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Crystal Structure and Activity Studies of the C11 Cysteine Peptidase from *Parabacteroides merdae* in the Human Gut Microbiome.

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ABSTRACT

Clan CD cysteine peptidases, a structurally related group of peptidases that include mammalian caspases, exhibit a wide range of important functions, along with a variety of activation specificities and mechanisms. However, for the clostripain family (denoted C11), little is currently known. Here, we describe the first crystal structure of a C11 protein from the human gut bacterium, Parabacteroides merdae (PmC11), determined to 1.7 Å resolution. PmC11 is a monomeric cysteine peptidase that comprises an extended caspase-like $\alpha/\beta/\alpha$ sandwich and an unusual Cterminal domain. It shares core structural elements with clan CD cysteine peptidases but

otherwise structurally differs from the other families in the clan. These studies also revealed a well-ordered break in the polypeptide chain at K147, resulting in a large conformational rearrangement close to the active Biochemical and kinetic analysis revealed K147 to be an intramolecular processing site at which cleavage is required for full activation of the enzyme, suggesting an auto-inhibitory mechanism for self-preservation. PmC11 has an acidic binding pocket and a preference for basic substrates, and accepts both Arg and Lys substrates and does not require Ca²⁺ for activity. Collectively, these data provide insights into the mechanism and activity of PmC11 and a detailed

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framework for studies on C11 peptidases from

other phylogenetic kingdoms.

INTRODUCTION

Cysteine peptidases play crucial roles in the virulence of bacterial and other eukaryotic pathogens. In the MEROPS peptidase database (1), clan CD contains groups (or families) of cysteine peptidases that share some highly conserved structural elements (2). Clan CD families are typically described using the name of their archetypal, or founding, member and also given an identification number preceded by a 'C', to denote cysteine peptidase. Whilst seven families (C14 is additionally split into three subfamilies) have been described for this clan, crystal structures have only been determined from legumain (C13)(3),four: caspase (C14a)(4). paracaspase (C14b(P)(5),metacaspase (C14b(M)(6), gingipain (C25)(7) and the cysteine peptidase domain (CPD) of various toxins (C80)(8). No structural information is available for clostripain (C11), separase (C50) or PrtH-peptidase (C85).

Clan CD enzymes have a highly conserved His/Cys catalytic dyad and exhibit strict specificity for the P₁ residue of their substrates. However, despite these similarities, clan CD forms a functionally diverse group of enzymes: the overall structural diversity between (and at times within) the various families provides these peptidases with a wide variety of substrate specificities and activation mechanisms. Several members are initially expressed as proenzymes, demonstrating self-inhibition prior to full activation (2).

The archetypal and arguably most notable family in the clan is that of the mammalian caspases (C14a), although clan CD members are distributed throughout the entire phylogenetic kingdom and are often required in fundamental biological processes (2). Interestingly, little is known about the structure or function of the C11 proteins, despite their wide spread distribution (1) and its archetypal member, clostripain from Clostridium histolyticum, being first reported in the literature in 1938 (9). Clostripain has been described as an arginine-specific peptidase with a requirement for Ca²⁺ (10) and loss of an internal nonapeptide for full activation; lack of structural information on the family appears to have prohibited further investigation.

As part of an ongoing project to characterize commensal bacteria in the microbiome that inhabit the human gut, the structure of C11 peptidase, PmC11, from **Parabacteroides** merdae was determined using the JCSG HTP structural biology pipeline (11) and now has analysed provide been to the structure/function correlation for C11 peptidase.

EXPERIMENTAL PROCEDURES

Cloning, expression, purification, crystallization, and structure determination of PmC11 were carried out using standard JCSG (http://www.jcsg.org) protocols (11) as follows.

Cloning - Clones were generated using the Polymerase Incomplete Primer Extension (PIPE) cloning method (12) The gene encoding PmC11 (SP5111E) was amplified by polymerase chain reaction (PCR) from Parabacteroides merdae genomic DNA using PfuTurbo DNA polymerase (Stratagene), using I-PIPE primers that included sequences for the predicted 5' and 3' ends (shown below). The expression vector, pSpeedET, which encodes an amino-terminal tobacco etch virus (TEV) protease-cleavable expression and purification tag (MGSDKIHHHHHHHENLYFQ/G), was **PCR** amplified with V-PIPE (Vector) primers. V-PIPE and I-PIPE PCR products were mixed to anneal amplified **DNA** fragments together. the Escherichia coli GeneHogs (Invitrogen) competent cells were transformed with the I-PIPE / V-PIPE mixture and dispensed on selective LBagar plates. The cloning junctions were confirmed by DNA sequencing. The plasmid encoding the full-length protein was deposited in the PSI:Biology Materials Repository at the DNASU plasmid repository (PmCD00547516). structure determination, using the PIPE method for soluble protein expression, the gene segment encoding residues M1-N22 was deleted since these residues were predicted to correspond to a signal peptide using SignalP (13).

Protein Expression and Selenomethionine-incorporation (SeMet) – The expression plasmid for the truncated PmC11construct was transformed into E. coli GeneHogs competent cells and grown in minimal media

supplemented with SeMet and 30 μg mL⁻¹ kanamycin at 37 °C using a GNF fermentor (14). A methionine auxotrophic strain was not required as SeMet is incorporated via the inhibition of methionine biosynthesis (15,16). Protein expression was induced using 0.1% (w/v) L-arabinose and the cells left to grow for a further 3 h at 37 °C. At the end of the cell culture, lysozyme was added to all samples to a final concentration of 250 μg mL⁻¹ and the cells harvested and stored at -20 °C, until required. Primers used in this section are as follows:

I-PIPE(forward):

CTGTACTTCCAGGGCGAGACTCCGGAACCCCGGACAACCCGC,

I-PIPE (reverse):

AATTAAGTCGCGTTATTCATAAACTGCC TTATACCAGCCGAC,

V-PIPE (forward):

TAACGCGACTTAATTAACTCGTTTAAAC GGTCTCCAGC,

V-PIPE (reverse):

GCCCTGGAAGTACAGGTTTTCGTGATGATGATGATGATGAT.

Protein purification for crystallisation. Cells were resuspended, homogenized and lysed by sonication in 40 mM Tris (pH 8.0), 300 mM NaCl, 10 mM imidazole and 1 mM Tris (2carboxyethyl) phosphine hydrochloride (TCEP) (Lysis Buffer 1) containing 0.4 mM MgSO₄ and 1 μL of 250 U μL⁻¹ Benzonase (Sigma). The cell lysate was then clarified by centrifugation (32,500 g for 25 min at 4 °C) before being passed over Ni²⁺-chelating resin equilibrated in Lysis Buffer 1 and washed in the same buffer supplemented with 40 mM imidazole and 10% (v/v) glycerol. The protein was subsequently eluted in 20 mM Tris (pH 8.0), 150 mM NaCl, 10% (v/v) glycerol, 1 mM TCEP and 300 mM imidazole and the fractions containing the protein pooled.

To remove the His-tag, PmC11 exchanged into 20 mM Tris (pH 8.0), 150 mM NaCl, 30 mM imidazole and 1 mM TCEP using a PD-10 column (GE Healthcare), followed by incubation with 1 mg of His-tagged TEV protease per 15 mg of protein for 2 h at room temperature and subsequent incubation at 4°C. The sample was centrifuged to remove any precipitated material (13,000 g for 10 min at 4°C) and the supernatant loaded onto Ni²⁺-chelating resin equilibrated with 20 mM Tris (pH 8.0), 150 mM NaCl, 30 mM

imidazole and 1 mM TCEP and washed with the same buffer. The flow-through and wash fractions were collected and concentrated to 13.3 mg mL⁻¹ using Amicon Ultra-15 5K centrifugal concentrators (Millipore).

Crystallisation and Data Collection. PmC11 was crystallized using the nanodroplet diffusion method using standard **JCSG** crystallization protocols (11). Drops were comprised of 200 nl protein solution mixed with 200 nl crystallization solution in 96-well sittingdrop plates, equilibrated against a 50 µl reservoir. Crystals of PmC11 were grown at 4°C in mother liquor consisting of 0.2 M NH₄H₂PO₄, 20% PEG-3350 (JCSG Core Suite I). Crystals were flash cooled in liquid nitrogen using 10% ethylene glycol as a cryoprotectant prior to data collection and initial screening for diffraction was carried out using the Stanford Automated Mounting system (SAM) (17) at the Stanford Synchrotron Radiation Lightsource (SSRL, Menlo Park, CA). Single wavelength anomalous dispersion (SAD) diffraction data were collected using a wavelength of 0.9793 Å, at the Advanced Light Source (ALS, beamline 8.2.2, Berkeley, CA) on an ADSC Quantum 315 CCD detector. The data were indexed and integrated with XDS (18) and scaled using XSCALE (18). The diffraction data were indexed in space group $P2_1$ with a = 39.11, b = 108.68, c = 77.97 Å and β = 94.32°. The unit cell contained two molecules in the asymmetric unit (ASU) resulting in a solvent content of 39% (Matthews' coefficient (Vm) of 2.4 Å³ Da⁻¹).

Structure Determination—The structure was determined by the SAD method using an X-ray wavelength corresponding to the peak of the selenium K edge. Initial phases were derived using the autoSHARP interface (19), which included density modification with SOLOMON (20). Good quality electron density was obtained at 1.7 Å resolution, allowing an initial model to be obtained by automated model building with ARP/wARP (21).Model completion and refinement were iteratively performed with COOT (22) and REFMAC (23,24) to produce a final model with an R_{cryst} and R_{free} of 14.3% and 17.5%, respectively. The refinement included experimental restraints in the form of Hendrickson-Lattman coefficients, TLS refinement with one TLS group per molecule in the ASU, and NCS restraints. The refined structure contains residues

24-375 and 28-375 for the two molecules in the crystallographic ASU. Structural validation was carried using the JCSG Quality Control Server (http://smb.slac.stanford.edu/jcsg/QC/) analyses both the coordinates and data using a variety of structural validation tools to confirm the stereochemical quality of the model (ADIT (25), MOLPROBITY (26), and WHATIF 5.0 (27) and agreement between model and data (SGCHECK (28) and RESOLVE (29). All of the main-chain torsion angles were in the allowed regions of the Ramachandran plot and the MolProbity overall clash score for the structure was 2.09 (within the 99th percentile for its resolution). The atomic coordinates and structure factors for PmC11 have been deposited in the Protein Data Bank (PDB, www.wwpdb.org) with the accession code 3UWS. Data collection, model and refinement statistics are reported in Table 1.

Structural Analysis—The primary sequence alignment with assigned secondary structure was prepared using CLUSTAL OMEGA (30) and ALINE (31). The topology diagram was produced with TOPDRAW (32) and all 3D structural figures were prepared with PyMol (33) with the electrostatic surface potential with additionally calculated **APBS** (18)contoured at ±5 kT/e. Architectural comparisons with known structures revealed that PmC11 was most structurally similar to caspase-7, gingipain-K and legumain (PBD IDs 4hq0, 4tkx, and 4awb, respectively). The statistical significance of the structural alignment between PmC11 and both caspase-7 and gingipain-K was reported to be equivalent (Z-score of 9.2) with legumain giving a very similar result (Z-score of 9.1). Of note, the β -strand topology of the CDP domains of Clostridium difficile toxin B (family C80; TcdB; PDB ID 3pee) is identical to that observed in the PmC11 β -sheet but the Z-score from DaliLite was notably less at 7.6. It is possible that the PmC11 structure is more closely related to the C80 family than other families in clan CD, and appear to reside on the same branch of the phylogenetic tree based on structure (2).

Protein production for biochemical assays—The PmCD00547516 plasmid described above was obtained from the PSI:Biology Materials Repository and used to generate a cleavage site mutant PmC11^{K147A} and an active-site mutant

PmC11^{C179A} using the QuikChange Site-directed Mutagenesis kit (Stratagene) as per the manufacturer's instructions using the following primers:

K147A mutant (forward):

CAGAATAAGCTGGCAGCGTTCGGACAGG ACG.

K147A mutant (reverse):

 $\begin{array}{c} \mathsf{CGTCCTGTCCGAACGCTGCCAGCTTATTC} \\ \mathsf{TG} \end{array}$

C179A mutant (forward):

CCTGTTCGATGCCGCCTACATGGCAAGC: C179A mutant (reverse):

GCTTGCCATGTAGGCGGCATCGAACAGG. The expression plasmids containing PmC11 were transformed into *E. coli* BL21 Star (DE3) and grown in Luria-Bertani media containing 30 μg mL⁻¹ kanamycin at 37 °C until an optical density (600 nm) of ~0.6 was reached. L-arabinose was added to a final concentration of 0.2% (w/v) and the cells incubated overnight at 25 °C.

Compared to the protein production for crystallography, a slightly modified purification protocol was employed. Initially, the cells were resuspended in 20 mM sodium phosphate (pH 7.5), 150 mM NaCl (Lysis Buffer 2) containing an EDTA-free protease inhibitor cocktail (cOmplete, Roche). Cells were disrupted by three passages (15 KPSI) through a One-Shot cell disruptor (Constant Systems) followed by centrifugation at 20,000 g for 20 min at 4°C. The supernatant was collected and sterile-filtered (0.2 um) before being applied to a 5 mL HisTrap HP column (GE Healthcare) equilibrated in Lysis Buffer 2 containing 25 mM imidazole, and the protein was eluted in the same buffer containing 250 mM imidazole. The peak fractions were pooled and buffer exchanged into the assay buffer (20 mM Tris, 150 mM NaCl, pH 8.0) using a PD-10 column. When required, purified PmC11 was concentrated using Vivaspin 2 30 K centrifugal concentrators (Sartorius). Protein concentration was routinely measured using Bradford's reagent (BioRad) with a BSA standard.

Fluorogenic substrate activity assays—The release of the fluorescent group AMC (7-Amino 4-Methyl Coumarin) from potential peptide substrates was used to assess the activity of PmC11. Peptidase activity was tested using 20 μ g PmC11 and 100 μ M substrate (unless otherwise stated) in assay buffer to a final reaction volume of 200 μ L and all samples were

incubated (without substrate) at 37 °C for 16 h prior to carrying out the assay. The substrate and plate reader were brought to 37 °C for 20 min prior to the addition of the PmC11 and samples prepared without PmC11 were used as blanks (negative controls). The curves were plotted using the blank-corrected fluorescence units against the time of acquisition (in min). The assays were carried out in black 96-well flat-bottomed plates (Greiner). AMC fluorescence was measured using a PHERAstar FS plate reader (BMG Labtech) with excitation and emission wavelengths of 355 nm and of 460 nm, respectively.

To investigate the substrate specificity of PmC11, substrates Z-GGR-AMC, Bz-R-AMC, Z-GP-AMC, Z-HGP-AMC, Ac-DEVD-AMC (all Bachem), BOC-VLK-AMC and BOC-K-AMC (both PeptaNova) were prepared at 100 mM in 100% DMSO. The amount of AMC (μmoles) released was calculated by generating an AMC standard curve (as described (34)) and the specific activity of PmC11 was calculated as pmoles of AMC released per min per mg of the protein preparation.

The reaction rates (Vmax) and Km values were determined for mutants $PmC11^{K147A}$ and $PmC11^{C179A}$ by carrying out the activity assay at varying concentrations of Bz-R-AMC between 0 and 600 μ M. The blank-corrected relative fluorescence units were plotted against time (min) with $\Delta FU/T$ giving the reaction rate. The Km and Vmax of PmC11 and PmC11 K147A against an R-AMC substrate were determined from the Lineweaver-Burk plot as described (34), calculated using GraphPad Prism6 software. All experiments were carried out in triplicate \pm SD.

Effect of VRPR-FMK on PmC11— To test the effect of the inhibitor on the activity of PmC11, 25 μM of Z-VRPR-FMK (100mM stock in 100% DMSO, Enzo Life Sciences), 20 μg PmC11, 100 μM R-AMC substrate, 1 mM EGTA were prepared in the assay buffer and the activity assay carried out as described above. A gel-shift assay, to observe Z-VRPR-FMK binding to PmC11, was also set up using 20 μg PmC11, 25 μM inhibitor, 1 mM EGTA in assay buffer. The reactions were incubated at 37 °C for 20 min before being stopped by the addition SDS/PAGE sample buffer. Samples were analysed by loading the reaction mixture on a 10% NuPAGE Bis-Tris gel using MES buffer.

Effect of cations on PmC11—The enzyme activity of PmC11 was tested in presence of

various divalent cations: Mg^{2+} , Ca^{2+} , Mn^{2+} , Co^{2+} , Fe^{2+} , Zn^{2+} and Cu^{2+} . The final concentration of the salts ($MgSO_4$, $CaCl_2$, $MnCl_2$, $CoCl_2$, $FeSO_4$, $ZnCl_2$, and $CuSO_4$) was 1 mM and a control was set up without divalent ions but with addition of 1 mM EGTA as the control. The assay was set up using 20 mg PmC11, 1 mM salts, 100 μ M R-AMC substrate, and the assay buffer, and incubated at 37°C for 16 h. The activity assay was carried out as described above.

Size exclusion chromatography— Affinity-purified PmC11 was loaded onto a HiLoad 16/60 Superdex 200 gel filtration column (GE Healthcare) equilibrated in the assay buffer. The apparent molecular weight of PmC11 was determined from calibration curves based on protein standards of known molecular weights.

Auto-processing profile of PmC11—Auto-processing of PmC11 was evaluated by incubating the enzyme at 37°C and removing samples at one hour intervals for from 0 to 16 hours and placing into SDS/PAGE loading buffer to stop the processing. Samples were then analysed on a 4-12% NuPAGE (Thermo Fischer) Novex Bis-Tris gel run in MES buffer.

Auto-processing cleavage site analysis— To investigate whether processing is a result of intra or inter-molecular cleavage the PmC11^{C179A} incubated mutant was with increasing concentrations of activated PmC11 (0, 0.1, 0.2, 0.5, 1, 2 and 5 µg). The final assay volume was 40 µL and the proteins were incubated at 37°C for 16 h in the PmC11 assay buffer. To stop the reaction, NuPAGE sample buffer was added to the protein samples and 20 µL was analysed on 10% NuPAGE Novex Bis-Tris gel using MES buffer. These studies revealed no apparent cleavage of PmC11^{C179A} by the active enzyme at low concentrations of PmC11 and that only limited cleavage was observed when the ratio of active enzyme (PmC11: PmC11^{C179A}) was increased to ~1:10 and 1:4.

RESULTS

Structure of PmC11—The crystal structure of the catalytically active form of PmC11 revealed an extended caspase-like $\alpha/\beta/\alpha$ sandwich architecture comprised of a central nine-stranded β -sheet, with an unusual C-terminal domain (CTD), starting at residue K250. A single cleavage was observed in the polypeptide chain at K147 (Figs 1A, B), where both ends of the cleavage site are fully visible and well-ordered in the electron density. The central nine-stranded

β-sheet (β1–β9) of PmC11 consists of six parallel and three anti-parallel β-strands with $4_{\uparrow}3_{\downarrow}2_{\uparrow}1_{\uparrow}5_{\uparrow}6_{\uparrow}7_{\downarrow}8_{\downarrow}9_{\uparrow}$ topology (Fig. 1A) and the overall structure includes 14 α-helices with six of these ($\alpha 1$ – $\alpha 2$ and $\alpha 4$ – $\alpha 7$) closely surrounding the β-sheet in an approximately parallel orientation. Helices $\alpha 1$, $\alpha 7$ and $\alpha 6$ are located on one side of the β -sheet with $\alpha 2$, $\alpha 4$ and $\alpha 5$ on the opposite side (Fig 1A). Helix α 3 sits at the end of the loop following β 5 (L5), just preceding the K147 cleavage site, with both L5 and α3 pointing away from the central β-sheet and towards the CTD, which starts with α 8. The structure also includes two short β-hairpins (βA– βB and $\beta D-\beta E$) and a small β -sheet ($\beta C-\beta F$), which is formed from two distinct regions of the sequence (β C precedes α 11, α 12 and β 9, while βF follows the βD-βE hairpin) in the middle of the CTD (Fig. 1B).

The CTD of PmC11 is composed of a tight helical bundle formed from helices $\alpha 8-\alpha 14$ and includes strands βC and βF , and β -hairpin βD - βE . The CTD sits entirely on one side of the enzyme interacting only with $\alpha 3$, $\alpha 5$, $\beta 9$ and the loops surrounding $\beta 8$. Of the interacting secondary structure elements (SSEs), $\alpha 5$ is perhaps the most interesting. This helix makes a total of eight hydrogen bonds with the CTD, including one salt bridge (R191–D255) and is surrounded by the CTD on one side and the main core of the enzyme on the other, acting like a linchpin holding both components together (Fig. 1C).

comparisons—PmC11 Structural expected, most structurally similar to other members of clan CD with the top hits in a search of known structures being caspase-7, gingipain-K and legumain (PBD IDs 4hq0, 4tkx, and 4awb, respectively) (Table 2). The C-terminal domain is unique to PmC11 within clan CD and structure comparisons for this domain alone does not produce any hits in the PDB (DaliLite, PDBeFold), suggesting a completely novel fold. As the archetypal and arguably most well studied member of clan CD, the caspases were as the basis to investigate structure/function relationships in PmC11, with caspase-7 as the representative member. Six of the central β -strands in PmC11 (β 1– β 2 and β 5– β8) share the same topology as the six-stranded β -sheet found in caspases, with strands β 3, β 4 and B9 located on the outside of this core structure (2) (Fig. 1B, box). H133 and C179 were found at locations structurally homologous to the caspase catalytic dyad, and other clan CD structures (2), at the C-termini of strands $\beta 5$ and $\beta 6$, respectively (Figs 1A, B and 2A). Additionally, a multiple sequence alignment of C11 proteins revealed that these residues are also highly conserved (data not shown), revealing the location of the C11 catalytic dyad.

Five of the α -helices surrounding the β -sheet of PmC11 (α 1, α 2, α 4, α 6 and α 7) are found in similar positions to the five structurally conserved helices in caspases and other members of clan CD, apart from family C80 (2). Other than its more extended β -sheet, PmC11 differs most significantly from other clan CD members at its C-terminus, where the CTD contains a further seven α -helices and four β -strands after β 8.

Auto-Processing of PmC11—Purification of recombinant PmC11 (MW = 42.6 kDa) revealed partial processing into two cleavage products of 26.4 and 16.2 kDa, related to the observed cleavage at K147 in the crystal structure (Fig. 2A). Incubation of PmC11 at 37 °C for 16 h, resulted in a fully processed enzyme that remained as an intact monomer when applied to a size-exclusion column (Fig. 2B). The single cleavage site of PmC11 at K147 is found immediately after a3, in loop L5 within the central β-sheet (Figs. 1A, B and 2A). The two ends of the cleavage site are remarkably well ordered in the crystal structure and displaced from one another by 19.5 Å (Fig. 2A). Moreover, the C-terminal side of the cleavage site resides near to the catalytic dyad with A148 being 4.5 and 5.7 Å from H133 and C179, respectively. Consequently, it appears feasible that the helix attached to K147 $(\alpha 3)$ could be responsible for steric auto-inhibition of PmC11 as K147 moves to form a covalent bond with A148. Prior to cleavage this would limit access to the active site; thus, the cleavage would be required for full activation of PmC11. To investigate this possibility, two mutant forms of the enzyme were created: PmC11^{C179A} (a catalytically inactive mutant) and PmC11K147A (a cleavage-site mutant). Initial SDS-PAGE and western blot analysis of both mutants revealed no discernible processing occurred as compared to active PmC11 (Fig. 2C). The PmC11K147A mutant enzyme had a markedly different reaction rate (Vmax) compared to WT, where the reaction velocity of PmC11 was 10 times

greater than that of PmC11^{K147A} (Fig. 2D). Taken together, these data reveal that PmC11 requires processing at K147 for optimum activity.

To investigate whether processing is a result of inter-molecular intra cleavage, the PmC11^{C179A} mutant was incubated with increasing concentrations of processed and activated PmC11. These studies revealed that there was no apparent cleavage of PmC11^{C179A} by the active enzyme at low concentrations of PmC11 and that only limited cleavage was observed when the ratio of active enzyme (PmC11: $PmC11^{C179A}$) was increased to \sim 1:10 and 1:4, with complete cleavage observed at a ratio of 1:1 (Fig. 2E). This suggests that the limited cleavage was most likely an effect of the increasing concentration of PmC11 and intramolecular cleavage. Collectively, these data suggest that a pro-form of PmC11 is autoinhibited by a section of L5 blocking access to the active site, prior to intra-molecular cleavage at K147. This cleavage subsequently allows movement of the region containing K147 and the active site to open up for substrate access.

specificity PmC11—The Substrate of autocatalytic cleavage of PmC11 at K147 (sequence KLK^A) demonstrates that the enzyme accepts substrates with Lys in the P₁ position. The substrate specificity of the enzyme was further tested using a variety of fluorogenic substrates. As expected, PmC11 showed no activity against substrates with Pro or Asp in P₁ but was active towards substrates with a basic residue in P₁ such as Benzoyl-L-Arg-4methylcoumaryl-7-amide (Bz-R-AMC). benzyloxycarbonyl-Gly-Gly-Arg-AMC GGR-AMC) and t-Butyloxycarbonyl-Val-Leu-Lys (BOC-VLK-AMC). The rate of cleavage was ~3-fold greater toward the single Arg substrate Bz-R-AMC than the other two (Fig. 2F) and, unexpectedly, PmC11 showed no activity towards BOC-K-AMC. These results confirm that PmC11 accepts substrates containing Arg or Lys in P₁ with a possible preference for Arg.

The catalytic dyad of PmC11 sits near the bottom of an open pocket on the surface of the enzyme at a conserved location in the clan CD family (2). The PmC11structure reveals that the catalytic dyad forms part of a large acidic pocket (Fig. 2G), consistent with a binding site for a basic substrate. This pocket is lined with the potential functional side chains of N50, D177

and T204 with the side chains of G134, D207 and M205 also contributing to the pocket (Fig. 2A). Interestingly, these residues are in regions that are structurally similar to those involved in the S_1 binding pockets of other clan CD members (shown in (2)).

Since PmC11 recognises basic substrates, the tetra-peptide inhibitor Z-VRPR-FMK was tested as an enzyme inhibitor and was found to inhibit both the auto-processing and activity of PmC11 (Fig. 3A). Z-VRPR-FMK was also shown to bind to the enzyme: a size-shift was observed, by SDS-PAGE analysis, in the larger processed product of PmC11 suggesting that the inhibitor bound to the active site (Fig. 3B). A structure overlav of PmC11 with the MALT1paracacaspase (MALT1-P), in complex with Z-VRPR-FMK (35), revealed that the PmC11 dyad sits in a very similar position to that of active MALT1-P and that N50, D177 and D207 superimpose well with the principal MALT1-P inhibitor binding residues (D365, D462 and E500, respectively (VRPR-FMK from MALT1-P with the corresponding PmC11 residues from the structural overlay is shown in Fig. 1D), as described in (5)). D177 is located near the catalytic cysteine and is conserved throughout the C11 family, suggesting it is the primary S_1 binding site residue. In the structure of PmC11, D207 resides on a flexible loop pointing away from the S1 binding pocket (Fig. 1D). However, this loop has been shown to be important for substrate binding in clan CD (2) and this residue could easily rotate and be involved in substrate binding in PmC11. Thus, N50, D177 and D207 are most likely responsible for the substrate specificity of PmC11. D177 is highly conserved throughout the clan CD C11 peptidases and is thought to be primarily responsible for the substrate specificity of the clan CD enzymes, as also illustrated from the proximity of these residues relative to the inhibitor Z-VRPR-FMK when PmC11 is overlayed on the MALT1-P structure (Fig. 3C).

Comparison with clostripain—Clostripain is the founding member of the C11 peptidases and contains an additional 149 residues compared to PmC11. A multiple sequence alignment revealed that most of the secondary structural elements are conserved between the two enzymes, although they are only ~23% identical (Fig 1A). Nevertheless, PmC11 may be a good model for the core structure of clostripain.

The primary structural alignment also shows that the catalytic dyad in PmC11 is structurally conserved in clostripain (36) (Fig. 1A). Unlike PmC11, clostripain has two cleavage sites (R181 and R190), which results in the removal of a nonapeptide, and is required for full activation of the enzyme (37) (highlighted in Fig. 1A). Interestingly, R190 was found to align with K147 in PmC11. In addition, the predicted primary S1-binding residue in PmC11 D177 also overlays with the residue predicted to be the P1 specificity determining residue in clostripain (38) (D229, Fig. 1A).

As studies on clostripain revealed addition of Ca²⁺ ions are required for full activation, the Ca²⁺ dependency of PmC11 was examined. Surprisingly, Ca²⁺ did not enhance PmC11 activity and, furthermore, other divalent cations, Mg²⁺, Mn²⁺, Co²⁺, Fe²⁺, Zn²⁺ and Cu²⁺, were necessary for PmC11 activity (Fig. 3D). In support of these findings, EGTA did not inhibit PmC11 suggesting that, unlike clostripain, PmC11 does not require Ca²⁺ or other divalent cations, for activity.

DISCUSSION

The crystal structure of PmC11 now provides, three-dimensional information for a member of the clostripain C11 family of cysteine peptidases. The enzyme exhibits all of the key structural elements of clan CD members, but is unusual in that it has a nine-stranded central βsheet with a novel C-terminal domain. The structural similarity of PmC11 with its nearest structural neighbours in the PDB is decidedly low, overlaying better with six-stranded caspase-7 than any of the other larger members of the clan (Table 3). The substrate specificity of PmC11 is Arg/Lys and the crystal structure revealed an acidic pocket for specific binding of such basic substrates. In addition, the structure suggested a mechanism of self-inhibition in both PmC11 and clostripain and an activation mechanism that requires auto-processing.

PmC11 differs from clostripain in that is does not appear to require divalent cations for activation.

Several other members of clan CD require processing for full activation including legumain (39), gingipain-R (40), MARTX-CPD (8) and the effector caspases, e.g. caspase-7 (41). To date, the effector caspases are the only group of enzymes that require cleavage of a loop within the central β -sheet. This is also the case in PmC11, although the cleavage loop is structurally different to that found in the caspases, it follows the catalytic His (Fig. 1A), as opposed to the Cys as in the caspases.

All other clan CD members requiring cleavage for full activation do so at sites external to their central sheets (2). The caspases and gingipain-R have both been shown to undergo intermolecular (trans) cleavage and legumain and MARTX-CPD are reported to perform intramolecular (cis) cleavage. In addition, several members of clan CD exhibit self-inhibition, whereby regions of the enzyme block access to the active site (2). Like PmC11, these structures show preformed catalytic machinery and for a substrate to gain access, movement and/or cleavage of the blocking region is required.

The structure of PmC11 gives the first insight into this class of relatively unexplored family of proteins and should allow important catalytic and substrate binding residues to be identified in a variety of orthologues. Indeed, insights gained from an analysis of the PmC11 structure revealed the identity of the Trypanosoma brucei PNT1 protein as a C11 cysteine peptidase with an essential role in organelle replication (42). The PmC11 structure should provide a good basis for structural modelling and, given the importance of other clan CD enzymes, this work should also advance the exploration of these peptidases and potentially identify biologically important substrates.

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Author contributions: KM, JSG DD, IAW, JCM designed research; KM, JSG, DD performed research; KM, JSG, DD, GHC, AS, MAE, and JCM analysed data; AG, SAL, AMD, MAE and IAW supervised various components of the JCSG structural genomics pipeline; MKG, AG, SAL, AMD and MAE contributed reagents/materials/analysis tools; and KM, JSG, GHC, MAE, IAW, and JCM wrote the paper.

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FIGURE LEGENDS

FIGURE 1. Crystal structure of a C11 peptidase from Parabacteroides merdae. (A) Primary sequence alignment of PmC11 (Uniprot ID: A7A9N3) and clostripain (Uniprot ID: P09870) with identical residues highlighted in grey shading. The secondary structure of PmC11 from the crystal structure is mapped onto its sequence with the position of the PmC11 catalytic dyad, auto-catalytic cleavage site (K147), and S₁ binding pocket Asp (D211) highlighted by a red star, a red downturned triangle, and a red upturned triangle, respectively. Connecting loops are colored gray, the main β -sheet is in orange, with other strands in olive, α - helices are in blue and the nonapeptide linker of clostripain that is excised upon auto-cleavage is underlined in red. Sequences around the catalytic site of clostripain and PmC11 align well. (B) Topology diagram of PmC11 colored as in B except that additional (non-core) β-strands are in yellow. Helices found on either side of the central β-sheet are shown above and below the sheet, respectively. The position of the catalytic dyad (H, C) and the processing site (K147) are highlighted. Helices (1–14) and β -strands (1–9 and A–F) are numbered from the N-terminus. The core caspase fold is highlighted in the grey box. (C) Tertiary structure of PmC11. The N and C termini (N, C) of PmC11 along with the central β-sheet (1-9), helix $\alpha 5$, and helices $\alpha 8$, $\alpha 11$ and $\alpha 13$ from the C-terminal domain, are all labelled. Loops are colored gray, the main β -sheet is in orange, with other β -strands in yellow, and α -helices are in blue.

FIGURE 2. Biochemical and structural characterization of PmC11. (A) Ribbon representation of the overall structure of PmC11 illustrating the catalytic site, cleavage site displacement, and potential S₁ binding site. The overall structure of PmC11 is shown in grey, looking down into the catalytic site with the catalytic dyad in red, The two ends of the autolytic cleavage site (K147 and A148, green) are displaced by 19.5 Å (thin black line) from one another and residues in the potential substrate binding pocket are highlighted in blue. (B) Size exclusion chromatography of PmC11. PmC11 migrates as a monomer with a molecular mass around 41 kDa calculated from protein standards of known molecular weights. Elution fractions across the major peak (1-6) were analyzed by SDS-PAGE on a 4-12 % gel in MES buffer. (C) The active form of PmC11 and two mutants PmC11^{C179A} (C) and PmC11^{K147A} (K) were examined by SDS-PAGE (lane 1) and Western Blot analysis using an anti-His antibody (lane 2) show that PmC11 auto-processes, whereas mutants, PmC11^{C179A} and PmC11^{K147A}, do not show auto-processing in vitro. (D) Cysteine peptidase activity of PmC11. Km and Vmax of PmC11 and K147A mutant were determined by monitoring change in the fluorescence corresponding to AMC release from Bz-R-AMC. Reactions were performed in triplicate and representative values \pm S.D. are shown. (E) Intramolecular processing of PmC11^{C179A} by PmC11. PmC11^{C179A} (20 µg) was incubated overnight at 37°C with

increasing amounts of processed PmC11 and analysed on a 10% SDS-PAGE gel. Inactive PmC11^{C179A} was not processed to a major extent by active PmC11 until around a ratio of 1:4 (5 μ g of active PmC11). A single lane of 20 μ g of active PmC11 (labelled 20) is shown for comparison. (F) Activity of PmC11 against basic substrates. Specific activity is shown \pm S.D. from three independent experiments. (*G*) Electrostatic surface potential of PmC11 shown in a similar orientation as Fig. S4, where blue and red denote positively and negatively charged surface potential, respectively, contoured at \pm 5 kT/e. The position of the catalytic dyad, one potential key substrate binding residue D177, and the ends of the cleavage site K147 and A148 are indicated.

FIGURE 3. *PmC11 binds and is inhibited by Z-VRPR-FMK and does not require* Ca^{2+} *for activity.* (A) PmC11 activity is inhibited by Z-VRPR-FMK. Cleavage of Bz-R-AMC by PmC11 was measured in a fluorometric activity assay with (+, purple) and without (-, red) Z-VRPR-FMK. The relative fluorescence units of AMC released are plotted against time (min) (n=3; ±SD). (B) Gelshift assay reveals that Z-VRPR-FMK binds to PmC11. PmC11 was incubated with (+) or without (-) Z-VRPR-FMK and the samples analysed on a 10% SDS-PAGE gel. A size-shift can be observed in the larger processed product of PmC11 (26.1 kDa). (C) PmC11 with the Z-VRPR-FMK from the MALT1-paracacaspase (MALT1-P) superimposed. A 3D structural overlay of Z-VRPR-FMK from the MALT1-P complex onto PmC11. The position and orientation of Z-VRPR-FMK was taken from superposition of the PmC11 and MALTI_P structures and indicates the presumed active site of PmC11. Residues surrounding the inhibitor are labelled and represent potentially important binding-site residues, labelled in black and shown in an atomic representation. Carbon atoms are shown in grey, nitrogen in blue, and oxygen in red. (C) Divalent cations do not increase the activity of PmC11. The cleavage of Bz-R-AMC by PmC11 was measured in the presence of the cations Ca²⁺, Mn²⁺, Zn²⁺, Co²⁺, Cu²⁺, Mg²⁺, Fe³⁺ with EGTA as a negative control, and relative fluorescence measured against time (min). The addition of cations produced no improvement in activity of PmC11 when compared in the presence of EGTA, suggesting that PmC11 does require metal ions for proteolytic activity. Furthermore, Cu²⁺, Fe²⁺, Zn²⁺ appear to inhibit PmC11.

TABLE 1. Crystallographic statistics for PDB ID 3UWS.

Data collection						
Wavelength (Å)	0.9793					
Spacegroup	P2 ₁ 39.11, 108.68, 77.97; β = 94.32°					
Unit cell dimensions a, b, c (Å); β°						
Resolution range (Å)	28.73–1.70 (1.79-1.70)					
Unique reflections	70,913					
R _{merge} on I (%)	10.2 (49.0) 11.0 (52.7) 4.1 (19.2) 15.6 (4.6) 15.9 99.6 (99.8)					
R _{meas} on I (%)						
R _{pim} on I (%)						
I/ σ ₁						
Wilson B (Å ²)						
Completeness (%)						
Multiplicity	7.3 (7.5)					
T V						
Model and Refinement						
Reflections (total/test)	70,883/3,577					
R_{cryst}/R_{free} (%)	14.3/17.5					
No. protein residues/atoms	700/5612					
No. water/EDO molecules	690/7					
ESU based on $R_{free}(\mathring{A})$	0.095					
B-values (Å ²)						
Average isotropic B (overall)	20.0					
Protein overall	18.8					
All main/side chains	16.7/20.8					
Solvent/EDO	29.4/35.6					
R.M.S deviations						
Bond lengths (Å)	0.01					
Bond angles (°)	1.6					
Ramachandran analysis (%)						
Favoured regions	97.0					
Allowed regions	3.0					
Outliers	0.0					

Values in parentheses are for the highest resolution shell. $R_{merge} = \Sigma_{hkl}\Sigma_{i}|I_{i}(hkl) - \langle I(hkl)\rangle|/\Sigma_{hkl}$ $\Sigma_{i}(hkl)$; $R_{meas} = \Sigma_{hkl}[N/(N-1)]^{1/2}\Sigma_{i}|I_{i}(hkl) - \langle I(hkl)\rangle|/\Sigma_{hkl}\Sigma_{i}I_{i}(hkl)$; and and R_{pim} (precision-indicating $R_{merge}) = \Sigma_{hkl}[(1/(N-1))]^{1/2}\Sigma_{i}|I_{i}(hkl) - \langle I(hkl)\rangle|/\Sigma_{hkl}\Sigma_{i}I_{i}(hkl)$ (43), where N is the multiplicity of reflection hkl, and $I_{i}(hkl)$ and $I_{i}(hkl)$ are the intensity of the ith measurement and the average intensity of reflection hkl, respectively (44). R_{cryst} and $R_{free} = \sum_{i} |F_{obs}| - |F_{calc}| / \sum_{i} |F_{obs}|$ for reflections in the working and test sets respectively, where F_{obs} and F_{calc} are the observed and calculated structure-factor amplitudes, respectively. R_{free} is the same as R_{cryst} but for 5% of the total reflections chosen at random and omitted from structural refinement. ESU is the estimated standard uncertainties of atoms and the average isotropic B includes TLS and residual B components.

TABLE 2. Summary of PDBeFOLD (45) superposition of structures found to be most similar to PmC11 in the PBD based on DaliLite (46). The results are ordered in terms of structural homology (Q_H), where %SSE^{PC-X} is the percentage of the SSEs in the PmC11 that can be identified in the target X (where X=Caspase-7 (47), Legumain (3) Gingipain (48), and TcdB-CPD (49) %SSE^{X-PC} is the percentage of SSEs in X (as above) that can be identified in PmC11 (as above); % Seq. ID is the percentage sequence identity after structural alignment; N_{align} is the number of matched residues and RMSD is the root mean squared deviation on the Cα positions of the matched residues.

Enzyme	Family	PDB ID	Q ^H	Z-score	%SSE ^{PC-X}	%SSE ^{X-PC}	% Seq. ID	Nalign	RMSD (Å)	N _{Strands}
PmC11	C11	3UWS	1.00	33.4	100	100	100	352	0.00	9
Caspase-7	C14A	4HQ0	0.16	4.3	38	79	14	162	3.27	6
Legumain	C13	4AW9	0.13	5.5	31	53	13	161	2.05	6
TcdB-CPD	C80	3PEE	0.10	4.9	28	50	12	138	3.18	9
Gingipain	C25	4TKX	0.07	5.4	28	27	12	153	2.97	10

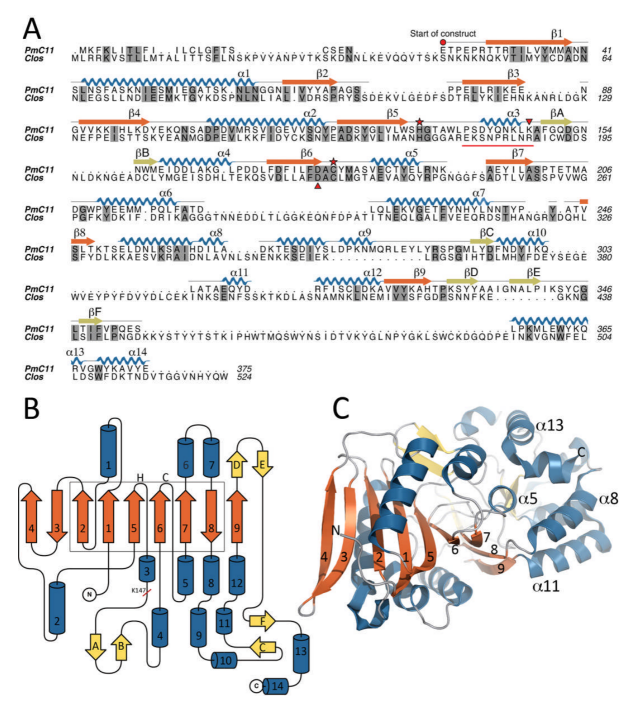


Figure 1

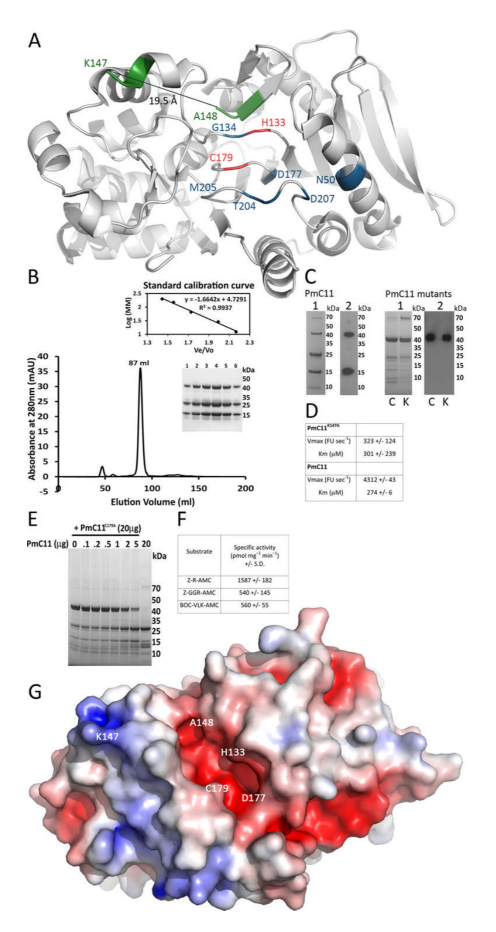


Figure 2

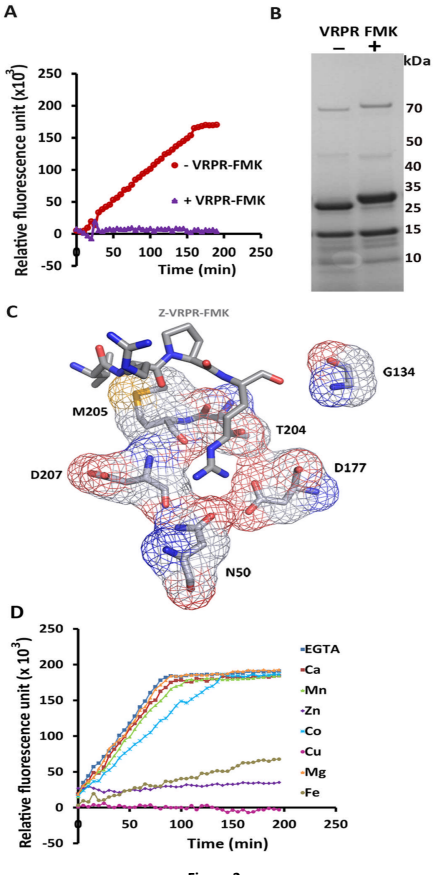


Figure 3