Universal capability of 3-ketosteroid Δ¹-dehydrogenases to catalyze Δ¹-dehydrogenation of C17- substituted steroids

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Abstract

Background: 3-Ketosteroid Δ1-dehydrogenases (KSTDs) are the enzymes involved in microbial cholesterol degradation and modification of steroids. They catalyze dehydrogenation between C1 and C2 atoms in ring A of the polycyclic structure of 3-ketosteroids. KSTDs substrate spectrum is broad, even though most of them prefer steroids with small substituents at the C17 atom. The investigation of the KSTD’s substrate specificity is hindered by the poor solubility of the hydrophobic steroids in aqueous solutions. In this paper, we used 2-hydroxypropyl-β-cyclodextrin (HBC) as a solubilizing agent in a study of the KSTDs steady-state kinetics and demonstrated that substrate bioavailability has a pivotal impact on enzyme specificity.

Results: Molecular dynamics simulations on KSTD1 from Rhodococcus erythropolis indicated no difference in ΔG_{bind} between the native substrate, androst-4-en-3,17-dione (AD; −8.02 kcal/mol), and more complex steroids such as cholest-4-en-3-one (−8.40 kcal/mol) or diosgenone (−6.17 kcal/mol). No structural obstacle for binding of the extended substrates was also observed. Following this observation, our kinetic studies conducted in the presence of HBC confirmed KSTD1 activity towards both types of steroids. We have compared the substrate specificity of KSTD1 to the other enzyme known for its activity with cholest-4-en-3-one, KSTD from Sterolibacterium denitrificans (AcmB). The addition of solubilizing agent caused AcmB to exhibit a higher affinity to cholest-4-en-3-one (Ping-Pong bi bi K_m_A = 23.7 μM) than to AD (K_m_A= 529.2 μM), a supposedly native substrate of the enzyme. Moreover, we have isolated AcmB isoenzyme (AcmB2) and showed that conversion of AD and cholest-4-en-3-one proceeds at a similar rate. We demonstrated also that the apparent specificity constant of AcmB for cholest-4-en-3-one (k_cat/K_m_A= 9.25 · 10^6 M^{-1} s^{-1}) is almost 20 times higher than measured for KSTD1 (k_cat/K_m_A= 4.71 · 10^5 M^{-1} s^{-1}).
Conclusions: We confirmed the existence of AcmB preference for a substrate with an undegraded isooctyl chain. However, we showed that KSTD1 which was reported to be inactive with such substrates can catalyze the reaction if the solubility problem is addressed.

Keywords: 3-ketosteroid dehydrogenase, KSTD, cholest-4-en-3-one \( \Delta^1 \)-dehydrogenase, 3-ketosteroids, 1,2-dehydrogenation, \( \Delta^1 \)-dehydrogenation, cholest-4-en-3-one, diosgenone, cholesterol metabolism
Background

Steroids are widespread compounds in the natural environment. They are essential components of eukaryotic (animals, yeast and fungi) and plant cells. Cholesterol, the parent substance of all sterols, is one of the most commonly occurring. Hence bacteria have evolved to use it as a carbon and energy source [1]. The microbial degradation of cholesterol proceeds under either aerobic or anaerobic conditions and the current stage of knowledge was recently reviewed by Rohman et al. [2]. 3-Ketosteroid Δ¹-dehydrogenases (KSTDs) initiate the crucial step of the degradation – steroid nucleus decomposition and as a result are found in anaerobic, facultative anaerobic and strictly anaerobic microorganisms [3–5]. KSTDs catalyze regio- and stereoselective dehydrogenation between C1 and C2 atoms of steroid ring A (Fig. 1) [6]. The stage at which the double bond is introduced during cholesterol degradation is still debated and may differ between bacterial species [2].

The wide KSTDs substrate spectrum, the regioselectivity of the catalyzed reaction and mild conditions of the synthesis make those enzymes widely applied for the biologically active Δ¹-dehydrosteroids production [2]. The 1,2-dehydrogenation process can be carried out with either a whole-cell system (native and recombinant bacteria are used) or isolated enzymes [7, 8]. The poor solubility of steroids in aqueous solutions is usually circumvented by the application of pseudo-crystallofermentation technique (i.e., steroid substrate in the solid-state) or by the addition of organic solvents and various solubilizing agents, e.g., vegetable oils, phospholipids, glycols, dextrans or cyclodextrins [8–14]. The biocompatibility of cyclodextrins makes them valuable solubilizing agents for scale-up microbial steroid biotransformation. Importantly, they do not exert a detrimental influence on the stability of the purified enzymes [15, 16].

Although KSTDs substrate scope is generally broad, most of the enzymes prefer steroids with small substituents on the C17 position [2]. Only a few of them exhibited activity towards more branched substrates, such as cholest-4-en-3-one, 3-oxo-5β-cholan-24-oic acid, the methyl ester
of 3-oxo-4-cholen-24-oic acid, cortisone acetate, hydrocortisone acetate or 21-acetoxy-pregna-4,9(11),16-triene-3,20-dione [17–22]. The comparative study of substrate specificity is additionally hindered by poor or very bad solubility of steroids in the water phase in which all kinetic enzyme assays are conducted [8].

3-Ketosteroid Δ1-dehydrogenase from *Streolibacterium denitrificans* (AcmB) is a good example of the enzyme for which the native substrate remains an open question [23]. So far two 3-ketosteroids have been proposed as the AcmB natural substrates. Cholest-4-en-3-one (2 in Figure 1) was initially proposed as the AcmB native substrate [20,24,25]. However, subsequent research demonstrated a high affinity of the enzyme for C19–C21 steroids and indicated that AcmB is located on the cytoplasmic side of the inner membranes of *S. denitrificans* [8,26]. These results suggested that androst-4-en-3,17-dione (AD; 1) may be the enzyme native substrate [27]. Nevertheless, AcmB’s ability to catalyze Δ1-dehydrogenation of steroids with the aliphatic side chain on the C17 position seemed interesting and quite unusual.

In this work, we demonstrate that substrate bioavailability can significantly influence the kinetic results and lead to misinterpretation of enzyme specificity. We used 2-hydroxpropyl-β-cyclodextrin (HBC) as a solubilizing agent to study the steady-state kinetics for AcmB and 3-ketosteroid Δ1-dehydrogenase from *Rhodococcus erythropolis* SQ1 (KSTD1) with AD and cholest-4-en-3-one. In contrast to previous reports [2,28], we have demonstrated never reported before KSTD1 activity towards cholest-4-en-3-one and even more complex 3-ketosaponin – diosgenone (3). Moreover, we proved that AcmB indeed exhibits a significantly higher affinity to cholest-4-en-3-one than to AD while KSTD1 indeed prefers smaller ketosteroids.
**Fig. 1** Structures of compounds tested in this study as substrates: androst-4-en-3,17-dione, (2) cholest-4-en-3-one, (3) diosgenone, (4) androstanolone, (5) progesterone, (6) testosterone propionate and (7) 6-dehydrotestosterone acetate.

**Methods**

**Materials**

All chemicals were purchased from Sigma-Aldrich, Tokyo Chemical Industry, Carl Roth or BioShop unless otherwise specified. (25R)-Spirost-1,4-dien-3-one (diosgenone) was a gift of Jacek Morzycki from the University of Bialystok.

**Proteins expression and purification**

The AcmB was cloned, expressed and purified as described previously at Wojtkiewicz et al. work [8].

A gene acmB2 was amplified from genomic *S. denitrificans* DNA using primers: forward (TACTTCCAATCAATGCCTATGAGCGGAGAAACTTTTCG) and reverse (TTATCCACTTCAAATTCATACCGCGCTCCG), Platinum™ Taq DNA Polymerase High Fidelity (ThermoFisher), 2.4% dimethyl sulfoxide (DMSO) and thermocycler (BioRad) program as follows: 94°C for 2 min, 32 cycles of (94°C for 15 s, 63°C for 30 s, 68°C for 120 s) with a final elongation at 68°C for 10 min. After gel electrophoresis a band of size ~1.7 kbp
was excised from agarose gel and purified using Gel-Out (A&A Biotechnology). Next, gene encoding AcmB2 (GenBank: SMB21450.1) was cloned into a pMCSG7 vector. 25 μL pMCSG7 vector (200 ng) was treated with 2.5 μL SspI in a total of 50 μL 1x Tango Buffer (Thermo Scientific) supplemented with 4 mM dithiothreitol (DTT) for 1.5 h at 37°C. The linearized, gel-purified vector was prepared for ligase-independent cloning. Namely, 200 ng of pMCSG7 vector were incubated in 40 μL 1x T4 Polymerase buffer (Novagen) supplemented with 4 mM DTT, 1 μL of T4 Polymerase (Novagen) and dGTP (Promega) for 40 min at room temperature, which was followed by a 10 min inactivation of the polymerase at 75°C. Similarly, the acmB2 gene was prepared, however, the nucleotides were substituted with dCTP (Promega). Then 3.5 μL of vector preparation was reacted with 6 μL of gene preparation and incubated on ice for 3 h. The mixture was transformed into Escherichia coli XL10-Gold Ultracompetent Cells (Agilent Technologies) and the cells were grown overnight in 2% (w/v) Lennox Broth (LB) supplemented with ampicillin 100 μg mL⁻¹ and 34 μg mL⁻¹ chloramphenicol. Afterward, the plasmid was extracted using Plasmid Miniprep DNA Purification Kit (EURx), sequenced and transformed into calcium chloride chemically competent E. coli BL21(DE3)Magic (Creative Biolabs). Bacteria cultivation, harvesting and disruption, as well as enzyme purification were conducted as described for AcmB [8].

A gene encoding 3-ketosteroid Δ¹-dehydrogenase from R. erythropolis (KSTD1) was cloned into pET15b vector [29] and used to transform E. coli BL21(DE3)Magic cells. An overnight culture of the transformed cells was grown in 2% (w/v) LB supplemented with 100 μg mL⁻¹ ampicillin and 50 μg mL⁻¹ kanamycin at 37°C, 180 rpm. The preculture was hundred times diluted in 1 L of fresh LB containing 0.5 M D-sorbitol and antibiotics and grown at 37°C, 180 rpm. When the OD₆₅₀ reached 0.6, the temperature was reduced to 16°C and the culture was induced with 100 μM isopropyl β-D-1-thiogalactopyranoside (IPTG). After 48 h the E. coli cells were centrifuged for 1 h at 4500 g, 4°C. The harvested cells were resuspended in a 1:5
ratio (w/v) in the 50 mM Tris-HCl pH 8.5 buffer, 100 mM NaCl, 10% (w/v) glycerol, 5 mM β-mercaptoethanol (BME) and 10 mM imidazole and supplemented with 100 μM phenylmethylsulfonyl fluoride (PMSF). The cells were disrupted by sonication (Sonics Vibra-Cell VCX500, 3 s on, 5 s off, 5 min, 40% amplitude, 150 000 J). The cell-free extract was obtained by centrifugation at 40 000 g, for 1 h at 4°C. Purification of KSTD1 was carried out on 5 mL HisTrap HP (GE Healthcare) column using FPLC system (BioRad NGC Quest 10 Plus) and linear imidazole gradient 10 – 300 mM in the previously described buffer. The fractions that absorbed UV-light at wavelength 450 nm were collected, desalted by dialysis and stored at –20°C.

The concentration of purified AcmB, AcmB2 and KSTD1 was determined using a free FAD extinction coefficient of 11 300 M⁻¹ cm⁻¹ at 450 nm.

**Spectrophotometric activity assay**

The steady-state kinetic parameters (Kₘ, Vₘₐₓ, kₙₐₜ, kₐₜₐₜ/Kₘ) for steroids were determined in a spectrophotometric activity assay using UV-2700 spectrophotometer (Shimadzu) in 0.5 mL quartz cuvettes with 10 mm path length. The measurements were carried out in 0.1 M K₂HPO₄/KH₂PO₄ buffer pH 6.5 (AcmB) or 0.1 M K₂HPO₄/KH₂PO₄ buffer pH 8.0 (KSTD1) with 0.029 – 0.174 mM 2,6-dichloroindophenol (DCPIP), 2% (w/v) 2-hydroxypropyl-β-cyclodextrin (HBC; 14.54 mM), 5 to 500 μM concentrations of the steroids dissolved in 2-methoxyethanol (EGME, 2% (v/v)) and 0.30 μM of AcmB or 0.23 μM of KSTD1, respectively.

The reduction of DCPIP was followed at 700 nm (εₚₜ₇₀ nm, HBC = 10 691 M⁻¹ cm⁻¹ or εₚₜ₇₀ nm, HBC = 11 627 M⁻¹ cm⁻¹). All measurements were performed in triplicates at 30°C. The linear slopes were obtained with linear regression fitted to the initial parts (5 s) of the kinetic curves. The ping-pong bi bi (non-sequential) mechanism model was fitted to the obtained data for AcmB.

The linear function was fitted to the data obtained for KSTD1. All data processing was conducted using OriginPro 2019 software.
**HPLC analysis**

To confirm KSTD1 activity with diosgenone mini reactor tests were performed. The reaction mixture (1 mL) consisted of 50 mM Tris-HCl buffer pH 8.0, 0.15 mM DCPIP, 2% (w/v) HBC, 0.1 mM diosgenone in dioxane (the final concentration 2% (v/v)) and 87.7 nM KSTD1. The reaction was carried out in duplicate at 30°C, in the thermoblock at 800 rpm for 20 min. To determine the conversion of the substrate, the reaction mixtures were diluted 1:1 with acetonitrile (ACN), centrifuged at 15 000 g for 5 min and analyzed with HPLC-DAD (Agilent 1100) according to [8].

To confirm AcmB2 activity with AD, cholest-4-en-3-one and diosgenone mini reactor tests were performed. The reaction mixture (1 mL) consisted of 50 mM Tris-HCl buffer pH 8.0, 0.15 mM DCPIP, 2% (w/v) HBC, 0.1 mM substrate in EGME (the final concentration 2% (v/v)) and 6.55 nM AcmB2 for AD and cholest-4-en-3-one or 65.5 nM for diosgenone. The reaction was carried out in duplicate at 30°C, in the thermoblock at 800 rpm for 960 h. To determine the conversion of the substrates, the samples were prepared and analyzed as described above.

**Determination of HBC/steroid stability constant**

The stability constants (Ks) of the inclusion complex between HBC and steroids were determined according to the modified protocol described by Ma et al [30]. The mixtures of various concentrations of HBC (from 0 to 98 mM), 2% (v/v) EGME, and excess of AD or cholest-4-en-3-one were incubated in the thermoblock at 30°C, 1000 rpm for 48 h. Afterward, the suspensions were filtered through 0.45 μm membrane filters, diluted 0 -200 times with ACN, centrifuged at 15 000 g for 15 min, and concentration of soluble steroid was quantified by HPLC-DAD using calibration on external standards [8]. The linear function was fitted to the data obtained for AD. The stability constant for AD:HBC 1:1 complexes (K_{1:1}) was calculated from equation (1), where S₀ is an y-intercept [31,32].
\[ K_{1:1} = \frac{\text{slope}}{S_0(1 - \text{slope})} \]  

The quadratic function \([S] = a[HBC]^2 + b[HBC] + S_0\) was fitted to the data attained for cholest-4-en-3-one. The stability constants for cholest-4-en-3-one, HBC 1:1 \((K_{1:1})\) and 1:2 \((K_{1:2})\) complexes, were calculated from the equation (2) and (3), respectively [32].

\[ K_{1:1} = \frac{b}{S_0} \]  

\[ K_{1:2} = \frac{a}{S_0 K_{1:1}} \]  

**MD simulations**

The structure of KSTD1 was taken from Protein Data Bank (PDB ID: 4c3y [33], resolution 2.3 Å). 1,4-androstadiene-3,17-dione (ANB) and FAD cofactor are already present in the active site of the crystal structure. KSTD1 was determined to be monomeric in solution [33] and, thus, we used the monomeric structure of the enzyme to perform MD simulations. To place structures of androst-4-en-3,17-dione (1), cholest-4-en-3-one (2) and diosgenone (3), androstanolone, progesterone, testosterone propionate and 6-dehydrotestosterone acetate in the enzyme active site, the Kabsch method was used [34] with ANB as a template. The geometry of each substrate was obtained by optimization in the gas phase at B3LYP/6-31G(d,p) level of theory with Gaussian16 [35]. PropKa ver 3.1 [36,37] and H++ packages [38] were used to determine the protonation states of titratable amino acids. The optimal pH for enzyme activity was reported to be about 8.0, so this value was selected for model construction. We assumed that Tyr318 is deprotonated due to the mechanistic hypothesis. AMBER parameters for ketosteroids and tyrosyl anion were obtained with Gaussian 16 software package [35] (at B3LYP/6-31G(d,p) level of theory) and ANTECHAMBER program from AmberTools [39]. Parameters for FAD were obtained from the RESP ESP charge DataBase (R.E.DD.B) [40]. A total amount of 33 sodium cations were added to neutralize the system. Finally, the system was soaked in a 93.5 × 76.5 × 72.0 Å³ box with TIP3P [41] water molecules.
To prepare the model for MD simulation, the system was optimized with Amber package [42] with ff03 force field [43], heated from 0 to 303 K over 100 ps with NVT conditions (canonical ensemble), and equilibrated for 100 ps with NPT (isothermal-isobaric) ensemble. Finally, 60 ns of NPT molecular dynamic simulation at 303 K was performed.

The ΔG of substrate binding was estimated using the MM-PBSA algorithm in a Poisson Boltzmann (PB) solvation [44] for 500 frames (snapshots from every 10 ps) from selected 5 ns of each MD simulation. Ionic strength was set to 0.15 M. To select a fragment of trajectory, distances describing the position of steroid were calculated: d(O\text{keto}-\text{OH}_{\text{Tyr487}}), d(H\text{2β}-\text{OH}_{\text{Tyr318}}) and d(H\text{1α}-N_{\text{FAD}}). The first of these parameters is describing the hydrogen bond between the 3-keto group of ketosteroid and Tyr487, the next two are related to the postulated mechanism of the catalyzed reaction, which postulate that H2β and H1α are transferred to Tyr318 and FAD respectively. For MMPBSA calculation we selected 5 consecutive ns, during which the sum of the standard deviation of these parameters was minimal.

**Results**

**Theoretical prediction of substrate specificity**

In the crystal structure of KSTD1 in complex with the ligand 1,4-androstadiene-3,17-dione (cf. 1), the steroid ring core (gonane ring system) of the ligand was buried in the active site pocket of the enzyme with its D-ring pointing towards the solvent-accessible area [33]. Likewise, the MD simulation of KSTD1:cholest-4-en-3-one complex revealed that the enzyme binds the steroid ring core while the isoocyt substituent at C17 position protrudes out of the active site and generally does not prevent it from the active binding (Fig. 2).
During the simulation, we also did not observe any significant clashes of the isooctyl side chain with the protein which would explain lack of the enzyme activity toward this class of substrate. From the reaction point of view, the most important interactions between substrate and the enzyme are limited to the 3-keto group of steroid ring A and its closest neighborhood (especially vicinity of C1 to FAD and C2 to Tyr318). Therefore we decided to check if the binding free energy could explain reported in the literature lack of activity [28]. Free energies of substrate binding were estimated for KSTD1 with MM-PBSA (Table 1). All studied compounds bound favorably to the active site with $\Delta G_{\text{bind}}$ in the range of $-11.0$ to $-7.9$ kcal/mol. Interestingly, there was no difference in $\Delta G_{\text{bind}}$ between the native substrate of KSTD1 from *R. erythropolis*, androst-4-en-3,17-dione ($-8.02 \pm 0.13$ kcal/mol, 1), and cholest-4-en-3-one ($-8.40 \pm 0.19$ kcal/mol, 2). Based on this result we formed the hypothesis that in principle it should be possible to convert with KSTD1 not only 3-ketosteroids with a degraded side chain (1, 4, 5-7) but also steroids such as 2 or even 3-ketosaponins as 3 (Fig. 2).

**Table 1 The estimated free energy of the binding of steroids to the KSTD1 active site**
| Ligand                                      | $\Delta G_{\text{bind}}$ [kcal/mol] | Std. err. of $\Delta G_{\text{bind}}$ [kcal/mol] |
|--------------------------------------------|-------------------------------------|-----------------------------------------------|
| Androst-4-en-3,17-dione (1)                | -8.02                               | 0.13                                          |
| Cholest-4-en-3-one (2)                     | -8.40                               | 0.19                                          |
| Diosgenone (3)                             | -6.17                               | 0.15                                          |
| Androstanolone (4)                         | -10.97                              | 0.15                                          |
| Progesterone (5)                           | -8.23                               | 0.17                                          |
| Testosterone propionate (6)                | -7.85                               | 0.15                                          |
| 6-Dehydrotestosterone acetate (7)          | -8.53                               | 0.15                                          |

**HBC/steroid inclusion complex formation**

The phase solubility diagrams of AD (1) and cholest-4-en-3-one (2) with HBC were determined in the HBC concentration range from 0 to 98 mM at 30°C, i.e., the temperature of catalytic assays (Fig. 3). As expected, the solubility of both steroids increased with higher HBC concentration. However, the solubility of more hydrophobic 2 in 98 mM HBC solution turned out to be over 10-times lower than the AD.

![Phase solubility diagrams](image)

**Fig. 3** Phase solubility diagrams for androst-4-en-3,17-dione (gray squares) and cholest-4-en-3-one (red circles).

The concentration of AD in the function of HBC concentration was fitted with the linear function ($R^2 > 0.998$) corresponding to $A_L$-type phase diagram. The linear course of phase solubility diagram with the slope less than 1 ($0.527 \pm 0.009$) indicates that HBC with AD forms the 1:1 inclusion complex. Meanwhile, the cholest-4-en-3-one/HBC phase solubility diagram
turned out to be the $A_P$-type and the data were fitted with quadratic function ($R^2 > 0.998$). $A_P$-type phase solubility diagram may indicate the formation of higher-order complexes with respect to cyclodextrin at higher HBC concentrations [30,31]. The fit parameters were presented in Table S1 of the Supporting Information, while the values of $K_{1:1}$ and $K_{1:2}$ were shown in Table 2.

**Table 2 Solubility parameters of steroids in the solution of HBC and 2% EGME at 30°C**

| Substrate                        | Solubility curve type | $K_{1:1}$ [M$^{-1}$] | $K_{1:2}$ [M$^{-1}$] |
|----------------------------------|-----------------------|-----------------------|-----------------------|
| Androst-4-en-3,17-dione (1)      | $A_L$                 | 7714.4                | -                     |
| Cholest-4-en-3-one (2)           | $A_P$                 | 1277.1                | 56.8                  |

The obtained constants were used to calculate the percentage of free and complexed steroids in the reaction medium. For androst-4-en-3,17-dione 99% of the substrate has been complexed by HBC. The ratio of free and complexed (1:1 and 1:2) cholest-4-en-3-one has varied slightly depending on the initial steroid concentration (Table S2) and averaged 2.9% of the free form, 53.5% of the steroid in 1:1 complex and 43.6% of the steroid in 1:2 complex.

**AcmBs native substrate**

To determine the substrate specificity of AcmB we conducted steady-state kinetic studies for androst-4-en-3,17-dione and cholest-4-en-3-one. 2,6-Dichloroindphenol was used as an artificial electron acceptor. Since the solubility of cholest-4-en-3-one in aqueous solutions is close to zero, measurements were carried out with 2% HBC and 2% EGME. The reaction velocities were presented in Figure 4. The Ping-Pong bi bi (non-sequential) mechanism was fitted to the obtained data (4) [45].

$$v = \frac{V_{max}[S]_t[DCPIP]}{K_{mS}[DCPIP] + K_{mDCPIP}[S]_t + [S]_t[DCPIP]}$$  \hspace{1cm} (4)
Due to the presence of HBC in the system, the substrate was available to the enzyme mostly in the form of 1:1 and/or 1:2 complexes (99% for S(HBC) AD complex, and 53.52 and 43.55% for S(HBC) and S(HBC)_2 complexes with cholest-4-en-3-one, see Table S2 for details). As a result, only in the case of AD, we could assume that the observed kinetics was mostly coming from the S(HBC) form of the substrate while in the case of cholest-4-en-3-one the kinetics of E:S complex formation was much more complicated and proceeds according to eq (5).

\[
\begin{align*}
E + S(HBC) & \xrightarrow{k_1} E:S + S(HBC) \\
E + S(HBC) & \xrightarrow{k_1} E:S + S(HBC) \\
E + S & \xrightarrow{k_2} E + S(HBC) \\
E + S & \xrightarrow{k_4} E + S(HBC)
\end{align*}
\]

(5)

E – enzyme (KSTD)
E’ – enzyme in its reduced form
S – substrate 1 (steroid)
HBC – 2-hydroxypropyl-β-cyclodextrin
DCPIP – substrate 2 (artificial electron acceptor)

Although it is possible to derive the kinetic equation that takes into account all forms of the substrate present in the solution, the correct fitting of all constants requires collecting kinetic data for different concentrations of HBC, steroid and DCPIP. Our experimental data turned out to be significantly underdetermined to handle such a complex non-linear fit. However, as the main purpose of our study was to compare different 3-ketosteroid dehydrogenases under the same conditions (here concentration of HBC which determines the distribution of S:HBC complexes) we have decided to present the kinetic parameters in the function of the total substrate concentration. Due to that, kinetic curves were fitted to the concentration of the total substrate.
Fig. 4 Results of steady-state kinetics for dehydrogenation reaction of androst-4-en-3,17-dione (A) and cholest-4-en-3-one (B) catalyzed by AcmB.

Under these conditions, the maximum velocity of reaction for AcmB ($V_{\text{max}}$) was over 8 times higher for AD than for cholest-4-en-3-one. However, AcmB affinity ($K_{\text{mS}}$) to cholest-4-en-3-one turned out to be over 20-times higher compared with AD. As a result, the enzyme catalytic efficiencies ($k_{\text{cat}}/K_{\text{mS}}$) for cholest-4-en-3-one was more than two-fold higher than that determined for AD.

**Table 3 Apparent kinetic parameters of the purified AcmB**

| Substrate                  | $K_{\text{mS}}$ [μM] | $K_{\text{mDCPIP}}$ [μM] | $V_{\text{max}}$ [μM min$^{-1}$] | $k_{\text{cat}}$ [s$^{-1}$] | $k_{\text{cat}}/K_{\text{mS}}$ [M$^{-1}$ s$^{-1}$] |
|----------------------------|----------------------|---------------------------|---------------------------------|-----------------------------|--------------------------------------------------|
| Androst-4-en-3,17-dione (1) | 529.2 ± 32.5         | 223.5 ± 16.9              | 859.2 ± 45.5                    | 46.1 ± 2.3                  | (0.87 ± 0.05) ∙ 10$^5$                              |
| Cholest-4-en-3-one (2)     | 23.7 ± 1.16          | 37.3 ± 3.3                | 104.6 ± 2.7                     | 5.8 ± 0.2                   | (2.45 ± 0.01) ∙ 10$^5$                              |

It should be noted here that DCPIP is also complexed by HBC. This fact however does not impact the comparisons of $K_{\text{mS}}$ values, as experiments were conducted for both substrates with the same type and concentrations of oxidation agent (DCPIP) and the HBC concentration was 30 to 84-times higher than DCPIP concentration. The influence of DCPIP complexation (such as decrease of HBC concentration) on obtained kinetic parameters was negligible.
In the genome of *S. denitrificans* we have identified two other putative AcmB homologs, which biological role has not been yet established. For this study, we selected gene encoding “3-oxosteroid 1-dehydrogenase” (GenBank: SMB21450.1) which we named AcmB2 due to its high sequence similarity to AcmB (44% identity and 62% similarity). The recombinant enzyme was expressed in *E. coli* BL21(DE3)Magic and purified on Ni-NTA matrix. To check if AcmB2 also exhibited activity with C17-substituted steroids we performed mini reactor tests with 100 μM of AD, cholest-4-en-3-one or diosgenone with the addition of 2% HBC (Fig. S8). In a reactor with diosgenone, we introduced a 5-fold higher concentration of AcmB2 than for other mini reactors to account for the observed lower rate of the initial tests. After 16 h of reaction, we observed significant conversions of all substrates i.e., 77.2 ± 1.5% for AD, 80.2 ± 2.7% for cholest-4-en-3-one and 53 ± 3.5% for diosgenone. These results show that other 3-ketosteroid dehydrogenase from *S. denitrificans* exhibits similar substrate specificity as AcmB.

**KSTD1 substrate spectrum**

The kinetic measurements for KSTD1 were performed at the same conditions as described previously for AcmB, except for the pH of the buffer solution. Surprisingly, the affinity of KSTD1 from *R. erythropolis* to AD and cholest-4-en-3-one turned out to be significantly lower than this exhibited by AcmB and it was not possible to reach enzyme saturation conditions in the presence of HBC. The solubility of steroids was limited even with solubilizing additives. On the other hand, a further increase of the HBC concentration results in an apparent *K_m* shift toward higher values as a consequence of substrate sequestration (i.e., decrease of free substrate concentration and stabilization of the substrate in S(HBC)_n form). Nevertheless, we have decided to estimate the first-order rate coefficients (*V_{max}/K_m*) for the low substrate concentrations. The linear function was fitted to the reaction velocities determined for 5 – 100 μM steroids concentrations (Fig. 5). To obtain comparable kinetic parameters, the same procedure was applied to AcmB (Fig. S9).
Fig. 5 Results of steady-state kinetics for dehydrogenation reaction of androst-4-en-3,17-dione (A) and cholest-4-en-3-one (B) catalyzed by KSTD1.

For the small concentrations of the first substrate \( S \ll K_m \) the velocity equation becomes invariant to the concentration of the second substrate (DCPIP) and simplifies to a first-order linear equation:

\[
v = \frac{V_{\text{max}} [S]}{K_m}
\]

(6)

The apparent catalytic efficiency \( k_{\text{cat}}/K_m \) of AcmB and KSTD1 for AD was of the same order. KSTD1 proved to be a slightly (22%) better catalyst for the AD \( \Delta^1 \)-dehydrogenation than AcmB. In the case of the reaction with cholest-4-en-3-one, the \( k_{\text{cat}}/K_m \) value measured for AcmB was almost 20-times higher than for KSTD1. The KSTD1 was 10-times better in the \( \Delta^1 \)-dehydrogenation of AD vs cholest-4-en-3-one.

The obtained results confirmed undeniably the ability of KSTD1 to catalyze \( \Delta^1 \)-dehydrogenation of steroids with the aliphatic chain on the C17 position. Due to the low cholest-4-en-3-one bioavailability and the lower KSTD1 affinity to this steroid, it was not possible to determine the actual KSTD1 substrate spectrum without the addition of solubilizing agent [28].
Table 4 Apparent kinetic parameters of the purified AcmB and KSTD1

| Substrate                                    | AcmB                  | KSTD1                 |
|----------------------------------------------|-----------------------|-----------------------|
| Androst-4-en-3,17-dione (1)                  | \((4.00 \pm 0.04) \cdot 10^6\) | \((5.17 \pm 0.03) \cdot 10^6\) |
| Cholest-4-en-3-one (2)                       | \((9.25 \pm 0.17) \cdot 10^6\) | \((4.71 \pm 0.03) \cdot 10^5\) |

Based on the results of MD simulations for KSTD1 and diosgenone ES complex and experimentally confirmed the activity of AcmB and AcmB2 with diosgenon [8] we decided to investigate if KSTD1 is also able to catalyze the reaction with 6-6-member ring steroids. The mini reactor tests proved that in presence of 2% HBC 100 μM diosgenone can be converted with excellent 97.3 ± 0.8% conversion in 20 min.

**Discussion**

The results of theoretical MD simulations for 3-ketosteroid Δ^1-dehydrogenase from *R. erythropolis* SQ1 have shown that neither isooctyl side chain of cholest-4-en-3-one nor polycyclic substituent of diosgenone is a steric hindrance for the reaction catalyzed by KSTD1. Only the gonane ring is recognized by KSTD active site and the additional substituents or rings are protruding into the solvent, outside of the enzyme. This hypothesis was successfully confirmed by experimental kinetic studies and reactor tests. Nevertheless, the affinity and the catalytic efficiency of KSTD1 from *R. erythropolis* for cholest-4-en-3-one turned out to be significantly lower than that determined for AcmB from *S. denitrificans*. Additionally, a more hydrophobic character of the AcmB surface when compared to KSTD1 may also contribute to its higher affinity to more hydrophobic substrates [46].

Our results are in line with reports of Wang et al. [21], who showed that KSTD from *R. erythropolis* WY 1406 strain can also convert substrates with extended C17 substituent, like 21-acetoxy-pregna-4,9(11),16-triene-3,20-dione. In addition to AcmB and the new AcmB2 [8,20], there were some literature reports about KSTDs activity towards steroids with a more
complex side chain on the C17 position [17–19,22,47]. Zhang et al. described five KSTD isoenzymes from *Gordonia neofelisaecis* of which four exhibited activity toward cholest-4-en-3-one. Importantly, the authors conducted experiments with 0.05% Tween 80 acting as a solubilizing agent [19].

Poor solubility of steroids with the aliphatic or polycyclic substituents at the C17 atom results in their low bioavailability to the enzyme. Compounded with a low affinity of most KSTDs to more complex steroids, these two factors may lead to inaccurate conclusions regarding substrate specificity. The results obtained for KSTD1, AcmB, AcmB2, as well as for KSTDs from *G. neofelisaecis* [19] indicate that the use of cyclodextrins or detergents can influence enzymes substrate specificity analysis. Such analysis, however, requires determination of the HBC/steroid stability constants which in turn help in the determination of the real concentration of the substrate, but complicate the observed kinetics due to different $G_s$ of E:S formation for S, S(HBC) or S(HBC)$_2$ forms of the substrate. Furthermore, it should be underlined here that there are several different methods for the determination of these constants (HPLC, NMR, FT-IR, XRD or calorimetry-based, etc.) and they usually differ in the delivered values [48,49]. Despite these difficulties, the application of solubilizing agent seems indispensable for the determination of the active substrates, especially when KSTD affinity to the compounds with an aliphatic chain on the C17 position is high, as in AcmB (apparent $K_{m,S}$ of 23.7 μM). Our results indicate that AD may not be a native substrate of AcmB. The established affinity toward AD is, however, the affinity toward S(HBC) complex (apparent $K_{m,S}$ of 529.2 μM) as we know from our previous kinetic studies that $K_m$ value measured in the assay without HBC was an order of magnitude lower (apparent $K_m$=59.6 ± 3.0 μM at 200 μM DCPIP and 30°C) [8] than in the assay with HBC (Table 3). The $K_{m,S}$ determined in this work for cholest-4-en-3-one is lower than those observed for AD without HBC. Furthermore, as 97% of the cholest-4-en-3-
one was in the form of complexes with HBC we can assume that the real $K_{ms}$ for cholest-4-en-3-one (if it was possible to measure it without HBC) is even lower.

The difference in the affinity of KSTDs to C17-substituted steroids might result from their structural differences. The homology modeling of KSTD from *Arthrobacter simplex* (active towards cholest-4-en-3-one) showed the presence of an additional loop close to the enzyme active site, non-observed in the crystal structure of KSTD1 form *R. erythropolis* [33,50]. Interestingly, the same feature was noticed in the AcmB homology model [51].

Our findings have implications for the understanding of the physiological role of AcmB. Initial studies on the anoxic cholesterol metabolism of *S. denitrificans* reported new metabolites, cholest-1,4-diene-3-one and 25-hydroxycholest-1,4-dien-3-one, and suggested that the double bond between the C1 and C2 atoms have had to be introduced at the early stage of cholesterol mineralization. As a result, cholest-4-en-3-one was considered as the AcmB native substrate [24,25]. However, subsequent kinetic studies demonstrated the AcmB preference for C19 – C21 steroids [20], which was supported by the localization of the enzyme on the cytoplasmic side of the bacterial inner membrane [26]. Due to that fact, Lin *et al.* deduced that the AcmB native substrate is rather the AD [26]. Kinetic studies indicating AcmB localization were estimated based on the enzyme activity measured in particular fractions, not by e.g., antibody technique and AcmB specific activity was calculated for the total protein extracted from the membrane. Importantly, the KSTD activity was still detected significantly in the peripheral-periplasmic fraction [26]. In this work, we demonstrated, that a study of the enzyme-substrate specificity and affinity in the case of hydrophobic steroids without the solubilizing agent addition may lead to misinterpretation of achieved results. In the presence of HBC, the AcmB affinity toward cholest-4-en-3-one turned out to be significantly higher than for AD (even taking into consideration all complications introduced by substrate complexation by HBC).
Finally, one has to remember about two other AcmB sequence homologs, from which we confirmed 3-ketosteroid Δ\(^1\)-dehydrogenase activity for one of them, AcmB2. Therefore it may well be that *S. denitrificans* deploys different KSTD enzymes on both sides of its inner membrane, which specialize in converting different types of 3-ketosteroids (as reported already for KSTD isoenzymes from *R. erythropolis* [52,53]). This aspect definitely requires further studies.

**Conclusions**

Our results indicate that it is necessary to test KSTD substrates in the presence of solubilizing agents such as HBC, to discern truly inactive steroids from those which under experimental conditions are in too low concentrations. Unfortunately, the presence of HBC introduces additional complications to the kinetic model due to the presence of different forms of the substrate (S and S(HBC)\(_n\)). Our modeling studies suggest that in principle all KSTDs, which share structural characteristics with KSTD1 should be able to convert more hydrophobic substrates with extended ring system or C17 substituent. This conclusion has an important impact on the biotechnological application of those enzymes.

**Additional files**

**Abbreviations**

AcmB: 3-ketosteroid Δ\(^1\)-dehydrogenase 1 from *Stereolibacterium denitrificans*; AcmB2: 3-ketosteroid Δ\(^1\)-dehydrogenase 2 from *Stereolibacterium denitrificans*; ACN: acetonitrile; AD: androst-4-en-3,17-dione; AMBER: assisted model building with energy refinement; AND: 1,4-androstadien-3,71-dione; BME: β-mercaptoethanol; DAD: diode array detector; DCPIP: 2,6-dichloroindophenol; dCTP: deoxycytidine triphosphate; dGTP: deoxyguanosine triphosphate;
DMSO: dimethyl sulfoxide; DTT: dithiothreitol; EGME: 2-methoxyethanol; ESP: Electrostatic Potential; FAD: flavin adenine dinucleotide; HBC: 2-hydroxypropyl-β-cyclodextrin; HPLC: high performance liquid chromatography; IPTG: isopropyl β-D-1-thiogalactopyranoside; kbp: kilobase pare; $k_{\text{cat}}$: catalytic constant/turnover number; $K_m$: Michaelis constant; $K_S$: HBC/steroid complex stability constant; KSTD(s): 3-ketosteroid $\Delta^1$-dehydrogenase(s); KSTD1: 3-ketosteroid $\Delta^1$-dehydrogenase from *R. erythropolis* SQ1; LB: Lennox Broth; MD: molecular dynamics; MM-PBSA: molecular mechanics energies combined with the Poisson–Boltzmann and surface area continuum solvation; MS: mass spectrometry; NPT: Isothermal-isobaric ensemble; NVT: Canonical ensemble; PDB: Protein Data Bank; PMSF: phenylmethylsulfonyl fluoride; RESP: Restrained Electrostatic Potential; $V_{\text{max}}$: maximal reaction velocity

**Authors’ contributions**

M. Sz. and P. W. designed research; M. G. conducted theoretical modeling; P. W. and A. R. conducted research on KSTD1; P. W. conducted research on AcmB; A. W. conducted research on AcmB2; P. W., M. G., A. W. and M. Sz. analyzed data; P. W., M. G. and A. W. wrote the methods; P. W. and M. Sz. wrote the introduction, results and discussion. M. Sz. and A. W. funded the research. All authors edited and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

All data generated and analyzed during this study are included in this published article and its additional files. Additional raw data (MD trajectories, raw kinetic data) are available on request.

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Not applicable.

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