The structure and function of deubiquitinases: lessons from budding yeast

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Protein ubiquitination is a key post-translational modification that regulates diverse cellular processes in eukaryotic cells. The specificity of ubiquitin (Ub) signalling for different bioprocesses and pathways is dictated by the large variety of mono-ubiquitination and polyubiquitination events, including many possible chain architectures. Deubiquitinases (DUBs) reverse or edit Ub signals with high sophistication and specificity, forming an integral arm of the Ub signalling machinery, thus impinging on fundamental cellular processes including DNA damage repair, gene expression, protein quality control and organelar integrity. In this review, we discuss the many layers of DUB function and regulation, with a focus on insights gained from budding yeast. Our review provides a framework to understand key aspects of DUB biology.

1. Introduction

Ubiquitination is a reversible post-translational modification (PTM) that governs a wide variety of cellular processes including protein degradation and sorting, cell signalling and gene expression [1]. Ubiquitination of target substrates occurs through the sequential activity of three enzymes, a ubiquitin (Ub)-activating enzyme (E1), a Ub-conjugating enzyme (E2) and a Ub ligase (E3) [1–5]. This enzyme cascade directs the attachment of Ub chains onto target proteins—the length and specific configuration of Ub chains dictates their functional consequences [1–5]. The ubiquitination state of protein substrates is also determined by deubiquitinases (DUBs) which hydrolyse Ub-substrate and Ub–Ub isopeptide bonds [1–5]. DUBs have three major cellular functions: (i) the generation of free Ub moieties from linear fusions of Ub [1–5]; (ii) trimming of existing polyubiquitin chains; and (iii) reversal of Ub signalling by removal of Ub chains from target proteins [1–5]. Thus, like ubiquitination enzymes, DUBs are master regulators within the Ub system, and impinge on diverse cellular processes. Importantly, mis-regulation of DUB function has been associated with the onset or progression of numerous diseases, including some cancers and Alzheimer’s disease [6–9]. Owing to the importance of DUBs in eukaryotic cell biology, there has been a concerted effort to understand their structure, mechanisms of substrate and polyubiquitin chain-type recognition and modes of regulation.

Key insights into DUB biology have emerged from work in model organisms such as the budding yeast, *Saccharomyces cerevisiae*. The yeast model system offers several key advantages for studying DUB function, including a compact genome, the conservation of the proteome among eukaryotes and the ease of maintenance under laboratory conditions [10]. In addition, yeast is highly amenable to a variety of genetic manipulations and exists in stable haploid and diploid states. In this review, we focus on the current state of knowledge about the structure and function of yeast DUBs and discuss the implications of insights gained.
from work in yeast for understanding DUB function in more complex organisms such as humans.

2. Deubiquitinase families in *Saccharomyces cerevisiae*

The yeast genome encodes 22 putative DUBs that are subdivided into classes based on their sequence and structural similarity. The roster of DUBs encoded by the human genome is more extensive with approximately 100 DUBs that are divided into seven families: the ubiquitin-specific proteases (USP/UBP); ubiquitin C-terminal hydrolases (UCHs); ovarian tumour proteases (OTUs); JAB1/MPN/MOV34s (JAMMs); Josephins, zinc finger with UFM1-specific peptidase domain protein protease (ZUFSP) and the recently discovered MINDY. These families are further subcategorized according to their mode of catalysis: USPs, UCHs, OTUs and MINDYs are cysteine proteases while the JAMMs are metalloproteases. Five of the seven DUB families (USP/UBP, UCH, OTU, JAMM and MINDY) are conserved in yeast (figure 1) [11–13], and below we summarize our

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**Figure 1.** The yeast DUBs. Shown are schematics of the DUB proteins in *S. cerevisiae* categorized by family, illustrating key functional domains (legend at base of the figure). Any human homologues are noted to the right of the protein diagrams. The size of the DUB proteins is indicated on the left of the schematics and domain sizes are indicated according to scale. ‘aa’ refers to amino acids.
understanding of the functions and mechanisms of action of these conserved DUB families.

2.1. Ubiquitin C-terminal hydrolases

As noted above, the majority of yeast DUBs, including USPs, UCHs, OTUs and MINDY, are cysteine proteases, which in general have a catalytic triad comprising Cys, His and Asp/Asn residues [5]. The nearby His residue lowers the $p_K$ of the catalytic Cys facilitating a nucleophilic attack, with the Asp/Asn residue functioning to stabilize the catalytic His residue [5].

UCHs were the first DUB family to be structurally characterized. The UCH DUBs possess a catalytic domain spanning 230 amino acid residues that adopts a core fold and catalytic triad resembling papain [14]. UCHs frequently contain additional C-terminal extensions that may assist in substrate or target recognition [14–16]. In yeast, Yuh1 is the sole member of the UCH family and closely resembles UCH-L3, a small human DUB (figure 1) [17]. Yuh1 preferentially processes small leaving groups and is inactive against larger Ub fusions [17]. Yuh1 appears to play a key role in the processing of the Ub-like protein Rub1p/NEDD8, a function that is conserved in its human counterpart [17].

2.2. Ubiquitin-specific proteases

The USP/UBP constitute the largest family of DUBs, with 16 members in yeast and 56 members in humans (figure 1 and table 1). USP/UBPs have a core catalytic domain and often possess insertions or terminal extensions that assist in substrate or target recognition [1,25,26]. Although the USP/UBP catalytic domains vary considerably in size and sequence, they all adopt a highly conserved structure resembling a right hand with three subdomains, including a finger, palm and thumb [1]. A cleft formed between the palm and thumb subdomains forms the catalytic centre with the thumb and palm containing the catalytic Cys and His, respectively [1], while the finger subdomain interacts with Ub to facilitate its positioning in the catalytic centre. Substrate recognition by USP/UBPs typically involves either their variable sequence regions or scaffolds and substrate adaptors in multi-protein complexes [1]. The overall complexity and diversity seen in the USPs underlies their diverse substrate profile and cellular roles.

2.3. Ovarian tumour proteases

The OTU family of DUBs were identified based on their homology to the ovarian tumour gene (OTU) in fruit flies. In yeast, the OTU DUB family contains two members, Otu1 and Otu2, while in humans, there are 16 (figure 1 and table 1). Crystal structures of yeast Otu1, and several human OTUs, reveal a conserved catalytic OTU domain consisting of five β-strands sandwiched between helical domains that vary in size [1,5,27]. The active site of the OTU domain contains an unusual loop not seen in other thiol-DUBs, and may lack the catalytic Asp/Asn residue, such that several human OTU DUBs are predicted to be catalytically inactive [3,5]. OTUs display remarkable specificity for different poly-Ub-chain linkages. For example, the human OTU, OTUB1 is highly specific for K48-linked chains, while OTUD2 [1,5] shows specificity for atypical Ub chains such as K11, K27 and K33 [28,29]. Some evidence suggests that the substrate and target specificities of OTU DUBs are evolutionarily conserved. For example, the yeast orthologue of OTUD2, Otu1, shares substrate preference and specificity for atypical Ub-chain linkages with OTUD2, and both DUBs function in endoplasmic reticulum-associated protein degradation (ERAD) [28].

2.4. MINDY

The recently discovered MINDY family of DUBs is highly conserved, with a core catalytic domain distinct from other DUBs [29]. Structural analysis of the human MINDY-1 in complex with Ub revealed that the C-terminus of Ub sits in the MINDY-1 catalytic groove in a highly conserved hydrophobic pocket. Like many other DUBs, the L73 residue of Ub is required for this interaction and thus catalysis. MINDY-1 has a strong specificity for longer K48-linked Ub chains and prefers to trim Ub from the distal end of chains [30]. Several MINDY DUBs exist in humans, with only two family members in yeast, one of which, Miy1, has demonstrated catalytic activity (figure 1 and table 1). Although Miy1 is the yeast orthologue of MINDY-1, it does not discriminate between positions for Ub-chain cleavage and, unlike human MINDY-1, is capable of cleaving shorter Ub chains [30]. The Ub-binding domains of MINDY, motif interacting with Ub (MIU), are responsible for MINDY chain preference and are discussed in further detail below.

2.5. JAB1/MPN/MOV34 metalloenzymes (JAMM/MPN±)

JAMM DUBs are conserved metalloproteases with a Glu-x [N]-His-x-His-x[10]-Asp motif which coordinates binding of two Zn\(^{2+}\) ions [1,3]. Catalysis requires a nucleophilic attack by the DUB on the carbonyl carbon of the isopeptide bond in the Ub chain, mediated by an activated water molecule [3]. Rpn11, a proteasome-associated DUB, is the sole representative of the JAMM DUBS in yeast [31,32] (figure 1 and table 1). By contrast, the human genome encodes 14 putative JAMM DUBs.

3. Substrate recognition

DUBs select their targets through binding to a protein substrate that they deubiquitinate, enabling regulation of specific cellular pathways and processes, or by direct recognition of select Ub-chain types, allowing control of the abundance of distinct Ub linkages (figure 2). For example, several DUBs in the UBP/USP family bind target proteins through additional protein–protein interaction (PPI) domains and will cleave an Ub chain from a specific target regardless of chain architecture [33,34]. By contrast, other DUBs, as discussed in more detail below, display remarkable chain specificity and in some cases will cleave only one linkage type [28,29,33,34].

3.1. Linkage specificity

Ubiquitination is a highly versatile PTM because it occurs as eight distinct types of multi-Ub chains, which convey unique structural and functional information. To form multi-Ub chains, Ub moieties are linked to the C-terminal Gly of
| DUB   | systematic name | localization | site of ubiquitination | site of phosphorylation | functions                                                                 | references |
|-------|-----------------|--------------|------------------------|-------------------------|---------------------------------------------------------------------------|------------|
| UBP1  | YDL122 W        | ER, cytoplasm| K378, K423, K524       | S154, S310, S311, S315, S319, S322, S324, S326, S329, S333, S334, S336, S360, S364, S400, S668, T1106, T1337 | prERAD                                              | [18–22]    |
| UBP2  | YDR124C         | cytoplasm    | K1065, K980, K991      | S614, S876, S901, S907, S917, T825                                    | endocytosis, degradation of misfolded proteins                           | [18–20]    |
| UBP3  | YER151C         | cytoplasm    | K173, S242, S240, S239, S232, S235, S339, S341, S360, S364, S400, S668, T1106, T1337 | degradation of misfolded proteins, ribophagy, proteophagy, mitophagy, formation of stress granules, regulation of histone methylation | [18–22]    |
| DQA4  | YDR069C         | cytoplasm, endosome | S414, S443, S498, T500, T502 | multi-vesicular body formation, ubiquitin recycling                      | [18–20]    |
| UBP5  | YER144C         | bud neck     | S316, S319             | unknown                                           | [18]        |
| UBP6  | YFR010 W        | nucleus, cytoplasm | K190, K242, K253, K308, K378 | S298, S383, S430, T389 | proteasome-mediated protein turnover                                       | [18–20,23]|
| UBP7  | YIL156 W        | cell periphery| K549                   | S155, S193, S235, S235, S486, S488, S582, S863, S868, S924, S925, T120, T113 | endocytosis                                             | [18–20,22]|
| UBP8  | YMR223 W        | nucleus      | K45                    | S45, S54, SSS, S640                                           | regulation of transcription                                             | [18,23]    |
| UBP9  | YER098 W        | cytoplasm    | K45                    | S528, S529, S60, S620, S63, T383                              | regulation of ribosome biogenesis, chromatin organization and cell cycle | [21,22,24]|
| UBP10 | YNL186 W        | nucleus and nucleus | S228, S229, S323, S327, S326, T507, T526 | unknown                                           | [18,22]    |
| UBP11 | YKR098C         | mitochondria | S1160, S1179, S12, S1204, S1226, S1252, S400, S484, T1113, T1202, T1227, T26, T486 | unknown                                           | [18–20]    |
| UBP12 | YIL197 W        | nucleus and cytoplasm | K469, K571 | S198, S217, S337, S339, S340, S354, S356, S357, S45, S465 | unknown                                           | [18–21]    |
| UBP14 | YBR058C         | nucleus and cytoplasm | K132, K170, K398, K500, K670, K770 | ubiquitin recycling | [18]        |
| UBP15 | YMR304 W        | cytoplasm    | K1127, K1143, K303, K508, K771 | endocytosis and cell cycle                                            | [18]        |
| UBP16 | YPL072 W        | mitochondria | unknown               | unknown                                           | [19]        |
| YNL1  | YJR099 W        | cytoplasm    | S134                   | ER-associated degradation                                         | [18,19]    |
| OTU1  | YFL044C         | nucleus and cytoplasm | K267 | S134 | unknown                                           | [18,19]    |
| OTU2  | YNL013C         | cytoplasm    | K100, K233             | unknown                                           | [18]        |
| RPN11 | YFR004 W        | nucleus and cytoplasm | K12, K205, K233 | S243, T262 | proteasome-mediated protein turnover, formation of proteasome storage granules | [18]        |
| MIY1  | YPL191C         | cytoplasm    | K348, K56              | S343                                               | [18–22]    |
| MIY2  | YGL082 W        | cytoplasm    | K162, K181, K352       | S187                                               | [18,19]    |
Figure 2. Mechanisms of substrate recognition by DUBs. (a) Overview of DUB recognition of Ub chains. A generic substrate ‘X’ with an attached Ub chain is illustrated on the left, with the proximal Ub being substrate bound, while the distal Ub is unmodified. The cartoons in the middle of the panel show that DUBs can recognize their substrates either through interaction with a specific target protein that they deubiquitinate or through interactions with the Ub chain itself. Schematics on the right illustrate endo or exo-cleavage activity associated with DUBs (the exo activity can originate at the distal or proximal end of a Ub chain). (b) Types of Ub chain linkages. The various types of Ub chain linkages are illustrated. The M1 and K63 chains, which adopt open conformations, and K6, K11, K27, K29, K33 and K48-linked chains, which adopt closed conformations, are grouped together. Mono-ubiquitination and branched Ub chains may also occur. (c–h) DUB mechanisms for substrate recognition. (c) Ub–S1 site interaction. DUBs that show little or no specificity for Ub chain linkages, such as Rpn11 and several USP/Ubp DUBs, contact Ub through an S1 site only. (d) Ub–S1–S1′ interactions. DUBs may contact Ub through both the S1 site and an S1′ site which can impart additional linkage specificity. For example, the RPN11-related DUB AMSH-LP achieves linkage specificity for K11-linked Ub chains through contacts made by its S1′ site. (e) S1′ site interaction with target. DUBs such as Ubp8 contain an S1′ site that recognizes a target protein, an interaction that does not involve Ub. (f) Ub–S1–S1′–S2 interactions. OTUD2 is an example of a DUB with an S2 domain in its catalytic site, which can accommodate longer K11-linked Ub chains. (g) DUB–Ub interactions outside of catalytic domain. Some DUBs such as MINDY-1 contain Ub-binding sites outside of their catalytic domain. The diagram illustrates the case of MINDY-2, in which tandem MIU (tMIU) domains work together to mediate the recognition of longer K48-linked Ub chains. (h) Multiple Ub-binding domains. IsoT achieves substrate specificity for K48-linked chains through several interactions with Ub chains using its four Ub-binding domains: ZnF-UBP, USP/UBP domain and two UBA domains, which are inserted into its catalytic domain.
another Ub protein via one of seven Lys residues (Lys6, Lys11, Lys27, Lys29, Lys33, Lys48, Lys63) or the amino terminus (Met1) (figure 2b) [35]. Poly-Ub chains have specific conformational and dynamic properties that are recognized by distinct Ub-binding domains; for example, K48, K6, K11 and K29-linked Ub chains are generally thought to adopt a compact conformation, while M1 and K63-linked Ub chains have an open conformation (figure 2b) [35]. These varied linkages help determine specific functions for Ub chains within the cell, including roles for the typical Lys48- and Lys63-linked chains in protein degradation and cell signalling respectively, with atypical chain linkages more likely to be involved in a wide variety of cellular processes including cell cycle regulation and transcription, among others [35].

A significant factor in the diversity of Ub-chain function reflects the selectivity of DUBs for Ub chains of varying lengths and linkages. For example, in yeast, the Ubp8 DUB strongly prefers the mono-ubiquitinated histone component, H2B, Ubp12 and Ubp15 favour K48-linked Ub chains [36–39], while the proteasome-associated Rpn11 DUB, shows little substrate preference [31,32]. All DUBs contact Ub through at least one Ub-binding site called the S1 site in the catalytic domain (figure 2) [31]. The S1 site is considered a major determinant of DUB specificity as it contacts 20–40% of Ub’s surface and contributes significantly towards correct positioning of the Ub C-terminal tail in the DUB’s catalytic core [3]. Other Ub contacts may occur through the so-called S1’ site of some DUBs, which further aids in correctly orienting the modified Lys or Met residue towards the catalytic centre (see below, figure 2c) [33,34]. Other Ub-binding sites (S2, S2’, S3, S3’ etc.) may also be present in the catalytic domain or on other Ub-binding domains in specific DUBs. Together, these diverse Ub-binding sites contribute to the mode of binding of an individual DUB to its substrate and as such its specificity [33,34].

3.2. Substrate specificity via the catalytic domain

One simple mechanism for DUB substrate specificity involves interactions with the proximal Ub on the target substrate, which determines the Lys linkage presented to the catalytic centre. This mechanism is beautifully illustrated by structural studies of Rpn11, a proteasomal-associated DUB that displays promiscuity towards Ub chains, and a related DUB, AMSH-LP, which displays specificity to K63-linked di-Ub chains (figure 2c,d) [31,32,40]. AMSH-LP forms extensive contacts with residues that neighbour K63 in the proximal Ub [40] only when K63 linkage occurs, thus ensuring that AMSH-LP is specific to K63-linked Ub chains (figure 2) [40]. Rpn11 does not contact the proximal Ub and instead appears to contact the distal Ub exclusively (figure 2) [31,32], a property that is thought to contribute to its lack of substrate specificity [31,32,40].

3.3. Substrate specificity via ubiquitin-binding domains

As previously mentioned, DUBs can contact Ub through sites outside of their catalytic domain, and we discuss key examples of this class of DUBs below.

3.3.1. Ubp14/IsoT

The Ubp14 DUB disassembles unanchored poly-Ub and recognizes K29, K48 and K63-linked Ub chains. The loss of Ubp14 function in S. cerevisiae results in defects in proteasomal degradation and the accumulation of unanchored Ub. While the mechanism of action of Ubp14 remains poorly understood, its general function and substrate preference is conserved in its human orthologue, IsoT, which has been studied extensively [41–44]. IsoT contains four putative Ub-binding domains, a ZnF-UBP domain, a USP/UBP domain and two UBA domains, which are inserted into the catalytic domain, and each of these domains form extensive interactions with different Ubs in a Ub chain [45]. The ZnF-UBP contains the S1’ site and as such recognizes the C-terminus of the proximal Ub, while the S1 pocket in the USP/UBP, the UBA1 and the UBA2 domains recognize the second, third and fourth Ubs in the chain, respectively (figure 2h) [45]. Interestingly, it appears that the ZnF-UBP and USP domain enable discrimination of K48-linked Ub chains, while the UBA domains cannot [44,45]. Taken together, these studies offer insight into how a DUB can use different Ub-binding domains to determine substrate specificity.

3.3.2. MINDY-1

Members of the MINDY family of DUBs are thought to recognize Ub chains through tandem MIU motifs (tMIU) that function cooperatively to mediate substrate recognition. Like the Ubp14 example above, the substrate specificity of the yeast MINDY DUB, MIY1, remains unexplored relative to its human orthologue, MINDY-1, which specifically binds K48-linked Ub chains with a preference for longer chains [30,46]. MINDY-1 contains two tandem MIU domains, tMIU1 and tMIU2, that are separated by a linker sequence (figure 2) [30,46]. tMIU2 imparts substrate specificity to MINDY-1 by interacting with three Ub moieties in a K48-linked chain through distinct side chains [46], while tMIU1 has no poly-Ub-binding capacity but instead appears to enhance the affinity of tMIU2 domain for Ub [46]. Recent crystal structures of the MINDY-1 tMIUs in complex with a cyclic K48-tetra Ub chain revealed that the K48-tetra Ub is wrapped around two helices [46]. In this arrangement, a single tMIU2 makes simultaneous interactions with three Ub moieties through three distinct binding sites (figure 2) [46]. The first is formed through hydrophobic interactions and hydrogen bonds between tMIU2 and the middle Ub in the Ub chain [46]. The second occurs between tMIU2 and the proximal Ub and is mediated through hydrophobic interaction between tMIU2 and the H44 hydrophobic patch of Ub. However, this interaction strategy may not be broadly used, as the residue mediating this interaction is poorly conserved among the tMIUs [46]. The third binding interface is formed between tMIU2 and the hydrophobic patch of the distal Ub [46]. The MINDY-1 structure further revealed that tMIU2 binds Ub in an orientation that is exhibited only by longer K48-linked Ub chains, thus explaining its linkage specificity (figure 2).

Taken together, the mechanism of substrate recognition by MINDY-1 demonstrates how Ub-binding domains can work in tandem to achieve substrate recognition and specificity [46].
3.3.3. Otu1/OTUD2

As previously discussed, the OTU DUBs display remarkable specificity for their target substrates. The two yeast OTU DUBs, Otu1 and Otu2, are both orthologues of human OTUD2. Otu1 cleaves atypical Ub linkages (K11, K27, K29 and K33), as well as K48-linked Ub chains [28]. Unlike the examples discussed above, crystal structures of Otu1 failed to reveal the mechanism by which Ub-linkage specificity is achieved, although other evidence suggests that Otu1 substrate specificity may be mediated through interactions with one of its targets, Cdc48, and its associated adaptors [28].

Like Otu1, human OTUD2 also cleaves atypical Ub linkages with a preference for longer K11 Ub chains, but differs from Otu1 in that it fails to act on K48 linkages [28]. OTUD2 contacts Ub chains using its S1 and S1’ sites which are located in its catalytic domain. The S1’ site appears to be particularly important for the activity of OTUD2 on K11 linkages, as mutations in this region abolish the K11 linkage specificity of the enzyme (figure 2) [28]. OTUD2 additionally binds Ub chains using an S2 site in its catalytic domain, in an orientation that favours longer K11 Ub chains (figure 2) [28]. While this site is conserved in higher eukaryotes, it is not conserved in S. cerevisiae, and whether Otu1 uses a similar mechanism to achieve substrate specificity remains to be determined [28].

4. Regulation of deubiquitinase activity

Because DUBs have a wide range of roles in the cell, a series of different regulatory mechanisms have evolved to tightly control DUB activity. In this section, we discuss how PTMs, regulatory domains and PPIs regulate DUB activity in yeast.

4.1. Substrate-mediated activation

Several DUBs have evolved allosteric mechanisms to regulate their activity. For example, structural analysis of UCH-L3, the human orthologue of the yeast DUB Yuh1, revealed that the active site undergoes significant conformational changes upon Ub binding. When unbound, UCH-L3 contains a disordered loop that covers its active site, which is stabilized upon Ub binding into an α-helix which forms contacts with Ub. This loop is conserved in several DUBs including Yuh1 [44,45], suggesting that a similar allosteric mechanism may be a feature of a subset of DUBs. Comparisons of Ub-bound Otu1 to the OTU domain of human DUB Otubain 2 alone revealed a disordered loop, which is positioned to sterically clash with a bound Ub. In Ub-bound Otu1, this region forms an ordered β-strand, suggesting that Ub binding to Otu1 may induce a conformation change that stabilizes the disordered region in a β-strand, removing this steric hindrance [47].

Another common mechanism of DUB regulation involves the catalytic residues of DUBs adopting a non-active state. For example, binding of Ub to MINDY-1, the human homologue of Miy1, induces the movement of its Cys loop and allows the catalytic residues to adopt an active conformation. When not bound to Ub, the catalytic His residue in MINDY is flipped away from the catalytic Cys residue, producing an inactive state [30]. Whether this mechanism is conserved in Miy1 is currently unknown.

4.2. Complex-mediated activation

Several DUBs are activated through complex binding, including the yeast DUBs Ubp8 and Ubp6. Ubp8 is part of the histone-modifying complex, SAGA, and is specifically responsible for the deubiquitination of mono-ubiquitinated histone component H2B chains [35–38]. Free Ubp8 is generally in an inactive state and activation is induced through interactions with three SAGA complex components, Sgf73, Sgf11 and Sus1. Sgf11 and Sgf73 use their ZnF domains to interact with two distinct regions of the catalytic domain of Ubp8 to promote an active catalytic core, while Sus1 functions to stabilize interactions involving Ubp8, Sgf11 and Sgf73 [35–38]. Strikingly, Sgf11 forms part of the Ubp8 catalytic lobe, creating a highly conserved extended interface with the Ubp8 catalytic domain that aids in target and substrate recognition [35–38]. Crystal structures of Ubp8 bound to a mono-ubiquitinated nucleosome suggest that Ubp8 may have evolved a histone-specific S1’ site in its catalytic domain [37] (figure 2e). The Ubp8 complex with mono-ubiquitinated nucleosomes provides one of the first views of a DUB-non-Ub substrate complex and may serve as a model for how other mono-Ub targeting DUBs might recognize their substrates.

Ubp6 is a proteasome-associated DUB, whose activity is dramatically increased upon binding to the 26S proteasome [26]. Structural analysis of free Ubp6 revealed two surface loops blocking access to the catalytic site [26], leading to the hypothesis that binding to the proteasomal component Rpt1 induces an active conformational state by relieving this steric hindrance [26]. Human proteasome-associated DUBs, UCH37, USP14 and POH1, all demonstrate similar proteasome-induced activation.

5. Deubiquitinase functions in protein turnover and ubiquitin homeostasis

Selective spatio-temporal regulation of unwanted or damaged protein is central to cellular homeostasis, adaptation to stress conditions and development of multicellular organisms. Cells have evolved multiple Ub-dependent machineries and processes for protein degradation, including the proteasome, selective autophagy and endocytosis. Below, we review how DUBs function within these proteostasis bioprocesses (summarized in figure 3).

5.1. Proteasome-associated deubiquitinases

The proteasome is an approximately 2.5 MDa protein complex consisting of two large subcomplexes, a barrel-shaped core particle (CP) and the regulatory particle (RP) that binds to the axial end of the CP. The RP in turn comprises two subcomplexes of its own, the lid and the base [48], which is proximal to the CP and contains six ATPases that form a ring structure that abuts the CP (figure 3b). The proteolytic active sites of the proteasome are buried within the barrel-shaped CP [49], with access restricted by narrow gates at either end of the CP, which are modulated by the Rpt2 subunit of the base [50]. While this ‘restricted access’ arrangement may help prevent the unregulated destruction of intracellular proteins, it also imposes a requirement for substrate unfolding to pass through the narrow opening of
the translocation channel leading to the CP, effected by the six ATPases of the base [51]. The Ub chain(s) on substrates being targeted for proteasomal degradation impose a steric constraint preventing translocation through the narrow opening into the catalytic core and must be removed to facilitate substrate degradation. Two DUBs, Rpn11 and Ubp6, are responsible for DUB activity at the proteasome [52–59] and have an interesting functional interplay. Mutation of RPN11 causes decreased rates of substrate degradation, while the loss of UBP6 is reported to result in the stabilization of some substrates and destabilization of others, thus indicating distinct and shared function at the proteasome [50,53–57,60–62]. Ubp6 and its human orthologue Usp14 was recently shown to interact preferentially on Ub–cyclin B conjugates with multiple sites of ubiquitination in a chain-type independent fashion and this might explain the differential effects of Ubp6 on substrate degradation [59]. Ubp6 non-catalytically and allosterically affects the rates of proteasomal degradation by interfering with critical substrate deubiquitination by Rpn11, stimulating 20S gate opening, thus increasing access to the degradation chamber and enhancing the rates of ATP hydrolysis by the base ring [58,60,63,64]. While Ubp6 was believed to trim Ub chains at the proteasome, recent evidence indicates that like Rpn11, Ubp6 is also able to cleave Ub chains en bloc [59,60] (figure 3a).

Ubp6 and Rpn11 also play a key role in the biogenesis and stability of the RP subcomplex of the proteasome [65,66]. RP biogenesis is an elaborate process involving dedicated chaperones [67]. However ubiquitinated substrates, with or without substrate shuttling factors, compete with RP
subunits for interaction surfaces on precursor RPs that are en route to form a functional complex [66]. The deubiquitinating activity of proteasome-associated Ubp6 (and perhaps other DUBs) destabilizes such unproductive interactions thus promoting proteasome maturation [66] (figure 3b).

5.2. Deubiquitinase-mediated regulation of the core endocytic machinery and cargo turnover

Ub-dependent sorting of plasma membrane (PM) proteins is an important and conserved regulatory mechanism. In yeast, the Rsp5 E3 ligase targets PM proteins for ubiquitination (mono-ubiquitination or K63-linked polyubiquitination) and subsequent internalization by endocytosis (figure 3c). The resulting vesicles containing ubiquitinated proteins then mature into multi-vesicular bodies before their contents are degraded after fusion with the vacuole [68]. Endocytosis requires the coordinated actions of more than 60 different proteins at the PM [69,70] and several core endocytic adaptors, including Ent1, Ent2, Ede1 and Sla1, harbour Ub-binding domains or interaction motifs [71] and many others are ubiquitinated by the Ned4-family ligase, Rsp5 [72–76]. Two DUBs, Ubp7 and Ubp2, deubiquitinate Ede1, which is functionally important as ubp2Δubp7Δ cells exhibit extended endocytic coat lifetimes and abnormal trafficking of endocytic proteins to early endosomes. Thus, DUBs facilitate initiation of endocytosis and subsequent recruitment of the endocytic coat [77].

The Rsp5 ubiquitin ligase also targets defective PM proteins for degradation [78,79], and promotes the down-regulation of properly folded transporters and receptors in response to specific biological stimuli. A family of adaptor proteins termed arrestin-related trafficking adaptors (ART) target specific PM proteins by recruiting Rsp5, which modifies both the cargoes and the ARTs (figure 3c). Thus, ARTs and Rsp5 together provide a cargo-specific quality control pathway that mediates endocytic downregulation of diverse PM proteins in response to specific stimuli [80], maintaining the integrity of PMs.

The Ubp2 DUB interacts with Rsp5 via the adaptor protein Rup1 [81], to regulate Rsp5 activity at multiple levels. First, Ubp2 positively regulates Rsp5-mediated ubiquitination signalling by removing the inhibitory auto ubiquitination of Rsp5 [82]. Second, Ubp2, together with Ubp15, deubiquitinates ARTs, thus preventing their degradation via the proteasome (figure 3c). Third, Ubp2 deubiquinates Rps5 substrates at the PM and thereby antagonizes the degradation of Rps5 substrates both in vivo and in vitro [81,83,84]. Which of the opposing effects that Ubp2 has on Rps5 activity dominates to determine whether a substrate is degraded is probably substrate-dependent and modulated by other cellular factors or stress conditions.

5.3. Deubiquitinase regulation of multi-vesicular body formation and function in ubiquitin homeostasis

Ubiquitinated PM protein cargoes are processed through the endocytic pathway through a complex system that involves the ESCRT 0–III complexes which bind and sort ubiquitinated substrates into multi-vesicular bodies (MVBs) that ultimately fuse with the vacuole [85]. However, prior to being engulfed in the MVBs, Bro1, an ESCRT III adaptor protein, recruits the DUB Doa4 to endosomes, and stimulates its enzymatic activity resulting in deubiquitination of the targets aiding in the formation of MVBs [86–89]. MVBs in turn fuse with the vacuole to empty their contents and facilitate degradation of their cargo in the vacuole [86–89] (figure 3d). This example illustrates a key role for a DUB in determining the cargoes of the MVBs and thereby their degradation.

The endosome-associated Doa4 DUB discussed above, as well as the proteasome-associated DUBs Ubp6 and Rpn11, also have major roles in trimming and/or en bloc removal of Ub chains from target substrates, thereby promoting the recycling of Ub [90,91]. In logarithmically growing cells, Ubp6 exists in a mixed population of: (i) monomers; (ii) unanchored chains; and (iii) anchored chains of different linkages on substrates [3,92]. Given the short half-life of Ub (2 h) [93], recycling of Ub from unanchored and anchored polyubiquitin chains is key for restoring the ubiquitination capacity of cells, particularly under stress conditions such as heat shock [41,54,90]. Ub depletion leads to specific transcriptional upregulation of UBP6 as a compensatory mechanism to replenish Ub levels by increasing the proteasomal capacity to deubiquitinate its substrates prior to degradation [94]. Similarly, under heat shock conditions that necessitate increased Ub recycling, DOA4 transcription is upregulated. Doa4 activity is further enhanced by selective downregulation of a Doa4 inhibitor, Rfu1, both at the transcriptional level and at the protein level by Rsp5-mediated selective degradation by the proteasome (figure 3, inset) [87,90,95]. Thus, the Ubp6 and Doa4 DUBs regulate Ub recycling using a highly coordinated and regulated circuitry (figure 3, inset). Other DUBs also contribute to Ub homeostasis. In particular, Ubp14 cleaves unanchored polyubiquitin chains to replenish cells with monomers of Ub [41], while Ubp3, Ubp8 and Ubp10 contribute to recycling of ubiquitin from polyubiquitin chains, though the significance of their action in Ub homeostasis remains poorly understood [90,96].

6. Deubiquitinase-mediated regulation of protein quality control

As discussed above, proteotoxic stress caused by heat or other environmental insults induces global protein unfolding/misfolding [97] which causes a profound and multi-component cellular response. First, quality control systems mediated by chaperones and the Ub proteasome or autophagy systems are triggered, to either refold or degrade toxic non-native species. Alternatively, proteins may remain in an unfolded state, and accumulate as protein aggregates at discrete sites in the cell [98–102]. Finally, as a compensatory mechanism, translation is shut down upon severe heat shock to prevent accumulation of additional misfolded species [103–105]. Thus, living systems have evolved elaborate mechanisms for protein quality control (PQC), and DUBs contribute to these mechanisms in several ways, as outlined below.

6.1. Regulation of cytosolic protein quality control

Upon proteasomal inhibition, misfolded proteins tend to partition into two compartments—the juxta/intra-nuclear quality control compartment (JUNQ/INQ), and the insoluble protein deposit (IPOD) that is situated near the vacuole and colocalizes with the pre-autophagosomal structure
A variety of studies implicate DUBs in PQC involving these compartments. First, over-expression of Doa4 causes partitioning of a reporter for misfolded proteins, Ubc9ts-GFP, to the IP0D upon proteasomal inhibition, while the reporter normally localizes to both the JUNQ/INQ and IP0D compartments in wild-type cells [106]. This phenotype may reflect the direct action of Doa4 on the misfolded model substrate or may be an indirect consequence of the role of Doa4 in Ub recycling. Second, overproduction of Ubp3 also positively regulates the degradation of Ubc9ts-GFP and can offset a deficiency in Hsp70 chaperone activity both during heat shock and ageing [109]. Ubp3 and its cofactor Bre5 are recruited to heat-induced protein aggregate-stress granule complexes [103,104] and stationary phase-induced stress granules [110,111], consistent with a direct role of Ubp3 in quality control of functional and misfolded aggregates. Third, the E3 ligase Rsp5, with assistance from the Hsp70 co-chaperone Ydj1, recognizes interaction motifs in misfolded proteins upon severe heat shock that are otherwise hidden resulting in their K63 polyubiquitination [112]. However, K63-polyubiquitinated substrates are poorly recognized by the proteasome and thereby not efficiently degraded [113]. Ubp3, along with Ubp2, associates with Rsp5 to reverse the K63 polyubiquitin chains that Rsp5 builds on its misfolded substrates and instead promotes K48 polyubiquitination by an unknown E3 ligase, targeting them for proteasomal degradation [114]. Thus, the chain editing functions of DUBs can facilitate the degradation of misfolded proteins (figure 4a), in collaboration with other enzymes. The mechanism of recognition of the severity of protein aggregation/damage by DUBs, and the interplay with the Hsp70/Hsp40/Hsp104 tri-chaperone system and the proteasome, remains largely unexplored.

6.2. Regulation of quality control of tail-anchored proteins

While misfolded proteins represent a major class of PQC substrates, a substantial fraction of the secretome transiently resides in the cytosol before translocating into the ER, and represents another class of substrates that needs to be controlled by the proteostatic machinery. Cells have evolved a mechanism termed pre-insertional endoplasmic reticulum-associated degradation that involves recognition of the C-terminal hydrophobic glycosylphosphatidylinositol (GPI) anchoring sequences of non-inserted tail-anchored proteins that have a prolonged residence time on the cytosolic leaflet of the ER by the Doa10 E3 ligase, which targets them for degradation via the proteasome [115]. Ubp1, an ER-associated DUB [116], antagonizes the activity of Doa10 by deubiquitinating its substrates and thus extending their residence time at the cytosolic leaflet of the ER [115]. Thus, the interplay between Doa10 and Ubp1 acts as a ‘timer’ determining the fate of secreted proteins (figure 4b, left panel).

6.3. Regulation of endoplasmic reticulum-associated protein degradation

Protein homeostasis in the ER is maintained by a quality control system called the ERAD pathway that retains misfolded proteins in the ER, and promotes their ultimate retrotranslocation into the cytosol, where they are polyubiquitinated, and degraded by the proteasome [117,118]. The recognition of a misfolded protein in the ER lumen is best understood for misfolded glycoproteins [117,119]. The N-linked glycan on these proteins is trimmed to generate α-1,6-mannose residue that is recognized by the multi-subunit HRD complex, which includes the Hrd1 ubiquitin ligase. Once a segment of the substrate reaches the cytoplasmic side of the ER membrane, it is polyubiquitinated by Hrd1 in concert with the E2, Ubc7 and its activator Cue1 [118]. A complex containing the Cdc48 ATPase is recruited to the Hrd1 complex via recognition of the polyubiquitin chain by its cofactor Ufd1/Npl4, and the polyubiquitinated substrate is then pulled through the central pore of Cdc48. The DUB Otu1 interacts with Cdc48 to trim the polyubiquitin chain (but does not completely remove the chain) facilitating further threading of the substrate through the central pore of Cdc48. Thus, Otu1 assists Cdc48 in the extraction of misfolded proteins across the ER membrane and targets the substrates that continue to be ubiquitinated for proteasomal degradation [120]. The relationship between Otu1 and Cdc48 provides another illustration of how DUB-mediated polyubiquitin chain editing plays a crucial regulatory role in coordinating protein turnover (figure 4b, right panel).

7. Deubiquitinase-mediated regulation of ribophagy and proteaphagy

Ribophagy is an autophagic mechanism in which the 60S ribosomal subunits are selectively degraded upon nitrogen starvation [121]. In normal conditions, the 60S ribosomal subunits are protected from degradation by ubiquitination of the Rpl25 component by the ribosome-associated E3 ligase, Ltn1 [122]. Upon nutrient limitation, the DUB Ubp3 deubiquitinates L25 and targets 60S subunits for degradation via autophagy in a Cdc48-Ufd3-dependent manner [123] (figure 5a). Hence, in this context, Ubp3 activity favours degradation of its substrate. Additionally, under conditions of nitrogen starvation, Ubp3 confers selectivity in the degradation of other factors involved in translation (eIF4GI, eRF3) and mRNA degradation (Dcp2, and Pop2) via autophagic and proteasomal degradation, respectively [124]. Therefore, by regulating the fate of both the translational and mRNA-degradation apparatus, Ubp3 seems to play a key role in reprogramming protein expression in response to nutritional stress.

Nitrogen starvation also induces selective autophagy of the both the RP and CP subunits of the proteasomes, a process termed proteaphagy [125,126]. Nitrogen starvation-induced proteaphagy is dependent on Atg1, a serine-threonine kinase and master regulator of autophagy as well as the core autophagy machinery [125,126]. However, degradation of the CP alone seems to be further fine-tuned by Ubp3 [126], placing Ubp3 at the crossroads of two major proteolytic pathways in the cell—proteasome-mediated degradation and autophagy (figure 5b). While the mechanistic details remain to be explored, these results highlight the pivotal roles for DUBs in maintaining protein homeostasis.
8. Roles for deubiquitinases in regulation of organelle morphology

8.1. Mitochondrial homeostasis

Mitochondria form a dynamic network that is constantly remodelled by their fusion and fission. While fusion promotes mixing of mitochondria, protects against loss of mitochondrial DNA and supports an optimal bioenergetic activity, their fission promotes their distribution and inheritance [127,128]. The mitofusin Fzo1 is a conserved dynamin-related GTPase that resides in the mitochondrial outer membrane and mediates the fusion of mitochondria. Guanosine triphosphate (GTP)-binding promotes Fzo1 homodimerization and further oligomerization occurs upon tethering of two mitochondria [127]. Subsequent GTP hydrolysis probably triggers a conformational change in Fzo1 allowing initial ubiquitylation of Fzo1 at K464, mediated by the SCF Mdm30 E3 ligase [129–132]. This modification induces Ub chain formation on K398 of a neighbouring Fzo1 molecule that further promotes outer membrane fusion by enhancing intermolecular interactions. The DUB Ubp12 deubiquitinates Fzo1 at K398 resulting in reduced oligomerization of Fzo1, thus antagonizing mitochondrial fusion. Fzo1 is degraded by the proteasome.
through ubiquitination at other lysine residues, whose modifications are antagonized by the deubiquitinating activity of Ubp2. Thus, by acting through two independent pathways, Ubp2 and Ubp12 regulate Fzo1 stability and oligomerization to control mitochondrial fusion and integrity [132] (figure 5c).

Recently, a method termed synthetic quantitative array technology was developed to identify modulators of selective autophagic degradation of mitochondria, a process termed mitophagy, on a genome-wide level [133]. The DUB, Ubp3, complexed with its cofactor, Bre5, was identified as a negative regulator of mitophagy. Additionally, the Ubp3–Bre5 complex, which is typically cytosolic, translocates dynamically to mitochondria under conditions that trigger mitophagy (target of rapamycin inhibition by rapamycin treatment) [133–135]. The molecular mechanism by which Ubp3–Bre5 exerts its influence on mitophagy is unknown.

### 8.2. Regulation of COPI and COPII vesicles

Trafficking of proteins and lipids between the ER and Golgi takes place through the COPI and COPII vesicles, respectively. Regulated polymerization of COPII coatomers, including Sec23 onto the ER deforms the lipid bilayer and results in formation of COPII-coated vesicles. Likewise,
COPI vesicles are formed at the Golgi by polymerization of COPI coatomers that include Sec27 [136]. Both Sec23 and Sec27 are ubiquitinated and targeted to the proteasome by the Rsp5 E3 ligase and the Cdc48 complex, and this activity is antagonized by the Ubp3–Bre5 DUB [137–139]. The biological importance of this DUB activity is illustrated by a massive expansion of the ER seen in ubp3Δ and bre5Δ mutant cells. Hence, efficient protein trafficking involves a regulatory balancing act between the E3 Rsp5 and the DUB complex, Ubp3–Bre5.

9. Deubiquitinase-mediated regulation of chromosome segregation

Cse4 is a centromere-specific histone variant, which contributes to the unique attributes of centromeric chromatin that enable attachment to the mitotic spindle. Mislocalization of Cse4 to regions outside the centromere is deleterious and causes aberrant chromosome behaviour and mitotic loss. Non-kinetochore Cse4 is ubiquitinated by the Psh1 E3 ubiquitin ligase and targeted for degradation via the proteasome. At kinetochores, the stability of Cse4 is modulated by the Ubp8 DUB [140]. In ubp8Δ cells, Cse4 accumulates a short oligomeric chain of Ub (that is Psh1 independent) and is rapidly degraded. Moreover, Cse4 seems to be massively mislocalized in ubp8Δ cells leading to chromosomal instability [140]. The E3 ligases Ubr1, Sh5 and SCF (through the Rcy1 F-box protein) also regulate the ubiquitination and degradation of Cse4 [141]. How these multiple E3 ligases work with the DUB Ubp8 to ensure appropriate Cse4 degradation remains poorly understood.

10. Roles for deubiquitinases in regulation of transcription

10.1. Proteolytic regulation of RNA polymerase II

Stalling of RNA polymerase II (RNAPII) transcription causes polyubiquitination of Rpb1, the largest subunit of RNAPII, and its subsequent degradation by the proteasome leading to disassembly of the RNAPII complex and cessation of RNAPII stalling. The machinery used for ubiquitination of Rpb1 depends on whether or not the RNAPII stalling reflects the cellular response to DNA damage [142–149]. RNAPII stalling induced by DNA damage involves association of the Elc1–Cul3 E3 ligase with Def1, an RNAPII degradation factor, to ubiquitinate Rpb1. By contrast, during DNA damage-independent RNAPII stalling, there is an interplay between the Rsp5 and Elc1/Cul3 E3 ligases along with Def1 to ubiquitinate Rpb1 [144]. In addition, the Cdc48/ Ufd1/Npl4 complex and its Ub-binding adaptors Ubx5 and Ubx4 are required for ultraviolet-induced DNA damage-dependent degradation of Rpb1 in stalled RNAPII [149]. Whether this group of regulators is also involved in DNA damage-independent degradation of Rpb1 is not known.

In both DNA damage-dependent and -independent cases, Rpb1 is degraded as a ‘last resort’, when RNAPII is persistently stalled. As a first line of defence, the Ubp3 and Ubp6/ Ubp2 DUBs [144] deubiquitinate Rpb1 during DNA damage-dependent and -independent RNAPII stalling, respectively, allowing RNAPII to recover from stalling. Also, Ubp2/ Ubp6 action helps reverse K63 polyubiquitin chains built on Rpb1 by Rsp5, and instead promotes Elc1/ Cul3-mediated K48 polyubiquitination, targeting it for proteasomal degradation. Thus, the DUBs Ubp2/ Ubp6 coordinate the ‘tag-teaming’ of two E3 ligases via their ubiquitin editing functions to overcome transcriptional stalling of RNAPII [143].

10.2. Regulation of RNA polymerase I and transcription factor stability

The stability of Rpa190, the largest subunit of RNAPI, is regulated by Ubp10. Ubp10 deubiquitinates Rpa190 preventing its proteasomal degradation, thus controlling rRNA production and coordinating cell growth [150]. Also, Tbp1, the TATA-binding protein, is essential for transcriptional activation mediated by Gal4 and Gcn4. The Ubp3–Bre5 complex deubiquitinates Tbp1 directly at promoters, antagonizing its proteasomal degradation [151]. Thus, Ubp3 is required for transcriptional activation of Tbp1-dependent genes.

10.3. Regulation of chromatin dynamics (histone H2B ubiquitination)

Chromatin plays a crucial role in regulating transcription and Ub modification of histones plays a significant role in chromatin compaction [152,153]. Histone H2B is ubiquitinated at K123 by the Rad6/Bre1 E3 ligase which in turn primes H3 for repressive methylation (H3K4me3 and H3K79me3) via recruitment of the COMPASS and Dot1 histone methylases [154]. Ubp8 and Ubp10 are both histone H2B DUBs [155–157], but they have very different functions. Ubp8 catalyses H2B deubiquitination in vitro and loss of Ubp8 increases the global level of H2B ubiquitination in vivo [155,156], suggesting that Ubp8 is the major H2B DUB. Ubp8 colocalizes with H3K4me3, while Ubp10 binds to H3K79me3-enriched sites, as well as telomeres and the rDNA locus [157,158]. Furthermore, as a component of the SAGA acetylation complex, Ubp8 is required for the transcription of SAGA-regulated genes [154]. H2B-Ub deubiquitination by Ubp8 promotes Pol II CTD phosphorylation, which is a hallmark of transcription elongation and is required for co-transcriptional mRNA processing [159] (figure 5f). Thus, Ubp8 and Ubp10 act as modulators of crosstalk between different PTMs on histones.

11. Cell signalling: interplay between deubiquitination and other post-translational modifications

Multiple interconnected signalling networks allow yeast cells to adjust their metabolism, gene expression, mating and developmental programmes in response to internal and external stimuli [160]. DUBs are versatile components of signalling networks in yeast, often influencing the activity of key regulators such as protein kinases. For example, Ubp8 positively regulates signalling by deubiquitinating the AMP-activated protein kinase, Snf1, and promoting its stability [161]. Additionally, via an obscure mechanism, Ubp8 and Ubp10...
together influence the activity of Snf1 through modulation of its phosphorylation status [162]. By contrast, deubiquitination of other kinases and their regulators by Ubp3 promotes their proteosomal degradation, causing inhibition of downstream signalling events including: (i) the cell wall integrity pathway kinase Pkc1 during cell wall stress [163]; (ii) the mating activated protein kinase Ste7 [164,165] during the pheromone response and; (iii) the RasGAP, Ira2, that functions in the protein kinase A pathway [166].

The yeast Hog1 protein is a stress-activated protein kinase that physically interacts with and phosphorylates Ubp3 at Ser695. Phosphorylated Ubp3 is recruited to osmo-responsive genes and facilitates efficient initiation of transcription by deubiquitinating Rpb1, the largest subunit of RNApolII [167]. Several additional phosphorylation sites have been identified in other yeast DUBs (summarized in table 1) whose functional relevance remains to be explored. The DUB Ubp2 is modified by oxidation of its catalytic cysteine following exposure to hydrogen peroxide, inactivating its DUB activity. This inactivation leads to accumulation of K63-linked polyubiquitin chains on ribosomal proteins and stabilization of the assembled 80S ribosome and the polysome to help cells cope with oxidative stress [168]. As noted earlier, several DUBs have cysteine as their catalytic residue, thus signalling-based redox regulation of DUBs may be a widespread theme in DUB biology.

12. Roles of deubiquitinases in regulation of the cell cycle

Ubiquitination of cell cycle regulators is integral to ensuring appropriate regulation of cell division, and specific roles for DUBs in cell cycle control have been discovered. For example, ubiquitination of proliferating cell nuclear antigen (PCNA), a sliding clamp protein with many roles in DNA replication, plays a key role in the tolerance to DNA damage in eukaryotes. Ubp10 forms a complex with PCNA resulting in its deubiquitylation during S-phase and loss of UBP10 differentially alters the interaction of PCNA with DNA polymerase ζ-associated protein Rev1 and with an accessory subunit Rev7. Specifically, while mutation of UBP10 enhances PCNA–Rev1 interaction, it decreases Rev7 binding to the sliding clamp [169]. Additionally, Ubp10 prevents degradation of Dbf4, the regulatory subunit of the Cdc7 kinase complex that initiates DNA replication [62]. The G1/S-phase transition of the cell cycle is also regulated by Ubp15, which stabilizes the S-phase cyclin Clb5, thus promoting entry into S-phase.

13. Perspectives

We are beginning to understand the general principles of DUB function, but many questions about the mechanisms of substrate and polyubiquitin chain recognition by DUBs and their condition-specific functions remain to be explored. Also, the extensive PTM of DUBs, including ubiquitination, sumoylation and phosphorylation, suggest that DUBs are regulated by and involved in many signalling pathways in the cell, yet the function of most PTMs on DUBs is unknown. Advances in mass spectrometric methods for detecting PTMs on proteins promise to help advance our understanding of DUB regulation and help in the identification of new DUB targets. Given the high degree of conservation of biological processes between yeast and humans, systematic studies in yeast should provide a reference map for unravelling human DUB function.

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