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# Molecular basis for the biosynthetic divergence of anthracyclines kosinostatin and chartreusin

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Kosinostatin and chartreusin are both anthracycline-type antitumor agents. The structure of kosinostatin incorporates a spirally fused pyrrolopyrrole moiety attached to a five-membered ring of its anthraquinone core, whereas chartreusin possesses a distinct, rearranged pentacyclic aromatic dilactone aglycone. Biosynthetic divergence in kosinostatin and chartreusin is governed by highly homologous proteins, KstA15 and ChaU, which catalyze phenol hydroxylation and hydration, respectively. In this report, we successfully engineered a functional swap between ChaU and KstA15 *via* directed evolution. The crystal structure of the ChaU mutant along with molecular dynamics simulations elucidated the structural determinants of its catalytic specificity, revealing the critical residue interactions and conformational dynamics. Furthermore, a ubiquitous distribution of KstA15-like proteins across microbial genomes has been demonstrated through gene mining. These findings deliver key insights into the evolutionary trajectory of functionally divergent enzymes and create a robust framework for rationally engineering biosynthetic pathways to produce novel anthracycline derivatives with potential pharmaceutical applications.

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## Introduction

Anthracyclines are microbially derived natural products defined by a broad antitumor spectrum, potent efficacy, and proven clinical effectiveness, which underpin their extensive use in cancer treatment regimens.<sup>1–3</sup> Representative anthracycline drugs include doxorubicin, daunorubicin, and aclarubicin.<sup>4–8</sup> In terms of biosynthetic origin, anthracyclines are aromatic type II polyketides. Their carbon skeleton is assembled through nine iterative rounds of Claisen condensation between malonyl-CoA and either an acetyl-CoA or propionyl-CoA starter unit. The resulting polyketide chain is subsequently processed by conserved enzymes including ketoreductase, cyclase, and aromatase to form the characteristic tetracyclic anthracyclinone core.<sup>9–11</sup> The structural diversity observed in anthracyclines primarily arises from a series of tailoring redox reactions that

introduce differential modifications onto this common scaffold.<sup>12–18</sup>

Among anthracycline derivatives, two highly oxidized molecules, kosinostatin and chartreusin, exhibit significant structural differences.<sup>19–22</sup> Nevertheless, both originate from the common precursor auromycinone (3), and their biosynthetic pathways bifurcate at a critical step regulated by two homologous ester cyclases, KstA15 and ChaU (Fig. 1).<sup>23,24</sup> KstA15 and its partner protein KstA16 are encoded within the *kst* gene cluster responsible for kosinostatin biosynthesis, whereas ChaU and its corresponding protein ChaX are found in the *cha* gene cluster dedicated to chartreusin construction (Fig. 1).<sup>24–26</sup> The KstA15-KstA16 complex utilizes oxygen as a substrate to catalyze C8 phenyl ring hydroxylation, whereas the ChaU-ChaX pair promotes D-ring dehydration *via* a hydration step (Fig. 1). In these two catalytic systems, KstA16 and ChaX are both members of the short-chain reductase (SDR) family, sharing high sequence homology and performing identical enzymatic functions in the reduction of the *para*-quinone moiety in anthracyclines.<sup>24</sup> While both KstA15 and ChaU belong to the nuclear transport factor 2 (NTF2) superfamily, they exhibit a high degree of sequence homology but markedly divergent biological functions (Fig. 1).<sup>24</sup> The crystal structures of ChaU and KstA15 (along with its functional homolog SnoaL2) have been determined.<sup>23,24,27</sup> Site-directed mutagenesis studies demonstrated that the catalytically essential residues in ChaU include D119, R50, E38, and S37. In contrast, the active sites in SnoaL2 are identified as H35, E106, H119, and D121, which correspond to

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Fig. 1 ChaU and KstA15: two annotated ester cyclases determining the biosynthetic branching points to chartreusin and kosinostatin.

H49, E119, H131, and D133 in KstA15 (Fig. 1).<sup>23,24</sup> Despite their high structural similarity, the molecular basis underlying the distinct catalytic functions of ChaU and KstA15 remains elusive. Here, we identified the critical amino acid combinations that govern the catalytic functional specificity of ChaU and KstA15, thereby facilitating a switch between C8 phenol hydroxylation and their hydration activities. Through integration of the crystal structure of a hydroxylation-competent ChaU mutant with molecular dynamics (MD) simulations, we elucidated the molecular basis that drives the functional divergence toward hydration *versus* phenol hydroxylation in this enzyme family. Furthermore, our work reveals that homologous proteins capable of catalyzing phenol hydroxylation are widely distributed among diverse microorganisms, whereas enzymes within this class that retain hydratase activity are scarcely represented, showing the occurrence of proteins annotated as “ester cyclases” in the NTF2 superfamily.

## Results and discussion

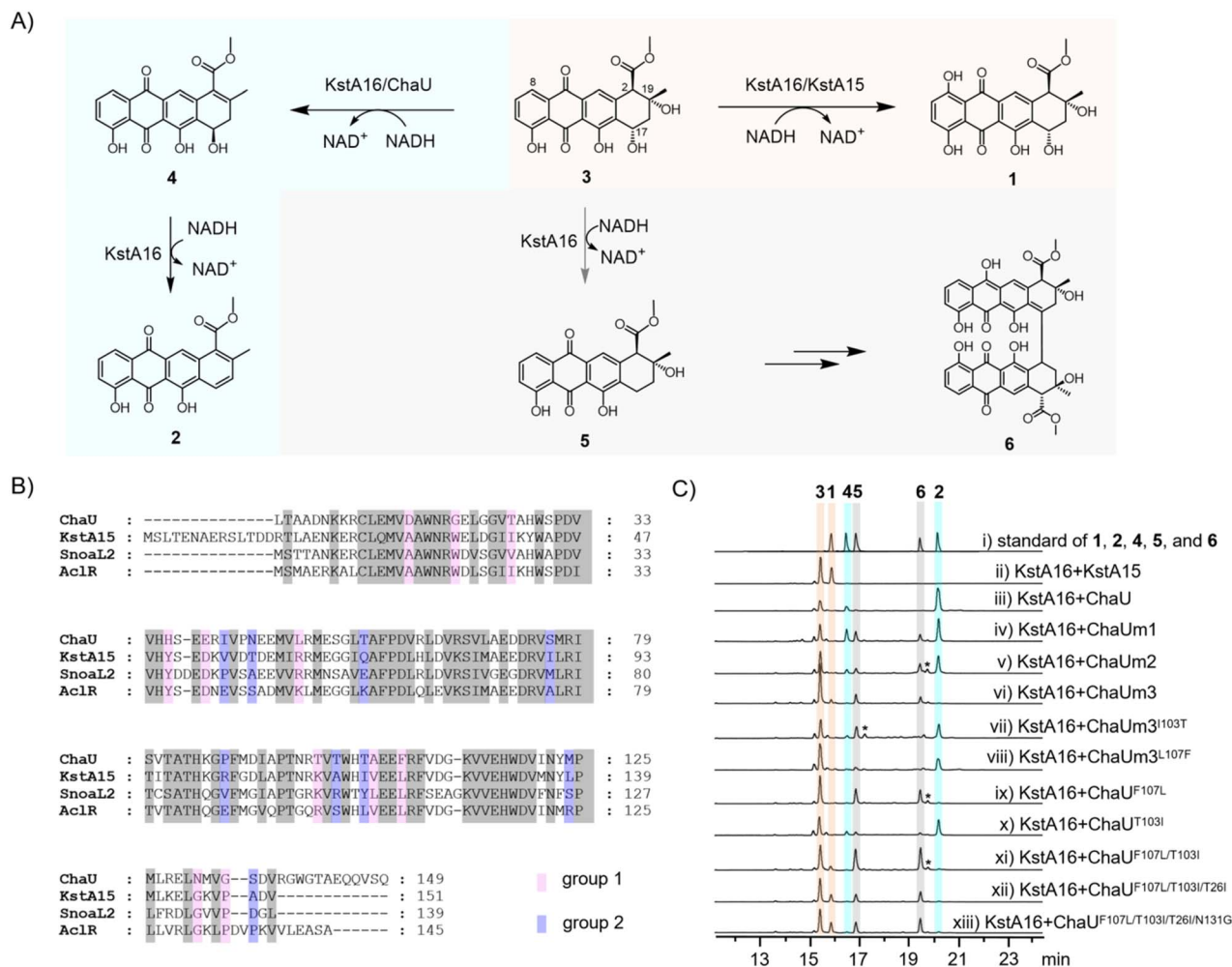
### Elucidating the core set of amino acids that bridge the catalytic properties of ChaU and KstA15

KstA15 and KstA16 jointly catalyze the hydroxylation of **3** at the C-8 position to form **1**, while ChaX and ChaU together mediate the conversion of **3** to yield **2**, proceeding through intermediate **4** *via* a (de)hydration reaction (Fig. 2A). Individually, either KstA16 or ChaX catalyzes the 17-dehydroxylation of **3**, producing **5** and a dimer (**6**), whose structure has been proposed based on molecular weight (Fig. 2A). Given the higher expression level of KstA16 relative to its isoenzyme ChaX, KstA16 was selected as the interacting partner for all subsequent catalytic reactions involving ChaU or KstA15-like proteins (Fig. S1). To pinpoint the specific amino acid residues that

account for the divergent enzymatic functions of ChaU and KstA15, we chose to subject ChaU to directed evolution for phenol hydroxylation. Thus, we performed sequence alignment of ChaU with its homologous proteins (KstA15, Snoal2, and AclR) known to catalyze such reactions (Fig. 2A and B).<sup>23</sup> This analysis identified non-conserved residues, which we systematically categorized into two groups: “Relatively non-conserved” (a common amino acid in the homologs that differs in ChaU, group 1), and “Completely non-conserved” (divergent amino acids across all four proteins, group 2) (Fig. 2B). The initial substitution of group 1 amino acid residues in ChaU with their counterparts from KstA15 produced the mutant ChaUm1, which contains 11 mutations (Fig. 2B and S2). However, ChaUm1 failed to gain phenol hydroxylation activity and instead retained its original hydration activity to convert **3** to **4**, **2**, **5** and **6**, the structures of which were determined by comparing their <sup>1</sup>H NMR and HR-MS data with those previously reported (Fig. 2C(i–iv), S16–S24 and Tables S6–S9),<sup>14,24</sup> indicating that key residues essential for phenol hydroxylation are not fully conserved among KstA15, Snoal2, and AclR. Building upon this mutant, we performed a sequential and pairwise replacement of the “group 2” amino acids in ChaU with their KstA15 equivalents, starting from the C-terminus of the protein (Fig. 2B and S2). Surprisingly, while ChaUm2 (ChaUm1<sup>S135A/M124L</sup>) remained incapable of hydroxylating phenol (Fig. 2C(v)), the ChaUm3 (ChaUm2<sup>T103I/T100A</sup>) mutant successfully acquired the catalytic function of KstA15, allowing it to cooperate with KstA16 to convert **3** into **1** (Fig. 2C(vi), S14–S15 and Table S5).

Given that the ChaUm3 mutant carries 15 amino acid substitutions, it is probable that not all residues contribute equally to its functional repertoire for C8 phenol hydroxylation. Therefore, we systematically generated a series of revertant





**Fig. 2** Rational design of ChaU to endow phenol hydroxylation activity. (A) The biosynthetic pathway from auramycinone (**3**) to resomycin C (**2**) and 8-hydroxyauramycinone (**1**). (B) Amino acid sequence alignment of ChaU with its homologous proteins KstA15, SnoaL2, and AcLR. (C) HPLC analysis of the enzymatic reactions *in vitro* catalyzed by KstA16 and its partners. ChaUm1 denotes the undecimutant ChaU<sup>D15A/G20W/T26I/H36V/E39D/L49R/T98K/A104V/F107L/N131G/G134P</sup>, ChaUm2 represents ChaUm1<sup>S135A/M124L</sup> and ChaUm3 stands for ChaUm2<sup>T103I/T100A</sup>. All assays were performed using **3** as the substrate and NADH as the cofactor, with a reaction duration of 30 minutes. UV = 440 nm. Peaks marked with an asterisk (\*) are unidentified.

mutants, in which each of the fifteen substituted residues was individually changed back to its wild-type counterpart in ChaU (Fig. S2). Specifically, both the ChaUm3<sup>I103T</sup> and ChaUm3<sup>L107F</sup> revertants lost phenol hydroxylation function (Fig. 2C(vii–viii)), thereby establishing that these two specific residues are essential for determining the catalytic divergence between hydration and C8 phenol hydroxylation among ChaU-like enzymes. Subsequently, we introduced single (ChaU<sup>F107L</sup> and ChaU<sup>T103I</sup>) and double (ChaU<sup>F107L/T103I</sup>) mutations into the wild-type ChaU. The data demonstrated that only the ChaU<sup>F107L/T103I</sup> double mutant could catalyze C8 phenol hydroxylation (Fig. 2C(ix–xi)), further underscoring that these two residues constitute the minimal determinant for shifting the catalytic activity of ChaU from hydration to phenol hydroxylation.

Given the limited catalytic activity of the dual mutant ChaU<sup>F107L/T103I</sup> toward phenol hydroxylation at this stage, we proceeded to generate a series of triple and quadruple mutants

through its combinatorial substitution with each amino acid residue from group 1 (Fig. S2). The results demonstrated a substantial enhancement in the efficiency of phenol hydroxylation for the triple mutant ChaU<sup>F107L/T103I/T26I</sup> and quadruple mutant ChaU<sup>F107L/T103I/T26I/N131G</sup> (Fig. 2C(xii–xiii)). To confirm that these four residues are indeed critical for phenol hydroxylation, we performed site-directed mutagenesis at the equivalent positions in KstA15. Mutation of the four amino acids in KstA15 to their corresponding residues in ChaU revealed that KstA15<sup>I117T</sup> and KstA15<sup>L121F</sup> completely lost phenol hydroxylation activity, while that of KstA15<sup>L40T</sup> and KstA15<sup>G145N</sup> was substantially diminished (Fig. S3). Notably, all four single mutants acquired the ability to catalyze the hydration reaction to yield **2** (Fig. S3). This suggested that phenol hydroxylation and hydration reactions are intricately related and the former necessitates the contribution of more key amino acid residues to retain its catalytic integrity, compared to the simpler hydration catalysis. If so, the amino acid residues critical for

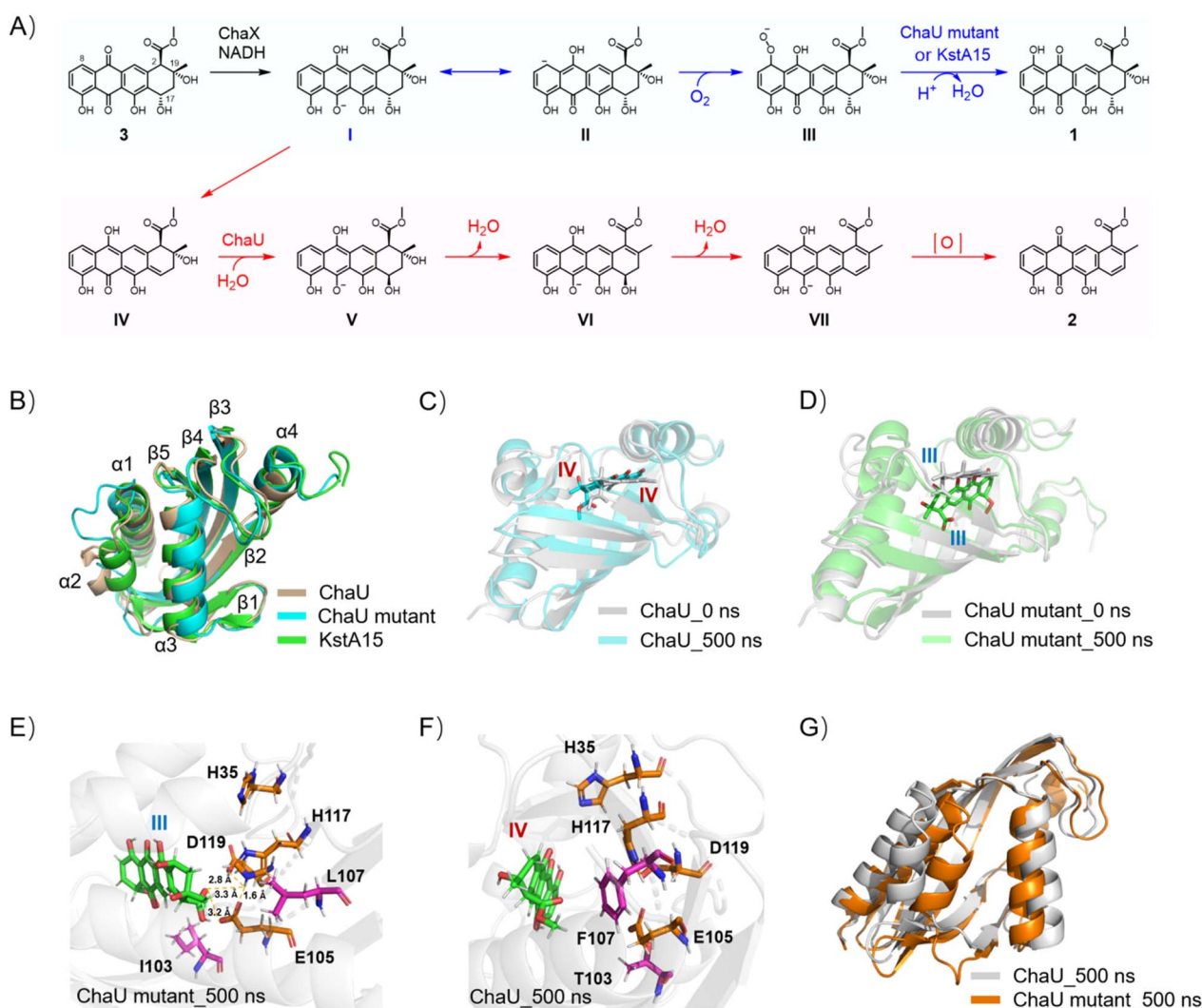


hydration may also impact C8 phenol hydroxylation. This hypothesis was tested by substituting the residues in KstA15 that align with ChaU's key hydration sites with alanine.<sup>24</sup> As a result, the mutants KstA15<sup>S51A</sup>, KstA15<sup>E52A</sup>, KstA15<sup>R64A</sup> and KstA15<sup>D133A</sup> indeed lost 50–60% catalytic activity towards C8 phenol hydroxylation and retained no hydratase function (Fig. S4). These data underscored the precision of evolutionarily refined enzymatic strategies that direct specific transformations in anthracycline antibiotic biosynthesis.

### Insights into the catalytic divergence between ChaU and the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant

Given that the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant shares the same catalytic function for phenol hydroxylation as KstA15, we first obtained the crystal of this quadruple mutant as a tetramer, and determined its structure at a resolution of 2.70 Å (Fig. S5 and

Table S4). A comparison of the crystal structures of ChaU, ChaU<sup>F107L/T103I/T26I/N131G</sup> and KstA15 indicated a conserved overall tertiary fold, while marked conformational deviation is localized to the  $\alpha$ 2-helix region (Fig. 3B). To delineate the mechanistic basis for the catalytic divergence between ChaU and ChaU<sup>F107L/T103I/T26I/N131G</sup> mutants, we performed molecular docking followed by molecular dynamics (MD) simulations. As shown in Fig. 3, a pronounced contrast in substrate-binding dynamics was observed between wild-type ChaU and ChaU<sup>F107L/T103I/T26I/N131G</sup> mutants *via* MD simulations. While the substrate **IV** (the putative direct substrate of ChaU<sup>24</sup>) retained a pose similar to its initial configuration with stability in WT ChaU (Fig. 3A and C), the mutant system exhibited a marked shift: the substrate **III** (the presumed direct substrate of KstA15 (ref. 14 and 23)) reoriented substantially around 80 ns, adopting a distinct binding mode that remained stable



**Fig. 3** Mechanistic investigations of catalytic divergence between ChaU and ChaU<sup>F107L/T103I/T26I/N131G</sup> mutants. (A) The intermediates involved in the 8-hydroxylation and 17-hydration reactions. (B) The alignment of the crystal structures of ChaU (PDB: 8KAA), ChaU mutant (PDB: 22JH) and KstA15 (PDB: 8R20). (C–D) Binding conformations of the substrate and protein in WT ChaU (C) and ChaU mutants (D), captured at the initial (0 ns) and final (500 ns) time points of the MD simulation. (E and F) Hydrogen-bonding interactions of residues H35, E105, H117, and D119 with the substrate in ChaU mutant (E) and WT ChaU (F) at 500 ns of the MD simulation. (G) Protein channel conformational analysis of WT ChaU and ChaU mutants at 500 ns of the MD simulation. ChaU mutant here specifically refers to the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant.



thereafter (Fig. 3A and D and S6–S7). To elucidate the structural determinants of this divergence, we analyzed hydrogen-bond interactions between the substrate and four ChaU residues (H35, E105, H117 and D119), which were selected based on their spatial positioning and functional analogy to catalytic residues in KstA15.<sup>23</sup> Notably, substrate **III** in the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant sustained at least one hydrogen bond with these residues throughout most of the simulation (Fig. 3E and S8). Conversely, equivalent hydrogen-bond interactions were essentially absent in the WT ChaU system (Fig. 3F and S8). These results suggested that the mutation alters the substrate-binding microenvironment of ChaU, leading to the stabilization of a substrate conformation that is proximal to key reactive residues. This reoriented pose thereby satisfies the spatial necessary for a KstA15-type reaction, providing a plausible mechanistic basis for the observed functional shift in the ChaU mutant, which was further supported by the marked decrease in phenol hydroxylation efficiency exhibited by the variants ChaU<sup>F107L/T103I/T26I/N131G/E105A</sup> and ChaU<sup>F107L/T103I/T26I/N131G/H117A</sup> (Fig. S4). Consistent with this model, conformational analysis indicated that the mutant enzyme preferentially adopts a more closed conformation upon substrate binding compared to the WT ChaU (Fig. 3G and S9), presumably diminishing the spatial distance between the

substrate and catalytic residues, which in turn consolidates the hydrogen-bonding network and thereby facilitating the establishment of a preorganized, catalytically competent architecture. We noted that in wild-type ChaU, the conformation of E105 is stabilized by a hydrogen bond with T103 and significant steric hindrance from F107 (Fig. S10). In the ChaU mutant, however, the substitution to I103 abolishes this hydrogen bond, and the replacement with L107 eliminates the steric restriction (Fig. S10). This releases E105 from its constrained state, allowing it to move closer to the incoming substrate and thereby tightening the binding cavity. The resultant contraction of the  $\alpha$ -helical structure further enables W100 to form a hydrogen bond with the substrate (Fig. S10). Such changes synergistically promote hydroxylation at the C8 position of the substrate, driving the reaction forward.

To enable a more direct comparison of the phenol hydroxylation mechanisms between the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant and KstA15, molecular docking and MD simulations of substrate **III** bound to KstA15 were also performed. Notably, a difference in substrate-binding conformation is observed between the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant and KstA15 (Fig. 3E and S11), despite their identical catalytic functions.<sup>23</sup> Within KstA15, H131 (the counterpart of H117 in the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant) is positioned sufficiently

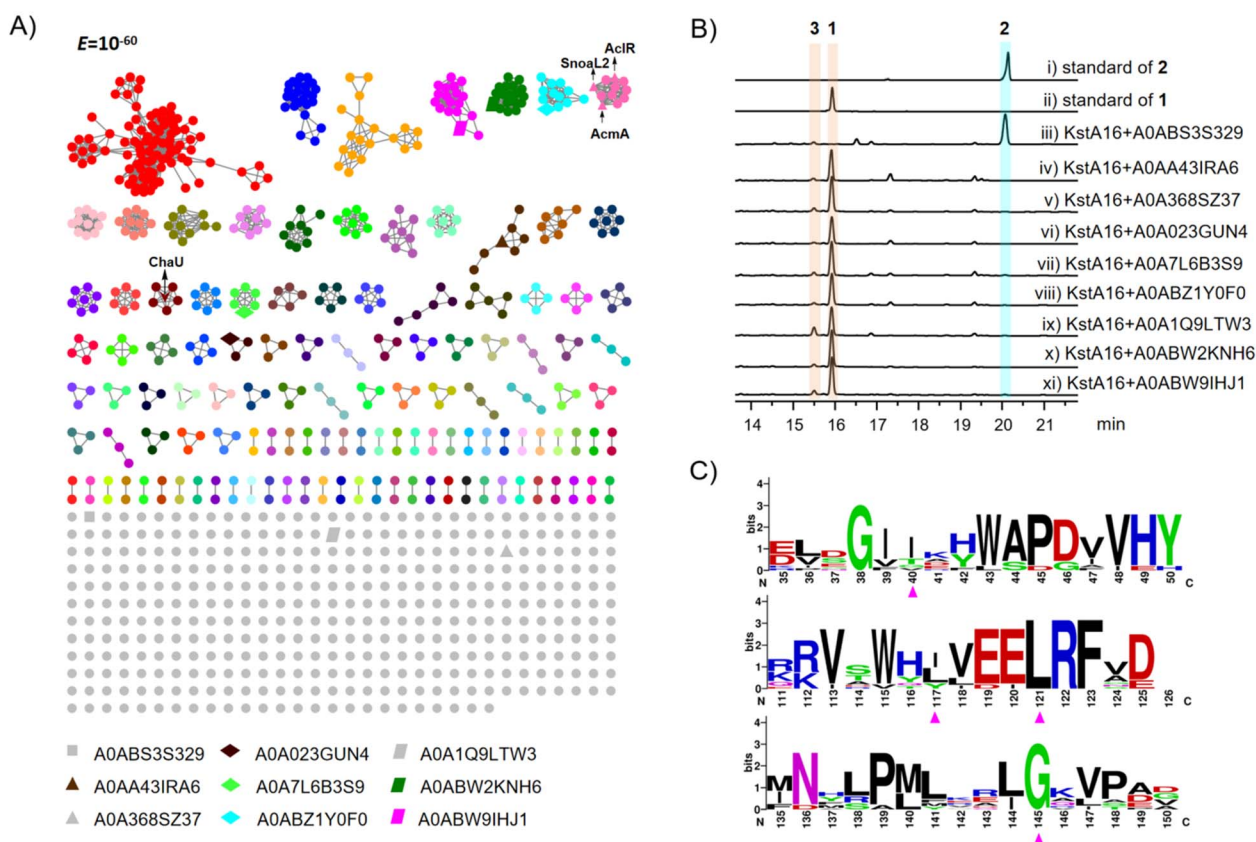


Fig. 4 Distribution and characterization of ChaU-like and KstA15-like proteins. (A) The SSN analysis of NTF2 superfamily ester cyclases. A0ABS3S329, A0A023GUN4, A0A1Q9LTW3, A0AA43IRA6, A0A7L6B3S9, A0ABW2KNH6, A0A368SZ37, A0ABZ1Y0F0, and A0ABW9IHJ1 are derived from *Actinomadura violacea*, *Micromonospora okii*, *Actinokineospira bangkokensis*, *Thermoleophilum bacterium*, *Micromonospora robiginosa*, *Marinactinospora rubrisoli*, *Marinitenerispora sediminis*, *Streptomyces althioticus* and *Streptomyces galilaeus*, respectively. (B) Comparative HPLC profiles of reactions catalyzed by ChaU and KstA15 homologs. All assays were conducted with **3** as the substrate and NADH as the cofactor for 2 hours. UV = 440 nm. (C) Sequence logo of conserved motifs distinguishing KstA15-like from ChaU-like proteins.



close to the peroxide anion, especially during the first 80 ns, to allow direct transfer of HE2 or HD1 in H131 to the peroxide anion moiety (Fig. S11 and S12). However in the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant, H117 and E105 are located adjacent to the carboxyl group of III (Fig. 3E and S11), likely contributing to substrate stabilization. This hinted the presence of another proton transfer channel in the ChaU<sup>F107L/T103I/T26I/N131G</sup> mutant catalytic systems, potentially accounting for the approximate three-fold higher KstA15 catalytic activity relative to ChaU<sup>F107L/T103I/T26I/N131G</sup> (Fig. 2C(ii, xiii) and S4).

Collectively, these findings indicated that the mutation orchestrates a coordinated reorganization across multiple levels—including substrate binding, residue interaction networks, and global conformational dynamics—rather than simply enhancing local substrate interactions. These integrated, multiscale changes collectively promote the emergence of structural and dynamic properties in the ChaU mutant that resemble those essential for KstA15-like catalysis, thereby providing a mechanistic basis for its altered reactivity.

### Broader natural occurrence of KstA15-like versus ChaU-like proteins

Despite the high sequence homology between KstA15 and ChaU, KstA15-like proteins, including SnoaL2, AclR, and SwaR1, have been identified in various anthracycline biosynthetic gene clusters,<sup>23,28,29</sup> while, ChaU represents the first NFT2 superfamily member reported to possess a hydratase function.<sup>24</sup> To explore the distribution of ChaU-like proteins across the microbial genome, we attempted to mine databases for them. Thus, using ChaU as a query, we searched against microbial proteomes in the UniProt database and retrieved 946 homologous sequences. A sequence similarity network (SSN) was constructed with these sequences (Fig. 4A).<sup>30,31</sup> Application of an E-value cutoff of  $10^{-60}$  resolved ChaU-like and KstA15-like enzymes into two distinct clusters within the network (Fig. 4A).

In an attempt to identify ChaU-like enzymes within other clades, we randomly selected 9 proteins from distinct branches of diverse microorganisms for heterologous expression in *Escherichia coli* (Fig. 4A). Co-catalytic assays with KstA16 demonstrated that only one protein (UniProt ID: A0ABS3S329) exhibited hydratase activity, whereas all others catalyzed specifically the C8-hydroxylation reaction (Fig. 4B). Aligning the 9 selected proteins against ChaU, KstA15, AclR, and SnoaL2 revealed that the ChaU residues T103 and F107 are functionally conserved in homolog A0ABS3S329 capable of catalyzing the hydration reaction (Fig. S13). A similar conservation pattern was observed for the corresponding residues (I/L117 and L121) in KstA15 and its functionally homologous proteins (Fig. 4C and S13). Residues T26 and N131 in ChaU are not conserved across its isozyme A0ABS3S329 (Fig. S13). Intriguingly, A0ABS3S329 shares the identical amino acid at position 26 with the corresponding residue in KstA15 (Fig. S13). Although position 131 is also variable between ChaU and A0ABS3S329, the corresponding site in KstA15 (position 145) and its isozymes is consistently a conserved glycine (Fig. 4C and S13). These findings reinforce

the conclusion that residues 103 and 107, along with residue 131 in ChaU, play a pivotal role in dictating its functional specificity as either a hydratase or a hydroxylase.

## Conclusions

In conclusion, we pinpointed two key amino acid residues responsible for the distinct catalytic activities of ChaU and KstA15 through a combined experimental approach involving sequence alignment and site-directed mutagenesis. This critical finding allowed us to rationally convert ChaU from a hydratase into a phenyl ring hydroxylase. Furthermore, protein crystallography, along with molecular dynamics simulations, ultimately elucidated the precise molecular basis for their catalytic divergence. Our study offers a paradigm for elucidating the evolutionary mechanisms of natural enzymes and establishes a foundational framework for the rational engineering of biosynthetic pathways.

## Author contributions

Y. S. W. designed the research and wrote the manuscript; R. X. T. provided financial support; J. Z. directed the protein crystallographic analysis; L. L. F., C. H. J., J. Y. M., H. Z. G. and X. M. performed the biochemical experiments; Y. S. W., W. B. and L. X. analyzed the data.

## Conflicts of interest

The authors declare no conflicts of interest.

## Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information: Tables S1–S9, Fig. S1–S24, NMR spectra and further experimental details. See DOI: <https://doi.org/10.1039/d6sc00451b>.

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