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Structure and Catalytic Mechanism of 3-Ketosteroid-Δ4-(5α)-dehydrogenase from Rhodococcus jostii RHA1 Genome

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Background: Ketosteroid dehydrogenases are enzymes of biotechnological relevance that introduce a double bond into steroids as a first step toward their degradation.

Results: First structures of a 3-ketosteroid-Δ4-(5α)-dehydrogenase combined with mutational analysis allowed the identification of residues essential for catalysis.

Conclusion: Tyr-319, Tyr-466, and Ser-468 have essential roles in catalysis.

Significance: These structures may facilitate the development of better catalysts for steroid conversion.

3-Ketosteroid Δ4-(5α)-dehydrogenases (Δ4-(5α)-KSTDs) are enzymes that introduce a double bond between the C4 and C5 atoms of 3-keto-(5α)-steroids. Here we show that the ro05698 gene from Rhodococcus jostii RHA1 codes for a flavoprotein with Δ4-(5α)-KSTD activity. The 1.6 Å resolution crystal structure of the enzyme revealed three conserved residues (Tyr-319, Tyr-466, and Ser-468) in a pocket near the isoalloxazine ring system of the FAD co-factor. Site-directed mutagenesis of these residues confirmed that they are absolutely essential for catalytic activity. A crystal structure with bound product 4-androstene-3,17-dione showed that Ser-468 is in a position in which it can serve as the base abstracting the 4β-proton from the C4 atom of the substrate. Ser-468 is assisted by Tyr-319, which possibly is involved in shuttling the proton to the solvent. Tyr-466 is at hydrogen bonding distance to the C3 oxygen atom of the substrate and can stabilize the keto-enol intermediate occurring during the reaction. Finally, the FAD N5 atom is in a position to be able to abstract the 5α-hydrogen of the substrate as a hydride ion. These features fully explain the reaction catalyzed by Δ4-(5α)-KSTDs.

Rhodococcus is a genus of aerobic Gram-positive bacteria closely related to Mycobacterium and Corynebacterium (1). Its members show a broad catabolic diversity and a wide range of unique enzymatic capabilities, among which is the ability to degrade naturally occurring sterols. Intermediates of the phytosterol degradation pathway find application in the production of bioactive steroids, making Rhodococcus a particularly interesting organism for steroid bioconversion (2).

An important step in the microbial degradation of various naturally occurring sterols is the desaturation of the steroid A ring (Fig. 1). Several different 3-ketosteroid dehydrogenases (KSTDs) are known to promote this desaturation, such as 3-ketosteroid Δ1-dehydrogenase (Δ1-KSTD), 3-ketosteroid Δ4-(5α)-dehydrogenase (Δ4-(5α)-KSTD), and 3-ketosteroid Δ4-(5β)-dehydrogenase (Δ4-(5β)-KSTD), which are all FAD-containing enzymes. Although Δ1-KSTDs desaturate the C1-C2 position, Δ4-KSTDs introduce a double bond at the C4-C5 position (3, 4).

Amino acid sequence analysis of (putative) KSTDs showed that Δ1-KSTDs and Δ4-KSTDs are clearly distinct, belonging to different branches of the phylogenetic tree. They display 20–24% sequence similarity to fumarate reductases (5). The highest homology is found in the FAD-binding domain; the substrate-binding domain shows lower sequence similarity. The residues involved in the dehydrogenation of the substrate by KSTDs are currently unknown, although chemical modification and mutagenesis studies on Δ1-KSTDs have implicated histidine and tyrosine residues in catalysis (6, 7). Mutational studies confirming the involvement of these residues in Δ4-(5α)-KSTDs have not been reported so far.

KSTDs also have an important function in Mycobacterium tuberculosis where they are part of the metabolic pathway for cholesterol degradation. In M. tuberculosis this pathway is central to the unusual ability of the bacterium to survive inside macrophages (8). Interestingly, transposon mutagenesis has suggested that the Δ4-KSTD homologue Rv1817 is involved in the pathogenicity of M. tuberculosis (9), and it has been proposed that KSTDs could be valuable targets for the development of antituberculosis drugs (8).

In the Rhodococcus jostii RHA1 genome (1), a single Δ4kstD gene (ro05698) is present. It codes for a protein of 490 amino acid residues, with an estimated molecular mass of 51.9 kDa.

3 The abbreviations used are: KSTD, ketosteroid dehydrogenase; 5α-AD, 5α-androstane-3,17-dione; 1-(5α)-AD, 1-(5α)-androstene-3,17-dione; 4-AD, 4-androstene-3,17-dione; ADD, 1,4-androstadiene-3,17-dione; RMSD, root mean square deviation; DCPIP, dichlorophenolindophenol.

http://www.jbc.org/content/suppl/2012/07/24/M112.374306.DC1.html

Supplemental Material can be found at:
The gene has been cloned, and the enzyme has been overexpressed as a His$_6$-tagged protein in *Escherichia coli* and purified (10). Spectroscopic analysis confirmed that it binds FAD, and the purified enzyme was shown to have 3-ketosteroid Δ4-(5α)-dehydrogenase activity. The enzyme has been crystallized, and its preliminary x-ray analysis has been published (10).

Here we present the crystal structure of Δ4-(5α)-KSTD (Ro05698) from *R. jostii* RHA1 in its steroid-free and product-bound form. These first structures of a KSTD enzyme allowed the identification of the active site residues involved in the dehydrogenation of 3-ketosteroid substrates and its reaction mechanism, which was confirmed through mutational analyses.

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains and Growth Conditions**—Rhodococcus strains were cultivated in LBP medium containing 1% (w/v) bacto-peptone (Difco, Detroit, MI), 0.5% (w/v) yeast extract (BD Biosciences) and 1% (w/v) NaCl at 30 °C and 200 rpm. *E. coli* strains were grown in Luria-Bertani broth (Sigma) at 37 °C unless stated otherwise.

**KSTD Activity Staining**—Initial Δ4-(5α)-KSTD activity was established by a native gel-based assay. Cell-free extracts of three *Rhodococcus* strains (*Rhodococcus erythropolis* SQ1, *Rhodococcus rhodochrous*, and *R. jostii* RHA1) were prepared as described (5). These were loaded on native PAGE gels and stained for KSTD activity following published procedures (5).

**Protein Expression, Purification, and Characterization**—Δ4-(5α)-KSTD was heterologously expressed in *E. coli* strain BL21(DE3) and subsequently purified and crystallized as described before (10). UV-visible spectra of purified enzyme measured in the range of 200 to 700 nm (Cary 100, Varian) with and without substrate were used to characterize the protein as a flavoprotein and to detect the reduction of the FAD upon addition of substrate.

Noncovalent binding of the flavin co-factor was determined as follows. Purified Δ4-(5α)-KSTD was applied to a SDS-PAGE gel. Following electrophoresis, the gel was treated with 5% acetic acid solution, and flavin was visualized by 254-nm UV irradiation. Lack of co-migration indicated noncovalent binding of the flavin co-factor to the Δ4-(5α)-KSTD protein.

**Activity Assays and Product Identification**—Because *E. coli* does not code for any KSTD enzymes, cell-free extracts were used for product formation assays as described before (5). Briefly, enzyme activities were determined spectrophotometrically at 30 °C in 50 mM Tris-HCl buffer, pH 7.4, using 80 μM dichlorophenolindophenol (DCPIP) as an artificial electron acceptor and 0−200 μM 1-(5α)-androsterone-3,17-dione (1-(5α)-AD) as substrate (Steraloids). The initial reaction rates were determined from the absorbance change at 600 nm during the first 20 s of the reaction. The products were identified by HPLC following a reaction containing purified protein (50−200 μg), 200 μM 1-(5α)-AD dissolved in ethanol, and 200 μM DCPIP. No activity was detected in control reaction mixtures lacking steroids or purified enzyme or in extracts of *E. coli* with an empty pET15b expression vector.

**Crystallization, Data Collection, and Processing**—Δ4-(5α)-KSTD was crystallized at 293 K as described before, with a precipitant consisting of 200 mM NH$_4$-acetate, 100 mM sodium citrate, pH 5.6, and 30% (w/v) PEG 4000 (10). Crystals were prepared for data collection by soaking them for ~30 s in mother liquor, supplemented with 20% (w/v) glycerol, followed by cryo-cooling in liquid nitrogen. The 4-AD soaks were prepared by soaking crystals for ~18 h in cryo-mother liquor, which had the 200 mM NH$_4$-acetate replaced by 200 mM NH$_4$Cl, and to which 10 μl of a saturated solution of 4-AD in ethanol had been added per 500 μl of cryo-mother liquor.

The diffraction data were collected at 100 K at the beamlines of the European Synchrotron Radiation Facility (Grenoble, France). The intensity data were processed using the programs MOSFLM (11) and SCALA (12) from the CCP4 package (13). A summary of the data collection statistics is shown in Table 1.

**Phasing**—The FFAS03 server (14) was used for identifying suitable models for molecular replacement. The structure with the highest sequence identity (24%) was that of flavocytochrome c$_3$ fumarate reductase from *Shewanella frigidimarina* in the open conformation (Protein Data Bank entry 1qo8) (15). Because of the low sequence identity, the molecular replacement model was enhanced by incorporating all-serine structural information from various homologous proteins (17−24% identity; Protein Data Bank entries 1d4c (16), 1e39 (17), 1kf6 (18), 1nek (19), and 1zoy (20)). This ensemble of all-serine models was used to solve the structure of Δ4-(5α)-KSTD by molecular replacement at 2.5 Å resolution using the program Phaser (21). Several model building cycles, consisting of manual model building using COOT (22) and density modification with Resolve (23), were performed to improve the model obtained from Phaser until an R factor of 0.44%. Because of low sequence similarity and variable positions of residues 291−426 (substrate-binding domain) in the homologous structures, no inter-
**Table 1**

Data collection statistics

<table>
<thead>
<tr>
<th></th>
<th>Steroid-free</th>
<th>4-AD</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>European Synchrotron</strong></td>
<td>BM16</td>
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</tr>
<tr>
<td><strong>Radiation Facility</strong></td>
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</tr>
<tr>
<td><strong>Wavelength (Å)</strong></td>
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<td><strong>Space group</strong></td>
<td>C222</td>
<td>C222</td>
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<tr>
<td><strong>Unit cell parameters</strong></td>
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<td></td>
</tr>
<tr>
<td>a (Å)</td>
<td>99.2</td>
<td>99.8</td>
</tr>
<tr>
<td>b (Å)</td>
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<td>116.1</td>
</tr>
<tr>
<td>c (Å)</td>
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<td>110.2</td>
</tr>
<tr>
<td>α (°)</td>
<td>90</td>
<td>90</td>
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<td>β (°)</td>
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<td>90</td>
</tr>
<tr>
<td>γ (°)</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td><strong>Resolution (Å)</strong></td>
<td>1.6 (1.69–1.60)</td>
<td>1.6 (1.69–1.60)</td>
</tr>
<tr>
<td>R пп,рп</td>
<td>0.081 (0.319)</td>
<td>0.110 (0.373)</td>
</tr>
<tr>
<td>R пп,лп</td>
<td>0.023 (0.138)</td>
<td>0.055 (0.201)</td>
</tr>
<tr>
<td><strong>Total number of observations</strong></td>
<td>888,741 (80,282)</td>
<td>80,072 (8,955)</td>
</tr>
<tr>
<td><strong>Total number of unique reflections</strong></td>
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<td><strong>Mean I/σ(I)</strong></td>
<td>22.6 (4.7)</td>
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<tr>
<td><strong>Completeness (%)</strong></td>
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<td>95.2 (74.1)</td>
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<tr>
<td><strong>Multiplicity</strong></td>
<td>11.9 (6.1)</td>
<td>4.9 (4.3)</td>
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</tbody>
</table>

* Data collection statistics according to van Oosterwijk et al. (10).

**Table 2**

Refinement statistics

<table>
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<tr>
<th></th>
<th>Steroid-free</th>
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</thead>
<tbody>
<tr>
<td>Resolution (Å)</td>
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<td>1.6</td>
</tr>
<tr>
<td>Average B-factor</td>
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<td>R пп</td>
<td>15.8</td>
<td>15.9</td>
</tr>
<tr>
<td>R пп,рп</td>
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<td>18.1</td>
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<tr>
<td>RMSD from target geometry</td>
<td>Bond lengths (Å)</td>
<td>0.012 (0.007)</td>
</tr>
<tr>
<td>Bond angles (°)</td>
<td>1.44</td>
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</tr>
<tr>
<td>Total number of atoms</td>
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<td>4353</td>
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<tr>
<td>Number of amino acids</td>
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<td>482</td>
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<tr>
<td>Number of Cl ions</td>
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<td>1</td>
</tr>
<tr>
<td>Number of acetate molecules</td>
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<td></td>
</tr>
<tr>
<td>Number of glycerc molecules</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Active site ligand</td>
<td>4-AD</td>
<td></td>
</tr>
<tr>
<td>Number of FAD molecules</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Number of water molecules</td>
<td>609</td>
<td>618</td>
</tr>
<tr>
<td>Ramachandran angles (%)</td>
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<td>95.5</td>
</tr>
<tr>
<td>Favored</td>
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<td>4.1</td>
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<tr>
<td>Allowed</td>
<td>0.8</td>
<td>0.4</td>
</tr>
<tr>
<td>Disallowed</td>
<td></td>
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</table>

*Ramachandran statistics were obtained from COOT (22).**

### Site-directed Mutagenesis

Site-directed mutagenesis was performed using the QuikChange™ site-directed mutagenesis kit (Stratagene) with plasmid pBluescript-ro05698 as a template. The PCR product of ro05698 (10) was cloned into the EcoRV site of pBluescript (II) KS (Fermentas). Primers used for site-directed mutagenesis are listed in supplemental Table S1. Successful mutagenesis was confirmed by DNA sequencing. The resulting mutant genes were subcloned following Ndel/BamHI digestion into the Ndel/BamHI-digested pET15b (Novagen) for protein expression and protein purification. Mutation S468A in Ro05698 was obtained by ordering a synthetic DNA fragment (487 bp, AGOWA) of ro05698 containing the desired mutation. The synthetic fragment was digested with NotI/BamHI and subcloned into the NotI/BamHI-digested pET15b-Ro05698 construct to obtain the full-length gene with the desired mutation.

### RESULTS AND DISCUSSION

**Ro05698 from R. jostii RHA1 Is a Δ4-(5α)-KSTD Enzyme**

The genome of R. jostii RHA1 contains at least three probable Δ1-KSTD enzymes but only one single putative Δ4-(5α)-KSTD enzyme, encoded by ro05698, based on its 31% sequence identity to the Δ4-(5α)-KSTD of R. erythropolis (1, 5). Indeed, cell-free extracts of R. jostii RHA1 and two other *Rhodococcus* strains showed Δ4-(5α)-KSTD activity in a native PAGE assay with 1-(5α)-AD as substrate (supplemental Fig. S1). In *R. jostii* RHA1, this activity presumably originates from the product of the ro05698 gene.

To firmly establish that ro05698 codes for an enzyme with Δ4-(5α)-KSTD activity, the gene was expressed in *E. coli*, and the Ro05698 protein was purified and incubated with the steroid substrate 1-(5α)-AD. HPLC analysis of the product confirmed the formation of 1,4-androstanediene-3,17-dione (ADD) (Fig. 2), indicating that ro05698 indeed codes for a Δ4-(5α)-KSTD. As expected, no activity was observed with the product 4-androstene-3,17-dione (4-AD) as substrate.

The absorption spectrum of purified Δ4-(5α)-KSTD protein is typical of that of a flavoprotein (31), with maxima at 461 and
3-Ketosteroid-$\Delta^4$-(5$\alpha$)-dehydrogenase Reaction Mechanism

The Three-dimensional Structure of $\Delta^4$-(5$\alpha$)-KSTD—The crystal structure of $\Delta^4$-(5$\alpha$)-KSTD—The crystal structure of $\Delta^4$-(5$\alpha$)-KSTD was elucidated by molecular replacement using an ensemble of six structures with 17–24% sequence identity to $\Delta^4$-(5$\alpha$)-KSTD as the starting model, and the resulting solution was refined to an $R$ factor of 15.8% at 1.6 Å resolution (for details see Table 2). The crystals contain one $\Delta^4$-(5$\alpha$)-KSTD molecule per asymmetric unit. $\Delta^4$-(5$\alpha$)-KSTD has an ellipsoid shape with dimensions of $80 \times 40 \times 35$ Å (Fig. 4). The overall fold resembles that of $p$-hydroxybenzoate hydroxylase (33), a fold often observed in flavoenzymes (34).

The protein consists of two domains connected via a two-stranded antiparallel $\beta$-sheet (Fig. 4). One domain (F) contains a noncovalently bound FAD co-factor; it consists of a central five-stranded parallel $\beta$-sheet flanked on one side by four additional $\beta$-strands and on the other side by three $\alpha$-helices. The other domain, or substrate-binding domain (S), residues 292–489) is an insert into domain F and divides it into two parts, F$_1$ (residues 7–291) and F$_2$ (residues 426–489). It consists of a four-stranded antiparallel $\beta$-sheet flanked on both sides by two $\alpha$-helices. The active site is located at the interface between the F and S domains next to the isoalloxazine ring system of the FAD co-factor.

The $\Delta^4$-(5$\alpha$)-KSTD structure is well defined, with the exception of the His$_6$ tag, the 19 residues of the linker, residues 1–6, and the C-terminal residue 490. In addition, two loops near the active site (residues 170–182 and 256–268; supplemental Fig. S3) are poorly visible in the electron density map, which may indicate flexibility. These loops may have a function in catalysis by shielding the FAD and substrate from the solvent (see below).

Comparison with Structurally Similar Proteins—A DALI (Distance-matrix ALIgnment) search (35) revealed that $\Delta^4$-(5$\alpha$)-KSTD is structurally most closely related to cytochrome $c_1$ fumarate reductase from S. frigidimarina (Z = 42.6, 27% sequence identity), cytochrome $c$ fumarate reductase from Shewanella putrefaciens MR-1 (Z = 42.2, 28% identical residues), quinol-fumarate reductase from E. coli (Z = 32.6, 22% identical residues), and L-aspartate oxidase from E. coli (Z = 32.5, 21% sequence identity), which are all FAD-binding proteins with a $p$-hydroxybenzoate hydroxylase-like fold. $\Delta^4$-(5$\alpha$)-KSTD has less, but still significant structural similarity to cholesterol oxidase from Brevibacterium sterolicum (1coy, Z = 16.2, 12% sequence identity), an enzyme that catalyzes a similar reaction on a similar steroid substrate as $\Delta^4$-(5$\alpha$)-KSTD.

The highest structural similarity is found for the F domain with root mean square differences (RMSDs) on C$_\alpha$ atoms of 1.6–2.0 Å. The S-domain is less similar in structure, and only the fumarate reductases of S. frigidimarina and S. putrefaciens, two highly homologous proteins (59% sequence identity to each other), have a similar S domain with RMSDs of 1.9–2.0 Å (116 residues of 135). Nevertheless, although the overall fold of all these proteins is highly similar and all of them display a similar FAD binding mode, their catalytic centers show a huge variation, consistent with the large variation of their substrates, ranging from small compounds (e.g., fumarate) to sugars and steroids.

FAD Binding—The FAD molecule binds noncovalently in a long cleft in the F-domain (supplemental Fig. S2). Its ADP part is bound by a motif resembling the $\beta$\beta$\beta$\beta$$ dinucleotide binding motif (Rossman fold), frequently observed in flavoproteins (36). The adenine moiety is bound by two hydrogen bonds to the backbone amide and carbonyl group of Val-205. The two hydroxyl groups of the adenosine ribose are hydrogen-bonded to Glu-51. The two phosphate groups have hydrogen bonding interactions with the backbone amides of Ile-31, Ala-32, Ala-59, Thr-60, and Arg-456 and with the hydroxyl group of Thr-60. In addition there is a salt bridge between one of the phosphates and His-268. The D-ribitol binds in an extended conformation and is hydrogen-bonded to several water molecules and to the side chains of Ser-471 and Thr-60.

The isoalloxazine ring system is located close to the interface of the F and S domains. Its N5 nitrogen atom is hydrogen-bonded to the backbone amide of Gly-64, and the main chain NHs of Ser-471 and Leu-472 have hydrogen bonding interactions with the O$_2$ oxygen of the pyrimidine ring. Ser-471 and Leu-472 are at the N-terminal end of the $\sim$25 Å long C-terminal $\alpha$-helix formed by residues 470–489, and the dipole moment provided by this helix may stabilize the negative charge on the O$_2$ oxygen atom in the anionic hydroquinone form of the FAD (37). Finally, the side chain hydroxyl group of Ser-471 points away from the N1 nitrogen of the flavin, but in a different rotamer conformation it would be at hydrogen bonding distance to the N1 nitrogen and could be involved in prototating it.
The isoalloxazine ring system is not planar; the butterfly angle between the pyrimidine and dimethylbenzene rings is 168°, similar to that observed in cholesterol oxidase (38). The color of the crystals and the spectrum of the protein sample used for crystallization indicate that the oxidized form of FAD is present in the structure.

**The Active Site**—In the steroid-free Δ4-(5α)-KSTD structure, the active site is open to the solvent and contains a well ordered network of water molecules and two acetate ions. A chloride ion is located 3.8 Å behind the dimethylbenzene ring of the isoalloxazine ring system (Fig. 5 and supplemental Fig. S2). It is bound in a partially hydrophobic environment (Phe-427 and the dimethylbenzene ring), and it is held in place by the backbone nitrogen of Ala-264. A negative charge close to the dimethylbenzene ring of the FAD may push electrons toward the pyrimidine ring and thereby lower the redox potential (39).

**Product Binding**—The 1.6 Å resolution structure of Δ4-(5α)-KSTD with bound 4-androstene-3,17-dione (4-AD; Table 2) shows that 4-AD binds at the si face of the isoalloxazine ring system, displacing three water molecules and the two acetate molecules (Fig. 5). Binding of 4-AD does not cause significant changes in the overall protein structure; the all-atom RMSD between the structures is only 0.31 Å. The side chains of Trp-136 and Ser-320, which display a correlated double conformation in the steroid-free structure, become ordered upon binding of the steroid molecule. Trp-136 has a hydrophobic stacking interaction with the steroid, but it is conserved in only a few putative Δ4-KSTDs (5). A W136F substitution caused only a 3-fold reduction in apparent activity, but a W136A mutation resulted in inactive enzyme (Table 3), indicating that an aromatic residue capable of stacking is required at this position. Concomitantly with the rigidification of the Trp-136 side chain, also the Ser-320 side chain becomes ordered.

4-AD binds with its C3 keto group at the same location as one of the acetate oxygen atoms in the steroid-free structure, forming a hydrogen bond to Tyr-466 (Fig. 5). Acetate is more often observed to mimic the binding of an oxygen atom of the substrate (40). The C4 carbon atom of 4-AD is near the Ser-468 hydroxyl group. Surprisingly, the C17 keto moiety of 4-AD has...
no interactions with the protein, but it is hydrogen-bonded to two water molecules. Apparently, the substituent at this position is not a very critical factor in substrate recognition, although preliminary results show that an aliphatic tail at C17, as in 5α-cholestan-3-one, is not accepted at this position.

Tyr-466 Is the Putative Catalytic Base—The position of Tyr-466 and its hydrogen bonding interaction with the C3 keto group of 4-AD suggest that this residue may function as the catalytic acid that protonates the C3 keto group, promoting keto-enol tautomerization and labilization of the C4 hydrogen atoms. Formation of the enolate at the C3 keto group has previously been suggested as the initial catalytic step for these proteins (41, 42). Tyr-466 is conserved in all Δ4- and Δ1-KSTD enzymes. Site-directed mutagenesis (Y466A and Y466F) resulted in catalytically dead Δ4-(5α)-KSTD enzymes (Table 3), confirming the importance of Tyr-466 and showing that the hydroxyl group of this tyrosine for activity (Table 3). In contrast, a E290Q mutation did not affect the catalytic activity of the enzyme, indicating that this residue is not essential for proton relay and that other relay routes may exist.

Ser-468 Is the Putative Catalytic Base—The Ser-468 hydroxyl group is at 3.1 Å from the C4 atom of 4-AD, a position compatible with a role of Ser-468 as the catalytic base, which abstracts a proton from C4. The C4 protons have been labilized because of keto-enol tautomerization of the keto group of the substrate upon interaction with the Tyr-466 hydroxyl group. The Ser-468 hydroxyl group is part of an extensive hydrogen bonding system, which can function as an effective system relaying protons in a concerted way to the solvent and which includes the Tyr-319 hydroxyl group, a water molecule, and the side chain of Glu-290 (Fig. 6). Because such concerted proton transfers do not involve high energy alkoxide intermediates (43), the proton can easily be transferred from the C4 atom to the solvent.

Ser-468 is conserved in Δ4-KSTD enzymes, including Rs1817 from M. tuberculosis H37Rv, but not in Δ1-KSTDs. Its hydroxyl group is essential for activity, because a S468A mutation abolished all catalytic activity, whereas a S468T mutant still retained activity. However, the apparent $K_{\text{m}}$ of this latter mutant protein was drastically increased, and the apparent $V_{\text{max}}$ was somewhat lower than that of wild-type enzyme (Table 3).

A Y319F mutant enzyme had also lost all activity on 1-(5α)-AD as substrate, showing the functional importance of the hydroxyl group of this tyrosine for activity (Table 3). In contrast, a E290Q mutation did not affect the catalytic activity of the enzyme, indicating that this residue is not essential for proton relay and that other relay routes may exist.

The Role of the FAD Co-factor in Catalysis—Flavoproteins involved in dehydrogenation reactions often display a few recurrent features, like the distance and angle between the FAD N5 nitrogen atom and the site of oxidative attack and the pres-
ence of a hydrogen bond donor close to the N5 nitrogen (44). In Δ4-(5α)-KSTD the N5 nitrogen is hydrogen-bonded to the backbone amide of Gly-64. It is at a distance of 3.9 Å from the C5 atom of the product, and the N10-N5-C5 angle is 108°. Both the distance and angle are in agreement with the values observed in other proteins (44). These values clearly suggest that the site of attack by the FAD is the C5 atom of the substrate. The position of the FAD N5 atom above and of Ser-468 below the 4-AD is in agreement with trans-dehydrogenation of the substrate (42).

In many flavoproteins, catalysis takes place at a location shielded from the solvent, which enhances the strength of polar interactions and is instrumental to substrate activation (34, 44). In some enzymes with the p-hydroxybenzoate hydroxylase fold loops move in upon substrate binding to cover the active site, and in others the complete S domain may relocate (15, 45). In Δ4-(5α)-KSTD two loops close to the active site (supplemental Fig. S3) show somewhat less defined electron density and may be flexible. However, in the 4AD-bound structure, no significant conformational differences are seen between the steroid-free and product-bound states. At present we cannot exclude that crystal contacts prevent large conformational changes or that the loops move to restrict solvent access to the FAD. Binding of the steroid substrate may also already provide sufficient shielding. Further research will be needed to resolve this question.

Comparison with Cholesterol Oxidase—Δ4-(5α)-KSTD is functionally and structurally related to cholesterol oxidase (45). The structures of the two proteins can be superimposed with an RMSD of 2.6 Å for 213 of 483 Ca atom positions with the highest similarity in the F-domain. Although Δ4-(5α)-KSTD catalyzes the dehydrogenation of the C4-C5 bond of steroids, cholesterol oxidase catalyzes the oxidation of steroids containing a 3β-hydroxyl group with concomitant isomerization of the Δ5 double bond to the Δ4 position. However, whereas in Δ4-(5α)-KSTD the substrate/product binds at the si face of the isoalloxazine ring system, in cholesterol oxidase the substrate binds in a pocket in front of the N5 nitrogen. This different substrate-binding mode positions the substrate such that in Δ4-(5α)-KSTD the C5 hydrogen atom is near the FAD N5 nitrogen, but in cholesterol oxidase it is the C3 hydroxyl group that is near the N5 atom of the FAD. This explains the different reaction specificity and regioselectivity of the two enzymes.

Cholesterol oxidase also contains an extra domain, which binds the apolar aliphatic tail connected to the C17 atom of the cholesterol. In the 4AD-bound structure of Δ4-(5α)-KSTD, the C17 keto group of the substrate/product is exposed to the solvent, in agreement with the wider range of substituents at this position that are accepted by Δ4-(5α)-KSTD.

The Catalytic Mechanism—The oxidation of 5α-AD or 1-(5α)-AD requires the abstraction of two hydrogen atoms and the transfer of two electrons as a hydride ion to the FAD. It is not known whether these proton and hydride transfers occur in a concerted or stepwise manner. The two electrons will finally be transferred to a currently unknown electron acceptor, for example menaquinone (46).

The crystal structure indicates Ser-468 as the putative base that abstracts the C4 α-hydrogen proton from the substrate and relays its own proton to the solvent via Tyr-139 (Fig. 7). The resulting deprotonated state of the substrate can be stabilized by the delocalization of the negative charge over the C3 keto group. In addition, Tyr-466, functioning as an acid, can stabilize the ensuing enolate by hydrogen bonding to the C3 oxygen.

FIGURE 7. Proposed reaction mechanism of Δ4-(5α)-KSTD. Panel 1, catalysis is initiated by the interaction of the O3 keto group of the substrate with the hydroxyl group of Tyr-466, which promotes keto-enol tautomerization and stabilization of the C3 keto group. Ser-468, acting as a base, abstracts a proton from the C4 atom of the substrate, with Tyr-319 serving as a proton relay system to the solvent. Panel 2, next, the FAD abstracts a hydride ion from the C5 carbon of the substrate, which, with a concomitant rearrangement, results in the formation of a double bond between C4 and C5. Panel 3, the product is formed and leaves the active site, perhaps only after the oxidizing substrate enters the active site. The negative charge on the reduced FAD is stabilized by the dipole moment of the C-terminal helix. Finally, the oxidized FAD is regenerated by an as yet unknown electron acceptor.
atom. The FAD N5 atom is in a good position to abstract a hydride ion from the C5 atom of the enolate intermediate. In synchrony, the lone pair electrons of the negatively charged C3 oxygen atom move back toward C3 and the double bond between C3 and C4 shifts to the C4-C5 position, generating the product. The negative charge on the N5 of FAD can be delocalized over the N1-C = O2 region, but also the rest of the isoaallo-
za
inge ring may contribute (47). Two backbone amides stabilize the negative charge on O2 by hydrogen bonding, and the dipole moment of the C-terminal helix is also directed toward O2. The generated reducing equivalents may be donated to a respiratory chain, similar to what has been proposed for the Δ1-KSTD of Arthrobacter globiformis (48), but further studies are required to confirm this.

The Importance of Ketosteroid Dehydrogenases for Pathogenic Organisms—Δ4-(5α)-KSTD is the first ketosteroid dehydrogenase of which a crystal structure has been elucidated. The enzyme has an important role in the desaturation of the steroid A ring, which is a key step in the microbial degradation of saturated steroids. It has been proposed that saturated steroid intermediates are formed during cholesterol catabolism. Several pathogenic bacteria, including M. tuberculosis, Rhodococcus equi, and Mycobacterium bovis, contain a cholesterol catabolic pathway similar to that of R. jostii RHA1 (8, 49). In M. tuberculosis this pathway and in particular the Δ1-KSTD enzyme (Rv3537, which shows 28% sequence identity to R. jostii RHA1 Δ4-(5α)-KSTD) have been implicated to be important for growth of the intracellular pathogen in the hostile environment of macrophages (50, 51). This makes the cholesterol catabolic pathway a promising target for the development of therapeutic agents to combat M. tuberculosis (8). Our R. jostii RHA1 Δ4-(5α)-KSTD structure may facilitate the design of potent ketosteroid dehydrogenase inhibitors, as a first step toward the development of new antituberculosis drugs.

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