A Conserved Lysine in β-Lactam Synthetase Assists Ring Cyclization: Implications for Clavam and Carbapenem Biosynthesis

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β-Lactam synthetase (β-LS) is the paradigm of a growing class of enzymes that form the critical β-lactam ring in the clavam and carbapenem antibiotics. β-LS catalyzes a two-stage reaction in which N⁵-(2-carboxyethyl)-L-arginine is first adenylated, and then undergoes intramolecular ring closure. It was previously shown that the forward kinetic commitment to β-lactam formation is high, and that the overall rate of reaction is partially limited to a protein conformational change rather than to the chemical step alone of closing the strained ring. β-Lactam formation was evaluated on the basis of X-ray crystal structures, site-specific mutation, and kinetic and computational studies. The combined evidence clearly points to a reaction coordinate involving the formation of a tetrahedral transition state/intermediate stabilized by a conserved Lys. The combination of substrate preorganization, a well-stabilized transition state and an excellent leaving group facilitates this acyl substitution to account for the strong forward commitment to catalysis and to lower the barrier of four-membered ring formation to the magnitude of a protein conformational change.

Introduction

How the strained four-membered ring of the β-lactam antibiotics is formed is a central question in their biosyntheses. For penicillin and cephalosporins, which are derived from penicillin, an oxygenase requires ferrous ion and one molecule of dioxygen to remove four hydrogens from a tripeptide precursor to create the bicyclic core of isopenicillin N (Scheme 1, 1) at a single active site. The considerable strain energy imparted to penicillin during its formation is accounted for thermodynamically in the concomitant reduction of oxygen to two molecules of water for every bicyclic β-lactam formed.[11] A mechanistically distinct solution, however, has evolved in the biosynthesis of the potent β-lactamase inhibitor clavulanic acid[2,3] (2) and the clinically important carbapenem antibiotics, for example, thienamycin[4,5] (3) and its simplest relative, carbapenem-3-carboxylic acid[6,7] (4). The first committed step to clavulanic acid is the thiamin-dependent condensation of α-glyceraldehyde-3-phosphate (G3P) and L-arginine to N⁵-(2-carboxyethyl)-arginine[8–10] (5, CEA). In the critical β-lactam-forming reaction, this β-amino acid is activated with ATP (adenylation) by β-lactam synthetase (β-LS), which then carries out a cyclization reaction to deoxyguanadinoprolavaminic acid (6, DGPC) with loss of AMP[11] (Scheme 2). A similar process occurs in the biosynthesis of carbapenem-3-carboxylic acid (4) during which (2S,5S)-5-carboxymethyl proline (7) is transformed to (3S,5S)-carpen-2-am-3-carboxylate (8) by carbapenem synthetase (CPS).[12] Here, the energetic cost of β-lactam formation is paid by the hydrolysis of ATP. Against expectation, however, protein conformational changes were discovered to be substantially rate-controlling in these two enzyme-catalyzed reactions rather than solely the chemical step of four-membered ring closure.[13,14] In fact, in wild-type β-LS, and presumably CPS by comparison, it was shown that the forward rate of β-lactam formation is so great at high pH that acyl-adenylation is rendered effectively irreversible.[14] We were intrigued by this outcome and have turned to computational methods, site-directed mutagenesis, and isotope incorporation experiments to understand the mechanism of β-lactam closure mediated by β-LS, and to illuminate how the energy barrier to this process is lowered in

Scheme 1. β-Lactam-containing natural products.
the enzyme to be comparable to a protein conformational change.

Steady-state kinetic methods have been applied to β-LS to show that an ordered Bi-Ter mechanism takes place in which ATP is the first substrate of two to bind, and pyrophosphate is the last of three products to dissociate after DGPC formation.[11] A series of crystallographic snapshots has been obtained at key points in the reaction coordinate. In addition to structures of 1) the resting enzyme and 2) β-LS-ATP, 3) a ternary complex of β-LS, CEA and the nonreactive analogue of ATP, AMP-CPP (β-LS-CEA-AMP-CPP) was obtained. The latter revealed a high degree of substrate preorganization for in-line attack of the CEA carboxylate on ATP, and a favorable gauche binding geometry of the substrate for β-lactam formation.[15, 16] 4) The lower homologue of CEA, carboxymethyl-L-arginine (CMA) fortuitously underwent reaction with ATP in the crystal to reveal the adenylated intermediate and PPI coordinated to two Mg$^{2+}$ ions. 5) Finally, a fifth structure was obtained with β-LS bound to all three products DGPC, PPI and AMP.[17] In these structures there is a disordered loop (residues 444–453), which becomes organized over the active site when it contains the adenylated intermediate or the final products. A tyrosyl-glutamyl catalytic dyad visible in these X-ray snapshots has been shown to deprotonate the secondary amine of the β-amino acid substrate of not only β-LS but also CPS to initiate cyclization of the four-membered rings.[17] Moreover, two additional residues potentially important for the chemical step of β-lactam formation were apparent in the β-LS structures. At the N terminus of the mobile loop lies Lys443, the ε-ammonium of which is in close proximity to the carbonyl of the adenylated CMA (N – O = 3.5 Å) and the product, DGPC (N–O = 3.0 Å). The second, His447, is located on the catalytic loop, and its ε-ammonium is near the carboxymethyl side chain C1 of the adenylated CMA intermediate suggesting its possible role as a catalytic base.

Studies of the mechanism of acyl substitution reactions are well-developed in the literature.[18–20] We wished to consider all possible routes that could account for the low-energy path acting in enzyme-catalyzed β-lactam formation. For highly reactive acyl groups, for example, acyl halides, or, by analogy, acyl-adenylates in the present instance, elimination to a ketene is a possible reaction path (Scheme 3, path A).[23, 24] The efficient intramolecular attack of a secondary amine onto a ketene in this manner has been exemplified in a close precedent.[25] In the X-ray structure of β-LS-CMA-AMP-PPI a distance of 4.7 Å can be measured from the histidine nitrogen to the α-carbon of the activated carboxymethyl side chain. The distance to this methane hydrogen will be greater than, and therefore will overestimate, the actual distance to the ethylene hydrogen of the native β-LS substrate, CEA (5), which contains an additional methylene. The crystal structure of β-LS and the final products showed the relevant histidine nitrogen further from the α-carbon of the β-lactam ring in DGPC (6; 5.6 Å). The placement of His447 prior to β-lactam formation suggests that this active site base could be well-positioned to catalyze such an elimination process. Moreover, in-plane attack by the β-nitrogen in CEA can be visualized to be geometrically attainable on the sp-hybridized central carbon of the hypothetical ketene intermediate for β-lactam formation.

Alternatively, a highly reactive acyl intermediate could ionize directly to an acylium ion (Scheme 3, path B).[26–28] Like a ketene, this highly electrophilic species would similarly present an in-plane orbital arrangement for cyclization. It is known in the acid-catalyzed hydrolysis of β-lactams that N-protonation leads to the reversible scission of the lactam bond, which results in a secondary amine and an acylium ion. To complete the overall hydrolysis reaction, water then adds to the acylium intermediate.[29, 30] In the present instance, however, ionization of the intermediate acyl-adenylate would give the acylium ion in the presence of a secondary amine. These species would lead to the formation of the protonated β-lactam but in the added presence of AMP dianion, which would rapidly deprotonate the N-protonated β-lactam (pK_a = 0 to −3)[30] driving the equilibrium in the direction of ring closure, the microscopic reverse of this step in hydrolysis.

More conventional routes to β-lactam formation would invoke nitrogen addition to the acyl-adenylate and the transient formation of a tetrahedral intermediate (Scheme 3, path C) followed by AMP loss, or a more concerted process involving only a tetrahedral transition state (Scheme 3, path D). From the X-ray structures it can be hypothesized that Lys443 could stabilize the formation of such a tetrahedral transition state or intermediate by hydrogen bonding or proton donation to the carbonyl oxygen.

Results and Discussion

Consideration of a ketene and acylium ion intermediates

To probe the possible involvement of a ketene mechanism (path A), a deuterium-incorporation experiment was carried out. Recombinant β-LS was diluted approximately tenfold in 99.9% D_2O and used to initiate synthesis in a preincubated
assay mixture containing ATP/Mg$^{2+}$, DTT, and CEA at pD 7.8. These assay components were exchanged in 99.9% D$_2$O and lyophilized three times prior to the addition of enzyme to ensure >95% deuterium in the reaction mixture. Previous solvent isotope effect experiments ensured that the active site of β-LS is accessible for isotopic exchange.$^{[17]}$ As a control, a reaction in 100% H$_2$O (pH 7.8) was performed in parallel. ESI-MS analysis from isolated DGPC in both reactions gave clear parent ions (m/z 229) accompanied by indistinguishable isotopic clusters indicating no incorporation of deuterium during β-lactam formation.

A pH versus log $k$ _cat_ profile of β-LS revealed catalysis to be dependent on the deprotonation of an ionizable group with a pK$_a$ of 8.1.$^{[14]}$ To further test if His447 is a catalytic base and responsible for the observed pK$_a$ in the wild-type enzyme, this residue was mutated to alanine. The mutant remained catalytically active showing a 20-fold decrease in $k$ _cat_. The kinetic data were fitted to Equation (1):

$$\log k_{cat} = \log ([x]_{max} + [x]_{min} \times 10^{pK_a}) / (1 + 10^{pK_a})$$

where ($x$)$_{max}$ and ($x$)$_{min}$ correspond to the upper or lower limits of the linear fit, respectively, to give a pK$_a$ 7.9 ± 0.2 in a pH–$k$ _cat_ profile (Figure 1)—a value virtually identical to that of the general base revealed in the wild-type enzyme. His447 is, therefore, a catalytic base.

Scheme 3. Four proposed mechanisms (A–D) from the acyl-adenylate intermediate to β-lactam formation in β-LS. R represents the functional groups found in vivo, while the groups shown in parentheses represent those used in DFT calculations.

Figure 1. The pH versus log $k$ _cat_ profiles at 25 °C of wild-type β-LS (ref. [14]; ■) and His447Ala (▲). Error bars are shown for all data points and do not exceed the size of the data points when not visible.
fore, not a catalytic base and, combined with the absence of deuterium uptake into the product, the possible intermediacy of a ketene in the β-lactam closure step can be excluded.

The four potential reaction pathways depicted in Scheme 3 were also evaluated by DFT methods (see below and the Supporting Information). The ketene (path A) and aciylum (path B) mechanisms were both found to be high-energy alternatives for β-lactam formation and, in accord with the biochemical results above, could be removed from further consideration.

**Lysine-assisted β-lactam formation**

The placement of Lys443 in β-LS crystal structures suggested its importance in β-lactam formation. While its mutation to alanine and methionine in CPS afforded no detectable activity,[21] in vitro kinetic assays of the Lys443Arg β-LS mutant at the wild-type pH optimum showed a large drop in $k_{cat}$ along with a diminished forward commitment to catalysis after acyladenylation.[19]

In the present study, upon pH variation with the Lys443Arg mutant, it became clear that the pH optimum of this mutant had shifted significantly higher than that of wild-type β-LS and lay above the experimental limit of the buffer system (pH 10.5). While the change in the $k_{cat}$ of Lys443Arg relative to that of the wild type optimal activity (1.1 s$^{-1}$) is 1400-fold lower at pH 7.8, this ratio shrinks to 80-fold at pH 10.5; this illustrates that factors governing DGPC formation become significantly more favorable in the Lys443Arg mutant as pH is increased (Table 1). From the kinetic parameters it was also clear that the $K_{M,CEA}$ of this mutant steadily increased from 120 μM at pH 7.8 to 950 μM at pH 10.5, which implies that the protonated form of Lys443 is more favorable for substrate capture, presumably owing to substrate charge stabilization as suggested previously in the X-ray structural analysis.[31]

The pH dependence of enzyme kinetic parameters reveals ionizable group(s) important for catalysis, binding, and enzyme conformation. While the pH–(1/$k_{cat}/K_{M}$) profile is relatively pH independent (slope = 0.17), and could represent numerous groups contributing to substrate capture in the Lys443Arg mutant, the pH versus log $k_{cat}$ profile is affected by group(s) that influence catalysis (Figure 2). The Lys443Arg pH–log $k_{cat}$ profile was best fit to Equation (1) with an $R^2$ of 0.996, from which was calculated a single $pK_a$ of 10.13 ± 0.06. This value was two $pK_a$ units higher than that of wild type ($pK_a = 8.1$). The significantly increased $pK_a$ is consistent with the newly introduced arginine residue being titrated, rather than the original lysine. In water, a lysine residue ($pK_a = 10.5$) is predicted to have a $pK_a$ two units lower than that of arginine ($pK_a = 12.5$). While a $pK_a$ of 8.1 is lower than the reference value of 10.5 in water, the $pK_a$ values of residues in enzyme active sites typically vary from their standard values owing to a lower dielectric environment in the protein interior.[31,32] This will cause the $pK_a$ values of bases to be lower than expected. In the case of β-LS, the catalytic loop shields the active site from the surrounding aqueous medium during catalysis and a perturbation in $pK_a$ would be expected. Together, these considerations support Lys443 as the ionizable group the deprotonation of which is essential to β-LS catalysis with the observed change in $pK_a$ consistent with the substitution of arginine for lysine at this position. The ability of arginine to act as a base in enzymatic reactions is preceded in several other systems.[23]

To account for the behavior of the pH–$k_{cat}$ profile in β-LS, it is possible that the β-LS mechanism is similar to an isomerase,[34] as previously suggested.[14] In this way, proton assistance by Lys443 would occur during β-lactam ring closure and its charge rebound or reproto- nation is coupled to its rate-determining conformational change (i.e., relaxation of the active-site loop).[14] This sort of active site “recharging” is preceded in kinetic studies of aspartyl proteases[35–37] and triose-phosphate isomerase.[38,39] Perhaps the mutation of Lys443 to a residue with a higher $pK_a$ (i.e., arginine) disrupts this loop relaxation, causing four-membered ring formation to now be fully rate determining. Since vis-

![Figure 2](https://www.chembiochem.org/)

**Figure 2.** The pH-rate profiles of Lys443Arg mutant. A) The pH versus log $k_{cat}$ and B) $k_{cat}/K_{M}$ profiles at 25 °C of wild-type β-LS (see ref. [14]) and Lys443Arg. Error bars are shown for all data points and do not exceed the size of the data points when not visible.

**Table 1.** Comparison of pH—dependent kinetic parameters$^{[a]}$ of β-LS and mutant Lys443Arg.

<table>
<thead>
<tr>
<th>pH</th>
<th>Wild-type</th>
<th>Lys443Arg</th>
<th>Wild-type</th>
<th>Lys443Arg</th>
<th>Fold change $\Delta k_{cat} + \Delta K_{M}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.8</td>
<td>0.43 ± 0.01</td>
<td>0.00079 ± 0.00001</td>
<td>0.057 ± 0.007</td>
<td>0.12 ± 0.01</td>
<td>540 ± 2</td>
</tr>
<tr>
<td>8.8$^{[a]}$</td>
<td>0.82 ± 0.01</td>
<td>0.00161 ± 0.00001</td>
<td>0.044 ± 0.002</td>
<td>0.19 ± 0.02</td>
<td>510 ± 4</td>
</tr>
<tr>
<td>9.3$^{[a]}$</td>
<td>1.01 ± 0.03</td>
<td>0.0036 ± 0.0001</td>
<td>0.082 ± 0.007</td>
<td>0.24 ± 0.02</td>
<td>280 ± 3</td>
</tr>
<tr>
<td>9.8</td>
<td>1.08 ± 0.03</td>
<td>0.00615 ± 0.000004</td>
<td>0.26 ± 0.02</td>
<td>0.40 ± 0.01</td>
<td>170 ± 1.5</td>
</tr>
<tr>
<td>10.0</td>
<td>1.09 ± 0.05</td>
<td>0.0083 ± 0.0002</td>
<td>0.57 ± 0.06</td>
<td>0.37 ± 0.02</td>
<td>130 ± 1.5$^{[a]}$</td>
</tr>
<tr>
<td>10.2</td>
<td>n.a.</td>
<td>0.0120 ± 0.0003</td>
<td>n.a.</td>
<td>0.57 ± 0.05</td>
<td>n.a. n.a.</td>
</tr>
<tr>
<td>10.5</td>
<td>n.a.</td>
<td>0.0140 ± 0.0004</td>
<td>n.a.</td>
<td>0.95 ± 0.07</td>
<td>n.a. n.a.</td>
</tr>
</tbody>
</table>

$^[a]$ Wild-type parameters at all pH values and mutant parameters at pH 8.8 and 9.3 are from previously published kinetic data (ref. [14]). $^[b]$ Value reflects a fold decrease in $K_{M,CEA}$ of Lys443Arg relative to wild type. Approximate fold decreases or increases are shown as $\Delta$ or $-\Delta$, respectively, with wild-type kinetic values divided by mutant kinetic parameters to give the values shown; n.a.: not applicable.
cosity variation at pH 9.3 \(^{[14]}\) and 10.4 (see the Supporting Information) had no significant effect on the \(k_{cat}\). It ensured that the chemical step alone is rate-determining throughout the relevant pH range for this mutant and not a viscosity-dependent conformational change.\(^{[40]}\) This observation is unlike the large viscosity dependence determined for wild type at basic pH values.\(^{[14]}\)

In accord with these kinetic measurements, the catalytic role of Lys443 can be visualized in the crystal structures of the β-LS active site. This series of static images illustrates that prior to chemistry, Lys443 is in closer proximity to the oxygen of the α-phosphate in AMP-CPP than to the carboxylate awaiting amidation. After reaction with ATP, the Lys443 primary amine is then 3.5 Å away from the activated carbonyl oxygen of the acyl-adenylate.\(^{[15]}\) In X-ray snapshots of the β-LS-DGPC-AMP-PP, complex, this nitrogen moves even closer to the β-lactam oxygen.\(^{[15]}\)

**Computational analysis of β-lactam formation**

Computational methods were guided by the X-ray data and allowed the prediction of geometries and high-energy intermediates, which were not obtainable from experimental methods alone. Since density functional theory (DFT) has been commonly used to answer questions related to β-lactams, we turned to this type of theoretical analysis to evaluate the role of Lys443 in DGPC formation using the model 4-(methylamino)butan-2-one-1-(methyl)-hydrogenphosphate (9, Figure 3) to represent the acyl-adenylate of CEA. The adenylated-CEA model bears similarity to CEA (5), but with a truncated structure for ease of computation. Since the arginine side chain of CEA and the adenosine of AMP are constrained in all collected β-LS crystal structures and do not participate in the catalytic chemistry, these moieties were abbreviated in the CEA model.

β-LS-CMA-AMP-PPi and β-LS-DGPC-AMP-PPi structures revealed that Lys-N–O1 distances were in the range of 3–4 Å. Using these observations, the potential energy surface was computed, which considered an accurately positioned lysine relative to 9. The donation of a lysyl proton to the model showed that this event was accompanied by a reduction of electron density at O1 and O2 as revealed by natural bond order (NBO) charges. The reduced negative charge suggests that proton assistance from lysine is not dominated by electrostatics, but rather by proton transfer. For example, the NBO charge\(^{[41]}\) on O1 of the CEA model prior to proton transfer (−0.64) is reduced compared to its corresponding charge (−0.55) after this event (see the Supporting Information). Compound 10 represents the resulting complex that contains the interacting proton donated from lysine (see the Supporting Information).

Since Mg\(^{2+}\) is thought to aid in stabilizing the phosphate negative charge of the adenylated CEA in β-LS, a control for 10 was studied (10b), which replaced the proton on the oxygen of P1 that was originally used to stabilize this negative charge in the calculations. The similar values in Table 2 imply that model 10 is qualitatively reasonable with respect to 10b. Moreover, the energetics involved in the initial step from 10 and 10b leading to a β-lactam are comparable and within 1–4 kcal mol\(^{-1}\). The B3LYP calculations by using the 6-31G(d) basis set and the 6-311+G(d,p) basis set with diffuse functions (see the Supporting Information).

The data in Table 2 also show that the B3LYP/6-31G(d) level describes well the gauche conformation as illustrated in 10. The 10 and 10b structures were compared to the X-ray crystal

![Figure 3. ChemDraw (left) and GaussView (right) generated structures for 9 (bond distance is in Å).](image)

**Table 2. Calculated and experimental values of truncated CEA structural parameters.**\(^{[a]}\)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Bond distance [Å]</th>
<th>Bond angle [degrees]</th>
<th>Dihedral angle [degrees]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C1–O1</td>
<td>C1–N4</td>
<td>C1–C2</td>
</tr>
<tr>
<td>10</td>
<td>1.20</td>
<td>2.71</td>
<td>1.50</td>
</tr>
<tr>
<td>10b</td>
<td>1.20</td>
<td>3.09</td>
<td>1.50</td>
</tr>
<tr>
<td>CEA crystal structure</td>
<td>(1.26)</td>
<td>(3.03)</td>
<td>(1.54)</td>
</tr>
</tbody>
</table>

\(^{[a]}\) Experimental values are in parentheses.
data of the preorganized CEA, which is positioned for in-line attack on the α-phosphorus of the ATP analogue, AMP-CPP.\cite{20}
The gauche conformation is best shown when comparing the calculated C1–C2–C3–N4 dihedral angle of 10 (−49.8°) and 10b (−69.4°), which are similar to the X-ray value (−53.9°). While the calculated O1–C1–C2 bond angles in 10 (128.5°) and 10b (129.1°) are slightly overestimated compared to that of the CEA crystal structure (118.7°) the bond distances match well the preorganization of CEA observed in the X-ray data.

The binding of CEA in the gauche conformation in the β-LS active site reduces a significant energetic cost compared to its anti form. DFT calculations estimated that this high-energy conformer of CEA is about 24 kcal mol\(^{-1}\) less stable than the corresponding anti form. The large energy difference suggests that this conformational modification imparted by β-LS acts in concert with Lys443 to reduce the energetic barrier to β-lactam formation.

The potential energy surface from 10 to the monocyclic β-lactam shown in 13 was computed and is illustrated in Figure 4. The DFT-calculated energy profile agrees with the experimental evidence, in which β-lactam formation in β-LS is not a high-energy process, but rather an overall exothermic process. This landscape also revealed that the interacting proton initially resides on O2, since the extent to which the proton is transferred is greater at P1=O2 than carbonyl C1=O1. However, an energetically low-lying process is computed from 10 to 11 representing an intramolecular proton transfer from O2 to O1 with an activation barrier of 4 kcal mol\(^{-1}\).

Conservation of the catalytic lysine in β-lactam synthetases
Clavulanic acid, (5R)-carbapen-3-em carboxylate, and thienamycin biosyntheses constitute pathways distinct from that of the well-studied penicillin/cephalosporin.\cite{42,43} The sequence homology among these β-lactam-closing enzymes suggests that these ATP-driven steps share common mechanistic features. The genes bla and thn encode the β-lactam synthetases in clavulanic acid and thienamycin biosyntheses, respectively. Both carA and cpmA genes encode the β-lactam-synthesizing enzymes in the formation of (5R)-carbapen-3-em carboxylate.\cite{7,44} Interestingly, the gene cluster of the monocyclic tabtoxin harbors tblS (orf10)—the corresponding protein of which also has sequence homology to β-LS (Bls2; 24% identity, 42% similarity).\cite{45}

From multiple-sequence alignments of the β-lactam synthetases in clavam, carbapenem, and tabtoxin biosynthesis it is evident that the catalytic lysine (Lys443 in β-LS) is strictly conserved (Figure 5). The data suggest that the role of Lys443 in stabilizing the transition state of β-lactam formation in β-LS is similarly retained among all of its homologues, and this point is underscored by mutagenesis studies of Lys443 in CPS\cite{13} and β-LS.\cite{14}

Moreover, it is noteworthy that Lys443 lies immediately before the catalytic loop in β-LS (residues 444–453), and is part of the corresponding loop in CPS. Hence, the protonation state of the conserved lysine could influence the interconversion of the open and closed forms of this active-site feature. In addition to structural evidence from β-LS and CPS, primary-sequence alignments suggest that this loop is also conserved in their homologues CpmA and ThnM (Figure 5).

Figure 4. B3LYP/6-31G(d) and PCM/B3LYP/6-31G(d) calculated potential energy surface for the conversion of intermediate 12 to β-lactam 15. The plotted diagram represents the relative energies calculated by using B3LYP/6-31G(d). Gas-phase energies are in brackets, aqueous-phase energies are in parentheses, bond distances are in Å, and energies are in kcal mol\(^{-1}\).
Conclusions

Site-specific mutation, pH, kinetic and computational studies implicate Lys443 in the activation of N^2-(2-carboxyethyl)-arginine (5, CEA) to facilitate formation of the β-lactam deoxyguaninoprolavaminic acid (6, DGPC). Isotope incorporation and kinetics experiments exclude ketene or acylimidate intermediates (Scheme 3, paths A and B). Simulations of these chemically precluded but high-energy pathways served as controls for the DFT methods employed in this paper, which identified a significantly lower energy process in which intramolecular acyl substitution by Lys443 act in a coordinated fashion to lower the barrier of the catalytic dyad, and key transition state stabilization by a catalytic strategy is general among the members of this family of β-lactam-synthesizing enzymes. While ATP activation affords an excellent leaving group, substrate preorganization, and participation of a catalytic dyad, and key transition state stabilization by Lys443 act in a coordinated fashion to lower the barrier of β-lactam cyclization to become energetically comparable to that of a conformational change involving an active-site loop to allow product release.

Experimental Section

Materials: Most assay components and buffers were purchased from Sigma Chemical, Co. (St. Louis, MO, USA). D₂O was purchased from Cambridge Isotope Laboratories (Andover, MA, USA). CEA and DGPC were synthesized as described previously. Overlap extension was used to introduce desired β-LS mutations in PCR amplifications with appropriate PCR primers for His447Ala (Table S1 in the Supporting Information). The pET29b/Lys443Arg construct along with overproduction and purification techniques are described elsewhere.

Steady-state continuous assays: Initial velocities were measured by using a coupled-enzyme assay that quantitates the formation of authentic DGPC standard by using previously published protocols. The supernatant was flash frozen with liquid nitrogen in 100–200 μL aliquots and stored at −20 °C for later HPLC analysis.

Deuterium incorporation: In the β-LS-catalyzed reaction, possible deuterium incorporation at the α-carbon of β-lactam DGPC was analyzed in a reaction containing >95% D₂O.

Steady-state kinetic analysis: All kinetic data were fit by using Kaleidograph 4.0. Initial velocity patterns were fit to Equation (2) with A representing CEA.

\[ \frac{v}{[E_0]} = \frac{(k_{cat}A)}{(K_m + A)} \]
Theoretical methods: DFT calculations were performed with the use of the Gaussian 03 program package.[47] Geometry optimizations, vibrational frequencies, and intrinsic reaction coordinates (IRCs) were conducted by using DFT with the exchange correlation of B3LYP along with the Pople basis set 6-31G(d).[48–50] The B3LYP functional is not without deficiencies,[51–53] for example, barrier heights can be underestimated by 4–5 kcal mol\(^{-1}\) based on a database analysis of 76 diverse barrier heights, which is due to a self interaction error in local DFT.[54,55] The average errors of B3LYP in thermochemistry calculations are –3.6 kcal mol\(^{-1}\) (from a database of 177 reactions).[56–58] Nonetheless, DFT calculations have been successfully applied to \(\beta\)-lactam systems for predicting structures and conformations, and energetics of reactions, such as ring-opening and decarboxylation.[59–61] The DFT calculations reproduce the gauche conformation relevant in the crystal data of (\(\beta\)-LS-CEA-AMP-CPP; PDB ID: 1JGT). A proton is used to replace magnesium present in \(\beta\)-LS crystal structures, which stabilizes its negative charge in the active site of \(\beta\)-LS. Comparison of this model to one with magnesium showed a similar energetic pathway to \(\beta\)-lactam formation (see the Supporting Information). While the substrate resides in a protein environment that is not expected to be similar to bulk aqueous solution, from ordering of an active site loop,[62] solvent effects were also calculated with the polarized continuum model (PCM) and self-consistent reaction field (SCRF) single point calculations at the B3LYP/6-31G(d) level. The compounds optimize to minima or maxima on the potential energy surface. Molecular structures were viewed with the GaussView program.[64]

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**Keywords:** beta-lactam · biosynthesis · clavulanic acid · density functional theory · pH-rate profiles


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