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Structural and Biochemical Studies of the Hedamycin Type II Polyketide Ketoreductase (*HedKR*): Molecular Basis of Stereo- and Regio-Specificities

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Abstract

Bacterial aromatic polyketides that include many antibiotic and antitumor therapeutics are biosynthesized by the type II polyketide synthase (PKS), which consists of 5 – 10 stand-alone enzymatic domains. Hedamycin, an antitumor antibiotic polyketide, is uniquely primed with a hexadienyl group generated by a type I PKS followed by coupling to a downstream type II PKS to biosynthesize a 24-carbon polyketide, whose C9 position is reduced by hedamycin type II ketoreductase (*hedKR*). *HedKR* is homologous to the actinorhodin KR (*actKR*), for which we have conducted extensive structural studies previously. How *hedKR* can accommodate a longer polyketide substrate than the *actKR*, and the molecular basis of its regio- and stereospecificities, is not well understood. Here we present a detailed study of *hedKR* that sheds light on its specificity. Sequence alignment of KRs predicts that *hedKR* is less active than *actKR*, with significant differences in substrate/inhibitor recognition. *In vitro* and *in vivo* assays of *hedKR* confirmed this hypothesis. The *hedKR* crystal structure further provides the molecular basis for the observed differences between *hedKR* and *actKR* in the recognition of substrates and inhibitors. Instead of the 94-PGG-96 motif observed in *actKR*, *hedKR* has the 92-NGG-94 motif, leading to *S*-dominant stereospecificity, whose molecular basis can be explained by the crystal structure. Together with mutations, assay results, docking simulations, and the *hedKR* crystal structure, a model for the observed regio- and stereospecificities is presented herein that elucidates how different type II KRs recognize substrates with different chain lengths, yet precisely reduce only the C9-carbonyl group. The molecular features of *hedKR* important for regio- and stereospecificities can potentially be applied to biosynthesize new polyketides via protein engineering that rationally controls polyketide ketoreduction.

Polyketides represent a large class of natural products that are diverse both in terms of chemical structure and bioactivity (1). Many of these compounds are pharmaceutically important and can act, for example, as antibiotic, anticancer and antihypercholesterolemic agents (2-4). The ensemble of enzymes contributing to polyketide biosynthesis is known as the polyketide synthase (PKS), which is genetically, structurally and functionally similar to fatty acid synthase (FAS) (5). There are generally three classes of PKS (6) (type I, type II,

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The atomic coordinates have been deposited in the Protein Data Bank (accession code 3SJU).

and type III). In the type I system, enzyme domains are covalently linked together, whereas in type II systems the enzymes are present as discrete proteins. The focus of this study is on enzymes of the type II PKS that are found mainly in bacteria and biosynthesize aromatic polyketides such as the antibiotics actinorhodin and tetracenomycin (Figure 1) (2, 7). In a typical type II PKS system (5) (Figure 2A), a linear poly- β -keto chain is synthesized by the minimal PKS (ketosynthase/chain length factor [KS/CLF] and acyl carrier protein [ACP]) (8). The chain is initiated by an acetate unit derived from decarboxylation of a malonyl-ACP thioester, or by an “initiation module” that supplies an acyl unit ranging from 3 – 6 carbons (9), and iteratively elongated by decarboxylative condensation of malonyl-CoA, to yield the full-length polyketide intermediate. After first-ring cyclization, typically between C7-C12 or C9-C14 (10), an optional ketoreductase (KR) can then catalyze the regiospecific reduction of a single carbonyl group (typically at C9) to a hydroxyl group (11). The KR is an important component of the PKS. Previous studies indicate that KR may direct the C7-C12 first-ring cyclization and introduce stereochemistry in the polyketide intermediate upon reduction (12, 13). The molecular determinants of stereospecificity in various KR domains from type I PKS systems have been characterized through site-directed mutagenesis and biochemical analyses (14-16). However, it is unclear whether the conclusions drawn from type II *actKR* studies (11, 17) may also be applied to other type II KRs.

Hedamycin is a pluramycin-type antitumor antibiotic produced by *Streptomyces griseoruber* that mediates its biological activity in part through intercalation in DNA (18-20) (Figure 1). One of the unique properties of this aromatic polyketide stems from its biosynthesis. In the actinorhodin (*act*) system and many other type II PKSs, polyketide chain synthesis is initiated, or primed, by a two-carbon acetate unit and the chain is elongated by the iterative condensation of malonate-derived units (Figure 2A). The polyketide precursor of hedamycin, however, is primed by a unique 2,4-hexadienyl unit that is produced by type I PKSs, HedT and HedU (21), which pass the starter unit to a downstream type II PKS to extend the chain through nine elongation steps and ultimately synthesize a 24-carbon polyketide (Figure 2B). Previous studies on the hedamycin (*hed*) type II PKS suggest that as part of the type II PKS, *hedKR* regiospecifically reduces the C9-carbonyl of the polyketide substrate, similar to *actKR* (22). However, *in vivo* heterologous expression also implied that *hedKR* is important for chain length determination, the molecular basis of which is unclear. The goal of this work is to elucidate the biochemical properties of *hedKR* and determine a possible structural basis for C9-ketoreduction regiospecificity, given the unique hedamycin polyketide substrate that is eight carbons longer than that of *actKR*. Our previous structural, assay and docking studies of *actKR* led to the identification of a residue motif (94-PGG-96) that acts as a determinant of stereospecificity (23). However, without further information in the form of crystal structures of additional type II polyketide KRs, it is not possible to assess whether this motif may play a similar role in other type II KRs or to establish a common molecular basis for regiospecific ketoreduction. The HedA KR (hereafter referred to as *hedKR*) from the hedamycin PKS represents a new target for further structural and biochemical analyses of type II polyketide KRs. Herein we present many interesting differences between *actKR* and *hedKR* in regard to substrate/inhibitor specificity, stereospecificity, and ACP-KR interactions. Additionally, the structural and functional studies presented help bridge the current knowledge gap on chain length control in which the type II polyketide KRs may be involved.

MATERIALS AND METHODS

Materials

trans-1-Decalone, *S*-(+)-tetralol, *R*-(-)-tetralol, cofactor NADP(H), and emodin were purchased from Sigma, and were the highest grade available. Quercetin was purchased from

Acros Organics. Custom oligonucleotides were purchased from Operon Biotechnologies. All organic solvents including DMSO were HPLC grade or better and purchased from Fisher.

DNA Manipulation

Plasmid pAD224 consists of the wild type *hedA* gene cloned into a pET28 expression vector (Novagen). The N92L and N92P single mutations were generated using the QuikChange II Site-Directed Mutagenesis Kit (Stratagene) with pAD224 as the template. *E. coli* NovaBlue cells were used for plasmid amplification. All mutations were confirmed by sequencing.

HedKR Protein Expression and Purification

Recombinant wild type or mutant *hedKR* was expressed in *E. coli* strain BL21(DE3). Following transformation, cells were grown at 37 °C in 1 L Luria-Bertani media supplemented with 50 µg/mL kanamycin to an OD₆₀₀ of 0.6-0.8. At that point, protein expression was induced by the addition of 0.1 mM IPTG at 18 °C overnight. Cells were harvested by centrifugation (5000 rpm × 30 min), resuspended in 50 mL buffer A (50 mM Tris-Cl pH 7.5, 300 mM NaCl, and 10% glycerol) at 4 °C, and lysed on ice by sonication (5 × 30 sec pulses). Cell debris were removed by centrifugation (14000 rpm × 45 min) followed by binding of lysate supernatant to 3 mL nickel IMAC resin (Bio-Rad) in batch-mode at 4 °C. Bound protein was washed with 10 mL of 10 mM imidazole in buffer A and then with 20 mL of 20 mM imidazole in buffer A. *HedKR* was eluted at 40, 60, 80, 100, 150, and 250 mM imidazole in buffer A. The last four fractions were pooled together and dialyzed at 4 °C overnight against 4 L buffer A. The protein was concentrated to ~10 mg/mL with Pierce iCON 9000 MWCO protein concentrators.

In Vitro Tetralol Assay for HedKR Stereospecificity

Steady-state kinetic parameters for wild type and mutant *hedKR* were determined by monitoring the oxidation of *S*-(+)- or *R*-(-)-tetralol in the presence of NADP⁺. The change in absorbance from the conversion of NADP⁺ to NADPH was monitored at 340 nm ($\epsilon_{340} = 6220 \text{ M}^{-1} \text{ cm}^{-1}$) on a DU 800 Spectrophotometer (Beckman Coulter) over 10 minutes. All assays were performed in 400 mM KPi buffer (pH 7.4) and 5% DMSO at 30 °C, and were initiated with the addition of enzyme at a concentration of 1.0-1.7 µM. The Michaelis-Menten constants K_m and k_{cat} for each substrate were obtained by varying the substrate concentration in the presence of 375 M NADP⁺. Data were fitted to the Michaelis-Menten equation using the program KaleidaGraph (Synergy).

ActKS/CLF Expression and Purification

The expression of *actKS/CLF* from *Streptomyces coelicolor* CH999/pRJC006 spores has been described previously (24). Briefly, spores were grown in 50 mL Super YEME containing 50 µg/mL kanamycin for 3 days at 30 °C shaking at 250 rpm. The mycelia were then transferred to 500 mL Super YEME containing 50 µg/mL kanamycin and grown as before for 2 days. Protein expression was induced by the addition of 5 µg/mL thiostrepton, and the cell growth continued as before for 1 day. Cells were harvested by centrifugation (5000 rpm × 30 min), resuspended in 40 mL lysis buffer (100 mM KPi pH 7.5, 0.1% Triton X-100, 5 mM TCEP, 1.5 mM benzamidine, 1 tablet EDTA-free protease inhibitor cocktail [Roche], and 10% glycerol), and lysed on ice by sonication (8 × 1 min pulses). Cell debris were removed by centrifugation (14000 rpm × 30 min) followed by binding of lysate supernatant to 3 mL nickel IMAC resin (Bio-Rad) in batch-mode by spinning at 4 °C for 2 hrs. Protein was eluted with increasing concentrations of imidazole in 100 mM KPi pH 7.5, 500 mM NaCl, and 10% glycerol. Fractions containing *actKS/CLF* were collected at 150 and 500 mM imidazole, pooled, and buffer-exchanged to 100 mM KPi pH 7.2 and 20% glycerol.

Holo-ActACP (C17S) and MAT Expression and Purification

E. coli BAP1 cells (25) expressing pTLF-569 (C17S *actI*-ORF3 in pET28b, provided by Robert W. Haushalter of the Burkart research group [Department of Chemistry and Biochemistry, University of California, San Diego]) were grown at 37 °C in 1 L Luria-Bertani media supplemented with 50 µg/mL kanamycin to an OD₆₀₀ of 0.6-0.8. At that point, protein expression was induced by the addition of 0.1 mM IPTG at 18 °C overnight. Cells were harvested by centrifugation (5000 rpm × 30 min), resuspended in 40 mL buffer D (25 mM K₂HPO₄ pH 7.5, 100 mM NaCl, and 10% glycerol) at 4 °C, and lysed on ice by sonication (5 × 30 sec pulses). The cell debris were removed by centrifugation (14000 rpm × 45 min) followed by binding of lysate supernatant to 5 mL nickel IMAC resin (Bio-Rad) in batch-mode at 4 °C. Protein was eluted with increasing concentrations of imidazole in buffer D. A fraction containing pure holo-*actACP* (C17S) was eluted at 500 mM imidazole and buffer exchanged to buffer D. *S. coelicolor* MAT was expressed and purified from *E. coli* BL21(DE3)/pGFL16 by nickel IMAC as described previously (26).

In Vitro PKS Reconstitution Assay

To determine whether *in vitro* *hedKR* can regiospecifically reduce a 16-carbon polyketide substrate at the C9-carbonyl to produce mutactin, 50 µM *hedKR* was incubated with 10 µM *actKS*/CLF, 50 µM holo-*actACP* (C17S), 1 µM MAT, 5 mM malonyl-CoA, and 2 mM NADPH. The total reaction volume was brought to 250 µL by the addition of 100 mM Tris-Cl pH 7.0 and incubated at room temperature overnight in the dark. The mixture was extracted once with 300 µL of 94% ethyl acetate, 5% methanol, and 1% acetic acid. Solvent was evaporated, products resuspended in 100 µL DMSO, and 20 µL subjected to reverse phase HPLC on a Synergi Hydro-RP analytical column (Phenomenex, 4 µ, 150 × 4.6 mm). For comparison, the same assay was conducted using *hedKR* mutants N92L and N92P, as well as WT *actKR* purified as described previously (17), in place of *hedKR*.

Protein Crystallization and Data Collection

Crystals of wild type *hedKR* were grown in hanging-drops at 15 °C by vapor diffusion. Drops were generated by mixing 1 µL protein with 1 µL well buffer above a well solution of 500 µL. The protein solution consisted of 5 mg/mL *hedKR*, 2 mM NADPH, and 1 mM DMAC as a potential inhibitor. Crystals were flash-frozen in liquid nitrogen after soaking in 30% glycerol. Data were collected on beamline 8.2.2 at the Advanced Light Source (ALS) to 2.4 Å. Diffraction intensities were indexed, integrated and scaled using HKL-2000 (27).

Phasing by Molecular Replacement and Refinement

The wild type *hedKR* structure was solved by molecular replacement using Phaser in CCP4i (28). The search model consisted of a homology model of *hedKR* prepared by SwissModel (29), based on the previously reported wild type *actKR* structure (PDB accession code 1X7H). Following an initial round of refinement with Refmac5 (30), manual rebuilding was performed and waters were added using Coot (31). Subsequent rounds of refinement in Refmac5 and rebuilding in Coot were conducted until the final *R* and *R*_{free} were 0.186 and 0.248, respectively. The quality of the final structure was analyzed with Procheck (32). All crystallographic statistics are listed in Table 3.

Inhibition Kinetics of HedKR

To determine whether quercetin is an inhibitor of *hedKR*, enzyme activity was measured by varying *trans*-1-decalone concentration in the presence of 375 µM NADPH and 0, 25, 50, or 75 µM quercetin. Activity was also measured by varying NADPH concentration in the presence of 2 mM *trans*-1-decalone and 0, 12.5, 25, or 50 µM quercetin. All assays were

performed in 400 mM KPi buffer (pH 7.4) containing 2% DMSO at 30 °C and were initiated with the addition of the enzyme at a concentration of 690 nM.

RESULTS AND DISCUSSION

Sequence Alignment Suggests that HedKR and ActKR Have Different Structures and Activities

HedKR is predicted to contain a short-chain dehydrogenase/reductase (SDR) fold (33), similar to other type II polyketide KRs. The *hedKR* protein sequence was aligned to those of other type II polyketide KRs including *actKR*, with sequence identity ranging from 61-66% (Figure 3). The characteristic sequence motifs found in SDR enzymes (Table 2 in ref (33)) are conserved in *hedKR*, such as the cofactor-binding motif TGxxxGxG (*hedKR* residues 10-17), as well as the active tetrad residues Asn112, Ser142, Tyr155, and Lys159. Notably, the NNAG motif in SDR enzymes becomes 87-NSAG-90 in *hedKR*. The NNAG motif was proposed to stabilize the central β -sheet (34), and is highly conserved in the other type II polyketide KRs except *hedKR*. In the bacterial 3β -hydroxysteroid dehydrogenase, an alanine mutation of N87 (equivalent to S88 in *hedKR*) lowered the reductive and oxidative k_{cat}/K_m values by 39 and 83%, respectively (34). Therefore, sequence alignment suggests that *hedKR* may have lower enzymatic activity than *actKR* due to the change of the NNAG motif to NSAG (detailed below in the *in vitro* assay sections).

In the active site, > 70% of the residues that define the substrate-binding pocket (based on *actKR*-emodin crystal structures (17)) are conserved between *actKR* and *hedKR*, such as Q147, V149, A152, M192, and V196. However, Phe189 of *actKR*, which interacts with both emodin and cofactor, becomes Tyr187 of *hedKR*, indicating subtle changes near the active site. Moreover, the “arginine patch” of *actKR*, proposed to be the docking point of incoming ACP and PPT-phosphate (23, 35), is not conserved in *hedKR*. Specifically, Arg65 of *actKR* becomes Thr63 of *hedKR*. Therefore, sequence alignment predicts that the interaction between the *hedACP* and *hedKR* differs from the corresponding interaction in the *act* PKS. Corroborating the above hypothesis, the *hedACP* differs significantly from the actinorhodin ACP (*actACP*): while the significant majority of type II polyketide ACPs exist as stand-alone proteins, the *hedACP* is covalently linked to the aromatase/cyclase as the bifunctional protein HedE (21). The difference between the *hedKR* and *actKR* arginine patches may therefore be intimately related to the different domain arrangement of *hedACP*.

In terms of stereospecificity, mutational studies of *actKR* showed that the “PGG” motif is important for the observed *S*-dominant stereospecificity, with a 3:1 (*S*:*R*) preference *in vitro* for the oxidation of *S*- versus *R*-tetralol (23). In *hedKR*, “PGG” becomes “NGG,” suggesting that the stereospecificity of *hedKR* may differ from that of wild type *actKR*. Altogether, the sequence alignment results show that *hedKR* has the SDR fold, and its structural features are highly conserved with those of *actKR*; however, enzymatic activity, interactions with ACP, and stereospecificity of *hedKR* may differ from those of *actKR*, reflecting the differences in substrates, domain architecture, and stereochemical requirement of hedamycin versus actinorhodin PKSs.

In Vitro Assays Support that HedKR is Much Less Active than ActKR

To determine *in vitro* reductase activity, *hedKR* was assayed with the substrate *trans*-1-decalone, as had been previously done with *actKR* (17). The steady-state kinetic parameters for reduction of *trans*-1-decalone by *hedKR* or *actKR* can be found in Table 1. The results show that *hedKR* can reduce *trans*-1-decalone *in vitro*. However, while the K_m values are similar, the k_{cat} of *hedKR* is ~23 times less than that of *actKR*. Similarly, when we assayed *hedKR* activity from the reverse direction, through the oxidation of *S*- and *R*-tetralol, we

found no activity for *R*-tetralol, and the k_{cat}/K_m of *hedKR* for *S*-tetralol is 20-fold less than that of *actKR* (Table 2). The above *in vitro* assay result is consistent with sequence-based prediction, that the change of the “NNAG” motif of *actKR* to “NSAG” of *hedKR* reflects a change in active site geometry, resulting in the decrease in enzymatic activity.

During hedamycin biosynthesis, *hedKR* reduces a 24-carbon substrate that is primed with a six-carbon starter unit (Figure 2B). To evaluate the substrate specificity and regiospecificity of *hedKR*, we sought to assess whether *hedKR* can reduce the 16-carbon octaketide intermediate synthesized by the actinorhodin minimal PKS (*act* min PKS), and if the ketoreduction still occurs at the C9-carbonyl group. To determine whether the C9-specificity is promoted by *hedKR* itself, we conducted *in vitro* PKS reconstitution assays, in which purified *actKS*/CLF, *S. coelicolor* MAT, and holo-*actACP* (C17S) were incubated with or without *hedKR* or *actKR*. Here, holo-*actACP* (C17S) is used instead of the wild type *actACP* due to the tendency of wild type *actACP* to dimerize via disulfide bond formation that interferes with experiments. The reconstituted minimal PKS will produce the 16-carbon octaketide chain, but it was unknown whether there would be sufficient effective interactions between *hedKR*, *actACP* and *actKS*/CLF for substrate transfer and ketoreduction to occur. HPLC analysis of the reconstitution products demonstrates that *act* min PKS + *hedKR* produces the 16-carbon, C9-reduced mutactin, albeit at a much lower level than in the presence of *actKR*, presumably due to weaker interactions between *hedKR*, *actACP* (C17S), and *actKS*/CLF (Figure 4). Based on the areas under the peaks corresponding to mutactin, in the same amount of incubation time, *hedKR* produces ~fivefold less mutactin than *actKR* does. The reconstitution and HPLC results are consistent with our *in vitro* assay result of *trans*-1-decalone, as well as the heterologous expression results by Das and Khosla, in which *hedKR* is functionally interchangeable with *actKR* (22). It should be noted that the previous work was based on *in vivo* protein expression and product characterization; the present study indicates *hedKR* may not be as active as *actKR* *in vitro*, which is consistent with sequence analysis of the SDR “NSAG” motif. Nevertheless, the assays do affirm that *hedKR* is functional *in vitro* and is capable of regiospecific C9-ketoreduction of a 16-carbon polyketide chain. The above result supports that the C9-regiospecificity of type II polyketide KR is not closely related to the number of carbons (referred to as “chain length” throughout the text) of the incoming polyketide substrate.

Inhibition Kinetic Assays Suggest that Quercetin, but not Emodin, is an Inhibitor of HedKR

We previously conducted extensive inhibition kinetic assays of emodin, which serves as a competitive inhibitor of *actKR*. Emodin contains three conjugated aromatic rings. We further solved the ternary structures of WT or P94L *actKR* bound to NADPH and the inhibitor emodin (17). Analyses of these structures revealed that emodin-binding changes the protein conformations of *actKR* (17). Guided by the insights drawn from the critical structural analysis, we sought to identify an effective inhibitor of *hedKR* by screening the inhibitory effect of four plant secondary metabolites on the reduction of *trans*-1-decalone (Figure 5A). Of particular interest were emodin and compounds from green tea extract such as epigallocatechin gallate (EGCG), which has been shown to inhibit the fatty acid KR in bacteria (36). The sequence, structural and functional similarities between fatty acid synthase (FAS) and type II PKS KRs (5) suggests that green tea extract compounds may also act as inhibitors of *hedKR*. Single-point inhibition kinetic studies showed that the flavonoid quercetin (Figure 5A) displayed the most potent inhibitory effects of the four compounds tested. Further in-depth assays were conducted to identify the mode of inhibition of quercetin (Figure 5B). The compound acts as a competitive inhibitor of *trans*-1-decalone, with a K_i of approximately 114 μM . Significantly, *actKR* is inhibited by emodin but not quercetin, while the inhibitory effect of quercetin is much higher than that of emodin toward

hedKR. Therefore, although *actKR* and *hedKR* were predicted to have similar structures, both sequence alignment and inhibition kinetics predict that the substrate/inhibitor-binding pockets will be different.

HedKR Crystallizes under Markedly Different Conditions than ActKR

To establish a structural basis for the observed substrate/inhibitor- and regiospecificities of *hedKR*, we sought to solve the *hedKR* crystal structure. Our initial attempts to crystallize *hedKR* based on conditions that led to effective *actKR* crystallization were unsuccessful (11, 17, 23). To help stabilize any possible protein flexibility, *hedKR* was incubated with the cofactor NADPH. After extensive screening, diffraction-quality *hedKR* crystals were produced in a condition completely different from that of *actKR*, with different crystal morphology. The differences in growth conditions and crystal morphology between *hedKR* and *actKR* suggest that the structural features of the two enzymes, such as protein conformation, may differ as well.

HedKR Overall Architecture is Similar to that of ActKR, but the Substrate Pockets Differ

The 2.4-Å crystal structure of wild type *hedKR* allows for detailed analysis and comparisons to the previously solved *actKR* structures. Although *hedKR* crystallizes in a different space group than *actKR*, the asymmetric unit similarly consists of two monomers, and the natural *hedKR* tetramer can be generated across the two-fold symmetry axis (Figure 6A). Each monomer consists of the characteristic Rossmann fold (37) for binding nucleotide cofactors. There is clear electron density for NADPH in each monomer of the crystal structure. An overlay of the two *hedKR* monomers shows that the catalytic tetrad residues (N112, S142, Y155, and K159) and cofactors adopt the same position and conformation between monomers. A significant difference is that Met192 in monomer B is more extended than in monomer A. This residue lies opposite Asn92 on the other side of the substrate pocket cleft, resulting in an effective closed conformation for monomer B versus the open conformation of monomer A, (Figure 6B).

The overall *hedKR* structure is very similar to that of the previously solved *actKR* structures, namely WT-NADPH, emodin-bound WT-NADPH, P94L-NADPH, and emodin-bound P94L-NADPH (11, 17, 23). The RMS deviations between the *hedKR* structural dimer and these *actKR* structures are, respectively, 0.73, 0.75, 0.77, and 0.76 Å. As with the *actKR* structures, the $\alpha 6$ - $\alpha 7$ loop region in the *hedKR* structure is disordered and displays weak $2F_o - F_c$ map electron density, although this region in monomer A is better defined than in monomer B. A comparison of the catalytic tetrad residues and cofactors between each *hedKR* and *actKR* structure shows that the positions and conformations of these components are similar, suggesting that the observed differences in *in vitro* enzyme activity between *hedKR* and *actKR* are not the result of catalytic residue misalignment, but likely stem from differences in substrate-binding due to different substrate pocket shapes.

Consistent with the above analysis, although the overall structures are similar, we found many structural features in the substrate pockets that differentiate *hedKR* from *actKR* (Figure 6C). Across from the $\alpha 6$ - $\alpha 7$ loop region in *hedKR* are three residues with longer sidechains than the corresponding residues in *actKR*: Glu96, Met150 and Tyr151 (*actKR* residues Ala98, Val152 and His153) (Figure 6D), which change the substrate pocket shape. Further, *hedKR* residues Tyr187 and Phe215 point into the substrate pocket, resulting in a more constricted pocket size in *actKR*. This narrowing of the substrate pocket is especially pronounced when *hedKR* monomer B is compared to the corresponding WT-NADPH *actKR* monomer, where the phenyl ring of Phe215 protrudes into the substrate pocket, which can potentially hinder access to the active site from the opening of the enzyme cleft. This difference is readily apparent when the *hedKR* structure is aligned with P94L-NADPH-

emodin *actKR*, where the *actKR* Phe189 sidechain points away from the substrate-binding pocket, resulting in a wider pocket for *actKR*. Therefore, the difference in substrate pocket shape may likely translate to differences between *actKR* and *hedKR* in the binding motifs of substrates, inhibitors and transition states. The above analysis corroborates the differences in k_{cat} and k_{cat}/K_m between *actKR* and *hedKR* for the *in vitro* reduction of *trans*-1-decalone and oxidation of *R*- and *S*-tetralols (Tables 1-2), as well as differences in inhibitor binding (detailed below).

The conformation of *hedKR* Met192 versus *actKR* Met194 differs as well. Compared to both the binary and ternary WT *actKR* structures, Met192 of *hedKR* monomer A has a more extended conformation than the corresponding Met194 of *actKR*. Met192 in *hedKR* monomer B, however, has a very similar conformation to Met194 in WT-NADPH-emodin *actKR* monomer B. As previously reported, monomer B of the ternary WT *actKR* structure has a distinctly closed conformation relative to monomer A (11, 17, 23). The observation that *hedKR* monomer B also has a closed conformation suggests that in *hedKR* the methionine opposite the “NGG motif” is important for the open/closed conformation by interacting with Asn92 (Figure 6B). This notion is also supported by a comparison of the *hedKR* structure with P94L-NADPH *actKR*: in both structures, monomer A is open and monomer B is closed, while the corresponding methionine residues in both structures have the same side-chain conformation.

Another important difference between *hedKR* and *actKR* is the presence of Asn92, which juts into the active site nearly the same distance as Leu94 in the P94L *actKR* structures does. Based on this observation, it is likely that the stereospecificity of WT *hedKR* will correspond to that of P94L *actKR*, as opposed to WT *actKR*. The structural comparisons between *hedKR* and *actKR* indicate that despite a similar overall architecture, the substrate-binding pockets of *actKR* and *hedKR* adopt distinct shapes, which offers a strong structural basis to interpret the observed differences in *in vitro* activity, substrate/inhibitor specificity, and stereospecificity.

Molecular Basis of Inhibitor Specificity between HedKR and ActKR

To understand how different inhibitors can be specifically recognized by *actKR* versus *hedKR*, we simulated quercetin binding at the *hedKR* active site by using the docking program GOLD (38). With no constraint bias, we obtained highly consistent docking solutions that have similar protein conformations, inhibitor positions, and orientations within the substrate pocket (Figure 7A). One of the quercetin hydroxyl groups is oriented within hydrogen-bonding distance of the side chain hydroxyl groups of active site Tyr155 and Ser142, which form the catalytic oxyanion hole. The hydroxyl-bearing carbon of quercetin is also within hydride-transfer distance of C4 of the nicotinamide ring. When quercetin was docked to the P94L-NADPH *actKR* structure, the docking solutions were not as consistent as within the WT *hedKR* structure. Reciprocally, when we tried docking the *actKR* inhibitor emodin into the WT *hedKR* structure, we also could not obtain a solution whose emodin-binding motif is consistent with those from the reported *actKR*-NADPH-emodin crystal structures. Moreover, the emodin molecules from the WT and P94L *actKR* structures did not overlay with any of the top-ten *hedKR*-emodin docking solutions at all (Figure 7B). In essence, the inhibitors emodin and quercetin serves as “molecular probes”: the wider, tricyclic emodin can inhibit the wider pocket of *actKR*, but can barely bind the narrower pocket of *hedKR*, while the reverse is true for *hedKR*. The above docking results support that the observed differences in inhibitor binding, substrate specificity, and enzymatic activity between *hedKR* and *actKR* are reflected by pocket size and shape differences.

Based on the proximity to the docked quercetin, N92, G186, Y187, V196, and L256 are predicted to define the binding pocket and likely affect the stereospecificity of *hedKR*

(Figure 7C). Therefore, the *hedKR* crystal structure and docking simulations lead to a structure-based hypothesis on KR stereospecificity that is consistent with the one observed with *actKR*, that potential steric interference by the active site residues lead to an *S*-dominant stereospecificity in type II polyketide KRs (11, 17, 23).

Wild Type HedKR Displays High Stereospecificity in Comparison to Wild Type ActKR

To assess the stereospecificity of *hedKR*, we utilize the ability of a type II KR to catalyze the reverse reaction, namely the oxidation of *S*- and *R*-tetralol (39). Previously, we showed that the “PGG” motif in *actKR* is a key determinant of stereospecificity. While wild type *actKR* has a threefold preference (k_{cat}/K_m) for *S*- versus *R*-tetralol, P94L *actKR* is highly specific for only *S*-tetralol (Table 2). Because the “PGG” motif of *actKR* becomes 92-NGG-94 in *hedKR*, with Asn92 jutting into the active site and creating a similar pocket shape as that of P94L *actKR*, we predicted that wild type *hedKR* would have similar stereospecificity to that of P94L *actKR*, namely *S*-dominant. Supporting this hypothesis, when assayed separately with *S*- and *R*-tetralol, *hedKR* does displays a dominant stereospecificity for *S*-tetralol (Table 2). Hence, steric interference is likely to occur in WT *hedKR* just as in P94L *actKR* and suggests that the “XGG” motif may be a general determinant of stereospecificity for most type II polyketide KRs.

Analysis of Mutant HedKR Stereospecificity

To determine whether N92 in the “NGG” motif is the most dominant factor that determines the stereospecificity of *hedKR*, two single mutants were prepared: N92L and N92P *hedKR*. The rationale for the N92L mutant is to mimic P94L *actKR*, whereas the N92P mutant was constructed to mimic WT *actKR* and, in theory, would display the same threefold preference for *S*- over *R*-tetralol. The *hedKR* mutants were subjected to the *in vitro* assay as before (Tables 1-2). Surprisingly, both mutants displayed very low levels of activity in the presence of *R*-tetralol. Moreover, based on the catalytic specificity constant, k_{cat}/K_m , N92L is about 18 times more active toward *S*-tetralol than N92P, while both N92L and N92P are much more active than the WT *hedKR*. It is likely that in N92L a favorable hydrophobic interaction between the Leu side chain and the *S*-tetralol cyclohexane ring may account for the higher activity; in comparison, the Asn residue in WT *hedKR* cannot participate in such non-polar interactions, resulting in an elevated K_m and hence low k_{cat}/K_m . Alternatively, the N92L substrate pocket may adopt a different shape than the WT pocket, perhaps due to non-polar interactions between Leu92 and M192, resulting in increased oxidative activity toward *S*-tetralol.

The N92L and N92P mutants of *hedKR* were also assayed through *in vitro* PKS reconstitution (Figure 4). Based on the HPLC chromatogram peak area corresponding to mutactin, both mutants actually produced more mutactin, relative to the SEK4 and SEK4b peak areas, than WT *hedKR*. Specifically, N92L produced almost as much mutactin as SEK4 and SEK4b combined. These results correlate with the tetralol kinetic assay results, suggesting that the presence of a hydrophobic residue at position 92 (such as proline or leucine), results in favorable non-polar interactions between residue 92 and M192. In part, these interactions could contribute to the forces that stabilize the closed conformation of a *hedKR* monomer in order to catalyze polyketide ketoreduction.

The N92P mutant of *hedKR* was constructed to mimic WT *actKR*, which displays measurable activity toward *R*-tetralol. The lack of activity toward *R*-tetralol by the N92P *hedKR* mutant may be due to the rate detection limit of the assay, but this result also implies that the “PGG” motif is not a sole determinant of stereospecificity in *hedKR*. In light of the structural differences between *hedKR* and *actKR*, particularly the distinct substrate pocket shapes, it is likely that there are long-ranged effects responsible for guiding the

stereospecificity of *hedKR*. Namely, stereospecificity may be determined by the shape of the enzyme active site pocket as a whole rather than by specific residues. It has been shown that the tropinone reductases, TR-I and TR-II, have the same overall fold, but opposite reaction stereospecificities (40). Nakajima *et al.* attributed the difference to distinct amino acids lining the substrate pockets of the two enzymes (40). Moreover, it has been suggested that residue side chains across the substrate cavity of 3 α -hydroxysteroid dehydrogenase determine the orientations of substrates and inhibitors (41). Therefore, many residues (as opposed to a single motif) that define the overall shape of the *hedKR* active site pocket and interact with substrates may collectively affect the stereospecificity of a KR. Nevertheless, due to high sequence conservation of pocket residues in type II polyketide KRs, there are likely long-ranged motifs that are important for stereospecificity. Future site-directed mutagenesis studies that focus on such residues can shed light on the molecular basis of long-ranged stereospecificity determinants of *hedKR*.

Biological Significance: C9 Regiospecificity Versus Substrate Chain Length

The number of carbons in a given type II polyketide, referred to as “chain length,” is closely associated with the specificity of the KS/CLF of the corresponding type II PKS (42). This hypothesis was strongly supported by mutational studies of *act*CLF residues, in which single mutations completely changed the product chain length, thus validating the dominant influence of KS/CLF (42). However, in the *hed* type II PKS, both the *hedKR* and ARO/CYC (HedE) are also important for chain length determination, as concluded from *in vivo* expression experiments of hedamycin type II PKS in a heterologous host. For example, the expression of *hedKS/CLF*, the bifunctional HedE (ACP + ARO/CYC), and *hedKR* resulted in four products (Figure 2B): the full-length, **10** (24 carbons), **11** and **12** (22 carbons), and **13** (8 carbons). In the absence of *hedKR*, the expression of *hedKS/CLF* and HedE resulted in five unreduced products with varying chain lengths: **18** (24 carbons), **17** (22 carbons), **16** (10 carbons), **15** (8 carbons), and **14** (6 carbons). Similar results have been reported for the actinorhodin PKS, that in addition to the full-length product (16 carbons), the *act* PKS also generates truncated products **1-3** (12 carbons) (Figure 2A) (43). In our previous work, we found that the presence of a truncated product is closely associated with the presence of an active *actKR* (23), and it was hypothesized that the truncated products **2** and **3** result from competition of *actKR* with *actKS/CLF*, causing premature reduction of a much shorter chain. In this work, the diverse chain lengths observed in *hed* PKS offer an excellent opportunity to examine the above hypothesis. The *in vivo* expression experiments of *hed* PKS can be explained by two possible reasons: (1) that the pocket size of *hedKR* is a partial determinant of the polyketide chain lengths, or (2) that similar to *act* PKS, the chain length is a result of competition between chain elongation (by *hedKS/CLF*) and ketoreduction (by *hedKR*). The *hedKR* crystal structure and enzyme kinetics help distinguish these two possibilities. In this work, we showed that *hedKR* can reduce both 16- and 24-carbon polyketide substrates. In concert with this result, docking simulations consistently docked the C9-carbonyl group of both 16- or 24-carbon, mono-cyclized polyketides into the *hedKR* active site. Therefore, while the substrate pockets of *hedKR* and *actKR* are different in shape (reflected by their different inhibitor specificities, Figure 6B), the KR active site itself is not specific to the chain length of an incoming substrate. Consequently, the *hedKR* crystal structure and docking simulations of PPT-tethered polyketides (either 16 or 24 carbons, Figure 8A-B) confirm that the C9-specificity arises from the prerequisite that the first ring is cyclized between C7-C12. Supporting this observation, and consistent with the *in vivo* expression results by Das and Khosla, docking simulations consistently docked carbon 9 of a C7-C12 cyclized (Figure 8A-B), or carbon 7 of a C5-C10 cyclized polyketide (Figure 8C) to the active site of *hedKR*. Combining the above together, the *hedKR* structural and functional studies confirmed that the C9-specificity requires the constraint of a pre-formed ring, and that the observed truncated products reflect a competition between chain

elongation (by KS/CLF), first-ring cyclization (by either KS/CLF or KR), and ketoreduction (by KR). Note, also, that the chain length is required to be at least 10 carbons or longer to reach from the surface arginine patch (the ACP-PPT docking site) into the *hedKR* active site. Therefore, the presence of the 8-carbon, mono-cyclized **13** requires that the ring is pre-formed, thus freeing the chain from the ACP-PPT tether, before reaching the *hedKR* active site (Figure 7D). Finally, mutational analyses of *hedKR* confirmed the importance of the “XGG” motif for stereospecificity, but also suggested that more long-ranged motifs may be involved. The above results pave a foundation for future work toward stereospecific engineering of type II PKSs to generate new polyketides.

Acknowledgments

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Abbreviations

KR	ketoreductase
FabG	β -Ketoacyl (Acyl Carrier Protein) Reductase
act	actinorhodin
hed	hedamycin
PKS	polyketide synthase
NADP	nicotinamide adenine dinucleotide phosphate
NADPH	reduced nicotinamide adenine dinucleotide phosphate
SDR	short-chain dehydrogenase/reductase
ACP	acyl carrier protein

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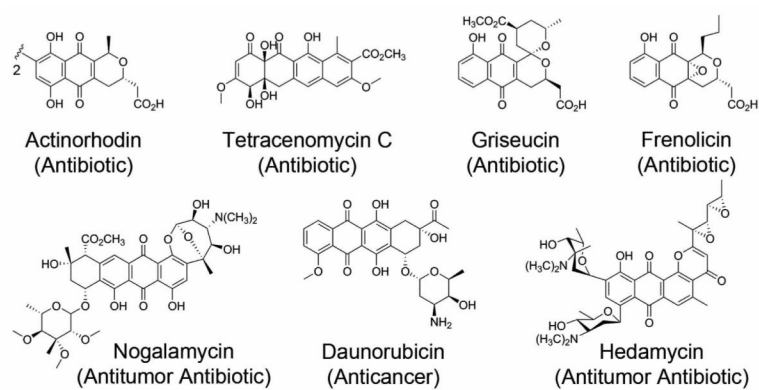
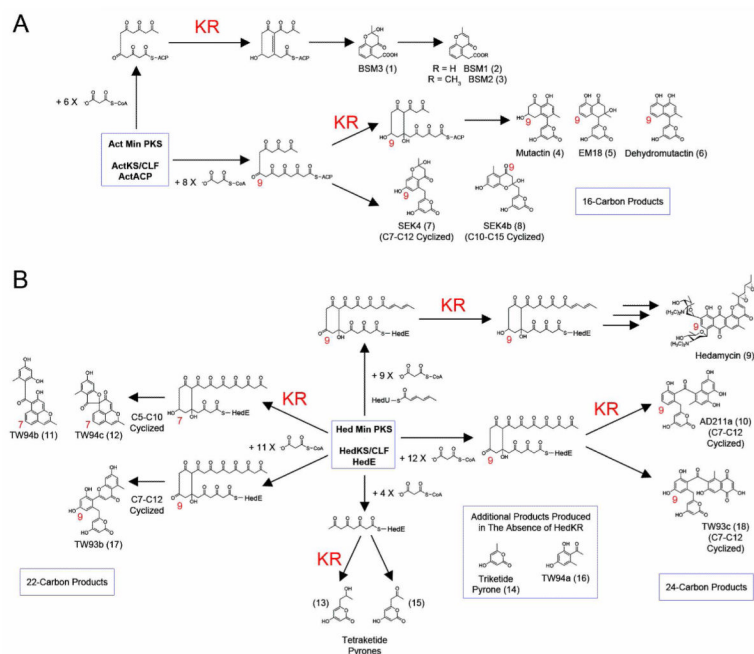


Figure 1.
Representative type II polyketides.

**Figure 2.**

Type II PKS pathway intermediates in the presence or absence of KR. (A) The actinorhodin system, in which the polyketide chain is initiated by an acetyl group and the presence of *actKR* can lead to 6- or 8-carbon products. (B) The hedamycin system, in which the unique 6-carbon type I polyketide starter unit initiates polyketide chain biosynthesis. In the presence of *hedKR*, 24-, 22- and 8-carbon products are formed, while 6-, 8-, 10-, 22-, and 24-carbon products are formed in the absence of *hedKR*. Red numbers are used to denote which carbon of an intermediate can undergo ketoreduction, and are shown for the final products as well.

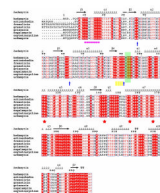


Figure 3.

Sequence alignment among various type II PKS KRs. Sequences included hedamycin, actinorhodin, frenolicin, granaticin, griseucin, nogalamycin, oxytetracycline, and urdamycin KRs. Key: magenta circles, SDR cofactor-binding motif; blue arrow, arginine patch residue; yellow rectangles, SDR motif involved in the stabilization of the central β -sheet; green-tinted box, "PGG" motif; red stars, catalytic residue.

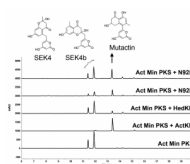


Figure 4.

HPLC analysis of products from *in vitro* reconstitution assays, demonstrating that *hedKR* is able to reduce an octaketide intermediate (produced by the *act* min PKS) at the C9-position to form mutactin (**4**). *HedKR* mutants N92L and N92P produce more mutactin, relative to the amount of SEK4 and SEK4b produced, than WT.

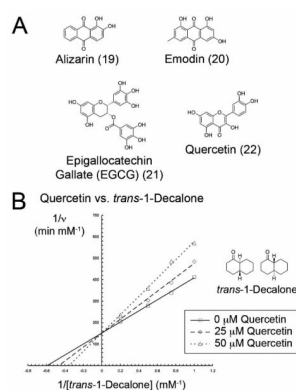


Figure 5.

Inhibition of *hedKR*-catalyzed *trans*-1-decalone reduction. (A) The four plant secondary metabolites tested as potential inhibitors of *hedKR*. (B) Inhibition kinetic assays demonstrate that quercetin (**22**) is a competitive inhibitor of *trans*-1-decalone reduction.

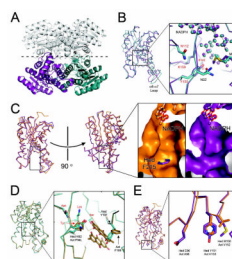


Figure 6.

Structural analysis of *hedKR*. (A) The *hedKR* crystallographic tetramer is formed by rotation of monomers A (teal) and B (purple) about the two-fold symmetry axis shown as a dashed line. (B) Overlay of *hedKR* monomers A (teal) and B (purple). The zoomed view displays the NADPH cofactor (ball and stick model) and important active site residues (sticks). M192 is more extended in monomer B, possibly interacting with N92 through hydrogen-bonding. (C) Comparison of WT *actKR* (orange) and WT *hedKR* (purple) substrate pockets. Portions of the NADPH cofactors are hidden by residues, but outlined in red here for viewing clarity. The *hedKR* pocket is more constricted and narrower than the *actKR* pocket, with F215 forming a ridge that juts into the pocket. The F215 sidechain is overlaid on the *actKR* surface for comparison. (D) Overlay of monomers A of P94L *actKR*-emodin (gold) and WT *hedKR* (teal). Key residues are shown as stick models. *HedKR* Y187 is shown piercing through the emodin molecules, demonstrating how this residue differentiates the *hedKR* active site from that of *actKR*. (E) Overlay of WT *actKR* (orange) and WT *hedKR* (purple), displaying residues across from the $\alpha 6$ - $\alpha 7$ loop region that differ between the two KRs.

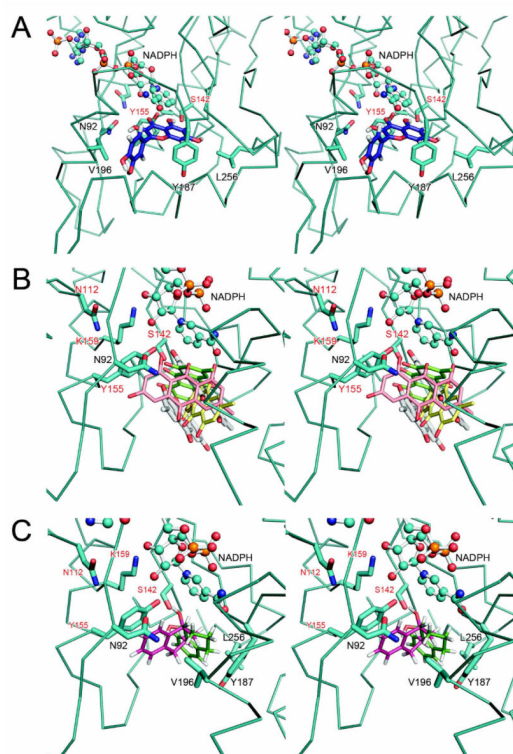


Figure 7.

Stereo views of *hedKR* *in silico* docking analyses. *HedKR* monomer A is portrayed in teal in each panel. (A) Quercetin (blue) docking. A quercetin hydroxyl group is within hydrogen-bonding distance of the catalytic S142 and Y155 sidechains, as well as within hydride-transfer distance from C4 of the NADPH nicotinamide moiety. The residues modeled as sticks are targets for site-directed mutagenesis studies to determine long-ranged substrate pocket motifs that affect *hedKR* stereospecificity. (B) Three examples of emodin docked in *hedKR* (green, pink and yellow), overlaid with the emodin molecules from the previously solved P94L *actKR* cocrystal structure (white). The *hedKR*-emodin docking solutions were not consistent and do not overlap with the emodins in the *actKR* crystal structures. (C) *S*-tetralol (green) and *R*-tetralol (magenta) docked within the *hedKR* active site. The rigid benzene ring of *R*-tetralol is sterically hindered by the N92 sidechain that juts into the pocket.

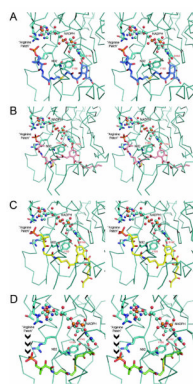


Figure 8.

Stereo views of various PPT-linked polyketides docked to *hedKR*, with monomer A portrayed in teal in each panel. (A) C7-C12 cyclized, 16-carbon polyketide (blue); (B) the putative C7-C12 cyclized, 24-carbon *hedKR* polyketide substrate (pink); and (C) C5-C10 cyclized, 22-carbon polyketide (yellow). These *in silico* docking analyses demonstrate that *hedKR* can accommodate the polyketide substrates portrayed in (A)-(C). (D) None of the 8-carbon polyketide (green) docking solutions placed the PPT phosphate within the “arginine patch,” which is the proposed docking site for ACP. These results suggest that the reduced tetraketide compound (**13**) is formed after release of the thioester-bound polyketide from holo-ACP, spontaneous pyrone ring cyclization, and C7-reduction of the molecule by *hedKR*.

Table 1

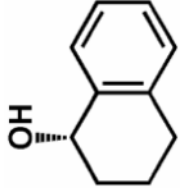
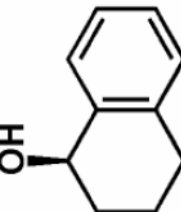
Kinetic Parameters for the Reduction of *trans*-1-Decalone by Wild Type and Mutants of Type II Polyketide Ketoreductases (*ActKR* and *HedKR*)

	k_{cat} (s^{-1})	K_{m} (mM)	$k_{\text{cat}}/K_{\text{m}}$ ($\text{s}^{-1} \text{mM}^{-1}$)
	<i>ActKR</i>		
WT	2.55 ± 0.121	0.790 ± 0.0850	3.23 ± 0.318
P94L	0.720 ± 0.193	0.709 ± 0.367	1.02 ± 0.592
	<i>HedKR</i>		
WT	0.109 ± 0.00758	0.591 ± 0.147	0.184 ± 0.0475
N92L	ND		
N92P	ND		

ND: No detectable activity

Table 2

Kinetic Parameters for the Oxidation of *S*-(+)-Tetralol and *R*-(-)-Tetralol by Wild Type and Mutants of Type II Polyketide Ketoreductases (*ActKR* and *HedKR*)

	<i>S</i> -(+)-Tetralol			<i>R</i> -(-)-Tetralol		
						
	k_{cat} (s^{-1})	K_m (mM)	k_{cat}/K_m ($s^{-1} mM^{-1}$)	k_{cat} (s^{-1})	K_m (mM)	k_{cat}/K_m ($s^{-1} mM^{-1}$)
<i>ActKR</i>						
WT	0.24 ± 0.01	6.91 ± 1.2	0.035 ± 0.006	0.06 ± 0.004	6.47 ± 0.90	0.010 ± 0.001
P94L	0.24 ± 0.03	6.57 ± 1.8	0.036 ± 0.011	ND	>>40	ND
<i>HedKR</i>						
WT	0.168 ± 0.0109	108 ± 7.98	0.00156 ± 0.000153	ND	ND	ND
N92L	0.169 ± 0.00653	4.32 ± 0.521	0.0391 ± 0.00496	ND	ND	ND
N92P	0.0323 ± 0.00166	14.1 ± 1.36	0.00229 ± 0.000249	ND	ND	ND

ND: No detectable activity

Table 3*HedKR* Crystallographic Statistics

	<i>HedKR</i> -NADPH
A. Crystallization	0.1 M imidazole pH 6.2, 50% MPD
B. Crystallographic Data	
Space group	C121
Cell dimension (Å)	116.70 57.39 82.11
	$\alpha = \gamma = 90^\circ$, $\beta = 131.61^\circ$
Resolution (Å)	61.39-2.40
Mosaicity	1.00
No. of observations	114703
No. of unique reflections	15891
Completeness % (last shell)	99.0 (89.0)
I/ σ (I) (last shell)	22.9 (6.7)
R _{merge} % (last shell)	8.5 (23.5)
C. Refinement	
Resolution (Å)	2.40
No. of reflections	15078
No. of protein atoms	3696
No. of cofactor atoms	96
No. of water atoms	36
R _{free} %	24.8
R _{crys} %	18.6
D. Geometry	
RMS bonds (Å)	0.016
RMS angles (°)	1.77
RMS B main chain	0.859
RMS B side chain	2.335
Ramachandran plot (%)	
Most favored	89.4
Favored	10.6
Generously allowed	0