Metal Ion Promiscuity and Structure of 2,3-Dihydroxybenzoic Acid Decarboxylase of Aspergillus oryzae

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Broad substrate tolerance and excellent regioselectivity, as well as independence from sensitive cofactors have established benzoic acid decarboxylases from microbial sources as efficient biocatalysts. Robustness under process conditions makes them particularly attractive for preparative-scale applications. The divergent metal-dependent enzymes are capable of catalyzing the reversible non-oxidative (de)carboxylation of a variety of electron-rich (hetero)aromatic substrates analogously to the chemical Kolbe-Schmitt reaction. Elemental mass spectrometry supported by crystal structure elucidation and quantum chemical calculations verified the presence of a catalytically supported metal complexed in the active site of 2,3-dihydroxybenzoic acid decarboxylase from Aspergillus oryzae (2,3-DHBD_Ao). This unique example with respect to the nature of the metal is in contrast to mechanistically related decarboxylases, which generally have Zn²⁺ or Mn²⁺ as the catalytically active metal.

The use of carbon dioxide as a C₂ reagent for the production of valuable chemicals, such as urea, (poly)carboxylates and phenolic acids, has recently sparked growing attention.[1] Of particular interest is the carboxylation of C nucleophiles yielding carboxylic acids.[2] A perfect atom economy of 100% and the low (no) cost of the reagent CO₂ renders carboxylation a highly attractive method. However, in traditional chemical protocols harsh reaction conditions require significant energy input, which often causes limited (regio)selectivities.[3] In this context, biocatalytic alternatives, by making use of decarboxylases in the reverse carboxylation direction, offer an elegant alternative.[4] Decarboxylases from secondary metabolic pathways are particularly attractive due to their relaxed substrate portfolio.[5] Three main types of enzymes, which greatly differ in their catalytic mechanism and cofactor requirement and consequently act on completely different substrates, have been elucidated so far:[6] i) ortho-carboxylation of phenols in analogy to the Kolbe-Schmitt process[7] is catalyzed by metal-dependent o-benzoic acid decarboxylases (o-BDs), which excel not only due to their high stability, but also by an unusually broad substrate portfolio.[8] ii) Side-chain carboxylation at the vinyl group of p-hydroxystyrenes, for which no chemical protocol exists, is feasible with metal-independent phenolic acid decarboxylases.[9] iii) More recently, the (ATP-independent) p-carboxylation of phenols and the decarboxylation of electron-rich heterocyclic and acrylic acid derivatives was shown to be catalyzed by prenylated FMN-dependent decarboxylases.[8] Although some of these enzymes display a remarkable substrate acceptance, stability-problems of their prenylated FMN-cofactor impose limitations on their large-scale use.[9]

Due to their independence from sensitive cofactors and their excellent stability, o-BDs are most attractive for large-scale applications.[10,11] In order to facilitate their applicability, computational methods are increasingly applied for the prediction of substrate-structure activities aiming to minimize time-consuming and expensive trial-and-error wet-lab experiments. Although the mechanism of o-BDs is basically well understood,[12,13] conflicting data exist concerning the nature of their catalytically essential divalent metal, for example, Zn²⁺, Mn²⁺ or Mg²⁺, which aggravates computational studies leading to inaccuracies in substrate-binding and energy pathways depending on the ionic radius and Lewis acidity[14] of the metal involved. In this study, we investigated the metal dependence of 2,3-dihydroxybenzoate decarboxylase from Aspergillus oryzae (2,3-DHBD_Ao)[15] by determination of its crystal structure, metal ion analysis by inductively coupled plasma tandem mass spectrometry (ICPMS/MS) and the catalytic energy profile by quantum chemical calculations.

All of the metal-dependent decarboxylases identified so far are members of the amidohydrolase superfamily, which share significant structural and mechanistic similarities, such as a (β/
αβ-barrel fold harboring one catalytically essential divalent metal ion in the active site.[16] Although the overall sequence similarity between subclasses is low, several amino acid residues involved in metal binding and catalysis are conserved (Table 1).[12] The mechanism follows an electrophilic aromatic substitution in analogy to the Kolbe-Schmitt reaction: First, a divalent metal ion chelates the carboxylate and phenol group in the α-position, which facilitates protonation at the ipso-carboxylate position by a highly conserved Asp residue, thereby breaking aromaticity as the rate-determining step. Subsequent cleavage of the C–C bond yields CO₂ and phenol as products (see figure in Table 1).[40] The identity of the electrophile CO₂ as co-product/co-substrate (as opposed to the nucleophile bicarbonate) has recently been unambiguously clarified.[17]

α-Benzonic acid decarboxylases (α-BDs) and close relatives display an interesting variety of divalent metal ion requirement. Initially Zn²⁺ was considered as the dominating metal, but recently the picture became more complex (Table 1). Elemental mass spectroscopy (ICPMS/MS) and density functional theory (DFT) calculations identified Mn²⁺ (rather than Zn²⁺ as previously assumed) as essential metal in iso-orotate decarboxylase from Cordyceps militaris (IDCase_Cm).[14] In accordance, characterization of 2,3-DHBD_Ao revealed that it does not contain Zn²⁺ as reported for a homologous enzyme from Fusarium species,[18] but is also catalytically active with Mn²⁺.

More surprisingly, 2,3-DHBD_Ao has a significantly improved turnover rate with Mg²⁺. To the best of our knowledge, this is the first example of a Mg²⁺-dependence of α-BDs. These findings are corroborated by quantum chemical calculations, which revealed a reduced activation barrier by 2 kcal/mol in the rate determining step (see below).

The structure of recombinant 2,3-dihydroxybenzoic acid decarboxylase of A. oryzae was solved by X-ray crystallography to 1.2 Å (Table S1 in the Supporting Information). In accordance with the recently published homologue from F. oxysporum[40] which shows 74% sequence identity and related decarboxylases, the enzyme features a distorted (β/α)₂-barrel fold with its active site at the centre of the barrel (Figure 1A). The catalytic metal shows well defined density complexed by Glu8, His167, the proton donor Asp293 and three water molecules. Interpretation of this density as the expected Zn²⁺ (Figure 1B) vastly overestimated the electron density. In contrast, Mg²⁺ gave a perfect fit (Figure 1B). Comparison (Zn²⁺ vs Mg²⁺) of the electron density based on the F. oxysporum crystal structure (PDB ID: 6M53[18]) reflects our findings (Figure S1). Crystals grown in the absence of Mg²⁺ showed less electron density, indicating a depletion of metal. In this case, the active site was either never correctly reconstituted with the metal during expression, or it was lost during purification and crystallization due to the weak trivalent complexation of the ion. Likewise, lack of the preferred ion, such as Zn²⁺ or Mn²⁺, would be due to their low abundance in growth medium at high protein expression levels.

To exclude that the occurrence of Mg²⁺ was a crystallographic artefact, ICPMS/MS analysis as well as activity measurements were performed.

Protein samples obtained from size exclusion chromatography (SEC) incubated with either Mg²⁺, Mn²⁺, Zn²⁺ (or a buffer devoid of these ions as control) were split into two parts to determine steady state turnover frequencies (TOFₘₐₓ, Figure 2) and their metal occupancy. SEC coupled to ICPMS/MS showed that the control sample contained no Zn²⁺ and only low amounts of Mg²⁺ and Mn²⁺ (around 17 and 10% occupancy, respectively, Table S2, Figure S2). Incubation with Mg²⁺ or Mn²⁺ showed uptake of either ion with differences in affinity and competition. Mn²⁺ was able to displace the Mg²⁺ originally

Table 1. Simplified general mechanism of α-BDs; catalytically active metal ions and their ligands among α-BDs listed by increased sequence identity to 2,3-DHBD_Ao.

| α-BD      | Catalytic Asp | M⁺⁺ Ligands | M⁺⁺ Sequence identity | PDB Ref |
|-----------|---------------|--------------|------------------------|---------|
| IDC_Cm    | Asp323        | His12, His14, His195, Asp323 | Mn⁺⁺; (Zn⁺⁺)[44] | 19 % | 4HK5 [19,12] |
| LigW_Sp   | Asp296        | Glu7, His173, Asp296 | Mn⁺⁺ | 26 % | 4ICM [20] |
| LigW_Na   | Asp314        | Glu19, His188, Asp314 | Mn⁺⁺ | 29 % | 4QRN [20] |
| 2,6-DHBD_Rs | Asp287      | Glu8, His10, His164, Asp287 | Zn⁺⁺ | 42 % | 2DVU [21] |
| 2,6-DHBD_Fs | Asp287      | Glu8, His10, His164, Asp287 | Mn⁺⁺ | 42 % | 4QRO [22] |
| SAD_Tm    | Asp298       | Glu8, His169, Asp298 | Zn⁺⁺ | 51 % | 6JQW [23] |
| 2,3-DHBD_Fo | Asp291      | Glu8, His167, Asp291 | Zn⁺⁺ | 74 % | 6M53 [18] |
| 2,3-DHBD_Ao | Asp293      | Glu8, His167, Asp293 | Mg⁺⁺ | 100 % | 7A19 [10] |

[a] Previous assumption[15,16] IDC_Cm = iso-orotate decarboxylase from C. militaris[14] LigW_Sp and LigW_Na = 5-carboxyvanillate decarboxylase from Sphingomonas paucimobilis and Novosphingobium aromaticivorans,[25,26] 2,6-DHBD_Rs and 2,6-DHBD_Fs = 2,6-dihydroxybenzoic acid decarboxylase[27] from Rhizobium and Polaromonas species;[13,15] SAD = salicylic acid decarboxylase from Trichosporon moniliiforme;[26,28] 2,3-DHBD_Fo and 2,3-DHBD_Ao = 2,3-dihydroxybenzoic acid decarboxylase from Fusarium oxysporum and A. oryzae.[15,16]

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found and eluted from the SEC column in stoichiometric ratio with the protein. On the other hand, Mg\(^{2+}\) was unable to replace Mn\(^{2+}\) and was only recovered with 60\% occupancy. This reduced ratio is likely due to a significant off rate of Mg\(^{2+}\) from the active site during SEC at 30°C in Mg\(^{2+}\)-free buffer, rather than an inability to completely saturate the enzyme, especially considering the complete occupancy in the crystal.

Zn\(^{2+}\) caused complete precipitation of the enzyme even in sub-millimolar concentrations. Enzyme-bound zinc could not be detected in any of the samples.

Additionally, we determined the relative activity of 2,3-DHBD\(_{Ao}\) in the decarboxylation of the enzyme’s natural substrate 2,3-dihydroxybenzoic acid (2,3-dhba) with respect to the metal ion in the active site. The highest rate was found with protein incubated with Mg\(^{2+}\), which was approximately 2.5 times higher than the one treated with Mn\(^{2+}\) and twice as high as the control sample without any additional metal ions (Figure 2). As mentioned above, the control samples were not completely devoid of Mg\(^{2+}\). This explains the slightly higher activity in the non-metal-treated preparation versus the Mn\(^{2+}\)-treated sample where most of the Mg\(^{2+}\) ions were replaced by Mn\(^{2+}\). In order to complete the data set, protein samples were also treated with Zn\(^{2+}\), which led to complete loss of enzyme activity due to precipitation.

Kinetic parameters of 2,3-DHBD\(_{Ao}\) loaded with Mg\(^{2+}\) or Mn\(^{2+}\) are well within the range of other metal-depending decarboxylases with different substrate preferences (Table S3). The \(k_{cat}/K_m\) value for Mg\(^{2+}\) containing 2,3-DHBD and the (putatively) Zn\(^{2+}\) occupied enzyme from literature is virtually identical.

Considering that the presence of Mg\(^{2+}\) is unprecedented in this enzyme family, we set out to corroborate these findings by quantum chemical calculations.

To gain insights into the reaction mechanism of 2,3-DHBD\(_{Ao}\), density functional theory (DFT) calculations were performed using a large cluster model of the active site (see the Supporting Information for computational details and model design). It was found that 2,3-DHBD\(_{Ao}\) follows a similar mechanism as previously established for LigW\(^{[17]}\) and 2,6-DHBD\(^{[13]}\), both in terms of the sequence of steps and also the associated energy barriers (Figure 3). For the Mg-enzyme complex, the protonation of the substrate by Asp293 was found and eluted from the SEC column in stoichiometric ratio with the protein. On the other hand, Mg\(^{2+}\) was unable to replace Mn\(^{2+}\) and was only recovered with 60\% occupancy. This reduced ratio is likely due to a significant off rate of Mg\(^{2+}\) from the active site during SEC at 30°C in Mg\(^{2+}\)-free buffer, rather than an inability to completely saturate the enzyme, especially considering the complete occupancy in the crystal. Zn\(^{2+}\) caused complete precipitation of the enzyme even in sub-millimolar concentrations. Enzyme-bound zinc could not be detected in any of the samples.

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to be rate-limiting, with a calculated barrier of 15.8 kcal/mol (TS\textsubscript{1Mg}), and the formed 2,4-dienone intermediate (Int\textsubscript{Mg}) lies 8.8 kcal/mol above the enzyme-substrate complex (E:S\textsubscript{Mg}). The following decarboxylation was calculated to have a low barrier (TS\textsubscript{2Mg}), 2.9 kcal/mol higher than Int\textsubscript{Mg}. The activation barrier for the Mn-enzyme complex using the same active site model is calculated to be 2.0 kcal/mol higher than Mg-enzyme (see the Supporting Information), which agrees well with the experimentally measured trend discussed above. A final mechanistic note worth mentioning here is that the monodentate substrate binding mode, that is, with only the carboxylate group of 2,3-dihydroxybenzonate being coordinated to the metal, is less favored than the bidentate binding shown in Figure 3 (see also the Supporting Information).

To summarize, crystal structure elucidation and metal analysis by ICPMS/MS supported by mechanistic quantum chemical calculations have identified Mg\textsuperscript{2+} as the catalytically relevant metal in the active site of 2,3-dihydroxybenzoic acid decarboxylase from A. oryzae. This finding represents a unique example within metal-dependent o-benzoic acid decarboxylases, which generally depend on Zn\textsuperscript{2+} or Mn\textsuperscript{2+}, and is essential for future structure-activity predictions.

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**Conflict of Interest**

The authors declare no conflict of interest.
Keywords: biocatalysis • computational chemistry • enzyme structure • metal-identity • ortho-benzoic acid decarboxylase

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