Evolution of Enzymatic Activity in the Enolase Superfamily: Structural and Mutagenic Studies of the Mechanism of the Reaction Catalyzed by o-Succinylbenzoate Synthase from Escherichia coli

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ABSTRACT: o-Succinylbenzoate synthase (OSBS) from Escherichia coli, a member of the enolase superfamily, catalyzes an exergonic dehydration reaction in the menaquinone biosynthetic pathway in which 2-succinyl-6-hydroxy-2,4-cyclohexadiene-1-carboxylate (SHCHC) is converted to 4-(2′-carboxyphenyl)4-oxobutyrate (o-succinylbenzoate or OSB). Our previous structural studies of the Mg$^{2+}$-OSB complex established that OSBS is a member of the muconate lactonizing enzyme subgroup of the superfamily: the essential Mg$^{2+}$ is coordinated to carboxylate ligands at the ends of the third, fourth, and fifth $\beta$-strands of the $(\beta/\alpha)\beta$-barrel catalytic domain, and the OSB product is located between the Lys 133 at the end of the second $\beta$-strand and the Lys 235 at the end of the sixth $\beta$-strand [Thompson, T. B., Garrett, J. B., Taylor, E. A., Meganathan, R., Gerlt, J. A., and Rayment, I. (2000) Biochemistry 39, 10662–76]. Both Lys 133 and Lys 235 were separately replaced with Ala, Ser, and Arg residues; all six mutants displayed no detectable catalytic activity. The structure of the Mg$^{2+}$-SHCHC complex of the K133R mutant has been solved at 1.62 Å resolution by molecular replacement starting from the structure of the Mg$^{2+}$-OSB complex. This establishes the absolute configuration of SHCHC: the C1−carboxylate and the C6−OH leaving group are in a trans orientation, requiring that the dehydration proceed via a syn stereochemical course. The side chain of Arg 133 is pointed out of the active site so that it cannot function as a general base, whereas in the wild-type enzyme complexed with Mg$^{2+}$-OSB, the side chain of Lys 133 is appropriately positioned to function as the only acid/base catalyst in the syn dehydration. The $\epsilon$-ammonium group of Lys 235 forms a cation−$\pi$ interaction with the cyclohexadienyl moiety of SHCHC, suggesting that Lys 235 also stabilizes the enediolate anion intermediate in the syn dehydration via a similar interaction.

The members of the enolase superfamily catalyze mechanistically diverse reactions initiated by the abstraction of a proton from a carbon adjacent to a carboxylate group ($\alpha$-proton) by an active site base to generate an enediolate anion intermediate. This intermediate, which is stabilized by coordination to an essential Mg$^{2+}$ ion, is then processed to the product by an active site acid (1). The members of the superfamily share a two-domain structure, with the active sites located at their interface. The larger central domain consists of a $(\beta/\alpha)\beta$-barrel domain and encompasses the three ligands for the Mg$^{2+}$ at the ends of the third, fourth, and fifth $\beta$-strands as well as one or more acid/base catalysts at the C-terminal ends of the second, third, fifth, sixth, or seventh $\beta$-strands. The smaller capping domain, which exhibits a mixed $\alpha/\beta$ motif, is more variable and is formed from both the N- and the C-termini of the polypeptide. This domain typically contains residues involved in substrate binding.

Excluding the ubiquitous enolases, over 500 nonredundant members of the superfamily now can be identified in the sequence databases (J. A. Gerlt, unpublished observations). These can be partitioned into two groups based on the identities of the acid/base catalysts. The mandelate racemase (MR)/ subgroup is defined by a conserved His−Asp dyad, with the His and Asp residues located at the C-terminal ends

1 This research was supported by Grants GM-52594 (J. A. G. and I. R.) and AR-35186 (I. R.) from the National Institutes of Health. Use of the Argonne National Laboratory Structural Biology Center beam line at the Advanced Photon Source was supported by the U.S. Department of Energy, Office of Energy Research, under Contract W-31-109-ENG-38.
2 The X-ray coordinates of the OSBS K133R-2-succinyl-6-hydroxy-2,4-cyclohexadiene-1-carboxylate complex have been deposited in the Protein Data Bank with accession number 1ROW.
3 To whom correspondence should be addressed. (I. R.) Phone: (608) 262-0437. Fax: (608) 262-1319. E-mail: Ivan_Rayment@biochem.wisc.edu. (J. A. G.) Phone: (217) 244-7414. Fax: (217) 244-6538. E-mail: j-gerlt@uiuc.edu.
4 University of Wisconsin.
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$^*$ This abbreviation is defined in the sequence databases (J. A. Gerlt, unpublished observations).

$^1$ Abbreviations: APS, Advanced Photon Source at Argonne National Laboratory; MePEG, methyl ether poly(ethylene glycol); MLE, muconate lactonizing enzyme; MR, mandelate racemase; OSBS, o-succinylbenzoate synthase; SBC, The Structural Biology Center at the Advanced Photon Source, Argonne, IL; SHCHC, 2-succinyl-6-hydroxy-2,4-cyclohexadiene-1-carboxylate; OSB, 4-(2′-carboxyphenyl)-4-oxobutyrate or o-succinylbenzoate; rms, root-mean-square; rmsd, root-mean-square deviation.
of the seventh and sixth β-strands, respectively. In addition, many, but not all members of this subgroup contain a Lys-X-Lys motif at the end of the second β-strand. The muconate lactonizing enzyme subgroup is defined by both a conserved Lys-X-Lys motif at the end of the second β-strand as well as a Lys at the end of the sixth β-strand. The C-terminal ends of the second and sixth/seventh β-strands are located on opposite faces of the active site, so these can catalyze α-proton abstraction and delivery in reactions proceeding by either syn or anti stereocchemical courses.

α-Succinylbenzoate synthases (OSBSs) are members of the MLE subgroup and catalyze an exergonic dehydration reaction in which 2-succinyl-6-hydroxy-2,4-cyclohexadien-1-carboxylate (SHCHC) is converted to 4-(2′-carboxyphenyl)-4-oxobutyrate (α-succinylbenzoate or OSB) (Scheme 1).

An abundant number of sequences for OSBSs is available (>70) since this enzyme is required for the biosynthesis of menaquinone (2), an essential cofactor for anaerobic growth in eubacteria and some archaea. Although these are highly diverged, with pairwise sequence identities often <15%, virtually all orthologs are predicted to contain a Lys-X-Lys motif at the end of the second β-strand and a Lys at the end of the sixth β-strand. In the case of the OSB from *Escherichia coli*, this active site architecture has been confirmed by our structural studies of a complex with Mg2+. OSB that show that the carboxylate group of OSB is coordinated to the Mg2+ and the α-carbon is sandwiched between the ε-ammonium groups of Lys 133 at the end of the second β-strand and Lys 235 at the end of the sixth β-strand (3).

Despite the availability of a high-resolution structure for the Mg2+-OSB complex, the mechanism of the OSBS-catalyzed reaction is uncertain. In particular, the configuration of the α-carbon of the SHCHC substrate is unknown, thereby confounding a detailed understanding of the roles of Lys 133 and Lys 235 in catalysis. Without this knowledge, the stereocchemical course, syn or anti, of the dehydration reaction likewise cannot be defined (Scheme 1). SHCHC is prepared enzymatically from chorismate by the coupled actions of isochorismate synthase and SHCHC synthase (4), the latter a thiamin diphosphate (TDP)-dependent enzyme that uses isochorismate and α-ketoglutarate (α-KG) as substrates (Scheme 2).

The configuration of C6 that bears the OH-leaving group is unchanged from that in isochorismate (2S in isochorismate, 6R in SHCHC); however, the absolute configuration of C1 bearing the α-proton is established in the reaction catalyzed by SHCHC synthase and has not been experimentally determined.

In this paper, we establish that OSBS from *E. coli* catalyzes a syn dehydration reaction. Replacements of either Lys 133 or Lys 235 with Ala, Ser, or Arg residues yield proteins with no detectable catalytic activity. We solved the structure of the Mg2+-SHCHC complex of the K133R mutant, thereby allowing the configuration of the substrate to be established. The 6-OH leaving group and the 1-carboxylate group are trans substituents of the C1–C6 bond, requiring that the dehydration reaction proceed via a syn stereocchemical course. The inactivity of the K133R mutant is attributed to the inability of the Arg 133 side chain to fit in the active site, so it extends away from the bound SHCHC and cannot abstract the α-proton. The ε-ammonium group of Lys 235 forms a cation–π interaction with the cyclohexadienyl moiety of SHCHC, thereby providing the opportunity for enhanced stabilization of an enediolate anion intermediate. Such assistance in catalysis is a novel catalytic role for an active site functional group in the enolase superfamily.

**MATERIALS AND METHODS**

*E. coli* strains XL1-Blue and BL-21 (DE3) were obtained from Stratagene. Taq DNA polymerase and restriction enzymes were purchased from Invitrogen. Oligonucleotide primers were synthesized by Operon (Alameda, CA) or by Biosynthesis (Lewisville, TX). Deuterated reagents were obtained from Cambridge Isotopes Ltd. All other reagents were of the highest quality grade commercially available.

NMR spectra were recorded using Varian Unity Inova 500 and 600 MHz NMR spectrometers. Spectrophotometric assays were performed with a Perkin-Elmer Lambda-14 spectrophotometer. PCR reactions were performed using a MJ Research PTC-200 Peltier Thermal Cycler.

**Biosynthesis of 2-Succinyl-6-hydroxy-2,4-cyclohexadien-1-carboxylate (SHCHC).** SHCHC was synthesized from barium chorismate, α-ketoglutarate (α-KG), thiamin diphosphate (TDP), and the coupled actions of isochorismate mutase (encoded by the *entC* gene from *E. coli*) and SHCHC synthase (encoded by the *menD* gene from *E. coli*) and purified as described previously (4). ESI-MS 247.1 (corresponds to monolithiated, monoprotonated species); *λ*max = 298 nm, ε = 4000 M–1; HPLC elution time from a C8 reverse phase column with isocratic elution with 95% of an aqueous solution of 0.1% TFA, 5% acetonitrile. ~7 min; 1H NMR (500 MHz, D2O) δ = 2.29 (m, 2H), δ = 2.91 (m, 2H), δ = 3.58 (d, 1.6 Hz, 1H), δ = 4.42 (dd, 1.7, 5.4 Hz, 1H),  δ = 6.18 (d, 5.4, 9.3 Hz, 1H), δ = 6.25 (dd, 5.7, 9.5 Hz, 1H), δ = 7.16 (d, 5.7 Hz, 1H).
Site-Directed Mutagenesis. The template for mutagenesis was the 0.96 kb menC gene from E. coli MG1655 subcloned using Ndel and XhoI cloning sites into a modified, 10 histidine-tagged pET15-b vector. The QuickChange site-directed mutagenesis method was utilized to introduce the desired mutations. The mutations were confirmed by sequencing performed at the W. M. Keck Biotechnology Center (Urbana, IL).

Protein Expression and Purification. Mutant proteins of OSBS were expressed in E. coli strain BL-21 (DE3) and purified as described previously for the wild-type enzyme (3). Fractions containing protein were identified by OD280 and SDS–PAGE. Electrospray ionization mass spectrometry (ESI-MS) was used to confirm the expected masses: predicted, 37,736; observed, 37,718 (within the expected 0.1% error).

Circular Dichroism. Wild-type and mutant versions of OSBS were subjected to circular dichroism (CD) studies at the Laboratory of Fluorescence Dynamics at the University of Illinois. The various proteins were dialyzed against 50 mM Tris-HCl, pH 8.0, containing 0.1 mM MnCl2; the samples were adjusted to protein concentrations of ~0.3 mg/mL. The CD spectra were obtained with a Jasco 720 spectropolarimeter using 0.1 cm quartz (QS) cuvettes. The samples were scanned five times at room temperature over the 190–350 nm wavelength range (0.5 nm increments), and the scans were averaged. The CD spectra were recorded in the absence of any ligands.

Spectrophotometric Assay. OSBS activity was measured at 25 °C using varying concentrations of 2-hydroxyl-6-succinyl-2,4-cyclohexadiene (SHCHC), 50 mM Tris-HCl, pH 8.0, and 0.1 mM MnCl2 by quantitating the decrease in absorbance at 310 nm (Δε = −2400 M−1 cm−1).

Kinetic Studies using 1H NMR Spectroscopy. Wild-type and mutant proteins were exchanged into D2O using an Amicon stirred cell. A 10 mL aliquot of protein was concentrated to 2 mL; 8 mL of D2O was added, and the protein solution was again concentrated to 2 mL. This process was repeated three times. Samples for 1H NMR analysis (800 μL) contained 1 mM SHCHC, 0.5 mg of protein, 50 mM [d1]Tris-DCl, pH 8.0, and 0.1 mM MnCl2 in D2O. Samples were monitored for formation of the OSB product as well as predicted, 37,736; observed, 37,718 (within the expected 0.1% error).

Table 1: Data Collection and Refinement Statistics

<table>
<thead>
<tr>
<th>Space Group</th>
<th>P21/P21/P21</th>
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<tbody>
<tr>
<td>Unit Cell, Å</td>
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<tr>
<td>Wavelength, Å</td>
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<td>Resolution, Å</td>
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<td>Total Reflections</td>
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<td>Unique Reflections</td>
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<td>Redundancy</td>
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</tr>
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<td>Average I/σ (raw/scaled)</td>
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<td>Completeness, %</td>
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</tr>
<tr>
<td>Rmerge, %</td>
<td>4.6 (29.3)</td>
</tr>
</tbody>
</table>

Refinement and Model Statistics

| NO. of Protein Atoms | 2484 |
| NO. of Heteroatoms | 361 |
| R Factor | 16.7 (19.5) |
| Rmerge Factor | 20.0 (24.6) |
| Wilson B Value, Å2 | 25.9 |
| Average B Factor, Å2 | 17.9 |
| Ramachandran, % | most favored 92.5 |
| additionally allowed 7.5 |
| disallowed 0.0 |
| rmsd in bond distance, Å | 0.016 |
| rmsd in bond angle, ° | 1.62 |
| disordered side chains | Arg20, Arg22, Ala38, Pro125, Gly126, Glu127 |

*The value in parentheses give the statistics for the highest resolution shell that extends from 1.68 to 1.62 Å. Rmerge = (ΣI–B[I])/ΣI, where the average intensity I is taken over all symmetry equivalent measurements, and B[I] is the measured intensity for a given reflection.

C circular dichroism, D2O, and SDS–PAGE. Electrospray ionization mass spectrometry (ESI-MS) was used to confirm the expected masses: predicted, 37,736; observed, 37,718 (within the expected 0.1% error).

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Crystals and X-ray Data Collection. The K133R mutant of OSBS was concentrated to 15 mg/mL, dialyzed against 5 mM Tris-HCl, pH 8.3 containing 2 mM MgCl2, drop frozen as small pellets in liquid nitrogen, and stored at −80 °C. Crystals were grown at 20 °C by small-scale batch experiments by combining 15 μL of protein solution and 15 μL of a solution containing 12–13% MePEG 5000, 100 mM sodium acetate, 60 mM MgCl2, at pH 5.5 (5). SHCHC was included in the crystallization at a final concentration of approximately 2.5 mM. These conditions were similar to those used to obtain the structure of the product complex (3). Crystals grew spontaneously on occasions but were generally grown by streak seeding 24 h after mixing. Typically, the crystals attained a maximum size of ~0.5 × 0.5 × 0.05 mm after two weeks. They belong to space group P21212 with cell dimensions of a = 72.1, b = 82.9, and c = 56.2 Å with a monomer in the asymmetric unit and diffracted to at least 1.6 Å resolution.

Crystals grown in the presence of SHCHC were transferred to a cryo-protectant solution containing 20% MePEG 5000, 15% ethylene glycol, 100 mM MgCl2, 5 mM SHCHC and then were flash-frozen in liquid nitrogen. Initially, the crystals were equilibrated in 20% MePEG 5000, 100 mM MgCl2, 5 mM SHCHC and then transferred to the final solution in four equal steps of increasing ethylene glycol concentration.

A native data set was recorded at BioCARS beam line 14BM-D at the Advanced Photon Source, Argonne National Laboratory with a single scan composed of 270 frames each of width 0.7° with an exposure of 20 s/frame at a wavelength of 0.979 Å. The data were integrated and scaled with HKL2000 and Scalepack (6) (Table 1).

Structure Determination and Refinement. The structure was determined by molecular replacement with the program MOLREP starting from the coordinates for the product complex (PDB accession code 1FH, ref 3). Thereafter, the structure was refined with the program Refmac (7, 8). Water molecules were added to the coordinate set with ARP/WARP and subsequent manual verification (7, 9). Iterative cycles of maximum likelihood refinement and manual model building reduced the R factor to 16.7% for all measured X-ray data from 30.0 to 1.62 Å resolution. The R free was 20.0% for 5% of the data that was excluded from the refinement. Refinement statistics are presented in Table 1. Analysis of the coordinates with the program PROCHECK (10) revealed that 92.5% of the residues lie in the most favored regions of the Ramachandran plot, whereas the remaining 7.5% of the residues lie in additionally allowed areas. No residues are located in the disallowed regions.

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RESULTS AND DISCUSSION

The experiments described in this manuscript were designed to determine the functions of Lys 133 and Lys 235 in the OSBS-catalyzed reaction. Our hypothesis, based on previous structural studies of the Mg\(^{2+}\)-OSB complex and the location of C6 of the OSB product in the active site, was that Lys 235 would function as the catalytic base, and Lys 133 would function as the catalytic acid in an anti dehydratation; it was hypothesized that mutagenesis of one or both of these residues would result in a catalytically impaired mutant that might be used for the determination of the structure of the Mg\(^{2+}\)-SHC complex.

Biosynthesis of SHCHC. Although this laboratory has described the enzymatic synthesis of SHCHC (Scheme 2), the absolute configuration of the carbon from which the proton is abstracted had not been determined (4). The synthesis of SHCHC is the first committed step in the menaquinone biosynthetic pathway. In vivo, isochorismate, the substrate for SHCHC synthase, is generated from chorismate by MenF, one of two homologous isochorismate synthases encoded by the \(E. \text{coli}\) genome. In our enzymatic synthesis, we utilized EntC, the second isochorismate synthase. The absolute configuration of isochorismate had been previously characterized, with no change in the absolute configuration of C2 (C6 in SHCHC), the OH-bearing carbon, occurring in the isochorismate mutase reaction. Conversion of isochorismate to SHCHC is an uncharacterized two-step process: the addition of the succinyl side chain derived from isochorismate by MenF, one of two homologous isochorismate synthases encoded by the \(E. \text{coli}\) genome. In our enzymatic synthesis, we utilized EntC, the second isochorismate synthase. The absolute configuration of isochorismate had been previously characterized, with no change in the absolute configuration of C2 (C6 in SHCHC), the OH-bearing carbon, occurring in the isochorismate mutase reaction. Conversion of isochorismate to SHCHC is an uncharacterized two-step process: the addition of the succinyl side chain derived from \(\alpha\)-KG in a process using TDP followed by the elimination of pyruvate. In this reaction, the absolute configuration of C6 also is unchanged, but the absolute configuration of C1, the carbon from which the proton is abstracted in the OSBS-catalyzed reaction, has not been determined. In an attempt to resolve this uncertainty, we analyzed the \(^1\text{H NMR}\) spectrum of SHCHC (data not shown) and realized that the coupling constants involving the protons of C1 and C6 (\(J_{5,6} = 5.5\ \text{Hz}, J_{1,6} = 1.8\ \text{Hz}\)) are consistent with either a cis or a trans relationship of the C1-carboxyl group and the C6-OH group. As a result, we have been unable to specify the stereochemical course of the OSBS-catalyzed reaction. However, because enzyme-catalyzed \(\beta\)-elimination reactions involving carboxylate anion substrates usually, but not always, proceed via an anti stereochemical course (11), we had assumed this stereochemical course and a cis relationship of the C1-carboxyl group and the C6-OH group. The structural studies reported in this paper resolve this ambiguity and demonstrate that this assumption is incorrect.

Kinetic Characterization of Mutant Enzymes. Six mutant enzymes were constructed and purified to homogeneity: K133A, K133S, K133R, K235A, K235S, and K235R. Using a spectrophotometric assay, no detectable activity was observed for any of the mutants. The value of \(k_{\text{cat}}\) for each of the mutants was estimated as \(<3 \times 10^{-5}\ \text{s}^{-1}\) (the dehydration rate in the absence of enzyme), a 630 000-fold decrease from the value of 19 s\(^{-1}\) measured for wild-type OSBS.

Because the reactions catalyzed by members of the enolase superfamily are initiated by the abstraction of an \(\alpha\)-proton to generate an enediol anion intermediate, it was possible that the mutants might catalyze the exchange of the \(\alpha\)-proton despite the lack of detectable dehydration activity (e.g., in a mechanism involving both a general base and a general acid, substitution of the general acid that facilitates the departure of the C6-OH group might allow the exchange of the \(\alpha\)-proton without dehydration). Each mutant was separately incubated with SHCHC in D\(_2\)O, and both formation of the OSB product and exchange of the \(\alpha\)-proton were monitored by \(^1\text{H NMR}\) spectroscopy. Neither dehydration nor exchange of the \(\alpha\)-proton was observed.

Effects of Mutations on the Circular Dichroism. CD spectra were obtained for wild-type OSBS and the six mutants to determine whether the mutations altered the secondary and tertiary structures. The spectra of the mutants were identical to those obtained for the wild-type enzyme (data not shown).

Active Site Structure. Crystals were obtained for the catalytically inactive K133R mutant protein in the presence of SHCHC. The electron density for the Mg\(^{2+}\)-SHCHC complex is essentially continuous for the entire length of the polypeptide chain with the exception of Arg 20, Ala 38, Pro 25, and Gly 126. The electron density starts at Met 1 and terminates at Leu 320. Likewise, almost all of the side chains are well-ordered (a list of disordered residues is given in Table 1).

The tertiary structure of the Mg\(^{2+}\)-SHCHC complex of the K133R mutant protein is essentially identical to that of the Mg\(^{2+}\)-OSB product complex of the wild-type enzyme described earlier. The rms difference between the positions of the \(\alpha\)-carbon atoms is 0.30 Å for 306 residues. In contrast, the rms difference between the tertiary structure of the Mg\(^{2+}\)-SHCHC complex of the wild-type enzyme and the unliganded protein is 0.90 Å, where the larger difference is due to the rotation of the small domain relative to the \((\alpha/\beta)\)8 barrel when a ligand is present in the active site. This indicates that the domain movements associated with the active site closure are similar regardless of whether the substrate or product are bound.

The electron density for the K133R mutation shows that, in contrast to the earlier structures where the Lys 133 extended toward the putative location of C1 in the substrate, the side chain for arginine extends to the solvent, where the electron density suggests that the side chain is quite mobile (Figure 1). The displacement of the arginine side chain from the active site is consistent with the increased size and polarity of the guanidinium moiety as compared to the amino group of lysine.

Examination of the active site reveals the electron density for a well-ordered molecule of SHCHC (Figure 2). This unequivocally defines the stereochemistry of the substrate and shows that the C1-carboxylate and the C6-OH leaving group have a trans orientation, and the absolute configurations of both carbons are \(R\). The absolute configuration of C1 differs from that assumed in the earlier structural studies of the wild-type enzyme, which leads directly to a different assignment for the catalytic acid and base as discussed below.

The C1 carboxylate of SHCHC forms a bidentate interaction with the magnesium ion. The remainder of the coordination sphere of the magnesium ion is completed by monodentate interactions with the carboxylates of Asp 161, Glu 190, and Asp 213 and a water molecule. All of the bond distances to the magnesium ion lie in the range of 1.9–2.1 Å, which is consistent with normal geometry for this ion.
The overall coordination of the magnesium ion can be described as distorted octahedral. The arrangement of ligands around the magnesium ion in the complex with SHCHC is different from that observed in the complex with OSB. In the latter case, OSB only utilized a single interaction between a carboxylate oxygen and the magnesium ion. This results in a different position for SHCHC and OSB in the active site (Figure 4) and is expected based on the change in hybridization at C1 and C6 from sp$^3$ to sp$^2$. Because of the movement of the carboxylate on C1 into the plane of the aromatic ring, the metal coordination of the carboxylate changes from bidentate in SHCHC to monodentate in OSB. This results in a translation of the remainder of the molecule by approximately 0.8 Å. This change allows the hydrophobic component of the substrate and product to remain in approximately the same plane. Apart from the change in the configuration of C1, the overall arrangement of atoms in both the substrate and the product are very similar. The movement of the product relative to the substrate requires only a minor rearrangement of the side chains in contact with the ligands (data not shown) so that ligand interaction diagram is essentially identical to that described earlier. Interestingly, the movement of the succinyl carboxylate group of SHCHC relative to that in OSB is accompanied by a rotation of the e-amino group of Lys 131, which serves to maintain an ionic interaction that stabilizes both substrate and product complex. A shift of ~0.8 Å between substrate and product is readily accommodated by nonpolar interactions, whereas such a movement would be highly unfavorable for an ionic interaction without a compensatory change.

**Structural Mechanism.** The kinetic effects of mutations at Lys 133 and Lys 235 in combination with the structure of the K133R mutant in the presence of SHC support a mechanism in which Lys 133 serves as the catalytic base for the initial proton abstraction event as depicted in Figure 5. The orientation of the C6 hydroxyl suggests that Lys 133 also serves as the catalytic acid responsible for protonation of the departing water molecule. Lys 235, although essential for enzyme activity, is on the opposite side of substrate and...
FIGURE 3: Stereoviews of the active site for and its coordination of SHCHC. Panel A shows the geometry of the metal coordination site, which is approximately octahedral with its associated bond distances. Panel B shows the ligand interactions of SHCHC with o-succinylbenzoate synthase. This illustrates that Lys 235 is on the opposite side to the proton that is abstracted by Lys 133 and the C6—OH leaving group and thus cannot function as either the catalytic base or the acid.

FIGURE 4: Stereo overlap of SHC in the K133R mutant protein and OSB in the wild-type structure (3). The coordinates for the product complex were taken from the RSCB with accession number 1FHV and superimposed with the program Align (14).
is unable to abstract the proton as indicated by the stereochemistry of C1. Given its position, Lys 235 may stabilize the negative charge on the enediolate intermediate through a cation—π interaction (Figure 3B). The inability of substitutions for Lys 235 to catalyze either dehydration or exchange of the α-proton can be explained by the loss of this stabilizing interaction.

This mechanism is different from that proposed earlier on the basis of the structure of the OSB product complex (3). In that work, Lys 235 was proposed to be the catalytic base, whereas Lys 133 was projected to function as the catalytic acid. This proposal was derived based on the position of OSB, at a time when the absolute configuration of the substrate was unknown, so that the only constraint was the position of C1 and C6. The suggestion that Lys 133 would serve as the required base and Lys 235 would function as the acid seemed entirely acceptable given the conservation and function of these residues in the MLE subgroup of the enolase superfamily. However, it is now realized that, like the syn dehydration catalyzed by (D)-glucarate dehydratase (GlucD), a single residue can catalyze both proton abstraction and departure of the leaving group. In GlucD, a member of the MR subgroup, His 339 at the end of the seventh β-strand is the dual function catalyst.

CONCLUSIONS

This study provides a structural mechanism for the reaction catalyzed by the OSBS from E. coli, which is expected to apply to all orthologs. Importantly, Lys 133 serves both as the catalytic base and as the acid, whereas the conserved Lys 235 that lies on the opposite side of the substrate appears to provide a positive charge that stabilizes the enediolate intermediate. Such assistance in catalysis is a new function for an active site functional group in the enolase superfamily, thereby expanding the potential catalytic functions of conserved active site residues in the enolase superfamily. The present studies highlight the importance of high-resolution structural studies of enzyme–substrate complexes, to both define the geometry of the bound substrate and to identify the roles of individual residues. Finally, this study illustrates the difficulty of assigning functions to individual residues based on the structure of a complex with a product that differs significantly in both structure and energy from the transition state.

REFERENCES