-chain of K86 protrudes towards cW_L-P_L-U radical and drives its rotation and shift towards cW_L-P_L-E . On the other side, the V288P mutation compresses the active site, slightly pushing and rotating cW_L-P_L-E (cyan sticks in Fig. 4c). The dual changes of cW_L-P_L-U radical and cW_L-P_L-E reach a conformation where the DKP ring of cW_L-P_L-U radical becomes more rigid. In such a conformation, the C2-C3-C12-N10 dihedral angle is positive with high probability, favoring the attack of N10 to the Re face of the indole ring, to generate an intermediate leading to NAS-B. This is accompanied by a low probability event, where the dihedral angle is negative to allow for the formation of an intermediate leading to NAS-C. The sign

distribution of the dihedral angle is supported by the product profile of the A86K-V288P double mutant.

Discussion

Cytochrome P450 (CYP) enzymes are among the most exquisite and versatile biocatalysts in nature to synthesize and modify natural products^{18,19}. P450s and their engineered variants are continuously exploited as biocatalysts to functionalize natural products or potential drug leads²⁰. P450-catalyzed reactions can be broadly categorized into two groups: common and unusual²¹. Common P450 reactions generate minor structural alterations, such as C-H, N-H hydroxylation, and epoxidation on C=C double bonds. The mechanisms for those reactions are clear and represented by a canonical P450 catalytic cycle while the mechanisms for unusual P450 reactions are often unknown or elusive. Along with uncharacterized mechanism for uncharted chemistry, unusual P450 reactions may catalyze an enigmatic and/or dramatic structural transformation. Those features of unusual P450 reaction are of special research interests.

As an unusual P450-catalyzed reaction, the reaction of NascB was assumed to involve radical generation at N1 and migration, intramolecular Mannich reaction to form the pyrroloindoline C3 radical, and radical addition to the other molecule of DKP to form the HTDKP framework⁶. Although our previous DFT calculations and experiments preferred the N1-initiation over N10-initiation mechanism⁶, there was a lack of direct proof. Based on the crystal structures, we can now clearly see that the N1 of cW_L-P_L-U is indeed much closer than the N10 to the heme-ligating water molecule (Fig. 3b). In addition, the cW_L-P_L-U is in a U-shaped, folded conformation. Its indole and prolyl entities are close to each other, providing a viable distance for the intramolecular Mannich reaction to form the pyrroloindoline C3 radical. As Nas_{F5053} shows no structural evidence to accommodate the second copy of the pyrroloindoline C3 radical, the radical dimerization mechanism proposed in fungal TDKP biosynthesis can also be excluded^{7,8}. Furthermore, the three well superimposed complex structures (Fig. 3c) suggest a conserved starting conformation and reaction initiation steps in the formation of ASP-A, NAS-C, and NAS-B, although differentiating conformational dynamics of substrates develop in wild-type Nas_{F5053} and the three mutants, leading to the formation of different products. Therefore, all our structural evidence solidly supports the assumed reaction mode of HTDKPs⁶. Except for the type V HTDKP formation through a N1-radical addition, the intramolecular and intermolecular radical cascade mechanism⁶ thus can be rationalized to be a common paradigm for the biosynthesis of other bacterial HTDKPs.

Our dynamics simulation analyses indicate that the stereo-specificity and regio-specificity of P450 is indeed controlled by a sophisticated interaction of the substrates with the protein. This observation is consistent with the previous results for NascB, which can generate various HTDKP products with type I–IV frameworks upon feeding different substrates⁶. In contrast to the substrate-based approach, engineering the specificity conferring-residues is more appealing for biocatalysis to generate structural diversity of HTDKPs. Although the outcome of the reaction specificity cannot be readily predicted solely based on the crystal structures, the identified four specificity-conferring residues can serve as targets for protein engineering. Through screening a small library of mutations on the four residues, the product specificity of Nas_{F5053} was able to be shifted between different frameworks, which enables Nas_{F5053} to predominantly produce NAS-B, NAS-C, ASPA, or even NAS-E. This approach makes it very convenient for biocatalysis to efficiently produce the desired types of HTDKPs. In addition to the five identified types of HTDKPs, engineering the specificity-conferring residues also has the potential to generate diverse frameworks; screening more mutants for finding different specificities is currently in progress.

Besides the regio-specificity and stereo-specificity to generate frameworks, the limited tolerance of P450s for substrates is another factor that restricts their application. Previously, we found NascB has a very limited freedom in accepting substrates at the cW_L-P_L-E site⁶. From the structure, cW_L-P_L-E is surrounded by the bulky residues E73, F387, and L77; especially E73 is very close to the substrate. These residues, constituting the “ceiling” of the pocket, may form a constraint that hinders accepting bulkier substrates. In the P450 dimers for a few of HTDKP-like products, which contain heterodimerized nucleobase-DKP frameworks^{22–24}, E73 is replaced by the larger residue Tyr in GutD and P450_{NB5737}^{22,23}. As nucleobases are smaller than DKPs, this bulky residue may act as a gatekeeper, to restrict the second copy of DKP entering the pocket and to force the enzyme to catalyze a hetero-dimerization between the nucleobase and DKP. Therefore, engineering these residues may be able to control the space of the binding pocket and subsequently enable the enzymes to accept

either larger or smaller substrates in the prolyl position of cW_L-P_L-E , and currently such attempts are in progress. At the bottom of the binding pocket, the U-shaped molecule has more freedom as observed in our previous study⁶, by extending its proline moiety to the tunnel entrance lined by another two gatekeeper residues, V236 and L77. Engineering these two residues has the potential for further broadening of the substrate scope in the “bottom” cavity. By combining the engineering in reaction specificity and pocket space, the P450 reactions are believed to be able to generate more varied molecular diversity of HTDKPs.

The reaction specificities of P450s are determined by the sophisticated and orchestrated enzymatic environments and therefore it is difficult to identify the specificity-conferring residues solely from the crystal structures, especially for Ala86, which is ~6 Å away from cW_L-P_L-U (Fig. 3b). Through repeated construction and evaluation of sequence chimeras, we provided a strategy to decipher the sequence-product relationships of HTDKP-producing P450 enzymes. This approach proves to be effective in identifying pivotal residues governing product specificity between two or more homologous proteins. Based on these discoveries, we were able to alter the P450s' specificity through protein engineering. For enzymatic reactions with complicated catalytic mechanisms, relying solely on structural analysis can easily miss important information. Therefore, it is better to incorporate the investigation of the sequence-product relationships and our approach provides an option for this purpose.

In conclusion, through discovery, identification, and functional characterization, we have identified a suite of P450s (Nas_{bB} , Nas_{F5053} , and Nas_{S1868}) that share high sequence similarities but generate unique overlapping product profiles across all the five types of bacterial dimeric DKP frameworks. Our systematic mutagenesis studies on the promiscuous Nas_{F5053} and the versatile NascB identified four key residues, Q65, A86, S284, and V288, which play critical roles in controlling product regio-configurations and stereo-configurations. We demonstrate that the engineering of these residues is able to alter the product ratio and even generate an interesting framework, which has not previously been observed for the substrate cW_L-P_L . To obtain insights into the structural basis for regio-specificity, stereo-specificity, and chemical versatility, we further determined high-resolution crystal structures of wild-type Nas_{F5053} in its substrate-free and substrate-bound form, and of two Nas_{F5053} mutants (Q65I-A86G and S284A-V288A) in their substrate-bound forms. The binding mode of cW_L-P_L revealed by the complex structures supports the previous proposed intramolecular and intermolecular radical cascade addition mechanism. Molecular dynamics simulations were employed to uncover the specificity-conferring mechanism of these residues, based on the crystal structures. Therefore, our biochemical, structural, and computational characterizations across this representative group of HTDKP-forming P450s provide a clear mechanism of how these sophisticated catalytic mechanisms take place, which expands our knowledge on the chemical diversity of cytochrome P450-catalyzed natural products and enables the rational engineering of this group of P450s and other homologs to obtain different HTDKP frameworks.

While this manuscript was undergoing revision after review, Shende and Co-workers published the structural and functional characterization of NzeB¹⁷, the synonym of Nas_{F5053} . Their structural data are consistent with our data. The active site residues that they identified are also covered by four key residues revealed in our manuscript.

Methods

Protein expression, purification, and enzyme assay. P450 genes with codon optimized for *E. coli* were cloned into pET28a (nas_{F5053}) and pMS1 (nas_{bB} and

nasS1868), which were overexpressed in *E. coli* BL21 (DE3) and *M. smegmatis* mc² 155, respectively (Supplementary Fig. 15). The in vitro biochemical reactions using all the P450s mentioned in this study were performed in a 100 μ L reaction system containing 0.1 μ M P450, 1 mM cW_L-P_L, 1 μ M spinach ferredoxin (Fd), 1 μ M ferredoxin reductase (FdR), 2 mM NADP⁺, 2 mM glucose, and 2 mM glucose dehydrogenase (GDH) in 50 mM HEPES buffer, 100 mM NaCl, at pH 7.5. After incubating at 4 °C for 24 h, the reactions were quenched and extracted with ethyl acetate (2 \times 200 μ L). Then the combined organics were concentrated in vacuo, which were re-dissolved in HPLC-graded methanol and the resulting solutions were filtered through 0.45 μ m membrane and finally analyzed by UHPLC-MS. A Diamonsil (C18, 2 μ m, 2.1 \times 50 mm, Shim-pack GIST) was used with a flow rate at 0.3 mL min⁻¹ and a PDA detector over a 23 min gradient program with water (eluent A) and methanol (eluent B): *T* = 0 min, 40% B; *T* = 10 min, 40% B; *T* = 15 min, 70% B; *T* = 18 min, 40% B; *T* = 23 min, 40% B.

Protein crystallization and crystal structure determination. Initial crystals were obtained in 0.2 M CaCl₂, 20% (w/v) polyethylene glycol (PEG) 3350, pH 7.5 at 20 °C using the hanging drop vapor diffusion technique with the addition of 5% glycerol to the protein stock. The initial crystals were subsequently crushed for seeding by using the Seed Bead Kit (Hampton Research). Final crystals were obtained using the micro-seeding technique in 0.2 M CaCl₂, and 22% (w/v) PEG 3350, pH 7.5 at 4 °C. Substrate-bound protein crystals were obtained by soaking the substrate-free crystals in the mother liquor containing 2.5 mM cW_L-P_L (diluting from 50 mM stock solution in DMSO) for 24 h or co-crystallization after mixing 0.13 mM protein with 2.5 mM cW_L-P_L. Both methods produce identical complex structures. The complex structure from soaking was chosen for structural analysis and presentation, due to a better overall quality, including resolution, of the diffraction data collected from the soaked crystals.

Crystals were mounted onto CryoLoops (Hampton Research) and soaked in a cryoprotection solution containing 0.2 M CaCl₂, 22% (w/v) PEG 3350, pH 7.5, and 20% (v/v) glycerol prior to flash freezing in liquid. For the substrate-bound protein crystals, the cryoprotection solution also contained 2.5 mM cW_L-P_L. The X-ray diffraction data were collected at the Australian Synchrotron MX beamlines. The collected data were indexed and integrated using XDS²⁵ and scaled and merged using Aimless²⁶. A partial initial model of the holo-structure was obtained by the molecular replacement technique with Phaser in Phenix²⁷ using the crystal structure of CYP121 from *Mycobacterium tuberculosis* (PDB accession code: 5WP2) as the search model. The initial model was improved by using the Morph Model tool in Phenix²⁸ and manually modified in COOT²⁹. The substrate-bound structure was solved by the molecular replacement technique using the holo-structure as the search model. The structures were refined using Phenix.Refine³⁰ and manually modified in COOT iteratively. The graphic presentations of protein structures were prepared with Pymol.

NMR spectroscopy. The NMR spectra were recorded on a Bruker Avance III spectrometer at a ¹H frequency of 400 MHz. Lyophilized samples (varying from 1 to 7 mg) were dissolved in 280 μ L DMSO-*d*₆ (Cambridge Isotope) and all spectra were recorded at 25 °C (298 K). ¹H and ¹³C resonances were assigned through the analysis of 1D ¹H, 1D ¹³C, 2D ¹H-¹H ROESY, 2D ¹H-¹³C HSQC, and 2D ¹H-¹³C HMBC (optimized for long-range heteronuclear couplings of 6 Hz). ¹H and ¹³C chemical shifts were calibrated with reference to the DMSO solvent signal (2.50 and 39.5 ppm for ¹H and ¹³C, respectively). NMR experiments were processed with Bruker Topspin program (version 3.57) and analyzed with MestReNova software.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The sequences of the *nasB*, *nasS1868*, and *nasF5053* reported in this work are available under existent accession numbers MW196742 (this hyperlink is currently on hold and will be released upon publication), WP_030881046.1 and WP_030888003.1 in Genebank, respectively. Their synthetic DNA sequences are listed in the supplementary information. All other data are also available upon request from the corresponding authors. The pdb coordination files for substrate-free F5053 and F5053 in complex with cW_L-P_L, NaF₅₀₅₃-Q65I-A86G in complex with cW_L-P_L and NaF₅₀₅₃-S284A-V288A in complex with cW_L-P_L were deposited in Protein Data Bank with the accession number of 6W0S, 6VXV, 6VZA, and 6VZB, respectively. Source data are provided with this paper.

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Author contributions

X.Q., X.J., B.K., and Z.D. conceived this project; C.S., W.Z., W.T., H.P., and Zhi Lin performed the biochemical experiment; Z.L. and X.J. solved the crystal structures; X.Q., X.J., C.S., and Z.L. analyzed data; and X.Q., X.J., C.S., and Z.L. wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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