Mechanistic role of each metal ion in *Streptomyces* dinuclear aminopeptidase: Peptide hydrolysis and $7 \times 10^{10}$-fold rate enhancement of phosphodiester hydrolysis

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The dinuclear aminopeptidase from *Streptomyces griseus* (SgAP) and its metal derivatives catalyze the hydrolysis of the phosphoester bis(p-nitrophenyl) phosphate (BNPP) and the phosphonate ester p-nitrophenyl phenylphosphonate with extraordinary rate enhancements at pH 7.0 and 25°C [A. Ercan, H. I. Park, L.-J. Ming, Biochemistry 45, (2006) 13779–13793.], reaching 6.7 billion-fold in terms of the first-order rate constant of the di-Co(II) derivative with respect to the autohydrolytic rates. Since phosphoesters are transition state-like inhibitors in peptide hydrolysis, their hydrolysis by SgAP is quite novel. Herein, we report the investigation of this proficient alternative catalysis of SgAP and the role of each metal ion in the dinuclear site toward peptide and BNPP hydrolysis. Mn(II) selectively binds to one of the dinuclear metal sites (M1), affording Mn-SgAP with an empty (E) second site for the binding of another metal (M2), including Mn(II), Co(II), Ni(II), Zn(II), and Cd(II). Peptide hydrolysis is controlled by M2, wherein the $k_{\text{cat}}$ values for the derivatives MnM2-SgAP are different yet similar between MnCo- and CoCo-SgAP and pairs of other metal derivatives. On the other hand, BNPP hydrolysis is affected by metals in both sites. Thus, the two hydrolytic catalyses must follow different mechanisms. Based on crystal structures, docking, and the results presented herein, the M1 site is close to the hydrophobic specific site and the M2 site is next to Tyr246 that is H-bonded to a coordinated nucleophilic water molecule in peptide hydrolysis; whereas a coordinated water molecule on M1 becomes available as the nucleophile in phosphodiester hydrolysis.

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1. Introduction

Enzymes can effect remarkable rate accelerations by stabilizing the transition-state (TS) of the substrates [1]. The use of TS analogues for the production of catalytic antibodies [2] and inhibition of peptidases and esterases by different phospho-centers [3–6] support the TS theory. However, some phosphoesters and fluorophosphates can be hydrolyzed by serine proteases and esterases via nucleophilic attack by the active-site Ser [7], which nevertheless produces an indefinitely stable dead-end complex with the phospho-center covalently attached to the Ser. Moreover, the phosphoester bis(p-nitrophenyl) phosphate (BNPP) and the phosphonate ester p-nitrophenyl phenylphosphonate can be effectively hydrolyzed by the dinuclear aminopeptidase (AP) from *Streptomyces griseus* (SgAP) with activities comparable to some native phosphoesters [8–10]. Since phospho- and phosphono-esters are TS-like molecules and can inhibit peptide hydrolysis [11–13], their hydrolysis by SgAP is novel and must take place according to a unique catalytic pathway of the enzyme.

SgAP (30 kDa) is a Ca$^{2+}$-influenced extracellular enzyme of high thermal stability with a catalytic specificity toward hydrophobic substrates [14,15]. It has a di-metal center (3.65 Å apart) bound to the protein through the side chains of His85 and Asp160 in one metal site and His247 and Glu132 in another site, along with a bridging Asp97 [16,17]. SgAP and the AP from *Aeromonas proteolytica* (ApAP, 32 kDa) have 29.6% sequence identity, identical metal-binding ligands, and a similar three-dimensional structure [18]. However, ApAP [19] along with mammalian AP-P and *Escherichia coli* Met AP [20–22] require only one metal to activate, and a second metal to modulate, its activity. Conversely, SgAP requires two metal ions for catalysis on the basis of crystallographic, NMR, and kinetic studies [8–10,19,23–25].

The active site of metalloenzymes can be investigated with spectroscopic and kinetic methods by the use of various metal ions [26–29]. Different homo- and hetero-dinuclear derivatives of dinuclear APs, including Leu-specific APs from bovine lens (blLAP) [29]...
and porcine kidney (pkLAP) [30] and ApAP [19], were constructed and investigated to gain further insight into the function of each metal ion in catalysis [19,31]. For example, in bLAP, the first metal-binding site shows higher effect on $k_{cat}$ while the second metal is more influential on $K_m$ [29]. On the other hand, the first Zn$^{2+}$ affects $K_m$ while the second metal affects $k_{cat}$ in pkLAP [30] and both $k_{cat}$ and $K_m$ values are affected by the two metals in ApAP [32]. The variations of the kinetic parameters with the different metal derivatives of these enzymes signifies the importance and the function of each metal ion in their dinuclear active center.

Different metal derivatives of ApAP were prepared and their activity and inhibition by 1-butaneboronic acid investigated, from which a mononuclear mechanism was proposed [33–35]. The crystal structure of ZnZn-ApAP upon binding with this inhibitor shows that the inhibitor binds one metal through two oxygen atoms, one of which may interact with the second metal [36]. Unlike ApAP, SgAP is inactive with only one metal and becomes fully active upon binding of the second metal [23]. In this case, selective Co$^{2+}$ binding has been verified with NMR, showing distinct hyperfine-shifted $^1$H NMR signals for Co,E- and Co,Co-SgAP [23]. Since selective metal-binding of SgAP occurs, different metal ions can be introduced to construct various hetero-dinuclear derivatives. We present herein further investigation of the role of each metal ion in the dinuclear active site in the action of SgAP by the use of various hetero-dinuclear derivatives MnM2-SgAP ($M_2 = Zn^{2+}, Co^{2+}, Ni^{2+}$, or Cd$^{2+}$). The mechanisms of this enzyme toward the hydrolysis of peptides and the alternative hydrolysis of BNPP are proposed.

2. Experimental section

2.1. Materials and reagents

The protease mixture Pronase, the buffers, HEPES, MES, NaH$_2$PO$_4$, EDTA, 1,10-phenanthroline, phenylglyoxylic monohydrate, DEAE-Sephadex and Sephadex G-50, the substrate Leu-$p$-nitroanilide (Leu-$p$NA), Met-$p$NA, Val-$p$NA, Ala-$p$NA, Gly-$p$NA, and BNPP were purchased from Sigma–Aldrich (St. Louis, MO) and the atomic absorption standards (>99.99%) of Zn$^{2+}$, Co$^{2+}$, Mn$^{2+}$, Cd$^{2+}$, Ni$^{2+}$, and Cu$^{2+}$ from Fisher Scientific (Pittsburg, PA). Deionized water of >18 M$^\Omega$ obtained from a MilliQ system (Millipore, Bedford, MA) was used to prepare all the solutions. All the glassware and plastic ware were treated with 2 mM EDTA solution and rinsed with deionized water prior to use.

2.2. Purification, demetalization, and characterization of SgAP

SgAP was purified according to the published procedures [14,23]. The fractions of the first peak from the DEAE-Sephadex column with AP activity were combined. Purified SgAP (0.05–0.1 mM, 10 ml) was dialyzed for 12 h against 250 ml of 20 mM Tris/HCl buffer at pH 7.5 with 100 mM NaCl for four times, with 2.0 mM EDTA and 2.0 mM 1,10-phenanthroline for the first two times and 2.0 mM 1,10-phenanthroline for latter two, followed by dialyzing against 250 ml of 20 mM MES at pH 6.0 for four times to remove the chelators. The AP concentration was determined according to the absorption $E_{280}^{35} = 15$ [14,23] and by titration with atomic-absorption-grade Co$^{2+}$ solution of known concentrations.

The formation of the various MnM-SgAP derivatives was verified by their activities and after prolonged incubation to ensure the retention of the activities (otherwise, migration of metal ions in the two sites may take place). Paramagnetically shifted $^1$H NMR features were also used as the fingerprints for identification of the Co$^{2+}$-containing derivatives [37–39]. The $^1$H NMR spectra of the paramagnetic derivatives were acquired on a Varian INOVA500 spectrometer (at 500 MHz $^1$H resonance) with a 5-mm bio-TR (triple resonance) probe by the use of the build-in polynomial 1–3–3–1 pulse sequence for samples in H$_2$O buffers and a pre-saturation pulse for samples in D$_2$O buffers with a 90° pulse of ~9 µs over 200 ppm spectral width and processed with a line-broadening of 40 Hz, followed by spline baseline correction.

2.3. Enzyme kinetics and inhibition

The kinetics were carried out in 0.1 M HEPES buffer at pH 8.0 containing 0.1 M NaCl and 10 mM CaCl$_2$ at 30 °C and the data were analyzed with the Michaelis–Menten model to derive the turn-over $k_{cat}$ and the Michaelis constant $K_m$ with non-linear fitting of the rate with respect to substrate concentration. Fluoride inhibition was carried out under the same conditions with different inhibitor concentrations, but in the absence of Ca$^{2+}$ to avoid the formation of the very insoluble CaF$_2$. Each inhibition study was performed at least twice and fitted to the Michaelis–Menten equation with non-linear regression and the inhibition constants $K_i$ for different inhibition patterns are determined accordingly.

2.4. pH Profiles and thermostability

Catalytic parameters ($k_{cat}$ and $K_m$) toward hydrolysis of Leu-$p$NA and BNPP and inhibitions were measured at different pHs (acetate at pH 5.0, MES at 5.5–6.5, HEPES at 7.0–8.0, TAPS at 8.5–9.5, and CAPS at 10.0). Thermostability of apo-SgAP and different metal derivatives of SgAP was determined on the basis of their activities toward Leu-$p$NA hydrolysis. Herein, SgAP and its various metal derivatives with or without 5 mM CaCl$_2$ were incubated at various temperatures for 1.0 min followed by incubation on ice for 5 min. Then, an excess amount of corresponding metal was added and activity determined.

3. Results and discussion

3.1. Mn$^{2+}$ binding and hetero-dinuclear active site of SgAP

The binding of Mn$^{2+}$ to apo-SgAP at pH 6.0 requires >50 equivalents to fully activate the enzyme (Fig. 1A), wherein the activity is significantly enhanced by Ca$^{2+}$ (5 mM). At pH 8.0 without Ca$^{2+}$, Mn$^{2+}$ binds to one site exclusively without showing activity (o, Fig. 1B and inset), and reaches full activation at >20 equivalents. Mn$^{2+}$ binding becomes less selective and the enzyme is 44% less active in the presence of Ca$^{2+}$ (●, Fig. 1B and inset). The binding of Mn$^{2+}$ to just one metal-binding site in the active center at one equivalent (denoted the M1 site) allows the introduction of another metal ion to the second site (M2) to construct several catalytically active hetero-dinuclear derivatives in the form of Mn,M2-SgAP (where M2 = Co$^{2+}$, Zn$^{2+}$, Ni$^{2+}$, or Cd$^{2+}$) (Fig. 1C). Conversely, the binding of Zn$^{2+}$, Cd$^{2+}$, or Cu$^{2+}$ to the enzyme exhibit a non-selective pattern [23,24,40], preventing the preparation of the corresponding M1,E-SgAP derivatives with an empty (E) M2 site. The first equivalent of Mn$^{2+}$ must bind to the active site of SgAP. Otherwise, the subsequent addition of one equivalent of Co$^{2+}$ (●, Fig. 1C) would not generate an active derivative since CoE-SgAP is inactive [23].

The derivative MnCo-SgAP shows paramagnetically shifted $^1$H NMR features in the downfield region ~30–80 ppm (Fig. 2, bottom) different from those of CoE- [23] and CoZn-SgAP [41], indicating a new Co$^{2+}$ derivative. The spectrum is also different from that of CoCo-SgAP which indicates that Co$^{2+}$ is not bound to both metal site of the protein (by replacing the Mn$^{2+}$). The NMR signals of the protons around mononuclear Mn$^{2+}$ cannot be detected due to signal broadening by the slowly relaxing unpaired electrons of Mn$^{2+}$. However, magnetic coupling between Mn$^{2+}$ and a metal
ion of fast electron relaxation, such as high-spin Co²⁺ and Fe²⁺, in a dinuclear center can afford hyperfine-shifted 1H NMR signals due to both the Mn²⁺ and the other metal sites [37–39]. Thus, the observed hyperfine-shifted 1H NMR signals herein can only result from the dinuclear MnCo-SgAP. Two solvent exchangeable signals are detected at 59.8 and 62.5 ppm which disappear in the sample prepared in a D₂O buffer (Fig. 2, top). These signals are consistent with the imidazole NH proton of coordinated His residues, one in each metal site (His85 and His247) [16,17]. The broad signals at ~45 ppm seems to be solvent exchangeable; however, the broadness of this signal prevents its assignment at this stage.

In order to use these hetero-dinuclear derivatives for kinetic studies, it is essential to prove that the two metal ions do not switch their binding positions with each other during the time span of the experiment. If metal ions in M1, M2-SgAP would switch their positions, different k_cat and K_m values would be expected. To verify this, the kinetic parameters were determined immediately after MnCo-SgAP was prepared and a day later, which showed no difference. Moreover, 1H NMR does not reveal any hyperfine-shifted 1H NMR signals characteristic of CoE-SgAP or CoCo-SgAP within a week which also supports the conclusion that there is no metal exchange between the two sites in MnCo-SgAP.

A few equivalents of Cd²⁺ are required to saturate the active site of SgAP to exhibit full activity [24]. However, one equivalent of Cd²⁺ is able to activate MnE-SgAP (Fig. 1C), suggesting that binding of Mn²⁺ may organize the active site for better binding of the second metal ion to exhibit activity. Nevertheless, the Cd²⁺ affinity toward MnE-SgAP is still ~400 and ~6000 times weaker than Zn²⁺ and Co²⁺ affinity (Fig. 1C).

The various metal derivatives of SgAP presumably should exhibit thermostability to different extents, thus can be further characterized. Apo-SgAP in the absence of CaCl₂ is the least stable, showing 50% denaturation (T₅₀) at ~60 °C within 1.0 min determined from activity measurements (Fig. 3A). The influence of transition ions on the stability is quite pronounced, wherein the one-metal derivatives CoE- and MnE-SgAP show T₅₀ at ~68 °C; MnCo-SgAP, ~80 °C; and other dimetal derivatives, >85 °C. In the presence of 5.0 mM CaCl₂, all forms of SgAP are dramatically stabilized. For example, apo-SgAP shows T₅₀ at ~83 °C while MnMn-SgAP is still fully active even at 85 °C for a minute. The results support the conclusion from metal-titration experiments that Mn²⁺ indeed binds to apo-SgAP to form the derivative MnE-SgAP which exhibits a thermo-denaturation pattern different from those of MnMn-SgAP and apo-SgAP without Ca²⁺ (Fig. 3A). If the lack of activity of the enzyme with <1.0 equivalent Mn²⁺ (Fig. 1C) is due
to that the metal is not bound to the enzyme, there should be little activity at ~70 °C as that of apo-SgAP, which is not the case.

3.2. Fluoride inhibition

The inhibition by F– toward Leu-pNA hydrolysis by MnMn-SgAP exhibits an uncompetitive inhibition pattern (Fig. 4A), indicating F– binding to the ES complex. Uncompetitive F– inhibition of metalloenzymes has been suggested to be associated with the status of the nucleophilic water [25,42–47]. In the case of urease, purple acid phosphatase, and tyrosinase, F– is proposed to replace the bridging hydroxide. The F– inhibition toward Leu-pNA hydrolysis is uncompetitive at pH 7.5 to 10.0 with a gradual increase in \( K_i \) from 0.24 to 19.4 mM (Fig. 5C), showing an ionization constant \( pK_a \) of 9.2 in the pK\(_{\text{a}}\)-vs.-pH plot. A similar \( pK_a \) is also observed in the pH profile of \( k_{\text{cat}}/K_m \) (discussed below). Because F– replaces prospective nucleophile, the \( pK_a \) profile should reveal the ionization of the amino acid that influences the generation of the nucleophile, likely Tyr246 which has been implied to interact with a coordinated water [16]. Moreover, ApAP does not contain a corresponding Tyr and does not decrease in activity associated with a \( pK_a \) in the alkaline range [48].

F– also inhibits various metal derivatives of SgAP in an uncompetitive manner, wherein the inhibition constant is significantly influenced by the metal ions in the active site (Table 1), more so by M2 as in the influence on the catalysis by M2. For example, the \( K_i \) values of MnMn-, MnNi-, and MnCo-SgAP are quite different, but closer for the counterparts M1, Ni- and Mn1, Co-SgAP (Table 1). Evidences from NMR, EPR, and resonance Raman studies of purple acid phosphatase (PAP) show that binding of F– interrupts the metal-metal interaction, indicating that F– interacts with both metal ions and the nucleophile in PAP (and urease as well) is proposed to be the bridging H\( _2 \)O/OH [44,45]. However, since only one metal (M2) is significantly associated with F– inhibition in the case of SgAP, the nucleophile in this enzyme must be a terminal OH–.

In a recent report, both F– (0–80 mM) and phosphate (0–50 mM) were observed to exhibit noncompetitive inhibition toward native SgAP [49], different from what we observed previously [10] and described herein. The addition of CaCl\(_2\) to the enzyme can activate the enzyme significantly [14], while phosphate or F– therein can presumably remove the added 1 mM Ca\(^{2+}\) due to the very low \( K_{\text{eq}} \) of Ca(HPO\(_4\))/Ca\(_3\)(PO\(_4\))\(_2\) and CaF\(_2\). The removal of Ca\(^{2+}\) would decrease the enzyme activity in a noncompetitive manner. Thus, the study cannot reveal clear inhibition patterns of phosphate and fluoride toward the enzyme itself and is not comparable to what we previously reported [10] and presented herein without the activator Ca\(^{2+}\).

The pH dependence of \( k_{\text{cat}} \) in Leu-pNA hydrolysis by MnMn-SgAP in the range of pH 7–10.5 is controlled by only one ionization constant \( pK_{\text{eq}} = 10.2 \) (Fig. 5A, left scales; Table 2), likely attributed to Tyr246 since heat of ionization (28.5 kJ/mol) for this \( pK_a \) is much higher than that of a coordinated nucleophilic water in the mononuclear metallopeptidases carboxypeptidase A (6.3) [50], thermolysin (5.1) [51], and serralysin (5.74) [52]. This comparison suggests that this \( pK_a \) may not be due to a coordinated water, but most likely the amino group of the substrate (\( pK_a = 7.94 \) [10]) which is consistent with that deprotonation and binding of the amino group of the substrate controls the activity as suggested in the catalysis by biLAP [53]. Moreover, this \( pK_a \) is much higher than those values of the few mononuclear metallopeptidases above, suggesting that it is not likely to be attributed to a bridging water since the latter is expected to be lower due to binding to two Lewis acidic metal centers.

The pH dependence of \( k_{\text{cat}} \) in BNPP hydrolysis shows two deprotonation constants \( pK_a \) of 7.8 and 10.0 in the range of pH 7 to 10.5, while \( k_{\text{cat}}/K_m \) is controlled by only one \( pK_a \) of 9.5 (Fig. 5, o, right scales; Table 2) which is comparable to one in Leu-pNA hydrolysis by MnMn- and ZnMn-SgAP [10]. A comparison of the pH profiles for the hydrolysis of Leu-pNA and BNPP indicates that the decrease in
6.3 dramatically different, suggesting different mechanisms for the pH profile of Leu-
1-2 substrates. For example, ZnZn- and MnZn-SgAP show similar kcat (101 vs. 92.8 s−1) and KM (3.27 vs. 3.31 mM) values; whereas MnMn-SgAP affords 27.5 s−1 and 0.875 mM, noticeably different from the values of MnZn-SgAP (Table 1). The kcat and KM values of CdCd- and MnCd-SgAP, CoCo- and MnCo-SgAP, and NiNi- and MnNi-SgAP are comparable to each other, whereas those of MnM2-SgAP are significantly different (Table 1). The results support the signficance of M2 in SgAP catalysis.

There is no detectable exchange between the two metal ions in the derivatives M1M2-SgAP during the measurements since the activities of the derivatives remain the same after a day. The kcat and KM values are controlled by the second metal M2 toward the hydrolysis of Leu-pNA, e.g., ZnZn- and MnZn-SgAP show similar kcat (101 vs. 92.8 s−1) and KM (3.27 vs. 3.31 mM) values; whereas MnMn-SgAP affords 27.5 s−1 and 0.875 mM, noticeably different from the values of MnZn-SgAP (Table 1). The kcat and KM values of CdCd- and MnCd-SgAP, CoCo- and MnCo-SgAP, and NiNi- and MnNi-SgAP are comparable to each other, whereas those of MnM2-SgAP are significantly different (Table 1). The results support the significance of M2 in SgAP catalysis.

Comparison of kcat and KM toward the hydrolysis of BNPP by various M1M2-SgAPs is not as clear-cut as in peptide hydrolysis (Table 3). For example, kcat and KM are 0.45 s−1 and 4.5 mM for ZnZn-SgAP, 0.103 s−1 and 3.76 mM for MnZn-SgAP, and 0.21 s−1 and 12 mM for MnMn-SgAP. Similar results are observed for the pairs of Co2+, Cd2+, and Ni2+ derivatives, wherein kcat and KM are significantly affected by both metal ions, as opposed to the case of Leu-pNA hydrolysis where M2 plays a more significant role in catalysis and substrate binding (Table 1). The results suggest that the mechanisms for peptide hydrolysis and BNPP hydrolysis are different, with M2 showing more influence on Leu-pNA hydrolysis than on BNPP hydrolysis.

3.5. Mechanism for peptide hydrolysis

The mechanisms for peptide hydrolysis by ApAP, E. coli Met AP, and bIAP vary from mononuclear to dinuclear peptide hydrolysis; however, a few other APs show that both metal ions are required for TS stabilization and catalysis [19–25,29–35]. On the basis of the results presented here, a dinuclear mechanism is proposed for SgAP (Scheme 1, Fig. 6).
3.5.1. Binding mode of peptide substrate

There are three functional groups in the substrate which can potentially interact with SgAP [19–25,29–35]: the carbonyl of the scissile peptide bond with M1 (and a dipole–charge interaction with M2), the amino terminus with M1, and the hydrophobic side chain with Phe219 (Scheme 1A). To visualize substrate and inhibitor binding to SgAP, the TS/C224 analogue (α-aminoisopentylphosphonate; red ball-and-stick structure) is superimposed onto the active center of SgAP (red ball-and-stick structure; Fig. 6) which reveals that the inhibitor binding modes of these two enzymes are similar, wherein the N-terminal hydrophobic residue is pointing toward Phe219 in SgAP (blue; Fig. 6) and the amino group of the TS/C analogue is within bonding distance to M1. A substrate (Leu–Phe) is superimposed onto the bound inhibitor to reveal its interaction with the hydrophobic site and the metal center. The N-terminal amino group and the nucleophilic water (red sphere) are as labeled.

Scheme 1.

Fig. 6. Relaxed-eye stereo plot of plausible binding modes for a peptide substrate (cyan; Leu–Phe) and a TS/C analogue (α-aminoisopentylphosphonate; red ball-and-stick structure) in the active site of SgAP. The bound inhibitor is obtained from the crystal structure of bILAP which is superimposed onto the dinuclear center of SgAP and show appropriate hydrophobic interaction. Then the substrate is superimposed onto the TS/C inhibitor to reveal its interaction with the hydrophobic site and the metal center. The N-terminal amino group and the nucleophilic water (red sphere) are as labeled.

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There are two possible orientations for the substrate to bind to the active-site metal ions [10]: (a) The scissile carbonyl group is bound to M1 (and M2 via charge–dipole interaction) which mimics the bridging oxygen of the bound α-aminoisopentylphosphonate inhibitor or (b) the carbonyl group is in place of the inhibitor oxygen that is H-bonded to the Tyr246. In the former case, the carbonyl oxygen is bound to M1 at the favorable syn position with the peptide bond facing the plausible nucleophile on M2 (see below),
whereas the substrate is in an unfavorable binding situation in the latter case with the metal situated out of the enzyme’s active site.

3.5.2. Nucleophile in peptide hydrolysis

The substrate binding mode Section 3.5.1 discussed above renders the coordinated water that is H-bonded to Tyr246 that is facing the scissile –CO– group. This water that is H-bonded to Tyr246 and facing the scissile –CO– group is known to perform nucleophilic attack (Scheme 1A). Crystal structure study of this enyme suggests that Tyr246 is involved in H-bonding with a plausible coordinated water at the position of one oxygen of a coordinated dibasic phosphate. The role of Tyr246 as a general acid thus can further enhance the Lewis acidity of the water, resulting in lowering in the pkₐ and increasing in the nucleophilicity of the water for effective nucleophilic attack. The role of a general acid to lower the pkₐ and enhance the nucleophilicity of a coordinated water and a Cys nucleophile, respectively, was also proposed in the case of metallocoulenases [54,55] and glutaredoxin [56]. The crystal structures of several dinuclear hydrolytic enzymes with a metal–metal distance 3.5 Å reveal a bridging hydroxide which has been proposed to be the nucleophile or the precursor of the nucleophile, e.g., ApAP [46], some PAP [57], urease [58], and pyrophosphatase [59]. If this would be the case in SgAP, both metal ions should influence the activity; which is not the case. The uncompetitive inhibition of F⁻ suggests that metal-bond nucleophilic OH⁻ becomes available for replacement by F⁻ only upon substrate binding. The substrate binding mode described herein suggests that the bridging OH⁻ is replaced by the bound substrate. This binding may induce changes in the active site and proper orientation of the scissile peptide bond relative to the attacking nucleophile. The F⁻ inhibition constants are significantly influenced by the second metal ion M2, indicating that the nucleophilic coordinated water is on M2.

Glu131 may further activate the nucleophile and/or stabilize the TS⁺ [49], analogues to Glu270 in carboxypeptidase A which is also involved in deprotonation of the nucleophile and protonation of the amino terminus of the leaving group [60]. The pH profile of F⁻ inhibition reveals a pkₐ at 9.3 which may be attributable to Tyr246, and is consistent with its role in the interaction and proper orientation of the nucleophile as proposed in accordance with crystal structures. It is thus most likely that the “first metal” is near the hydrophobic site (i.e., M1 in Fig. 6) while M2 hosts the nucleophilic water that is H-bonded to Tyr246 and facing the scissile –CO– group. Nucleophilic attack by this water affords the TS⁻ as in the structure of blLAP with a bound TS⁻ analogue (red ball-and-stick structure, Fig. 6 and Scheme 1B).

3.5.3. Stabilization of the TS⁻

The kinetic results suggest that M2 plays a major role in stabilizing TS⁻ and performing nucleophilic attack since kcat/Km varies more with this metal ion. However, the crystal structure of blLAP with a bound TS⁻-like α-aminoalkylphosphate inhibitor [3] shows that both metal ions are involved in TS⁻ binding and stabilization (Fig. 6). Upon nucleophilic attack, the oxygen on the scissile carboxyl group changes from sp² hybridization to sp³ hybridization in the gem-diolate-like intermediate which allows the oxygen to serve as a better bridging ligand between the two metals. The pH dependence of kcat toward the hydrolysis of Leu-pNA by different metal derivatives of SgAP shows a pkₐ value of ~9.2, assignable to Tyr246 (near M2) which is suggested to activate the nucleophilic water and stabilize the gem-diolate-like TS⁻. The results from phosphate inhibition toward the hydrolysis of Leu-pNA by native and phenylglyoxal-modified ZnZn-SgAP and from NMR relaxation studies of Co²⁺ derivatives suggest that Arg202 may also be involved in TS⁻ stabilization [41] (Scheme 1B).

3.5.4. Release of the products

The TS⁻ breaks down (Scheme 1C) to release the C-terminal main protein body, while the N-terminal amino acid presumably remains bound (Scheme 1D) as suggested by crystallographic studies of various amino acid–bound SgAP [61]. The N-terminal amino group is supposed to detach from the metal after the substrate is hydrolyzed owing to increase in its pkₐ after bond cleavage (Scheme 1C). Crystal structures of SgAP with a bound amino acid (Met, Leu, or Phe) show that one carboxyl oxygen of the bound amino acid is coordinated to M2 and H-bonded to Tyr246 while the other oxygen binds to both metal ions [61]. Moreover, the amino group of the bound amino acid product is detached from M1, and is presumably H-bonded to Glu131 (Scheme 1C and D). The pH profile for Leu inhibition toward Leu-pNA hydrolysis shows a weak inhibition and no clear pkₐ value at pH 6.0 to 10.0 [10]. The release of the amino acid product can thus spontaneously occur without the assistance of a general acid.

3.6. Catalytic promiscuity

Dinuclear hydrolyses such as AAs have been investigated by numerous methods to reveal the reaction mechanisms [6,45,62,63], e.g., the role of metal ions in activity [19,31,64], and stabilization of the tetrahedral TS⁺, and binding of TS⁻-inhibitors [3,65]. Although a phospho-center has been known to serve as a TS⁻ analogue of peptide substrates, the phosphodiesther BNPP and the phosphonate ester P-nitrophenylphosphonate can be effectively hydrolyzed by SgAP and its metal derivatives with a catalytic proficiency of ~40 billion-fold for the native enzyme and 67 billion-fold for the di-CO²⁻ derivative toward BNPP hydrolysis under physiological conditions [9,10]. Moreover, Cu²⁺-SgAP exhibits a significant catechol oxidase activity, showing ~10% catalytic efficiency compared to a native catechol oxidase [40]. Revealing the key factors involved in these alternative catalyses are essential to gain further insight into dinuclear catalysis as well as “enzyme catalytic promiscuity [66–70].”

The catalytic efficiencies in terms of the first-order rate constants [71,72] (i.e., kcat/Km with k₁ the uncatalyzed rate constant) for Leu-pNA hydrolysis by the several metal-substituted derivatives of SgAP at pH 8.0 are quite significant, ranging from 0.44 × 10⁶ for the di-Ni derivative to 6.7 × 10⁴ for the native enzyme with respect to k₁ of 9.8 × 10⁻⁸ s⁻¹ determined under the same conditions (Table 1), but are much smaller for the less specific peptide substrates [10]. The kcat/Km value for Leu-pNA hydrolysis by the native enzyme represents an enormous G² value of ~57 kJ/mol at 303 K. These values of catalytic proficiency for peptide hydrolysis serve as the criteria for the evaluation of the promiscuous catalysis toward BNPP hydrolysis.

Phosphoesters resembles the gem-diolate-like TS⁻ of peptides during hydrolysis, which explains why BNPP can serve as an inhibitor toward peptide hydrolysis by SgAP and ApAP [9,10]. Moreover, BNPP and a few phosphoesters and fluorophosphates are also known to be TS⁻-inhibitors against serine proteases and esterases such as trypsin, chymotrypsin, and acetylcholine esterase through covalent linkage with the nucleophilic serine in the active site [7,11–13] to afford Ser–O–P(O)₂⁻–OR with concomitant cleavage of a phosphoester or the P–F bond that resembles the “initial burst” kinetics of serine proteases. However, the phosphoester Ser–O–P(O)₂⁻–OR is indefinitely stable, thus has only one turn-over for the phosphoester bond cleavage by these enzymes. On the other hand, sulfamate bond cleavage by subtilisin is a novel observation of catalytic promiscuity in the serine protease family [73]. Likewise, the cleavage of BNPP by SgAP and its metal derivatives is also catalytic with enormous catalytic efficiencies relative to the autohydrolytic rate constant [74–76] (Table 3). The second-order rate constants of BNPP hydrolysis by SgAP and its
metal derivatives (100 and 0.5–78 M$^{-1}$ s$^{-1}$, respectively) are considerably higher than those of many synthetic chemical model systems of Zn$^{2+}$ complexes (e.g., in the range of $k$2 $\sim$ 10$^{-6}$ to $10^{-4}$ M$^{-1}$ s$^{-1}$) and other chemical systems performed at $>$35 °C and pH 8.5 [77–85]. Thus, the high activity of SgAP cannot simply be attributed to the high Lewis acidity of the metal ions as in the chemical model systems. The activities of SgAP and its metal derivatives toward BNPP hydrolysis are significantly higher than or comparable to those of some native phosphoester-hydrolyzing enzymes and their derivatives, such as alkaline phosphatase (0.05 M$^{-1}$ s$^{-1}$ [86]) and Burkholderia phosphonate monoester hydrolase (11.4 M$^{-1}$s$^{-1}$ [87]) and ethyl(p-nitrophenyl)phosphate hydrolysis by metal derivatives of Pseudomonas phosphotriesterase (1.1–7.2 M$^{-1}$ s$^{-1}$ [88]). The specific activities toward the hydrolysis of 1.0 mM BNPP by SgAP and its derivatives (0.87–154 nmol min$^{-1}$ mg$^{-1}$ derived from Table 3) are also in the range of those of several phosphodiesterases and phosphatases (0.3–2450 and $\sim$2–40 nmol min$^{-1}$ mg$^{-1}$, respectively) [89,90]. However, native phosphodiester-specific enzymes can have much higher activities, such as the phosphodiesterase gene product of Eluc shows $k_{cat}$ of 59 s$^{-1}$ and $k_{cat}$/$K_{m}$ = 1480 M$^{-1}$s$^{-1}$ [91]. Nevertheless, this comparison concludes that the phosphodiesterase activity of SgAP and derivatives toward BNPP hydrolysis are indeed quite significant.

3.7. Mechanism for BNPP hydrolysis

The hydrolysis of phosphoesters has different mechanistic requirements from peptide hydrolysis, e.g., a tetrahedral gem-diole-like TS$^i$ for the former and a trigonal bipyrindal TS$^i$ for the latter. One the basis of the different (a) influences by M1 and M2, (b) inhibition patterns, (c) pH–activity profiles, and (d) pKa values between BNPP and peptide hydrolysis by SgAP, we propose a mechanism for the proficient BNPP hydrolysis below (Scheme 2), wherein BNPP is bound to the active site analogous to the TS$^i$ of peptide during hydrolysis.

3.7.1. Binding mode of BNPP

The variation in the $k_{cat}$ and $K_{m}$ values among SgAP derivatives on BNPP hydrolysis (Table 3) indicates the hydrolytic reaction is influenced by both M1 and M2. M2 plays a significant role in peptide hydrolysis, but is shown herein to have a different role in BNPP hydrolysis. The binding mode of BNPP is proposed to be similar to the TS$^i$ in peptide hydrolysis (cyan; Fig. 7) where two oxygens of phosphate bind to M2 with one bridging to M1 (cf. red ball-and-stick inhibitor in Fig. 6; Scheme 2A). Phosphate inhibits BNPP and Leu-pNA hydrolysis in competitive and noncompetitive patterns, respectively, it competes with BNPP binding to SgAP, but not with Leu-pNA. The combination of the pH profiles, kinetic, and $^{31}$P NMR relaxation studies suggests that phosphate binds to Arg202 [16,34,41]. This Arg is supposed to bind the TS$^i$-like BNPP (Scheme 2A). In addition, the similar pKa values for BNPP and Leu-pNA hydrolysis at the alkaline side indicates the involvement of the same amino acid for catalysis and/or TS$^i$ stabilization, suggested to be Tyr246.

3.7.2. Generation of nucleophile

The different F$^\text{–}$ inhibition patterns toward BNPP (no inhibition) and Leu-pNA (uncompetitive) hydrolysis reflect different mechanisms for the generation of the nucleophile in these two hydrolytic reactions. In addition, the steric hindrance imposed by the active site on the leaving group and the requirement of an in-line attack on phosphoester substrates [92,93] by the nucleophile during hydrolysis suggest that M1 may possess the nucleophile (red HO in Fig. 7; Scheme 2A) and the leaving group is near the open-end of the active site (i.e., the group on the left-hand side of the substrate, Fig. 7). The nucleophile in the peptide hydrolysis is integrated into the substrate at the TS$^i$, thus is not available for the hydrolysis of the TS$^i$-like BNPP, which as a consequence must have a different nucleophile.

3.7.3. Stabilization of the TS$^i$

The pH dependence of $k_{cat}$ is controlled by a pKa value of 9.2, suggested to be due to Tyr246. The $k_{cat}$ and $K_{m}$ values of BNPP hydrolysis by SgAP are dependent on both metal ions, indicating both metal ions are involved in BNPP hydrolysis. Since phosphate and fluoride inhibitions indicate that BNPP binds to SgAP as a TS$^i$ analogue of a peptide substrate (red inhibitor in Fig. 6), and the nucleophile is expected to be generated by M1 (OH in Fig. 7) rather than M2 in peptide hydrolysis since the nucleophile in pep-
tide hydrolysis is now part of the TS\(^1\). This nucleophile is able to perform SN2 in-line attack \([92,93]\) in BNPP hydrolysis from the opposite site of the leaving group to form a trigonal bipyramidal TS\(^1\) (pink; Fig. 7). Herein, Arg202 was suggested to stabilize the TS\(^1\) (whereas a counterpart is not present in AP\(^\text{AP}\)), along with Tyr246, Glu131, and both metal ions (Scheme 2B).

3.7.4. The release of the products
The phosphomonoester product has been shown to be a good inhibitor toward BNPP hydrolysis (\(K_i = 0.9 \text{ mM}\)) \([10]\). Thus, upon detachment of the leaving group \(p\)-nitrophenol (Scheme 2C), an enzyme–product complex is formed which adopts a configuration analogous to the bound phosphate in the enzyme \([16]\) with one O originated from the nucleophile bound to M1, one bridging O, and the other one H-bonded with Tyr246 and may bind to M2 (Scheme 2D). Release of the phosphomonoester and placement of coordination sphere with water molecules complete the catalytic cycle.

4. Concluding remarks
Although enzymes are frequently ascribed to possess catalytic specificity, most enzymes are known to catalyze the conversion of a family of substrates and analogues with similar structures, such as the hydrolysis of peptides and carboxylesters by pep-tidases. Several enzymes have also been demonstrated to perform catalyses toward substrates of different families, exhibiting enzyme catalytic promiscuity \([66–70]\). However, it would be less expected for an enzyme to exhibit an enormous alternative catalysis that is normally carried out by an evolutionarily and structurally unrelated enzyme. We describe in this report the hydrolysis of a phosphodiester substrate by SgAP and its metal derivatives, reaching a dramatic 67-billion-fold rate enhancement relative to the non-catalyzed reaction at neutral pH and room temperature and having rate constants comparable to several native phosphoester-hydrolyzing enzymes.

Dinuclear centers of various metal ions are found in many phosphodiesterases, including urease (Ni), nucleases (Zn), arginase (Mn), prolidase (Mn), and AP (Zn or Co) \([6]\), as well as in a number of Fe-, Mn-, and Cu-containing oxidases and oxygenases \([94]\). The catalytic versatility of dinuclear SgAP and its metal derivatives entitle this enzyme to serve as a "natural dinuclear model system\([9]\" for further investigation of various dinuclear catalyses. Indeed, the di-Cu derivative of SgAP has recently been demonstrated to exhibit a significant activity toward the oxidation of catechol and derivatives \([40]\), despite its very different active-site structure from that of catechol oxidase. Future studies of this unique natural dinuclear model system by means of physical methods and molecular biology are expected to provide further information to gain better understanding of various dinuclear catalyses in chemical and biological systems and further knowledge about how to use this and other metalloenzymes as scaffolds to build artificial metalloenzymes of novel catalytic activities.

5. Abbreviations
- AP: aminopeptidase
- ApAP: Aeromonas proteolytica AP
- blLAP: Leu-specific AP from bovine lens
- BNPP: bis(\(p\)-nitrophenyl)phosphate
- DEAE: diethylaminoethyl
- HEPES: N-[2-hydroxyethyl] piperazine-NN-2-ethanesulfonic acid
- MES: 2-[N-morpholino]ethanesulfonic acid
- PAP: purple acid phosphatase
- pkLAP: Leu-specific AP from porcine kidney
- pNA: \(p\)-nitroaniline
- SgAP: Streptomyces griseus aminopeptidase
- TS\(^1\): transition-state

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