Tuned by metals: the TET peptidase activity is controlled by 3 metal binding sites

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TET aminopeptidases are dodecameric particles shared in the three life domains involved in various biological processes, from carbon source provider in archaea to eye-pressure regulation in humans. Each subunit contains a dinuclear metal site (M1 and M2) responsible for the enzyme catalytic activity. However, the role of each metal ion is still uncharacterized. Noteworthy, while mesophilic TETs are activated by Mn$^{2+}$, hyperthermophilic TETs prefers Co$^{2+}$. Here, by means of anomalous x-ray crystallography and enzyme kinetics measurements of the TET3 aminopeptidase from the hyperthermophilic organism *Pyrococcus furiosus* (PfTET3), we show that M2 hosts the catalytic activity of the enzyme, while M1 stabilizes the TET3 quaternary structure and controls the active site flexibility in a temperature dependent manner. A new third metal site (M3) was found in the substrate binding pocket, modulating the PfTET3 substrate preferences. These data show that TET activity is tuned by the molecular interplay among three metal sites.

Protein degradation occurs in the three life domains and it is involved in a wide variety of cellular processes such as protein quality control, amino acids pool renewal and as a carbon source provider$.^{1,2}$ Cytosolic protein degradation is carried out by different classes of proteases that are substrate specific or that are self-compartmentalized to shield the active site to avoid unspecific proteolysis$.^{3}$ Among self-compartmentalized proteases, a new class of dinuclear metalloaminopeptidases complex was recently discovered and named TET due to its tetrahedral shape$.^{1}$ TETs are dodecameric particles of 500kDa and they were first discovered in archaea and then identified in bacteria and eukaryotes$.^{3-7}$ To date, three types of archaeal TETs have been identified and characterized based on their substrate specificity. TET1 is a glutamyl-aminopeptidase, TET2 has a broad specificity against neutral amino acids and TET3 shows a preference for positively charged residues such as lysine and arginine$.^{3,8-10}$ The physiological role of these dodecameric particles is not completely understood, although it has been found that TET homologues, DNPEP and PmM18AAP, are involved in blood pressure regulation in humans$.^{5,11,12}$ and participates in the hemoglobin degradation in *P. falciparum*$.^{13}$, respectively.

TETs dodecamers are characterized by a tetrahedral shape showing four openings on the facets. Each apex of the tetrahedron is formed by three subunits arranged along a three-fold axis. Each subunit has an active site that hosts two metal ions (M1 and M2) forming a dinuclear metal center. The classification of each metal site was derived from the nomenclature of Schechter and Berger*.^{14}$ M1 lays in the mouth of the PfTET3 active site, while M2 is adjacent to the specificity binding pocket (P1) that hosts the P1 side chain. P1 electrostatic potential is adapted to welcome positively (TET3), negatively (TET1) or neutral (TET2) charged residues$.^{3,8,10}$ TETs monomers are all formed by a proteolytic domain and a PDZ-like oligomerization domain. Each subunit interacts through two large interfaces. The dimerization interface is mediated by the PDZ-like domains that form the dimeric building block. The oligomerization interface is supported by both the proteolytic domain and the PDZ-like domain, driving the association of the dimeric building blocks into the dodecameric assembly$.^{15,16}$ It has been recently shown that TET dimers are present in vivo* and that their activity against long peptides is considerably lower compared to the dodecameric particle*, suggesting a fine regulation of TET activity by controlling its oligomerization.

A key factor in TET oligomerization is the dinuclear metal site. In particular, when TET particle is treated with a high concentration of chelating agent (EDTA) associated with basic pH (pH 10) at room temperature, the
dodecamer dissociates into dimers\textsuperscript{18}. Indeed, the dinuclear metal site is close to the oligomeric interface and the depletion of metals may destabilize the H-bonds interactions between two adjacent subunits resulting in the deoligomerization of the TET particle\textsuperscript{18}. The dinuclear metal center has not only a structural role in the TET catalytic activity, but it is also responsible for the TET catalytic activity. To date, there is no characterization of the role played by each metal in the active site of TET, although x-ray crystallography and extended x-ray absorption fine structure (EXAFS) studies have shown that the dinuclear metal center of TET can be occupied by Zn\textsuperscript{2+}, Co\textsuperscript{2+} and Mn\textsuperscript{2+} ions which catalyse the aminopeptidase activity with different yields\textsuperscript{38-10}. Interestingly, while the three thermophilic archaean TETs require Co\textsuperscript{2+} to reach the optimal activity, their mesophilic counterparts Streptococcus pneumoniae PepA (SpPepA), DNPEP and PIM18AAP are activated by Co\textsuperscript{2+} and/or Mn\textsuperscript{2+}. The molecular basis of Co\textsuperscript{2+} and/or Mn\textsuperscript{2+} activation has not been elucidated yet.

In this study, to get insights on the metal specificity of TET and in particular to address the specific role of each metal in the active site of the enzyme, the TET3 aminopeptidase from the hyperthermophilic organism Pyrococcus furiosus (PfTET3) was characterized by means of x-ray crystallography and UV-spectroscopy in the presence of Zn\textsuperscript{2+}, Co\textsuperscript{2+} and Mn\textsuperscript{2+}. In particular, crystallogenesis with controlled Co\textsuperscript{2+} concentration and subsequent characterization by x-ray anomalous experiment and enzymatic assays revealed that Co\textsuperscript{2+} replaces the M1 site and occupies a new third metal site (M3) in the specificity binding pocket. Importantly, the M2 site was not replaced by Co\textsuperscript{2+}. Moreover, enzymatic assays performed on a wide range of temperature (20°C–85°C) have allowed determining the role of each metal ion. These results highlight a complex metal interplay at the base of the TET3 enzyme activity, establishing that the M1 site controls the flexibility of the active site, M2 hosts the activated water molecule and M3 modulates the substrate specificity for the enzyme. Finally, the molecular basis of Co\textsuperscript{2+} activation is depicted, showing that Co\textsuperscript{2+} increases the activity of PfTET3, by stabilizing the active site, only at high temperature.

Results
X-ray crystal structure of PfTET3 at 2.5 Å resolution reveals that M2 is the catalytic metal. Previously, PfTET3 was assigned to M18 family and it was reported as a homotetramer based on gel filtration elution profile\textsuperscript{10}. However, PfTET3 displays 90% of sequence identity with TET3 from Pyrococcus horikoshii (PhTET3), whose x-ray crystal structure was determined, showing a dodecameric assembly\textsuperscript{10}. To unambiguously characterize PfTET3, we determined its crystal structure by SAD at 2.5 Å resolution. Purified PfTET3 was crystallized in HEPES 0.1M pH 7.7, NH\textsubscript{4}CH\textsubscript{3}COO 0.2 M, 2 mM CoCl\textsubscript{2}, MPD 44% and Gd-DO\textsubscript{3}A\textsuperscript{20} using hanging drop vapor diffusion method. Crystals appeared after 1 month and X-ray data collection was performed at ID29 beamline at the European Synchrotron Radiation Facility (ESRF-Grenoble). PfTET3\textsubscript{Gd} crystal was collected at Gd LIII absorption edge. A strong anomalous signal relative to the Gd atoms extends up to 3.0 Å, allowing determining the Gd substructures. These latter were used to solve the phase problem relative to PfTET3. Data collection and refinement statistics are reported in Table 1.

The asymmetric unit contains three monomers forming one apex of the PfTET3 tetrahedron (Fig. 1A). By applying the crystal symmetry operators, PfTET3 dodecamer is reconstituted (Fig. 1B). Each monomer is formed by a proteolytic domain and a PDZ-like domain, known as TET dimerisation domain. Residues 120–128 are disordered and they were not modeled. The proteolytic domain comprises a central \( \beta \)-sheet surrounded by seven \( \alpha \)-helices and accommodates the catalytic pocket. This latter contains two strong positive peaks of electron density that corresponds to the two metal ion sites. Each metal site was classified based on the nomenclature of Schechter and Berger\textsuperscript{41}. M1 is coordinated by residues His314, Asp177, Glu208 and two water molecules, while M2 is coordinated by Asp254, His63 and a chloride ion (Fig. 1C). In P1 pocket, Asp254 establishes H-bond with water molecules. The purified PfTET3 contains two \( \text{Zn}^{2+} \) ions in the active site. However, since the PfTET3 crystal was present in the crystallization conditions, it can be inferred that a metal exchange occurred. Indeed, M1 displays an octahedral coordination, while M2 displays a trigonal bypyramidal coordination. By looking to the preferred coordination number of \text{Co}^{2+} and \text{Zn}^{2+}, it appears that \text{Co}^{2+} adopts preferentially an octahedral geometry (coordination number of 6), while \text{Zn}^{2+} is often found in a tetrahedral coordination (coordination number 4)\textsuperscript{42}. These data suggest that M1 is occupied by \text{Co}^{2+} while M2 is occupied by \text{Zn}^{2+}. The two metals are separated by 3.3 Å in the three monomers present in the asymmetric unit (Fig. 1D). By comparison, in the PhTET3 structure the two Zn ions are separated by 3.4 Å (PDB 2WZN) while in SpPepA crystallographic structure, 12 monomers were found in the asymmetric unit (PDB 3KL9) and the distance between the two Zn ions varies between 3.1 Å – 3.5 Å, suggesting a dynamic positioning of the metal ions in the active site.

In the PfTET3 structure, M2 encompasses a Cl\textsuperscript{−} ion in his coordination sphere (2.8 Å). Indeed, it was reported that a carboxylate rich environment favors a hard acid behavior of the two metal ions in the active site resulting in the binding of halides in the order F\textsuperscript{−} > Cl\textsuperscript{−} > Br\textsuperscript{−} > I\textsuperscript{−}\textsuperscript{22}. The catalytic mechanism of aminopeptidase involves a hydroxide ion, which is formed by the polarization of a water molecule bound to the two metal ions. Indeed, the Nb and Ne of His63 are involved in the weak H-bonds network with the O\text{6} of Asp65 (3.1 Å) and the O\text{5} of Glu207 (3.1 Å), respectively. Glu207 is the general base responsible for the deprotonation of the water molecule. It has been proposed that such an environment results in a decreasing of the acidity of the M2 metal\textsuperscript{22} allowing the polarization of the nucleophilic water molecule. It was previously reported that F\textsuperscript{−} ions act as a noncompetitive inhibitor by displacing the hydroxide ion from the metal ion\textsuperscript{24}. Thus, the Cl\textsuperscript{−} ion can mimic the position of the hydroxide. In PfTET3\textsubscript{Gd} structure, Cl\textsuperscript{−} ion is in the coordination sphere of M2, indicating that M2 is the catalytic metal.

Interestingly, M1 is coordinated by Glu208, which lays on a loop involved in PfTET3 oligomerization interface. In particular, Gln206 and Arg212 from one subunit, form H-bonds with the carbonyl carbon of Pro257 and Asn292 and Gln295 side chains of the adjacent subunits. This suggests that M1 may have a stabilizing role of PfTET3 oligomerization interface.
where n represents the 

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Furthermore, below Co2+ absorption edge, sulphur displays significant anomalous signal (f' for S is 0.62 e−). Met313 is close to M1 site and it can be used as a probe to compare the anomalous signal. Indeed, ANODE found two small peaks corresponding to M2 site and Met313 in the active site (Fig. 2D and Table 2). These data show that Co2+ replaces the M1 site and occupies an additional M3 site, while M2 retains the Zn2+ ion recovered from the cell culture medium.

### Table 1. Data collection and refinement statistics for PfTET3Gd, PfTET3Zn, PfTET3Co, PfTET340eV

| Structure | PfTET3Gd peak | PfTET3Zn peak | PfTET3Co peak | PfTET340eV peak |
|-----------|---------------|---------------|---------------|-----------------|
| Beam line | ID29 (ESRF)   | Proxima II (SOLEIL) | Proxima II (SOLEIL) | Proxima II (SOLEIL) |
| Space Group | Tetragonal I4_122 | Tetragonal I4_122 | Tetragonal I4_122 | Tetragonal I4_122 |
| Unit Cell Constants (Å) | a = 203.91 b = 203.91 c = 112.44 | a = 201.7 b = 201.7 c = 113.2 | a = 201.7 b = 201.7 c = 113.4 | a = 202.6 b = 202.6 c = 113.6 |
| Resolution (Å) | 48.06 – 2.50 (2.60 – 2.50) | 49.37 – 3.35 (3.62 – 3.35) | 49.42 – 3.25 (3.56 – 3.25) | 49.53 – 3.30 (3.47 – 3.30) |
| Wavelength (Å) | 1.71081 | 1.604970 | 1.282290 | 1.614380 |
| Rmerge (%) | 19.2 (124.8) | 7.2 (77.8) | 5.7 (72.5) | 8.7 (122.9) |
| Rfree (%) | 4 (42.2) | 2.4 (25.5) | 2.3 (24.8) | 2.9 (40.3) |
| Completeness (%) | 96.0 (77.4) | 99.7 (99.1) | 99.8 (99.7) | 99.8 (99.4) |
| Unique Reflections | 39432 (3500) | 17047 (3429) | 18701 (4389) | 18113 (3664) |
| Anomalous multiplicity | 11.5 (4.6) | 5.3 (5.2) | 5.3 (5.2) | 5.2 (5.2) |
| FOM | 0.73 |

**R** workf (%) = 17.8

M1 site is replaced by Co2+ in the PfTET3 active site. In the previous paragraph, PfTET3 de novo x-ray structure suggested that M2 is the catalytic metal. The three TETs from *Pyrococcus horikoshii* display enhanced enzymatic activity in the presence of Co2+, implying that M1 and/or M2 are exchanged. To determine which metal site is exchanged by Co2+, anomalous x-ray scattering experiments were performed on PfTET3. Crystals of PfTET3 were grown at 20 °C in the presence of 2 Co2+ equivalents per monomer. For the sake of the clarity, hereafter, PfTET3 enzyme treated with Co2+ is reported as PfTET3Co, while the recombinant PfTET3 is named PfTET3Zn. PfTET3Co crystals appear after 7–10 days, they were flash-frozen and x-ray data were collected on the same crystal at two different wavelengths corresponding to the Co2+ and Zn2+ absorption edges, respectively. The anomalous x-ray data for both Co2+ and Zn2+ were processed using XDS/Aimless/SHELXC/ANODE softwares. The positions of the anomalous scatterers (Co2+ and Zn2+) were determined using ANODE. ANODE calculates an anomalous difference Fourier by applying a 90° phase shift to the protein phase that is calculated from a PDB model. We used the de novo PfTET3Gd structure reported in this paper as a PDB model. At Zn2+ absorption edge, ANODE identifies four main peaks. Three peaks correspond to the M2 site of each monomer (A, B, C) and one corresponding to the M1 site in monomer C. Although in monomer A and B ANODE did not identify a clear peak for M1 site, residual electron density protruding from the M2 site is observed (Fig. 2B and Table 2). Indeed, at Zn K absorption edge, f' of Zn2+ is 3.9 e− while f' of Co2+ is 2.4 e−. Conversely, at Co absorption edge, ANODE identifies three main peaks corresponding to M1 site for A, B, C, monomers. Interestingly, other three smaller peaks were also identified by ANODE in each monomer in the asymmetric unit, corresponding to a third metal site M3, coordinated by residues Glu281, Asp254 and Thr232 (Fig. 2C and Table 2). These data show that Co2+ replaces the M1 site and occupies an additional M3 site, while M2 retains the Zn2+ ion recovered from the cell culture medium.
M1 site in PfTET3 active site is implicated in the oligomeric interface stabilization. In the presence of Co\(^{2+}\), the PfTET3\(\text{Zn}\) M1 site is exchanged by Co\(^{2+}\), while M2 remains occupied by Zn\(^{2+}\). However, the respective role(s) of M1-M2 in the active site of PfTET3\(\text{Zn}\) remain to be determined. It was previously reported for Aeromonas proteolytica aminopeptidase (AAP) and methionine aminopeptidase (MetAP)\(^{23,26}\) that 50–80% of the enzymatic activity can be obtained with only one metal in the active site. To evaluate the role of M1 and M2 in the PfTET3\(\text{Zn}\) dinuclear metal site, an EDTA titration was performed at 85 °C on the PfTET3\(\text{Zn}\) enzymatic activity using 5 mM Lys-pNa as substrate (Fig. 3A). The EDTA concentration used spans from 5 nM to 20 mM. Interestingly, the data points are fitted with a biphasic dose response curve (correlation coefficient \(= 0.97\)), indicating the presence of two distinct metal binding sites displaying different affinities. The first inflection point corresponds to an EC\(_{50}\) = 13 \(\mu\)M, while EC\(_{50}\) is 3.8 mM. After the first inflection point, PfTET3\(\text{Zn}\) activity is still at 75%, while after the second inflection point it falls down to 30%. Based on the crystallographic studies reported above and the EDTA titration, the M1 site is the lower affinity site exchanged by Co\(^{2+}\), while the M2 is the high affinity site responsible for the catalytic activity of PfTET3\(\text{Zn}\).

To evaluate the effect of the depletion of each metal ion on the PfTET3\(\text{Zn}\) quaternary structure at its physiological temperature, 1 \(\mu\)M PfTET3\(\text{Zn}\) was heated at 85 °C for 5 minutes in the absence of EDTA, in the presence of 15 \(\mu\)M EDTA (to selectively remove the M1 site) and in the presence of 6 mM EDTA (to remove both M1 and M2 sites), respectively. The samples were then loaded on a gel filtration column (Superose 6) and the UV profiles were analysed (Fig. 3B). PfTET3\(\text{Zn}\), heated in the absence of EDTA is eluted as a dodecamer. The sample treated with 15 \(\mu\)M EDTA shows two peaks, corresponding to the dodecameric PfTET3\(\text{Zn}\) and to the monomeric form of PfTET3\(\text{Zn}\) based on column calibration. The dodecamer represents 60% of PfTET3\(\text{Zn}\), while the monomer accounts for the remaining 40%. The sample treated with 6 mM EDTA shows a major peak, corresponding to the monomeric PfTET3\(\text{Zn}\). These results show that the removal of the M1 site is sufficient to destabilize the quaternary structure of PfTET3\(\text{Zn}\) that partially dissociates into monomers. The proximity of the M1 site to the oligomeric interface of PfTET3\(\text{Zn}\) strongly suggests that the M1 site modulates the molecular interactions between adjacent subunits.

Co\(^{2+}\) enhances PfTET3 activity only at Pyrococcus physiological temperature. It has been shown that mesopholic TETs (DNPEP, SpPepA, PfM18) are activated by Co\(^{2+}\) and/or Mn\(^{2+}\), while Zn\(^{2+}\) being...
Interestingly, while mesophilic TETs display enhanced activity in the presence of Co\(^{2+}\)/Mn\(^{2+}\), hyperthermophilic TETs were activated only by Co\(^{2+}\). This observation prompted us to study the effect of temperature over the metal preference of PfTET3. PfTET3\(_{Zn}\) enzymatic activity was monitored by following the release of pNa from Lys-pNa substrate in the presence of two different concentrations (0.1 mM and 1 mM) of Co\(^{2+}\), Mn\(^{2+}\) and Zn\(^{2+}\) and at three different temperatures (\(T = 20\ °C, 50\ °C, 85\ °C\)). At 20\ °C and 50\ °C, the addition of each type of metal ions inhibited the enzymatic activity of PfTET3\(_{Zn}\). However, Mn\(^{2+}\) resulted the less inhibitory metal at \(T < 50\ °C\) (Table 3). At 85\ °C, PfTET3\(_{Zn}\) activity is enhanced by Co\(^{2+}\) addition while being strongly inhibited by Mn\(^{2+}\) and Zn\(^{2+}\). These results indicate a specific effect of Co\(^{2+}\) on PfTET3 enzyme; activator at physiological temperature and inhibitor at \(T < 50\ °C\). Furthermore, Table 3 shows that at 85\ °C and 1 mM Co\(^{2+}\), PfTET3\(_{Zn}\) activity is less enhanced, revealing that Co\(^{2+}\) concentration is also important to regulate the enzyme activity. These results prompted us to perform a Co\(^{2+}\) titration of PfTET3\(_{Zn}\) activity at 20\ °C, 50\ °C and 85\ °C. The results

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**Table 2. Height (\(\sigma\)) of the anomalous peaks determined by ANODE at Zn\(^{2+}\)-edge, Co\(^{2+}\)-edge and 40 eV below Co\(^{2+}\)-edge for monomers A, B and C in ASU respectively.**

| Absorption edge | M1 site | M2 site | M3 site |
|-----------------|---------|---------|---------|
| Zn K-edge      | -/-, 13\(\sigma\) | 21\(\sigma\), 15.4\(\sigma\), 14.2\(\sigma\) | -/-, -/- |
| Co K-edge      | 13.5\(\sigma\), 12.2\(\sigma\), 10.4\(\sigma\) | -/-, -/- | 9.4\(\sigma\), 5.9\(\sigma\), 5.5\(\sigma\) |
| Pre-Co-edge    | -/-, 5\(\sigma\), 4\(\sigma\) | -/-, -/- | -/-, -/- |

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Inhibitory\(^{4,5,9,27}\). Interestingly, while mesophilic TETs display enhanced activity in the presence of Co\(^{2+}\)/Mn\(^{2+}\), hyperthermophilic TETs were activated only by Co\(^{2+}\). This observation prompted us to study the effect of temperature over the metal preference of PfTET3. PfTET3\(_{Zn}\) enzymatic activity was monitored by following the release of pNa from Lys-pNa substrate in the presence of two different concentrations (0.1 mM and 1 mM) of Co\(^{2+}\), Mn\(^{2+}\) and Zn\(^{2+}\) and at three different temperatures (\(T = 20\ °C, 50\ °C, 85\ °C\)). At 20\ °C and 50\ °C, the addition of each type of metal ions inhibited the enzymatic activity of PfTET3\(_{Zn}\). However, Mn\(^{2+}\) resulted the less inhibitory metal at \(T < 50\ °C\) (Table 3). At 85\ °C, PfTET3\(_{Zn}\) activity is enhanced by Co\(^{2+}\) addition while being strongly inhibited by Mn\(^{2+}\) and Zn\(^{2+}\). These results indicate a specific effect of Co\(^{2+}\) on PfTET3 enzyme; activator at physiological temperature and inhibitor at \(T < 50\ °C\). Furthermore, Table 3 shows that at 85\ °C and 1 mM Co\(^{2+}\), PfTET3\(_{Zn}\) activity is less enhanced, revealing that Co\(^{2+}\) concentration is also important to regulate the enzyme activity. These results prompted us to perform a Co\(^{2+}\) titration of PfTET3\(_{Zn}\) activity at 20\ °C, 50\ °C and 85\ °C. The results
are shown in Fig. 3C. The plots confirmed the activator role of Co$^{2+}$ only at 85 °C, while at 20 °C and 50 °C Co$^{2+}$ inhibits PfTET3 enzymatic activity. The data points at 85 °C describe a growing sigmoidal curve up to 230% of PfTET3Zn activity, while representing a decreasing curve from 230% to 150% of the PfTET3Zn activity. The data points at 85 °C were fitted with two dose-response functions, one from 0% up to 230% of PfTET3Zn activity and the second from 230% to 150% of PfTET3Zn activity. These two dose-response functions allowed determining two EC50 values, EC501 of 7.4 μM and EC502 of 550 μM. EC501 represents the Co$^{2+}$ concentration value at which half of the maximal enzyme activity is reached. Interestingly EC501 value of the Co$^{2+}$ titration (7.4 μM) is similar to the EC501 found in the EDTA titration (13 μM). Based on the structural studies reported above as well as on the EC501 values of the Co$^{2+}$/EDTA titration, the Co$^{2+}$ mediated enhancement of the PfTET3 activity is dependent from the M1 site. Furthermore, it suggests that the maximum activity of PfTET3 is reached by a hybrid dinuclear metal site, with M1 and M2 filled with Co$^{2+}$ and Zn$^{2+}$, respectively.

Table 3. PfTET3 activity at different metal-temperature using 5 mM Lys-pNa.

| Metal   | % Relative activity T = 20 °C | % Relative activity T = 50 °C | % Relative activity T = 85 °C |
|---------|-------------------------------|-------------------------------|-------------------------------|
| none    | 100                           | 100                           | 100                           |
| Zn$^{2+}$ 0.1mM | 55.5                         | 61.6                         | 52.3                          |
| Co$^{2+}$ 0.1mM | 50.3                         | 67.0                         | 229.4                         |
| Mn$^{2+}$ 0.1mM | 82.8                         | 75.1                         | 38.0                          |
| Zn$^{2+}$ 1mM  | 13.5                         | 15.5                         | 35.4                          |
| Co$^{2+}$ 1mM  | 14.3                         | 35.6                         | 107.2                         |
| Mn$^{2+}$ 1mM  | 47.5                         | 27.2                         | 15.9                          |

Co$^{2+}$ at the M1 site modulates the PfTET3 active site flexibility. To shed light on the role played by Co$^{2+}$, the kinetic parameters of PfTET3 were measured at 85 °C in the presence/absence of 300 μM Co$^{2+}$ and using Lys-pNA as substrate. Results are reported in Table 4. Interestingly, in the presence of Co$^{2+}$, there is a strong reduction of Km compared to the recombinant PfTET3Zn (0.6 mM vs 7 mM). Conversely, kcat is similar with or
To quantify the thermodynamic contribution of Co²⁺ on PfTET³ activity, the kcat/Km ratio of each PfTET³ variant were inserted in the Haldane equation, that allow to determine the difference in transition state free energies (ΔΔGES) between PfTET³Co/PfTET³Zn. ΔΔGES resulted of −6.7 kJ/mol. These results demonstrate that Co²⁺ allows the stabilization of the transition state at the physiological temperature of the PfTET³ enzyme.

To get more insights on this phenomena, the kinetic parameters of PfTET³Zn were measured at five additional temperatures (20 °C, 27.5 °C, 35 °C, 50 °C, 67.5 °C) using Lys-pNA as substrate and an Arrhenius plot was built. It has been reported that thermophilic enzymes may display cold inactivation and they display a non-linear Arrhenius plot. Recently, a non-linear Arrhenius plot for the tetrameric thermophilic enzyme, alcohol dehydrogenase was published. Based on site-directed mutagenesis at the active site of the alcohol dehydrogenase and the thermodynamic parameters determined (ΔH‡, TΔS‡ and the Arrhenius pre-factor Aobs), the authors proposed a direct link between temperature and protein-protein interfaces flexibility, suggesting that at low temperature, the interfaces becomes steeper resulting in the impairment of the alcohol dehydrogenase activity. Interestingly, our data reported above have shown that the M1 site exchanged by Co²⁺, determines a stabilization of the transition state of the PfTET³ enzyme. Furthermore, the M1 site is close to the dimeric and oligomeric interfaces of PfTET³ (Fig. 4). These observations lead us to build a Arrhenius plot for PfTET³Zn by plotting the kcat vs 1/T.

|       | Km (mM) | kcat (s⁻¹) | kcat/Km (s⁻¹M⁻¹) |
|-------|---------|------------|-------------------|
| PfTET³Co, 85 °C | 0.62 ± 0.07 | 2436 ± 191 | 3.9*10⁶ |
| PfTET³Co, 85 °C | 7.0 ± 1.41 | 2890 ± 412 | 4.1*10⁵ |
| PfTET³Zn, 67.5 °C | 8.7 ± 0.6 | 3500 ± 183 | 4.0*10⁵ |
| PfTET³Zn, 50 °C | 4.0 ± 0.6 | 959 ± 101 | 2.4*10⁵ |
| PfTET³Zn, 35 °C | 5.0 ± 0.8 | 707 ± 81 | 1.4*10⁵ |
| PfTET³Zn, 20 °C | 4.0 ± 0.9 | 45.7 ± 0.37 | 1.1*10⁴ |

Table 4. Temperature dependence of PfTET³ kinetic parameters.
The plot resulted non-linear, with a break at 35 °C (Fig. 5A). The data points were fitted by two straight lines, the first covering the points 20 °C < T < 35 °C and the second covering the points 35 °C < T < 85 °C. Both straight lines are described by equations of the type ln(kcat) = −Ea/R*1/T + lnAobs, where Ea is the energy of activation, R is the gas constant and Aobs is the Arrhenius pre-factor. From the slope of both straight lines Ea has been calculated. Interestingly, in the range 20 °C < T < 35 °C, Ea is high (118 kJ/mol) while in the range 35 °C < T < 85 °C, Ea value decreases to 25.8 kJ/mol. The values of the enthalpy of activation (ΔH‡), the entropy of activation (TΔS‡) and of the Aobs are listed in Table 5. It is noteworthy that the Aobs below 35 °C adopts a value of 1022 while above 35 °C, Aobs is 107. The high value of Aobs below 35 °C indicates an unusual temperature dependence compared to the Aobs ≈ 1013 observed for a number of chemical reactions modeled by transition state theory32. Furthermore, in the range 20 °C–35 °C, ΔH‡ adopts a high value (116 KJ/mol). Noteworthy, this high energy barrier is compensated by a high positive TΔS‡, thus implying an increased flexibility of the PfTET3 active site. Indeed, it has been reported that flexibility may lower the energy barrier of the reaction by increasing the number of enzyme conformational substates, resulting in a higher probability for the reaction to occur33,34. Conversely, in the range 35 °C–85 °C, ΔH‡ dropped to 23 KJ/mol, while TΔS‡ became negative (−41 KJ/mol) reflecting an increased rigidity of the active site.

### Table 5. Activation parameters for PfTET3Zn

| Range       | ΔH‡ (kJ/mol) | TΔS‡ (kJ/mol) | Log (Aobs) (s⁻¹) | ΔG‡ (kJ/mol) |
|-------------|--------------|--------------|-----------------|--------------|
| 20 °C–35 °C | 116          | 53           | 10²            | 59.7         |
| 35 °C–85 °C | 23           | −41.5        | 10⁷            | 64.5         |

Table 6. Glu207 pKa values of PfTET3Zn and PfTET3Co at 20 °C and 85 °C.

| pKa          | 20 °C | 85 °C |
|--------------|-------|-------|
| PfTET3Zn     | 5.9   | 4.75  |
| PfTET3Co     | 5.6   | 4.45  |

The plot resulted non-linear, with a break at 35 °C (Fig. 5A). The data points were fitted by two straight lines, the first covering the points 20 °C < T < 35 °C and the second covering the points 35 °C < T < 85 °C. Both straight lines are described by equations of the type ln(kcat) = −Ea/R*1/T + lnAobs, where Ea is the energy of activation, R is the gas constant and Aobs is the Arrhenius pre-factor. From the slope of both straight lines Ea has been calculated. Interestingly, in the range 20 °C < T < 35 °C, Ea is high (118 kJ/mol) while in the range 35 °C < T < 85 °C, Ea value decreases to 25.8 kJ/mol. The values of the enthalpy of activation (ΔH‡), the entropy of activation (TΔS‡) and of the Aobs are listed in Table 5. It is noteworthy that the Aobs below 35 °C adopts a value of 10² while above 35 °C, Aobs is 10⁷. The high value of Aobs below 35 °C indicates an unusual temperature dependence compared to the Aobs ≈ 10¹³ observed for a number of chemical reactions modeled by transition state theory32. Furthermore, in the range 20 °C–35 °C, ΔH‡ adopts a high value (116 KJ/mol). Noteworthy, this high energy barrier is compensated by a high positive TΔS‡, thus implying an increased flexibility of the PfTET3 active site. Indeed, it has been reported that flexibility may lower the energy barrier of the reaction by increasing the number of enzyme conformational substates, resulting in a higher probability for the reaction to occur33,34. Conversely, in the range 35 °C–85 °C, ΔH‡ dropped to 23 KJ/mol, while TΔS‡ became negative (−41 KJ/mol) reflecting an increased rigidity of the active site.

However, a change in rate-limiting step may be at the base of the non-linear Arrhenius plot. In the aminopeptidase enzymatic reaction, the rate limiting step is governed by the general base that deprotonates the water molecule between the two metal ions. In the case of PfTET3, Glu207 is the general base. To evaluate the effect of temperature on the rate limiting step of the enzymatic reaction catalysed by PfTET3, the pKa value of Glu207 was determined at 20 °C and 85 °C in the presence/absence of 300 μM Co²⁺ in the pH range 4–8.7. By plotting the pH on the x-axis and the log(kcat) on the y-axis, a graph was built (data not shown). The data points were fitted with a sigmoidal function, allowing the calculation of the inflection point corresponding to the Glu207 pKa. Interestingly, at 20 °C Glu207 pKa is 5.9 and 5.6 for PfTET3Zn and PfTET3Co, respectively (Table 5). At 85 °C, Glu207 pKa decreases down to 4.7 in PfTET3Zn and to 4.4 in PfTET3Co (Table 6). This means that at low temperature, the PfTET3 enzymatic reaction is chemically unfavored in both variants. The acidity of Glu207 is affected by the amino acids in its surroundings. Interestingly, in the PfTET3Gd structure here reported, Glu207 establishes...
repulsive interactions with the carbonyl carbon of Gly288 (3.3 Å) as well as with the carboxylic side chain of Asp230 (2.7 Å) (Fig. 5B). These interactions diminishes the acidity of Glu207 resulting in an elevated pKa at 20 °C. At 85 °C, the pKa value of Glu207 suggests that the distances with residues Gly288 and Asp230 are increased, resulting in a more efficient enzymatic reaction. Furthermore, the pKa value changes only slightly in the presence of Co2+, suggesting that the observed Co2+ effect on PfTET3 activity is not directly linked to the pKa variation of Glu207. Indeed, the kcat values for PfTET3Zn and PfTET3Co are similar, while their Km differs by a factor of 10. These results support the idea that M1 site is devoted to the regulation of the flexibility of the PfTET3 active site to optimize the enzyme activity at the physiological temperature.

**Figure 6. The M3 site broaden the PfTET3 substrate preferences.** (A) P1 pocket of PfTET3 in electrostatic surface representation. Blue spheres are Co2+ ions, red sphere is a Zn2+ ion. For the sake of the clarity, only the residues interacting with the M3 site are highlighted. The presence of M3 modify the electrostatic properties of P1 allowing the degradation of glutamate. (B) Substrate specificity of PfTET3 is expanded towards glutamate in the presence of Co2+ (blue rectangles) at 85 °C. In the absence of Co2+ (red rectangles) PfTET3 is almost inactive. Error bars are standard deviations of three independent measurements.

The specificity binding pocket (P1) of PfTET3 is devoted to the substrate side-chain accommodation and it is formed by Thr232, Asp254, Glu281, Thr285 and Thr287. This pocket is negatively charged and it allows the degradation of positively charged residues such as lysine and arginine. The evidence of the Co2+ bound in P1 pocket (M3) may change the substrate preferences of PfTET3Co (Fig. 6A). In particular, the presence of the cation could allow the coordination of negatively charged side chains and their subsequent degradation. To examine this possibility, PfTET3 activity was tested at 85 °C by using 5 mM Glu-pNa and Asp-pNa as substrates, in the presence/absence of 300 μM Co2+. Results are reported in Fig. 5B, showing that only PfTET3Co degrades Glu-pNa (Fig. 6B). The activity of PfTET3Co on Glu-pNa represents nearly 30% of the PfTET3Co activity on Lys-pNa. Interestingly, aspartate is not degraded efficiently. The distance between the M1-M2 sites and M3 is 9.5 Å. This length can be covered by a long side-chain, such as lysine, arginine and glutamate. Conversely, aspartate side chain is too short and thus cannot be accommodated efficiently in P1. These results suggest that the M3 site allows extending the type of substrates that can be degraded by PfTET3Co.

**Discussion**

One third of all proteins contain metal ions that are devoted to catalysis and structural stability. Among these, metallopeptidases cover a large portion of the metalloproteins. Metallopeptidases are present in a wide variety of oligomeric states, from monomeric (methionine aminopeptidase, Aeromonas proteolytica Aminopeptidase) to higher oligomeric states (leucine aminopeptidase, tricorn peptidase complex and TET). They all contain one or two metals in the active site that support the catalytic activity of the peptidase. Moreover, it has been shown that different metals in the active site of some of these proteases lead to dramatic variations of the enzyme kinetic parameters. While monomeric peptidases have been extensively studied to determine the role of each metal ion in PfTET3 structure, there is a lack of information regarding the role of each metal ion in oligomeric peptidases. Here, by means of anomalous X-ray crystallography coupled to enzyme kinetics, we have characterized the respective role of each metal ion in the hyperthermophilic PfTET3 aminopeptidase and determined the molecular basis of Co2+ activation.

In the x-ray structure of PfTET3aCl, a Cl− ion is coordinated by the M2 site (Fig. 1D). Similarly, it was reported for the monomeric Streptomyces dinuclear aminopeptidase that F− ion replaces the hydroxide ion on the catalytic metal, resulting in the enzyme activity inhibition. Thus, the position of the Cl− ion in PfTET3aCl structure strongly suggest that M2 is the catalytic metal hosting the nucleophilic hydroxide. This observation is confirmed by the anomalous x-ray experiments performed on PfTET3 crystals grown in the presence of 2 Co2+ equivalents. In this case, the Zn2+ present in the M2 site is not replaced by Co2+ (Fig. 2). Indeed, the kinetics measurements show that the higher catalytic efficiency kcat/Km of PfTET3Co compared to the kcat/Km of PfTET3Zn is due to a lower Km more than to a higher kcat Table 4).

Conversely, the anomalous x-ray experiments on PfTET3 crystals grown in the presence of Co2+ have shown that the M1 site is replaced by Co2+. Interestingly, the M1 site is coordinated by Asp177, Glu208 and His314.
Noteworthy, Glu208 and His314 residues lay on loops implicated in H-bonds between adjacent subunits, mediating the oligomeric and the dimeric interfaces, respectively (Fig. 4C,D). Indeed, the removal of the M1 site by EDTA is sufficient to partially dissociate PfTET3 into monomers (Fig. 3A,B). Furthermore, next to Glu208, there is Glu207 which is the general base that deprotonates the water molecule which in turn performs the nucleophilic attack on the C-N bond of the incoming substrate. Our data revealed that the pKa of Glu207 varies with temperature, adopting optimal values for catalysis at the physiological temperature of the PfTET3 enzyme (Table 6). Moreover, the pKa of Glu207 varied only slightly by Co2⁺ addition. This is in agreement with the slight variation of kcat values observed between PfTET3Co and PfTET3Zn variants (Table 4). Furthermore, it is noteworthy that the ratio kcat/Km of PfTET3Co is ten times higher than that of PfTET3Zn. The kcat/Km parameter relates not only on the enzyme-substrate complex (as kcat) but also on the free enzyme43. This means that PfTET3Co active site is more stable and catalytically active at 85 °C than the PfTET3Zn counterpart. Thus, the M1 site shows a critical role in maintaining the quaternary structure of TET and modulating the flexibility of the enzyme active site. It is noteworthy that the dodecameric PfTET3 particle has twelve M1 sites that are involved in the stabilization of the overall macromolecular architecture. This is demonstrated by the non-linear Arrhenius plot here reported (Fig. 4A). It has been suggested that such a plot may arise for thermophilic enzymes and it represents a reversible distribution of the protein conformations displaying inactive states at low temperature30,31. Although the observed break in the Arrhenius plot here reported may be due to the higher pKa observed for Glu207 at 20 °C compared to 85 °C, it is noteworthy that at both temperatures the addition of Co2⁺ has a slight effect on the Glu207 pKa value, while it has a strong impact on the catalytic efficiency (kcat/Km) of PfTET3 at its physiological temperature. Furthermore, the addition of Co2⁺ at low temperature inhibits PfTET3 activity at a similar concentration at which Co2⁺ enhances PfTET3 activity at 85 °C (Fig. 3C). This means that the replacement of the M1 site by Co2⁺ at 20 °C (≤ 50°C) may impair the equilibrium at the active site and at the interfaces of PfTET3 dodecamer. At low temperature, the hyperthermophilic PfTET3 enzyme has a high enthalpic energy barrier (116 KJ/mol) to overcome. This is attenuated by the high positive value of ΔTS* (53 KJ/mol) in the range 20 °C–35 °C. However, the addition of Co2⁺ may decrease the flexibility of the active site in the range 20 °C–35°C, that is needed to counterbalance the high ΔH* in this temperature range. Conversely, in the range 35 °C–85 °C, ΔH* is lower (23KJ/mol) and ΔTS* adopts a negative value, reflecting a higher conformational rigidity. In this case, the addition of Co2⁺ should favor the enzymatic reaction. Indeed, PfTET3 enzymatic activity is enhanced by Co2⁺ at 85 °C (Table 3 and Fig. 3C). The observed effects of Co2⁺ are physiologically relevant, since recent studies on the metallome of P. furiosus revealed a high intracellular content of Co2⁺, compared to E. coli44.

Because our thermodynamic data showed that PfTET3 transition state is stabilized by Co2⁺ at 85 °C and our structural data identified the M1 site as the metal site exchanged by Co2⁺, we propose that the M1 site has a stabilization role on the PfTET3 active site. These data indicate that PfTET3 shows the optimal activity with an hybrid dinuclear metal site, with M2 and M1 occupied by Zn2⁺ and Co2⁺, respectively. Interestingly, in monomeric aminopeptidases such as AAP and MetAP the M1 and M2 sites appear reversed compared to PfTET3 and the enzymes do not need two metals in their active site to be active45. Conversely, the oligomeric leucyl aminopeptidase strictly needs the presence of two metals for its activity.

The x-ray data collected at Co2⁺ absorption edge highlighted two metal sites occupied by Co2⁺, the M1 site and a new M3 site. The M3 site is coordinated by residues of the P1 pocket (Fig. 5A). Interestingly, an M3 site has been observed at the dimeric interfaces in PfTET3Co, and is coordinated by Glu, Asp and Thr side chains and in addition to three water molecules. In the structure of PfTET3Co here presented, M3 is coordinated by the same protein ligands. Moreover, a M3 site has been observed in methionine aminopeptidases where it is coordinated by two His residues, three water molecules and by a Pro-Leu inhibitor molecule26,36,44. Furthermore, in endonucleases, a functional equivalent metal binding site has been proposed to act as stabilizing the negative charge of the phosphate transition state, by coordinating a water molecule that protonate the 3’ oxygen of the leaving group57,48. In this study we report for the first time a new function for the M3 site. Indeed, by modifying the electrostatic properties of the P1 pocket of the dodecameric aminopeptidase PfTET3, M3 allows the degradation of a negatively charged substrate (glutamate). Furthermore, it is noteworthy that M3 does not interfere with positively charged substrates (Fig. 5B). Indeed, at the same Co2⁺ concentration by which PfTET3 degrades glutamate, it also processes lysine. This observation suggests that M3 may also be responsible for the stabilization of the negatively charged transition state, as occurs in endonucleases and alkaline phosphatase. The presence of M3 has important biological implications by considering the recently characterized hetero dodecameric assembly in Pyrococcus horikoshii formed by six TET2 subunits and six TET3 subunits, identified in vivo and characterized in vitro16,17. This TET2_TET3 hetero dodecamer resulted more efficient in degrading complex peptides than the two separated homo-dodecamers of TET2 and TET3. The M3 site here presented represents a further upgrade in peptide processing, allowing the TET2_TET3 hetero dodecamer to be active on a wide variety of peptides.

In summary, this study allowed the characterization of the role of each metal site in the TET enzymes. The three metals are key players for the effective catalytic activity of TET as well as for the flexibility modulation of the active site. In particular, Co2⁺ may ensure the optimal geometry of the PfTET3 active site at high temperature. To our knowledge, it is the first time that such molecular interplay among three metal ions is observed for an oligomeric aminopeptidase. Furthermore, these data depict a connection between the organism living temperature and the metal selection in the TET active site. We speculate that the choice of metal in the active site of TET may be determined by the organism living temperature. It is noteworthy that the optimal concentration of Co2⁺ (300μM) enhancing PfTET3 activity in vitro reported in this paper is compatible with the range of in vivo Co2⁺ concentration (from 85μM to 9 mM) found in the surface soil of Volcano island in Sicily, where Pyrococcus furiosus was first isolated49. Intriguingly, mesophilic TET aminopeptidases such as DNPEP and PmM18 are preferentially activated by Mn2⁺, while Co2⁺ promotes their activity to a lower extent. It could be of interest to study the molecular basis that lead the mesophilic TET to select Mn2⁺ instead of Co2⁺.

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Materials and Methods

Production and purification of PfTET3. PfTET3 was cloned in vector pET-41c by RoBioMol (RoBioMol platform at the IBS (CEA/CNRS/UMR), Grenoble). Expression and purification were carried out in the same way as for PfTET3\(^{30}\). The final step is a gel filtration in buffer Tris 20 mM pH 7.5, NaCl 300 mM. PfTET3 was concentrated in Amicon Ultra 100 kDa at a final concentration of 8 mg/ml for crystallization.

Crystallization of PfTET3. Initial crystal hits were obtained by using the HTX lab platform at the EMBL, Grenoble\(^{30}\). The crystal conditions were optimized with hanging drop vapour diffusion method. The final crystal conditions were HEPES 0.1 M pH 7.7, MPD 44%, NH\(_4\)CH\(_3\)COO 0.2 M. Then, three types of PfTET3 crystals were prepared: PfTET3\(_{Gd}\) to determine the PfTET3 x-ray structure de novo, PfTET3\(_{Co}\) to evaluate stoichiometric metal exchange in the active site and PfTET3\(_{Zn}\) to show the presence of two metal sites after purification. PfTET3\(_{Gd}\) crystals were grown in HEPES 0.1 M pH 7.7, MPD 44%, NH\(_4\)CH\(_3\)COO 0.2 M and CoCl\(_2\) 2 mM and the drops were produced by adding 1.5 μl PfTET3 8 mg/ml + 1.5 μl reservoir + 1.5 μl 300 mM Gd-DOD3A. Gd-DOD3A is a lanthanide complex designed to obtain high-phasing power heavy atom derivatives. In this complex, a Gd\(^{3+}\) ion is coordinated by a tetraazacyclododecane macrocycle and two water molecules. By interacting with a carboxylic group from a macromolecule (aspartate or glutamate), the lanthanide complex loses one water molecule and establishes a complex with the macromolecule. CoCl\(_2\) was added because it ameliorates the quality of the crystals. PfTET3\(_{Co}\) crystals were grown in HEPES 0.1 M pH 7.7, MPD 44%, NH\(_4\)CH\(_3\)COO 0.2 M and CoCl\(_2\) 0.4 mM and the drops contained 1.5 μl PfTET3 8 mg/ml + 1.5 μl reservoir.

Crystal structure determination of PfTET3\(_{Gd}\). PfTET3\(_{Gd}\) crystals were cryo-cooled with liquid nitrogen in mother liquors. X-ray diffraction intensities were collected on ID29 at the European synchrotron radiation facility, in Grenoble (ESRF) at Gd LIII wavelength, after energy scan. Data were processed with XDS\(^{51}\) and Aimless\(^{52}\). Phase shift (α) estimation of Gd atoms was made using SHELXC\(^{53}\). The resolution was initially cut at 3.0 Å based on the strength of the anomalous signal estimated by d''/sig (d''/sig as implemented in shelx\(^{53}\) software represents the anomalous differences divided by their estimated standard deviation). These initial substructures were input in Phaser, using SAD setup\(^{54}\) extending resolution up to 2.5 Å. Nine additional substructures were found and initial PfTET3 phases were determined by SHELXE\(^{53}\). Three cycles of density modification were performed by cparrot\(^{55}\), obtaining an initial FOM = 0.73. Initial model building was performed by BUCCANEER\(^{56}\) and the model was completed manually using COOT\(^{57}\). Model refinement was performed by PHENIX\(^{58}\). To calculate the R\(_{free}\), 5% of the reflections were excluded throughout the refinement process. Data collection statistics are listed in Table 1.

Table 1. Images were prepared using CCP4mg software\(^{59}\).

Determination of the heavy atoms sites with ANODE of PfTET3\(_{Gd}\), PfTET3\(_{Zn}\) and PfTET3\(_{Co}\). PfTET3\(_{Gd}\) crystals were cryo-cooled with liquid nitrogen in mother liquor. X-ray diffraction intensities were collected on PROXIMA II beamline at the french synchrotron SOLEIL, in Paris. Data were collected at Co\(^{2+}\)-edge and Zn\(^{2+}\)-edge. The absorption edges were evaluated with a fluorescence spectrum. The intensities were processed with XDS\(^{51}\) and Aimless\(^{52}\). Data collection and reduction revealed the same tetragonal space group (98) observed in de novo PfTET3\(_{Gd}\) pdb model. Matthews's coefficient proposes three monomers (A, B and C) in the ASU at both Zn\(^{2+}\)-edge and Co\(^{2+}\)-edge. In both cases molecular replacement solutions confirmed Matthews's coefficient estimation. Then, phase shift (α) estimation of cobalt and zinc atoms were successively calculated by SHELXC\(^{53}\) cutting the resolution at 3.35 Å and 3.25 Å for Co\(_2\)\(^{2+}\) edge and Zn\(_2\)\(^{2+}\) edge. In both cases molecular replacement solutions confirmed Matthews's coefficient estimation. Nine additional substructures were found and input in ANODE together with the PDB model of PfTET3Gd depleted of metal ions. The files produced by SHELXC were then input in ANODE. The number of sites found by ANODE for both datasets is listed in Table 2. Data collection statistics are listed in Table 1. Images were prepared using CCP4mg software\(^{59}\).

PfTET3\(_{Zn}\) titration by Co\(^{3+}\) or EDTA. 10 nM PfTET3\(_{Zn}\) (relative to monomer) was titrated with CoCl\(_2\). The enzymatic activity was measured using 5 mM Lys-pNa as substrate at T = 20 °C, 27 °C, 35 °C, 50 °C, 67.5 °C and 85 °C. The reaction mixture contained 0.1 M HEPES pH 7.2, KCl 316 mM and 0 mM – 5 mM CoCl\(_2\). At T = 20 °C, 100 nM PfTET3\(_{Zn}\) was used due to the lower activity of PfTET3\(_{Zn}\) enzyme at this temperature. The same procedure was adopted for the EDTA titration, but the measurements were executed only at 85 °C and using 0 mM- 20 mM EDTA. All buffer solutions were pre-warmed and the pH is referred to the working temperature. Measurements were performed in triplicate. The reported specific activities were calculated as described above. 100% of the activity corresponds to the activity of the PfTET3\(_{Zn}\) enzyme.

PfTET3\(_{Zn}\) oligomerization in the presence of EDTA. Three reaction tubes of 500 μl of HEPES pH 7.2 (at working temperature), KCl 316 mM containing 1 μM PfTET3\(_{Zn}\) alone, 1 μM PfTET3\(_{Zn}\) + 15 μM EDTA and 1 μM PfTET3\(_{Zn}\) + 6 μM EDTA respectively, were heated at 85 °C for 5 minutes. The tubes were subsequently cooled at room temperature and loaded on a Superose 6 column, preequilibrated with the same buffer used for the activity assay, without EDTA.

Kinetics of PfTET3 enzymatic activity on chromogenic substrates. PfTET3\(_{Zn}\) and PfTET3\(_{Co}\) enzymatic activities were determined by monitoring the release of pNa at λ = 405 nm from different substrates (lysine, glutamate, aspartate) at T = 20 °C, 27 °C, 35 °C, 50 °C, 67.5 °C and 85 °C. The reaction mixture contained 10 nM PfTET3 (relative to monomer), 0.1 M HEPES pH 7.2, KCl 316 mM, 5 mM substrate at a final volume of 100 μl. pH is referred to the working temperature. All buffer solutions were pre-warmed. Only for the measurements at 20 °C, PfTET3 was at 100 mM final concentration. Measurements were performed in triplicate by using a multi-cuvette JASCO V-630 UV-visible spectrophotometer. Temperature was controlled by a Peltier system. The enzymatic reaction was monitored along 10 minutes at 20 °C and for 5 minutes at higher temperatures. To calculate the specific activity, the linear slope (A/t) of the enzymatic reaction corresponding to the steady-state
of the reaction was determined by using Spectra analysis software provided by Jasco. The slope was converted to \( V_0 \) (M/s) using the pNA extinction coefficient (\( \varepsilon \)) (10,000 M\(^{-1}\) cm\(^{-1}\)). The specific activity was then calculated by dividing \( V_0 \) per PfTET3 added volume and PfTET3 mg/ml concentration. Plots were drawn using Origin software version 8.5. The kinetic parameters (Km, kcatal and kcat/Km) were obtained by plotting \( V_0 \) as a function of substrate concentration (0.2 mM – 20 mM). Data points were fitted by the Hill equation provided with Origin 8.5 (\( y = V_{\max} \times (x^n)/(Km+x^n) \)) and fixing term \( n = 1 \) to get the Michaelis–Menten equation.

Effect of metal cations on PfTET3\(_{\text{Zn}}\) activity as function of temperature. Purified PfTET3\(_{\text{Zn}}\) enzyme activity was measured using 5 mM Lys-pNa as substrate at T = 20 °C, 50 °C and 85 °C in the buffer 0.1 M HEPES pH 7.2, KCl 316 mM and individually supplied with 0.1 mM / 1 mM of CoCl\(_2\), MnCl\(_2\) and ZnCl\(_2\). All buffer solutions were pre-warmed and the pH is referred to the working temperature. Measurements were performed in triplicate. The reported specific activities were calculated as described above.

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were deposited in SBGrid60.

The authors declare no competing financial interests.

Data availability: The coordinates of PfTET3Gd model and their experimental intensities are deposited in the PDB database (PDB 4X8I). The original diffraction images collected for PfTET3Co, PfTET3Zn and PfTET340eV were deposited in SBGrid60.

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