Regioselective para-Carboxylation of Catechols with a Prenylated Flavin Dependent Decarboxylase

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Abstract: The utilization of CO₂ as a carbon source for organic synthesis meets the urgent demand for more sustainable production of chemicals. Herein, we report on the enzyme-catalyzed para-carboxylation of catechols, employing 3,4-dihydroxybenzoic acid decarboxylases (AroY) that belong to the UbID enzyme family. Crystal structures and accompanying solution data confirmed that AroY utilizes the recently discovered prenylated FMN (prFMN) cofactor, and requires oxidative maturation to form the catalytically competent prFMN<sup>iminium</sup> species. This study reports on the in vitro reconstitution and activation of a prFMN-dependent enzyme that is capable of directly carboxylating aromatic catechol substrates under ambient conditions. A reaction mechanism for the reversible decarboxylation involving an intermediate with a single covalent bond between a quinoid adduct and cofactor is proposed, which is distinct from the mechanism of prFMN-associated 1,3-dipolar cycloadditions in related enzymes.

Carboxylation reactions have received considerable attention in view of the use of CO₂ as an abundant C source for sustainable chemical production. However, to date, only a few examples of CO₂ fixation reactions have been realized on industrial scale, mainly owing to the high energy input required for substrate activation. In recent years, biocatalysts have been exploited as attractive alternatives to chemical methods to catalyze carboxylation reactions under mild, aqueous conditions. Whereas the biocatalytic carboxylation of aldehydes (TPP-dependent pyruvate decarboxylases), epoxides (epoxide carboxylases from Xanthobacter sp.),[4] and heteroaromatic compounds, such as pyrroles (pyrrole-2-carboxylate decarboxylase from Bacillus megaterium)[5] and indoles (indole-3-carboxylate decarboxylase from Arthrobacter nicotianae),[6] exhibited narrow substrate specificity, promising results were obtained in the biocatalytic carboxylation of phenols and styrenes. ortho-Benzonic acid decarboxylases and phenolic acid decarboxylases show a relaxed substrate specificity for the ortho-carboxylation of phenols[7] and the β-carboxylation[8] of styrenes, respectively, whilst maintaining their exquisite regioselectivity.

To expand the toolbox for biocatalytic carboxylation, we searched for enzymes enabling the regiocomplementary para-carboxylation of phenols. Most of the already characterized enzymes either require an ATP-consuming activation (phosphorylation) step prior to carboxylation (phenylphosphate carboxylases),[9] suffer from a rapid loss of activity under aerobic conditions, especially after purification (4-hydroxybenzoate[10] and 3,4-dihydroxybenzoate decarboxylases[12]) or have not been extensively examined in vitro yet,[12] which limits their usability for biotransformations. Based on a literature survey and a preliminary activity screen of hetero-
logously expressed potential \textit{para-}carboxylases (see the Supporting Information). 3,4-dihydroxybenzoic acid decarboxylases from \textit{Enterobacter cloacae} (EcAroY) and \textit{Klebsiella pneumoniae} (KpAroY, 89\% identical) were selected for further studies.\textsuperscript{[3a,13]} Both enzymes belong to the UbiD family and are related to a ferulic acid decarboxylase (Fdc1), which has been shown to facilitate the (de)carboxylation of cinnamic acids in the presence of a recently discovered prenylated flavin (prFMN) cofactor.\textsuperscript{[14]}

Owing to the natural occurrence of the UbiD-associated prenyltransferase UbiX in the \textit{E. coli} expression host, lyophilized whole cells displayed high decarboxylation activity in initial screenings (see the Supporting Information). However, upon purification of the decarboxylase from the \textit{E. coli} host, only little enzyme activity could be detected for either of the two enzymes, despite additional co-expression with the UbiD-associated prenyltransferase UbiX to provide sufficient prFMN in vivo. Upon in vitro reconstitution with reduced prFMN (Figure 1a),\textsuperscript{[14,15]} decarboxylation activity could be detected with 3,4-dihydroxybenzoic acid (3,4-DHBA, 1) following brief exposure to oxygen to generate the active prFMN$^\text{iminium}$ form (Figure 1b). The lack of activity for anaerobically reconstituted protein clearly demonstrates the requirement for oxidative maturation of the prFMN cofactor. EPR and UV/Vis spectroscopy revealed the presence of a radical semiquinone intermediate following in vitro reconstitution and oxidation, which is reminiscent of an intermediate observed with the related \textit{E. coli} UbiD\textsuperscript{[15]} (see the Supporting Information). After reconstitution and maturation, the enzyme activity has a half-life of only 5–8 min under aerobic conditions but remained unchanged for at least 14 h under anaerobic conditions (Figure 1c).

A comparison of the \textit{Kp}AroY and \textit{Ec}AroY crystal structures revealed only small differences, in accordance with the similar activities observed in solution experiments. The AroY monomer consists of an N-terminal prFMN-binding domain (residues 1–339), an oligomerization domain (residues 340–475), and a C-terminal $\alpha$-helix (residues 476–495; Figure 2a). A comparison with the previously reported structures of the fungal Fdc1 and \textit{E. coli} UbiD\textsuperscript{[14]} shows that AroY structures adopt an “open” conformation, where the position of the prFMN domain is more akin to that observed for UbiD, compared to the more closed conformation observed for Fdc1 (see the Supporting Information). This open conformation is observed for all crystallographically independent AroY monomers, and in the 4.6 Å cryo-EM solution structure of apo-EcAroY (Figure 2a).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1}
\caption{a) Established in vitro reconstitution of AroY with prFMN generated by UbiX from FMN and DMAP and subsequent aerial oxidation. b) AroY requires oxygen for activation; the activity was measured using 150 $\mu$M 3,4-DHBA (1) and is relative to the highest activity detected. c) Decay of oxidized KpAroY and EcAroY when kept under aerobic or anaerobic conditions. Measured with 150 $\mu$M 1. $t_{1/2}$ = 8.1 ± 1.3 min for KpAroY and 5.3 ± 1.2 min for EcAroY.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2}
\caption{AroY structure and suggested mechanism based on calculations. a) Hexameric quaternary structure of EcAroY. Dimer pairs are shown in ribbon representation in green and blue. The cryo-EM envelope is shown as a gray translucent surface. A detailed view of the EcAroY monomeric structure (in the circle) showing the prFMN-binding domain (blue), oligomerization domain (green), and C-terminal helix (red) is also given. b) Optimized structure of the active-site model employed in the computational study. Atoms marked with asterisks were fixed during the geometry optimization. The prFMN cofactor is shown in green and the substrate in salmon ball-and-stick presentation. Distances are given in Å. For clarity, only polar hydrogen atoms and the hydrogen atoms on the substrate are shown. c) Reaction mechanism suggested on the basis of the calculations.}
\end{figure}
The active site of AroY is clearly defined by the prenylated isoalloxazine moiety of the cofactor and the presence of key conserved residues (Figure 2b). It is situated near the hinge point of the prFMN-binding domain motion, and at the cleft between the oligomerization and the prFMN-binding domains. Two water molecules are clearly defined in the active site; one of them forms hydrogen bonds to His327 and Lys363 while the other one interacts with Lys363, His436, and Arg188. We hypothesize that the two water molecules mimic the two hydroxy groups at the meta- and para-positions of the substrate protocatechueic acid (1). Given the relative rigidity of the substrate, we superimposed a catechol moiety onto the two water molecules, and the carboxylate moiety was positioned in close proximity to the prFMN iminium group. While this has some similarity to the structure of the Fdc1:substrate complex (PDB-ID: 4ZA7),[^14] the exact relative position of the substrate carboxylate moiety and the prFMN N5–C1' iminium linkage is different. In Fdc1, the substrate α-carbon atom adjacent to the carboxyl group is located directly above the prFMN C1' atom whereas in AroY, the α-carbon atom is located above the isoalloxazine N5 atom (see the Supporting Information).

Exchanging several amino acids within the putative catechol-binding motif (Arg188, His327, Lys363 to Ala and His436 to Lys or Thr) led to a complete loss of activity. The same effect was observed upon exchange of Arg181 or Glu289, which are located near the carboxylate group of the substrate, to Ala. The analogous Glu282 in Fdc1 is proposed to be required for the donation of a proton to the covalently bound intermediate,[^14a,16] which hints at a similar role for Glu289 as a catalytic acid in AroY.

Based on the available structural, mutational, and kinetic data as well as DFT calculations (see the Supporting Information), we propose a reaction mechanism involving a quinoid intermediate (Figure 2c). This intermediate has chemical similarity to that proposed for the phenolic acid decarboxylases (PAD)^[^16] and is different from the 1,3-dipolar cycloaddition mechanism proposed for Fdc1.[^14a,16–18] The calculations, employing a large model of the active site with 283 atoms (Figure 2b), suggested that the generation of a cycloadduct is unlikely in the case of AroY as it would require the formation of a very strained intermediate (Figure S29). The latter is not required in the case of the cinnamic acid like substrates of Fdc1, for which previous calculations have validated the proposed 1,3-dipolar cycloaddition mechanism to the exocyclic alkene.[^14a,16b] Instead, the mechanism involving a quinoid intermediate was calculated to have feasible energy barriers, which are significantly lower than those of the 1,3-cycloaddition mechanism. The calculated energy profile and optimized structures of all intermediates and transition states along the reaction pathway are given in Figures S30–S32.

In view of synthetic applicability, the substrate scope of EcAroY was investigated with different protocatechuic acid derivatives (Scheme 1a, 1–3). EcAroY exhibited high decarboxylation activity with 3,4-dihydroxybenzoic acid (1, > 99% after 1 h; Scheme 1a) and gallic acid (2), which reacted at a slightly lower rate than 1 (ca. 83% conv. after 1 h). For its isomer 2,3,4-trihydroxybenzoic acid (3), however, no reaction was observed. Very similar kinetic parameters for the decarboxylation of 1 with purified and in vitro reconstituted enzymes were determined for both AroY enzymes (KpAroY: vmax = 4.7 ± 0.4 s⁻¹, Kapp = 96 ± 23 µM; EcAroY: vmax = 4.6 ± 0.4 s⁻¹, Kapp = 61 ± 17 µM). These parameters are reported as apparent values given the very low prevalence of inactive species (i.e., enzyme-bound FMN or radical prFMN).

To force the reaction equilibrium towards the carboxylation of suitable substrates, we subjected phenols 4–8 to a 3 M potassium bicarbonate buffer as a CO₂ source (Scheme 1b). Whereas the carboxylation of simple phenols was not successful (nonsubstrates are shown in Table S7), EcAroY catalyzed the regioselective para-carboxylation of catechol (4) in the presence of either bicarbonate or pressurized CO₂ (30 bar). Apart from 4, EcAroY also accepted pyrogallol (5) as a substrate, and carboxylation with bicarbonate predominately occurred in the para-position to the central hydroxy group to give 2. Carboxylation in vicinity to the peripheral hydroxy groups occurred only to a minor extent to give 3.

As the catechol scaffold appears to be crucial for substrate acceptance, the carboxylation of substituted catechols 6–8 with bicarbonate was tested (Scheme 1b). Small electron-withdrawing (3-F, 6) to medium-sized electron-donating (3-OME, 7; 3-Me, 8) substituents were tolerated in the 3-position, and carboxylation occurred exclusively at the 5-position, that is, in para-position to the central hydroxy group, which was confirmed by NMR spectroscopy (see the Supporting Information).

In conclusion, our data contribute to a better understanding of the ATP-independent para-carboxylation of phenolic substrates. Crystal structures and in vitro reconstitution data unambiguously demonstrate that prFMN is employed as a cofactor, and that oxidative maturation is required for activity. The exact mechanism of oxidative maturation and the cause of the observed oxygen sensitivity remain unclear at this stage. Owing to the preference of AroY for catechols over simple phenols, a second hydroxy group seems to be mandatory for ideal substrate positioning (with...
a hydrogen bond between OH and His327; Figure 2b) in the active site of the enzyme. The second hydroxy group further enhances the nucleophilicity of the (catechol) substrate to facilitate the nucleophilic addition step onto prFMN. Electron-withdrawing and -donating groups are tolerated in the meta-position relative to the carboxylation site whereas substitution in the ortho-position was not tolerated owing to steric hindrance in the active site.

In the context of the wider UbiD family, the metal-assisted binding and the associated oxidative maturation of prFMN are common to all biochemically and structurally characterized enzymes (the fungal Fdc1, E. coli UbiD, and AroY). The substrate binding specificity is distinct for each of these enzymes, but in all cases appears to be largely governed by residues derived from the oligomerization domain that are involved in binding to the non-carboxylate substrate moiety. The carboxylate group, on the other hand, is bound near the conserved Glu-Arg-Glu/Asp triad of ionic residues, which is located near the prFMN N5=Cl− iminium moiety. A key difference is the relative position of the oligomerization and prFMN domains, and thus the relative position of the substrate-binding and carboxylate-binding motifs. These are considerably closer in the fungal Fdc1 structure than in bacterial UbiD and AroY. A putative domain motion might allow the UbiD/AroY enzymes to adopt a more Fdc1-like conformation, but this has not been directly observed. While the quinoid-based mechanism proposed for AroY suggests an alternative to 1,3-dipolar cycloadditions in the case of aromatic substrates, it does not explain how the (de)carboxylation of non-phenolic substrates[19] is achieved. A better understanding of the UbiD enzyme family will require further studies of these and additional family members.[20]

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Conflict of interest

The authors declare no conflict of interest.

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