Who with whom: functional coordination of E2 enzymes by RING E3 ligases during poly-ubiquitylation

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Abstract

Protein modification with poly-ubiquitin chains is a crucial process involved in a myriad of cellular pathways. Chain synthesis requires two steps: substrate modification with ubiquitin (priming) followed by repetitive ubiquitin-to-ubiquitin attachment (elongation). RING-type E3 ligases catalyze both reactions in collaboration with specific priming and elongating E2 enzymes. We provide kinetic insight into poly-ubiquitylation during protein quality control by showing that priming is the rate-determining step in protein degradation as directed by the yeast ERAD RING E3 ligases, Hrd1 and Doa10. Doa10 cooperates with the dedicated priming E2, Ubc6, while both E3s use Ubc7 for elongation. Here, we provide direct evidence that Hrd1 uses Ubc7 also for priming. We found that Ubc6 has an unusually high basal activity that does not require strong stimulation from an E3. Doa10 exploits this property to pair with Ubc6 over Ubc7 during priming. Our work not only illuminates the mechanisms of specific E2/E3 interplay in ERAD, but also offers a basis to understand how RING E3s may have properties that are tailored to pair with their preferred E2s.

Keywords E2 conjugating enzyme; ER-associated protein degradation; linchpin; RING E3 ligase; ubiquitin

Subject Categories Post-translational Modifications & Proteolysis; Structural Biology

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Introduction

The post-translational modifier ubiquitin (Ub) controls virtually every process in eukaryotic cells. Protein modification with Ub is carried out by three sequentially acting enzymes. The E1 enzyme activates the Ub C-terminus for transfer to the active site of an E2 enzyme to form an E2–Ub thioester conjugate (in this text, “~” signifies the thioester linkage). E3 ligases are generally attributed with two functions: They bring an E2–Ub conjugate and a suitable substrate into proximity (“recruitment”), and they catalyze the transfer of Ub from the E2 to the substrate thus ensuring spatiotemporal control of the process (“stimulation”). All eukaryotic organisms from yeast to human have a hierarchy of E1/E2/E3 enzymes, with 1–2 human E1s, up to 36 E2s, and hundreds of E3s. The hierarchy dictates that a given E2 must work with numerous E3s, but it is also true that numerous E2s can work with a given E3. For the largest family of E3s, the RING E3s, the identity of the collaborating E2 defines the product of the reaction (i.e., mono- or poly-ubiquitylation) and, therefore, the biological outcome (Christensen et al., 2007). How an E3 pairs with a specific E2—from the set of enzymes it can physically interact with—to determine a distinct functional outcome remains an open question.

The ability to stimulate E2 activity is rooted in the way RING E3 ligases function. They bind an E2–Ub conjugate and stimulate Ub transfer to a substrate without participating directly in the reaction. In solution, Ub conjugated to an E2 is highly flexible, creating a dynamic conformational equilibrium that is associated with low transfer activity (Fig 1A; Pruneda et al., 2011). RING E3s work allosterically by restricting the flexible Ub toward so-called closed conformations that involve non-covalent interactions between a hydrophobic surface patch on Ub and a surface proximal to the E2 active site (Reverter & Lima, 2005; Saha et al., 2011; Wickliffe et al., 2011; Dou et al., 2012; Plechanovová et al., 2012; Pruneda et al., 2012; Brown et al., 2014; Kelly et al., 2014; Branigan et al., 2020). In general, RING E3s achieve this shift in E2–Ub conformational equilibria through a conserved residue, the allosteric linchpin, that engages both the Ub and the E2 to restrict their relative orientations. A shift toward closed conformations is associated with a dramatic increase in Ub transfer activity from an E2 (Pruneda et al., 2012; Branigan et al., 2020). Structural data reveal that a RING linchpin engages Ub and the E2 through hydrogen

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bonding (Fig 1B; Dou et al., 2012; Plechanovová et al., 2012; Pruneda et al., 2012; Branigan et al., 2015). In a survey of yeast and human RING-type E3s, arginine, with its multiple hydrogen bond donor groups, is most commonly found at the linchpin position (Figs 1C and EV1 for yeast and human, respectively). Surprising, at least half of all human and yeast RINGs feature a residue other than arginine at the linchpin position. Some have been reported as functional, but less efficient linchpins than arginine (Yin et al., 2009; Pruneda et al., 2012; Scott et al., 2014; Stewart et al., 2017); some are residues that can potentially act as a hydrogen bond donor, but have not been experimentally verified as functional linchpins; some are residues that lack hydrogen bond potential altogether. Prominent examples include yeast RING E3s Rad16 (histidine) and Rad18 (leucine), both involved in DNA damage pathways, and Rbx1 (asparagine), the common RING module of the large family of cullin-RING ligases (CRLs). How these RINGs stimulate the ubiquitylation activity of their paired E2s is largely unknown.

The best-understood types of protein ubiquitylation involve the formation of poly-Ub chains of specific linkages (e.g., Ub K48- or K11-linked chains serve as signals for proteasomal degradation) (Komander & Rape, 2012; Metzger et al., 2013b). For RING-type E3s, substrate modification with a poly-Ub chain involves two types of Ub transfer reactions: (i) a “priming” step in which a single Ub is transferred to a protein substrate and (ii) “elongation” reactions in which Ub is attached to the substrate-modified Ub to generate a poly-Ub chain (reviewed in Stewart et al., 2016; Deol et al., 2019). The priming step is biochemically diverse due to the varied nature of potential substrates and Ub attachment sites, whereas chain elongation entails highly specific and repetitive reactions. The different biochemical requirements of priming and elongation have led to the recognition of dedicated E2s for each step. Dedicated chain-elongating E2s that specialize in the synthesis of specific Ub linkages such as K11-, K48-, and K63-linked chains are well known. However, the identities and mechanisms of specific mono-ubiquitylating priming E2s and of E2s that appear to carry out both functions are less well understood. For example, members of the UbcH5 (UBE2D) family are highly promiscuous in vitro and potentially function in both priming and elongation capacities. It is likely that their functional action, i.e., priming or elongation, is dictated by the particular E3 with which such E2s cooperate in a given case (summed up by Stewart et al., 2016), but how such a selection is achieved is not known.

Examples of RING E3s that are confirmed to collaborate with separate priming and elongation E2s to catalyze attachment of poly-Ub chains to substrates continue to be reported (Rodrigo-Brenni & Morgan, 2007; Saha & Deshaies, 2008; Kleiger et al., 2009; Parker & Ulrich, 2009; Pierce et al., 2009; Williamson et al., 2009; Wu et al., 2010a; Kelly et al., 2014; Scott et al., 2014, 2016; Dove et al., 2016; Weber et al., 2016; Hill et al., 2019). Still, our understanding of how E3 ligases control the interplay of E2 enzymes during the two steps of poly-ubiquitylation is rather limited. To understand how RING E3s utilize E2s for priming and elongation reactions, we focused on a well-defined molecular poly-ubiquitylation system. ER-associated protein degradation (ERAD) is a highly conserved protein quality control pathway that targets misfolded ER-resident proteins and marks them for proteasomal degradation (Christiansen & Ye, 2014). Yeast ERAD relies on two RING E3 ligases, Hrd1 and Doa10, and two E2 enzymes, Ubc6 and Ubc7, to modify substrates with K48-linked Ub chains (Fig 1D). Ubc6 harbors a C-terminal transmembrane domain and is thought to function primarily as a mono-ubiquitylating priming E2 (Weber et al., 2016). Ubc7 is recruited to the membrane by the accessory protein Cue1; binding to the Ubc7-binding region (U7BR) of Cue1 renders Ubc7 competent for Ub transfer, while the Cue1 domain of the protein serves to align a growing poly-Ub chain for K48-specific elongation by Ubc7 (Kostova et al., 2009; Bagola et al., 2013; Metzger et al., 2013a; von Delbrück et al., 2016). The E3 Doa10 employs Ubc6 and Ubc7 for priming and elongation, respectively (Weber et al., 2016), while Hrd1 relies predominantly on Ubc7 (Bays et al., 2001) (Fig 1E). Hrd1-targeted ubiquitylation is largely unaffected by the absence of Ubc6, implying that Ubc7 can carry out both priming and elongation with Hrd1. Nevertheless, no direct evidence for Ubc7’s priming activity has been reported to date. Notably, Doa10 features a potentially suboptimal histidine as its linchpin, while Hrd1 harbors a canonical arginine (Fig 1F). This feature implies different requirements for E3-mediated E2 stimulation in the two RING E3s, but this notion has not been directly addressed experimentally.

Our comparisons of the two non-redundant ERAD E3 ligases and their interacting E2 enzymes lead to several key observations and conclusions. First, we present direct in vitro evidence that chain-elongating Ubc7 can work as a priming E2. Second, E3-mediated E2 stimulation contributes to in vivo substrate degradation rates in the priming step, but is dispensable in the chain elongation step. Third, Hrd1 and Doa10 exploit different modes of E2 stimulation that are tailored to the properties of their preferred priming E2. Hrd1 has a high affinity for Ubc7 and relies on its canonical allosteric linchpin to stimulate the E2 in the priming reaction. Ubc6 has high basal activity and does not absolutely rely on stimulation by a functional linchpin forRING-mediated ubiquitylation. This enables Doa10 with its non-canonical linchpin residue to use Ubc6 and not Ubc7 for priming, while not affecting the E3’s use of Ubc7 for chain elongation to a degree that impinges on protein degradation rates. Altogether, this study provides insight into how E3s functionally pair with their potential E2s to enable specific functional outcomes.

**Results**

**Hrd1 selects Ubc7 through high binding affinity**

To understand the E2 preferences of Hrd1 and Doa10, we characterized the catalytically relevant combinations of each E3 ligase with Ub conjugates of either E2. Isothermal titration calorimetry (ITC) was performed using the E3 RING domains, a C-terminally truncated version of Ubc6, and Ubc7 in complex with the U7BR of Cue1, which renders the E2 competent for Ub transfer (Bagola et al., 2013). As the native Ubc6–Ub thioester is susceptible to hydrolysis (see also later Fig 5C), we engineered stable mimics of E2–Ub thioester conjugates by generating a covalent disulfide linkage between a Ub(G76C) mutant and the E2 active site cysteine ([Lorenz et al., 2016]; “SS–” signifies the disulfide linkage for these conjugates).

The Hrd1 RING binds the U7BR/Ubc7-SS-Ub conjugate more than one order of magnitude stronger than it binds the Ubc6-SS-Ub conjugate, with a $K_D$ in the low $\mu$M range (Fig 2A). This implies that Hrd1 selects between Ubc6 and Ubc7 on the basis of K48-linked Ub chain (Fig 1D).
arginine - functional linchpin residue with hydrogen bond donor potential
residue without hydrogen bond donor potential

Figure 1. Ubiquitylating enzymes in yeast ERAD and mechanism of RING-mediated E2 stimulation.
A Current model of RING E3-mediated E2 activation.
B RING linchpin-mediated interactions in an E2-Ub/RING complex. Crystal structure of UbcH5a-Ub/RNF4 (PDB: 4AP4) highlighting the hydrogen bond interactions involving the RING linchpin arginine to side chain and backbone atoms on both the E2 and Ub.
C Histogram of amino acid frequencies at the linchpin position of all 42 yeast RING domains. The linchpin position was defined as the residue at the n + 1 position after the final Zn$^{2+}$ ligand coordinating residues. "–" refers to the atypical RING of Pib1, which lacks the loop containing the linchpin. The SP-RING domains of the two yeast SUMO E3 ligases Siz1 and Siz2 as well as the RING1 domains of the two yeast RBR E3 ligases Hel1 and Itt1 were excluded from this analysis.
D Summary of the relevant properties of E2 enzymes and E3 ligases of the yeast ERAD system.
E Cartoon showing ERAD E2 pairings for both E3 ligases during priming and elongation (see text for details).
F Sequence alignment of Hrd1 and Doa10 for the final Zn$^{2+}$-binding loops (C-X-X-C) showing the linchpin position.
substrate degradation (Bays et al., 2001). In contrast, the Doa10 RING binds each E2 conjugate with similar weak affinity, with K_Ds in the high μM range, consistent with this E3 cooperating with both E2s. We note that the dissociation constants measured for the soluble versions of the RING domains and E2 conjugates reflect the intrinsic affinity for these functional protein–protein interactions and, therefore, the probability of an E3 engaging a particular E2–Ub. Tethering of the relevant components in the ER membrane (see Fig 1F) may overcome the relatively low intrinsic affinity by providing high local concentrations of an E3 and its E2–Ub conjugates.

Hrd1 is presumed to collaborate with Ubc7 for both priming and chain elongation steps, but direct evidence of substrate priming by Ubc7 has not been reported. We implemented an in vitro substrate ubiquitylation assay in which a sequence derived from bovine RNase A (“S-peptide”) is genetically fused to an E3 (e.g., “S-Hrd1”) to serve as a high-affinity substrate recruiter for an RNase A variant that lacks its S-peptide sequence (referred to here as “RNase”; Bays et al., 2001). The Ub(K48R) mutant was used in reactions containing Ubc7 to limit reactions to the priming step by preventing K48-linked poly-Ub chain formation. Reactions that contain a functional E2/E3 pair will generate a product that corresponds to RNase modified with Ub. The pairing of Ubc7 and S-Hrd1 yields a robust RNase-Ub band and a fainter RNase-Ub2 band, which likely reflects double mono-ubiquitylated RNase (Fig 2B). In the absence of S-Hrd1, Ubc7-mediated ubiquitylation is negligible. Importantly, this result provides direct evidence that the efficient chain-elongating E2 Ubc7 is able to prime a substrate. The pairing of Ubc6 and S-Hrd1 also generated modified RNase, albeit to a lesser extent, and this reaction does not depend entirely on the presence of S-Hrd1. Importantly, the stimulatory effect of S-Hrd1, i.e., the difference of RNase modification in the presence of the E3 compared to its absence, is much greater in the case of Ubc7 than Ubc6. Altogether, the results indicate that Ubc6/S-Hrd1 can, in principle, form a functional E2/E3 pair for priming, but is much less efficient than the Ubc7/S-Hrd1 pair.

Although the in vitro ubiquitylation assay provided insights into the capability of Ubc6 and Ubc7 to prime a substrate, RNase is an artificial substrate that interacts with the E3 ligase in a non-native way. To obtain a quantitative measure of E2 activity and the potential of E3s to stimulate it without confounding contributions from a substrate, we monitored the kinetics of Ub transfer from preformed E2–Ub conjugates to a small nucleophile (see Fig EV2A; Pickart & Rose, 1985; Wenzel et al., 2011). Nucleophile is present at huge molar excess thus circumventing the need for a specific substrate interaction. Ub discharge from Ubc6–Ub and U7BR/Ubc7–Ub was followed as a function of time (Fig 2C), and rates were extracted (Fig 2D), providing a metric for substrate-independent E2 discharge activity. While U7BR/Ubc7–Ub undergoes aminolysis like most E2s, Ubc6 has been reported to be hydroxy-reactive (Wang et al., 2009; Weber et al., 2016). We therefore used ethanalamine as the nucleophile, as it provides both an amino and a hydroxy group. While this experimental design allows for uniform reaction conditions, a direct comparison of the E2s’ discharge rates is inadvisable due to the inherent reactivity profile differences. Also, the discharge rates reported here are not directly transferrable to ubiquitylation of substrate in vivo. Despite these caveats, the assays allow assessment of the stimulatory effect of an E3 for a given E2, which we report as the ratio of the discharge rates with and without E3 (Fig 2E). U7BR/Ubc7–Ub discharges very slowly in the absence of an E3 and Hrd1 greatly accelerates the process. In contrast, Ubc6–Ub discharges quite rapidly on its own and is only mildly stimulated by Hrd1. Quantitatively, Hrd1 is roughly an order of magnitude more potent at stimulating discharge from U7BR/Ubc7 than from Ubc6. These results are consistent with the observations made in the in vitro substrate ubiquitylation assays and in line with the difference in binding affinity. They suggest that the difference observed in the in vitro ubiquitylation assay (see Fig 2B) is due to Hrd1’s more effective stimulation of Ubc7. Altogether, the binding and functional assays indicate that the known functional in vitro preference of Hrd1 for Ubc7 over Ubc6 for priming is dictated mainly by binding affinity.

RING linchpin plays a key role in Ubc7 stimulation

Having established that Ubc7 can carry out priming with Hrd1, we wondered why it reportedly does not do so with Doa10, especially as the competing E2 for priming, Ubc6, binds with comparably low affinity to Doa10 (see Fig 2A). Hrd1 and Doa10 differ in their linchpin residues (Arg and His, respectively; see Fig 1F), suggesting the answer might lie in how the two E3s stimulate Ubc7. To investigate the contribution of linchpins in the stimulation of Ubc7, we generated Hrd1 and Doa10 RING variants that harbor one of four amino acids at their linchpin positions (arginine—potent hydrogen bond donor and linchpin of Hrd1; histidine—potential hydrogen bond donor and linchpin of Doa10; alanine—no hydrogen bond potential; or glutamate—hydrogen bond acceptor and charge reversal of arginine).

A strong dependence on linchpin identity was observed in Ub discharge assays for U7BR/Ubc7 acting with either E3 ligase (Fig 3A): For both, arginine is most effective, histidine and alanine are much less effective, and glutamate provides only minor stimulation of U7BR/Ubc7 activity over a “no E3” control. Wild-type Hrd1 (400R) elicits the highest Ub discharge rates from U7BR/Ubc7 with rates achieved by wild-type Doa10(94H) being roughly three times lower. Notably, Doa10 with an arginine linchpin (94R) is as effective as wild-type Hrd1. These observations imply that Hrd1-mediated Ubc7 stimulation is driven by the identity of the linchpin.

The mono-Ub to di-Ub step is the slowest of Ub chain elongation, but not rate-determining for protein degradation

Doa10’s suboptimal histidine linchpin and Ubc7’s strong linchpin dependence offer a rationale for why this E3 does not rely on Ubc7 for priming. How this affects Doa10’s preferred use of Ubc6 for priming will be discussed later. However, Doa10 does engage Ubc7 for chain elongation, begging the question how this is carried out. We implemented an in vitro assay in which one Ubc7-dependent elongation step can be monitored at a time (Fig EV3A; von Delbrück et al., 2016). The first three reactions of chain elongation were recapitulated by adding Alexa 488-labeled Ub to mono-Ub, di-Ub, or tri-Ub to generate di-Ub, tri-Ub, or tetra-Ub, respectively. Reactions were carried out in the absence or presence of Hrd1 or Doa10 or their linchpin variants, respectively, as well as a variant of Cue1 that contains both the U7BR and the Cue1 domain. Reaction rates for each step were derived from quantification of the time courses (Fig 3B).
A

| Kinetic Parameters | U7BR/Ubc7-SS-Ub | Ubc6-SS-Ub |
|--------------------|------------------|------------|
| Hrd1               | $(1 \pm 0.2) \times 10^{-5}$ | $(5 \pm 3) \times 10^{-4}$ |
| Doa10              | $(2 \pm 0.2) \times 10^{-4}$  | $(7 \pm 6) \times 10^{-4}$  |

B

**Figure 2.**

- **Panel A:** Table showing kinetic parameters for Hrd1 and Doa10.
- **Panel B:** Graph showing percentage of RNase-Ub relative to RNase + RNase-Ub.
- **Panel C:** Gel images comparing Ubc7 and Ubc6 before and after charging.
- **Panel D:** Graphs showing time-dependent change in % Ubc7~Ub/(Ubc7 total) and % Ubc6~Ub/(Ubc6 total).
- **Panel E:** Bar graph showing fold stimulation by Hrd1 compared to no E3 control.
Figure 2. Hrd1 selects Ubc7 through high binding affinity.

A Dissociation constants for ERAD E2 SS-Ub/E3 pairs. $K_D$ values were determined by ITC titration of Hrd1 or Doa10 to Ubc6-SS-Ub and U7BR/Ubc7-SS-Ub, respectively. Due to the weak binding, errors are high, so the constants are reported to a single significant figure.

B In vitro substrate ubiquitylation assay for Hrd1 with Ubc7 (left) and Ubc6 (right). Ubc7 reactions contained equimolar amounts of Cue1 and were performed with Ub(X48R). Top: representative immunoblot using a poly-clonal α-RNase A antibody; “no E3” reactions do not contain S-Hrd1, “no ATP” reactions do not contain ATP, but S-Hrd1. The RNase-Ub band co-migrates with a non-specific band (Δ) common to all samples. Bottom: Quantification of RNase-Ub signals is shown. Values are reported as means ± standard deviation ($n = 3$). Significances for pairwise comparisons were determined by one-way ANOVA test; *$P < 0.05$. For clarity, only significances related to the “no E3” control of a given E2 are shown.

C In vitro Ub nucleophile discharge assays for Hrd1 with U7BR/Ubc7 (top—yellow) and Ubc6 (bottom—green) with ethanolamine as nucleophile. Representative Coomassie gels are shown.

D Quantification of Ub nucleophile discharge assays for Hrd1 with U7BR/Ubc7 (left) and Ubc6 (right). Plots of E2–Ub discharge (dots) as a function of time with first-order reaction models fitted to the discharge data (lines) are shown. Values for each time point are reported as means ± standard deviation ($n = 3$). Insets show reaction rates derived from these fits. Significances were determined by Student’s t-test; *$P < 0.05$.

E Stimulation of U7BR/Ubc7 and Ubc6 discharge activities by Hrd1. E2 stimulation is reported as the ratio of rates derived from RING-catalyzed reactions and “no E3” controls in D. Values are reported as means ± standard deviation ($n = 3$). Significance was determined by Student’s t-test; *$P < 0.05$.

Source data are available online for this figure.

Similar to Ubc7 discharge assays, the rates of chain elongation reactions showed a clear dependence on the linchpin identity for both E3 ligases. As before, the E3 variant with an arginine linchpin yielded the highest reaction rates, followed by histidine, alanine, and glutamate; the latter again gave rates close to the “no E3” control. However, unlike in discharge assays, both wild-type E3s provide very similar stimulation of Ubc7-mediated elongation in each reaction step despite the difference in their linchpins (compare rates with black frames in each step). Thus, the difference in allosteric Ubc7 stimulation afforded by each E3 does not impact elongation reactions where the substrate is Ub at the end of a growing chain.

A comparison of the individual reaction steps revealed that the first reaction, i.e., mono-Ub to di-Ub, is the slowest with the next two reactions (di-Ub to tri-Ub and tri-Ub to tetra-Ub) showing generally comparable and higher rates (see Fig EV3B and C). The Ubc7 co-factor, Cue1, aligns the end of a growing Ub chain for modification by binding to the penultimate Ub moiety (von Delbrück et al., 2016). To parse out the contributions of Cue1 and the RING linchpin in Ubc7-mediated Ub chain formation, reactions were carried out with the Cue1(RGA) variant that carries a mutation in its Cue1 domain and thus renders it Ub binding-deficient and unable to align the growing Ub chain (Bagola et al., 2013). Similar to previous studies, the Cue1(RGA) mutant shows a large decrease in rates of reactions using chains of at least di-Ub as substrate (see di-Ub to tri-Ub and tri-Ub to tetra-Ub in Fig EV3B and C). In contrast, the Cue1 mutant has no effect on the reactions that generate di-Ub (see mono-Ub to di-Ub in Fig 3B; von Delbrück et al., 2016). Notably, in reactions carried out by each wild-type E3, the Cue1(RGA) mutation consistently reduces rates to the level of the mono-Ub to di-Ub reaction (highlighted as “minimal wt rates” in Fig 3B). The results demonstrate that, for elongation reactions beyond di-Ub, a defect caused by a RING linchpin mutation in the presence of wild-type Cue1 is smaller or equal to that of Cue1(RGA) in the presence of wild-type RINGs (in Fig 3B, compare rates of linchpin variants in di-Ub to tri-Ub reactions with rates of wild-type E3 in Cue1(RGA) background). The data also reveal that the first reaction of the elongation process is its kinetic bottleneck.

How are these behaviors reflected in vivo? We were unable to follow Ub chain formation on ERAD substrates in vivo in a sufficiently quantitative manner, so we focused on its ultimate biological outcome, i.e., ERAD substrate degradation. Of course, substrate ubiquitylation (priming and chain formation) entails only two steps of many in this process (Fig 3C), but if one or both are rate-determining, changes in them are predicted to affect the observed rate of degradation. Pulse-chase and CHX decay assays were performed in yeast to follow ERAD-mediated protein degradation of vettted model substrates for each E3 (Prα*–3xHA and Hmg2-6xMyc for Hrd1; Deg1-eGFP2 and FLAG-Sbh2 for Doa10—see Figs 3D, and EV4E and EV5E). Relative to strains expressing wild-type Cue1 and Hrd1 or Doa10, degradation of respective model substrates is markedly slower in strains lacking the respective E3 or Cue1. Given the well-documented integral role of the two E3s and of Cue1 in degradation of these substrates, the deletion strains report the stability of these substrates in the absence of ERAD-mediated degradation. In particular, the Hrd1 model substrate Hmg2-6xMyc (Fig EV4E) and the Doa10 model substrate Deg1-eGFP2 (Fig 3D—right panel) are not completely stabilized in the Cue1, Hrd1, and Ao10 strains, respectively, indicating that these substrates can be degraded through other pathways.

Remarkably, in the presence of the relevant wild-type E3, strains that express the Cue1(RGA) mutant show similar model substrate degradation kinetics to those expressing wild-type Cue1 (Figs 3D and EV4E). Only in the case of the Doa10 model substrate FLAG-Sbh2 does the Cue1(RGA) mutation have a negative effect on protein degradation kinetics (Fig EV5E). This effect is, however, only modest when compared to those in the Ao10 and Cue1 strains, and by the end of the assay, protein levels in the Cue1(RGA)-expressing strain were similar to those in the strain expressing the wild-type protein. The simplest explanation for this overall lack of an effect of the Cue1(RGA) mutant is that chain elongation is not rate-determining for ERAD protein degradation (see Fig 3C). Furthermore, we can conclude that the first reaction of chain elongation (mono-Ub to di-Ub) is not rate-determining for protein degradation in vivo at either E3 ligase because this step was the slowest in vitro and the Cue1(RGA) mutation slowed down all elongation steps to the rate of the first step. If the first step contributed to the experimentally observed rate of degradation, one would expect that additional reactions occurring at similar speeds (rather than much faster) would contribute to the overall rate. As this is not what we observed,
we conclude that none of the steps of chain elongation are rate-determining for substrate degradation as observed in our assays. This also suggests that the rates observed in vitro for the first step (“minimal wt rates”) are a lower limit above which effective protein degradation at both E3s under the conditions assayed in vivo is guaranteed.

Figure 3.
Linchpin-mediated Ubc7 stimulation during chain elongation is not crucial for ERAD substrate degradation

The revelation that Ub chain elongation is not rate-determining for substrate degradation implies two possibilities: (i) either the E3-dependent priming step is rate-determining or (ii) a process other than substrate poly-ubiquitylation and independent of an E3 is (see Fig 3C). To distinguish the two possibilities, linchpin mutants of Hrd1 and Doa10 were introduced into yeast. As Ubc7 stimulation depends on the linchpin (see Fig 3A), linchpin variants might allow us to uncouple the recruitment function of each E3 ligase from their E2 stimulation function. In pulse-chase experiments tracking Hrd1 model substrate degradation, we observed a dependence of substrate degradation on the identity of the Hrd1 linchpin (Figs 4A —left panel and EV4A) with similar trends to those in vitro: Substrate degradation occurs most rapidly in the strain expressing wild-type Hrd1 with its arginine linchpin, followed by histidine and alanine linchpin mutants. Expression of Hrd1 with a glutamate linchpin results in degradation impaired to a degree comparable to a strain that lacks Hrd1 altogether.

The fact that substrate degradation rates are detectably altered by introduction of Hrd1 linchpin mutations suggests that there is a Hrd1-dependent step that is rate-determining. In in vitro chain elongation assays, Hrd1 linchpin mutations slow the first step below the lower limit that would guarantee unimpeded protein degradation (see Fig 3B—left panel). Therefore, we can unambiguously conclude from experiments with the linchpin variants that linchpin-mediated stimulation of Ubc7 by Hrd1 is involved in a rate-determining step of protein degradation. But we cannot say whether this is the priming step or the first step of chain elongation (or both) based on the data for Hrd1 alone. As Doa10 uses different E2s for the two steps in question, we turned to Doa10 for clarification.

The effects of Doa10 linchpin mutations on protein degradation were assessed by introducing individual Doa10 variants in the genomic DOA10 locus and analysis of protein degradation by CHX decay assays. In stark contrast to the Hrd1 case, degradation rates of Doa10 substrates are not altered for Doa10 linchpin variants (Fig 4A —right panel). Similar to the case for the Cue1 mutant, the model substrate FLAG-Sbh2 appears to be an exception, as its degradation rates are decreased in strains carrying either Doa10(94R) or (94A).

However, degradation was still substantially faster than in the strain deleted for Doa10 (Fig EV5A). The overall lack of effects in observed degradation rates for Doa10 linchpin mutants implies that these mutations do not slow chain elongation to an extent that adversely affects protein degradation. But the Doa10 linchpin mutations substantially impede the rate of the slowest mono-Ub to di-Ub reaction in in vitro chain elongation reactions to well below the threshold “minimal wt rates” (see Fig 3B—right panel). This means that our previously specified lowest rate estimate for chain elongation can be seriously underestimation in an observed change in the rate of protein degradation observed in our assays. Importantly, the rates observed for Hrd1 linchpin mutants are similar to those of Doa10 linchpin mutants for the first step of elongation (see Fig 3B). Altogether, the results allow us to infer that Hrd1 linchpin mutants do not impede degradation rates through effects on chain elongation. We therefore propose that priming is the E3-dependent rate-determining step in protein degradation in the presence of Hrd1 linchpin variants.

Linchpin-mediated E2 stimulation is crucial for priming at Hrd1, but not at Doa10

To test our hypothesis further, Hrd1 linchpin mutants were tested for their ability to prime in the in vitro substrate ubiquitylation assay (Fig 4B). In contrast to the robust RNase modification observed for Ubc7 with wild-type S-Hrd1(400R), only faint or undetectable RNase-Ub bands appeared with Hrd1 linchpin mutants. The concurrence of Hrd1 linchpin effects in both in vivo substrate degradation assays and in vitro ubiquitylation assays is strong evidence that priming of substrates and not Ub chain elongation is the slower and thus rate-determining step for Hrd1-dependent ERAD substrate degradation. The results also reveal that linchpin-mediated E2 stimulation by Hrd1 is crucial for priming, but is dispensable for elongation in vivo. This is consistent with the ability of Doa10 with its suboptimal linchpin to function effectively with Ubc7 in the chain elongation stage of its substrate ubiquitylation process.

The apparent linchpin independence of Doa10 substrate degradation rates might suggest that its priming step is also linchpin-independent. To test this hypothesis, we performed in vitro Ub discharge assays of Doa10 linchpin mutants with Doa10’s preferred priming E2, Ubc6. No clear correlation between linchpin identity and ability to stimulate Ubc6 was observed (Fig 4C). Indeed, while...
the non-native arginine linchpin is most effective at stimulating Ubc6 discharge, the wild-type histidine linchpin is least effective and alanine and glutamate linchpins appear slightly more effective than the wild-type. Moreover, Ubc6 stimulation does not show any correlation with linchpin identity, when interacting with Hrd1 (Fig EV2F and G), thus strongly supporting linchpin insensitivity as

![Graph A: Hrd1 substrate degradation](image)

![Graph B: Ubc7 + S-Hrd1](image)

![Graph C: Ubc6 + Doa10](image)

![Graph D: Ubc7 + S-Doa10 and Ubc6 + S-Doa10](image)

**Figure 4.**
an intrinsic feature of Ubc6. While this lack of linchpin dependency appears congruent with the observation that degradation of Doa10-dependent substrates is unaffected by Doa10 linchpin mutations (see Fig 4A—right panel), we cannot rigorously conclude that the priming step of Doa10-catalyzed substrate ubiquitylation is rate-determining. The formal possibility that a step that does not involve Doa10 is rate-determining for its substrate degradation cannot be ruled out by the combination of in vivo degradation and in vitro discharge assays for Doa10. Based on the fact that poly-ubiquitylation and, specifically, priming by Ubc7 are rate-determining for Hrd1-dependent degradation, it is likely that the same is true for Doa10-dependent priming by Ubc6. Importantly, however, for both Doa10 and Hrd1 substrate degradation, the Ubc7-dependent chain elongation is not rate-determining.

Data presented thus far provide a rationale for why Ubc7 is not the functional priming partner for Doa10. The question remains how Doa10 is able to function with Ubc6. To assess how Ub transfer to a substrate (as opposed to nucleophile) is affected by the Doa10 linchpin, we performed in vitro assays with both E2s and S-Doa10 linchpin variants (Fig 4D). When paired with Ubc7, only the hyper-active Doa10(94R) variant produces product at a level significantly above the “no E3” reaction. Thus, in this proxy for a priming reaction, wild-type Doa10 is ineffective at stimulating Ubc7 to transfer Ub to a substrate. In Ubc6 reactions, product bands of comparable intensity were detected in all reactions containing Ubc6, including the reaction with no E3 present. Hence, the assay does not provide clarity regarding the linchpin contribution, or lack thereof, for priming reactions carried out by Ubc6. Nevertheless, the assay does recapitulate the strong dependence of Ubc7 on an optimal linchpin. Given the robust RNase modification observed for the S-Doa10(94R) variant with Ubc7, it appears that the suboptimal wild-type histidine linchpin of Doa10 actively disfavors use of Ubc7 for priming, leaving the way clear for pairing with Ubc6.

In the substrate ubiquitylation assays (see Fig 4D), reactions carried out in the absence of E3 reveal a difference between the ability of Ubc6 and Ubc7 to transfer Ub without E3 stimulation. While no to little product was generated by Ubc7, product generated by Ubc6 was clearly detected. Moreover, similar amounts of product were generated in reactions that included Doa10. These observations suggest a scenario where priming can be carried out by Ubc6 without an absolute requirement that its Ub transfer activity be stimulated by an E3. In contrast, regulation of the priming function of Ubc7 appears to depend critically on its linchpin-dependent stimulation by an E3 (as shown for Hrd1—see Fig 2).

**The Ubc6–Ub conjugate populates closed conformations more than other E2–Ub conjugates**

The functional preference of Doa10 for Ubc6 over Ubc7 during priming could be determined solely by Ubc7’s dependence on an optimal linchpin. But it raises the question of how Ubc6 circumvents the disadvantage of Doa10’s suboptimal histidine linchpin. Several observations point to a high basal (i.e., non-E3-stimulated) activity for Ubc6. First, Ubc6 has higher intrinsic ubiquitylation activity on the proxy substrate RNase (compare “no E3” lanes between Ubc6 and Ubc7 in Figs 2B and 4D). Second, the absolute rates of Ub discharge onto ethanolamine in the absence of an E3 are higher for Ubc6 compared to U7BR/Ubc7 (compare “no E3” controls in Figs EV2C and D with EV2E and F). This observation must be interpreted with care due to the bi-functional nature of ethanolamine and the different reactivity profiles of the two E2s. Third, the rates of hydrolysis (i.e., Ub discharge via H2O) of Ubc6 and U7BR/Ubc7 conjugates are vastly different (Fig 5A). And although reactivity profile differences may play a role in this case too, stability toward hydrolysis has been used as a general measure of E2 reactivity (Pruneda et al, 2011; Plechano-vová et al, 2012). In sum, the observations indicate that Ubc6–Ub has higher intrinsic reactivity than Ubc7–Ub.

To understand the source of Ubc6’s higher intrinsic activity, disulfide-linked Ub conjugates of both E2s were characterized by NMR. We focused on the spectral properties of the common subunit, i.e., Ub, in the two species, collecting NMR spectra on samples in which only the Ub subunit was isotopically labeled.
Figure 5.
(E2-SS-(15N)Ub). The spectra of Ub in the context of each conjugate were compared to the spectrum of free Ub, and differences are reported as chemical shift perturbations (CSPs). It has been established for other E2- Ub species that the more Ub inhabits "open" conformations, the closer its spectrum will resemble that of free Ub (i.e., small CSPs) (Hamilton et al., 2001; Pruneda et al., 2011, 2012; Wickliffe et al., 2011; Choi et al., 2015; Dove et al., 2016). As the populations or lifetimes of closed conformations increase, observed CSPs on Ub increase due to the altered chemical environment at the E2/Ub interface. Ub CSPs that center around Leu8, Ile44, Val70, and the C-terminal tail are observed in both the Ubc6 and Ubc7 conjugates (Fig 5B). Leu8, Ile44, and Val70 define the Ub hydrophobic patch that interacts with an E2 in reactive closed conformations (see Fig 1B), and the C-terminal residues link Ub to the E2 active site. The striking difference between the two conjugates is the very large magnitude of Ub CSPs in the Ubc6 conjugate (Fig 5B and Appendix Figs S3 and S4), which are easily among the largest published to date. Seven peaks are either lost due to exchange broadening or shift to an extent that they cannot be assigned with confidence in the spectrum of Ubc6-SS-(15N)Ub as compared with only one such perturbation in the U7BR/Ubc7-SS-(15N)Ub spectrum. The data reveal that Ub conjugated to Ubc6 experiences environments that are more dissimilar to free Ub than Ub in conjugates of Ubc7 or other E2s studied to date (see spectral insets in Fig 5B and Appendix Fig S3 and examples in Choi et al., 2015; Dove et al., 2016; Pruneda et al., 2011). This is strong evidence that a greater population of Ubc6-Ub resides in closed conformations compared to other E2- Ub conjugates and particularly Ubc7-Ub in the absence of an E3 ligase.

15N relaxation parameters were measured as an independent assessment of Ub in different conjugates. Ratios of 15N relaxation times, T1/T2 (Fig 3C), provide a metric for how flexible Ub is in each conjugate (Pruneda et al., 2011). For globular proteins, T1/T2 ratios scale with molecular weight, as they report on molecular tumbling time. Despite Ubc6 having a lower molecular weight than U7BR/Ubc7, their conjugated Ub's average T1/T2 values are similar (39.5 ± 8.2 vs. 36.6 ± 9.8, respectively; Fig 3C). For context, T1/T2 ratios were also determined for Ub attached to human Ubc13 which is much closer in molecular weight to Ub than U7BR/Ubc7. Here, the much smaller average ratio (18.2 ± 3.9) is congruent with a highly flexible Ub that predominantly inherits a myriad of open conformations, as established previously by other techniques (Pruneda et al., 2011). Taken together, these values are strong evidence that Ub attached to Ubc6 spends more time in conformations in which the conjugate tumbles as a single unit rather than two tethered but independent molecular entities. In combination with the large CSPs observed at Ub's hydrophobic patch, the relaxation data strongly support the notion that the Ubc6–Ub conjugate is more biased toward closed conformations than those of other E2s.

A bias toward closed conformations is important for two reasons: first, it is consistent with the greater intrinsic Ub transfer activity we observed for Ubc6–Ub, as closed conformations are generally associated with increased reactivity. In line with this is the observation that E3s only modestly stimulate Ubc6 from its already high basal activity as compared to Ubc7 (see Fig EV2G). Second, the closed conformation bias of Ubc6–Ub helps explain why the conjugate is less dependent on allosteric stimulation via a RING linchpin than Ubc7–Ub. The results thus provide an explanation for why Ubc6 does not strictly rely on a linchpin and, consequently, how Doa10 with its suboptimal linchpin is able to pair functionally with Ubc6 for priming.

Versatility in ERAD priming E2 usage

Despite their established E2 preferences, we have shown that both ERAD E3s can work with their non-preferred priming E2, at least in vitro. This raises the question as to whether there is some crossover of priming E2 utilization in vivo. To address this point, we examined ERAD substrate turnover in cells that lack either Ubc6 or Ubc7. Consistent with the essential role of Ubc7 in the Ub chain elongation step, degradation of most substrates monitored is abolished in cells lacking Ubc7 to levels of cells deleted for the E3 ligases (Figs 6A–C and EV4C—compare solid and dotted curves). Hmg2-6xMyc (Fig EV4C) and Deg1-eGFP2 (Fig 6B) degradation still occurs in Aucb7 strains, as it does in Mhrd1 and Amdo10 strains, respectively (see also Fig 3D), suggesting an alternative pathway of protein quality control for these substrates. As predicted, deletion of Ubc6 has no effect on degradation of Hrd1 model substrates (Figs 6D and EV4D—compare solid and dashed blue curves). However, in strains expressing a suboptimal, Doa10-like Hrd1 linchpin mutant (400H), deletion of Ubc6 has a slight adverse impact on the degradation of the Hrd1 model substrate PrA*-3xHA (Fig 6D—compare solid and dashed green curves). In this Hrd1 hypomorphic setting, the absence of Ubc6 impairs degradation to the level of strains lacking Hrd1 altogether. A similar, though less pronounced trend was detected for Hmg2-6xMyc degradation (Fig EV4D). These observations imply that Hrd1 can use Ubc6 for priming in vivo at least to a small degree and/or under extreme conditions.

To pursue the potential of Ubc6 priming at Hrd1 further, we turned to published evidence that genetically linked Ubc6 to Hrd1 for degradation of its model substrate, CPY* (Hiller et al., 1996). To
confirm involvement of Ubc6 unambiguously, we leveraged the ability of this E2 to transfer Ub to hydroxyl-containing side chains (Weber et al., 2016) by using a lysine-less version of CPY* (“CPY*-K0-HA”) (Balridge & Rapoport, 2016). We confirmed that CPY*-K0-HA is not a substrate for Doa10, as its degradation rate is not affected by deletion of DOA10 (Fig 6E—compare black vs. purple curves). Nevertheless, deletion of UBC6 is as detrimental as deletion of HRD1 or introduction of a catalytically dead version of Ubc7 (C895) (Fig 6E—compare green vs. blue and yellow curves).

Furthermore, simultaneous deletion of HRD1 and UBC6 does not result in an additional effect (Fig 6E—compare green vs. gray curves), suggesting Hrd1 and Ubc6 function in the same pathway for this substrate. Thus, the model Hrd1 substrate can be primed by Ubc6 in its hydroxyl-reactive mode.

In a similar vein, we asked whether Doa10 substrates can be primed by Ubc7, despite the suboptimal linchpin. As predicted by the functional pairing of Ubc6 and Doa10, we observed a negative impact on the degradation of Doa10 model substrates with deletion of UBC6 (Fig 6B and F—compare solid vs. dashed green curves). Degradation of the Doa10 model substrate Deg1-eGFP2 is impaired to similar levels in cells deleted for DOA10 or UBC6 (Fig 6B), illustrating that processing of this Doa10 substrate is strictly dependent on priming by Ubc6. However, cells that lack Ubc6 are still able to turn over FLAG-Sbh2, the other Doa10 model substrate studied here, albeit at a slower rate than when the favored priming E2 is present (Fig 6F—compare solid and dashed green curves). This suggests that Ubc7 can carry out priming at Doa10 E3 ligase. However, when we repeated the experiment in a yeast strain that lacks Ubc6 and expresses the Ubc7-activating Doa10(94R) variant, we did not observe improved FLAG-Sbh2 turnover (Fig 6F—compare dashed green and blue curves). Curiously, in a strain that expresses Ubc6 and the hyperactive Doa10(94R) variant, FLAG-Sbh2 degradation is even slightly impaired (Fig 6F—compare solid green and blue curves) suggesting that in vivo the situation is more complex and Doa10 cannot switch seamlessly between priming E2s depending on its linchpin. Nevertheless, the data illustrate that “suboptimal” E2/E3 pairings are possible and can carry out priming reactions in cells.

As it has been so thoroughly studied, the ERAD poly-ubiquitylation system provided a powerful way to address the question of how an E3 pairs with its preferred E2(s) for priming and elongation. Several features of the ERAD system were key. First, its two E3 ligases have non-overlapping substrate specificities: Hrd1 mainly targets proteins with lesions in their luminal or membrane domains, while Doa10 predominantly degrades proteins with cytosolic lesions (Ruggiano et al., 2014). This allowed us to monitor the effects of perturbations on substrates of one or the other E3. Second, the two E3s use the same chain-elongating E2, Ubc7, but differ in their choice of priming E2. Third, both E3 ligases retain the ability to pair with the non-preferred E2 for priming (i.e., Hrd1 with Ubc6 and Doa10 with Ubc7). While this property can confound interpretation of in vivo studies, it provided important controls in our investigation.

Through a combination of in vitro and in vivo experiments, we have determined the mechanism how each of the ERAD RING E3 ligases utilizes their preferred E2 enzyme during priming. Hrd1 selects Ubc7 over Ubc6 through higher binding affinity: a mechanism we dub “affinity-driven pairing”. The greater than one order-of-magnitude difference in affinity guarantees that a given Hrd1 RING will engage Ubc7–Ub more frequently and with a longer lifetime than a similarly disposed Ubc6–Ub, particularly because both E2s appear to be expressed at roughly similar levels [according to PaxDb—(Wang et al., 2015) and (Clague et al., 2015)]. Furthermore, Hrd1 binding stimulates the Ub transfer activity of Ubc7 better than it does Ubc6 (see Fig 2E). Together, the affinity and subsequent stimulation provide a strong selective preference of Hrd1 for Ubc7 (Fig 7—top). This is in line with the previously reported differential stimulation of Ubc7 by Hrd1 and Doa10 (Cohen et al., 2015). Thus, Hrd1 seems to follow the classical E3 dualism of using its E2 recruitment and stimulation function to control the priming reaction.

In contrast, Doa10 collaborates with different E2s for priming and elongation and does not seem to differentiate between the two based on affinity. Our study reveals that Doa10’s use of Ubc6 during priming is established through a mechanism we call “rate-driven pairing” (Fig 7—bottom). It relies on two features: (i) Doa10’s suboptimal histidine linchpin residue, with which the E3 largely abdicates its E2 stimulation function and thus disfavors collaboration with Ubc7 and (ii) Ubc6’s relative linchpin insensitivity, which allows its utilization by Doa10 despite its suboptimal linchpin. The linchpin insensitivity stems from Ubc6’s high basal activity due to the adoption of reactive closed conformations of its conjugate; hence, the E2 undergoes only moderate stimulation by either E3 in vitro. High E3-independent Ubc6 activity might pose a danger to a cell through ubiquitylation events that are outside of E3 control (as exemplified in vitro for the E3-independent ubiquitylation of RNase by Ubc6—see Fig 2B). We speculate that the reported constitutive auto-ubiquitylation of Ubc6 is a symptom of this as well as a means to keep this danger in check, as the auto-modification is Doa10-independent (Weber et al., 2016) and leads to rapid Ubc6 degradation in the absence of an ERAD substrate (Walter et al., 2001). Such a regulatory mechanism would fit with our finding that Ubc6 is a highly active E2 and underscores the importance of the substrate recruitment function of Doa10 as opposed to its stimulation function as means of spatiotemporal control of the priming reaction on an E3-bound substrate.
Figure 6.
Our investigation adds the ERAD system to those poly-ubiquitylation systems that have been studied to a level of detail that provides kinetic insights on the different steps. Most prominent among these systems are the mitotic master-regulator RING E3 complex APC/C (Rodrigo-Brenni & Morgan, 2007; Jin et al., 2008; Summers et al., 2008; Garnett et al., 2009; Williamson et al., 2009; Wu et al., 2010b; Brown et al., 2014, 2015; Kelly et al., 2014; Chang et al., 2015; Lu et al., 2015) and the SCF RING E3 ligases complexes (Kleiger et al., 2009; Wu et al., 2010a; Emberley et al., 2012; Pierce et al., 2013). For the latter example, millisecond kinetic measurements revealed that most enzyme-substrate encounters with the SCF ligase are non-productive due to the low rate of the first Ub transfer, while rates of sequential Ub additions were markedly faster than substrate dissociation (Pierce et al., 2009). Qualitatively, this kinetic dichotomy is reminiscent of the one we found in the ERAD system.

As seen previously, we find that Hrd1’s selection of Ub7 and Doa10’s pairing with Ubc6 during priming is essential for robust substrate degradation in vivo (Figs 6 and EV4F; Weber et al., 2016). We also found that the ERAD E3s retain the ability to collaborate with their non-preferred E2 for priming. Our work provides a
mechanistic framework for how preferred E2/E3 pairings are established but does not answer the question why these preferences are functionally important for ERAD substrate degradation. Using the linchpin dependencies uncovered, we could alter the E2/E3 pairings in vitro. For example, switching the Doa10 linchpin to arginine creates an E3 that can efficiently ubiquitylate the substrate proxy RNase via Ubc7 (see Fig 4D). Likewise, swapping Hrd1’s linchpin to histidine disables its ability to stimulate Ubc7 (see Fig 3A), while its Ubc6 stimulation remains virtually unchanged (see Fig EV2G). But the same mutations do not translate into a seamless switch of priming E2 preference when viewed through the lens of protein degradation in vivo. While the Hrd1(400H) variant renders Hrd1 substrate degradation somewhat Ubc6-dependent (see Figs 6D and EV4D), it was still less efficient than with wild-type Hrd1(400R), which entirely depends on Ubc7. Similarly, degradation of Doa10 model substrates did not become Ubc6-independent in strains that carry the Ubc7-amiable Doa10(94R) mutant (see Fig EV5D and Appendix Fig S2D). Clearly, there is more to priming E2 preference than the effectiveness of an E3’s linchpin, e.g., ternary interactions between the E3 ligase, its preferred E2 conjugate, and a specific substrate. Thus, correct E2/E3 pairing for priming fundamentally matters for efficient substrate degradation. As to the ability of both E3s to potentially use their non-preferred E2 for priming, we speculate that it might work as a failsafe mechanism under extreme conditions, e.g., Hrd1’s use of Ubc6 for substrates with few or inaccessible lysines (see Fig 6E).

To summarize, we have provided a mechanistic and conceptual framework for how ERAD RING E3s exploit different stimulation requirements of their cognate E2s to pair with them for a specific function. Although the rate-driven pairing model described here is based on the idiosyncratic properties of Ubc6 and might therefore be specific to the ERAD system, our work sheds light into an important and largely overlooked question: Why do a substantial fraction of RING domains feature a residue at the linchpin position that is suboptimal to arginine or cannot hydrogen bond (60% in yeast—Fig 1C, 53% in humans—Fig EV1), when such an interaction is thought to be crucial for stimulating a collaborating E2? Our work suggests a picture where the E2 stimulation function of RING E3s may not be absolutely crucial to control ubiquitylation in all instances. E2s may have different stimulation requirements thus providing RINGs with the opportunity to regulate pairing with specific E2 counterparts for a desired functional outcome.

Most E2s characterized to date have conjugates that exist predominantly in open (less reactive) conformations, thereby implying linchpin sensitivity. But a majority of E2–Ub species have yet to be characterized (Stewart et al., 2016). Given the large fraction of RING E3s that harbor suboptimal linchpins, it is not unreasonable to propose a spectrum of E2–Ub behaviors that span from predominantly open and highly linchpin-dependent, like in the case of Ubc7 (shown here) or human UbcH5 (Pruneda et al., 2011) to more closed conjugates, like that of Ubc6 (shown here) or human Ube2G1 (Choi et al., 2015), which may show higher basal activity and be linchpin-insensitive. RING E3s will have evolved to take advantage of such differences and to “thread the needle” if their biological functions require them to pair with more than one E2. For example, Rbx1, the RING module of all CRLs, has asparagine as its linchpin in both yeast and humans. CRLs control the degradation of about a fifth of all proteasome-dependent degradation (Soucy et al., 2009) raising the question as to why their RING does not provide the most efficient linchpin. Previous work showed that the asparagine linchpin in Rbx1 is actually superior to arginine in its ability to stimulate both the idiosyncratic chain building E2, Cdc34, and the functionally essential NEDD8-specific E2, Ubc12 (Scott et al., 2014). However, the asparagine linchpin is, as expected, less efficient than arginine at stimulating Ub transfer activity from UbcH5. CRLs have been shown to stimulate E2s of the UbcH5 family in addition to Cdc34 (Duda et al., 2008; Saha & Deshaies, 2008; Yamoha et al., 2008) and potentially use the former for priming (Wu et al., 2010a). But in an additional wrinkle, the priming reaction for many CRL substrates in vivo is carried out by the RBR (RING-between-RING) E3 ligase ARIH1 and its cognate E2 UbcH7 (Scott et al., 2016; Dove et al., 2017), which rely on a different molecular mechanism from RING E3s and their E2s (Wenzel et al., 2011). Analogous to the ERAD cases studied here, we speculate that the non-optimal linchpin of Rbx1 disfavors priming reactions catalyzed by Rbx1, thus promoting the preferred priming action of the ARIH1/UbcH7 pair.

Finally, we further extend our insights to another class of E3 ligases that choose from the same pool of E2s as RING E3s, specifically, the aforementioned RING-Between-RING (RBR) E3s. The idea that RINGs have evolved suboptimal linchpins for functional reasons has been invoked in the context of RBR E3 ligases. Despite containing an E2-recruiting RING domain, RBR E3s work via an E3–Ub thioester intermediate that subsequently modifies a substrate (Wenzel et al., 2011). Here, the RBR RING1 domain sterically disfavors closed E2–Ub conformations (Dove et al., 2016). Maintaining the bound E2–Ub in an open, unreactive conformation prevents premature Ub discharge from the E2 onto nearby (E3) lysine residues. Indeed, no human or yeast RBR features an arginine at the linchpin position. We expect there are variations on the themes revealed from our work and look forward to future studies that shed light on other E3 ligases and particularly on E2s to better understand their functional pairing mechanisms.

Materials and Methods

Plasmids, protein expression, and purification

Plasmid constructs, expression, and purification of most proteins have been previously described (see Appendix Table S1). Ubc7 containing only the catalytic cysteine (“C89only” = C39A/C141S), and the U7BR domain of Cue1 (aa 150–203) and Hrd1 (aa 325–412) were cloned into a pET28a vector containing human His6-tagged SUMO3, respectively (Meulmeester et al., 2008); Doa10 (aa 19–102) was cloned into a pGEX-6p1 vector. Using constructs described above, S-peptide-tagged variants of all RING E3 ligases were obtained by cloning. All indicated mutants were created by site-directed mutagenesis. For expression, plasmids were transformed in BL21/DE3 bacteria and grown either in rich LB medium, TB medium, or minimal MOPS medium supplemented with [15N]ammonium chloride. Proteins were expressed overnight at 16°C after induction at OD 0.6–0.8 with 1 mM IPTG. Recombinant proteins were enriched via affinity chromatography using Glutathione Sepharose or Ni-NTA for GST-tagged or His6-tagged proteins, respectively. Depending on the construct, proteins were eluted by
imidazole or cleavage with GST-HR-V-3C protease, SUMO protease (His_{6}-Ulp1 or GST-tagged, catalytic fragment of hSENP1), or thrombin (Sigma). U7BR was purified by incubation with thrombin and GST-hSENP1 after binding to Ni-NTA, followed by a washing step and imidazole elution. All proteins were subjected to Superdex 75 size exclusion chromatography as the final step of purification.

**Generation and purification of recombinant protein complexes and adducts**

Labeling of Ub(S20C) with Alexa Fluor 488 C5 maleimide was previously described (Bagola et al., 2013). K48-linked Ub chains (His_{6}-Ub and His_{6}-Ubs) were assembled as previously described (von Delbrück et al., 2016). The U7BR/Ubc7(C89 only) complexes used for NMR as well as the U7BR/Ubc7(wt) complexes used for *in vitro* Ub discharge assays were purified by mixing purified Ubc7 and U7BR in a 1:2 ratio and subjecting them to Superdex 75 size exclusion chromatography. Disulfide-linked E2-SS-Ub conjugates were obtained by incubating at least 200 μM of desired single-cysteine E2 variants (Ubc6 and Ubc13 naturally contain only one cysteine) with a twofold molar excess of a disulfide-linked adduct of Ub(G76C) and 2-nitro-5-thiobenzoate (TNB) for 45 min at RT followed by Superdex 75 size exclusion chromatography. The Ub(G76C)-TNB adduct was prepared prior to conjugate formation by incubating 1 mM Ub(G76C) with 10 mM DTNB (Ellman’s reagent—Sigma) followed by buffer exchange over Sephadex G-25 resin.

**Isothermal calorimetry**

Measurements were performed on a Microcal VP-ITC calorimeter (Malvern Panalytical) at 15°C. Proteins were dissolved in PBS (25 mM sodium phosphate pH 7.0, 150 mM NaCl). Binding information was obtained with 5 μL additions of titrant (Dna10 or Hrd1 at 3.5 mM) in 4-min intervals into 50 μM indicated E2-SS-Ub samples. Data were analyzed using Microcal PEAQ-ITC analysis software (Malvern Panalytical).

**In vitro substrate ubiquitylation assay**

The experimental setup was modified from Bays et al. (2001). In detail, reactions included E1 (300 nM), indicated E2 (5 μM each; for reactions that contain free Ubc7, 5 μM Cue1(wt) was added as well), indicated 5-peptide-tagged RING variants (5 μM), RNase S (5 μM), and Ub (wild-type or K48 for Ubc6 and Ubc7, respectively, at 50 μM) in *in vitro* reaction buffer (50 mM Tris–HCl, pH 8.0, 4 mM MgCl2, 0.5 mM DTT). Reactions were started by the addition of 4 mM ATP and incubated at 30°C for 30 min. Modification was analyzed by immunoblotting against RNase A. Signals were visualized on a Li-COR Odyssey system. Reactions were performed in triplicates. For each reaction, the RNase-Ub signal was quantified and normalized by the total signal, i.e., signal of RNase and RNase-Ub (RNase-Ub signals were not quantified as they are often negligible and coincide with an unspecific signal).

**In vitro Ub discharge assays**

*In vitro* Ub discharge assays were performed similar to the protocol described in Wenzel et al. (2011). The assay allows monitoring the kinetics of Ub transfer from a preformed E2–Ub conjugate onto a small nucleophile (Pickart & Rose, 1985; Wenzel et al., 2011). To enable investigation of both Ubc7 and Ubc6 activation, we chose ethanolamine as the nucleophile in light of the reported hydroxyl-reactivity of Ubc6 (Wang et al., 2009; Weber et al., 2016). Similar to substrate proteins encountered by these E2s, ethanolamine provides a primary amine and a hydroxyl group as potential nucleophiles. Although the bi-functional substrate renders direct quantitative comparison between Ubc6 and Ubc7 discharge kinetics difficult due to the potentially different products formed by the two E2s, it allows for uniform assay conditions and, for a given E2, for comparison of E2 stimulation by different E3 ligases. For the assay, indicated E2s (30 μM) were charged with Ub (120 μM) in the presence of human E1 (3 μM for Ubc6 or 500 nM for Ubc7) and 5 mM ATP at 37°C for 3 min. To assay Ubc6 activity, purified Ubc6 (aa 2–179) was used. To assay Ubc7 activity, purified complex of U7BR/Ubc7 was charged with Ub(K48R) to avoid intrinsic chain formation. Charging reactions were quenched with 10 mM EDTA and, to start discharge, directly diluted 1:1 into PBS containing ethanolamine (100 mM final concentration) and indicated E3 ligases (30 μM final concentration). Reactivity time courses were performed at RT, if not indicated otherwise. Samples were quenched in non-reducing sample buffer at indicated time points and visualized by Coomassie-stained SDS–PAGE. For each time point, the band intensity for the E2~Ub conjugate and the free E2 were determined by densitometry analysis using Gel Doc EZ Imager (Bio-Rad) and the Image Lab 6.0.1 software (Bio-Rad). Discharge was quantified by plotting over time the intensity of the E2~Ub conjugate band divided by the sum of the intensities of the E2~Ub conjugate and the free E2 band. Reaction rates were derived by fitting discharge to a first-order reaction model using xcrvfit. Reactions were performed in triplicates and presented rates given as the mean of all replicates ± standard deviation.

**In vitro chain elongation reactions**

Single-turnover chain elongation reactions included human E1 (150 nM), indicated Cue1 variants (2 μM), free Ubc7 (2 μM), indicated E3 ligase variants (2 μM), acceptor Ub (C-terminally His_{6}-blocked mono-Ub or indicated K48-linked Ub chains at 14.8 μM each), and donor Ub (fluorescent Ub(S20C)-CS at 200 nM) in *in vitro* reaction buffer. Reactions were started by the addition of 4 mM ATP and incubated at 30°C. Samples were quenched in reducing sample buffer after 0, 1, 2, 3, 4, 5, 10, 30, and 60 min and visualized by SDS–PAGE. Product formation over time was quantified by fluorescence at 488 nm using a Typhoon FLA9500 (GE Healthcare) and ImageQuant TL 7.01 software (GE Healthcare). Reaction rates were derived by fitting discharge to a first-order reaction model using GraphPad Prism 6 software. Reactions were performed in triplicates and presented rates given as the mean of all replicates ± standard deviation.

**Yeast strains**

All strains are haploid descendants of DF5 (trp1-1 (am)/trp1-1 (am), his3Δ200/his3Δ200, ura3-52/ura3-52, lys2-801/lys2-801, leu2-3, -112/leu2-3, -112, MATa/α) (Finley et al., 1987). Strains were generated by standard transformation or crossing protocols.
Appendix Tables S2 and S3 list the yeast expression plasmids and yeast strains, respectively, used in this study. Appendix Fig S5 shows a validation of yeast strains.

**Cycloheximide decay assay**

Cycloheximide decay assays were performed as previously described (Weber et al., 2016). In brief, log-phase growing yeast cells were resuspended in cycloheximide-containing SD medium (Sigma, final concentration: 0.33 mg/ml). At indicated time points, samples were collected and degradation was stopped by the addition of 10 mM sodium azide. Cell lysates were prepared by the addition of glass beads and vigorous shaking. Samples were analyzed by immunoblotting. Appendix Table S4 lists the antibodies used for immunoblotting. For means of quantification, fluorescently labeled secondary antibodies were used and visualized on a Li-COR Odyssey system. Experiments were performed in triplicates.

**Pulse-chase assay**

Pulse-chase assays were performed as previously described (Mehnert et al., 2014). In brief, log-phase growing yeast cells were resuspended in SD medium. Newly synthesized proteins were labeled of [18]C in triplicates. Fluorescently labeled secondary antibodies were used and visualized on a Li-COR Odyssey system. Experiments were performed in triplicates.

During monitoring of Hrd1 model substrate degradation by CHX decay, we experienced technical problems (signal detection/data quality needed for quantification). Hence, we switched to pulse-chase experiments for these substrates. Since sample preparation in pulse-chase assays is inherently more error-prone (immunoprecipitation of substrate out of every sample), we tried to reduce the use of this assay by analyzing Doa10 model substrate degradation via CHX analysis.

**NMR spectroscopy**

NMR spectra were recorded on a 500 MHz Bruker Avance II (University of Washington) at 23°C in 25 mM sodium phosphate pH 7.0, 150 mM NaCl, 10% D2O. 1H/15N-HSQC-TROSY experiments were collected on 200 μM labeled protein, if not indicated otherwise. Datasets were processed using NMRpipe/NMRDraw (Delaglio et al., 1995) and visualized with NMRView (Johnson & Blevins, 1994). Chemical shift perturbations observed by 2D HSQC-TROSY NMR were quantified in parts per million with the equation: Δδ(1H/15N) = [(δ(1H) - δ(1H)0)² + 0.04(δ(14N) - δ(15N)0)²]⁻¹/².

Nitrogen relaxation time analyses were performed with delays of 10, 40, 80, 120, 320, 640, and 1,000 ms for T1 and 8.48, 33.92, 42.4, 25.44, 59.36, 16.96, 50.8, and 67.84 ms for T2. Rates and their respective errors for each analyzed residue were quantified using the built-in “rate analysis” function of NMRView. Average T1/T2 ratios for the (15N)Ub(G76C) constructs shown in Fig 5C are presented as the respective means ± standard deviation for all analyzed residues.

**Data availability**

This study includes no data deposited in external repositories.

**Expanded View** for this article is available online.

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**Author contribution**

CL and TR carried out the experiments with support from MM and MKJ. REK and TR wrote the manuscript with support from CL, AW, MKJ, and TS. REK and TS supervised the project. All authors provided critical feedback and helped shape the research, analysis, and manuscript.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**

Bagola K, von Delbrück M, Dittmar G, Scheffner M, Ziv I, Glickman MH, Ciechanover A, Sommer T (2013) Ubiquitin binding by a CUB domain regulates ubiquitin chain formation by ERAD E3 ligases. Mol Cell 50: 528 – 539

Baldridge RD, Rapoport TA (2016) Autoubiquitination of the Hrd1 Ligase triggers protein retrotranslocation in ERAD. Cell 166: 394 – 407

Bays NW, Garner RG, Seelig LP, Joazeiro CA, Hampton Ry (2003) Hrd1p/Der3p is a membrane-anchored ubiquitin ligase required for ER-associated degradation. Nat Cell Biol 3: 24 – 29

Branigan E, Plechanova A, Jaffrey EG, Naismith JH, Hay RT (2015) Structural basis for the RING-catalyzed synthesis of K63-linked ubiquitin chains. Nat Struct Mol Biol 22: 597 – 602

Branigan E, Penedo JC, Hay RT (2020) Ubiquitin transfer by a RING E3 ligase occurs from a closed E2–ubiquitin conformation. Nat Commun 11: 2846

Brown NG, Watson ER, Weissmann F, Jarvis MA, VanderLinden R, Grace CRR, Frye Jj, Qiao R, Dubé P, Petzold C et al (2014) Mechanism of polyubiquitination by human anaphase-promoting complex: RING repurposing for ubiquitin chain assembly. Mol Cell 56: 246 – 260

Brown NG, VanderLinden R, Watson ER, Qiao R, Grace CRR, Yamaguchi M, Weissmann F, Frye Jj, Dube P, Li Cho S et al (2015) RING E3 mechanism...
for ubiquitin ligation to a disordered substrate visualized for human anaphase-promoting complex. Proc Natl Acad Sci USA 112: 5272–5279
Chang L, Zhang Z, Yang J, McLaughlin SH, Barford D (2015) Atomic structure of the APC/C and its mechanism of protein ubiquitination. Nature 522: 450–454
Chai Y-S, Lee Y-J, Lee S-Y, Shi L, Ha J-H, Cheong H-K, Cheong C, Cohen RE, Ryu K-S (2015) Differential ubiquitin binding by the acidic loops of Ube2 g1 and Ube2r1 enzymes distinguishes their Lys-48-ubiquitylation activities. J Biol Chem 290: 2251–2263
Christensen DE, Brzovic PS, Klevet RE (2007) E2-BRCAL RING interactions dictate synthesis of mono- or specific polyubiquitin chain linkages. Nat Struct Mol Biol 14: 941 – 948
Christianson JC, Ye Y (2014) Cleaning up in the endoplasmic reticulum: ubiquitin in charge. Nat Struct Mol Biol 21: 325 – 335
Clague MJ, Heride C, Urbé S (2019) The demographics of the ubiquitin system. Trends Cell Biol 29: 417 – 426
Cohen I, Wiener R, Reiss Y, Ravid T (2015) Distinct activation of an E2 ubiquitin-conjugating enzyme by its cognate E3 ligases. Proc Natl Acad Sci USA 112: 6625 – 6632
Delaglio F, Grzesiek S, Vuister GW, Zhu G, Pfeifer J, Bax A (1999) NMRpipe: a multidimensional spectral processing system based on UNIX pipes. J Biomol NMR 6: 277 – 293
von Delbrück M, Kniss A, Rogov VV, Pluska L, Bagola K, Löhr F, Güntert P, Hamilton KS, Ellison MJ, Barber KR, Williams RS, Huzil JT, McKenna S, Ptak C, Clague MJ, Heride C, Urbé S (2015) Structure of a conjugating enzyme-ubiquitin complex: a single-molecule study. EMBO J 34: 104863
Der Hiller MM, Finger A, Schweiger M, Wolf DH (1996) ER degradation of a misfolded luminal protein by the cytosolic ubiquitin-proteasome pathway. Science 273: 1725 – 1728
Jin L, Williamson A, Banerjee S, Philipp I, Rape M (2008) Mechanism of ubiquitin-chain formation by the human anaphase-promoting complex. Cell 133: 653 – 665
Johnson BA, Blevins RA (1994) NMR View: a computer program for the visualization and analysis of NMR data. J Biomol NMR 4: 603 – 614
Kelly A, Wickliffe KE, Song L, Fedrigo I, Rape M (2014) Ubiquitin chain elongation requires E3-dependent tracking of the emerging conjugate. Mol Cell 56: 232 – 245
Kleger G, Saha A, Lewis S, Kuhlman B, Deshaies RJ (2009) Rapid E2-E3 assembly and disassembly enable processive ubiquitylation of cullin-RING ubiquitin ligase substrates. Cell 139: 957 – 968
Komander D, Rape M (2012) The ubiquitin code. Annu Rev Biochem 81: 203 – 229
Kostova Z, Mariano J, Scholz S, Koenig C, Weissman AM (2009) A Ubc7p-binding domain in Cue1p activates ER-associated protein degradation. J Cell Sci 122: 1374 – 1381
Lorenz S, Bhattacharryya M, Feiler C, Rape M, Kuriyan J (2016) Crystal structure of a Ube2S-ubiquitin conjugate. PLoS ONE 11: e0147550
Lu Y, Wang W, Kirschner MW (2015) Specificity of the anaphase-promoting complex: a single-molecule study. Science 348: 1248737
Mehnert M, Sommer T, Jarosch E (2014) Der1 promotes movement of misfolded proteins through the endoplasmic reticulum membrane. Nat Cell Biol 16: 77 – 86
Metzger MB, Klevet RE, Weissman AM (2013b) RING-type E3 ligases: master manipulators of E3 ubiquitin-conjugating enzymes and ubiquitination. Biochim Biophys Acta 1834: 47 – 60
Meulmeester E, Kunze M, Hsiao HH, Urlaub H, Melchior F (2008) Mechanism and consequences for paralog-specific sumoylation of ubiquitin-specific protease 25. Mol Cell 30: 610 – 619
Parker JU, Ulrich HD (2009) Mechanistic analysis of PCNA poly-ubiquitylation by the ubiquitin protein ligases Rad18 and Rad5. EMBO J 28: 3657 – 3666
Pickart CM, Rose IA (1985) Functional heterogeneity of ubiquitin carrier proteins*. J Biol Chem 260: 1575 – 1581
Pierce NW, Kleger G, Shan S-O, Deshaies RJ (2012) Detection of sequential polyubiquitylation on a millisecond timescale. Nature 462: 615 – 619
Pierce NW, Lee JE, Liu X, Sweredoski MJ, Graham RJL, Larimore EA, Rome M, Zheng N, Clurman BE, Hess S et al (2013) Cand1 promotes assembly of new SCF complexes through dynamic exchange of F box proteins. Cell 153: 206 – 215
Plechanovova A, Jaffray EG, Taham MH, Naismith JH, Hay RT (2012) Structure of a RING E3 ligase and ubiquitin-loaded E2 primed for catalysis. Nature 489: 115 – 120
Pruneda JN, Stoll KE, Bolton LJ, Brzovic PS, Klevet RE (2011) Ubiquitin in motion: structural studies of the ubiquitin-conjugating enzyme-ubiquitin conjugate. Biochemistry 50: 1624 – 1633
Pruneda JN, Littlefield PJ, Soss SE, Nordquist KA, Chazin WJ, Brzovic PS, Klevet RE (2012) Structure of an E3E2-Ub complex reveals an allosteric mechanism shared among RING/U-box ligases. Mol Cell 47: 933 – 942
Reverter D, Lima CD (2005) Insights into E3 ligase activity revealed by a SUMO-RanGAP1-Ubc9-Nup358 complex. Nature 435: 687 – 692
Rodrigo-Brenni MC, Morgan DO (2007) Sequential E2s drive polyubiquitin chain assembly on APC targets. Cell 130: 127 – 139
Ruggiano A, Foresti O, Carvalho P (2014) Quality control: ER-associated degradation: protein quality control and beyond. J Cell Biol 204: 869 – 879
Saha A, Deshaies RJ (2008) Multimodal activation of the ubiquitin ligase SCF by Nedd8 conjugation. Mol Cell 32: 21 – 31
Saha A, Lewis S, Kleiger G, Kuhlman B, Deshaies RJ (2011) Essential role for ubiquitin-ubiquitin-conjugating enzyme interaction in ubiquitin discharge from Cdc34 to substrate. Mol Cell 42: 75 – 83
Scott DC, Sviderskiy VO, Monda JK, Lydeard JR, Cho SE, Harper JW, Schultman BA (2014) Structure of a RING E3 trapped in action reveals ligation mechanism for the ubiquitin-like protein NEDD8. Cell 157: 1671 – 1684
Scott DC, Rhee DY, Duda DM, Kelsall IR, Olszewski JL, Paulo JA, de Jong A, Ovaa H, Alpi AF, Harper JW et al. (2016) Two distinct types of E3 ligases work in unison to regulate substrate ubiquitylation. Cell 166: 1198 – 1214.e24
Soucy TA, Smith PG, Milholen MA, Berger AJ, Adhikari S, Brownell JE, Burke KE, Cardin DP, Critchley S et al. (2009) An inhibitor of NEDD8-activating enzyme as a new approach to treat cancer. Nature 458: 732 – 736
Stewart MD, Ritterhoff T, Klevit RE, Brzovic PS (2016) E2 enzymes: more than just middle men. Cell Res 26: 423 – 440
Stewart MD, Duncan ED, Coronado E, DaRosa PA, Pruneda JN, Brzovic PS, Klevit RE (2017) Tuning BRCA1 and BARD1 activity to investigate RING ubiquitin ligase mechanisms. Protein Sci 399: 475 – 483
Summers MK, Pan B, Mukhyala K, Jackson PK (2008) The unique N terminus of the Ubch10 E2 enzyme controls the threshold for APC activation and enhances checkpoint regulation of the APC. Mol Cell 31: 544 – 556
Walter J, Urban J, Volkwein C, Sommer T (2001) Sec61p-independent degradation of the tail-anchored ER membrane protein Ubo6p. EMBO J 20: 3124 – 3131
Wang X, Herr RA, Rabelink M, Hoeben RC, Wiertz EJHj, Hansen TH (2009) Ube2j2 ubiquitinates hydroxylated amino acids on ER-associated degradation substrates. J Cell Biol 187: 655 – 668
Wang M, Herrmann CJ, Simonovic M, Szklarczyk D, von Mering C (2015) Version 4.0 of PaxDb: protein abundance data, integrated across model organisms, tissues, and cell-lines. Proteomics 15: 3163 – 3168
Weber A, Cohen I, Popp O, Dittmar G, Reiss Y, Sommer T, Ravid T, Jarosch E (2016) Sequential poly-ubiquitylation by specialized conjugating enzymes expands the versatility of a quality control ubiquitin ligase. Mol Cell 63: 827 – 839
Wenzel DM, Lissounov A, Brzovic PS, Klevit RE (2011) UBCH7 reactivity profile reveals parkin and HHARI to be RING/HECT hybrids. Nature 474: 105 – 108
Williamson A, Wickliffe KE, Mellone BC, Song L, Karpen GH, Rape M (2009) Identification of a physiological E2 module for the human anaphase-promoting complex. Proc Natl Acad Sci USA 106: 18213 – 18218
Wickliffe KE, Lorenz S, Wemmer DE, Kuriyan J, Rape M (2011) The mechanism of linkage-specific ubiquitin chain elongation by a single-subunit E2. Cell 144: 769 – 781
Wu K, Kovacev J, Pan Z-Q (2010a) Priming and extending: a UbcHS/Cdc34 E2 handoff mechanism for polyubiquitination on a SCF substrate. Mol Cell 37: 784 – 796
Wu T, Merbl Y, Huo Y, Gallop JL, Tzur A, Kirschner MW (2010b) UBE2S drives elongation of K11-linked ubiquitin chains by the anaphase-promoting complex. Proc Natl Acad Sci USA 107: 1355 – 1360
Yamoh S, Oashi T, Sarikas A, Gazdou S, Osman R, Pan Z-Q (2008) Autoinhibitory regulation of SCF-mediated ubiquitination by human cullin 1’s C-terminal tail. Proc Natl Acad Sci USA 105: 12230 – 12235
Yin Q, Lin S-C, Lamothe B, Lu M, Lo Y-C, Hura G, Zheng L, Rich RL, Campos AD, Myszka DG et al. (2009) E2 interaction and dimerization in the crystal structure of TRAF6. Nat Struct Mol Biol 16: 658 – 666

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