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Analyzing the binding of Co(II)-specific inhibitors to the methionyl aminopeptidases from *Escherichia coli* and *Pyrococcus furiosus*

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Abstract

Methionine aminopeptidases (MetAPs) represent a unique class of protease that is capable of the hydrolytic removal of an N-terminal methionine residue from nascent polypeptide chains. MetAPs are physiologically important enzymes; hence, there is considerable interest in developing inhibitors that can be used as antiangiogenic and antimicrobial agents. A detailed kinetic and spectroscopic study has been performed to probe the binding of a triazole-based inhibitor and a bestatin-based inhibitor to both Mn(II)- and Co(II)-loaded type-I (*Escherichia coli*) and type-II (*Pyrococcus furiosus*) MetAPs. Both inhibitors were found to be moderate competitive inhibitors. The triazole-type inhibitor was found to interact with both active-site metal ions, while the bestatin-type inhibitor was capable of switching its mode of binding depending on the metal in the active site and the type of MetAP enzyme.

Keywords

Hydrolysis, Manganese, Electron paramagnetic resonance, Antibiotics, Electronic absorption

Introduction

Methionine aminopeptidases (MetAPs) represent a unique class of metalloprotease that is responsible for removing N-terminal methionine residues from polypeptide chains [1, 2]. On the basis of fumagillin affinity chromatography and mass spectrometry, MetAPs were identified as the molecular targets of ovalicin, fumagillin, and TNP-470 [3]. The activity of this sesquiterpiene epoxide containing molecule has been evaluated for the treatment of a variety of cancers such as Kaposi's sarcoma, cervical cancer, brain cancer, and renal cell carcinoma [4]. Furthermore, in vivo studies showed that one of the observed effects of the inhibition of MetAPs by antiangiogenesis agents is the failure to expose glycine residues at the N termini of certain signaling proteins involved in cell cycle regulation, preventing myristolyation [5]. Selective inhibition of tumor cells by targeting a common molecule required for signal transduction of numerous growth stimulators provides an effective approach to treat cancer [6, 7, 8, 9, 10]. MetAPs represent just such a target and, therefore, understanding the catalytic mechanism of MetAPs is critically important for the future development of drugs that prevent tumor vasculature formation, growth, and proliferation [11, 12, 13, 14]. In addition, sesquiterpiene epoxide containing molecules have been shown to inhibit the in vitro growth of the malaria parasites *Plasmodium falciparum* and *Leishmania donavani*; therefore, drug design efforts that target MetAPs may have additional applications in the treatment of malaria and leishmaniasis [15].

MetAPs are organized into two classes (types I and II) on the basis of the absence or presence of an extra 62 amino acid sequence (of unknown function) inserted near the catalytic domain. The type-I MetAPs from *Escherichia coli* (EcMetAP-I), *Staphylococcus aureus, Thermotoga maritime*, and *Homo sapiens* and the type-II MetAPs from *Homo sapiens* (HsMetAP-II) and *Pyrococcus furiosus* (PfMetAP-II) have been crystallographically characterized [16, 17, 18, 19, 20, 21]. All six display a novel "pita-bread" fold with an internal pseudo twofold symmetry that structurally relates the first and second halves of the polypeptide chain to each other. Both domains contribute conserved residues to the metallo-active site. In all six structures, a bis(μ -carboxylato)(μ -aquo/hydroxo)dicobalt core is observed with an additional carboxylate residue at each metal site and a single histidine bound to Co1. An X-ray crystal structure of EcMetAP-I was recently reported with partial occupancy (40%) of a single Mn(II) ion bound in the active site [22]. This structure was obtained by adding the transition-state analog inhibitor I-norleucine phosphonate, to impede divalent metal binding to the second site, and by limiting the amount of metal ion present during crystal growth. This structure provides the first structural verification that MetAPs can form mononuclear active sites and the single divalent metal ion resides on the H171 side of the active site as previously predicted by ¹H NMR and extended X-ray absorption fine structure spectroscopy [23, 24].

A current challenge facing the rational design of highly potent inhibitors of MetAPs is that some candidate compounds fail to have similar efficacy between in vitro and in vivo experiments [25]. Such differences between

in vivo and in vitro effectiveness have been partially ascribed to the lack of certainty about the identity of the in vivo metal ion for MetAPs and its stoichiometry [25, 26]. The dissimilarity in metal identity and its occupancy number within the active site may have subtle effects on the catalytic mechanism of MetAPs, which manifests itself in significant differences in the efficacy of potential inhibitors. It has been established that MetAPs can be activated by Co(II), Fe(II), and Mn(II), but that Zn(II) will not activate either type-I or type-II enzymes [23, 26, 27, 28]. However, a single study on the type-I MetAP from Saccharomyces cerevisiae suggested that Zn(II) ions, in the presence of 5 mM glutathione and 2 mM EDTA under aerobic conditions, provided an active MetAP enzyme, while the Co(II)-containing enzyme was largely inactivated under similar conditions and no activity could be observed in the presence of Fe(II) [29]. Even more recently, the physiologically relevant metal ion for HsMetAP-II was shown to be Mn(II) using Co(II)-specific synthetic inhibitors to probe intracellular HsMetAP-II (Fig. 1) [25]. To understand the mechanism of inhibitor binding to MetAP and to determine why a triazole-containing inhibitor, 3-((2-naphthylmethyl)sulfanyl)-4H-1,2,4-triazole (A-310840), strongly inhibits Co(II)-loaded HsMetAP-II but not Mn(II)-loaded HsMetAP-II, while a bestatin-like inhibitor, ((2RS,3R)-3-amino-2-hydroxy-5-ethylthio)pentanoyl-((S)-(-)-(1-naphthyl)ethyl)amide (A-311263), inhibits each metal-substituted HsMetAP-II enzyme equally well, we have investigated the binding affinities and structural aspects of each of these inhibitor interactions with EcMetAP-I and PfMetAP-II.



Fig. 1. Molecular structures of the triazole-like (*A310840*) and the bestatin-like (*A311263*) methionine aminopeptidase (MetAP) inhibitors

Materials and methods

Purification of MetAP enzymes

All chemicals used in this study were purchased commercially and were of the highest quality available. Recombinant EcMetAP-I was expressed and purified as previously described [23, 27, 28]. Purified EcMetAP-I exhibited a single band on sodium dodecyl sulfate polyacrylamide gel electrophoresis. Protein concentrations were obtained spectrophotometrically at 280 nm using an extinction coefficient $\varepsilon_{280 nm}$ of 16,445 M⁻¹ cm⁻¹. Apo-EcMetAP-I was washed free of methionine using Chelex-100-treated methionine-free buffer [25 mM *N*-(2-hydroxyethyl)piperazine-*N*'-ethanesulfonic acid (HEPES), pH 7.5, 150 mM KCI] and concentrated by microfiltration using a Centricon-10 (Amicon, Beverly, MA, USA) prior to all kinetic assays. Individual aliquots of apo-EcMetAP-I were routinely stored at -80 °C or in liquid nitrogen until needed.

PfMetAP-II was purified as previously reported and exhibited a single band on sodium dodecyl sulfate polyacrylamide gel electrophoresis [28]. Protein concentrations were estimated from the absorbance at 280 nm using an extinction coefficient $\epsilon_{280 \text{ nm}}$ of 21,650 M⁻¹ cm⁻¹. Metal-free PfMetAP-II was prepared by concentrating

purified PfMetAP-II to a volume of approximately 5 mL, after which EDTA was added to a final concentration of 10 mM. The resulting protein solution was dialyzed against 25 mM HEPES buffer (2 L, pH 7.5) containing 10 mM EDTA and 150 mM KCl at 4 °C for 2 days with two buffer changes per day. The protein solution was then dialyzed against Chelex-100-treated 25 mM HEPES buffer (2 L, pH 7.5) containing 150 mM KCl for 3 days against two buffer changes per day. The resulting PfMetAP-II was inactive and found to contain no detectable metal ions via inductively coupled plasma atomic emission spectrometry. This enzyme, deemed "apo-PfMetAP-II," was stored at –80 °C until needed.

Synthesis of the triazole MetAP inhibitor A-310840

The inhibitor A-310840 was synthesized as previously described [25]. Briefly, to a suspension of 3-mercapto-1,2,4-triazole (0.18 g, 1.8 mmol) and cesium carbonate (0.72 g, 2.2 mmol) in 5 mL of *N*,*N*-dimethylformamide was added 2-(bromomethyl)naphthalene (0.38 g, 1.7 mmol). The mixture was heated at 40 °C for 16 h. The volume was reduced by rotary evaporation, and the remaining mixture was shaken with water and methylene chloride and then filtered. The layers of the filtrate were separated, and the organic phase was dried over magnesium sulfate. Filtration and solvent removal gave a white solid (0.143 g). Mass spectrometry (desorption chemical ionization/NH₃) *m/z* 242 (M + H)⁺, 259 (M + NH₄)⁺; ¹H NMR (300 MHz, dimethyl-*d* ₆ sulfoxide, δ) 14.08 (s, 1H), 8.57 (bds, 1H), 7.88 (m, 4H), 7.49 (m, 3H), 4.51 (s, 2H); Anal. Calcd for C₁₀H₂ON₄S: C, 64.70; H, 4.59; N, 17.41. Found: C, 64.93; H, 4.58; N, 17.25.

Synthesis of the bestatin MetAP inhibitor A-311263

The inhibitor A-311263 was synthesized as previously described from d-methionine (Aldrich Chemical Co.) [25]. Mass spectrometry (electrospray ionization + quadrupole 1 MS) m/z 347 (M + H)⁺, 693 (2 M + H)⁺; ¹H NMR (300 MHz, dimethyl- d_6 sulfoxide, δ) 8.75–7.47 (m, 7H), 6.55 (br, 1H), 4.82–4.75 (m, 1H), 4.15 (d, 1H), 3.60–3.33 (br m, 3H), 2.68–2.34 (m, 3.6H), 2.18 (q, 0.4H), 1.87–1.68 (m, 2H), 1.58–1.53 (m, 3H), 1.13 (t, 0.6H), 0.98 (t, 0.4H). Anal. Calcd for C₁₉H₂₇N₂O₂S·HCl: C, 59.59; H, 7.11; N, 7.32. Found: C, 59.54; H, 7.13; N, 7.16.

Enzymatic assay of MetAPs

All enzymatic assays were performed under strict anaerobic conditions in an inert atmosphere glove box (Coy) with a dry bath incubator to maintain the temperature. Catalytic activities were determined with an error of \pm 5%. Enzyme activities for both EcMetAP-I and PfMetAP-II were determined in 25 mM HEPES buffer, pH 7.5, containing 150 mM KCl with the tetrapeptide substrate MGMM. The amount of product formation was determined by high-performance liquid chromatography (HPLC; Shimadzu LC-10A class-VP5). A typical assay involved the addition of 4 µL of metal-loaded MetAP to a 16 µL substrate—buffer mixture at 30 °C for 1 min. The reaction was quenched by the addition of 20 µL of 1% trifluoroacetic acid solution. Elution of the product was monitored at 215 nm following separation on a C8 HPLC column (Phenomenex, Luna; 5 µm, 4.6 cm × 25 cm), as previously described [23, 27, 28]. Enzyme activities are expressed as units per milligram, where one unit is defined as the amount of enzyme that releases 1 µmol of product in 1 min at 30 °C. Inhibition studies were performed by the addition of 8 mM.

Isothermal titration calorimetry

Isothermal titration calorimetry (ITC) measurements were carried out with a MicroCal OMEGA ultrasensitive titration calorimeter. The titrants (A-310840 and A-311263) and apo-MetAP solutions were prepared in Chelex-100-treated HEPES buffer at pH 7.5. Stock buffer solutions were thoroughly degassed before each titration. The enzyme solution (50–80 μ M) loaded with 2 equiv of metal ions [Co(II) or Mn(II)] was placed in the calorimeter cell and stirred at 200 rpm to ensure rapid mixing. Typically, 4–10 μ L of titrant was delivered over 7.6 s with a 6-min interval between injections to allow for complete equilibration. Each titration was continued until 4.5–6 equiv of inhibitor had been added to ensure that no additional complexes were formed in excess titrant. A

background titration, consisting of the identical titrant solution but only the buffer solution in the sample cell, was subtracted from each experimental titration to account for heat of dilution. The data were analyzed with a one- or two-site binding model by the Windows-based Origin software package supplied by MicroCal [<u>30</u>].

Spectroscopic measurements

Electronic absorption spectra were recorded using a Shimadzu UV-3101PC spectrophotometer. All apo-MetAP samples used in spectroscopic measurements were made rigorously anaerobic prior to incubation with Co(II) (CoCl₂ 99.999% or better purity; Strem Chemicals, Newburyport, MA, USA) for approximately 30 min at 25 °C. All Co(II)-containing samples were handled throughout in an anaerobic glove box (Ar/5% H₂, 1 ppm or less O₂; Coy Laboratories) until frozen. Electronic absorption spectra were normalized for the protein concentration and the absorption due to uncomplexed Co(II) (ε_{512} nm = 6.0 M⁻¹ cm⁻¹) [23]. Low-temperature EPR spectroscopy was performed using either a Bruker ESP-300E or a Bruker EleXsys E600 spectrometer, each equipped with an ER 4116 DM dual-mode X-band cavity operating at 9.63 GHz ($B_0 \perp B_1$) or 9.37 GHz ($B_0 \parallel B_1$) and an Oxford Instruments ESR-900 helium-flow cryostat. All spectra were recorded using 100-kHz magnetic field modulation. Other EPR running parameters are specified in the figure legends for individual samples. EPR simulations were carried out using matrix diagonalization (XSophe, Bruker Biospin), assuming a spin Hamiltonian $H = \delta g$ HS + SDS + SAI, with S = 3/2 and $D \gg \delta$ gHS (=hv). Enzyme concentrations for EPR were 1 mM.

Results

Determination of IC₅₀ values

Data on inhibition of MetAP activity by A-311263 and A-310840 (IC₅₀ values) towards the Co(II)- and Mn(II)loaded PfMetAP-II and EcMetAP-I enzymes are provided in Table <u>1</u>. A-310840, a triazole-containing inhibitor, binds potently to PfMetAP-II in the presence of Co(II) (IC₅₀ of 15 μ M) but is approximately 67-fold less potent for Mn(II)-loaded PfMetAP-II (IC₅₀ of 1,000 μ M). A similar trend is observed for EcMetAP-I, except that A-310840 binds to the Co(II)-loaded enzyme well (IC₅₀ of 44 μ M) but is approximately 23-fold less potent for Mn(II)-loaded EcMetAP-I (IC₅₀ of 1,000 μ M). In comparison, the bestatin-type inhibitor, A-311263, has little or no preference for the type of divalent metal ion that occupies the active site. A-311263 also inhibits PfMetAP-II and EcMetAP-I loaded with either Co(II) or Mn(II) similarly, with IC₅₀ values between 20 and 34 μ M.

Table 1. IC₅₀, inhibition constants (K_i), dissociation constants (K_d), stoichiometry (n), and thermodynamics parameters for 3-((2-naphthylmethyl)sulfanyl)-4H-1,2,4-triazole (A-310840) and ((2RS,3R)-3-amino-2-hydroxy-5-ethylthio)pentanoyl-((S)-(-)-(1-naphthyl)ethyl)amide (A-311263) binding to Co(II)- and Mn(II)-loaded type-II methionine aminopeptidase (MetAP) from *Pyrococcus furiosus (PfMetAP-II*) and type-I MetAP from *Escherichia coli (EcMetAP-I*)

Wild-type enzymes	n	IC₅₀ (μM)	<i>Κ</i> i (μΜ)	<i>Κ</i> _d (μΜ)	Δ <i>H</i> (kJ/mol)	<i>T∆S</i> (J/mol)	∆ <i>G</i> (kJ/mol)
A-310840 binding to wild-							
type enzymes							
[CoCo(PfMetAP-II)]	1	15	3	2.0 ± 0.1	-5.5 ± 2	27 × 10 ³	-32.5
[MnMn(PfMetAP-II)]	1	1,000	440	666 ± 10	-0.3 ± 0.1	17 × 10 ³	-18.1
[CoCo(EcMetAP-I)]	1	44	8	10 ± 0.1	-8.36 ± 0.3	20 × 10 ³	-28.5
[MnMn(EcMetAP-I)]	1	1,000	750	714 ± 10	-24.6 ± 3	-7×10^{3}	-17.9
A-311263 binding to wild-							
type enzymes							
[CoCo(PfMetAP-II)]	1	20	2	0.45 ± 0.11	-14.9 ± 2	21 × 10 ³	-36.2
[MnMn(PfMetAP-II)]	1	27	20	26 ± 5	-62.7 ± 5	-36×10^{3}	-26.1

[CoCo(EcMetAP-I)]	1	32	3	2.4 ± 0.1	1.57 ± 0.3	33 × 10 ³	-32
[MnMn(EcMetAP-I)]	1	34	20	50 ± 5	-19 ± 4	6 × 10 ³	-24.5

Kinetic determination of K_i values

Previous work indicated that A-310840 and A-311263 are strong inhibitors of HsMetAP-II on the basis of IC₅₀ values [25]. We have carefully examined the hydrolysis of the tetrapeptide substrate MGMM by both PfMetAP-II and EcMetAP-I in the presence of various concentrations of A-310840 and A-311263. K_i values for A-310840 and A-311263 binding to PfMetAP-II and EcMetAP-I were obtained by fitting the kinetic data to the Michaelis– Menten equation for competitive inhibition (Table 1). A-310840 is a much more potent competitive inhibitor towards Co(II)-loaded PfMetAP-II ($K_i = 3 \mu$ M) and EcMetAP-I ($K_i = 750 \mu$ M) enzymes. On the other hand, A-311263 inhibits Co(II)-loaded PfMetAP-II ($K_i = 2 \mu$ M) and EcMetAP-I ($K_i = 3 \mu$ M) by a magnitude similar to that of Mn(II)-loaded PfMetAP-II ($K_i = 20 \mu$ M) and EcMetAP-I ($K_i = 20 \mu$ M).

Isothermal titration calorimetry

The use of ITC to measure the binding of exogenous ligands to biomolecules relies on the fact that such an interaction is accompanied by a heat effect. The heat absorbed (endothermic) or released (exothermic) upon inhibitor–enzyme interaction is used to obtain the dissociation constant, K d, and the enthalpy of binding, ΔH . ITC measurements for the binding of A-310840 and A-311263 to Co(II)- and Mn(II)-loaded EcMetAP-I and PfMetAP-II were carried out with a MicroCal OMEGA ultrasensitive titration calorimeter at 25.0 \pm 0.2 °C. K d values were obtained by fitting these data, after subtraction of the background heat of dilution, via an interactive process using the Origin software package. This software package uses a nonlinear least-squares algorithm, which allows the concentrations of the titrant and the sample to be fit to the heat flow per injection, to an equilibrium binding equation for either one site or two noninteracting sites. The K_a value, the enzymemetal stoichiometry (n), and the change in enthalpy (ΔH) were allowed to vary during the fitting process. The relationship between K a and the dissociation constant is defined as $K_d = 1/K_a$. The best fits obtained for the binding of A-310840 to Co(II)-loaded PfMetAP-II and EcMetAP-I provided an overall n value of 1 and K d values of 2.0 and 10 μM, respectively, which indicate approximately 333-fold and approximately 71-fold stronger binding than is observed for A-310840 binding to Mn(II)-loaded PfMetAP-II (K_d = 666 μ M) and EcMetAP-I (K_d = 714 μ M). On the other hand, the best fits obtained for the binding of A-311263 to Co(II)-loaded PfMetAP-II and EcMetAP-I provided an overall n value of 1 and K d values of 0.45 and 2.4 μ M, respectively. These values indicate approximately 58-fold and approximately 21-fold tighter binding than is observed for Mn(II)-loaded PfMetAP-II and EcMetAP-I, respectively (Table 1).

The heat of reaction, ΔH , measured during an ITC experiment was converted into other thermodynamic parameters using the Gibbs free energy relationship:

$$\Delta G^0 = -\mathrm{RT} \ln K_{\mathrm{a}} = \Delta H^0 - T \Delta S^0$$
 (2)

The thermodynamic parameters for the binding of A-310840 and A-311263 to Co(II)- and Mn(II)-loaded PfMetAP-II and EcMetAP-I are listed in Table <u>1</u>. Binding of these inhibitors to both type-I and type-II MetAPs was found to be spontaneous on the basis of the observed negative Gibbs free energy (ΔG). In general, the ΔG values for the binding of both inhibitors to PfMetAP-II are slightly more negative than the ΔG values obtained for EcMetAP-I. The energy change associated with Co(II)-loaded EcMetAP-I with A-311263 is weakly endothermic, with a ΔH value of 1.57 kJ mol⁻¹, while in other cases ΔH was found to be weakly exothermic, with values ranging between -0.3 and -63 kJ mol⁻¹. It is noteworthy that the entropic factor ($T\Delta S$) for the binding of each inhibitor to PfMetAP-II and EcMetAP-I varies depending on the enzyme and metal ions present. For example, the $T\Delta S$ values

for the binding of A-310840 to Mn(II)-loaded EcMetAP-I and the binding of A-311263 to Mn(II)-loaded PfMetAP-II are negative, whereas in all other cases the observed $T\Delta S$ values are positive (Table <u>1</u>).

Electronic absorption spectroscopy

The electronic absorption spectra of Co(II)-loaded PfMetAP-II and EcMetAP-I were recorded in the absence and presence of A-310840 and A-311263 (Figs. 2, 3) and the absorption due to apoenzyme was subtracted in each case. The absorption spectra of the uncomplexed wild-type Co(II)-loaded PfMetAP-II and EcMetAP-I enzymes were identical to those previously reported [23, 28]. Upon the addition of 1 equiv of A-310840 to both Co(II)loaded PfMetAP-II and Co(II)-loaded EcMetAP-I, the molar absorptivities increased by approximately 140 M^{-1} cm⁻¹. The observed increase in absorbance at approximately 550 nm is similar in magnitude to that observed at 400 nm, raising the possibility that this increase may be to do to an underlying charge-transfer or $\pi \rightarrow \pi^*$ transition with $\lambda < 400$ nm, rather than a change in absorptivity of the d-d bands from Co(II). Upon closer inspection, the λ_{max} values of the absorption maxima show small blueshifts compared with those of the wild-type enzymes, indicating that A-310840 directly interacts with the Co(II) active sites of both EcMetAP-I and PfMetAP-II. Interestingly, the addition of 1 equiv of A-311263 to both Co(II)-loaded PfMetAP-II and Co(II)-loaded EcMetAP-I also perturbs the visible absorption spectra of Co(II)-loaded EcMetAP-I and PfMetAP-II, but in a way markedly different from that of A-310840. The maximum molar absorptivity decreases upon the addition of A-311263 to Co(II)-loaded PfMetAP-II by approximately 50 M⁻¹ cm⁻¹, but the observed λ_{max} value remains unaffected. The maximum molar absorptivity does not change upon the addition of A-311263 to Co(II)-loaded EcMetAP-I (approximately 90 M⁻¹ cm⁻¹); however, the λ_{max} value shifts from 580 to 560 nm.



Fig. 2. UV–vis spectrum of wild-type [CoCo(PfMetAP-II)] (*dashed line*), A-310840 bound to [CoCo(PfMetAP-II)] (*solid line*), and A-311263 bound to [CoCo(PfMetAP-II)] (*dotted line*) in 25 mM *N*-(2-hydroxyethyl)piperazine-*N*'- ethanesulfonic acid (HEPES) buffer, pH 7.5, and 150 mM KCl. The absorption due to apoenzyme was subtracted in each case. PfMetAP-II is type-II methionine aminopeptidase (MetAP) from *Pyrococcus furiosus*



Fig. 3. UV–vis spectrum of wild-type [CoCo(EcMetAP-I)] (*dashed line*), A-310840 bound to [CoCo(EcMetAP-I)] (*solid line*), and A-311263 bound to [CoCo(EcMetAP-I)] (*dotted line*) in 25 mM HEPES buffer, pH 7.5, and 150 mM KCl. The absorption due to apoenzyme was subtracted in each case. EcMetAP-I is type-II MetAP from *Escherichia coli*

EPR spectra of A-310840 and A-311263 bound to Co(II)- and Mn(II)-loaded PfMetAP-II and EcMetAP-I

EPR spectra for both Co(II)-loaded PfMetAP-II and Co(II)-loaded EcMetAP-I have been reported previously [23]. These spectra contain broad features and exhibit no resolvable ⁵⁹Co hyperfine splitting, presumably owing to extensive strains in q and/or zero-field-splitting parameters. The spectra observed for Co(II)-loaded PfMetAP-II and EcMetAP-I upon the addition of A-310840 or A-311263 are shown in Fig. 4 along with computer simulations. Spin Hamiltonian parameters derived from these simulations are provided in Table 2. In each case, the EPR spectra of the inhibited forms of Co(II)-loaded PfMetAP-II and EcMetAP-I are very similar to the spectrum from the corresponding unligated enzyme. The observed signals are dominated by an essentially axial component with E/D ~ 0.09. For Co(II)-loaded EcMetAP-I + A-310840 (Fig. 4, spectrum B) and Co(II)-loaded PfMetAP-II + A-311263 (Fig. 4, spectrum C), a low-field eight-line hyperfine pattern was resolved, indicating the presence of a second, minor species with markedly lower g_{eff} strain. This signal may also be present in the spectrum of Co(II)loaded PfMetAP-II + A-310840 (Fig. 4, spectrum D), but at very low concentrations, and was not observed for Co(II)-loaded EcMetAP-I + A-311263 (Fig. 4, spectrum A). In the EPR spectrum of Co(II)-loaded EcMetAP-I bound by A-310840 (Fig. 4, spectrum B), another feature associated with this second, minor species is observed at an inflection around 1,900 G. On the basis of this feature, $g_{x,y}$ and hence E/D could be reasonably estimated by simulation, resulting in an $M_s = \pm \frac{1}{2}$ species with E/D = 0.22. These parameters are typical for five-coordinate Co(II) centers bound within a rigid coordination sphere [31, 32, 33]. For Co(II)-loaded PfMetAP-II + A-311263 (Fig. 4, spectrum C), only the hyperfine pattern could be distinguished, at lower field than that observed for Co(II)-loaded EcMetAP-I + A-310840. The actual values of g_{xy} and E/D for this minor species are dependent on the identity of the ground-state doublet and therefore cannot be determined from these data.



Fig. 4. EPR spectra of MetAP species containing Co(II). In *A*–*D*, the upper spectrum is the experimental spectrum and the lower spectrum is a computer simulation: [CoCo(EcMetAP-I)] + A-311263 (*A*); [CoCo(EcMetAP-I)] + A-310840 (*B*); [CoCo(PfMetAP-II)] + A-311263 (*C*); [CoCo(PfMetAP-II)] + A-310840 (*D*). Trace *E* is of naked [CoCo(PfMetAP-II)]. Experimental spectra were recorded at 7 K, 0.2 mW (*A*–*C*); 9 K, 0.2 mW (*D*), and 5 K, 0.2 mW (*E*). Spin Hamiltonian parameters for the simulations are given in Table <u>2</u>

Table 2. Spin Hamiltonian parameters for the dominant axial and minor rhombic species obtained from thecomputer simulation of EPR spectra of Co(II)-loaded EcMetAP-I and PfMetAP-II in the presence of either A-310840 or A-311263

Wild-type enzymes	Axial			Rhombic		
	g(x, y)	g(z)	E/D	g(x, y) [A(y) (10 ⁻³ cm ⁻¹)]	g(z)	E/D
A-310840 binding to wild-type						
enzymes						
[CoCo(PfMetAP-II)]	2.21	2.09	0.09	ND	ND	ND
[CoCo(EcMetAP-I)]	2.316	2.650	0.095	2.52 [8.6]	2.56	0.22
A-311263 binding to wild-type						
enzymes						

[CoCo(EcMetAP-I)]	2.316	2.650	0.095	ND	ND	ND
[CoCo(PfMetAP-II)]	2.21	2.09	0.09	2.5–2.9 [10.6]	ND	ND

ND not determined

In addition to the S = 3/2 signals observed for each of the Co(II)-loaded MetAP-inhibited complexes, weak signals were observed between $g_{\text{eff}} = 10$ and 12 in parallel mode (not shown) for Co(II)-loaded PfMetAP-II and EcMetAP-I bound by A-311263 and Co(II)-loaded EcMetAP-I + A-310840. These data are consistent with S' = 2 and/or S' = 3 transitions from an S = 0, 1, 2, 3 spin ladder due to spin coupling between two S = 3/2 Co(II) ions. While quantification of broad, weak integer spin signals of indeterminate S' origin and with absorption into zero field is unreliable, integration of the S = 3/2 components based on simulation of the observed spectra suggested that approximately $85 \pm 10\%$ of the Co(II) ions are accountable for by the observed S = 3/2 species. The observed integer-spin signals are particularly pronounced for Co(II)-loaded PfMetAP-II + A-311263, and the same integer-spin transitions are likely due to the EPR absorption at very low field (0–500 G) observed in normal-mode EPR (Fig. 4, spectrum C; a small amount of such a signal was included in the simulation but parameters are not given as no attempt was made to fully characterize or quantify this signal). It is unlikely that these signals are dependent upon the inhibitor as similar low-field signals are observed for uncomplexed Co(II)-loaded PfMetAP-II (Fig. 4, spectrum E) and EcMetAP-I enzymes at high pH. Indeed, the lack of such signals from Co(II)-loaded PfMetAP-II + A-310840 may be of more significance.

The EPR spectra of Mn(II)-loaded EcMetAP-I and PfMetAP-II complexed with either A-310840 or A-311263 are presented in Fig. <u>5</u>. When Mn(II)-loaded EcMetAP-I and PfMetAP-II were incubated with A-310840, both enzymes displayed signals superficially similar to those of the corresponding wild-type enzymes. Characteristic central six-line patterns with approximately 90 G splitting and Curie law temperature dependence are observed, as well as additional features that demonstrate Boltzmann temperature dependence and an approximately 45 G splitting [25, 26]. The central six-line patterns observed for Mn(II)-loaded EcMetAP-I and PfMetAP-II bound by A-310840 are somewhat lower in intensity compared to those of wild-type forms of the enzymes, but the individual lines are better resolved. However, the resonance positions of the transitions in the 3,000–3,800-G region of the spectra are indistinguishable (Fig. <u>5</u>, spectra F, H). Both sets of spectra provide other features with Boltzmann temperature dependence that exhibit approximately 45 G splitting, characteristic of dinuclear Mn(II) centers (Fig. <u>5</u>, spectra I, K, L, N).



Fig. 5. EPR spectra from MetAP containing Mn(II). Traces A-E are 6,000-G (600-mT) experimental scans. Traces *F*, *G*, and *H* and *I*, *J*, and *K* are expanded views of selected regions of *A*, *B*, and *C*, respectively. Traces *L*, *M*, and *N* are derivatives of traces *I*, *J*, and *K*, respectively. The traces correspond to [MnMn(EcMetAP-I)] + A-310840 (*A*, *F*, *I*, *L*), [MnMn(EcMetAP-I)] + A-311263 (*B*, *G*, *J*, *M*), [MnMn(Pf(MetAP-II)] + A-310480 (*C*, *H*, *K*, *N*), and [MnMn(Pf(MetAP-II)] + A-311263 (*D*). Trace *E* is a computer simulation of trace *D* assuming *S* = 5/2, *g* = 2.0, *A* $I = 5/2 = 8.8 \times 10^{-3} \text{ cm}^{-1}$, $D = 86 \times 10^{-3} \text{ cm}^{-1}$, $\sigma D = 7 \times 10^{-3} \text{ cm}^{-1}$, E/D = 0, and $\sigma E/D = 0.14$. Experimental spectra were recorded at 30 K, 0.2 mW (*A*–*C*, *F*–*H*, *I*–*K*) or at 40 K, 0.2 mW (*D*)

In contrast, the addition of A-311263 to Mn(II)-loaded EcMetAP-I and PfMetAP-II provided EPR spectra that are distinctly different from those of wild-type Mn(II)-loaded enzymes. The spectrum of Mn(II)-loaded EcMetAP-I + A-311263 contained a central six-line pattern (Fig. 5, spectrum G) indistinguishable from the patterns of the A-310840 complexes and wild-type enzymes. However, this signal did not exhibit the Boltzmann-temperature-dependent "wings" at 2,200–2,600 G with approximately 45 G splittings that are characteristic of Mn(II)–Mn(II) spin–spin coupling (Fig. 5, spectra J, L). More dramatic differences were seen in the spectrum of Mn(II)-loaded PfMetAP-II + A-311263 (Fig. 5, spectrum D). A very complex spectrum was observed that had intense features across the field range 0–6,000 G. The form of this spectrum was essentially invariant from 5 to 40 K and exhibited Curie law temperature dependence, suggesting that the features are due to isolated S = 5/2 Mn(II) ions. Indeed, this signal could be well simulated assuming a single Mn(II) species with spin Hamiltonian parameters S = 5/2, g = 2.0, $A_{I=5/2} = 8.8 \times 10^{-3}$ cm⁻¹, $D = 86 \times 10^{-3}$ cm⁻¹, $\sigma D = 7 \times 10^{-3}$ cm⁻¹, E/D = 0, and $\sigma E/D = 0.14$ (Fig. 5, spectrum E).

Discussion

Type-II MetAPs have been identified as the molecular target for antiangiogenesis agents such as fumagillin and its derivatives [3, 34], whereas type-I MetAPs have been suggested to be potential antibacterial and/or antifungal targets [35, 36]. Therefore, developing a detailed understanding of how potential small molecule drug

candidates interact with the active sites of MetAPs will facilitate the rational design of new, potent MetAP inhibitors with improved in vivo stability and cytotoxicity [37]. As one might expect, most of the information regarding the binding of MetAP inhibitors has come from the inspection of X-ray crystal structures [22, 38, 39]. Although such crystallographic information is extremely valuable, it represents a static view that may not be representative of a dynamic enzyme. Thus, detailed solution characterization of MetAP drug candidates is critically important in the design and development of new drugs. Herein we have investigated the binding affinities and the spectroscopic properties of two inhibitors of PfMetAP-II and EcMetAP-I, namely, a triazole-containing inhibitor 3-((2-naphthylmethyl)sulfanyl)-4H-1,2,4-triazole (A-310840) and a bestatin-like inhibitor ((2RS,3R)-3-amino-2-hydroxy-5-ethylthio)pentanoyl-((S)-(-)-(1-naphthyl)ethyl)amide (A-311263) [25].

Examination of the inhibition patterns of both Co(II)- and Mn(II)-loaded EcMetAP-I and PfMetAP-II in the presence of A-310840 and A-311263 revealed that they are both moderately strong competitive inhibitors. A-310840 preferentially inhibits Co(II)-loaded EcMetAP-I and PfMetAP-II enzymes over the Mn(II)-loaded enzymes, whereas A-311263 shows little or no metal ion selectivity for either type-I or type-II enzymes. The observed K_{i} values follow the same trend as the IC₅₀ values obtained for EcMetAP-I, PfMetAP-II, and HsMetAP-II except that the IC_{50} values are substantially weaker for the two bacterial enzymes compared with the human type-II enzyme [25]. For competitive inhibitors, the observed K_i value is equal to the dissociation constant (K_d); therefore, ITC not only provides the opportunity to compare experimentally derived K d values with K i values, but also to obtain thermodynamic parameters for inhibitor binding to both the PfMetAP-II and EcMetAP-I enzymes in a single experiment. The K d values derived from ITC are in reasonable agreement with the kinetically derived K i values (Table 1). The observed enthalpies (ΔH) for the binding of A-310840 and A-311263 to Mn(II)-loaded PfMetAP-II and EcMetAP-I are negative. However, both A-310840 and A-311263 exhibited positive ΔH values for binding to Co(II)-loaded EcMetAP-I but negative ΔH values for binding to Co(II)-loaded PfMetAP-II. These data exemplify the active-site differences between type-I and type-II MetAPs and indicate that Mn(II)-loaded MetAPs more favorably bind these inhibitors, and possibly substrates. Interesting, the entropic factor ($T\Delta S$) follows an opposite trend, with inhibitor binding to the Co(II)-loaded enzymes being more entropically favored than inhibitor binding to Mn(II)-loaded MetAPs. Combination of the observed ΔH and T ΔS values for A-310840 and A-311263 binding to both the Co(II)-loaded and the Mn(II)-loaded PfMetAP-II and EcMetAP-I enzymes are all negative, indicating that inhibitor binding is spontaneous. However, the observed ΔG values favor inhibitor binding to the Co(II)-loaded PfMetAP-II and EcMetAP-I enzymes over the Mn(II)-loaded forms by nearly 30%.

To gain some structural insight into how A-310840 and A-311263 bind to both type-I and type-II MetAPs in the presence of either Mn(II) or Co(II), electronic absorption and EPR spectra were recorded in the presence of each inhibitor. The observed changes in electronic absorption spectra of the Co(II)-loaded forms of PfMetAP-II and EcMetAP-I upon A-310840 and A-311263 binding indicate that each ligand interacts with the active-site Co(II) ions. Moreover, the observed molar absorptivities upon A-310840 binding to Co(II)-loaded EcMetAP-I and PfMetAP-II suggest that the active-site Co(II) ions are five-coordinate, while A-311263 binding provides a Co(II) environment that is five- or six-coordinate [40]. Similarly, the observed EPR spectra of the Co(II)-loaded EcMetAP-I and PfMetAP-II enzymes in the presence of A-311263 are due to transitions in the $M_s = \pm \frac{1}{2}$ Kramers doublets and arise from either five- or six-coordinate Co(II) ions The broad, featureless EPR signals from the dominant axial components of the spectra indicate significant structural microheterogeneity in each case; thus, there is a great deal of flexibility in the ligand environment, perhaps related to coordination by water. For Co(II)loaded EcMetAP-I and PfMetAP-II enzymes in the presence of A-310840 a minor species, in some cases with well-resolved hyperfine splitting and significant rhombic distortion of the axial zero-field splitting, is observed, suggesting an alternative geometry with a rigid coordination environment and likely five-coordinate geometry [41, 42]. Similar species observed in the EPR spectra of Co(II)-substituted forms of the aminopeptidase from Aeromonas proteolytica have been shown to be pH-dependent [41, 42].

For a small percentage (less than 15%) of Co(II) ions bound to either the EcMetAP-I or PfMetAP-II enzymes in the presence of A-311263 and the Co(II)-loaded form of EcMetAP-I bound by A-310840, a weak integer spin signal in

the parallel mode is observed. However, for Co(II)-loaded PfMetAP-II bound by A-310840, no integer spin signal is observed. EPR studies on Co(II)-loaded EcMetAP-I revealed that at pH 7.5 there is no significant spin coupling between the two Co(II) ions, though a small proportion (approximately 5%) of the sample exhibited detectable spin–spin interactions at pH values above 9.6 [23]. Therefore, the integer spin signals observed for the EcMetAP-I-inhibited enzymes are likely due to inhibitor binding since no spin-spin interactions are observed in the absence of inhibitor. Conversely, the uncomplexed form of Co(II)-loaded PfMetAP-II exhibits detectable (approximately 15%) spin–spin interactions at pH 7.5. Thus, the observed integer spin signal for Co(II)-loaded PfMetAP-II bound by A-311263 is identical to that for the uncomplexed enzyme, suggesting that the binding of A-311263 does not provide a single atom bridge between the two Co(II) ions bound in the active site. However, the lack of an observable integer spin signal when A-310840 is complexed with Co(II)-loaded PfMetAP-II suggests that A-310840 may modulate antiferromagnetic coupling via a diazine bridge. Such a coupling interaction is consistent with the limited number of diazine-bridged dinuclear Co(II) model complexes [43, 44, 45]. The lack of intense parallel-mode signals is likely due to the weak K_d values observed for the second Co(II) binding event (0.35 and 2.5 mM for PfMetAP-II and EcMetAP-I, respectively), which provides enzyme samples containing a mixture of both mononuclear and dinuclear Co(II) centers with the majority of sites in the mononuclear state [23, 28]. For dinuclear Co(II) species, it appears that the binding of either A-311263 or A-310840 modulates the exchange interaction between the Co(II) ions by altering the precise nature of the electronic wavefunction overlap network (i.e., the bonding network) between these ions.

Both PfMetAP-II and EcMetAP-I exhibit distinct binding constants for two Mn(II) ions, and the higher of these, K_{d2} , dictates that distinct MetAP species containing either one or two Mn(II) ions will be present in samples prepared even at millimolar concentrations [22, 46, 47]. The dinuclear forms have been shown to contain two Mn(II) ions in close proximity and, for both PfMetAP-II and EcMetAP-I, features in the EPR spectra have been identified that clearly show spin–spin interaction between the two Mn(II) ions [43, 44, 45, 46]. As expected, EPR spectra obtained for the Mn(II)-loaded forms of both PfMetAP-II and EcMetAP-I in the presence of either A-311263 or A-310840 indicate the presence of magnetically isolated S = 5/2 Mn(II) ions coordinated by oxygen-and nitrogen-containing ligands [46]. The S = 5/2 components of the spectra with A-310840 were indistinguishable from those of the corresponding resting enzymes. The EPR spectra of PfMetAP-II and EcMetAP-I incubated with A-310840 exhibited features that are characteristic of spin–spin coupling between the Mn(II) ions; again, these features were very similar to those exhibited by the corresponding resting enzymes. The lack of significant perturbation of the electronic structure of the monomanganese and dimanganese species of these enzymes by A-310840 is likely related to the poor efficacy of this inhibitor for the manganese forms of PfMetAP-II and EcMetAP-II and EcMetAP-I.

Significant differences in the electronic structures of the Mn(II) complements of PfMetAP-II and EcMetAP-I were elicited by incubation with A-311263 in contrast to incubation with A-310840. In the case of Mn(II)-loaded EcMetAP-I + A-311263, no signals or temperature-dependent features corresponding to spin-coupled Mn(II)– Mn(II) were observed between 5 and 50 K. This observation suggested the loss of any spin—spin interaction. The loss of the S' = 1, 2,..., 5 signals would be expected to result in an increase in the S = 5/2 signal intensity due to the newly separated Mn(II) ions; the loss of the dipolar component of the coupling would, of course, entail physical separation of the two Mn(II) ions. Intriguingly, no significant enhancement of the central six-line pattern signal intensity was observed. Therefore, it is unclear whether no inter-Mn(II) interaction exists or whether the interaction is present but unobservable. An unusually high spin-coupling constant ($|J| \gg 35$ cm⁻¹) would prevent the population of S' > 0 levels owing to zero-field splitting within the non-Kramers doublets of populated S' > 0 states ($\Delta > 0.3$ cm⁻¹), which results from broadening by large increases in the values and/or distributions of the individual Mn(II) zero-field-splitting parameters. Alternatively, a relaxation-dependent phenomenon may be in play. Regardless, the inter-Mn(II) interaction in Mn(II)-loaded EcMetAP-I is significantly perturbed, either abolished or greatly enhanced, by A-311286.

The EPR spectrum obtained for Mn(II)-loaded PfMetAP-II + A-311286 looked completely different from any other EPR signal observed for a MetAP species but was very similar to the EPR signal observed for the *argE*-encoded *N*-acetyl-I-ornithine deacetylase from *E. coli* [47]. Despite the complexity of this signal, it was simulated as a single S = 5/2 Mn(II) environment, consistent with its temperature dependence. The lack of an intense central six-line pattern is due to a distribution in *E/D*; although the system is nominally axially symmetric with a mean *E/D* = 0 assumed for the sample, the distribution results in a significant proportion of the molecules in the sample exhibiting *E/D* \neq 0. Thus, it appears that a rhombic distortion is introduced into the electronic symmetries of the Mn(II) ions bound to PfMetAP-II upon complexation with A-311286. Similar to Mn(II)-loaded EcMetAP-I, no evidence is observed for inter-Mn(II) spin coupling in the EPR spectrum obtained for Mn(II)-loaded PfMetAP-II + A-311286, further supporting the contention that the active site is significantly perturbed by A-311286.

The X-ray crystal structure of A-311263 bound to HsMetAP-II revealed the presence of a μ -alkoxide bridge [48]. In addition, the X-ray crystal structure of a related purported substrate-analog inhibitor (3*R*)-amino-(2*S*) hydroxyheptanoyl-I-Ala-I-Leu-I-VaI-I-Phe-OMe (SK-V-126) bound to EcMetAP-I has been reported [18]. In this structure, the N-terminal amine coordinated Co2, the keto oxygen (analogous to the carbonyl carbon in a terminal peptide bond) is ligated to Co1, and the (2*S*)-hydroxyl group displaces the bridging water/hydroxide ion [1, 18]. Taken together with the EPR data, which primarily contain data for mononuclear enzyme species, this suggests the alkoxide moiety displaces the water (hydroxide) oxygen atom and is terminal on M1, while the N-terminal amine nitrogen of the inhibitor is bound by the carboxylate groups making out the purported second metal binding [22]. Inspection of the X-ray crystal structure of the dicobalt(II) form of type-I MetAP from *S. aureus* complexed with a 1,2,4-triazole-based inhibitor and the crystal structure of a 1,2,3-triazole-based inhibitor bound to the dicobalt(II) form of HsMetAP-II reveals that a common feature for the triazole derivative in enzyme recognition is the interaction of the nitrogen atoms in positions 1 and 2 with the dicobalt(II) center such that their N–N bond is nearly collinear with the metal–metal vector in the dinuclear site [49, 50, 51].

Combination of EPR and crystallographic data suggests that when two metal ions are bound to MetAPs, A-310840 binds via a diazine bridge (Fig. 6). On the other hand, when only one Co(II) ion is bound in a MetAP active site, binding of A-310840 clearly results in a more rigid five-coordinate environment. Conversely, the addition of A-310840 does not markedly perturb the EPR spectra of either Mn(II)-loaded PfMetAP-II or Mn(II)loaded EcMetAP-I. These data indicate nonbinding or a very weak binding interaction of the triazole ligand with the Mn(II) ions bound in the MetAP active sites. Therefore, the specificity of A-310840 for Co(II) over Mn(II) is likely due to Co(II)'s preference for softer donor ligands (nitrogen) over harder donor ligands (oxygen) [52, 53]. However, this explanation does not tell the entire story. Since MetAPs are likely a mixture of mononuclear and dinuclear enzymes under physiological conditions [54, 55] and the putative partially empty, second metal binding site is carboxylate-rich, the lone pair of electrons on the N2 atom of the triazole ligand will be repelled by the negative charge associated with the carboxylate groups in this site. Alternatively, the N-terminal amine moiety of A-311263 will form a zwitterionic interaction with this carboylate site, resulting in an additional binding interaction over the A-310840 molecule (Fig. 7). This hypothesis is consistent with the observed thermodynamic parameters for both A-310840 and A-311263 binding to Co(II)- and Mn(II)-loaded PfMetAP-II or EcMetAP-I, particularly in the decrease in ΔG observed for A-310840 binding to Mn(II)-loaded MetAPs compared with Co(II)-loaded enzymes.



Fig. 6. Proposed scheme for binding of A-310840 to the active sites of both Co(II)- and Mn(II)-loaded EcMetAP-I (residue numbering) and PfMetAP-II



Fig. 7. Proposed scheme for binding of inhibitor A-311263 to the active site of both Co(II)- and Mn(II)-loaded EcMetAP-I (residue numbering) and PfMetAP-II

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