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Spectroscopic and X-ray Crystallographic Characterization of Bestatin Bound to the Aminopeptidase from *Aeromonas (Vibrio) proteolytica*

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SUBJECTS:  
Peptides and proteins, Metals, Amines, Ions

Abstract

Binding of the competitive, slow-binding inhibitor bestatin ([(2S,3R)-3-amino-2-hydroxy-4-phenylbutanoyl]-leucine) to the aminopeptidase from Aeromonas proteolytica (AAP) was examined by both spectroscopic and crystallographic methods. Electronic absorption spectra of the catalytically competent [Co_(AAP)], [CoCo(AAP)], and [ZnCo(AAP)] enzymes recorded in the presence of bestatin revealed that both of the divalent metal ions in AAP are involved in binding bestatin. The electron paramagnetic resonance (EPR) spectrum of the [CoCo(AAP)]−bestatin complex exhibited no observable perpendicular- or parallel-mode signal. These data indicate that the two CoII ions in AAP are antiferromagnetically coupled yielding an S = 0 ground state and suggest that a single oxygen atom bridges between the two divalent metal ions. The EPR data obtained for [CoZn(AAP)] and [ZnCo(AAP)] confirm that bestatin interacts with both metal ions. The X-ray crystal structure of the [ZnZn(AAP)]−bestatin complex was solved to 2.0 Å resolution. Both side chains of bestatin occupy a well-defined hydrophobic pocket that is adjacent to the dinuclear ZnII active site. The amino acid residues ligated to the dizinc(II) cluster in AAP are identical to those in the native structure with only minor perturbations in bond length. The alkoxide oxygen of bestatin bridges between the two ZnII ions in the active site, displacing the bridging water molecule observed in the native [ZnZn(AAP)] structure. The M−M distances observed in the AAP−bestatin complex and native AAP are identical (3.5 Å) with alkoxide oxygen atom distances of 2.1 and 1.9 Å from Zn1 and Zn2, respectively. Interestingly, the backbone carbonyl oxygen atom of bestatin is coordinated to Zn1 at a distance of 2.3 Å. In addition, the NH₂ group of bestatin, which mimics the N-terminal amine group of an incoming peptide, binds to Zn2 with a bond distance of 2.3 Å. A combination of the spectroscopic and X-ray crystallographic data presented herein with the previously reported mechanistic data for AAP has provided additional insight into the substrate-binding step of peptide hydrolysis as well as insight into important small molecule features for inhibitor design.

Aminopeptidases catalyze the hydrolysis of a wide variety of N-terminal amino acids from proteins and polypeptide chains (1–3). Most enzymes in this group have broad substrate specificities and are widely
distributed in both plant and animal tissues (2). Their biological and medicinal significance is extensive because of their many roles in the degradation of proteins and biologically active peptides, including hormones. The importance of understanding the reaction mechanism of aminopeptidases is underscored by their central role in several disease states including stroke, diabetes, cancer, HIV, bacterial infections, and neuropsychiatric disorders associated with the dysregulation of glutamatergic neurotransmission, such as schizophrenia, seizure disorders, and amyotrophic lateral sclerosis (ALS) (4–6). Recently, it has been shown that several naturally occurring hydroxyethyl isostere dipeptide metalloaminopeptidase inhibitors (i.e., bestatin, leuhistin, and actinonin) inhibit matrix degradation and invasion of extracellular matrixes by fibrosarcoma cells as well as decrease HIV viral load (6, 7). For these reasons, several metallopeptidases have become the subject of intense efforts in inhibitor design (8–13).

Aminopeptidases that have dimetallic active sites can be split into two distinct groups based on their active site structures (8). The first group includes the leucine aminopeptidases from bovine lens (bLAP),1 porcine kidney, tomato, and *Escherichia coli* (PepA) (Figure 1), while the second group contains the leucine aminopeptidases from *Aeromonas (Vibrio) proteolytica* (AAP) (14) and *Streptomyces griseus* (SAP) (Figure 1A) (15). AAP, while not a specific pharmaceutical target at this time, contains a dinuclear active site that is superimposable on the vast majority of dinuclear metallopeptidases that are potential pharmaceutical targets such as SAP, the d-aminopeptidase from *Bacillus subtilis* (DppA), the *dapE*-encoded *N*-sucinyl-l,l-diaminopimelic acid desuccinylase (DapE), the *argE*-encoded *N*-acetyl-l-ornithine deacetylase (ArgE), the carboxypeptidase G2 from *Pseudomonas* sp. strain RS-16 (CPG2), and glutamate carboxypeptidase II (GCP-II), sometimes referred to as *N*-acetylated-α-linked-acidic dipeptidase (NAALADase) (8, 14–17. AAP has been structurally characterized to 1.8 Å resolution (1AMP) (18), but recently, its structure was determined to 1.2 Å resolution (19, 20). In both structures, AAP was shown to contain a single globular domain with a centrally located mixed β sheet sandwiched between α helices. The cocatalytic active site contains a (μ-aquo)(μ-carboxylato)dizinc(II) core with a terminal carboxylate and histidine residue at each metal ion resulting in symmetric coordination spheres for the two active site ZnII ions (Figure 1B). Both ZnII ions reside in distorted tetrahedral coordination geometries with a Zn–Zn distance of 3.5 Å. A glutamate residue, Glu151, forms a hydrogen bond with the bridging water molecule, while the second oxygen atom is 3.4 Å from the N of His97, which is a ligand to Zn1.
Aminopeptidase inhibitors with boronic acids (21–23), chloromethyl ketones (24), phosphonic acids (25–27), hydroxamate (28–30), and α-hydroxyamide (24) functional groups have been previously studied. Peptide analogue aminopeptidase inhibitors of microbiological origin have not been examined in detail. One of these, bestatin, is a naturally occurring dipeptide isolated from cultures of *Streptomyces olivoreticuli* (31). Bestatin has been shown to be a competitive, slow-binding inhibitor of both bLAP and AAP with $K_i^*$ values of 1.3 and 18 nM, respectively (32–34). For all known aminopeptidases, X-ray crystallographic data of a bestatin-bound analogue has only been reported for bLAP (35). The X-ray crystal structure of the [ZnZn(bLAP)]−bestatin complex revealed that the N-terminal amino group coordinates to Zn2, while the alkoxide moiety bridges between the two Zn$^{II}$ ions in the bLAP active site. Interestingly, the backbone carbonyl oxygen of bestatin is not bound to the dinuclear metal center in bLAP but is, instead, hydrogen-bonded to the positively charged terminal amine of an active site lysine residue.

In an effort to gain insight into the structure of AAP in the Michaelis complex, we have studied the binding of the competitive, slow-binding inhibitor bestatin to AAP using both spectroscopic and crystallographic methods. The spectroscopic and X-ray crystallographic data presented herein combined with the previously reported mechanistic data for AAP (14) provide additional insight into the substrate-binding step of peptide hydrolysis and important features for inhibitor design.
Materials and Methods

Enzyme Purification.
All chemicals used in this study were purchased commercially and were of the highest quality available. The aminopeptidase from *Aeromonas (Vibrio) proteolytica* was purified from a stock culture kindly provided by Professor Céline Schalk. Cultures were grown according to the previously published procedure (36) with minor modifications (37) to the growth media. Purified enzyme was stored at 77 K until needed.

Spectrophotometric Assay.
AAP activity was measured by the method of Prescott and Wilkes (36) as modified by Baker et al. (22). In this assay, the hydrolysis of 0.5 mM L-leucine p-nitroanilide (L-pNA) (10 mM Tricine at pH 8.0) was measured spectrophotometrically at 25 °C by monitoring the formation of p-nitroaniline. The extent of hydrolysis was calculated by monitoring the increase in absorbance at 405 nm (Δε405 value of p-nitroaniline of 10 800 M⁻¹ cm⁻¹).

One unit is defined as the amount of enzyme that releases 1 μmol of p-nitroaniline at 25 °C in 60 s. Depletion of enzyme-bound zinc or cobalt was prevented by the addition of 0.1 mM ZnSO₄ or CoCl₂ to the buffer. The specific activity of purified ZnII-bound AAP was typically found to be 120 units per mg of enzyme. This value is identical to that reported by Prescott and Wilkes (36). Enzyme concentrations were determined from the absorbance at 280 nm with the value ε₂₈₀ = 41 800 M⁻¹ cm⁻¹ (38). The accuracy of this value was checked by the Edelhoch method (39−41) using a 5:13:2 molar ratio mixture of N-acetyl-L-tryptophanamide/Gly-Tyr-amide/L-cysteine to model AAP. The molar absorptivity determined from this method, ε₂₈₀ = 43 950 M⁻¹ cm⁻¹ for AAP solubilized in 6 M guanidine hydrochloride, was in excellent agreement with the previously reported value by Prescott et al. (38).

Electronic Absorption and Electron Paramagnetic Resonance (EPR) Samples.
[CoZn(AAP)] and [ZnCo(AAP)] were prepared by dialysis for 72 h at 4 °C against 10 mM 1,10-phenanthroline monohydrochloride (o-phen) in 50 mM Hepes buffer at pH 7.5 (42). AAP was then exhaustively dialyzed against metal-free (chelexed) 50 mM HEPES buffer at pH 7.5. Any remaining metal ions were estimated by comparing the activity of the apo enzyme with a sample that had been reconstituted with ZnII. AAP incubated with o-phen typically had less than 5% residual activity after dialysis. Enzyme concentrations for UV−Vis and EPR samples were typically 1−2 mM. All buffers contained 20% 2-propanol to prevent aggregation at high protein concentrations (43, 44). Metal insertion was effected by the direct addition, with efficient mixing, of 1 equiv of MCl₂ (where M = Co or Zn; ≥99.999% CoCl₂, Strem Chemicals, Newburyport, Massachusetts; 99.999% ZnCl₂, Aldrich) followed by an incubation period for 30 min at 20−25 °C. The second metal was then inserted in the same manner, and the electronic absorption spectrum was recorded prior to freezing in liquid nitrogen for EPR spectroscopy. A 5-fold excess of bestatin was introduced onto the inside side wall of an EPR tube, and the enzyme sample was introduced above this as a plug of ~2 cm in length. As earlier work has demonstrated by optical methods (45), violently flicking the above system facilitates rapid and efficient mixing of the reagents and rapid freezing was achieved by plunging the tube into a beaker of a mixture of liquid and solid methanol over liquid nitrogen. These same EPR samples were then used to record electronic absorption spectra of the AAP−bestatin complex.

Spectroscopic Measurements.
All spectrophotometric measurements were performed on a Shimadzu UV-3101PC spectrophotometer equipped with a constant temperature holder and a Haake (Type 423) constant-temperature circulating bath. The use of 200-μL, 1-cm path-length microcuvettes (QS, Hellma) stoppered with rubber septa facilitated the recording of the optical spectra under anaerobic conditions. Subtraction of the absorption spectrum of apo AAP from those of the substituted enzymes was performed using Shimadzu UV-3101 software. Low-temperature dual-mode EPR
spectroscopy was performed using a Bruker ESP-300E spectrometer equipped with an ER 4116 DM dual-mode X-band cavity and an Oxford Instruments ESR-900 helium-flow cryostat as described previously (15). Background spectra recorded on an EPR tube containing buffer were aligned with and subtracted from experimental spectra as in earlier work (45). All AAP EPR samples contained 20% 2-propanol by volume. Signals because of oxygen were occasionally observed in both EPR modes. These signals routinely disappeared upon raising the temperature in the helium cryostat to 125 K for 5 min and recooling. All spectra were recorded at a modulation frequency of 100 kHz and modulation amplitude of 1.26 mT (12.6 G) with a sweep rate of 10 mT s\(^{-1}\). Parallel- and perpendicular-mode EPR spectra were recorded at microwave frequencies of about 9.37 and 9.65 GHz, respectively; precise microwave frequencies were recorded for individual spectra to facilitate \(g\) alignment. Other EPR recording parameters are specified in the figure captions for individual samples. EPR simulations were carried out using XSophe (46).

Crystallization.
AAP was cocrystallized with bestatin using the crystallization conditions reported for the native enzyme (18). Briefly, purified AAP (10 mg/mL) in 10 mM Tris at pH 8.0, 10 mM KSCN, 0.4 M NaCl, and a 4-fold molar excess of bestatin was crystallized by vapor diffusion using 100 mM Tris at pH 8.0, 100 mM KSCN, and 4.5 M NaCl as the precipitating solution. Crystals with dimensions 0.7 × 0.4 × 0.4 mm were obtained in 2 days and were shown to be isomorphous with the native crystals. The crystals belong to space group \(P6_122\) with the following unit cell dimensions: \(a = b = 107.8\ \text{Å},\ c = 102.6\ \text{Å},\ \alpha = \beta = 90^\circ,\ \gamma = 120^\circ\), and one monomer per asymmetric unit.

Data Collection and Processing.
Diffraction data were collected at 4 °C on an R-axis IIC area detector system mounted on a Rigaku RU-200B rotating anode generator operating at 45 kV and 120 mA. One crystal was used for the entire data collection period. The exposures for 25 min were taken with an oscillation step size of 0.5°. A 0.3-mm collimator was used, and the crystal–detector distance was 100 mm. The diffraction data were integrated and scaled using HKL software (47). Data collection and refinement statistics are outlined in Table 1. The data were collected with an extremely high redundancy; 94.2% of the reflections were measured 4 times or more. A total of 33% of the observed reflections were measured 9–12 times. Because the data are so redundant, the \(R_{\text{merge}}\) values were raised artificially. With greater redundancy, the correlation between \(I\) and \(I_{\text{avg}}\) becomes lower and the \(R_{\text{merge}}\) increases. The \(R_{\text{merge}}\) in the outer shell (2.07–2.0 Å) was 68.3%, and the overall \(R_{\text{merge}}\) was 14.8%. The \(I_{\text{avg}}/\sigma(I_{\text{avg}})\) was 3.0 in the outer resolution shell.

**Table 1: Data Collection and Refinement Statistics**

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<td>(b = 107.8)</td>
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<tr>
<td>(c = 102.6)</td>
<td>(a = 107.8)</td>
<td>(b = 107.8)</td>
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<table>
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<td>(I_{\text{s}}/I_{\text{avg}})</td>
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<td>Completeness, Overall (%)</td>
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<td>(R_{\text{merge}}) (Overall) (%)</td>
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<td>Highest Resolution Shell (Å)</td>
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<tr>
<td>Completeness, Outer Shell (%)</td>
<td>71.4</td>
<td>(R_{\text{merge}}) (Outer Shell) (%)</td>
</tr>
</tbody>
</table>
Transfer of the model into the coordinate file was performed using the original resolution range to determine the number of reflections, protein atoms, and water molecules. The resolution range was 10−2.0 Å, with a cutoff of $F / sF = 0$. The R factor was 19.5%, and the R free was 24.5%. The number of reflections was 21,037, the number of protein atoms was 2,211, the number of zinc ions was 2, the number of bestatin atoms was 22, and the number of water molecules was 136. The B factor model was individual. The RMSD from ideality was for bond lengths 0.006 Å, bond angles 1.19 deg, improper angles 0.63 deg, and dihedral angles 25.6 deg. The residues in most favored positions were 87.2%.

Structure Solution and Refinement. Because crystals of the bestatin-inhibited enzyme were isomorphous with those of the native enzyme, the phases from the published native structure (access code 1AMP) (18) were used as the starting model. In this process, the zinc ions and water molecules were omitted from the original coordinate file. All refinement procedures were carried out using the software package X-PLOR (48). An R free (49) data set was made prior to any refinement using 9% of the total reflections. The initial model was subjected to a rigid-body refinement using all reflections in the 20.0−4.0 Å resolution range. Subsequent rounds of positional refinement were carried out using higher resolution data incrementally to 2.0 Å resolution. The R factor and R free at this point were 27.8 and 32.0%, respectively. Difference electron density maps with amplitudes $2F_{\text{obs}} - F_{\text{calc}}$ and $F_{\text{obs}} - F_{\text{calc}}$ were then calculated and showed clear electron density in the active site for the bound inhibitor and the missing Zn$^{II}$ ions. The two zinc ions and several water molecules were added to the model, and the model was subjected to further rounds of positional refinement. A model for bestatin was built using Insight (Accelrys, Inc.) and fit to the observed electron density. Additional water molecules were added to the model using the WATERPICK protocol in X-PLOR. Further rounds of positional refinement as well as overall and individual B-factor refinement protocols resulted in a final structure with an R factor of 19.5% and an R free of 24.5%. In addition to the 291 amino acid residues, the final model contained 2 zinc ions, 136 water molecules, and 22 atoms of the inhibitor molecule. Simulated annealing omit maps, in which the active site region was omitted, were calculated, which confirmed the presence of the bound inhibitor.

Results

Electronic Absorption Spectroscopy.

The electronic absorption spectrum of [CoCo(AAP)] was recorded in both the absence and presence of bestatin (Figure 2), and the absorption because of apo AAP was subtracted in both cases. The spectrum recorded in the absence of the inhibitor is identical to those previously reported for [CoCo(AAP)] (34) and is characterized by a maximum molar absorptivity of $\sim 95 \text{ M}^{-1} \text{ cm}^{-1}$ at 545 nm. Upon the addition of 4 equiv of bestatin, the maximum molar absorptivity decreased slightly to $\sim 80 \text{ M}^{-1} \text{ cm}^{-1}$, indicating that the Co$^{II}$ coordination number does not change. However, three absorption bands at 495 nm (83 $\text{ M}^{-1} \text{ cm}^{-1}$), 525 nm (88 $\text{ M}^{-1} \text{ cm}^{-1}$), and 545 nm (87 $\text{ M}^{-1} \text{ cm}^{-1}$) appear upon bestatin binding, indicating that bestatin interacts with the dinuclear active site of AAP.
Electronic absorption spectra of [Co(AAP)] and [ZnCo(AAP)] were also recorded in the absence and presence of bestatin (Figure 3). Spectra recorded in the absence of bestatin are identical to those previously reported for [Co(AAP)] and [ZnCo(AAP)] after the subtraction of the absorption because of apo AAP (34). When bestatin was added to [Co(AAP)], the maximum molar absorptivity increased from $\sim 50$ to $58 \text{ M}^{-1} \text{ cm}^{-1}$ and the broad absorption band observed for [Co(AAP)] was replaced by three, resolved absorption bands at 495 nm ($54 \text{ M}^{-1} \text{ cm}^{-1}$), 520 nm ($58 \text{ M}^{-1} \text{ cm}^{-1}$), and 545 nm ($45 \text{ M}^{-1} \text{ cm}^{-1}$), similar to the changes observed for [CoCo(AAP)]. On the other hand, when bestatin is added to [ZnCo(AAP)], the overall shape of the absorption band did not change but the maximum molar absorptivity increased slightly from $\sim 30$ to $35 \text{ M}^{-1} \text{ cm}^{-1}$ and the absorption band is blue-shifted from 530 to 520 nm.

EPR Spectroscopy.

The EPR spectrum of [CoCo(AAP)] is shown in Figure 4. Upon the addition of 4 equiv of bestatin, the perpendicular-mode signal observed at 10 K resulting from the two $S = \frac{3}{2}$ Co$^{2+}$ ions is quenched. Examination of the parallel-mode EPR spectrum of [CoCo(AAP)]–bestatin revealed no detectable parallel-mode signal. The EPR spectrum of [CoZn(AAP)] and [ZnCo(AAP)] was also recorded with and without added bestatin (Figure 5). Noticeable differences are present in the spectra of both heterodimetallic forms of AAP upon binding bestatin, namely, the appearance of distinct hyperfine splitting in both samples. The spectra of [CoZn(AAP)] and [ZnCo(AAP)] were also examined at higher temperatures and powers. The maximum normalized signal intensity
was observed at low temperatures (∼5 K) and low microwave powers, indicating that the observed signals are likely the result of an $M_s = |\pm \frac{1}{2}|$ ground-state transition. The observed signals are distorted by saturation at higher powers (≥20 mW) and by relaxation at higher temperatures (≥20 K), and consequently, multiple species could not be reliably resolved by subtraction of spectra recorded under different conditions. However, computer simulation of the EPR signal exhibited by $[\text{CoZn(AAP)}]$−bestatin (Figure 5) suggests that two species are present. One species is axial with no resolved hyperfine structure. The EPR spectrum observed for this species was simulated assuming an isotropic $g$ factor and axial zero-field splitting ($H = SDS + 6\beta g S; S = \frac{3}{2}, g_{iso} = 2.25, D \gg h\nu, E/D = 0$). The second species was highly rhombic and exhibited clearly resolved $^{59}$Co hyperfine splitting ($H = SDS + 6\beta g S + SAl; S = \frac{3}{2}, I(^{59}\text{Co}) = \frac{7}{2}, g_{iso} = 2.38, D \gg h\nu, E/D = 0.24, A(^{59}\text{Co}) = 8.1 \times 10^{-3} \text{ cm}^{-1}$). On the other hand, the EPR spectra of $[\text{ZnCo(AAP)}]$ in the absence and presence of bestatin consisted of multiple unresolved features. Even so, some clear features are present in the EPR spectrum of the $[\text{ZnCo(AAP)}]$−bestatin complex that are absent in the EPR spectrum of $[\text{ZnCo(AAP)}]$. In particular, a high-field shoulder centered at $g_{eff} \sim 1.78$ (3840 G) and a clear hyperfine pattern centered at $g_{eff} \sim 7.8$ (945 G) suggest the formation of a distinct species, with very high rhombicity ($E/D \sim \frac{1}{3}$), low $g$ strain, and modest spin–orbit coupling ($g_{real} \sim 2.4$), upon the addition of bestatin.

![Figure 4](image)

**Figure 4** Perpendicular-mode EPR signal of $[\text{CoCo(AAP)}]$ in 50 mM HEPES buffer at pH 7.5 in the presence (—) and absence (---) of 4 equiv of bestatin. All spectra were recorded at 10 K, 0.2 mW, 1.26 mT modulation amplitude, and 10 mT s$^{-1}$ sweep rate with 20% added 2-propanol by volume.

![Figure 5](image)

**Figure 5** (A) Perpendicular-mode EPR spectra of (a) $[\text{CoZn(AAP)}]$, (b) $[\text{CoZn(AAP)}]$ + bestatin, and (c) computer simulation of b. (B) Perpendicular-mode EPR spectra of (a) $[\text{ZnCo(AAP)}]$, (b) $[\text{ZnCo(AAP)}]$ + bestatin, (c)
X-ray Structure of Bestatin-Bound AAP.
The X-ray crystal structure of the \([\text{ZnZn(AAP)}]−\text{bestatin}\) complex was solved to 2.0 Å resolution and is shown in Figure 6. Inhibitor binding does not introduce major conformational changes to the overall protein structure. The structures of \([\text{ZnZn(AAP)}]\) and the \([\text{ZnZn(AAP)}]−\text{bestatin}\) complex agree with an rms deviation of 0.19 Å for the 291 structurally equivalent \(\text{Ca}\) atoms. Both side chains of bestatin occupy the well-defined hydrophobic pocket adjacent to the dinuclear \(\text{Zn}^{\text{II}}\) active site made up of Leu155, Met180, Ile193, Cys223, Tyr225, Cys227, Met242, Phe244, Phe248, Tyr251, and Ile255. The leucine portion of bestatin extends back toward Leu155, while the phenylalanine ring extends toward Phe244, Tyr251, and Phe248. The C terminus of bestatin forms a hydrogen bond with Tyr225 (2.7 Å), while the two Cys residues at the back of the hydrophobic pocket (Cys223 and Cys227) form a disulfide bond.

![Figure 6 Stereo figure of the active site region of [ZnZn(AAP)] with bestatin bound. The His residues that act as Zn\(^{\text{II}}\) ligands have been omitted for clarity.](image)

The amino acid residues ligated to the dizinc(II) cluster in AAP are identical to those in the native structure with only minor perturbations to the bond lengths (Table 2; Figure 7). Zn1 of the \([\text{ZnZn(AAP)}]−\text{bestatin}\) complex exhibits a distorted trigonal bipyramidal geometry with the N\(^{\varepsilon}\) nitrogen of His256 and the alkoxide oxygen atom of bestatin making up the axial positions. On the other hand, Zn2 resides in a distorted trigonal bipyramidal geometry with the N\(^{\varepsilon}\) nitrogen of His97 and the N-terminal amine nitrogen atom of bestatin making up the axial positions. The angle including \(\text{Namine}−\text{Zn2}−\text{N}\(^{\varepsilon}\) is 161°. Glu152 is coordinated in an asymmetric bidentate fashion to Zn1 with the additional oxygen atom providing a potential sixth ligand at a distance of 2.3 Å. Similarly, Asp179 is coordinated to Zn2 in an asymmetric bidentate fashion with the second oxygen atom residing 2.4 Å from the metal center, providing a potential sixth ligand. The alkoxide oxygen of bestatin bridges between the two \(\text{Zn}^{\text{II}}\) ions in the active site displacing the bridging water molecule observed in the native \([\text{ZnZn(AAP)}]\) structure. The M–M distance for the AAP−bestatin complex is identical to that in native AAP (3.5 Å) with alkoxide oxygen atom distances of 2.1 and 1.9 Å from Zn1 and Zn2, respectively. Interestingly, the backbone carbonyl oxygen atom of bestatin is coordinated to Zn1 at a distance of 2.3 Å. In addition, the NH\(_2\) group of bestatin, which mimics the N-terminal amine group of an incoming peptide, binds to Zn2 with a bond distance of 2.3 Å. Two hydrogen bonds exist between Glu151 and the bestatin molecule. One of the Glu151 oxygen atoms is 3.0 Å from the alkoxide moiety of bestatin, while the second is 3.3 Å from the backbone amide of the inhibitor. Another important hydrogen-bonding interaction is also observed between an oxygen atom of Asp99 and the N\(^{\varepsilon}\) nitrogen of His97 (3.1 Å) forming an Asp-His-Zn triad (50). This distance is longer than those observed in \([\text{ZnZn(AAP)}]\) (2.8 Å), \([\text{ZnZn(AAP)}]−1\)-butanediol boronic acid (BuBA) (2.7 Å), and \([\text{ZnZn(AAP)}]−\text{l-leucine phosphonic acid (LPA)}\) (2.9 Å).
Table 2: Selected Bond Lengths (Angstroms) for the Zinc–Ligand Distances in the AAP–Bestatin Complex

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<th>LPA complex</th>
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Discussion
Electronic absorption spectra of the catalytically competent [Co(AAP)], [CoCo(AAP)], and [ZnCo(AAP)] enzymes recorded in the absence and presence of bestatin suggest that, like the bLAP–bestatin complex (35), both of the divalent metal ions in AAP are involved in binding bestatin. The molar absorptivities observed for each active site CoII ion are also consistent with five-coordinate metal ions; however, the first metal-binding site is much more perturbed than the second. Interestingly, the electronic perturbations observed for [Co(AAP)] and [ZnCo(AAP)],
upon the addition of bestatin, are essentially opposite to those observed for LPA binding to Co\textsuperscript{II}-substituted AAP (51, 52). The X-ray crystal structure of the [ZnZn(AAP)]−LPA complex revealed that LPA interacts with both metal centers, but a single oxygen-atom bridge is absent (Figure 1C) (52). However, the N-terminal amine of LPA binds to the second metal center with a bond distance of 2.1 Å. Thus, LPA binds to AAP as an η\textsubscript{1,2-μ}-phosphonate with one and two phosphate oxygen atom ligand to the first and second metal ions, respectively, along with the N-terminal amine nitrogen atom bound to Zn\textsuperscript{2}. On the basis of these data, the electronic absorption spectra of Co\textsuperscript{II}-substituted forms of AAP in the presence of bestatin are consistent with bestatin contributing two strongly bound ligands to the first metal ion but only one strong ligand to the second metal-binding site.

EPR spectra of [Co(AAP)] in the absence and presence of bestatin were recorded in both the perpendicular and parallel modes. In the absence of bestatin, the perpendicular-mode EPR spectrum reveals an $M_s = |±\frac{1}{2}\rangle$ ground-state transition with an isotropic $g_{\text{real}} = 2.25$ and $E/D = 0.095$, which accounts for only 13% of the expected spin density, suggesting that the two Co\textsuperscript{II} ions are spin-coupled in the majority of enzyme molecules, as previously reported (43). A parallel-mode signal is observed at neutral pH but is abolished at pH 10. These data were interpreted as being a consequence of there being a weakly bound μ-(H\textsubscript{2}O) bridge between the metal ions in the active site of the resting enzyme and a strongly bound μ-(OH) bridge in the inactive enzyme at high pH (20). The addition of bestatin to [CoCo(AAP)] completely abolished the perpendicular-mode signal. These data suggest that the two Co\textsuperscript{II} ions in AAP−bestatin complex are strongly spin-coupling; therefore, bestatin binds to both Co\textsuperscript{II} ions. Examination of the [CoCo(AAP)]−bestatin complex in the parallel mode provided no detectable signal over a broad range of temperatures and powers. These data suggest that the two Co\textsuperscript{II} ions in AAP are antiferromagnetically coupled yielding an $S = 0$ ground state. On the basis of the reported magnetic properties of several μ-aquo or μ-hydroxodicobalt(II) model complexes, one would expect weak to moderately strong antiferromagnetic coupling in systems containing a single oxygen-atom bridge (53−55). Conversely, systems lacking a strongly bound single atom bridge, such as bis(μ-carboxylato) and tetrakis(μ-carboxylato)dicobalt(II) cores, do not exhibit strong antiferromagnetic spin coupling (56−58). Interestingly, the (μ-phosphonato)dicobalt(II) core of the AAP−LPA complex exhibits ferromagnetic coupling (48, 49). A combination of these data suggests that a single oxygen-atom bridge is present in the bestatin−AAP complex that can be assigned to bestatin alkoxide group, similar to the bLAP−bestatin structure (35).

EPR data obtained on heterodimetallic forms of AAP ([CoZn(AAP)] and [ZnCo(AAP)]) serve both to confirm that bestatin interacts with both metal ions and to support the contention that a bestatin-derived single oxygen-atom bridge exists between the Co\textsuperscript{II} ions in solution as well as in the crystalline state. EPR spectra of [CoZn(AAP)] and [ZnCo(AAP)] consist of a pH-dependent mixture of two species (44). For each of these heterodimetallic forms of AAP, an axial species with low spin−orbit coupling and high $g$ strain dominates at pH 7−8, suggesting that the active nucleophile is present as a water molecule. At higher pH, a rhombic species with a high degree of spin−orbit coupling and very little $g$ strain, indicated by the well-resolved $^{59}$Co hyperfine structure, is the dominant species. This species is, therefore, electronically asymmetric and highly geometrically constrained as would be expected for a strongly bound bridging ligand such as μ-OH. Upon the addition of bestatin to [CoZn(AAP)] and [ZnCo(AAP)], there is clearly an increase in the proportions of the rhombic species exhibiting the well-resolved hyperfine structure relative to the poorly resolved axial species. These data indicate that bestatin interacts with both metal ions and strongly suggest the replacement of water with a more strongly bound ligand such as the alkoxide functionality of bestatin.

Spectroscopic studies suggest that bestatin bridges between the two active site metal ions of AAP in solution via a μ-alkoxide moiety but provides no detailed information regarding any interaction of the N-terminal amine with the second divalent metal ion in the active site. To further characterize the binding of bestatin to AAP, the enzyme was crystallized in the presence of bestatin and the X-ray crystal structure was determined to 2.0 Å resolution (Figures 6 and 7). The AAP−bestatin complex provides structural information that can be used as a
model for inhibitor design, as well as the enzyme–substrate complex. Like all of the structures reported for AAP–inhibitor complexes, the phenyl group of bestatin resides in a well-defined hydrophobic pocket adjacent to the dinuclear Zn$^{II}$ active sites (18, 59). This hydrophobic pocket has been shown to be the initial substrate recognition and binding point on the enzyme, which constitutes the first step in the catalytic mechanism (60). Bestatin binding to AAP does not alter the Zn$^{II}$–Zn$^{II}$ distance but does change the coordination number of both Zn$^{II}$ ions to five coordinate, consistent with the electronic absorption data.

No active site ligands are displaced upon bestatin binding except the water molecule that bridges the two Zn$^{II}$ ions in the wild-type structure. This bridging water molecule is displaced by the alkoxide oxygen of bestatin, consistent with EPR data obtained on the [CoCo(AAP)]$^{-}$–bestatin complex. Interestingly, the Zn1–O1 distance is 2.1 Å, while the Zn2–O distance is 1.9 Å. Therefore, the alkoxide oxygen atom appears to interact more strongly with Zn2 than with Zn1, suggesting that Zn2 plays a role in substrate activation as well as stabilization of the transition state. The bestatin carbonyl group (O2) binds to Zn1, albeit weakly, with a bond distance of 2.3 Å. Surprisingly, the N-terminal amine nitrogen of bestatin (N1) resides 2.3 Å from Zn2, suggesting that it is not coordinated or is very weakly interacting with Zn2. These data are consistent with both the electronic absorption and EPR data, indicating that the two metal ions in the AAP–bestatin complex are perturbed in a very similar manner but that the proportion of the “new” species is much higher for M1 than for M2. The EPR data also show that M2 of the AAP–bestatin complex is not as perturbed as M2 in the AAP–LPA complex. A comparison to the transition-state analogue inhibitor structure of [ZnZn(AAP)]$^{-}$–LPA suggests that the bestatin structure best represents the substrate-binding step and further supports the proposal that carbonyl binding occurs before N-terminal amine binding to the dinuclear active site (61, 62).

The orientation of the N-terminal amine group in the AAP–bestatin structure is similar to the AAP–LPA structure, but the electron orbital overlap observed between Zn1 and the N-terminal amine nitrogen atom in the AAP–bestatin complex is smaller, resulting in a longer M2–amine–N1 distance. However, the orientation of the N-terminal amine of bestatin is distinctly different than that observed for AAP bound by d-iodophenylalaninehydroxamate (d-IPH) (59). In the d-IPH complex, the N-terminal amine group is not present in the active site pocket but instead forms a hydrogen bond with Tyr-225 near the surface of the enzyme at the mouth of the active site pocket. Conversely, the N-terminal amine in the LPA complex is coordinated to Zn2 with a bond distance of 2.1 Å, while it resides 2.4 Å away from Zn2 in the bestatin structure. Therefore, the inhibitor conformation observed in the AAP–d-IPH complex is an inaccurate representation of the actual transition state of peptide hydrolysis, i.e., the stereochemistry of the d-IPH inhibitor is incorrect for an actual peptide because IPH resides in the d configuration. With this in mind, little mechanistic information can be gleaned from the [ZnZn(AAP)]$^{-}$–d-IPH structure because d-IPH is simply acting as a metal chelator, rather than mimicking either substrate binding or the transition state of the catalytic reaction. Consequently, any mechanistic conclusions derived from the AAP–d-IPH structure likely led to an incorrect assignment for the role of the second metal ion in the peptide hydrolysis reaction catalyzed by AAP (37, 43). On the basis of the structures of the AAP–bestatin and AAP–LPA complexes, Tyr225 is not involved in the binding the N-terminal amine of the peptide substrate. In fact, this residue is not likely involved mechanistically but, instead, may contribute only as another large hydrophobic residue used to line the active site pocket.

Glu151 of the active site pocket has been implicated in the catalytic mechanism based on the close proximity of its side chain to the bridging water molecule in the native enzyme (59) and to the bridging phosphonate moiety of LPA in the AAP–LPA complex (52). In the AAP–bestatin complex, there are two hydrogen-bonding interactions between Glu151 and the inhibitor. One Glu151 oxygen atom is 3.0 Å from the μ-alkoxide moiety of bestatin, while the second is 3.3 Å from the backbone amide of the inhibitor. Glu151 appears to be well-positioned to function as a proton shuttle in the hydrolytic reaction, where it could be responsible for removing and/or delivering the proton that is transferred to the leaving group (63). It should be noted, however, that bestatin
contains an additional hydroxyl group between the backbone amine and the carbonyl carbon as compared to the natural substrate. Therefore, interactions with Glu151 may differ for substrates versus bestatin, but the interactions observed in the AAP–bestatin complex suggest that two hydrogen-bonding interactions are possible with Glu151. Furthermore, the AAP–bestatin structure suggests that Glu151 can transfer a proton from the nucleophilic hydroxide moiety to the new N-terminal amine. Such a proton transfer must occur because the newly formed N-terminal amine must leave, at a minimum, as an NH$_2$ group.

In conclusion, we have spectroscopically and crystallographically characterized the binding of bestatin to AAP. Because bestatin has been shown to be an effective drug to inhibit matrix degradation and invasion of extracellular matrixes by fibrosarcoma cells as well as decrease HIV viral load in males (6, 7), the binding mode of bestatin to AAP provides a structural foundation for inhibitor design. In addition, mechanistic data pertaining to the substrate-binding step in peptide hydrolysis can also be gleaned. The data presented herein are consistent with backbone carbonyl binding to Zn1 followed by N-terminal amine binding to Zn2. Because AAP is 80% active with only one divalent metal ion, carbonyl binding likely occurs at Zn1. However, the short Zn2–O1 bond distance of 1.8 Å suggests that, if a second divalent metal ion is present, it likely assists in the activation of the backbone carbonyl, making the carbonyl carbon susceptible to nucleophilic attack. Furthermore, these data suggest that Glu151 is the active site acid/base and can transfer a proton to the newly formed N-terminal amine.

References


Taylor, A., Peltier, C. Z., Torre, F. J., and Hakamian, N. (1993) Inhibition of Bovine Lens Leucine Aminopeptidase by Bestatin: Number of Binding Sites and Slow Binding of This Inhibitor, Biochemistry32, 784.


Abbreviations: Hepes, 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; Tricine, *N*-tris[hydroxymethyl]methylglycine; bestatin, [[2S,3R]-3-amino-2-hydroxy-4-phenylbutanoyl]-leucine; LPA, l-leucine phosphonic acid; BuBA, 1-butaneboronic acid; AAP, aminopeptidase from *Aeromonas proteolytica*; bLAP, bovine lens leucine aminopeptidase; α-phen, 1,10-phenanthroline; EPR, electron paramagnetic resonance.