Turnip yellow mosaic virus protease binds ubiquitin suboptimally to fine-tune its deubiquitinase activity

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Abstract

Single-stranded, positive-sense RNA viruses assemble their replication complexes in infected cells from a multi-domain replication polyprotein. This polyprotein usually contains at least one protease whose primary function is to process the polyprotein into mature proteins. Such proteases also may have other functions in the replication cycle. For instance, cysteine proteases (PRO) frequently double up as ubiquitin hydrolases (DUB), thus interfering with cellular processes critical for virus replication. We previously reported the crystal structures of such a PRO/DUB from *Turnip yellow mosaic virus* (TYMV) and of its complex with one of its PRO substrates. Here we report the crystal structure of TYMV PRO/DUB in complex with ubiquitin. We find that PRO/DUB recognizes ubiquitin in an unorthodox way: It interacts with the body of ubiquitin through a split recognition motif engaging both the major and the secondary recognition patches of ubiquitin (Ile44 patch and Ile36 patch, respectively, including Leu8 which is part of the two patches). However, the contacts are suboptimal on both sides. Introducing a single point mutation in TYMV PRO/DUB aimed at improving Ub-binding led to a much more active DUB. Comparison with other PRO/DUBs from other viral families, particularly coronaviruses, suggests that low DUB activities of viral PRO/DUBs may generally be fine-tuned features of interaction with host factors.

Introduction

Host-pathogen relationships are complex. The outcome of pathogen infection depends on a subtle balance between host immune responses triggered by infection and pathogen replication aimed at promoting propagation. In recent years, ubiquitination and deubiquitination events have emerged as central processes in antiviral mechanisms and viral multiplication (1–5). Ubiquitination is the conjugation of ubiquitin (Ub), a highly conserved 76-residue protein, to a target protein, through the formation of an isopeptide bond between the C-terminal glycine residue of Ub to a Lys of the target protein (6). Targets of ubiquitination are cellular proteins mostly involved in host immune responses and/or viral proteins (4). In certain cases, ubiquitin-like modifiers such as SUMO, NEDD8 or Ub-like interferon-simulated gene 15 (ISG15) may also be covalently attached to various substrates (7). Substrates are often polyubiquitinated, *i.e.* a chain of multiple Ub moieties, each linked by an isopeptide bond, is formed. Depending on the linkage type between distal and proximal Ub, the fate of tagged proteins varies, from targeting to proteasome or other degradation pathways for degradation (8) to non-proteolytic events such as interaction with various partners (6). Ubiquitination is a reversible process. Deubiquitination is catalyzed by deubiquitinases (DUBs), which can cleave isopeptide bonds to either trim, degrade or edit polyUb chains from substrate proteins (7).

Since viruses strictly depend on the host to replicate and spread, they have evolved to circumvent or even hijack for their own advantage the ubiquitin-dependent responses triggered by entry of virus into the cell and subsequent replication (4, 9, 10). Indeed, a number of viruses have evolved DUBs (11, 12), either to counteract antiviral mechanisms or to favor their replication. The targets of viral DUBs can be cellular and/or viral proteins (11). As an example, deubiquitination of cellular proteins by viral DUBs can down-regulate the production of diverse antiviral molecules such as interferons or cytokines and allow viruses to evade host immune responses (12, 13). Another example is the deubiquitination of viral proteins by viral DUBs that avoids their targeting to the proteasome, a process that can be viewed as a rescue of these viral proteins. For some viruses an excess of certain viral proteins can be detrimental for viral replication (14, 15). These viruses use the deubiquitination step to modulate proteasome-dependent degradation to subtly control the level of the relevant proteins (9). For instance, adjusting the amount of RNA-dependent RNA polymerase (RdRp) may regulate the replication of some RNA viruses such as *Sindbis virus* (SINV) (16), *Turnip yellow mosaic virus* (TYMV) (14, 17) or *Hepatitis A virus* (HAV) (18).

DUBs are cysteine proteases or metalloproteases and are classified into seven families including two new families that have been recently defined (7, 19–21). These enzymes can specifically cleave one or several Ub linkage types or display a more general deubiquitinating activity. DUBs encoded
by some single-stranded, positive-sense RNA viruses ((+)-ssRNA viruses) such as arteriviruses, coronaviruses, picornaviruses and tymoviruses are actually bifunctional enzymes also responsible for the viral polyprotein maturation through a protease activity (PRO) that cleaves defined peptide bonds (22–27). The molecular determinants that regulate these dual activities remain largely unknown.

The dual PRO/DUB enzyme encoded by Turnip yellow mosaic virus (TYMV) is a valuable example to address these questions because it is known to tightly regulate the level of RdRp during viral replication (24, 28). TYMV encodes an essential 206 kDa replicative polyprotein called 206K, which contains sequence domains indicative of methyltransferase (MT), protease (PRO), NTPase/helicase (HEL), and RNA-dependent RNA polymerase (POL or RdRp) activities. The TYMV PRO domain first cleaves the 206K to give rise to an intermediate product called 140K (encompassing the MT, PRO and HEL domains) and the protein 66K (POL), after which it cleaves the 140K intermediate to release proteins called 98K (MT-PRO) and 42K (HEL) (29–31). The 66K polymerase is subject to phosphorylation and ubiquitination events triggered by the host, which ultimately target the modified protein to the proteasome where it is degraded (14, 32). Due to its DUB activity, the PRO domain of TYMV can counteract such degradation and inhibit 66K degradation (24). The whole process ensures a low level of 66K/POL in infected cells (33), the accumulation of which is deleterious for viral RNA replication (14). Although TYMV 66K is likely to be tagged with Lys48-linked polyUb chains and TYMV PRO/DUB is able to process in vitro Lys48- and Lys63-linked polyUb chain (24), little is known about the type and the composition of polyUb chains attached to the 66K polymerase. In addition, how TYMV PRO/DUB recognizes ubiquitinated 66K is unknown.

The structure of TYMV PRO/DUB (34) has shown that the protein is a DUB from the ovarian tumor (OTU) family (7) that evolved to acquire a PRO function (34). Strikingly, Tymoviridae PRO/DUBs are the only OTU DUBs that lack two elements of the canonical cysteine protease active site displayed by all other OTU DUBs. First, it has only a catalytic dyad (composed of Cys783 and His869) instead of the typical (Cys-His-Asp/Asn) triad of OTU DUBs. The Asp/Asn residue is replaced by a serine (Ser871 in TYMV PRO/DUB) that is conserved in the other members of the Tymoviridae family (34). Second, there is no pocket that could constitute the oxyanion hole that is formed during the catalytic mechanism (34). In contrast, Tymoviridae PRO/DUBs display a unique loop (G165P166P167) in close vicinity of the active site (34). We previously concluded this loop is involved in substrate recognition and contributes to align the side chains of catalytic residues (28). The mobility of this loop therefore would contribute to switching from the PRO activity to the DUB activity. In one of the TYMV PRO/DUB crystal structures, the protein has adventitiously self-assembled into the active form (35), leading to a physiologically relevant PRO/DUB:PRO complex that gives clues to the mechanism of the PRO function of the enzyme. Indeed, this structure provides a snapshot of how the enzyme recognizes the C-terminal extremity of another PRO domain during the PRO↓HEL cleavage event, which occurs in the course of polyprotein maturation (30, 34).

To better understand the DUB function of the TYMV PRO/DUB domain, we report its crystal structure in complex with ubiquitin. We supplemented the low resolution of the structure (3.7Å) with molecular dynamics (MD) simulations. We used this modeling approach to further probe the differences in molecular recognition between two of its substrates, i.e. PRO of the PRO↓HEL cleavage site and ubiquitin. A structure-guided mutagenesis study identified point mutants with an increased DUB activity, showing that the unusual recognition of Ub by TYMV PRO/DUB is suboptimal. Comparison of this PRO/DUB-Ub structure with that of the PRO/DUB:PRO complex that occurs during polyprotein processing (34) shows that these unrelated substrates are recognized by largely overlapping recognition surfaces.

Results

Overall structure of the covalent TYMV PRO – Ub complex

In order to solve the crystal structure of a TYMV PRO/DUB:ubiquitin complex and because the affinity of a single module of Ub for the enzyme is
low (24, 34), we used a modified form of Ub (Ub-VME) in which the C-terminal Gly76 is substituted with a vinyl methylester function that spontaneously and irreversibly forms a covalent linkage with the catalytic cysteine of DUBs in a Michael addition (36, 37). TYMV PRO/DUB and Ub-VME were incubated at 25°C, leading to the formation of a covalent complex as evidenced by SDS-PAGE (Supplementary Figure 1A), which was then purified by size exclusion chromatography (Supplementary Figure 1B). Crystals of the protein complex grew in a single drop after 120 days. Only a single crystal showed acceptable diffraction that allowed us to collect data. The structure was solved at 3.7 Å resolution by molecular replacement. The crystallographic asymmetric unit contains two PRO/DUB-Ub complexes, one of which is well ordered and could be modelled with confidence, except in a few places where density was ambiguous. We complemented this crystallographic model with molecular dynamics simulations that helped to resolve ambiguities and allowed an accurate view of the complex (see below for details). The second complex in the asymmetric unit was modeled from the first and the structure refined with tight non-crystallographic restraints with good statistics (Table 1). We will limit our analysis to the single well-ordered complex composed of chains A (TYMV PRO/DUB, ordered residues 732-876 by polyprotein numbering) and B (Ub-VME, residues 1-76 including the terminal glycyl-vinylmethylester covalently linked to the catalytic Cys783).

The interaction surface of the PRO/DUB-Ub complex measured by PISA server (38) buries 860 Å² (11%) and 908 Å² (19.5%) of solvent-accessible area for the TYMV PRO/DUB and Ub molecules, respectively, which is on the lower side of the reported values for other DUB:Ub complexes (39–41). As in these other complexes, the Ub-binding interface of TYMV PRO/DUB can be viewed as two distinct areas (Figure 1). First, the body of Ub is bound by a surface of TYMV PRO/DUB distant from the PRO/DUB active site and contributed on one side by its N-terminal lobe (residues 732-770) and on the other by the C-terminal lobe (residues 836-876. For a more detailed description of the three lobes, see (34)). Second, the C-terminal extremity of Ub inserts into the TYMV PRO/DUB catalytic cleft between the central lobe (residues 773-835) and the C-terminal lobe.

**TYMV PRO/DUB uses two polar loops to simultaneously engage the two major hydrophobic patches on the body of Ub**

Distant from the active site the interaction of TYMV PRO/DUB with Ub appears quite unusual: Ub plugs into a large groove at the surface of TYMV PRO/DUB, so that both of its major recognition patches (the so-called Ile44 and Ile36 patches) are bound simultaneously (Figure 1A). On one side of the groove, the Ile44 patch is contacted by the Tymoviridae-specific N-terminal lobe (residues 732-772), while on the other side the Ile36 patch is contacted by the C-terminal lobe that is common to all OTU DUBs.

**The Ile44 patch-interacting site**

The side chains of TYMV PRO/DUB Glu759 and Asn760, from the N-ter lobe of the protein, project directly towards the Ile44 patch of Ub, composed of residues Ile44, Leu8, His68 and Val70 (6) (Figure 2A). In previous work, based on a docking model of the complex and a subsequent mutagenesis study, we suspected the involvement of Glu759 and Asn760 in Ub recognition. We hypothesized the presence of a hydrogen bond between Asn760 and His68, and a salt bridge between Glu759 and Lys6 and/or His68 (34). Indeed the simultaneous replacement of these two residues by two Gly residues (mutation E759G/N760G) led to a small but significant decrease of DUB activity in vitro (34). No such interactions are seen in the crystal structure (Figure 2A). However, Lys6 of Ub is engaged in a strong crystal contact with Asp739 and Thr741 from a neighboring molecule (data not shown). This precludes any interaction with TYMV PRO/DUB in the asymmetric unit but does not exclude the existence of such an interaction in solution. Hence, in order to better understand the interaction network between TYMV PRO/DUB Glu759 and Asn760 residues and the Ile44 patch of Ub, we performed molecular dynamics simulations of the complex. For the starting model, we made two changes that depart from the crystal structure: we first replaced the C-terminal ubiquitin residue (a Gly substituted with a vinyl methylester group, see above) with an unmodified glycine. Thus the complexes we
simulated mimicked product-bound states, as in our previous structure of a PRO/DUB:PRO complex (34). Second, we modeled residues 727-731 that are not visible in the electron density map and we acetylated Ser727 to better model the native state of the TYMV PRO/DUB domain (that is, linked to the rest of the polypeptide by its N-terminus). When free from crystal contacts, Lys6 now points towards the side chain of Glu759. However, in two independent 50-ns simulations, the two residues never engaged in the formation of a stable salt bridge. This is readily shown by the distribution of distances between atom Nζ of Lys6 and atom Oε of Glu759, which shows only a minor peak at 2.8 Å (Figure 2B). Instead the simulations confirm a strong involvement in the interface of the aliphatic portions of the Glu759, Asn760, Thr761 and Thr763 side chains. They pack against the hydrophobic Ile44 patch of Ub (Figure 2C top). The only polar interaction between these polar residues and ubiquitin is a hydrogen bond between Thr763 and Gln49 (about 50% occupancy) (Figure 2C bottom).

Although our previous docked model of the TYMV PRO/DUB:Ub complex suggested the potential involvement of another part of the N-terminal lobe (including Leu732 and Leu765) in Ub binding (34), the crystal structure now shows these residues lying at the edge of the Ile44 patch in the vicinity of Ub residues Gln49, Glu51 and Asp52 (Supplementary Figure 2) and with at best a small contribution to the interface. Simulations consistently show that Leu732 and Leu765 actually tend to come away from ubiquitin (data not shown). This is in agreement with our previous report that Ala mutations of these residues (mutation L732A/L765A) showed no effect on DUB activity in vitro (34), ruling out their involvement in Ub binding.

**The Ile36-interacting site**
The Ile36 patch of Ub, the core of which is composed of residues Ile36 and Leu71 (6), is positioned against segment 840-847 of TYMV PRO/DUB (Figure 2D), with Ile847 also interacting with Ub Leu8 (see below). Arg844 is clearly the TYMV PRO/DUB residue closest to Ile36, but density for Arg844’s side chain fades beyond its Cγ. Thus, in order to obtain a better view of the Arg844 side chain, we analyzed its behavior in molecular dynamics simulations. The simulations show that the charged end of the Arg844 side chain is highly mobile and samples a large conformational space, where it finds several defined bound states. Indeed, Arg844’s guanidinium function alternatively makes a transient salt bridge with the Ub Glu34 side chain (Figure 2E) or hydrogen bonds with the Ub Gln40 side chain or main chain carbonyls of Ub Glu34 and Gly35 (data not shown). In contrast, Arg844’s aliphatic portion down to Cγ remains stably packed against the Ub Ile36 patch (Figure 2F). Thus we arrived at a similar picture as for the region of the Ub Ile44 patch, with polar or charged residues of TYMV PRO/DUB contacting the hydrophobic patches of Ub almost exclusively by their aliphatic portions.

**The Leu8-interacting site**
Leu8 of Ub is located between the two hydrophobic patches, in a flexible loop that connects the two first α-helices (42, 43). This loop can undergo conformational changes, from a « loop-out » to a « loop-in » position (44), which in turn enables it to be part of either the Ile44 or the Ile36 patch (44, 45) (Figure 2G). The flexibility of the loop that comprises Leu8 is now recognized to be important for recognition of ubiquitin-binding proteins (UBPs) (43). In the TYMV PRO/DUB-Ub complex, this loop adopts an intermediate position between the « loop-out » and « loop-in » positions and Leu8 points towards the bottom of the groove (Figure 2G). In this region, Ile847 and Phe849 of TYMV PRO/DUB make strong hydrophobic contacts with Leu8, Thr9, Val70, Leu71 and Leu73 of Ub. This centrality of Ile847 in an interaction network based essentially on hydrophobic interactions is consistent with our previous work. Indeed, mutating Ile847 to Ala, which conserves its apolar properties, has a significant but mild effect on DUB activity, both in vitro and in vivo, while addition of a negative charge in this region (mutation I847D) drastically decreased DUB activity (28, 34).

In summary, the crystal structure of the covalent complex between TYMV PRO/DUB and Ub, supplemented by molecular dynamics simulations, shows that Ub nestles in a cavity of TYMV PRO/DUB. This binding mode mimics a clamp that
holds Ub through hydrophobic interactions made, surprisingly, by TYMV PRO/DUB polar and charged residues, with one side of the clamp formed by the α2-β2 loop containing the P758E759N760T761A762T763 motif, and the other side of the clamp constituted essentially by residues belonging to β3 and β4 strands, centering on Arg844 in the β3-β4 loop (Figure 2H and 1B). The bottom of the Ub binding groove is composed of hydrophobic residues that are part of beta-strands β3 and β4. On its side, Ub engages three binding sites simultaneously, i.e. in addition to its C-terminus (see below), the two hydrophobic patches centered on Ile44 and Ile36 connected by the loop encompassing Leu8. Despite this three-part contact, the buried surface is on the small side compared to other viral DUB:Ub complexes.

TYMV DUB activity can be improved by point mutations that affect the atypical binding surface used to contact Ub

In order to probe the puzzling use of polar residues in TYMV PRO/DUB to bind the Ub hydrophobic patches and to assay several structure-guided point mutants for DUB activity. The activity of the mutant proteins produced in E. coli was measured using the general DUB substrate ubiquitin-7-amino-4-methyl coumarin (Ub-AMC) as described (24, 34). In this in vitro test, TYMV PRO/DUB is not saturated, even at the highest Ub-AMC concentrations attainable (34). It is therefore not possible to determine precisely $k_{cat}$ and $K_m$ parameters. Instead, the assay far from saturation allows the measurement of $K_{app}$ that approximates $k_{cat}/K_m$. First, we mutated polar residues that interact with the Ile44 hydrophobic patch of Ub. Replacement of Glu759 or Asn760 by alanine resulted in substantially increased DUB activity compared to the WT enzyme: 137% ± 7% and 135% ± 8% for E759A and N760A, respectively (Figure 3A). The charged carboxylate of Glu759 is thus detrimental to DUB activity. This is consistent with structural data that highlight the importance of the apolar portions of the interfacial residues and the flickering nature of polar interactions, such as the Glu759-Lys6 salt bridge. Finally, DUB activity can actually be increased by the removal of polar groups and maintaining only apolar side chains. In contrast, mutation of Thr763 to alanine showed a slightly decreased DUB activity (89% ± 8%, see Figure 1G), again in accordance with structural and sequence data (Figure 1B, Figure 2C bottom).

Second, we wanted to better understand the role of Arg844 side chain, which can not only make van der Waals contacts with the Ile36 hydrophobic patch of Ub, but also can form hydrogen bonds or a salt bridge with several Ub residues (Figure 2E). We thus replaced Arg844 by Ala. This mutation led to a dramatic 3-fold increase of DUB activity (320% ± 14%, see Figure 3A), an effect also observed with the double mutant N760A/R844A (344% ± 5%, see Figure 3A). This implies that Arg at position 844 of TYMV PRO/DUB is detrimental to DUB activity. Since this residue is located far away from the active site, it is likely that its polar side chain alters the binding to Ub rather than affects the turn-over of the enzyme. The observation that TYMV DUB activity can be substantially improved by point mutations prompted us to model the interaction of the R844A mutant with Ub. We performed molecular dynamics simulations of the complex in the same conditions as for the wild type and with the same initial models, albeit with the truncation of R844 side chain to mimic an alanine. In two independent replicates the complex shifted from its initial conformation to one in which A844 packs against the center of the Ile36 patch, as exemplified by the new van der Waals contacts of A844 C$\beta$ with Ile36 C$\gamma$1 (Figure 3B). In contrast, the catalytic dyad's dynamics were not affected, as shown by the continued rarity of the activating hydrogen bond between H869 and C783 (Figure 3C). These results show how the complex can easily adjust to the much smaller alanine side chain to effectively shield the Ile36 patch from solvent, without disturbing the active site. They confirm the centrality of the apolar contact between residue 844 and Ub and suggest the effect of R844A is indeed on ubiquitin binding rather than on enzyme turnover.

Altogether, these results reinforce the conclusion that TYMV PRO/DUB indeed binds both Ile44 and Ile36 patches of Ub suboptimally, contributing to a poor DUB activity. They establish that point mutations aimed at improving Ub binding do result in a considerably increased DUB activity.
Binding mode of the C-terminal tail of Ub: how TYMV PRO/DUB recognizes different C-terminal sequences

In addition to interactions that involve the body of Ub via its two conserved hydrophobic patches, a large portion of PRO/DUB:Ub interacting surface engages the five C-terminal residues of Ub inserted into the catalytic cleft of TYMV PRO/DUB (Figure 1A). As expected, the C-terminal tail of Ub adopts a beta conformation that creates a dense hydrogen-bonding network with TYMV PRO/DUB residues that belong to the substrate binding site (Figure 1B). These involve the backbone carbonyl oxygens and amide hydrogens of Leu822, Thr824 and Ser868, and side chains of Thr824 and Ser868 (Figure 4A). The strong electron density in the vicinity of Arg74 of Ub was difficult to interpret at this resolution, because Arg72 and Arg42’s side chains also point in the same direction. Again, molecular dynamics simulations were helpful in settling this ambiguity. Only the Cβ of the three arginines were initially modeled in the crystal structure. Alternate solutions for their side chains were then generated, all consistent with electron density, and simulations were performed. Simulations nicely converged to the same arrangement in that region, no matter the starting point. We kept this solution for refinement of the crystal structure (Figure 4B). The three arginines all point towards the acidic S5 pocket of TYMV PRO/DUB, comprised of Glu816 and Glu825 (34). Arg72 and Arg74 of Ub both make salt bridges with Glu816 and/or Glu825, a type of interaction often seen in other complexes involving Ub (37, 40, 41, 46–49). The side chain of Asp39 fromUb also points towards Arg74 (the two make a stable salt bridge in the simulations), making a ring of acidic residues around the three clustered arginines (Figure 4B).

We can now assess how the PRO/DUB catalytic cleft adjusts to different substrates. Indeed, TYMV PRO/DUB recognizes a consensus peptide substrate K/R-L-X-G-G/A/S (corresponding to positions P5-P4-P3-P2-P1), where X is any amino acid, corresponding to the HEL↓POL and PRO↓HEL cleavage sites (KLNGA↓ and RLLGS↓, respectively) and the C-terminal extremity of Ub (RLRGG↓). The requirements for this sequence can be explained by a comparison of the crystal structure of TYMV PRO/DUB-Ub complex with that of the PRO/DUB:PRO complex with the C-terminal extremity of a PRO from PRO↓HEL cleavage site inserted in the catalytic cleft of a PRO enzyme (see Figure 3 in ref (34)). Overall, the acidic S5 pocket of TYMV PRO/DUB, comprised of residues Glu816 and Glu825, highly conserved in the Tymoviridae family (see sequence alignment in Figure 1B), always accommodates a basic residue at position P5 (Lys or Arg, see above). In the specific case of Ub, the combination of the TYMV PRO/DUB acidic patch and an acidic residue from Ub (Asp39) perfectly accommodate the three Arg of Ub, two belonging to its C-terminus (Arg72 and Arg74) and one oriented towards its C-terminus (Arg42). The strict requirement for a Leu at position P4 is imposed by the hydrophobic S4 pocket, created by residues Val840, Ser842, Ile847 and His862 (Figure 4C), rather conserved in homologous PROs (Figure 1B). The absence of a real S3 pocket leads to a relaxed specificity at the P3 position, accommodating structurally unrelated residues such as Arg, Leu or Asn. The conserved Gly at position P2 fits in a pocket containing Ser868 and Phe870. These two residues, conserved in the Tymoviridae family (Figure 1B), constitute the so-called glycine specificity motif (GSM), a common feature of alphavirus PROs and PRO/DUBs (50, 51). Finally, limited specificity for a small side chain at position P1 is due to the flexible enzyme’s T864G865P866P867S868 loop which regulates the constriction of the S1 pocket and consequently substrate specificity and enzymatic activity (28). The GPP motif is a strictly conserved (Figure 1B) addition to the OTU DUB fold found in the Tymoviridae family.

In conclusion, the C-terminal residues of Ub assume an extended conformation and occupy the catalytic cleft of TYMV PRO/DUB. They do so by creating an intricate network of salt bridges, further strengthened by numerous hydrophobic contacts. The consensus sequence of the C-terminal extremity of PRO, HEL and Ub, composed of invariant residues (positions P4 and P2), conserved residues (positions P5 and P1) and non-conserved residues (position P3), eventually defines which residues are specificity determinants. These allow TYMV PRO/DUB to discriminate amongst its substrates. The TYMV PRO/DUB is actually a...
deubiquitinase that acquired a protease function to process its polyprotein (34) (see also discussion below). It is likely that the PRO’s substrate cleavage sites have evolved to mimic the C-terminal extremity of Ub. Such optimization of substrate sequences allows a single enzyme to perform several enzymatic reactions. While this may be a simplification, it supports the “genetic economy” concept that allows (+)ssRNA viruses to ensure numerous enzymatic functions despite a small and compact genome.

**TYMV PRO/DUB contacts the bodies of unrelated substrates through highly overlapping recognition patches**

By comparing the PRO/DUB-Ub and PRO/DUB:PRO complexes (Figure 5A,C), we show that the TYMV PRO/DUB recognition surfaces for two of its substrates overlap to a large extent (Figure 5B,D). Notably, as for Ub binding (Figure 5B), PRO binding involves residues E759-N760 on one side and R844 on the other, with I847 in the middle (Figure 5B). However, the N-terminus of TYMV PRO/DUB differentially recognizes the substrates. While the Pro733Ala734Pro735 motif is prominently involved in PRO recognition (Figure 5D), only Leu732 (Figure 5B) makes a tenuous contact to Ub in the crystal structure (Supplementary Figure 2), a contact that is not stable in simulations (see above). The Pro733Ala734Pro735 motif provides a strong additional apolar contact that makes the PRO/DUB:PRO complex less dependent on the hydrophobic bottom of the binding groove. Indeed, our structural data and mutagenesis studies (this work and references (28, 34)) establish Ile847 as a central residue for Ub recognition but with less of an impact on PRO binding. In addition, the enzyme harbors a single catalytic site, comprised of Cys783 and His869, for both its protease and deubiquitinase activities. PRO and Ub thus share the same TYMV PRO/DUB ligand binding site and bind in an orientation that exposes their C-terminal extremity towards the catalytic residues. Their interactions with TYMV PRO/DUB are therefore mutually exclusive and compete for binding to the enzyme. This regulates the dual PRO and DUB activities, both in time (proteolytic maturation of the polyprotein at early stages of infection, then regulation of the 66K RdRp amount in later stages) and in space (within the cytoplasm where the polyprotein is translated; then within the viral replication complexes where the viral RNA genome is replicated).

**Discussion**

Ub is a small molecule that interacts with many very different partners. Despite the wide variety of structural folds and functions encountered in ubiquitin-binding proteins (UBPs), Ub interacts with most of them through the same surface(s). In most of the Ub:UBP complexes, Ub engages a canonical protein interaction site known as the “hydrophobic Ile44 patch” (6, 52). A second hydrophobic patch of Ub, centered around Ile36, can also be targeted by UBPs (6). While the Ile44 patch is a well-known hot spot used by Ub to interact with its partners, fewer studies report an Ile36-based interface (44, 53). In addition, growing evidence shows the importance of Leu8 in UBP binding. Leu8 is located between the two hydrophobic patches and is usually considered to be part of the Ile44 patch (6, 52). However, Leu8 is located in a flexible loop that can undergo conformational changes (42, 43), shifting from a “loop-out” to a “loop-in” conformation (44). In turn, Leu8 can be displaced from the Ile44 patch to become a component of the Ile36 patch (44).

**An unusual mode of ubiquitin binding**

In TYMV PRO/DUB-Ub complex, Ub engages not only both of its two hydrophobic patches simultaneously, but also the loop that comprises Leu8 (Figure 2) a mode of binding without precedent thus far (see below). In order to score the relative importance of the residues interacting with the two hydrophobic patches, we designed and assayed non-conservative TYMV PRO/DUB mutations aimed at disrupting the binding interface. Mutation of residues that interact with Ile44 patch (E759A, N760A, T763A) had a mild effect on DUB activity (Figure 3A), probably because of their contribution in Ub recognition. Altering the central residue (mutation R844A, Figure 3A) in the interaction of TYMV PRO/DUB with Ub Ile36 patch dramatically improved DUB activity, a result which likely reflects improved Ub binding, as confirmed by molecular dynamics simulations. This binding interface thus appears far from
optimal for ubiquitin binding. Regarding the motif that interacts with Ub Ile8, we had previously shown the critical role played by TYMV PRO/DUB Ile847 in Ub recognition (28, 34). The crystal structure of the TYMV PRO/DUB-Ub complex presented here establishes that Ile847 and Phe849 engage in strong hydrophobic contacts with Ub Leu8 and Thr9 from the flexible loop (Figure 2G). In addition, this loop adopts a position where Leu8 no longer belongs to any hydrophobic patch but instead forms a distinct hydrophobic motif (Figure 2G,H). Altogether, our results show that the primary determinant of the TYMV PRO/DUB:Ub interaction is neither centered around Ile44 as usually observed, nor around the Ile36 patch. Instead, Leu8, located between the two hydrophobic patches, directly interacts with TYMV PRO/DUB Ile847, located between the two polar patches that sense the Ub patches. The Leu8:Ile847 pair therefore makes the major contribution to this interaction. Leu8 could thus be a major determinant in Ub involved in sensing its partners (43, 44, 54).

It is interesting to compare how Ub binds to different viral DUBs, including those that have the dual PRO and DUB activities. The other viral OTU DUBs for which the structures of complexes with Ub are available are encoded by CCHFV (55), DUGV (56) and EAV (25). The EAV PLP2 is an interesting case. It is an OTU PRO/DUB like TYMV’s. Furthermore EAV belongs to the order Nidovirales that also includes Coronavirus, members of which have caused three deadly epidemics in the 21st century including the current COVID-19 pandemic (57). All Nidovirales encode several proteases, at least one of which is a papain-like protease that doubles up as a DUB (58). Yet in Coronavirus this PRO/DUB does not belong to the OTU family as in EAV, but to the ubiquitin-specific protease (USP) family (7). This illustrates the capability of RNA viruses to acquire multiple cellular genes for the same function. It also underlines the major argument in favor of the view that TYMV PRO/DUB is a modified cellular DUB that secondarily acquired its processing protease function: it belongs to a family (OTU) of strict DUBs with no PRO activity, the only exceptions being a few viral OTU DUBs with dual PRO/DUB activity. We include in our structural comparison the USP PRO/DUBs encoded by the Coronavirus SARS-CoV (41), MERS-CoV (59) and MHV (unpublished structure). The comparison is also extended to cellular DUBs of the OTU family. We use cellular OTU DUBs from yeast (60) and human (61). The mode of interaction of Ub with TYMV PRO/DUB is thus seen to be very divergent (Figure 6). In all cases, other viral DUBs interact with the body of Ub only through its Ile44 patch (Supplementary Figure 3A). The Leu8-loop of Ub is most often found in the “loop-out” conformation (Supplementary Figure 3 insets), Leu8 being consequently part of Ile44 patch, including for cellular OTU DUBs. In the viral complexes with MHV PLP2, EAV PLP2 or CCHFV vOTU, the Ub’s Leu8-loop occupies the intermediate position observed in TYMV PRO/DUB-Ub complex (Supplementary Figure 3A insets). The loop adopts this intermediate position in all other crystal structures of complexes involving CCHFV vOTU and Ub (62, 63) (data not shown). These comparisons show that Ub Leu8 usually belongs to the Ile44 patch but also can be located between the two Ub hydrophobic patches to contact its partner. This intermediate position is found regardless of the enzyme considered, i.e. either a dual PRO/DUB or a DUB, either of viral or of cellular origin. Therefore, the function of Leu8 is not a hallmark of a DUB family but a specific feature of some enzymes, such as TYMV PRO/DUB.

Superimpositions of TYMV PRO/DUB with cellular OTU DUBs show that yeast OTU1 and human OTUD2 also interact simultaneously with the two hydrophobic patches of Ub (Supplementary Figure 3A), but engage mainly hydrophobic residues, together with one charged residue that structurally resembles Arg844 of TYMV PRO/DUB, i.e. E203 in yOTU1 or Arg245 in hOTUD2 (Supplementary Figure 3). From an evolutionary point of view, TYMV PRO/DUB appears to be a cellular OTU DUB that has acquired a PRO function by retaining the clamp that holds Ub, but losing important hydrophobic residues that interact with the two hydrophobic patches of Ub. This produces an enzyme with low DUB activity.

The low DUB activity of TYMV PRO/DUB may be an evolutionary compromise that ensures proper viral replication

TYMV PRO/DUB exhibits a significant but low deubiquitinase activity, its catalytic efficiency $k_{app}$
(which approximates $k_{\text{cat}}/K_m$) being around $2.5 \times 10^3 \text{ M}^{-1}\text{s}^{-1}$ (24, 28, 34), which is 10- to 1000-fold lower than that of other DUBs such as CCHFV vOTU, EAV PLP2, or MERS-CoV or SARS-CoV PLpro (25, 55, 56, 64). Several non-mutually exclusive hypotheses can be proposed that explain this low activity.

First, the crystal structure of TYMV PRO/DUB showed that the protein structurally belongs to the OTU superfamily of DUBs, but displays a peculiar active site. Indeed, TYMV PRO/DUB has no Asp or Asn that are usually part of the catalytic triad of OTU DUBs in combination with a Cys and a His, nor any oxyanion hole to stabilize the thioester intermediate of the catalytic mechanism (34). This results in an altered active site and explains the low DUB activity. In some cases, one or several catalytic functional groups are provided by the substrate, restoring a functional active site, a phenomenon called substrate-assisted catalysis (65). The crystal structure of the TYMV PRO/DUB-Ub complex presented in this work shows that Ub does not supply any residue that would restore a complete DUB active site. However, since Ub is located in the P side of TYMV PRO/DUB, it cannot be ruled out that the third residue may be provided by the substrate positioned in the P' side of the enzyme, i.e., the polyubiquitinated polymerase.

Second, the TYMV DUB activity is measured in vitro with a recombinant PRO/DUB domain and a single Ub molecule, while in vivo the enzyme is present in large macromolecular assemblies. Many deubiquitinases possess additional domains, built around a structurally conserved DUB scaffold, that are involved in substrate specificity and regulation of DUB activity. Additional protein domains of the TYMV replication protein may interact with the PRO/DUB domain and/or its substrate and thus contribute to the regulation of TYMV DUB activity. DUB activity in vivo is in fact carried by the 98K protein (24), a large multi-domain protein that comprises both the MT and PRO/DUB domains, separated by a region harboring the chloroplast targeting domain (CTD) (66) and a proline-rich region (PRR). This domain organization is similar to that found in the C-terminal part of nsP2 protein of numerous alphaviruses. This consists of an N-terminal protease subdomain (nsP2pro) and a C-terminal subdomain with a methyltransferase fold (MT-like), connected by a long loop. Crystal structures of the C-terminal part of nsP2 protein of Chikungunya virus (CHIKV) and Venezuelan equine encephalitis virus (VEEV) show several intramolecular interactions between the two subdomains and the involvement of the MT-like subdomain in nsP2pro function (51, 67, 68). Indeed, the nsP2pro active site is located at the interface between the two subdomains and its accessibility is regulated by the interdomain loop. Moreover, the MT-like subdomain actively participates in nsP2pro’s substrate recognition and binding. Although the nsP2pro subdomains of CHIKV and VEEV do not display DUB activity, these findings illustrate the opportunities for protease regulation inherent to the inclusion of the TYMV PRO/DUB domain in a larger protein. Crystal structures comprising full-length polyubiquitinated 66K and/or 98K protein should help to understand the role of the substrate and/or the other domains of 98K protein in TYMV DUB activity.

Third, Arg844 may contribute to the protease activities of TYMV PRO/DUB as it forms a minor contact to the substrate in the PRO/DUB:PRO complex (Figure 5CD). Analyses of the PRO\HEL and HEL↓POL cleavages in vivo indicate that processing of the polyprotein is not affected by the R844A mutation (Supplementary Figure 5). However, we cannot rule out that proteolytic activity also occurs on presently unknown cellular substrates, as with other viral proteases. In such a case, the presence of the non-optimal Arg844 to contact Ub could be a tradeoff to bind efficiently other substrates with unrelated surfaces.

Fourth, and this is our preferred hypothesis in view of the lack of discernible effect on processing protease activity of R844A, low DUB activity may be a fine-tuned feature of TYMV PRO/DUB. Indeed, viral proteases are usually highly specific enzymes whose activity depends not only on the particular sequence of a cleavage site, but also on the remainder of the substrate. The cleavage site, often located in a solvent-exposed flexible loop, is commonly recognized by proteases in an extended conformation that favors its perfect positioning into the catalytic cleft (69). Substrate specificity is
ensured by specific interactions between the body of the substrate and the enzyme. While Ub is usually recognized through its Ile44 hydrophobic patch only (6, 52), we show in this work that TYMV PRO/DUB, unlike other viral PRO/DUBs, has maintained the cellular OTU DUB mode of recognition, involving the two hydrophobic patches on Ub simultaneously. Nevertheless, the interacting surface is both small and suboptimal in its composition, as shown by mutants that improve the DUB activity (Figure 3A). Since recognition of the substrate body is usually the driving force that allows enzyme/substrate recognition, this observation is puzzling. Residues involved in Ub recognition are not conserved in the Tymoviridae family (Figure 1B and Supplementary Figure 4). We hypothesize that maintaining a low DUB activity may be an evolutionary compromise to ensure proper viral replication. Indeed, although it is initially produced in amounts equimolar to 98K and 42K (the two other products of the 206K polyprotein maturation process), 66K displays a transient accumulation in the viral replication cycle (33). The 66K polypeptide is degraded at a late stage of viral infection by the ubiquitin proteasome system (UPS) through polyubiquitination (14). Nevertheless, by harboring a DUB activity, TYMV possesses a rescue system to avoid complete degradation of the 66K protein. This ensures maintenance of the appropriate level of polymerase and safeguards efficient replication of the TYMV genome. This level may be reached with low DUB activity. In the case of TYMV, too high a level of 66K is actually detrimental to replication. We suggest that a finely tuned DUB activity may be a general feature of viruses that use deubiquitination to adjust the amount of (a) protein(s) that is/are critical for their replication. Indeed, Lei et al. found that the MERS-CoV PLpro Ub-binding surface is likewise suboptimal and nicely discussed the functional implications of this finding (40). Future experiments will be aimed at determining whether this applies also to the case of TYMV.

Materials and methods

Covalent coupling of Ub to TYMV PRO/DUB

TYMV recombinant PRO/DUB fused to an N-terminal 6His-tag (34) was expressed and purified as previously described (28), and diluted to a final concentration of 5 mg/mL in a fresh buffer composed of 10 mM Tris-HCl, 350 mM ammonium acetate, 1 mM DTT, pH 8. A C-terminally modified vinyl methyl ester variant of HA-tagged ubiquitin (HA-UB-VME) was prepared in 50 mM sodium acetate pH 4.5 essentially as previously described (36). In order to adjust the pH, HA-UB-VME was then diluted 10 fold in binding buffer (50 mM Tris-HCl, 150 mM NaCl, pH 8) and incubated 10 min at 25°C. Conjugation of both proteins was achieved by adding a two-fold molar excess of 6His-PRO/DUB to HA-UB-VME followed by incubation at 25°C during 30 min. Unreacted proteins were removed by size exclusion chromatography on a HiLoad 16/600 Superdex 75 pg column (GE Healthcare) with 50 mM Tris-HCl, 500 mM NaCl, pH 8 as elution buffer. Elution fractions were verified by 16.5% Tris-Tricine SDS-PAGE and those containing pure 6His-PRO/DUB-HA-Ub complex were pooled and dialyzed over-night at 4°C against binding buffer. Covalent complex was then concentrated to 24 mg/mL using ultrafiltration on Amicon Ultra with a cut-off of 10 kDa and frozen in liquid nitrogen for storage at -80°C.

Crystallization of 6His-PRO/DUB-HA-Ub complex

Crystallization conditions of 6His-PRO/DUB-HA-Ub complex were screened by a robot using commercial kits from QIAgen and the sitting-drop vapour-diffusion method. Some promising conditions were manually reproduced at 19°C in larger drop volumes (1 µL of 15 mg/mL complex solution plus 1 µL crystallization reagent equilibrated against a 0.5 mL reservoir volume) using the hanging-drop vapor-diffusion setup. Few crystals appeared after several months in 20% PEG-20K, 0.1 M MES-NaOH pH 6.5. Prior to data collection, these crystals were harvested, transferred to a cryo-protectant solution (21% PEG-20K, 0.1 M MES-NaOH pH 6.5, 20% glycerol) and flash frozen in liquid nitrogen.

Data collection and processing and structure determination

Data collection was performed at beamline PROXIMA-1 at French synchrotron SOLEIL. Only one crystal showed correct diffraction and a complete data set could be collected at 3.66 Å. Data
were processed and scaled with XDS (70). Since structures of individual TYMV PRO/DUB and human ubiquitin were available, the structure of 6His-PRO/DUB-HA-Ub complex was solved by molecular replacement. Calculation of Matthews coefficient (71) suggested two complex molecules in the asymmetric unit and several molecular replacement protocols were tested with Phaser (72). The good solution consisted to first locate one complex molecule using a C-terminally truncated version of TYMV PRO/DUB (PDB code 5LW5 chain A (28)) and ubiquitin (PDB code 1UBQ (73)) as search models, and second to use the resulting solution as an input to find the second complex molecule. The electron density was of sufficient quality to manually rebuild the model in COOT (74). Initial stages of refinement were done with REFMAC (75) and then with PHENIX (76). Due to the low resolution, no solvent molecules could be modeled. The final model thus consists of two TYMV PRO/DUB molecules (residues 732-876 in chain A, residues 732-876 in chain C; His-tags could not be modeled) and two HA-Ub-VME molecules (residues 1-76 in chain B, residues 1-76 in chain D; HA-tags could not be modeled). Data processing and refinement statistics are listed in Table 1.

Molecular dynamics simulations and structure visualization

Molecular dynamics simulations of a TYMV PRO/DUB:Ub product state complex and of a R844A mutant thereof were performed using the AMBER16 program suite (77) with the ff14SB force field. We noted in preliminary simulations comprising residues 732-876 of PRO/DUB that the first residues tended to interact with Ub, but this seemed to be influenced by the +1 charge spuriously added to L732 by taking it as the N-terminus. Thus we simulated a complex made of an N-acetylated PRO/DUB 727-876 (residues 727-731 were modeled stereochemically) and all ubiquitin residues (1-76). The LEaP program was used for preparation of the systems. Hydrogen atoms were added with default parameters. Complexes were neutralized with K+ cations and immersed in an explicit TIP3P water box with a solvation shell at least 12 Å-deep. The systems were then minimized and used to initiate molecular dynamics. All simulations were performed in the isothermal isobaric ensemble (p=1 atm, T=300 K), regulated with the Berendsen barostat and thermostat (78), using periodic boundary conditions and Ewald sums for treating long range electrostatic interactions (79). The hydrogen atoms were constrained to the equilibrium bond length using the SHAKE algorithm (80). A 2-fs time step for the integration of Newton’s equations was used. The nonbonded cutoff radius of 10 Å was used. All simulations were run with the SANDER module of the AMBER package. Each complex was simulated for 50 ns twice and the trajectories sampled every 10 ps. Analysis of the trajectories with cpptraj showed convergence within the first 5 ns as judged by stabilization of rmsd. The last 45 ns were kept for analyses.

All simulation trajectories and crystal structures were visualized and structural figures were made with PyMOL (81) (http://www.pymol.org). PyMOL was also used to mutate R844 to Ala prior to system preparation.

Deubiquitination assay in vitro

Point mutations were introduced in the bacterial vector encoding TYMV PRO/DUB (34) by using the quick Change II site directed Mutagenesis (Agilent) strategy. Recombinant PRO/DUB proteins were produced and purified as described previously (34). Prior to deubiquitination assay, purified proteins were dialyzed overnight at 4°C in buffer 50 mM HEPES-KOH, 150 mM KCl, 1 mM DTT, 10% glycerol, pH 8.0, adjusted to a concentration of 100 µM and kept at -80°C until use. The fluorogenic substrate Ub-AMC (Boston Biochem) dissolved in DMSO was diluted in assay buffer (50 mM HEPES-KOH, 10 mM KCl, 0.5 mM EDTA, 5 mM DTT, 0.5% Nonidet-P40, pH 7.8). DUB activity was assessed at room temperature in a Hitachi F2000 spectrofluorometer in assay buffer with a final concentration of DMSO adjusted to 2% to match the DMSO concentration in the highest Ub-AMC concentration assays. Reactions were initiated by the addition of enzyme to the cuvette and the rate of substrate hydrolysis was determined by monitoring AMC-released fluorescence at 440 nm (excitation at 380 nm) during 10 min. Enzyme concentrations were 125 nM for WT PRO/DUB and mutants. In order to determine the apparent $k_{cat}/K_m$ ($K_{app}$), the substrate concentration was kept at a concentration below 0.5 µM where the initial velocity is linear in substrate concentration and $K_{app}$
values were then determined according to the equation $V / [E] = K_{app} / [S]$ as described previously (24). Depending on the batch of Ub-AMC, the DUB activity of the WT enzyme displayed variability, with $K_{app}$ varying between $2,388 \pm 398 \text{ M}^{-1}\text{s}^{-1}$ and $2,824 \pm 213 \text{ M}^{-1}\text{s}^{-1}$. Hence, the activity of the WT protein was measured as a reference for each independent experiment, and $K_{app}$ of mutant proteins were normalized to that of the WT protein measured simultaneously. All experiments were performed at least in duplicates and data were expressed as the means and standard deviations of these independent experiments.

**Data availability:** The structure presented in this paper has been deposited in the Protein Data Bank (PDB) with the following code: 6YPT. All remaining data are contained within the article.

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**Footnotes.** The character “:" is used for a non-covalent complex (PRO/DUB:PRO), “-” for a covalent complex (PRO/DUB-Ub).

**Table 1.** Data collection and refinement statistics.

| Data collection       |          |
|-----------------------|----------|
| Wavelength (Å)        | 0.978    |
| Space group           | P 1 2 1 1|
| Unit cell (Å, °)      | 37.93 51.86 125.25 90 98.37 90 |
| Resolution range      | 37.46 - 3.679 (3.81 - 3.679) |
| Total reflections     | 17,522   |
| Unique reflections    | 5303     |
| Multiplicity          | 3.3 (3.3) |
| Completeness (%)      | 97.96 (85.85) |
| Mean I/sigma(I)       | 4.14 (0.97) |
| Wilson B-factor       | 106.81   |
| R-merge               | 0.215 (1.211) |
| CC1/2                 | 98.5 (41.5) |

| Refinement            |          |
|-----------------------|----------|
| Reflections used in refinement | 5,290 (467) |
| Reflections used for R-free     | 265 (24)  |
| R-work                 | 0.2057 (0.3269) |
| R-free                 | 0.2837 (0.3611) |
| RMS (bonds)            | 0.004    |
| RMS (angles)           | 0.51     |
| Ramachandran favored (%) | 92.13   |
| Ramachandran allowed (%) | 7.87    |
| Ramachandran outliers (%) | 0.00   |
| Rotamer outliers (%)   | 3.00     |
| Clashscore             | 7.30     |
| Average B-factor       | 175.69   |
| macromolecules         | 175.62   |
| ligands                | 190.88   |
| Number of TLS groups   | 4        |

Statistics for the highest-resolution shell are shown in parentheses.
**Figures**

**Figure 1.** Overall structure of the covalent TYMV PRO/DUB-Ub complex. A. Crystal structure of the covalent TYMV PRO/DUB-Ub complex. TYMV PRO/DUB is represented as molecular surface, with the N-ter lobe colored in yellow, the central lobe in magenta and the C-ter lobe in green. The enzyme’s catalytic dyad, composed of Cys783 and His869, is indicated in red. HA-Ub-VME is shown in ribbon diagram and colored in orange. Residues Ile36 and Ile44 are displayed in ball-and-sticks format and colored in cyan. Ubiquitin residues are labeled in italics and underlined. B. Sequence alignment of polyprotein processing endopeptidases belonging to the Tymoviridae family. The sequence of TYMV PRO/DUB was aligned with enzymes encoded by Chayote mosaic virus (ChMV), Physalis mottle virus (PhMV), Eggplant mosaic virus (EMV), Dulcamara mottle virus (DuMV), Okra mosaic virus (OkMV) and Kennedya yellow mosaic virus (KYMV) as in Lombardi et al. (34). Alignment was performed by CLUSTALW (82), then edited and displayed with ESPript3 (http://espript.ibcp.fr (83)). White characters in red boxes indicate identity and red characters in white boxes indicate homologous residues. Secondary structures of TYMV PRO/DUB (PDB code 4A5U (34)) are indicated on top. Black stars indicate the residues of enzyme interacting with Ub Ile44 hydrophobic patch, black circles the residues interacting with Ub Ile36 hydrophobic patch and black triangles the residues interacting with the C-ter extremity of Ub.
Figure 2. Interactions between TYMV PRO/DUB and Ub. A. Close-up view of the Ub’s Ile44 patch. Both proteins are shown in cartoon and residues involved in the interaction are shown in stick. Proteins are colored as in Figure 1, with O and N atoms in red and blue, respectively. Hydrogen bond between Gln49 from Ub and Thr763 from TYMV PRO/DUB is show as dotted line. B,C. Analysis of interactions around the Ub’s Ile44 patch by molecular dynamics simulations. The distances between three pairs of residues were measured during 90 ns of production time in two simulations and their frequency were plotted. Hydrogen bonds and electrostatic interactions, grey; hydrophobic contact, black. D. Close-up view of the Ub’s Ile36 patch. Proteins and residues are represented and colored as in A. E,F. Analysis of interactions around the Ub’s Ile36 patch by molecular dynamics simulations. The distances between two pairs of residues were measured during 90 ns of production time in two simulations and their frequency were plotted, as in B,C. G. Close-up view around Leu8 of Ub. Both proteins are displayed in cartoon-loop with some side chains shown in stick. The cavity of TYMV PRO/DUB that fits Ub is highlighted by grey molecular surface of the enzyme. Three crystal structures of Ub were superimposed to compare the position of the loop encompassing Leu8: “loop-out” conformation (PDB code 1UBQ (73)), purple; intermediate conformation
(this work), orange; “loop-in” conformation (PDB code 2G45 (84)), blue. H. Overall view of the two polar loops of TYMV PRO/DUB that bind the two hydrophobic patches of Ub.
**Figure 3. In vitro** DUB activity of structure-guided mutants of TYMV PRO/DUB. A. DUB activity of recombinant TYMV PRO/DUB (WT and structure-guided mutants) was measured by a fluorescence assay using Ub-AMC as substrate. $K_{\text{app}}$ was determined according to the equation $V / [E] = K_{\text{app}} [S]$, where $V$ is the initial velocity calculated from the kinetic data, and [E] and [S] are the corresponding enzyme and substrate concentrations. Values were expressed as the percentage of that of WT protein. B,C. Behavior of residue 844 side chain (B) and of the catalytic dyad (C783 and H869) (C) was investigated by performing molecular dynamics simulations of the product state complex, using wild-type TYMV PRO/DUB or R844A mutant. The R844A mutant was generated by truncating the Arg side chain at Cβ to mimic an alanine. Distances were measured along the same 90 ns in two simulations as in Figure 2 (WT, red histograms) and along 90 ns in two simulations for R844A (black histograms). B. Distance between the side chains of TYMV PRO/DUB residue 844 (Cβ atom) and Ub Ile36 (Cγ1 atom) C. Distance between TYMV PRO/DUB Cys783 (Sy atom) and His869 (Nδ1 atom). The minor peak at 3.5 Å signals alignment of the catalytic dyad.
Figure 4. Interactions network at the C-terminal tail of Ub. A. Detailed hydrogen bonding between the last five residues of Ub and TYMV PRO/DUB. The C-terminal extremity backbone of Ub (including Arg72 to Gly-VME76) is represented as sticks. The residues of TYMV PRO/DUB involved in the interaction with Ub are displayed as sticks. Hydrogen bonds are shown as dotted lines. B. Global hydrophobic and electrostatic interactions network between the last five residues of Ub and TYMV PRO/DUB. Both proteins are shown in cartoon, with residues involved in interaction depicted in sticks. The overall coloring scheme is the same as that in Figures 1 and 2.
Figure 5. Comparison of the binding interfaces in the PRO/DUB-Ub and PRO/DUB:PRO complexes. Covalent and non-covalent complexes between TYMV PRO/DUB and Ub (this work) (A) or TYMV PRO/DUB from the PRO↓HEL cleavage site (PDB code 4A5U (34)) (C), respectively, are shown in surface representation. The enzyme TYMV PRO/DUB is colored in grey, the substrates Ub and TYMV PRO/DUB are colored in orange and pink, respectively. The interacting surfaces used by the enzyme to bind its substrates (B, Ub; D, PRO/DUB) are colored in cyan and are shown after rotation of the protein.
Figure 6. Ub binding mode with TYMV PRO/DUB and other viral and cellular DUBs.
Comparison of overall Ub binding mode for viral PRO/DUBs (black lettering), viral OTU DUBs (blue lettering) and cellular OTU DUBs (magenta lettering). Whether each enzyme belongs to the OTU or USP family is indicated. Crystal structures of Ub in complex with viral or cellular PRO/DUBs or DUBs were aligned with Pymol. In each case, Ub is displayed as molecular surface and colored in orange, while the enzyme is shown in green cartoon. The two hydrophobic patches of Ub are colored in cyan. Ile44 and Ile36 of Ub, at the center of the two patches, are shown in blue, and Leu8, located in a loop between the two patches, is highlighted in red. The conformation of the Ub’s Leu8-loop in each complex is compared to the classical “loop-out” and “loop-in” conformations, and to the intermediate conformation found in TYMV PRO/DUB-Ub complex (inset, see also Fig.2G). We choose for comparison PRO/DUBs encoded by SARS-CoV (PDB code 4MM3 (41)), MERS-CoV (PDB code 4RF1 (59)), MHV (PDB code 5WFI, unpublished structure) and EAV (PDB code 4IUM (85)) (black), and DUBs encoded by CCHFV (PDB code 3PHW (55)) and DUGV (PDB code 4HXD (56)) (blue). We also compared DUBs from yeast (PDB code 3BY4 (60)) and human (PDB code 4BOS (61)) (magenta).
Turnip yellow mosaic virus protease binds ubiquitin suboptimally to fine-tune its deubiquitinase activity
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