Structural and Kinetic Characterization of the 4-Carboxy-2-hydroxymuconate Hydratase from the Gallate and Protocatechuate 4,5-Cleavage Pathways of Pseudomonas putida KT2440*

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The bacterial catabolism of lignin and its breakdown products is of interest for applications in industrial processing of ligno-biomass. The gallate degradation pathway of Pseudomonas putida KT2440 requires a 4-carboxy-2-hydroxymuconate (CHM) hydratase (GalB), which has a 12% sequence identity to a previously identified CHM hydratase (LigJ) from Sphingomonas sp. SYK-6. The structure of GalB was determined and found to be a member of the PIG-L N-acetylglucosamine deacetylase family; GalB is structurally distinct from the amidohydrolase fold of LigJ. LigJ has the same stereospecificity as GalB, providing an example of convergent evolution for catalytic conversion of a common metabolite in bacterial aromatic degradation pathways. Purified GalB contains a bound Zn$^{2+}$ cofactor; however the enzyme is capable of using Fe$^{2+}$ and Co$^{2+}$ with similar efficiency. The general base aspartate in the PIG-L deacetylases is wherein Glu-48 positions and activates the metal-ligated water for the hydration reaction and His-164 in the catalytic mechanism. A comparison with the PIG-L deacetylases led to a proposed mechanism for GalB wherein Glu-48 positions and activates the metal-ligated water for the hydration reaction and His-164 acts as a catalytic acid.

Lignin, a complex biopolymer found in the plant cell wall, represents a large carbon reservoir in the environment. The polymer is built from nonrepeating aromatic units connected via many different linkages. Bacterial species from the genera Pseudomonas, Comamonas, and Sphingomonas utilize the aromatic metabolites produced from lignin depolymerization, and their catabolic capabilities can therefore be harnessed for industrial processing of biomass to produce valuable products such as biofuels (reviewed in Ref. 1).

The atomic coordinates and structure factors (code 5CGZ) have been deposited in the Protein Data Bank (http://wwpdb.org).

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The abbreviations used are: PCA, protocatechuate; OMA, 4-oxalomesaconate; CHM, 4-carboxy-2-hydroxymuconate; HMG, 4-hydroxy-4-methyl-2-oxoglutarate; NTA, nitrilotriacetic acid; ICP-MS, inductively coupled plasma mass spectrometry; DDVA, 5,5′-dehydrodiva-nillic acid; cTCA, cyclohexane-1,2-dicarboxylic acid; PDB, Protein Data Bank.

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ase (TIM) barrel that is structurally distinct from other characterized divalent metal-dependent hydratases (13–15). The regions in the substrate binding pocket of LigJ are not modeled in the final structure, and accordingly, the catalytic mechanism of this enzyme has not been elucidated. Like LigJ, GalB has no sequence similarity to the previously characterized divalent metal-dependent hydratases. The initial characterization of the gallate pathway identified (2E,4E)-CHM as the substrate for GalB, whereas (2Z,4E)-CHM is the proposed substrate for LigJ (8, 9). The proposed differences in substrate specificity between the two enzymes would give a biological rationale for the two possibly distinct CHM hydratases. However, no structural or mechanistic investigations have been reported in the literature to elucidate the mechanism of GalB and its relationship to LigJ CHM hydratases.

Herein we report the first structure and in-depth kinetic characterization of the GalB CHM hydratase from Pseudomonas putida KT2440. GalB, unlike LigJ, adopts a Rossmann-like fold that resembles the PIG-L deacetylase family of hydrolytic enzymes. A comparative analysis of PmdE from Comamonas sp. strain E6 (herein referred to as LigJ<sub>C</sub>) and GalB indicates that both enzymes in fact utilize the same CHM isomer leading to the same (−)-CHA enantiomer product. LigJ<sub>C</sub> shares 63 and 80% sequence identity to LigJ from Sphingomonas sp. SYK-6 and R. palustris, respectively. We also show that both GalB and LigJ<sub>C</sub> catalyze the reversible hydration of 4-carboxy-2-hydroxypenta-2,4-dienoate (CHPD; a CHM analog lacking the C5 carboxylate) to HMG and propose a previously undescribed route for HMG production from the metabolism of the lignin metabolite 5,5′-dehydrodivanillate (DDVA). Together, our findings shed light on the analogous enzymes GalB and LigJ, which have evolved convergently to perform the same chemical reaction.

**Experimental Procedures**

*Chemicals*—L-Lactate dehydrogenase (rabbit muscle), L-malate dehydrogenase (porcine heart), and DOWEX<sup>®</sup> 1X8-200 ion-exchange resin were from Sigma-Aldrich. Restriction enzymes and *Pfu* polymerase were from Invitrogen or New England Biolabs. All other chemicals were analytical grade and were obtained from either Sigma-Aldrich or Fisher Scientific.
DNA Manipulations—pET29a plasmids harboring galA, galB, and galD from *P. putida* KT2440 were a generous gift from E. Díaz, Biological Research Center-CSIC, Madrid, Spain (9). GalB variants A16D, R21A, E48A, R67A, H127A, Y123A, H164A, and R216A were produced by site-specific mutagenesis from E. Díaz, Biological Research Center-CSIC, Madrid, Spain. Primer sequences utilized for mutagenesis and the creation of the GalBNt construct are available upon request. For study were recombinantly overexpressed in *galB* Supercenter (University of Guelph).

were harvested by centrifugation at 5000 *g* for 15 min. Protein Purifications—Buffers containing 20 mM sodium-HEPES, pH 7.5, were used throughout each purification procedure unless indicated otherwise. Each cell pellet was resuspended in buffer and disrupted by French press at 1200 p.s.i. Cell debris was removed by centrifugation (17,500 × g for 15 min) and supernatants filtered through a 0.45-μm filter.

The purification of GalA, GalB, and GalD variants was carried out by chromatography on an AKTA Explorer 100 (GE Healthcare Life Sciences). All column resins were from GE Healthcare Life Sciences, and column chromatography utilized a SourceTM 15Q anion-exchange column (2 × 13 cm), phenyl-SepharoseTM hydrodynamic interaction column (1 × 18.5 cm), and a HiLoad 26/60 Superdex 200 gel filtration column (2.6 × 60 cm).

For purification of GalB and variants, crude extract was loaded onto the anion-exchange column, and the column was washed with 2 column volumes of buffer followed by a linear gradient of NaCl from 0.0 to 0.3 M over 12 column volumes. GalB was eluted with ~0.15 M NaCl. Active fractions were pooled and concentrated to ~10 ml by ultrafiltration using a YM10 filter (Millipore, Nepean, Ontario, Canada). Ammonium sulfate was added to the concentrated protein extract to a final concentration of 1.0 M, and the extract was loaded onto the hydrophobic interaction column, which was pre-equilibrated with 1.0 M ammonium sulfate. A 12-column volume linear gradient from 1.0 to 0.0 M ammonium sulfate was applied, and GalB was eluted in ~0.5 M ammonium sulfate. Active fractions were pooled, concentrated to ~2 ml by ultrafiltration, and loaded onto the gel filtration column. Protein was eluted using an isocratic flow of buffer containing 0.15 M NaCl. The purified enzyme was dialyzed in the buffer to remove salt, concentrated to ~40 mg protein/ml by ultrafiltration, and stored at ~80 °C.

For purification of GalA, crude extract was loaded onto the anion-exchange column, and the column was washed with 2 column volumes of buffer containing 0.1 M NaCl followed by a linear gradient of NaCl from 0.1 to 0.45 M over 12 column volumes. GalA was eluted with ~0.25 M NaCl. Active fractions were pooled and concentrated to ~2 ml by ultrafiltration. GalA did not bind sufficiently to the hydrophobic resin, and the concentrated sample was loaded onto the gel filtration column with the protein eluted with an isocratic flow of buffer containing 0.15 M NaCl. The purified enzyme was dialyzed into 20 mM MES buffer, pH 6.5, containing 50 μM (NH₄)₂Fe(SO₄)₂ with 250 μM DTT and concentrated to ~20 mg protein/ml by ultrafiltration. The protein solution was flushed with N₂ and degassed before being flash-frozen in liquid N₂. The frozen protein was stored at ~80 °C.

For purification of GalD and LigJ*E*6, the crude extract was incubated with Ni²⁺-nitrilotriacetic acid (NTA) affinity resin containing 10 mM imidazole at 4 °C for 16 h with constant mixing. The mixture was then passed through a gravity column, retaining the Ni²⁺-NTA beads. The Ni²⁺-NTA resin was washed with 20 ml of buffer containing 15 mM imidazole, and protein was eluted with buffer containing 250 mM imidazole. The polyhistidine tag on LigJ*E*6 was removed by first changing the buffer to 50 mM sodium phosphate, pH 8.0, containing 100 mM NaCl and 10% glycerol by ultrafiltration using a YM10 filter. The His₆-LigJ*E*6 was diluted to ~10 mg/ml and incubated with 2 units of thrombin for 16 h at 15 °C. The thrombin was removed by incubation with p-aminobenzimide-agarose for 1 h at 15 °C, and the beads were subsequently removed by filtration through a 0.2-μm filter. The cleaved His₆ tags, and any tagged protein remaining after the thrombin incubation, were removed by incubation with fresh Ni²⁺-NTA for 1 h at 15 °C and subsequently passed through a 0.2-μm filter to remove Ni²⁺-NTA beads. GalD contains an internal thrombin cleavage site at residue 239 of the 361-residue protein, and thus removal of the His₆ tag was not attempted. The purified protein was concentrated by ultrafiltration and stored at ~80 °C in 20 mM HEPES, pH 7.5.

The molecular mass of the GalA, GalB, GalD, and LigJ*E*6 subunits, as determined by SDS-PAGE, was ~45, ~25, ~40, and ~40 kDa (data not shown), which is consistent with the predicted molecular mass values of 47.6, 27.5, 37.6, and 38.2 kDa, respectively, from previous reports (9, 10, 19, 20).
Structure and Characterization of the GalB CHM Hydratase

Determination of Protein Concentration, Purity, and Molecular Mass—Protein concentrations were determined by the Bradford assay using bovine serum albumin as the standard (21). SDS-PAGE was performed and stained with Coomassie Blue according to established procedures (22). The molecular weight of the GalB holoenzyme was determined by gel filtration using a HiLoad 26/60 Superdex 200 prep column (2.6 × 60 cm).

Crystallization and Structure Determination of GalB—Wild type GalB was screened using a Qiagen Classic suite and yielded crystals in which diffraction did not exceed 4.0 Å resolution. The secondary structural prediction using Phyre 2.0 indicated that the N-terminal four residues of the protein are likely to be unstructured and may contribute to disorder in the crystallization (23). An enzyme variant lacking the four N-terminal residues (GalBNt) was screened, and the conditions were optimized. The crystallization condition contained 15% glycerol in the reservoir solution with 2 μl of the reservoir solution mixed with 2 μl of 80 mg/ml purified GalBNt. Cubic crystals grew within a month at 4 °C and were diffracted to ~3.0 Å resolution. Crystals left for ~10 months to natively dehydrate at 4 °C were soaked in 50% glycerol for 1 min before being flash-frozen in liquid nitrogen. More than 20 of these crystals were screened with few of them diffracting beyond 2.5 Å resolution and only one diffracting to 2.1 Å resolution. The bestdiffraction crystal was utilized for both the native and anomalous data collections.

Data were collected at the Canadian Light Source. A fluorescence scan revealed a strong zinc signal and only minimal signals for other elements (data not shown). The zinc, iron, and cobalt K-edges were scanned; however, only the zinc K-edge showed scattering factors consistent with the presence of the metal in the crystal. A data set was collected to 2.0 Å at a wavelength of 0.97949 Å, on the Canadian Light Source 08ID-1 beamline. An attempt to solve the structure by molecular replacement using PIG-L deacetylase structures as search models was unsuccessful. Thus, an anomalous data set for phasing was collected up to 2.2 Å on the same crystal at the wavelength 1.2817 Å (zinc K-edge), on the Canadian Light Source 08B1-1 beamline. The crystal was of the cubic space group P4₁,32, with cell dimensions a = b = c = 201.66 Å. Diffraction data were processed in XDS, and the structure was solved in Phenix (24, 25). The crystal had significant radiation damage by the end of the native data set collection, but Autosol was able to find two zinc sites in the anomalous data set (26). An initial model was built and refined from the anomalous data in Phenix using Autobuild and Refine (27, 28). The native structure was determined by molecular replacement with Phaser using one protomer of the structure from the anomalous data as the search model (29). The structure was rebuilt in Coot, and further refinement was completed in Phenix Refine (28, 30). NCS restraints were not used during refinement, but TLS atomic displacement parameters (with the protein subdivided into five residue groups) were utilized in the final stages of refinement (31). Table 1 lists data collection and final model refinement statistics. The representations of the structure of GalB presented in Figs. 2–4 were prepared with PyMol v1.4.1 (Schrödinger, LLC).

Preparation of 4-Carboxy-2-hydroxyxymonconate—CHM was synthesized enzymatically from gallic acid using the gallate dioxygenase GalA and the OMA tautomerase GalD. 5-ml reactions were set up in 750 mM sodium phosphate buffer, pH 7.0, containing 150 mM gallic acid. 300 μg of both GalA and GalD was added every 5 min for 50 min with the solution being mixed constantly. After the final addition, the reaction was further incubated with mixing for an additional 30 min. Reactions took place in a closed cell, and throughout the course of the reaction the solution was kept oxygenated by bubbling oxygen through the solution. At the completion of the reaction the enzymes were removed by ultrafiltration through a YM10 filter. The solution was treated with Chelex, sodium form, for 10 min, which was subsequently removed by filtration through a 0.2-μm filter. By the end of the reaction >95% of the gallic acid was catabolized and >65% of the product was CHM, as determined by kinetic assays. CHM was purified by HPLC on an Åkta Explorer 100 (GE Healthcare Life Sciences) with 500-μl aliquots of the reaction applied to an Aminex fast acid ion-exchange column, HPX-87H (100 × 7.8 mm) and was separated with an isocratic elution using 100 mM H₂SO₄. Compounds were detected at 215 nm; using this method CHM, OMA, and gallic acid were resolved to retention volumes of 6.5, 8.0, and 22.5 ml, respectively. When purified CHM was incubated with GalB, converting the CHM to CHA, both the CHA and CHM co-eluted from the column using the isocratic flow of 100 mM H₂SO₄. Changing the procedure to an isocratic flow of 500 mM H₂SO₄ facilitated the separation of CHA from CHM, with the compounds having retention volumes of 6.7 and 9.8 ml, respectively.

Preparation of CHPD—CHPD was synthesized enzymatically with GalB. 5-ml reactions comprised 50 mM (R/S)-HMG, 50 μM CoCl₂, and 100 μg of GalB in 100 mM HEPES, pH 7.0. Reactions were incubated at 15 °C overnight with constant mixing. Reactions were stopped by removing the enzyme by ultrafiltration through a YM10 filter. The reaction was subsequently treated with Chelex, sodium form, for 1 h to remove metals, with the beads removed by filtration through a 0.2-μm filter. 500-μl samples of the reaction were separated by HPLC via an Aminex fast acid ion-exchange column using 100 mM H₂SO₄. HMG and CHPD were resolved to retention volumes of 6.4 and 9.0 ml, respectively.

GalB Metal Analysis—To assess the metal utilized by the enzyme in vivo, GalB expressed in and purified from P. putida KT2442 was analyzed by inductively coupled plasma mass spectrometry (ICP-MS). The enzyme was purified as described previously, and the purified protein was washed with Chelex (sodium form)-treated buffer several times in an ultrafiltration stir cell using a YM10 filter to remove exogenous metals. The final protein sample was concentrated to 5 ml in buffer before being diluted to 125 ml in HPLC-grade water (Fisher Scientific) and sent for ICP-MS analysis at ALS Global (Waterloo, Ontario, Canada). The final concentration of enzyme analyzed by ICP-MS was 24.1 μM. The detection limit for iron (9.0 μM) with the ICP-MS methodology utilized was too low relative to the stoichiometric amounts of enzyme supplied to draw conclusions about its presence. Thus the iron content of the GalB-purified proteins was assessed using a ferrozine assay (32, 33).

Protein Metal Ion Substitution—All buffers were treated with Chelex, sodium form, for 20 min to remove exogenous metals,
with the beads removed by filtration through a 0.45-μm filter. Metal-free apoenzymes were prepared by treating 50 mg of purified enzyme in 50 ml of 100 mM sodium-HEPES, pH 7.0, with 1 mM EDTA for 12 h at 4 °C with constant mixing. Using ultrafiltration with YM10 filter, the EDTA was removed from the enzyme solution by successive washes with buffer, and the final apoprotein was concentrated to a total volume of 600 μl for a minimum of 1 h at 15 °C prior to use in the kinetic assays. For incubation with Fe²⁺, a 200-μl solution comprising 0.92 μg/ml apoenzyme was transferred to a Baker Ruskinn Bugbox anaerobic chamber filled with nitrogen and degassed for 5 min. The Fe²⁺ enzyme solutions were made up in the anaerobic chamber to 50 μM (NH₄)₂Fe(SO₄)₂ from a freshly degassed stock of 5 mM (NH₄)₂Fe(SO₄)₂, prepared in 50 mM H₂SO₄ and incubated on ice for 1 h. Less than 3% of the specific activity was lost after uncapping and exposing the Fe²⁺ anaerobically incubated enzyme sample to air for 1 h.

**Zinc and Cobalt Dissociation Constant Determination**—Apoprotein of either the wild type or variants of GalB was incubated with varying concentrations of either ZnSO₄ or CoCl₂ in a total volume of 600 μl of 100 mM HEPES, pH 7.0, and was kept at 15 °C for 1 h with constant mixing. Binding reactions were then filtered by ultrafiltration using an Amicon Microcon YM10 to remove enzyme from the solution. Samples of the enzyme-free solutions were mixed with 250 μM freshly prepared 4-(2-pyridylazo)resorcinol and made up to 1 ml with Chelex (sodium form)-treated buffer. The amount of free Co²⁺ or Zn²⁺ present in the reaction was determined in at least duplicate spectrophotometrically at 505 nm for Co²⁺ and 497 nm for Zn²⁺ using extinction coefficients described previously (34). The concentration of enzyme-bound Zn²⁺ or Co²⁺ was calculated using Equation 1, and dissociation constants were determined by nonlinear regression in GraphPad Prism with Equation 2, where [M²⁺] is concentration of metal and C is the saturated concentration of the metal.

\[
[M^{2+}]_{\text{bound}} = [M^{2+}]_{\text{total}} - [M^{2+}]_{\text{free}} \tag{Eq. 1}
\]

\[
[M^{2+}]_{\text{bound}} = \frac{C[M^{2+}]_{\text{free}}}{K_d + [M^{2+}]_{\text{free}}} \tag{Eq. 2}
\]

**Enzyme Assays**—All of the kinetic assays were performed at least in duplicate at 25 °C using a Varian Cary 3 spectrophotometer equipped with a thermostatted cuvette holder. Assays for CHM hydration and CHA dehydroylation were completed similarly to that described previously (9). Briefly, the hydration of CHM to CHA, or the dehydration of CHA to CHM, was monitored by detecting the olefinic signal of CHM at 265 nm. The hydration of CHPD to HMG, or the dehydroylation of HMG to CHPD, was similarly monitored at 260 nm, where UV scans indicate the maximum absorbance of the CHPD olefinic signal. Standard assays were completed in 100 mM HEPES, pH 7.5, and the extinction coefficients of 9451 M⁻¹cm⁻¹ utilized for CHM and 8221 M⁻¹cm⁻¹ for CHPD were determined experimentally. All assays were carried out with apoGalB preincubated for at least 1 h with the stated metal at 50 μM prior to the assay. Metal ion specificity assays were completed in 100 mM HEPES, pH 7.0, with 50 μM CHM and 50 μM metal chloride, EDTA, or with enzyme prior to metal substitution.

The efficiency of the removal of bound metals by EDTA was assessed using enzyme activity assays. ApoGalB (1.5 μM) that had been preincubated with 50 μM ZnCl₂ was incubated with either 250 or 1000 μM EDTA, and the specific activity of the enzyme toward 50 μM CHM under standard conditions was measured at successive time points. The loss of activity in the enzyme solutions was fit to a one-phase exponential decay, Equation 3, where SA is the specific activity, SA₀ is the initial specific activity, k is the rate constant, and t is time.

\[
SA = (SA_0) e^{-kt} \tag{Eq. 3}
\]

The pH dependence of CHM utilization kinetics was determined with pH varying between pH 5.5 and 9.5 in a three-component constant ionic strength buffer containing 0.1 M Tris, 0.05 M acetic acid, and 0.05 M MES. Enzyme was preincubated with 50 μM ZnCl₂ or 50 μM CoCl₂ in the stated pH buffer for a minimum of 30 min before assays were completed. The extinction coefficient for CHM at each pH point was determined experimentally. All data were fitted by nonlinear regression in GraphPad Prism. The pH profile of k_cat shows a single ionization event and is fitted to Equation 4. The pH profile of K_m is bell-shaped with slopes of unity and is fitted to Equation 5. The pH profile of k_cat/K_m is bell-shaped, with the ascending limb of the curve having a slope of ~2 and a descending slope of unity, and is fitted to Equation 6.

\[
k_{\text{cat}} = \frac{(k_{\text{cat}})_{\text{max}}}{(1 + [H^+] / K_a)} \tag{Eq. 4}
\]

\[
K_m = \frac{(K_m)_{\text{max}}}{(1 + [H^+] / K_b)} \tag{Eq. 5}
\]

\[
k_{\text{cat}}/K_m = \frac{(k_{\text{cat}} / K_m)_{\text{max}}}{(1 + [H^+] / K_a)(1 + [H^+] / K_b)} \tag{Eq. 6}
\]

Polarimetry assays were completed in a Rudolph IV polarimeter using a 100-mm cuvette. Reactions of 7 ml contained 15 mM racemic CHA, 30 mM racemic HMG, or 5 mM CHM in 100 mM HEPES, pH 7.5, and 0.62, 8.7, or 0.10 μM GalB preincubated with 50 μM CoCl₂, respectively. Reactions were initiated with the addition of enzyme, and the solution was passed through a 0.2-μm filter prior to data acquisition.

**Results**

**Expression and Purification of GalB**—The gene for GalB from *P. putida* KT2440 was overexpressed both natively in *P. putida* KT2442 and recombinantly in *E. coli* BL21(ADE3) cells with typical yields of 15 and 22 mg of purified protein/liter of bacterial culture, respectively. The subunit molecular mass of the enzyme as determined by SDS-PAGE is 27 kDa, consistent with the predicted molecular mass (27,461 Da). GalB variants and the protein containing an N-terminal truncation were overex-
**Structure and Characterization of the GalB CHM Hydratase**

### Table 1

**GalB<sub>Nt</sub> data collection and refinement statistics**

Data shown in parentheses are for the highest resolution shell.

|                  | Native | Zn-K edge |
|------------------|--------|-----------|
| **Data collection** |        |           |
| Space group      | P4, 32 | P4, 32    |
| Unit cell: \(a = b = c\) (Å) | 201.66 | 201.66    |
| Wavelength (Å)   | 0.97949| 1.2817    |
| Resolution (Å)   | 48.83-2.10 (2.18-2.10) | 9.84-2.20 (2.28-2.20) |
| No. observed reflections | 638,492 (63,458) | 1,539,423 (92,226) |
| No. unique reflections | 81,109 (7,977) | 135,263 (10,008) |
| Redundancy       | 7.9    | 21.5      |
| Data completeness (%) | 99.8 (99.9) | 99.9 (99.9) |
| **Model refinement** |        |           |
| \(R_{work} / R_{free}\) (%) | 3.1 (37.5) | 1.9 (24.3) |
| Wilson average atomic displacement parameters (Å²) | 99.9 (79.0) | 99.9 (75.5) |
| Overall          | 35.5   | 37.5      |

**Overall Structure of GalB**—An initial substructure of GalB<sub>Nt</sub> was determined by zinc single-wavelength anomalous diffraction (SAD) phasing to 2.2 Å resolution, which was then subsequently utilized as a model for molecular replacement of the native data set (2.1 Å) (Table 1). There are two protomers in the asymmetric unit, which can be superimposed with a root-mean-square deviation of 0.087 Å, indicating only minor differences. Each protomer has an overall fold similar to the PIG-L family of N-deacetylases, described in more detail below, consisting of an \(\alpha\)-\(\beta\)-\(\alpha\) sandwich with the N-terminal portion forming a five-stranded Rossmann fold (residues 1–160) leading to a mixed \(\alpha/\beta\) C-terminal region (residues 161–234) and a \(\beta\)-strand (\(\beta_8\) residues 237–240) (Fig. 2A). The N-terminal Rossmann fold is composed of a \(\beta\)-sheet (\(\beta_1\)–\(\beta_3\)) wrapped by \(\alpha_4\) and \(\alpha_5\) on one face and \(\alpha_1\) and \(\alpha_2\) on the other. The C-terminal portion consists of two anti-parallel \(\beta\)-strands (\(\beta_6\)–\(\beta_7\)) interspersed with a helical hairpin (\(\alpha_6\) and \(\alpha_3\)). The \(\beta_6\) from an adjacent monomer packs parallel to \(\beta_6\) making a continuous \(\beta\)-sheet of \(\beta_2\)–\(\beta_3\)–\(\beta_4\) with the \(\beta_6\) anti-parallel to the rest. The protomer is composed of the Rossmann fold, the C-terminal region, and contributions by \(\alpha_6\)’ and \(\beta_6\)’ from adjacent units. The bound zinc ion is found at the bottom of a ~22 Å-deep, solvent-accessible pocket that has a total volume of ~1060 Å³ as determined by DogSiteScorer (35).

**Oligomeric State**—The interface between the two protomers in the asymmetric unit constitutes 125 Å² (~1%) of each molecules surface area, as determined by PISA, and is likely not a physiologically relevant interaction (36). Instead, each protomer acts as the asymmetric fraction of a distinct hexamer, point group \(D_5\) (Fig. 2B). The hexamer is assembled by interactions along three distinct interfaces: dimerization through packing of \(\beta_6\) on \(\beta_6\)' of an adjacent protomer along a 2-fold axis, which buries ~13% of the total surface area (Fig. 2C); dimerization through packing of \(\alpha_5\)/\(\alpha_5\)' with \(\alpha_6\)/\(\alpha_6\)' along a 2-fold axis, which buries ~9% of the total surface area (Fig. 2D); and packing of \(\alpha_3\) and the loop between \(\beta_5\) and \(\alpha_5\) against \(\alpha_2\)' and the loop region between \(\alpha_3\)' and \(\beta_4\)' related by a 3-fold axis, which buries ~6% of the total surface area (Fig. 2E). Together with additional interactions, the hexameric oligomerization buries ~50% of the total surface area on the protomer, indicating that the hexamer is the biological unit. Analysis of the GalB protein by gel filtration yields a mass of 156.2 kDa, consistent with a hexameric organization (164.8 kDa; data not shown).

**GalB Metal Binding Site**—Scans of the GalB crystal revealed fluorescence consistent with zinc being the only transitional metal present in the crystal. The GalB zinc ion is coordinated in a skewed trigonal bipyramidal fashion by His-14 N\(\delta\), Asp-17 O\(\delta\), Glu-48 O\(\varepsilon\), His-127 N\(e\), and a water molecule (Fig. 3).

Three conformers of the Glu-48 residue are seen in the two chains. The predominant conformer, with occupancy of ~60%, is found in both chains with the O\(\varepsilon\) of Glu-48 coordinating the zinc ion. The alternate conformation of the residue is slightly different between the two chains. In chain B the residue is flipped, having the O\(\varepsilon\) coordinating the zinc ion and the O\(\varepsilon\) positioned by interaction with Asn-124 N\(\delta\) and the Tyr-90 hydroxyl (Fig. 3A). In chain A the carboxylate rotated away from the metal ion, with the O\(\varepsilon\) interacting with Asn-124 N\(\delta\)...
and the hydroxyl of Tyr-90, whereas the Oe₂ projects out of the metal binding site (Fig. 3C). The average atomic displacement parameter for the Glu-48 is ~34.5 Å², which is similar to the atomic displacement parameter of the other metal binding residues, indicating that the residue is well ordered.

The coordinate distance of the water to the metal (2.3 Å in chain A and 2.8 Å in chain B) is longer than that of the protein ligands (~2.1 Å) to the metal (Fig. 3B). The metal ligating water is also positioned by interaction with the Oδ₂ of Asp-17 (2.8 Å). The water is in close proximity to the Glu-48 Oe₁ (2.1 Å), but the water is not in line with the atom for ideal interaction, suggesting that the Glu-48 does not interact directly with the water. The average atomic displacement parameter for the metal ligating water is 51.7 Å², indicating that the water is possibly present at less than full occupancy and its presence may be modified by the different conformers of the Glu-48.

Outside of the metal binding residues, the pocket is lined with hydrophilic residues with several residues (such as Arg-21, Arg-67, His-164, and Arg-216’ from the α8’ on the adjacent protomer) providing electropositive character to the pocket and likely facilitating the binding of the tricarboxylic acid substrate, CHM (Fig. 4A).

Structurally Related Proteins—A search of structural homologs to GalB using DALI revealed members of the PIG-L family of N-deacetylases as the closest structural homologs (37). There are currently eight structures of unique PIG-L family members in the protein database, and GalB aligns with these proteins with root-mean-square deviation values of <3.0 Å resulting in Z-scores of >20 and coverage of >70% of the deacetylase sequence. Members of this family, such as the N-acetyl-1-D-myo-inositol-2-amino-2-deoxy-α-D-glucopyranoside deacetylase from Mycobacterium tuberculosis (MshB), N,N'-diacetylchitobiose deacetylase from Pyrococcus species, and N-acetylglucosaminypseudoaglycone deacetylase from Actinoplanes steichomyceticus, are divalent metal-dependent hydrolytic enzymes with preference for zinc or iron (II) ions (38–40). Oligomerization differs across the PIG-L family of proteins with variability in the both GalB N- and C-terminal portions,
resulting in proteins predicted or observed to exist as hexamers, trimers, or monomers (40–43). Common among all of the PIG-L deacetylases and GalB is the single /H9251/H9252 domain with a Rossmann-like fold containing a solvent-accessible pocket. With the exception of the Glu-48 residue, zinc binding in these structural homologs is facilitated by a His-X_/Asp-X_/His motif similarly found in GalB (for GalB: His-14-X_/Asp-17-X_/His-127) (Fig. 3C). MshB is the only PIG-L family member for which the structure has been determined that contains a glutamate in the primary sequence aligned to GalB Glu-48. However, the
equivalent MshB residue is found on a loop region which is rotated away from the bound zinc ion and active site (43, 44).

The bound zinc ion in GalB is exposed on one face to a solvent-accessible pocket, which is similar to the PIG-L family and is the substrate binding pocket in these enzymes. Outside of the metal binding residues, only the GalB residues Arg-67 and Gln-196 are conserved in the binding pocket among all PIG-L N-deacetylases.

In MshB (and other PIG-L deacetylases), the active site residues His-144, Asp-15, and Tyr-142 are implicated as the general acid, general base, and oxyanion transition state stabilizer, respectively (45). The equivalent residues in GalB are Asn-124,
Structure and Characterization of the GalB CHM Hydratase

Ala-16, and Tyr-123, respectively (Fig. 4A), implying that the general base and acid roles of the PIG-L deacetylases at least are not recapitulated by these residues. The hydration of CHM to CHA is likely to proceed through deprotonation of the CHM enol, producing an enolate that is analogous to the oxynion transition state in the PIG-L deacetylases. The MshB oxynion stabilizing the Tyr-142 residue is found in a glycine-rich loop (Gly-140/Gly-141/Tyr-142/Gly-143), which provides conformational flexibility to allow the positioning of Tyr-142 with its side chain projecting into the active site pocket. The GalB Tyr-123, however, is not found in as flexible a region (Asp-121/Pro-122/Tyr-123/Asn-124), and the phenol side chain is observed rotated away from the active site pocket. The GalB Tyr-123 hydroxyl is stabilized by interaction with the Nε and Oε of Gln-217 with the tyrosine π electrons interacting with the Nε of Gln-165. Although some movement is likely to occur upon substrate binding, the lack of conformational freedom and interactions made by the GalB Tyr-142 suggests that it is unlikely to rotate into the active site pocket and fulfill the enolate stabilization role. Thus, although conserving the protein fold and metal ion binding, GalB lacks key residues that in PIG-L enzymes facilitate hydrolase activity, likely reflecting differences in the chemical reaction of the enzymes.

Metal and Kinetic Analysis of GalB—To ascertain the native metal ion utilized by GalB in vivo, protein purified from the native host, P. putida KT2442, was analyzed. When overexpressed and purified, using three chromatographic steps, the specific activity of the native protein was 1.02 μmol min⁻¹ μg⁻¹. The specific activity of the native protein increased 4-fold when 50 μM exogenous Co²⁺ was added, indicating that either the native enzyme was not fully saturated with metal during overexpression or else the metal was lost during purification. ICP-MS analyses on the native purified protein, without the addition of any exogenous metals, revealed the presence of zinc, cobalt, and copper in relative stoichiometric quantities of 23.0.1, and 0.1%, respectively. Because of the high background, amounts of iron could not be estimated by ICP-MS. Instead, Fe was determined by the ferrozine assay to be below 0.6%. Thus zinc is the predominant metal ion found in the enzyme through the aerobic purification methods utilized here.

The GalB N-terminal truncation variant used for crystallization maintained the same activity toward CHM as the wild type enzyme, indicating that the observed structure represents the fully active zinc bound enzyme. A metal cofactor is required for enzyme activity, and the half-life of Zn²⁺-saturated GalB enzyme activity when treated with either 250 μM or 1 mM EDTA was 32.3 or 17.1 min, respectively. After treatment with 1 mM EDTA for 12 h, the protein lacked detectable enzyme activity above the error of the assay conditions. CHM hydratase activity could be restored by incubating the apoenzyme with divalent metals (Table 2). The enzyme showed maximal activity with Fe²⁺ and Co²⁺. The presence of all other metal ions, including Zn²⁺, resulted in less than 30% of the activity observed with Fe²⁺.

The metal binding ligands observed in GalB are the same as those observed in the PIG-L deacetylase family, except that Glu-48 in GalB represents a potential extra ligand; however this residue shows alternative orientations, implying that its interactions with the metal ion are weaker than the other three ligating residues (Fig. 3). Both Zn²⁺ and Co²⁺ bind strongly to GalB with dissociation constants (K_d) of 0.63 ± 0.05 and 0.56 ± 0.009 μM, respectively (Table 3). The H127A variant showed a 28- and 1625-fold increase in K_d for Zn²⁺ and Co²⁺, respectively. The E48A variant, however, showed only a 5- and 26-fold increase in K_d for Zn²⁺ and Co²⁺, respectively. The effects of the Glu-48 substitution were less pronounced than that of the substitution of His-127, a conserved metal ligand in the PIG-L family, therefore suggesting a minimal role for Glu-48 in metal binding.

In the GalB CHM hydration steady state kinetics, the choice of metal ion (among Fe²⁺, Co²⁺, and Zn²⁺) affected the specificity constant (K_cat/K_m) by less than 3-fold (Table 4). However, the reaction rates with the metals at high substrate concentrations diverged markedly, with both K_cat and K_m being ~7-fold higher for Fe²⁺ or Co²⁺ versus Zn²⁺; the highest catalytic activity was therefore observed with Fe²⁺, but the increase in K_cat was offset by a parallel increase in K_m. GalB specificity was >200-fold lower for CHPD, indicating an important role for the CHM C5 carboxylate group in substrate binding. The enzymatic reaction is reversible, with the dehydration of CHA and HMG being catalyzed with kinetic parameters similar to the hydration reaction. Polarimetric analysis of the GalB-catalyzed hydration of CHM indicates that only the (−)-enantiomer of CHA was produced (Fig. 5). Similarly, the enzyme was found to dehydrate only the (−)-enantiomers of both CHA and HMG in

| Metal ion | Relative activity |
|-----------|------------------|
| Fe²⁺      | 100 ± 4.6        |
| Co²⁺      | 79 ± 4.0         |
| Native + Co²⁺ | 34 ± 1.5     |
| Mn²⁺      | 30 ± 1.9         |
| Zn²⁺      | 17 ± 0.66        |
| Fe²⁺      | 14 ± 2.2         |
| Ni²⁺      | 13 ± 0.33        |
| Ca²⁺      | 11 ± 1.0         |
| Mg²⁺      | 9.8 ± 0.66       |
| Native    | 8.6 ± 0.95       |
| Cd²⁺      | 7.4 ± 0.66       |
| Apo       | 4.0 ± 0.33       |
| Co²⁺      | 0.19 ± 0.10      |
| EDTA      | 0.030 ± 0.0033   |

| Protein  | Zn²⁺     | Co²⁺     |
|----------|----------|----------|
| Wild type| 0.63 ± 0.048 | 0.056 ± 0.0085 |
| A16D     | 0.75 ± 0.095 | 0.071 ± 0.0043 |
| E48A     | 3.2 ± 0.29  | 1.5 ± 0.19  |
| H127A    | 17.6 ± 1.4  | 90 ± 10    |
| H14A/D17A| NS        | NS        |
the reverse reaction, leaving the positive enantiomers, when exposed to racemic mixtures of the substrates.

**CHM and GalB Hydration pH Profiles**—The pH dependence of the GalB-catalyzed hydration of CHM was assessed in a three-component buffer. The extinction coefficient of CHM increases sigmoidally with increasing pH, likely because of the deprotonation of the enol group leading to increased electron delocalization and chromogenicity. Fitting the increase in extinction coefficient to an acid titration resulted in a $pK_a$ value of 7.3 ± 0.1 for the CHM substrate (Fig. 6A).

The $pK_{a}$ values for the other ionizable groups of the CHM (the C2, C4, and C5 carboxylates) were determined by titration. CHM was isolated by chromatography in 100 mM H$_2$SO$_4$ and could not be further purified, hampering the definition of $pK_a$ values that were close to the second $pK_a$ of sulfuric acid (~2.0). The titration of the CHM in H$_2$SO$_4$ indicated two ionizable groups with $pK_{a}$ values around 3.5 and two values of about 6 and 7 (data not shown). Considering the $pK_{a}$ values of 7.3 determined for CHM using absorption spectroscopy as described above, the $pK_{a}$ value for the other ionizable group fitted to the titration curve was estimated to be ~6.3.

Kinetic parameters for the CHM hydration reaction could not be determined for pH values below 5.5, and those above 9.5 could not be determined due to high $K_m$ values for CHM. The dependence of pH on the Zn$^{2+}$ and Co$^{2+}$ enzyme-catalyzed reactions was assessed, and the profiles for each were similar (Table 5). In both cases, the pH profile of log($k_{cat}$) shows a single ionization event in the enzyme-substrate complex, with a $pK_a$ value of 6.1 ± 0.1 and 6.2 ± 0.1 for the Zn$^{2+}$ and Co$^{2+}$ reactions, respectively (Fig. 6C).

The pH profiles of $K_m$ are bell-shaped, with the ascending and descending limbs having slopes of unity (Fig. 6D). The ascending $pK_a$ values for both metal cofactor-catalyzed reactions is 7.3 ± 0.2, with the descending $pK_a$ values being 8.7 ± 0.2 for Zn$^{2+}$ and 8.5 ± 0.3 for Co$^{2+}$. The close proximity of the ascending $pK_a$ and $pK_{a}$ values as well as the observation of the same ascending $pK_a$ in both the Zn$^{2+}$ and Co$^{2+}$ dependences, suggests that the $pK_a$ in the $K_m$ plot likely represents the free substrate rather than the free enzyme.

The pH profiles of log($k_{cat}/K_m$) are also bell-shaped but have the ascending limb of the curve with a slope of ~2 and a descending slope of unity (Fig. 6D). For both metal ions, fitting of the data to Equation 3 gives rise to two $pK_a$ values on the ascending limb with each being ~6.7 and one $pK_a$ on the descending limb being ~8.4. However, fixing one of the ascending $pK_a$ values as $pK_{a}^{mod}$, as observed for $K_m$, results in $pK_a$ values of 6.2 ± 0.2 and 8.6 ± 0.1 for the Zn$^{2+}$ cofactor profile and 6.4 ± 0.2 and 8.3 ± 0.2 for the Co$^{2+}$ cofactor profile. For both cofactor-supported reactions, the ascending $pK_a$ values are the same, and the descending $pK_a$ values are within close proximity to each other, suggesting that conserved groups contribute to these ionization values, which are only moderately affected by the metal cofactor. The two $pK_{a}$ values of ~7.3 and ~6.3 may correspond to the two $pK_a$ values observed in the ascending slope of the pH profile of log($k_{cat}/K_m$). However, the proximity of the lower $pK_a$ from the log($k_{cat}/K_m$) profile (~6.3) to the $pK_{a}^{mod}$ from the pH profile of log($k_{cat}$) (~6.2) suggests that the value represents an ionizable group on the free enzyme rather than the free substrate.

The lack of an ionizable group on the free substrate above pH 8 suggests that the $pK_a$ of the descending bell curves observed in the pH profile of $K_m$ (~8.7 for Zn$^{2+}$ and ~8.5 for Co$^{2+}$) and
substrate was reduced 105-fold relative to the wild type enzyme, and log

indicating the importance of a small nonpolar residue at this position for hydratase function. Neither the wild type GalB nor the A16D variant exhibited inhibition by glucosamine or N-acetyl glucosamine up to 1 mM in the CHM hydration reaction. In the PIG-L deacetylase MshB, Tyr-142 is proposed to act as an oxyanion stabilizing residue. As described previously, GalB Tyr-123 aligns with Tyr-142 of MshB in the primary sequence, but the GalB residue is differently positioned, with the phenolic side chain rotated away from the active site pocket (38, 44). Accordingly, the GalB Y123A variant resulted in no significant change in CHM utilization relative to the wild type protein (Table 4). Together, the results indicate that the catalytic mechanism and substrate binding mode of GalB is distinct from that of the PIG-L N-deacetylases.

The putative active site of GalB encompasses several hydrophilic residues that may support substrate binding and catalysis. Three arginine residues (Arg-21, Arg-67, and Arg-216) are found within 12 Å of the bound zinc ion and are candidates for ionic interaction with the three carboxylates on CHM (Fig. 4A). Each of the residues was substituted separately with alanine, and their activities were measured with both CHM and CHPD (Table 4). Each variant had decreased specific activities relative to the wild type enzyme for both CHM and CHPD hydration. The GalB R216A variant had a 142-fold decrease in \( k_{cat} \) for CHM but a 1.2-fold increase in \( k_{cat} \) for CHPD hydration. The E48A substitution has minimal effect on the metal binding capacity of the enzyme. The E48A substitution, however, resulted in a >7000-fold decrease in the catalytic efficiency of the enzyme for CHM, largely due to a >760-fold decrease in \( k_{cat} \) (Table 4). The pH dependence of the E48A-catalyzed hydration of CHM showed no significant change on the \( pK_a \) of the enzyme-substrate complex or on the descending limb of log \( k_{cat}/K_m \) (Fig. 7). However, the substitution causes the \( pK_a \) of the ascending limb of \( k_{cat}/K_m \) to shift from 6.2 ± 0.2 observed in the wild type to 5.7 ± 0.1 in the enzyme variant (Table 5). Together, the E48A investigations suggest a minimal role for Glu-48 in metal binding but that the residue is important to the enzyme catalytic mechanism by likely supporting proton transfer.

A histidine residue, His-164, is found in the GalB active site with the Cβ being 8.6 Å from the zinc ion (Fig. 7). Substitution of His-164 with alanine resulted in no significant change in \( K_m \) for the CHM hydration under standard assay conditions but there was a 56-fold decrease in \( k_{cat} \) (Table 4). The pH dependence of the H164A variant-catalyzed hydration of CHM showed no significant change in the \( pK_a \) of the enzyme-substrate complex or the ascending limb of the \( k_{cat}/K_m \) profile (Fig. 7).
shown are: dioxygenase genes (GalA, LigAB, and LigZ) in homologous to the GalB CHM hydratase are shown in.

Comparative Analysis of GalB and LigJ CsE6—Degradation of gallic acid through the gallate pathway can be monitored by coupling the production of oxaloacetate from GalA-GalD-GalC to NADH oxidation by L-malate dehydrogenase. When GalB is omitted from the coupled reaction, CHM is not hydrated to CHA and the NADH is not oxidized (data not shown). Substitution of LigJ CsE6 for GalB in the coupled reaction leads to the oxidation of NADH in a 1:1 stoichiometric ratio to the amount of gallic acid originally present, indicating that LigJ CsE6 utilizes the same CHM isomer as GalB. When the tautomerase GalD is omitted from either the GalB- or LigJ CsE6-coupled reactions, only a very slow NADH oxidation is observed and the rate is independent of hydratase concentration, demonstrating that only the enol form (CHM) and not the keto form (OMA) of the compound is utilized as the substrate by both hydratases. The slow rate is thus attributable to the rate-limiting spontaneous tautomerization of the OMA to CHM in the absence of GalD. Polarimetric analysis of the LigJ CsE6-catalyzed reactions confirms that the enzyme has the same substrate stereo requirement as the GalB enzyme, producing and utilizing only the (−)-CHA and (−)-HMG enantiomers (Fig. 5).

Comparison of GalB and LigI Active Sites—Although they exhibit different folds, GalB and LigI share some similarities in their active site organization. In the LigI structure from R. palustris (PDB code 2GWG) the bound zinc ion is ligated in a tetrahedral fashion by residues His-6, His-8, His-178, and Glu-248 (Fig. 4C). Analogous to the GalB Glu-48 residue, the LigI metal ligating Glu-248 is found to coordinate the zinc ion on the solvent-accessible face of the metal ion that separates the metal from the remainder of the active site pocket. The GalB Glu-48 is critical to enzyme function and may be functionally convergent on Glu-248 in LigI. There are two missing regions not modeled in the final R. palustris structure, which map close to the active site. The cysteine residues of the LigI from Pseudomonas ochraceae NG11, including Cys-183 and Cys-186 from the Thr-181–Gly-191 missing region of the R. palustris enzyme, are not involved in CHM catalysis but may be involved with substrate binding (10, 46). Similar to GalB, the LigI active site pocket is lined with several hydrophilic residues, which could support substrate binding and catalysis. However, to date, no investigations of the LigI active site residues have been completed, and the role of these residues remains unknown.

Operon Organizations—A MultiGeneBlast search was utilized to identify species and gene clusters containing homologs to the gal and lig operons. Clusters for both gal and lig genes were found across proteo (α, β, and γ)- and actinobacteria (Fig. 8). In general, gene clusters containing a galB homolog were less prevalent in β-proteobacteria, whereas those with a ligJ were found throughout all bacterial clades. The galB gene was not found to replace ligJ in any of the canonical Lig pathway gene clusters. In contrast, galD homologs are frequently found in lig operons and is found in the same gene cluster as the canonical protocatechuate 4,5-cleavage pathway of Sphingo-
bium sp. SYK-6 (9, 47). The presence of a GalD homolog in the Lig pathway suggests that the product of 2-pyrene-4,6-dicarboxylate (PDC) lactonase, LigI, is OMA rather than (2Z,4E)-CHM, which is subsequently tautomerized to (2E,4E)-CHM by the GalD homolog to be used by either GalB or LigI (Fig. 1). Thus the genetic organization is consistent with our experimental data suggesting that both GalB and LigI utilize the same isomer of CHM produced by GalD. It is, however, unclear why two evolutionarily distinct hydratases are recruited to perform the same reaction in these related pathways.

Discussion

The metabolism of gallic acid by P. putida through the gallate pathway is similar to the metabolism of other aromatic compounds such as hydroxycinnamylate (through the Hpa/Hpc pathway) and catechol (through the Xyl 2,3-cleavage pathway). In all of these pathways a dienol or dienolate compound is produced (either as a substrate or transition state intermediate), which is then hydrated to an aldol product to enable C–C cleavage by a subsequent aldolase that connects the products to central cellular metabolism (48–50). GalB is structurally distinct from the previously characterized hydratases, including the CHM hydratase LigI, and is instead related to the PIG-L family of deacetylases.

In common with the PIG-L deacetylases, GalB has a single α/β domain with a Rossmann-like fold containing a His-X₆-Asp-X₄-His divalent metal binding motif found at the bottom of a solvent-accessible pocket. As with members of the PIG-L family, GalB was initially described as a Zn²⁺-dependent enzyme (9). Kinetic analyses of the PIG-L family protein MshB identified the enzyme activity as maximal when utilizing Fe²⁺ as a cofactor, and Fe²⁺ was found to be the predominant ion when the gene is expressed and the product purified under anaerobic conditions (51). When expressed and purified aerobically, however, MshB is found to switch metals, resulting in predominantly Zn²⁺ bound in the purified enzyme. Unlike previous reports on GalB, here we showed that enzymatic activity could be rescued when the apoenzyme was incubated for at least 1 h with the exogenous metal ions (9). GalB exhibits the same trend as MshB in metal ion activity, with the highest turnover catalyzed by cofactors Fe²⁺ > Co²⁺ >> Zn²⁺ (51). Only small changes in CHM specificity are seen when GalB utilizes Fe²⁺, Co²⁺, or Zn²⁺ as a cofactor, and the increase in kcat for Fe²⁺ relative to Zn²⁺-supported reactions is tempered by a simultaneous increase in Km, where Zinc is found as the predominant metal ion when GalB is overexpressed in the native strain of Pseudomonas and aerobically purified, with iron and cobalt each found in less than 1% of the total enzyme. The identity of the metal utilized by the enzyme in vivo may not be zinc, however, as the lengthy and aerobic protein isolation strategies utilized in our study may have resulted in a loss of iron, similarly seen in MshB and other iron-dependent enzymes (51). GalB is able to bind both Zn²⁺ and Co²⁺ efficiently with submicromolar Kd values. As expected, enzyme variants of the metal ligating residues, except for the Glu-48 residue, resulted in substantial increases in the proteins Km for metal ion. The Glu-48 residue is not observed to coordinate the metal ion in any of the known PIG-L deacetylases, which contain an equivalent glutamate residue, and its inclusion in GalB is likely to fulfill more of a functional role than that of a metal ligand.

There is some uncertainty as to the isomeric form of CHM utilized by LigJ and GalB (8, 9). The initial characterization of the gal operon identified (2E,4E)-CHM as the product of the GalD-catalyzed reaction, describing the compound as the 2E isomer on the basis of the chemical shift of the C3 olefinic proton (δ 6.73 ppm) being lower than expected if it was the 2Z isomer (δ 7.38 ppm) (9). However, a recent crystal structure of the PDC lactonase LigI, which catalyzes the production of an unknown tautomer of OMA or CHM in the Lig pathway, was solved and found to contain (2Z,4E)-CHM bound to the enzyme. The potential difference in the isomeric forms of CHM produced through the two different pathways was initially hypothesized to be the result of differences in the substrate stereospecificity of the two hydratases, GalB and LigJ, from the distinct pathways. However, here we have shown that both GalB and LigJCatE6 can indeed utilize the same CHM isomer produced from GalD, and this utilization leads to the same (−)-enantiomer of CHA. Similarly, when reactions with either GalB or LigJCatE6 were incubated with racemic CHA, only the (−)-enantiomer was consumed by both of the hydratase enzymes. A galD homolog is commonly found in the protocatechuate 4,5-cleavage pathways, and previous studies have shown that GalB can rescue the growth of a Sphingomonas sp. SYK-6 derivative species (Sphingomonas sp. DL1), which lacks ligJ when grown on vanillate or syringate, further indicating that the GalB and LigI enzymes act on the same (2E,4E)-CHM isomer produced physiologically by GalD (9, 10). The assumed CHM product observed in the LigI structure may therefore be an artifact of crystallization arising from utilization of CHM/OMA prepared at pH 10, but when crystallized at pH 6.5, OMA underwent nonenzymatic tautomerization to (2Z,4E)-CHM, which may not be the physiological substrate for the enzyme (8).

Previous characterizations of the Galc/LigK gene product have identified both CHA and HMG as substrates for the aldolase enzyme, with the enzyme having stereo preference for the (R)-HMG and (−)-CHA enantiomers (12, 52–54). Physiological routes leading to CHA production, through both the gallate and PCA 4,5-cleavage pathways, have been identified for some time. However, there has yet to be a definition of a physiological route leading to HMG production. Characterizations of DDVA metabolism in Sphingomonas sp. SYK-6 have identified the genes LigXa and LigZ, which transforms DDVA into a meta-cleavage product (4-[2-(5-carboxy-2-hydroxy-3-methoxyphenyl)-2-oxoethyldiene]-2-hydroxypent-2-enoic acid) that is hydrolyzed by LigY into 5-carboxyvanillate and CHPD (5, 55, 56) (Fig. 1). The 5-carboxyvanillate is further metabolized to vanillate and then degraded through the protocatechuate 4,5-cleavage pathway. However, the subsequent metabolism of the CHPD has yet to be defined. Here we have shown that both GalB and LigJCatE6 utilize the (R)-HMG enantiomer from a racemic HMG mixture and that the dehydration reaction is reversible. GalB had a 10-fold increase and a 23-fold decrease in Km and kcat, respectively, for CHPD hydration relative to the CHM substrate. Although the kinetic parameters for CHPD are less ideal relative to the CHM substrate, it is reasonable to assign CHPD hydratase roles to both GalB and LigI homologs in the
LigXa/LigZ/LigY pathway, which gives a route to HMG production for the HMG/CHA aldolase that has yet to be identified.

Attempts to cocrystallize GalB or the E48A variant with either CHM or CHA did not yield new crystallization conditions from the crystallization screens used and yielded only poorly diffracting crystals in the holoenzyme condition. Soaking the crystals with their substrates or products resulted in cracking of the crystals and was not successful. Also, attempts to model the substrates and/or products of GalB into the active site by computational docking methods did not yield productive models that would support catalysis. The kinetic investigations carried out here have led to a proposal for key residues in the GalB active site and have facilitated the generation of a proposed model for substrate binding and catalysis (Fig. 4B).

Binding of the substrate is facilitated by the C1 carboxyl positioned by interaction with Ne of His-198, the C2 enolate positioned by interaction with Nε of Arg-67 and Nε of Gln-196, the C4 carboxylate positioned by interaction with Nη2 of Arg-21 and the hydroxyl of Tyr-203, and the C5 carboxylate positioned by interaction with Nη1 of Arg-216. The R216A variant had a ~50,000-fold decrease in the specificity constant for CHM but only a 3-fold decrease in the specificity constant for CHPD, which is consistent with the residue facilitating (2E,4E)-CHM binding through the C5 carboxylate. In the proposed mechanism the (2E,4E)-CHM enolate serves as the substrate for the enzyme and Glu-48 activates the metal-ligated water for the addition at the C4 with protonation by His-164 at the C5 (Fig. 9). Tautomerization of the enolate leads to abstraction of the proton from Glu-48 to the C3 position, thus completing the reaction cycle.

The enol form (CHM), and not the keto form (OMA), acts as the substrate for GalB, indicating that either the enol proton contributes to the catalytic mechanism or the enolate serves as the substrate. The CHM has a pKα of 7.3, which leads to an increase in the extinction coefficient for the substrate with increasing pH. This is consistent with the enol deprotonation leading to an increase in the chromogenicity of the compound, which would not arise from deprotonation of the carboxylate moieties of CHM. The enol pKα is observed in the enzyme pH dependence, with the deprotonation leading to increased GalB hydration activity, indicating that the substrate for the enzyme is in fact the enolate and that the enol proton is not likely to be involved in the catalytic mechanism.

The single ionization in the enzyme-substrate complex (pKα 6.1 ± 0.1) of GalB is analogous to the pKα observed in carbonic anhydrases, where the zinc metal cofactor in the carbonic anhydrides facilitates the decrease in the pKα of water, leading to a hydroxide primed for nucleophilic attack at physiological pH values (reviewed in Ref. 57). In the PIG-L N-deacetylases and some members of the metalloprotease family (such as carboxypeptidase A and thermolysin, which are structurally distinct from the PIG-L N-deacetylases but contain a similar metal ligand and act on similar chemical linkages), a similar Lewis acid mechanism is also proposed (38, 58, 59). In these enzymes an acidic residue (an aspartate in PIG-L and glutamate in the metalloproteases) in close proximity to the metal ligand is proposed to deprotonate the metal-ligated water to act as a nucleophile (45, 60–62). In both cases, after bond scission the protonated acidic residues relinquish the proton either to solvent or to the newly formed terminal atom. Similar to what is observed with the PIG-L N-deacetylases and metalloproteases, the GalB Glu-48 carboxylate ligates the metal ion and is in close proximity to the metal-ligated water. The GalB E48A variant has a 760-fold decrease in kcat for the CHM hydration reaction, and the pKα of the free enzyme in the variant-catalyzed reaction shifts down 0.6 unit (from ~6.3 in the wild type to ~5.7 in the variant) supporting the role of Glu-48 in proton transfer and acting as a general base. The abstracted proton from the metal-ligated water could be utilized to protonate the C3 during tautomerization or could be lost to solvent, with the solvent donating the proton to complete the reaction cycle.

His-164 is located in the active site with the imidazole ring facing the pocket and its Nδ making a hydrogen bond to the phenolic hydroxyl of Tyr-203. Rotation of the imidazole group of His-164 such that Nδ projects toward the active site results in
Nδ being 6.6 Å from the metal-ligated water (Fig. 4B). The H164A substitution results in a 56-fold decrease in $k_{cat}$ with a relatively negligible decrease on the $K_m$ of the CHM hydration reaction. The $pK_a$ of the free enzyme in the H164A variant increases ~0.6 unit, from 8.6 in the wild type to 9.2 in the variant, consistent with the proposal that the residue functions as a general acid in the reaction mechanism.

The effect of either Zn$^{2+}$ or Co$^{2+}$ as a cofactor on the pH dependence of the enzyme is minimal. The GaLB $pK_a$ values for both the enzyme-substrate complex and the free enzyme/substrate are unaffected by the metal cofactor utilized. Bell-shaped profiles for pH dependence on $k_{cat}/K_m$ for Zn$^{2+}$ cofactor reactions are observed in the PIG-L deacetylases MshB (p$K_a$ values 7.3 and 10.5) and BshB (p$K_a$ values 6.5 and 8.5), which are similar to GaLB (45, 63). In both MshB and BshB, a minimal or no effect is observed on either p$K_a$ when utilizing Co$^{2+}$ as the cofactor, and in MshB neither Fe$^{2+}$, Ni$^{2+}$, nor Mn$^{2+}$ affected the p$K_a$ values (45, 63). In the proposed GaLB and PIG-L deacetylase reactions, the metal contributes to the mechanism by positioning and likely lowering the p$K_a$ of the catalytic water. However, it appears that in both GaLB and the PIG-L deacetylases that the metal does not significantly affect the p$K_a$ of the catalyzed reaction, possibly the result of key residues in the enzymes having a more significant effect on the p$K_a$.

GaLB is a hydratase from the gallic acid utilization pathway that is structurally distinct from previously characterized hydratase enzymes from other aromatic catabolic gene clusters, including the CHM hydratase LigI. A comparison of the GaLB and LigI active site organizations indicates similar residues that may facilitate a common mechanism between the convergently evolved enzymes. The structure and kinetic characterization of GaLB presented here will help guide future investigations into both the LigI and GaLB CHM hydratases.

Author Contributions—S. M. did most of the experiments, crystallized and solved the GaLB structure, and wrote most of the paper. A. S. B. optimized the protocols to synthesize and purify CHM in addition to creating some of the GaLB variants. M. S. K. helped with the crystallography and critically appraised the manuscript. S. Y. K. S. coordinated the project and assisted S. M. with writing the manuscript. All authors reviewed the results and approved the final version of the manuscript.

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