

The structure of the catalytic domain of *Tannerella forsythia* karilysin reveals it is a bacterial xenologue of animal matrix metalloproteinases

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Summary

Metallopeptidases (MPs) are among virulence factors secreted by pathogenic bacteria at the site of infection. One such pathogen is *Tannerella forsythia*, a member of the microbial consortium that causes periodontitis, arguably the most prevalent infective chronic inflammatory disease known to mankind. The only reported MP secreted by *T. forsythia* is karilysin, a 52 kDa multidomain protein comprising a central 18 kDa catalytic domain (CD), termed Kly18, flanked by domains unrelated to any known protein. We analysed the 3D structure of Kly18 in the absence and presence of Mg²⁺ or Ca²⁺, which are required for function and stability, and found that it evidences most of the structural features characteristic of the CDs of mammalian matrix metalloproteinases (MMPs). Unexpectedly, a peptide was bound to the active-site cleft of Kly18 mimicking a left-behind cleavage product, which revealed that the specificity pocket accommo-

dates bulky hydrophobic side-chains of substrates as in mammalian MMPs. In addition, Kly18 displayed a unique Mg²⁺ or Ca²⁺ binding site and two flexible segments that could play a role in substrate binding. Phylogenetic and sequence similarity studies revealed that Kly18 is evolutionarily much closer to winged-insect and mammalian MMPs than to potential bacterial counterparts found by genomic sequencing projects. Therefore, we conclude that this first structurally characterized non-mammalian MMP is a xenologue co-opted through horizontal gene transfer during the intimate coexistence between *T. forsythia* and humans or other animals, in a very rare case of gene shuffling from eukaryotes to prokaryotes. Subsequently, this protein would have evolved in a bacterial environment to give rise to full-length karilysin that is furnished with unique flanking domains that do not conform to the general multidomain architecture of animal MMPs.

Introduction

Metallopeptidases (MP) are mostly zinc-dependent protein and peptide hydrolases with major endogenous roles in the physiology and pathology of living organisms. In addition, MPs have exogenous functions in the venoms used by predators, such as in poisonous snakes, scorpions and spiders, or, conversely, in poison-mediated defence strategies against predators such as in scorpion fish (Carrijo *et al.*, 2005; King, 2007; Fox and Serrano, 2009). Another source of exogenous MPs is bacteria, which secrete them as virulence factors during host infection. In this condition, MPs damage host tissues by cleaving cell-surface and tissue proteins, either directly or indirectly by activating host peptidases. MPs also inactivate key proteins in host defence, recruit nutrients, and activate other bacterial virulence factors. These strategies are required for bacterial invasion, survival, proliferation and colonization of the host in a hostile environment (Lantz, 1997; Miyoshi and Shinoda, 2000; Potempa and Pike, 2009). One such human infection is periodontitis, which causes chronic inflammation of the gums and affects 10–15% of adults worldwide, leading to gingival

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tissue destruction and tooth loss (Fox, 1992; Cutler *et al.*, 1995). Due to their inflammatory and infective character, severe forms of the disease further contribute to the development of systemic pathologies such as cardiovascular disease, stroke, rheumatoid arthritis and diabetes (Jordan, 2004; Pihlstrom *et al.*, 2005; Behle and Papanou, 2006; Persson, 2006; Wegner *et al.*, 2010). Current treatment and curettage of severe periodontal disease is only partially effective. Furthermore, it entails the intensive usage of antibiotics, which contributes to the spread of antibiotic resistance (Haffajee *et al.*, 2003; Pihlstrom *et al.*, 2005; Suvan, 2005). Consequently, there is a need for innovative and specific therapeutic approaches to this pathology, and detailed structural information of participating virulence factors may contribute to this development (Mittl and Grütter, 2006).

Periodontitis is caused by bacteria that grow on tooth surfaces and in the gingival crevice, and three species, *Porphyromonas gingivalis*, *Treponema denticola* and *Tannerella forsythia*, are the major periodontopathogens, collectively referred to as the red complex (Socransky *et al.*, 1998). These pathogens secrete a vast armamentarium of proteolytic enzymes, which has been thoroughly characterized for the first two organisms (Potempa *et al.*, 2000; Eley and Cox, 2003). In contrast, the only peptidases identified to date in *T. forsythia* are a cysteine proteinase, PrtH (Saito *et al.*, 1997), and the MP karilysin (Karim *et al.*, 2010). The latter is a 472-residue secretory protein comprising a 20-residue signal sequence, a 14-residue pro-peptide, an 18 kDa catalytic peptidase domain (CD) and a 30 kDa C-terminal domain of unknown function. The latter has been found only in other putative secretory proteins of *T. forsythia* (Karim *et al.*, 2010). The recombinant full-length enzyme gives rise to karilysin CD – hereafter Kly18 – through sequential autolysis. The first cut at Asn34–Tyr35 (numbering according to the complete gene sequence, see UniProt D0EM77) removes the pro-peptide and generates a fully active 48 kDa variant. Subsequent autolytic cleavage events happen downstream of the CD and do not affect activity (Karim *et al.*, 2010).

Whereas the flanking sequences do not resemble any other structurally or functionally analysed protein, Kly18 is similar to mammalian matrix metalloproteinases (MMPs) (see fig. 2 in Karim *et al.*, 2010). These constitute a separate family within the ‘metzincin’ clan of MPs (Bode *et al.*, 1993; Stöcker *et al.*, 1995; Gomis-Rüth, 2003; 2009; Visse and Nagase, 2003; Murphy and Nagase, 2008; Tallant *et al.*, 2010b) and participate in both broad-spectrum turnover of extracellular-matrix components and selective limited proteolysis to activate or inactivate other proteins and enzymes. When dysregulated, their degrading potential leads to uncontrolled proteolysis and tissue destruction, apoptosis and inflammation, as observed in periodontitis among other conditions (Overall and López-

Otín, 2002). MMPs have mainly been studied in mammals, and 23 paralogues are present in humans [see <http://degradome.uniovi.es/met.html#M10> (Puente *et al.*, 2003; Quesada *et al.*, 2009)]. They are secreted as soluble or membrane-bound mosaic proteins comprising several inserts and domains, which include a signal peptide for secretion, an ~80-residue zymogenic pro-domain including a conserved cysteine engaged in latency maintenance (Rosenblum *et al.*, 2007), a ~165-residue zinc- and calcium-dependent CD, a linker region and a ~200-residue haemopexin-like C-terminal domain for collagen binding, pro-MMP activation and dimerization (Gomis-Rüth, 2004; Maskos, 2005). In addition to mammals, MMPs have been studied (at least at the mRNA level) in several classes of animals and plants (Table 1). As to lower eukaryotes and prokaryotes, sequences similar to MMP CDs have been found in the genomes of fungi, viruses, archaea and bacteria, but to our knowledge they have not been studied (Gomis-Rüth, 2003). At the structural level, only human, mouse and pig MMP CDs have been analysed so far (see Tallant *et al.*, 2010b and <http://pfam.sanger.ac.uk/family/PF00413#tabview=tab8>).

In order to explore the mechanisms of action of karilysin and to provide a high-resolution scaffold for the design of specific inhibitors against this bacterial virulence factor, we studied structure and function of Kly18, in the absence and presence of alkaline-earth metal ions. In addition, phylogenetic studies and evolutionary considerations rendered a plausible explanation for the origin of this MP.

Results and discussion

Endopeptidolytic activity of Kly18

The distinct (autolytic) variants of *T. forsythia* karilysin have been previously characterized for their proteolytic potential (Karim *et al.*, 2010; Koziel *et al.*, 2010). These studies revealed that the enzyme requires calcium for activity and thermal stability and that it cleaves bonds preferentially N-terminally of bulky hydrophobic residues. It efficiently degraded elastin, fibrinogen and fibronectin, thus pointing to possible roles in periodontitis progression. In addition, karilysin inactivated the antimicrobial peptide, LL-37, which is a component of the immune system that targets bacteria and other microorganisms, thus further supporting a role of this MP in *T. forsythia* virulence. Here, we further studied the endopeptidolytic potential of Kly18 in front of two fluorogenic peptide substrates of seven and 10 residues respectively. These were developed to be cleaved by distinct mammalian MMPs, with k_{cat}/K_m values ranging between $1E4$ and $6E5\text{ M}^{-1}\text{ s}^{-1}$ (Knight *et al.*, 1992; Nagase *et al.*, 1994; Fields, 2001). These substrates were efficiently cleaved by Kly18, with k_{cat}/K_m values of 4895 and $14\,709\text{ M}^{-1}\text{ s}^{-1}$ for the two substrates, respectively, thus

Table 1. MMPs reported from non-mammalian organisms.Kingdom *Animalia**Amphibia*

African clawed frog *Xenopus laevis* (Yang *et al.*, 1997); bullfrog *Rana catesbeiana* (Gross and Lapière, 1962; Oofusa *et al.*, 1994); Japanese newt *Cynops pyrrhogaster* (Suzuki *et al.*, 2001); Eastern newt *Notophthalmus viridescens* (Vinarsky *et al.*, 2005); Mexican axolotl *Ambystoma mexicanum* (Yang and Bryant, 1994); green-striped burrowing frog *Cyclorana albogutta* (Hudson *et al.*, 2007); Pacific tree frog *Pseudacris regilla* (Veldhoen *et al.*, 2006)

Reptilia

Spectacled caiman *Caiman crocodilus apaporiensis* (Shintani *et al.*, 2007); tiger keelback snake *Rhabdophis tigrinus tigrinus* (Komori *et al.*, 2006)

Aves

Chicken *Gallus gallus* (Yang and Kurkinen, 1998); common canary *Serinus canaria* (Kim *et al.*, 2008)

Actinopterygii (fish)

Rainbow trout *Oncorhynchus mykiss* (Saito *et al.*, 2000); zebrafish *Danio rerio* (Wyatt *et al.*, 2009); Atlantic salmon *Salmo salar* (Skugor *et al.*, 2008); Japanese pufferfish *Takifugu rubripes* (Tsukamoto *et al.*, 2007); Amur catfish *Silurus asotus* (Cho *et al.*, 2002); medaka fish *Oryzias latipes* (Matsui *et al.*, 2000); channel catfish *Ictalurus punctatus* (Yeh and Klesius, 2008)

Hydrozoa

Hydra *Hydra vulgaris* (Leontovich *et al.*, 2000)

Echinoidea

Sea urchins *Paracentrotus lividus* and *Hemicentrotus pulcherrimus* (Gache *et al.*, 2004)

Secernentea (nematodes)

Microbivorous roundworm *Caenorhabditis elegans* (Wada *et al.*, 1998); parasitic nematode *Gnathostoma spinigerum* (Uparanukraw *et al.*, 2001); soybean cyst nematode *Heterodera glycines* (Kovaleva *et al.*, 2004); potato cyst nematode *Globodera rostochiensis* (Kovaleva *et al.*, 2004)

Insecta

Fruitfly *Drosophila melanogaster* (Llano *et al.*, 2000; 2002); wax moth *Galleria mellonella* (Altincicek and Vilcinskis, 2008); red flour beetle *Tribolium castaneum* (Knorr *et al.*, 2009)

Bivalvia (mollusks)

Eastern oyster *Crassostrea virginica* (M. Gomez-Chiarri, pers. comm.); abalone (sea snail) *Haliotis diversicolor* (Wang *et al.*, 2008)

Kingdom *Plantae*

Soybean *Glycine max* (Ragster and Chrispeels, 1979); thale cress *Arabidopsis thaliana* (Maidment *et al.*, 1999); cucumber *Cucumis sativus* (Delorme *et al.*, 2000); sugarcane *Saccharum* spp. (Ramos and Selistre-de-Araujo, 2001); barrel medic *Medicago truncatula* (Combier *et al.*, 2007); loblolly pine *Pinus taeda* (Ratnaparkhe *et al.*, 2009); tobacco plant *Nicotiana tabacum* cv. BY-2 (Schiemeyer *et al.*, 2009)

indicating that the bacterial enzyme and its mammalian counterparts evince comparable substrate specificity and efficiency.

Structure of Kly18

The structure of Kly18 was solved both in the absence and in the presence of magnesium, hereafter 'unbound' and 'magnesium-bound' variants respectively. These variants crystallized in distinct space groups and contained two and one protein molecules, respectively, in the crystallographic asymmetric unit. Calcium had been reported to be important for karilysin activity and thermal stability (Karim *et al.*, 2010), and so this cation was added to the protein prior to crystallization to produce a calcium-bound form. However, in the crystal structure, which was obtained to significantly higher resolution than the unbound form (Table 2), only one partially occupied magnesium site was found as inferred from electron density maps and distances to ligands, probably owing to the excess of the latter over calcium in the crystallization conditions (see *Experimental procedures*).

The Kly18 molecule is roughly spherical, with a diameter of ~40 Å. Carved into its frontal surface, a shallow

active-site cleft traverses the molecule left (non-primed side) to right (primed side) and divides it into a large N-terminal (NTS; Tyr35–Gly162) and a small C-terminal subdomain (CTS; Ile163–Ser201; see Fig. 1A and B). Both the N- and the C-terminus are on the lower left of the molecule surface. The N-terminal α -amino group, in the front, is salt-bridged to the side-chain of Asp187 within the 'C-terminal helix' α C of CTS and thus anchored to the peptidase moiety. The NTS consists of a twisted five-stranded β -sheet (strands β I– β V), whose strands are all parallel to a substrate bound to the cleft except for outermost strand β IV, which forms the upper rim of the crevice and runs antiparallel. Two helices, the 'backing helix' (α A) and the 'active-site helix' (α B), nestle into the concave side of the sheet (Fig. 1A and C). On the convex side of the latter, three loops protrude from the molecular surface: the loop segment connecting β II and β III (L β II β III), L β IV β V and L β III β IV. The latter, termed 'S-loop' due to its sinuous chain trace, encompasses a structural zinc binding site, at which the cation is tetrahedrally co-ordinated by three histidines and an aspartate (His102, Asp104, His117 and His133; Fig. 1D). The final stretch of the S-loop is termed 'bulge-edge segment' and it shapes the upper rim of the active-site cleft on its primed side (Fig. 1A). This loop

Table 2. Crystallographic data.

Data set	Kly18 with Mg ²⁺	Kly18 without Mg ²⁺
Space group/cell constants (<i>a</i> , <i>b</i> , <i>c</i> , in Å; β in ° if $\neq 90$)	14/82.79, 85.79, 53.48	C2/121.4, 53.1, 86.3, 134.6
Wavelength (Å)	0.9790	0.8726
No. of measurements/unique reflections	153 136/21 225	62 645/15 411
Resolution range (Å) (outermost shell) ^a	30.3–1.70 (1.79–1.70)	45.3–2.40 (2.53–2.40)
Completeness (%)	99.0 (95.9)	99.3 (95.9)
R_{merge}^b	0.034 (0.121)	0.088 (0.511)
$R_{\text{t.i.m.}} (= R_{\text{meas}}^b/R_{\text{p.i.m.}}^b)$	0.036 (0.139)/0.013 (0.065)	0.101 (0.608)/0.050 (0.322)
Average intensity ($\langle I \rangle / \langle I \rangle_{\text{hkl}}$)	37.1 (10.8)	14.8 (2.5)
B-factor (Wilson) (Å ²)/average multiplicity	18.9/7.2 (4.4)	38.4/4.1 (3.4)
Resolution range used for refinement (Å)	∞–1.70	∞–2.40
No. of reflections used (test set)	20 403 (820)	14 703 (708)
Crystallographic R_{factor} (free R_{factor}) ^b	0.158 (0.186)	0.189 (0.259)
No. of protein atoms/solvent molecules/ligands/ions	1328/196/ 1 tripeptide (AFT)/2 Zn ²⁺ , 1 Mg ²⁺ , 1 Cl [−]	2644/115/ 1 tris, 1 tetrapeptide (AFTS)/2 Zn ²⁺
<i>Rmsd</i> from target values		
Bonds (Å)/angles (°)	0.006/1.03	0.013/1.32
Bonded B-factors (main-chain/side-chain) (Å ²)	0.46/1.23	0.60/1.46
Average B-factors for protein/peptide ligand atoms (Å ²)	16.6/19.6	33.3/50.2
Main-chain conformational angle analysis ^c		
Residues in favoured regions/outliers/all residues	160/0/166	315/0/332

a. Values in parentheses refer to the outermost resolution shell.

b. For definitions, see table 1 in Mallorquí-Fernández *et al.* (2008).

c. According to MOLPROBITY (Davis *et al.*, 2007).

opens out into the active-site helix, which includes the first half of a zinc-binding consensus sequence, HEXXHXXG/NXXH/D (amino acid one-letter code; X for any residue), which is characteristic of MMPs in particular and metzincins in general (Gomis-Rüth, 2003; 2009; Tallant *et al.*, 2010b). The first two histidines of the motif, His155 and His159, bind the catalytic zinc ion through their N ϵ 2

atoms. Imbedded between them is the general-base/acid required for catalysis, Glu156. After this helix, the NTS ends at the glycine of the consensus sequence (Gly162), and the polypeptide enters the CTS. The latter residue allows for a sharp turn in the trajectory of the polypeptide to reach the third zinc ligand of the motif, His165. After this residue, Ser166 forms a hydrogen bond with Asp187,

Fig. 1. Structure of Kly18.

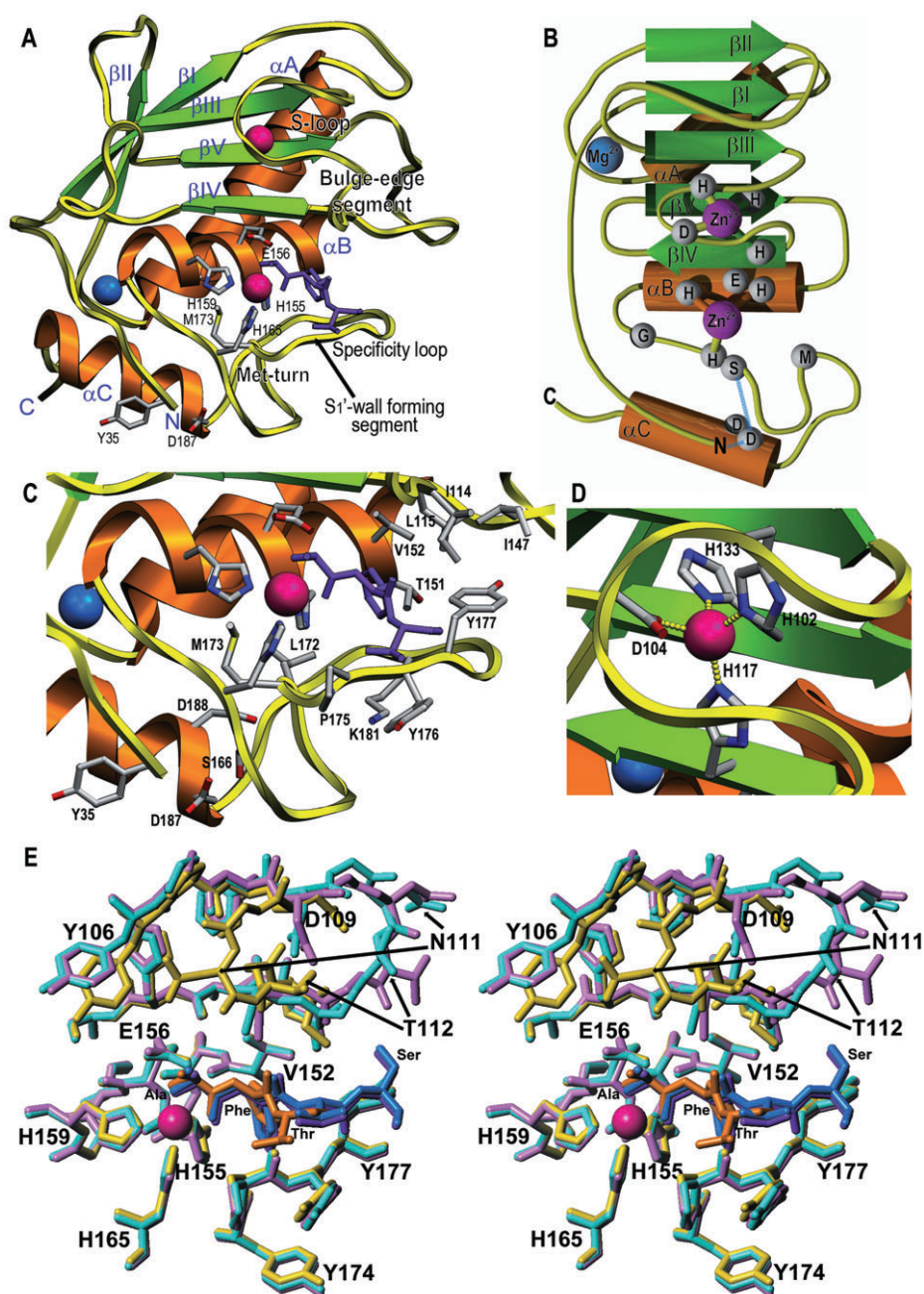
A. Richardson-type plot of Kly18 in the commonly accepted standard orientation for MPs, i.e. with the view into the active-site cleft, which runs horizontally from left [non-primed side; it accommodates substrate residues upstream of the cleavage bond at subsites S₁, S₂, S₃, etc. according to Schechter and Berger (1967)] to right (primed side; it binds downstream residues at S₁', S₂', S₃', etc.). The regular secondary structure elements are shown as green arrows (β -strands β I to β V) and orange ribbons (helices α A to α C) and labelled. Residues participating in catalytic-zinc binding, as well as the general base/acid glutamate, the Met-turn methionine, and the residues engaged in anchoring of the N-terminus to the protein moiety are displayed as sticks and labelled. The cleavage-product peptide bound to the primed side of the active-site cleft is shown as deep purple sticks. The zinc and magnesium ions are shown as magenta and blue spheres respectively. Chain segments characteristic of MMPs and discussed in the text are further pinpointed. Helix α A is inserted after strand β I and runs across the back surface of the molecule from top right to bottom left. Loop L β I α A is flexible at Ser54–His57 and was traced on the basis of weak electron density maps. Helix α B follows strand β V and runs horizontally across the centre of the molecule, contributing to the back wall of the active-site cleft. The two aforementioned helices contact each other through the side-chains of Ala70 (α A) and Ala154 (α B), which participate in a central hydrophobic core together with residues from the sheet. In addition, a significant part of the 'bulge-edge segment' was likewise flexible in Kly18 and was traced on the basis of weak electron density maps at Asp109–Thr112. After the S-loop and strand β IV, L β IV β V includes three prolines in a row (Pro121–Pro123), which adopt a 1,4-turn structure and are followed by an alanine and two glycine residues. Thereafter, the polypeptide enters strand β V and a large loop, L β V α B, which contributes to cleft delimitation and substrate binding.

B. Topology scheme of Kly18 illustrating the regular secondary structure elements and cations with the same colour code as in (A). Hydrogen bonds and salt bridges are displayed as blue dashed lines.

C. Close-up view of (A) showing the active site and residues framing the specificity pocket or S₁' subsite.

D. Close-up view of (A) centred on the structural zinc ion.

E. Superposition in stereo of the unbound (the two molecules in the asymmetric unit are shown as cyan and purple sticks respectively) and magnesium-bound (yellow sticks) Kly18 structures with the corresponding cleavage-product peptides as blue/deep purple and orange sticks respectively. Variation between the magnesium-bound and -unbound structures was found at L β I α A (slight variation) and within the bulge-edge segment of L β III β IV (large variation), which appears as essentially two conformers. The latter segment is folded inward in the magnesium-bound structure and outward in the unbound structure molecules, leading to a maximal displacement of ~13 Å (measured at Asn111 C α). As the bulge-edge segment contributes to substrate binding on the primed side of the cleft, an outswung conformation should enable binding of longer substrates. This resulted in that the peptide found in the cleft could be traced for one more residue, a tentative serine, in the unbound structure due to additional electron density.



which also binds the N-terminus of the molecule (see above). Thereafter, four residues lead to a tight 1,4-turn of type I (Ala171–Tyr174), termed the ‘Met-turn’ due to an invariable methionine found at the third turn position in all MMPs and metzincins (Gomis-Rüth, 2003; 2009; Tallant *et al.*, 2010a,b). This turn is stabilized by a double main-chain interaction with the side-chain of Asp188 within helix α C (Fig. 1C). Between the Met-turn and this helix, segment Pro175–Tyr177, termed ‘S₁’-wall forming segment’, shapes the lower rim of the cleft on its primed side. Thereafter the chain enters the ‘specificity loop’

(Gly179–Gln183), which delimitates the back of the S₁’ subsite or specificity pocket of Kly18 (Fig. 1A and C). At Asn186, the chain enters the C-terminal helix, α C, which leads into the C-terminus on the molecular surface.

Kly18 was obtained as a serendipitous complex with a peptide, which mimics a slightly shifted C-terminal cleavage product bound to the primed side of the cleft (Fig. 1A, C and E). The high resolution of the electron density maps enabled us to assign a tripeptide of sequence Ala–Phe–Thr in the magnesium-bound structure. The N-terminus of this peptide is bound to both the catalytic zinc ion and the

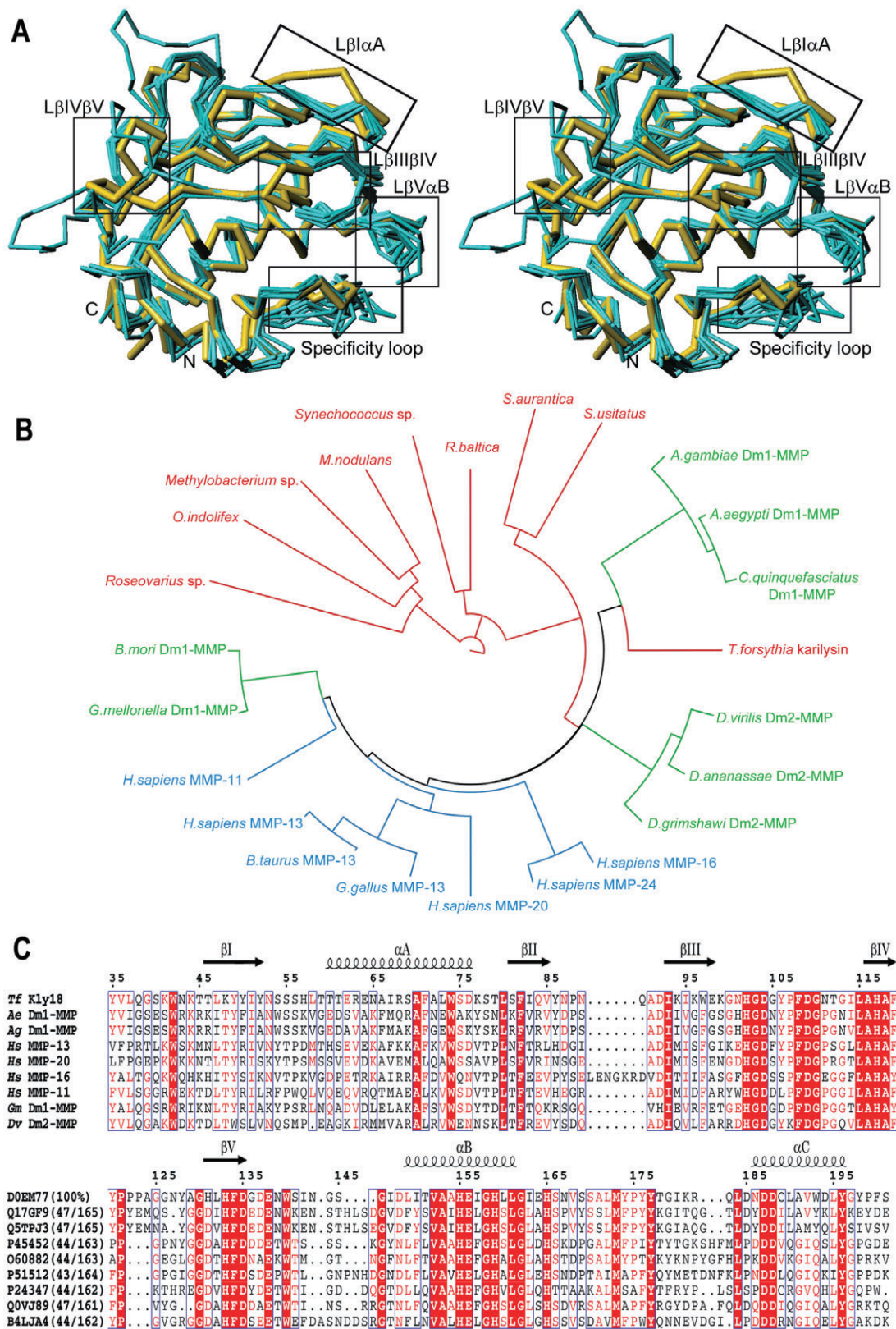


Table 3. Structural similarities of Kly18 with metzincins according to DALI (Holm and Sander, 1998).

Metzincin family	Z-score	rmsd (Å)	Common sequence stretch (residues)	Sequence identity (%)
MMPs	20.5–25.6	1.3–1.8	147–156	42–46
Serralysins	14.9–16.3	2.3–2.8	132–158	20–24
Snapalysin	13.5	2.3	119	20
Astacins	11.5–11.8	2.4–2.7	123–131	14–20
Adamalysins/ADAMs	9.8–11.4	2.6–3.4	129–141	13–19
Ulilysin	9.1–9.2	2.9–3.1	135–138	12–13
Leishmanolysin	8.7	3.2	129	17

general-base/acid glutamate, and the carbonyl oxygen of the first residue also binds the zinc. This arrangement is strongly reminiscent of hydroxamate-based MP inhibitors, which bind zinc through the oxygen atoms of an NH(OH)-CO warhead (Betz *et al.*, 1997). The downstream phenylalanine occupies the S₁' pocket, which is delimited by Leu115, Ile116, Thr151, Val152, His155, Leu172, Tyr177 and Lys181 (Fig. 1C). These residues confer an overall hydrophobic character to this rather large cavity and explain the preference of Kly18 for bulky hydrophobic residues in position P₁' of substrates (Karim *et al.*, 2010).

A magnesium site was found on the back of the molecule, close to the C-terminus. The ion is co-ordinated by Ser75 O of α A and Ser78 O of α A β II. Three solvent molecules defined by proper electron density and a position occupied by a bulk solvent complete an octahedral ligand sphere characteristic for alkaline-earth metal cations. Superposition of the magnesium-bound structure over the two crystallographically independent molecules of the unbound structure revealed that they are practically equivalent, with pairwise rmsd values of ~0.5 Å (Fig. 1E). No significant differences were observed at the magnesium site apart from the lack of the cation. In contrast, variations were observed within two flexible segments, L β I α A and the bulge-edge

segment (see Fig. 1E and its legend). However, as the magnesium site is far away from the active-site cleft and the latter segments (Fig. 1A and C), there is no evident structural explanation for the these differences or for the requirement of alkaline-earth ions for activity and stability (Karim *et al.*, 2010).

Structural similarities

As expectable, structural similarity searches identified mammalian MMP CDs as the closest relatives of Kly18 (Table 3). Structurally analysed MMP CDs comprise only mammalian representatives of MMP-1 to MMP-3, MMP-7 to MMP-14, MMP-16 and MMP-20, which all span 160–173 residues (Maskos, 2005; Rao, 2005; Tallant *et al.*, 2010b). Other MPs were identified as more distant relatives, among them serralysins, snapalysin, astacins, adamalysins/ADAMs, ulilysin and leishmanolysin. This is consistent with the general similarity observed across these families, which justifies that they are grouped in the metzincin clan of MPs (Gomis-Rüth, 2003; 2009). Superimposition of the structures of Kly18 onto several MMPs revealed similar chain traces for the overall CDs, including all regular secondary structure elements, the catalytic residues and the Met-turn (Fig. 2A). No disulphide bonds

Fig. 2. Structural, evolutionary and sequence relationships of Kly18.

A. Superposition in stereo of the α A chain traces of Kly18 (yellow) and the catalytic domains of MMP-1, -3, -7, -8, -9 (without fibronectin-type II insertions), -10, -11, -12, -13 and -16 (all in cyan). Kly18 segments showing significant deviations from the common chain trace are framed and labelled.

B. Twenty-four selected catalytic domain sequences of described and potential MMPs were subjected to phylogenetic analysis in relation with Kly18. Chosen sequences included: sequences of hitherto uncharacterized bacterial proteins of *Synechococcus* sp. (UniProt B4WJ70), *Rhodopirellula baltica* (Q7UJQ3), *Stigmatella aurantiaca* DW4/3-1 (Q098Y1), *Solibacter usitatus* strain Ellin6076 (Q029Q8), *Methylobacterium nodulans* strain ORS2060/LMG 21967 (B8IRC9), *Methylobacterium* sp. strain 4-46 (B0UK88), *Oceanibulbus indolifex* HEL-45 (A9EGF2), *Roseovarius* sp. 217 (A3W4J0); insect sequences [designated as either Dm1-MMP-like or Dm2-MMP-like on the basis of sequence comparison with the two *Drosophila melanogaster* proteins, the two only biochemically characterized insect forms (Llano *et al.*, 2000; 2002)] of *Galleria mellonella* (wax moth; Q0VJ89), *Drosophila grimshawi* (Hawaiian fruitfly; B4JVX6), *Drosophila virilis* (fruitfly; B4LJA4), *Drosophila ananassae* (fruitfly; B3MCI2), *Bombyx mori* (silk moth; B2CI49), *Aedes aegypti* (yellow fever mosquito; Q17FG9), *Anopheles gambiae* (African malaria mosquito; Q5TPJ3) and *Culex quinquefasciatus* (Southern house mosquito; B0W3F6); and vertebrate MMPs, i.e. *Homo sapiens* MMP-16 (MT3-MMP) (P51512), *H. sapiens* MMP-24 (MT5-MMP) (Q9Y5R2), *H. sapiens* MMP-20 (enamelysin) (O60882), *H. sapiens* MMP-13 (collagenase-3) (P45452), *H. sapiens* MMP-11 (stromelysin-3) (P24347), *Gallus gallus* (chicken) MMP-13 (O93363) and *Bos taurus* (bovine) MMP-13 (O77656). Sequences and tree branches are coloured red (bacteria), green (winged insects) and blue (vertebrates).

C. Sequence alignment of Kly18 with selected catalytic domains depicted in (B). The abbreviations are: Tf, *T. forsythia*; Ae, *A. aegypti*; Ag, *A. gambiae*; Hs, *H. sapiens*; Gm, *G. mellonella*; and Dv, *D. virilis*. The UniProt sequence access code, as well as the percentage of sequence identity with Kly18 and the common sequence (both in parenthesis), is shown for each sequence in the lower sequence block. The numbering and the regular secondary structure elements (strands β I– β V and helices α A– α C) correspond to the Kly18 structure.

are present in any structure. In addition, Kly18 possesses most of the structural details characteristic of MMP CDs that distinguish them from other metzincins: (i) the residue downstream of the third zinc-binding histidine is always a serine, and it hydrogen bonds the aspartate residue in helix αC that binds the N-terminus in physiologically relevant 'superactive' MMP variants (Reinemer *et al.*, 1994; Tallant *et al.*, 2010b), (ii) after this conserved serine, the downstream chain trace, which leads to the Met-turn and is characteristic for MMPs, and the latter turn itself, are perfectly superimposable, (iii) the Met-turn is stabilized through hydrogen bonds of its main-chain with a conserved aspartate residue immediately downstream of the aforementioned one within helix αC , (iv) the characteristic loop structures found in Kly18, the S-loop, the bulge-edge segment, the S_1' -wall forming segment – which includes the sequence Pro175–Tyr–Tyr177 – and the specificity loop, are essential hallmarks of mammalian MMP CDs (Tallant *et al.*, 2010b), (v) the S_1' pocket is hydrophobic and designed to accommodate bulky hydrophobic side-chains, (vi) in addition, Kly18 accomplishes with the structural requirements of another characteristic feature of the active-site cleft of MMPs, i.e. the selectivity for proline in position P_3 of a substrate, through an S_3 pocket framed by His117 and Phe119 of strand βIV and Tyr106 of the S-loop, (vii) the first part of the S-loop includes an essential structural zinc site (Williams and Murray, 1994) formed by residues contained in conserved sequence stretches equivalent to Pro107–His–Asp–Gly–(Xxx)₃–Phe–Asp–Gly110 of Kly18 (Fig. 2C), and finally (viii) the sequences found in Kly18 within the upper-rim strand βIV , Leu115–Ala–His–Ala–Phe–Xxx–Pro121, and within strand βV , His133–Phe–Asp135, are likewise conserved among MMPs.

There are, however, minor deviations from an average MMP structure in Kly18, but they are found mostly in regions that are also variable among mammalian structures (Fig. 2A). Kly18 does not have any of the three characteristic calcium binding sites featured by the bulge-edge segment *plus* the first part of $L\beta V\alpha B$, by $L\beta IV\beta V$ and by $L\beta I\alpha A/L\beta V\alpha B$, respectively, in mammalian MMPs. This is particularly noteworthy because Kly18 has most of the conserved residues participating in the first site (Fig. 2C), and this could explain the flexibility of the bulge-edge segment in the bacterial enzyme (see above and legend to Fig. 1). Here it must be mentioned that mammalian MMPs are variable as to these calcium sites: they may comprise just the bulge-edge segment site (MMP-11), two sites (MMP-7, MMP-8, MMP-16 and MMP-20) or three sites (MMP-3, MMP-9 and MMP-10). Some proteins have even been solved with either two or three calcium ions (MMP-1, MMP-2, MMP-12 and MMP-13). Further differences in length and conformation between Kly18 and mammalian MMP CDs are found within the specificity

loop. This element, however, also deviates widely among the latter structures and it results in hydrophobic S_1' pockets of varying depth, i.e. with the capacity to allocate P_1' chains of disparate size. The same scenario is observed at $L\beta V\alpha B$, which is highly variable among MMPs and contributes to substrate specificity. $L\beta I\alpha A$ also displays a slight variation, probably owing to the flexible nature of this segment in Kly18. Finally, $L\beta IV\beta V$ is unique to Kly18 due to a three-residue insertion, which gives rise to a stretch of sequence Pro121–Pro₂–Ala–Gly–Gly126 (Fig. 2C).

In the light of all these findings, we conclude that Kly18 is a true MMP CD, which like all its mammalian counterparts shows small hallmark deviations from the common average structure.

Phylogenetic and sequence similarity analyses of Kly18

Previous and current sequence similarity searches revealed that Kly18 is close to animal MMP sequences despite belonging to a bacterium from the phylum *Bacteroidetes* (Karim *et al.*, 2010). Highest sequence identities (> 40%) and shortest evolutionary distances are observed with respect to potential proteins from mosquitoes (order *Diptera*, family *Culicidae*) within the superorder *Endopterygota* (Fig. 2B and C). The former include the transmitting insect vectors of the parasite that causes malaria (*Anopheles gambiae*); of the viruses that cause dengue fever, Chikungunya and yellow fever (*Aedes aegypti*); and of West Nile virus (*Culex quinquefasciatus*). All these mosquitoes have intimate contact with mammalian hosts, on whose blood they feed, mostly in developing countries, regions that also present the highest incidence of odontopathogenic bacterial infections (Axelsson *et al.*, 2002). Next in evolutionary distance are vertebrate MMPs (e.g. MMP-11, -13, -16 and -20), sequences from other endopterygota including fruitflies (order *Diptera*, family *Drosophilidae*), and moths and butterflies (order *Lepidoptera*). Clearly farthest away are several putative bacterial proteins from cyanobacteria, planctomycetes, acidobacteria and proteobacteria (Fig. 2B).

Evolutionary implications

Horizontal or lateral gene transfer (HGT) entails shuffling of genes – xenologues – between different species and it contributes to the diversification of life by complementing the Darwinian tree-based pathway that is based on reproductive isolation between species (Gogarten, 1994; Koonin *et al.*, 2001). Such xenologues are defined as homologues of genes that have evolved within a donor species before being transferred to a receptor, i.e. they have not been inherited from the last universal common

ancestor and are thus incongruent with the species tree. HGT is only successful if the co-opted gene becomes incorporated into the genome of the acceptor and if selective advantage is conferred. Mechanisms of HGT include transformation, transduction and conjugation of nucleic acid chains through several pathways that may include viral and bacteriophage vectors. The direction of HGT probably followed the 'out of Africa' principle, which pinpoints the most crowded taxon as the most probable source (Cavalli-Sforza, 1998; Koonin *et al.*, 2001). HGT is widespread and well documented within and between bacteria and archaea, as exemplified by the sharing of antibiotic resistance plasmids, often across species barriers (Macrina and Archer, 1993). In contrast, the transfer from bacteria to eukaryotic cells is less common, and examples include the interaction between *Agrobacterium tumefaciens* and plant cells, which induces Crown-Gall disease in the latter (Sprague, 1991; Gomis-Rüth *et al.*, 2004). Finally, HGT from eukaryotes to bacteria is of particular importance given the possible roles of transferred genes in bacterial pathogenicity (Ochman *et al.*, 2000; Koonin *et al.*, 2001). This transfer is very rare and restricted to few cases proposed on the basis of phylogenetic and comparative genome analyses – 'lateral genomics' (Doolittle, 1999; Koonin *et al.*, 2001). Direct proof of such gene shuffling is unavailable for there is no footprint of such evolutionary events. For this reason, additional evidence as provided by structural relatedness may help to uncover new potential cases of xenology.

Karilysin is a unique protein of *T. forsythia* and its natural habitat is the gingival crevice of humans, where it contributes to periodontal disease. Its CD, Kly18, shows closest structural similarity and comparable S₁' subsite specificity to mammalian MMPs and shorter evolutionary distance to MMP sequences of winged insect and mammals than to other bacterial yet uncharacterized sequences. The MMP family is currently described only in animals and plants, with 23 paralogues found in humans. The existence of sequences in mosquitoes related to the only two insect forms present in *Drosophila melanogaster*, Dm1-MMP and Dm2-MMP (Llano *et al.*, 2000; 2002), and to sequences found in other fruitflies, moths and butterflies, strongly suggests that they likely correspond to expressed functional proteins and that MMPs within winged insects result from Darwinian evolution, as is the case for the presence of MMPs within vertebrates. Coincidentally, winged-insect MMPs follow the general domain architecture of mammalian MMPs, which is found in all MMPs studied, except for a protein from the nematode *Gnathostoma spinigerum*, and plant MMPs. Notwithstanding, the latter possess at least a pro-domain with a characteristic cysteine possibly participating in latency as in animal forms.

In contrast, karilysin has only a short pro-peptide between the signal peptide and the beginning of the mature CD and a C-terminal domain, and neither bears any relationship to any other characterized protein. The pro-peptide lacks the characteristic cysteine residue (Springman *et al.*, 1990; Van Wart and Birkedal-Hansen, 1990; Gonzales *et al.*, 2004; Rosenblum *et al.*, 2007), but contains an aspartate that could contribute to latency maintenance as in proastacin (Guevara *et al.*, 2010). In addition, no gene potentially encoding a Kly18 relative was found either in the other two red complex constituents, *P. gingivalis* and *T. denticola*, or in any bacterium of the phylum *Bacteroidetes*. Taken together, these facts strongly suggest that Kly18 is an MMP xenologue that has been co-opted from an external source. Likely donor candidates are mammals or mosquitoes feeding on human blood, due to the intimate coexistence between *T. forsythia* and the human blood-irrigated gingival crevice. This hypothesis is further backed by the existence of a sequence within the genome of invertebrate iridescent virus 3 (UniProt Q196W5), which infects mosquitoes of the genus *Aedes* and *Culex*, that shows 40% sequence identity to Kly18 within 146 common residues. This would suggest participation of a viral vector in the HGT. Subsequently, Kly18 would have evolved in a bacterial environment, where it was furnished with unique flanking domains to give rise to full-length karilysin. This proposed example of HGT entailing the shuffling of a metazoan MMP to a human pathogenic bacterium, which involves distinct domains of life, is both a short circuit of Darwinian evolution and a testimony to the compatibility of proteins and cellular mechanisms despite evolutionary divergence over billions of years.

Experimental procedures

Protein production and purification and endopeptidolytic assays

Karilysin CD (Kly18) was obtained by autolysis of full-length karilysin, which was produced by recombinant overexpression as a fusion construct with glutathione-S-transferase in *Escherichia coli* BL21 cells, as described elsewhere (Karim *et al.*, 2010). Activity of Kly18 in front of the fluorogenic peptide substrates, Mca-Arg-Pro-Lys-Pro-Val-Glu-Nva-Trp-Arg-Lys(Dnp)-NH₂ (Mca, 7-methoxycoumarin-4-acetate; Nva, norvaline; Dnp, dinitrophenyl) and Mca-Pro-Leu-Gly-Leu-Dap(Dnp)-Ala-Arg-NH₂ (Dap, L-diaminopropionyl; both from Bachem, with $\lambda_{\text{EX}} = 328$ nm and $\lambda_{\text{EM}} = 393$ nm), was assayed at 37°C in 100 mM Tris-HCl, 5 mM calcium chloride, pH 8.0 by using an enzyme concentration of 25 nM for the former substrate and 50 nM for the latter substrate. Fluorescence was recorded for 12 h in a fluorimetric plate reader (Tecan Infinite M200). Substrate concentrations varied between 1 and 8 μ M to avoid quenching effects due to high substrate concentra-

tions (Knight *et al.*, 1992; Nagase *et al.*, 1994; Fields, 2001). The k_{cat}/K_m parameter was estimated using the minimum concentration that permitted to reliably determine the associated velocity according to equation, $[E]_{\text{min}} = v_{\text{min}}/(k_{\text{cat}}/K_m \times [S])$, which is a valid approximation when $[S] \ll K_m$ (Knight *et al.*, 1992).

Crystallization and structure analysis

Crystallization assays were performed by the sitting-drop vapour diffusion method. Reservoir solutions were prepared by a Tecan robot and crystallization drops were dispensed on 96×2 -well MRC plates (Innovadyne) by a Cartesian (Genomic Solutions) nanodrop robot at the High-Throughput Crystallography Platform at Barcelona Science Park. Crystals suitable for structure analysis were obtained for unbound Kly18 in a Bruker steady-temperature crystal farm at 20°C from drops containing 100 nl of protein solution (at 9 mg ml⁻¹ in 5 mM Tris-HCl, pH 8.0, 0.02% sodium azide) and 100 nl of 45% 2-methyl-2,4-pentanediol, 0.2 M ammonium acetate, 0.1 M Tris-HCl, pH 8.5 as reservoir solution. Crystals of magnesium-bound Kly18 were obtained at 4°C by mixing protein solution pre-incubated with calcium (at 7.3 mg ml⁻¹ in 5 mM Tris-HCl, pH 8.0, 5 mM calcium chloride, 0.02% sodium azide) and 14% sodium polyacrylate 5100, 20 mM magnesium chloride, 0.1 M HEPES, pH 7.5 in a volume ratio of 1:3. The latter conditions were scaled up to the microlitre range with 24-well Cryschem crystallization dishes (Hampton Research). Crystals were cryo-protected with 50% 2-methyl-2,4-pentanediol, 15% glycerol, 0.2 M ammonium acetate, 0.1 M Tris-HCl, pH 8.5 (unbound Kly18) and 30% sodium polyacrylate 5100, 20% glycerol, 0.02 M magnesium chloride, 0.1 M HEPES, pH 7.5 (magnesium-bound Kly18). Complete diffraction data sets were collected for unbound and magnesium-bound Kly18 from liquid-N₂ flash-cryo-cooled crystals at 100 K (provided by an Oxford Cryosystems 700 series cryostream) at beam lines ID23-2 (equipped with a MarMosaic 225 CCD detector) and ID29 (with an ADSC 315R CCD detector), respectively, of the European Synchrotron Radiation Facility (ESRF, Grenoble, France) within the Block Allocation Group 'BAG Barcelona'. Crystals were either monoclinic (unbound Kly18) or tetragonal (magnesium-bound Kly18), with two or one molecules per asymmetric unit respectively. Diffraction data were integrated, scaled, merged and reduced with programs XDS (Kabsch, 2001) and SCALA (Evans, 2006) within the CCP4 suite of programs (CCP4, 1994) (see Table 2).

The structure of unbound Kly18 was solved by Patterson search with program PHASER (McCoy *et al.*, 2007) and the co-ordinates of human MMP-8 [Protein Data Bank access code (PDB) 1MNC] as searching model. These calculations rendered two unique solutions at, respectively, 328.1, 106.1, 166.2, -0.27, 1.00, -0.55; and 237.5, 143.7, 46.9, -0.28, 0.99, 0.01 (Eulerian angles α , β and γ , and x , y , z in fractional cell co-ordinates). The global value of log-likelihood gain was 384. The correspondingly rotated and translated searching models were used to compute initial phases, which were subjected to a run with program ARP/WARP (Perrakis *et al.*, 2001). This process significantly improved the electron density map and enabled straightforward tracing of the entire polypeptide chain on a Silicon Graphics Octane2 workstation

with the TURBO-Frodo program (Carranza *et al.*, 1999). Subsequent crystallographic refinement with REFMAC5 (Murshudov *et al.*, 1997), which included TLS refinement, alternated with manual model building until completion of the model. The latter comprised residues Tyr35–Phe200 (numbering according to UniProt database entry D0EM77) and two zinc ions for each of the two chains, A and B. In addition, 115 solvent molecules, a tetrapeptide of tentative sequence Ala–Phe–Thr–Ser, and a Tris molecule from the crystallization conditions were assigned. The structure of magnesium-bound Kly18 was solved with program PHASER by using the protein chain co-ordinates of Kly18 as searching model. A single solution was obtained at 43.0, 89.6, 89.8, 0.01, 0.01, 0.01 with a global log-likelihood gain of 2135. Phase improvement, model building and refinement proceeded as stated above. The final model spanned residues Tyr35–Ser201, two zinc ions, a tripeptide of sequence Ala–Phe–Thr, tentatively assigned chloride and magnesium ions, and 196 solvent molecules (see Table 2).

Phylogenetic analysis

Selected MMP sequences obtained from sequence similarity searches were pairwise aligned with Kly18 to delimit the CDs of the former with program MULTALIN (Corpet, 1988). These fragments were thereafter aligned altogether with Kly18 with MULTALIN, which computes parameters for a phylogenetic tree in rfd format. The latter was manually converted to dnd format and plotted as a circular tree with PHYLIP DRAWGRAM at <http://mobyle.pasteur.fr/cgi-bin/portal.py?form=drawgram>.

Miscellaneous

Figures were prepared with SETOR (Evans, 1993), GRASP (Nicholls *et al.*, 1993) and TURBO-Frodo. Structure similarities were determined with DALI (Holm *et al.*, 2006), and sequence alignments were performed with MULTALIN. Experimental model validation was performed with MOLPROBITY (Davis *et al.*, 2007) and the WHATCHECK routine of WHATIF (Vriend, 1990). The final co-ordinates of unbound and magnesium-bound Kly18 are available from the PDB at <http://www.pdb.org> (access codes 2XS3 and 2XS4).

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