Crystal Structure of a Novel Viral Protease with a Serine/Lysine Catalytic Dyad Mechanism

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The blotched snakehead virus (BSNV), an aquatic birnavirus, encodes a polyprotein (NH2-pVP2-X-VP4-VP3-COOH) that is processed through the proteolytic activity of its own protease (VP4) to liberate itself and the viral proteins pVP2, X and VP3. The protein pVP2 is further processed by VP4 to give rise to the capsid protein VP2 and four structural peptides. We report here the crystal structure of a VP4 protease from BSNV, which displays a catalytic serine/lysine dyad in its active site. This is the first crystal structure of a birnavirus protease and the first crystal structure of a viral protease that utilizes a lysine general base in its catalytic mechanism. The topology of the VP4 substrate binding site is consistent with the enzymes substrate specificity and a nucleophilic attack from the si-face of the substrates scissile bond. Despite low levels of sequence identity, VP4 shows similarities in its active site to other characterized Ser/Lys proteases such as signal peptidase, LexA protease and Lon protease. Together, the structure of VP4 provides insights into the mechanism of a recently characterized clan of serine proteases that utilize a lysine general base and reveals the structure of potential targets for antiviral therapy, especially for other related and economically important viruses, such as infectious bursal disease virus in poultry and infectious pancreatic necrosis virus in aquaculture.

Keywords: serine/lysine catalytic dyad; viral protease; polyprotein; birnavirus; lysine general base

Introduction

Proteases of viral origin have evolved to process virally encoded polyproteins during replication, in co- and post-translational steps. These proteases are essential for viral replication, assembly and infectivity and are therefore potential targets for antiviral drugs.1,2 Viruses of the Birnaviridae family infect animal species belonging to vertebrates, mollusks, insects and rotifers and are characterized by their bi-segmented double-stranded RNA genome (segments A and B).3 Segment A encodes two overlapping reading frames and the larger open reading frame encodes a polyprotein (NH2-pVP2-VP4-VP3-COOH; Figure 1). The polyprotein is processed through the proteolytic activity of VP4 to generate the proteins pVP2 and VP3.4 During virus assembly pVP2 is further processed by VP4 to generate the capsid protein VP2 and structural peptides. Recently, the crystal structure of the infectious bursal disease virus particle (IBDV, an avian birnavirus) was solved at a resolution of 7 A.5 VP2 was the only component seen in the structure of the virus icosahedral capsid.

The blotched snakehead virus (BSNV) was initially identified from a cell line developed from the blotched snakehead fish (Channa lucius) caudal peduncle.6 Abbreviations used: BSNV, blotched snakehead virus; IPNV, infectious pancreatic necrosis virus; IBDV, infectious bursal disease virus; DXV, Drosophila X virus; VP4, viral protein (protease) 4 in the polyprotein (pVP2-X-VP4-VP3) coded for in the birnavirus.

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and those involved in the processing of pVP2 can be described by the consensus Pro-X-Ala \((\text{Ala/Ser})\) motif\(^7\) (Figure 1).

Site-directed mutagenesis studies on VP4 protease have shown that the conserved catalytic residues serine 692 and lysine 729 in BSNV are essential for polyprotein processing.\(^7\) Similar results were found for the VP4 protease in infectious pancreatic necrosis virus (IPNV) (serine 633 and lysine 674)\(^9\) and IBDV (serine 652 and lysine 692).\(^5,9\)

The VP4 protease of the Birnaviridae family therefore are proposed to utilize a serine/lysine catalytic dyad mechanism to catalyze the processing of the polyprotein,\(^4,8,9\) which distinguish it from other viral proteases. Evolutionarily, VP4 protease belongs to the clan SJ and family S50 in the MEROPS protease database.\(^10\) Other characterized proteases with a Ser/Lys catalytic dyad mechanism include: signal peptidase,\(^11-13\) LexA,\(^14\) cl\(\lambda\),\(^15\) UmuD\(^\gamma\),\(^16\) and Lon protease.\(^17\)

Here we report the first structural characterization of a birnavirus protease and the first viral protease to utilize a lysine general base in its catalytic mechanism.

Results and Discussion

Structure solution

After optimization of the VP4 protease construct for crystallization we obtained single and ordered cubic crystals sufficient for diffraction data collection. The crystallizable construct is 217 residues in length (residues 558–773). We have solved the structure of BSNV to 2.2\(\text{Å}\) resolution using single-wavelength anomalous dispersion (SAD) phasing methods. The refined model includes residues 559 to 769 except for a small mobile loop (residues 637 to 648) located at the VP4 surface.

Overall protein architecture

BSNV VP4 protease is approximately 38\(\text{Å} \times 50 \text{Å} \times 36 \text{Å}\) in size with a predominantly negatively charged surface consistent with its calculated isoelectric point. VP4 displays a protein fold, which is made up of two separate domains. Domain I (residues 559 to 693; light blue in Figure 2) is predominately \(\beta\)-sheet in structure. Domain II (694 to 769; dark blue, purple and gold in Figure 2) is \(\alpha/\beta\) in structure. There are a total of 13 strands incorporated within two \(\beta\)-sheets and one \(\beta\)-hairpin. There are three \(\alpha\)-helices (gold in Figure 2) and one \(3_{10}\)-helix (612–614, not shown for clarity) in the structure. A mixed but mostly antiparallel twisted \(\beta\)-sheet contains eight strands (light blue), a parallel \(\beta\)-sheet contains three short strands (dark blue), and a short \(\beta\)-hairpin (purple) contains the remaining two strands. The largest \(\beta\)-sheet is highly twisted and forms an open \(\beta\)-barrel structure, which provides the scaffold for the substrate binding site. The \(\alpha\)-helices pack against the parallel three-stranded \(\beta\)-sheet and the \(\beta\)-hairpin in domain II. A sequence alignment of BSNV VP4 with the proteases from three other birnaviruses (IBDV, IPNV and Drosophila X virus (DXV)) is presented in Figure 3. BSNV VP4 is 23\% identical in sequence to IBDV VP4, 18\% identical in sequence to IPNV VP4 and 18\% identical in sequence to DXV VP4. There are a number of deletions in the VP4 from IBDV, IPNV and DXV in comparison to the BSNV VP4.

Active-site architecture

The structure of BSNV VP4 reveals that the putative nucleophile Ser692 \(\text{O}\) and proposed general base Lys729 \(\text{N}^\circ\) are within hydrogen bonding distance (3.1\(\text{Å}\)) and therefore consistent with previous site-directed mutagenesis studies\(^7\) which showed that these residues are essential for activity (Figure 4). Ser692 resides just before helix a1 and Lys729 resides within helix a2. Lys729 \(\text{N}^\circ\) is also coordinated by Thr712 \(\text{O}^{17}\) (4.0\(\text{Å}\) in molecule A, 3.3\(\text{Å}\) in molecule B) and Pro590 \(\text{O}\) (3.3\(\text{Å}\) in molecule A, 4.3\(\text{Å}\) in molecule B) (Figure 4), the position of the \(\text{N}^\circ\) being slightly different in the two molecules in the asymmetric unit. The \(\text{N}^\circ\) of Lys729 is almost completely buried. The accessible surface area (ASA) of Lys729 \(\text{N}^\circ\) is 5.5\(\text{Å}^2\) (average for molecules A and B) as compared to the average ASA for the \(\text{N}^\circ\) of other lysine residues in BSNV VP4, which is 51.4\(\text{Å}^2\) (average for ten other lysine residues in BSNV VP4, molecules A and B). When a cleavage site (substrate) binds into the binding pocket of VP4, the \(\text{N}^\circ\) of Lys729 would likely become completely buried. This suggests that the \(\epsilon\)-amino group of Lys729 is able to become deprotonated and function as a general base due to a lowering of its \(pK_a\) value resulting from its burial within the protein.

An important component of the catalytic machinery in serine proteases is the oxyanion-hole that works by neutralizing the developing negative charge on the scissile carbonyl oxygen atom during the formation of the tetrahedral intermediates.
Figure 2. The protein fold of BSNV VP4 protease. (a) A ribbon diagram of VP4. The N-terminal mostly antiparallel β-sheet domain (domain I) is in light blue. The C-terminal α/β-domain is shown in dark blue (three-stranded parallel β-
Typically oxyanion holes are formed by two main-chain amide hydrogen atoms that serve as hydrogen bond donors to the developing oxyanion. From the conservation seen in the sequence alignment with other birnavirus VP4 proteases (Figure 3), it would be likely that Ser692 and Gly690 would be the residues that contribute to the oxyanion hole. Yet based on the structure of VP4, with no substrate or inhibitor bound, the residues most likely to contribute atoms to the oxyanion hole are Ser692 and Gln691 whose main-chain NH are pointing towards the binding site (Figure 4). The Gly690 NH is pointing away from the binding site. It appears that the only other possible hydrogen bonding donor that could contribute to the oxyanion hole would be the Gln691 side-chain if it were able to make large changes in its $\gamma_1$ and $\gamma_2$ angles to positions its side-chain N$_3$ over the developing transition state oxyanion. Also it is possible that substrate binding may induce conformation changes in the binding site such that Gly690 N would be a contributor to the oxyanion stabilization.

A possible candidate for a deacylating (catalytic) water (Wat270 in molecule A, Wat134 in molecule B; cyan in Figure 4), is within 3.8 Å of Lys729 N$_z$. This water is 3.8 Å away from Thr712 O$_g$ (2.6 Å), Ser692 O$_g$ (2.7 Å), Ser692 O (2.9 Å) and Pro590 O (2.8 Å). The other ordered water molecules observed near the active site region would likely be displaced by the substrate. For a water to serve as a deacylating or catalytic water in the deacylation step of the hydrolytic reaction it needs to be positioned such that it is within hydrogen bonding distance to a general base for activation as well as at a suitable angle of approach with respect to the carbonyl of the ester intermediate (so-called Bürgi angle$^{18,19}$ of $\sim 107^\circ$).

By superimposing the active sites of the VP4 structure onto the active site of the inhibitor-bound acyl-enzyme structure of Escherichia coli type I signal peptidase, whose nucleophilic serine is covalently bound to a $\beta$-lactam inhibitor, we can approximate the distance to the ester carbonyl carbon and the Bürgi angle. Wat270 is equidistant between Lys729 N$_z$ (3.8 Å) and the ester carbonyl carbon (3.8 Å) of the inhibitor and has an angle of approach to the ester carbonyl carbon atom (Bürgi angle) of 150.2$^\circ$.

Surface analysis reveals that the largest pocket on the surface of VP4 corresponds to an extended bent crevice on the VP4 surface that incorporates the catalytic residues (Ser692 and Lys729) at one end. The dimensions of the crevice are consistent with the size of the side-chains at the P1 and P3 positions in the cleavage site (Schechter & Berger nomenclature$^{19}$). The position of every tenth residue is labeled, as well as the position of Ser692 and Lys729.

Figure 3. A sequence alignment of birnavirus VP4 proteases. The sequences were obtained from the Swiss-Prot data bank with the accession numbers BSNV (Q8AZM0), IBDV (P15480), IPNV (P90205) and DXV (Q96724). The secondary structure and numbering of BSNV is shown above the alignment. The secondary structure is colored the same as for Figure 2. The conserved and catalytic residues are boxed.

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ition of the binding-site relative to the nucleophilic serine hydroxyl group shows that Ser692 O\textsuperscript{γ} attacks from the si-face of the scissile bond. This unusual feature has also been seen in other Ser/Lys proteases such as signal peptidase\textsuperscript{13,20,21} Lex-A-like proteases\textsuperscript{14–16} and Lon protease.\textsuperscript{17}

Comparison to other proteases utilizing a Ser/Lys dyad mechanism

Despite limited sequence identity the VP4 protease shows similarities in the binding site/active site region with many of the other Ser/Lys proteases. It was previously predicted from bioinformatics analysis that there would be structural similarities between VP4 and Lon protease.\textsuperscript{7} VP4 (protease family S50) is different from Lon protease (protease family S16) in that it does not require ATP for activity and that it does not form hexameric complexes. The crystal structures of the proteolytic domain of Lon protease from \textit{E. coli} (PDB: 1RRE\textsuperscript{17}), \textit{Methanococcus jannaschii} (PDB: 1XHK\textsuperscript{22}) and \textit{Archaeoglobus fulgidus} (PDB: 1Z0B\textsuperscript{23}) have been solved. Interestingly, a search for similar structures using the DALI server\textsuperscript{24} reports only one protein with significant similarity in structure, the proteolytic domain of the \textit{A. fulgidus} Lon protease (PDB: 1Z0B) (Figure 5(a)). It has a positional root-mean-square
deviation of 2.4 Å on 121 superimposed Cα atoms. Figure 5(b) shows the superposition of the proteolytic domains of *E. coli* (PDB: 1RRE), *M. jannaschii* (PDB: 1XHK) and *A. fulgidus* (PDB: 1Z0B) Lon protease on BSNV VP4.

VP4 and Lon are similar in that the lysine general base comes from an α-helix (helix a2 in VP4; Figure 3) adjacent to the binding site rather than from the strand (strand β3 in VP4) that creates the mouth of the binding site, as seen in signal peptidase and Lex-like proteases. A comparison of the protein folds of signal peptidase and VP4 is shown by superposition of the *E. coli* signal peptidase active site on the VP4 active site (Figure 5(c)). Previous analysis has shown the catalytic core of UmuD' (and other LexA-like proteases) and signal peptidase to be similar.

Despite only 16% sequence identity, the *E. coli* Lon protease is the most similar of the Lon protease structures to VP4 in the active site region (Figure 6(a)), although a superposition of VP4 and *E. coli* Lon reveals interesting differences, such as the size of the binding pocket. The binding pocket of VP4 is significantly larger and more extended than that of Lon protease (Figure 6(a)). A direct comparison between the general base lysine conformation and environment in *E. coli* Lon protease and VP4 protease is made difficult by the fact that in the Lon structure the serine nucleophile has been mutated to an alanine and therefore cannot hydrogen bond with the lysine ε-amino group. In addition, the Lon structure has a sulfate ion, most likely from the crystallization conditions, interacting with the lysine ε-amino group, which could influence the conformation of the lysine side-chain.

Despite very different protein folds (Figure 5(c)), the strands that line the binding site region and represent most of the characterized hydrogen bonding interaction with inhibitors and substrates in signal peptidase and LexA repressor are superimposable with similar strands in the proposed binding site region of VP4 protease (Figure 6(b) and (c)).

A previous comparison of the micro-environment of the lysine general-base in signal peptidase and LexA-like proteases revealed a difference in the orientation of the Nε atom of the lysine. These
different conformations are stabilized by a hydrogen bond to a hydroxyl group of a second (non-nucleophilic) serine or threonine, which is located in a different orientation with respect to the N^ atom of the lysine. Interestingly, relative to the nucleophile position (Ser O^g), the VP4 lysine general base (Lys729, N^z) is in a position more similar to that seen in the LexA-like proteases than in signal peptidase.14–16,20 (Figure 6(d)).

Model of the X-VP4 cleavage site bound to the VP4 binding site

Because the binding site region of VP4 is similar to that of signal peptidase and LexA we are able to model in a peptide substrate (the X-VP4 cleavage site, P1^ (Ala558) to P3 (Pro555)) into the binding site of VP4 (Figure 7), using as a guide the co-crystal structures of signal peptidase with a non-covalently bound peptide-based inhibitor,20 a covalently bound inhibitor complex with signal peptidase21 and LexA protease with its substrate cleavage site in its binding site.14 This model helps in the description of potential cleavage-site/binding-site interactions. The model is consistent with the P1–P3 residues of the substrate cleavage site region making antiparallel β-sheet interactions with the main-chain atoms in the residues from β-strand b3 (residues Val592 and Phe594) as well as the main-chain atoms from residues in the loop that leads to the nucleophilic residue Ser692 (residues Ala689, Gln691 and Ser692). The hydrogen bonding contacts observed between the docked in and energy minimized X-VP4 cleavage site peptide and the VP4 binding site include: Ala557 O (P1)←Ser692 N, Ala557 O (P1)←Gln691 N, Ala557 N (P1)←Ala689...
O, Gln556 O (P2)···Val592 N, Gln556 N (P2)···Val592 O, and Pro555 O (P3)···Ala689 N. The side-chains of P1 and P3 would point into the S1 and S3 binding pockets. The side-chains for the P2 and P1 residues of the cleavage-sites would be pointing away from the VP4 surface and into solvent (Figure 7), consistent with the variability seen in these residues, for example the Phe at the P10 position at the 417/418 cleavage site (Figure 1). The binding cleft bends just after the location of the P3 residue, consistent with the presence of a proline at this position in the consensus cleavage site (Figures 1 and 7).

Protein–protein interaction in the crystal reveals an N-terminal β-strand cross-over and interactions with the active site

The VP4 crystal structure reveals that the two molecules in the asymmetric unit show an interesting extensive protein–protein interaction (Figure 8). The N-terminal β-strand (β1) makes a cross-over antiparallel β-sheet interaction with the neighboring VP4 molecule in the crystal (residues 562–569). In addition, a β-hairpin extension (β-strands b3 and b4, residues 592–600) from β-sheet 1 (light blue in Figure 2) reaches out to make contacts with the binding site of its neighboring molecule. The side-chain carboxylate from Asp596 sits right above the oxyanion hole of its neighboring molecule (Figure 8). Overall, the protein–protein interaction between molecules A and B within the asymmetric unit buries 1453 Å² of surface per molecule (14.4% of the total protein surface). The level of buried surface predicts that VP4 would exist as a dimer, yet attempts to detect a dimer in solution have so far been unsuccessful. Furthermore, preliminary mutagenesis experiments involving residues at the asymmetric unit interface show no effect on VP4 activity (data not shown).

Concluding Remarks

Structural analysis of the BSNV VP4 protease has allowed us to see for the first time a viral protease that has evolved the unusual serine/lysine dyad catalytic machinery, which has previously been seen in bacterial proteases and amidases. The structure now allows us to ask more detailed questions about the VP4 catalytic mechanism such as the role of Thr712 in its interaction with the lysine general base (Lys729). The solubility and stability of this construct of VP4 protease may make possible the measurement of the pKₐ value for the VP4 lysine general base (Lys729 ε-amino group) with and without bound peptides by NMR titrations. Co-crystallization with peptides will be instrumental in confirming the cleavage-site–binding site interactions we have observed from our docking and modeling studies. The structure of VP4 gives us the relative spatial arrangement for the VP4 amino and carboxy termini relative to the binding pocket orientation and may help in addressing the questions pertaining to the order and timing of cleavage events and also whether this protease catalyzes the cleavage of its viral polyprotein in an intramolecular (cis) or an intermolecular (trans) reaction. The extensive protein–protein interaction seen in the VP4 crystal packing has so far not been shown to be of functional significance but it will continue to be explored in future studies.
Materials and Methods

Cloning, mutagenesis, expression and purification

The cloning of the entire open reading frame of BSNV VP4 (residues 558–791) with a methionine N-terminal residue preceding Ala558 in pET-28b+ was described. To promote crystallization this original expression construct was altered by site-directed mutagenesis to change the codon encoding Pro774 to a stop codon thereby arresting translation at Arg773. The new construct therefore eliminated 18 amino acid residues at the C terminus of VP4. The Quick Change kit (Stratagene) was used for site-directed mutagenesis. The resulting expressed protein is a 217 residues long polypeptide (residues 558–773). Its calculated molecular mass is 23,441 Da and the theoretical pI value is 5.7.

The truncated vector construct described above (pET28_bsnvVP4_558-773) was transformed into the *E. coli* strain Tuner (DE3). For overexpression of selenomethionine incorporated protein, an overnight culture was prepared in M9 minimal media. 10 ml of this overnight culture was used to inoculate one liter of M9 minimal media with kanamycin (0.05 mg/ml) and grown for 7 h. The following amounts of amino acids were then added to the culture as solids (100 mg of lysine, phenylalanine, threonine; 50 mg of isoleucine, leucine, valine; 60 mg of l-selenomethionine). After 15 min the expression was induced by adding 0.5 ml of 1 M IPTG. Cells were induced for 3 h and the pellet was kept frozen at −80 °C.

Frozen cell pellets from one liter of induced culture (2 g wet weight) were completely resuspended in buffer A (50 mM Tris–HCl (pH 8.0), 10% (v/v) glycerol, 10 mM DTT, 7 mM magnesium acetate, 0.1% (v/v) Triton-X100). The cells were lysed by overnight incubation at 4 °C after adding lysozyme (0.2 mg/ml) and benzonase (1 unit/ml). The lysate was centrifuged at 14,000 rpm for 30 min and the clear supernatant was decanted for ammonium sulfate fractionation.

Ammonium sulfate (AS) fractionation was carried out at 30, 40, 50 and 60% (w/v) AS concentrations. After adding the AS, the supernatant was incubated on ice for 30 min, then, the precipitates were pelleted by centrifugation at 14,000 rpm for 30 min. Pellets were dissolved in approximately 1 ml of buffer B (20 mM Tris–HCl (pH 8.0), 50 mM NaCl, 1 mM EDTA, 10% glycerol, 10 mM DTT). The BSNV VP4_558-773 precipitated at the 40% and 50% AS concentrations. These fractions were pooled and dialyzed against buffer A.

The dialyzed sample (∼2 ml) was applied to a 0.7 ml Q-Sepharose anion exchange column equilibrated with buffer B. The protein was then eluted from the column using a NaCl step-gradient.

SDS-PAGE analysis of the collected fractions demonstrated a single band corresponding to VP4_558-773. The protein was concentrated to 23 mg/ml using a Millipore centrifugal filter (5000 cut off).

Crystallization

Crystallization was carried out by the sitting drop vapor diffusion method at 21 °C. 1 μl of the protein sample was mixed thoroughly with 1 μl of the reservoir solution and the mixture was equilibrated against 1 ml of the reservoir solution. The protein was at a concentration of 23 mg/ml suspended in buffer C. Optimized crystallization reservoir solution conditions consisted of 0.1 M Tris–HCl (pH 8.5), 25% (v/v) PEG2000 MME, 60 mM Mg(OAc)2. Cryo-solvent conditions were the same as the crystallization conditions supplemented with 20% glycerol.
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glycerol. Crystals were flash cryo-cooled in liquid nitrogen.

Data collection

Diffraction data were collected on the selenomethionine incorporated crystals at beamline X26C, at Brookhaven National Laboratory, National Synchrotron Light Source using a Q4 CCD detector and a Nonius CAD4 Goniometer. The crystal-to-detector distance was 200 mm. Data were collected with 1° oscillations, and each image was exposed 30 s. Data were processed to 2.2 Å resolution with the program HKL2000. See Table 1 for data collection statistics.

Structure determination and refinement

Single-wavelength anomalous dispersion (SAD) phasing was carried out using a data set collected at the peak wavelength (0.97804 Å) with the program autoSHARP. Values for $f^2$ and $f'$ were found to be $-6.16$ and $4.06$, respectively. Eight of the ten possible selenium sites were found. To improve the quality of the electron density maps we used the program SOLOMON, run within autoSHARP, to perform a density modification based on solvent flipping. The program Arp/Warp$^{27}$ automatically built approximately 80% of the polypeptide chains. The rest of the model was built and fit using the program XFIT within the suite XTALVIEW.$^{30}$ The structure has been refined using the programs CNS$^{31}$ and Refmac5$^{32}$ to an $R_{	ext{cryst}}$ of 22.9% and an $R_{	ext{free}}$ of 27.9%. The average $B$ value for all atoms is 27.6. There are 401 residues and 400 water molecules in the model with two molecules in the asymmetric unit. The refined model includes residues 559 to 769 for each molecule. There is no visible electron density in either molecule in the asymmetric unit for residues 637 to 648. These residues are located at the surface of the molecule and presumably form a mobile loop. There is also no electron density for the N-terminal three residues and the last five C-terminal residues. The stereochemistry of the structure was analyzed with the programs PROCHECK.$^{33}$ For further data collection, structure solution and refinement details see Table 1.

Structural analysis

The secondary structural and protein topology analysis was performed with the programs PROMOTIF$^{34}$ and HERA$^{35}$ respectively. The programs SUPERIMPOSE$^{36}$ and SUPERPOSE$^{37}$ were used to superimpose molecules for comparing the different molecules in the asymmetric unit and for comparison to other proteases. The program CONTACT within the program suite CCP4$^{38}$ was used to measure the hydrogen bonding and van der Waals contacts within the enzyme. The program CASTp$^{39}$ was used to analyze the molecular surface and measure the substrate binding site. The program SURFACE RACER$^{1.2}$ was used to measure the solvent accessible surface of the protein and individual atoms within the protein. A probe radius of 1.4 Å was used in the calculations. The Protein–Protein Interaction Server$^+$ was used to analyze the interactions between the molecules in the asymmetric unit in the VP4 crystal.

Modeling of the X-VP4 cleavage site into the VP4 protease binding site

A model of the X-VP4 cleavage site (P1’ (A558) to P4 (R554)) bound to the VP4 binding site was prepared using the structure of the covalently bound penem inhibitor complex with signal peptidase$^{13}$ (PDB: 1B12) to guide the position of the P1 Ala residue of the substrate. The positions of the P2, P3 and P4 residues were guided by the structure of the cleavage site of LexA$^{14}$ (PDB: 1JHE) and the co-crystal structure of a non-covalently bound peptide-based inhibitor with signal peptidase$^{21}$ (PDB: 1TT7D). The model was prepared using the program XFIT within the suite XTALVIEW.$^{30}$ The model was energy-minimized using the program CNS.$^{31}$

Figure preparation

Figures were prepared using the programs Molscript,$^{41}$ Raster3D$^{42}$ and PyMol.$^\ddagger$

† http://www.biochem.ucl.ac.uk/bsm/PP/server/
‡ http://pymol.sourceforge.net/

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Table 1. Data collection, phasing and refinement statistics

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<td>Anomalous multiplicity</td>
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C. Phasing statistics

| Number of sites       | 8              |
| Number of acentric reflections | 38,524 |
| Overall FOM acentric  | 0.21053        |
| Number of centric reflections | 1988    |
| Overall FOM centric   | 0.01738        |

D. Refinement statistics

| Protein molecules (chains) in au | 2 |
| Water molecules                 | 401 |
| Total number of atoms           | 3998 |
| $R_{	ext{cryst}} / R_{	ext{free}}$ (%) | 22.6/27.9 |
| Average B-factor (Å$^2$) (all atoms) | 27.6 |
| Estimated overall coordinate error (Å) | 0.150 |
| rms deviation on angles (deg.)  | 1.70 |
| rms deviation on bonds (Å)      | 0.016 |

The data collection statistics in parentheses are the values for the highest resolution shell (2.28 Å to 2.20 Å).

$^a$ $R_{	ext{merge}} = \frac{\sum |I_i - \langle I \rangle|}{\sum |I|}$, where $I$ is the observed intensity, and $\langle I \rangle$ is the average intensity obtained from multiple observations of symmetry-related reflections after rejections.

$^b$ FOM, figure of merit $= \frac{1}{N} \sum |P(\alpha) - \langle P \rangle|$, where $P(\alpha)$ is the phase angle and $P(\alpha)$ is the phase probability distribution.

$^c$ $R_{	ext{merge}} / R_{	ext{free}} = \frac{\sum |F_{\text{calc}} - |F_o| |}{\sum |F_o|}$, where $F_{\text{calc}}$ and $F_o$ are the observed and calculated structure factors, respectively.

$^d$ $R_{	ext{merge}}$ is calculated using 5% of the reflections randomly excluded from refinement.
Protein Data Bank accession codes

Atomic coordinates and experimental structure factors of BSNV VP4 protease have been deposited with the RCSB Protein Data Bank under the accession code 2GEF.

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