Protein Engineering in the Ubiquitin System: Tools for Discovery and Beyond

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Abstract—Ubiquitin (UB) transfer cascades consisting of E1, E2, and E3 enzymes constitute a complex network that regulates a myriad of biologic processes by modifying protein substrates. Deubiquitinating enzymes (DUBs) reverse UB modifications or trim UB chains of diverse linkages. Additionally, many cellular proteins carry UB-binding domains (UBDs) that translate the signals encoded in UB chains to cellular proteins and high throughput selection by phage and yeast approaches such as unnatural amino acid incorporation, therapeutic. The recent flourish of protein-engineering approaches such as unnatural amino acid incorporation, protein semisynthesis by expressed protein ligation, and high throughput selection by phage and yeast cell surface display has generated designer proteins as powerful tools to interrogate cell signaling mediated by protein ubiquitination. In this study, we highlight recent achievements of protein engineering.

ABBREVIATIONS: aGFP, against GFP; Aha, azido-homoAla; AVSN, vinyl sulfonamide linkage; BocK, Boc-protected -amino group; BoNT/A, type A botulinum neurotoxin; CHIP, carboxyl terminus of Hsc70-interacting protein; CRL, Cullin RING ligase; Dha, dehydroAla; DHFR, dihydrofolate reductase; diUB, di-ubiquitin; DUB, deubiquitinating enzyme; EPL, expressed protein ligation; ER, endoplasmic reticulum; FL, firefly luciferase; FRET, Förster resonance energy transfer; GOPAL, genetically encoded orthogonal protection and activated ligation; HECT, homologous to E6-AP C-terminus; IMD, immunomodulatory drug; MAPK, mitogen-activated protein kinase; MESSna, sodium β-mercapto-ethane sulfonate; MINDY, motif interacting with UB domain-containing novel DUB; NEMO, NFκB essential modulator; OUT, orthogonal UB transfer; PCNA, proliferating cell nuclear antigen; PROTAC, proteolysis-targeting chimera; PTM, post-translational modification; Pyl-RS, pyrro-Lys tRNA synthetase; RBR, RING-between-RING; RING, Really Interesting New Gene; RL, renilla luciferase; SCF, Skp1-Cullin-F box; SH2, Src homology 2; tRNA, transfer ribonucleic acid; TUBE, tandem UB-binding entity; UAA, unnatural amino acid; UB, ubiquitin; UBA, acceptor UB; UBA, UB-associated domain; UBC, UB-conjugating domain; UBD, UB-binding domain; UBL, UB-like protein; Ubv, UB variant; UbxD8, ubiquitin-domain-containing protein 8; UFD, UB-fold domain; UIM, UB-interacting motif; USP, UB-specific protease; wt, wild-type.
on mapping, probing, and manipulating UB transfer in the cell.

Significance Statement—The post-translational modification of proteins with ubiquitin alters the fate and function of proteins in diverse ways. Protein engineering is fundamentally transforming research in this area, providing new mechanistic insights and allowing for the exploration of concepts that can potentially be applied to therapeutic intervention.

I. Introduction

The 2018 Nobel Prize in Chemistry was awarded to Frances H. Arnold, George P. Smith, and Sir Gregory P. Winter for fundamental contributions to enzyme-directed evolution and protein engineering. Frances Arnold engineered enzymes by directed evolution to gain tolerance to high temperature or high concentration of organic solvent (Chen and Arnold, 1993; Zhao et al., 1998). She also engineered cytochrome P450 to catalyze challenging organic reactions (Kan et al., 2016, 2017; Hammer et al., 2017; Chen et al., 2018). Gregory Smith developed a phage display method for sorting through millions of peptides or proteins for desired molecular recognition with a target molecule through a process known as biopanning (Smith, 1985; Smith and Petrenko, 1997). Gregory Winter applied phage display to engineering humanized antibodies and optimizing their therapeutic efficacy (McCafferty et al., 1990; Clackson et al., 1991; Winter et al., 1994). The field of protein engineering pioneered by these scientists is constantly evolving and expanding. Designer proteins coming from the protein-engineering pipeline assume versatile roles not only as enzymes or antibodies with desired catalytic or binding capacities but also as powerful chemical tools to study cell biology. As examples, components of the protein translational machinery consisting of transfer ribonucleic acid (tRNA) synthetases and ribosomes were engineered for site-specific incorporation of unnatural amino acids (UAAs) into proteins (Liu and Schultz, 2010; Lang and Chin, 2014). The UAAs expand the chemical functionalities on the protein scaffold and generate precise acetylation, methylation, or phosphorylation patterns to reveal the roles of post-translational modifications (PTM) in cell signaling (Wang et al., 2001; Neumann et al., 2010). In contrast, PTM enzymes, including acetyltransferases, methyltransferases, and kinases, were engineered to append chemical labels to their cellular targets to enable their identification from the proteome (Shah et al., 1997; Islam et al., 2013; Yang et al., 2013).

Ubiquitin (UB) is a 76-residue protein that modifies other proteins to mediate signal transduction in the cell and is particularly amenable to protein engineering due to its compact size and stable fold. We will review the technical platforms for engineering UB transfer and a sampling of approaches to the design of UB, enzymes of the UB system, and targets of UB transfer to deduce the cellular signals encoded in this ubiquitous post-translational modification.

UB was first determined to be a post-translational protein modifier that targets eukaryotic proteins for proteolysis in the late 1970s (Ciechanover et al., 1978, 1980; Hershko et al., 1979; Wilkinson et al., 1980). This discovery was recognized by the awarding of the 2004 Nobel Prize in Chemistry to Aaron Ciechanover, Avram Hershko, and Irwin Rose. Until the mid-90s, ubiquitination was primarily studied as a signal that targets proteins for degradation in the 26S proteasome. Indeed, UB-mediated proteasomal degradation controls myriad critical cellular processes. The importance of proteasomal degradation has recently been underscored by the awarding of the 2019 Nobel Prize in Physiology or Medicine, in part, for the discovery of regulated degradation of hypoxia-inducing factor 1α (HIF1α) as a means of sensing changes in cellular oxygen (Maxwell et al., 1999; Ivan et al., 2001; Jaakkola et al., 2001). However, we now understand UB to have a number of non-proteasomal functions in, for example, endocytosis and lysosomal targeting, subcellular localization of proteins, autophagy, DNA repair, and kinase activation. Malfunction of the UB system plays causal roles in diseases such as cancer, inflammatory diseases, and neurodevelopmental and degenerative disorders. The reader is recommended to the many comprehensive reviews on the topics of cell regulation and disease pathogenesis associated with protein ubiquitination (Nakayama and Nakayama, 2006; Frescas and Pagano, 2008; Hoeller and Dikic, 2009; Schwartz and Ciechanover, 2009; Lipkowitz and Weissman, 2011), including a review in this journal summarizing neuronal functions supported by protein ubiquitination (Yi and Ehlers, 2007).

UB is conjugated to proteins through a multi-enzyme cascade that entails the sequential involvement of enzyme classes known as E1s or UB-activating enzymes, E2s or UB-conjugating enzymes, and E3s or UB protein ligases (Fig 1A). The outcome of this is generally the formation of an isopeptide bond between the C terminus of UB and the ε-amino group of a Lys residue. As will be discussed below, UB can be added to proteins not only as a single moiety but also, and perhaps more commonly, in chains of various linkages involving any of its seven Lys residues or its N-terminal amine on Met1. The nature of these chains is determined largely by specific E2s and E3s involved, which, together with E1, write the UB signal on proteins. Ubiquitination is a reversible process, and deubiquitinating enzymes (DUBs; deubiquitinases) can either completely reverse or alternatively edit substrate-bound chains by cleaving linkages with various degrees of specificity. The outcome of protein ubiquitination is largely a function of the UB chain linkage in the context of the substrate and its recognition by the many
Fig. 1. Writers, editors, and readers of cellular signals encoded in protein ubiquitination. (A) The UB-activating enzymes E1s, UB-conjugating enzymes E2s, and UB ligases E3s constitute the UB transfer cascade. E3s are classified into HECT and RBR UB ligases, which have catalytic Cys residues that form thioester intermediates with UB, and RING-type ligases that include the families of RING finger and U-box-containing proteins. RING-type E3s activate E2–UB to directly transfer UB to substrate proteins. E3s can also undergo autoubiquitination (shown for RING-type E3s). UB chains installed on the substrate proteins are recognized by DUBs that cleave or edit the chain. Chains are also recognized by UBD-containing proteins that, in many cases, recruit other cellular partner proteins. (B) Crystal structure of yeast E1 Uba1 in complex with UB with a C-terminal AVSN functionality. AVSN is covalently conjugated to the catalytic Cys residue of Uba1 to freeze its conformation at the moment of UB–E1 thioester formation [Protein Data Bank (PDB) ID 6O83] (Hann et al., 2019). (C) The E1-E2 complex linked by a disulfide bond between the catalytic Cys residues of yeast Uba1 and Ubc4. The conformation of E1 enables the catalytic Cys of E2 to attack the E1–UB thioester to form an E2–UB thioester linkage (PDB ID 4II2) (Olsen and Lima, 2013). (D) Tricomplex of the HECT domain of Rsp5 with UB and the Sna3 substrate peptide. Through a three-way linker, the catalytic Cys of the HECT domain is linked to the C terminus of UB and the substrate peptide to capture the conformation of the HECT domain at the moment of UB transfer to substrate (PDB ID 4LCD) (Kamadurai et al., 2013). (E) RING E3 Cbl-b in complex with UbcH5b–UB conjugate and the substrate peptide from the ZAP70 protein. UB adopts a closed conformation when the E2–UB conjugate is bound to the RING domain to prime the transfer of UB to substrates. (F) The catalytic domain of the DUB CYLD in complex with K63 diUB with the proximal UB in green and the distal UB in blue (PDB ID 2KDE) (Zhang et al., 2009). (G) Tandem UIMs of the Rpn10/SSa protein in a complex with K48 diUB (PDB ID 3WXG) (Sato et al., 2015). (H) Schematic view of open and closed conformations of UB in relative to E2 with and without the association with RING E3s. UB adopts a closed conformation upon the association of RING E3 to prime the transfer of UB to substrates.
proteins containing UB-binding domains (UBD), which translate or read the UB code on a particular protein.

Despite the intense focus on identifying the signaling pathways mediated by UB transfer, there are many important outstanding questions about the mechanism of UB transfer through the E1-E2-E3 cascade and the roles of individual enzymes in cell regulation (Hochstrasser, 2006; Deshaies and Joazeiro, 2009; Harper and Tan, 2012). Due to the roles of the UB system and of altered protein levels in disease, there is great interest in specifically enhancing, inhibiting, or redirecting components of the UB system with the goal of therapeutic development. Recently, some exciting work on engineering UB, its multi-enzyme E1-E2-E3 transfer cascade (the writers), DUBs (the editors), and UBDs (the readers) has provided powerful chemical tools to decode the protein ubiquitination signal in the cell. In this study, we first introduce the basic components of UB transfer cascades and the techniques of protein engineering and enzyme-directed evolution. We then devote much of the review to laying out the technical platforms for engineering protein ubiquitination and provide illustrative examples. It is our hope that recent progress in engineering UB and its transfer inspires the development of new tools to probe, map, and manipulate the cellular pathways of protein ubiquitination.

A. The E1-E2-E3 Cascade as the Writer, Deubiquitinating Enzyme as the Editor, and Ubiquitin-Binding Domain as Reader of Protein Ubiquitination Signals

1. E1: The Ubiquitin-Activating Enzyme. E1 initiates UB transfer by activating the carboxylate of the C-terminal Gly residue of UB with ATP to form a UB-AMP conjugate (Fig. 1A) (Hershko and Ciechanover, 1998; Pickart, 2001). A catalytic Cys residue of E1 reacts with UB-AMP to form a thioester linkage with UB. The UB–E1 thioester (“–E1”) designates the labile thioester bond) is reactive with the catalytic Cys residue of the E2 enzymes to initiate UB transfer. The human genome encodes two E1s, Uba1 (also known as Ube1) and Uba6, which pair with distinct sets of E2s to direct UB transfer to different pools of substrate proteins, with Uba1 being the E1 responsible for the large majority of cellular ubiquitination (Jin et al., 2007; Liu et al., 2017b). Each E1 has unique functions in cell regulation. The crystal structures of yeast E1 (the ortholog of human Uba1) reveal that E1 uses different binding sites to sequentially engage UB. A UB molecule first associates with the adenylation domain of E1 as a UB-AMP conjugate and then is transferred to the active site Cys in the thiolation domain, resulting in a UB–E1 conjugate (Fig. 1B) (Lee and Schindelin, 2008; Olsen and Lima, 2013; Hann et al., 2019). UB transfer from the adenylation to the thiolation domain would require a substantial rearrangement of the E1 enzyme to enable the reaction of the catalytic Cys residue of E1 with UB-AMP (Fig. 1C). The UB-binding site in the adenylation domain has been targeted by inhibitors to regulate protein ubiquitination in the cell (Hyer et al., 2018). A series of UB-like proteins (UBL) that are also protein modifiers have been identified. Included among these proteins are Nedd8, SUMO, and ISG15. These UBLs modify their downstream targets through a similar mechanism as UB and have specific E1s that activate transfer through unique enzymatic cascades (Schulman and Harper, 2009). Similar to the inhibitors of the UB E1, NEDD8 E1 inhibitors with promising anticancer activities have been developed (Soucy et al., 2009; Brownell et al., 2010; Bedford et al., 2011).

2. E2: The Ubiquitin-Conjugating Enzyme. E2s are central players in the ubiquitination cascade. Members of this ~40-member family have a conserved UB-conjugating domain (UBC) fold that is critical to interactions with both E1 and the various families of E3s (Ye and Rape, 2009; Wenzel et al., 2011). This fold includes a catalytic Cys through which UB is bound in a thioester linkage (E2–UB) after transesterification from E1. The N-terminal helix of UBC is a key element for interacting with the UB-fold domain (UFD) of the E1 enzyme (Lee and Schindelin, 2008; Olsen and Lima, 2013) (Fig. 1C). Besides the UBC, various E2s may have extended N or C termini that affect their pairing with various E1 and E3 enzymes. The function of these proteins is significantly more complex than initially appreciated. They can either serve as carriers of UB to substrate-specific E3s, in the case of homologous to E6-AP C-terminus (HECT) and RING-between-RING (RBR) E3s, or function as the final catalytic intermediates in substrate ubiquitination and chain elongations when employed by Really Interesting New Gene (RING) or U-box E3s (RING-type E3s), which constitute the large majority of known E3s (Fig. 1A and 1C).

Particularly when directly transferring UB to ubiquitin chains, E2s play important roles in defining the linkage types of UB chains generated. How specific Lys on substrates are targeted by the RING ligase:E2–UB complex is still poorly understood and clearly depends on the E3-substrate interaction. However, some general concepts in UB chain formation have emerged. Upon the covalent loading of donor UB (UBD) to E2, which results in the formation of UBDR–E2 thioester conjugate, E2s orient a specific Lys (or the α-amino group of Met1) of the acceptor UB (UBA) to enable its approach and nucleophilic attack on the UBDR–E2 thioester, resulting in an isopeptide (or peptide) bond between the amino group of UB and the C-terminal carboxylate of UB and the formation of di-ubiquitin (diUB). Repeating such a process with a specific Lys residue on a UB at the growing end of a substrate-bound (or free) UB chain would lead to the assembly of UB chains of a specific linkage. Because each of the seven Lys residues and the N-terminal amino group of UB could be the site of
iso peptide linkage with UB$^3$, it remains to be fully sorted out as to how various E2s specify the synthesis of different UB chains of defined linkages, which ultimately serve as the means of storing cellular signals in the context of the modified substrate. Various protein engineering platforms have a great opportunity to tease out subtle differences among various E2s in programming UB chain synthesis. In addition to UB, most UBL proteins, such as SUMO and NEDD8, have their own E2s that mediate the transfer of UBLs to their substrates (Walden et al., 2003; Huang et al., 2004; Yunus and Lima, 2006).

3. E3: The Ubiquitin Ligase. E3 enzymes bridge UB transfer from E2 to substrate proteins. There are more than 600 E3s encoded in the human genome. These proteins are largely responsible for providing the exquisite substrate specificity that characterizes the UB system. As noted above, they can be classified into two major types based on whether they form catalytic thioester intermediates with the C terminus of UB (E3–UB) after transfer from E2–UB, or alternatively serve to allosterically activate E2–UB for direct transfer to substrate. In humans, the former category includes the ~28 HECT E3s and 12 RBR E3s (Rotin and Kumar, 2009; Spratt et al., 2014). RING-type E3s include >500 RING E3s and ~7 structurally related U-box E3s (Hatakeyama and Nakayama, 2003; Deshaies and Joazeiro, 2009; Rotin and Kumar, 2009; Lipkowitz and Weissman, 2011; Spratt et al., 2014). They activate the direct transfer of UB from E2 to substrate proteins by orienting UB$^3$ bonded to E2 in a restricted closed conformation through interactions involving the RING or U-Box, E2, and UB and thereby facilitate nucleophilic attack from a Lys residue on substrate proteins or a substrate-bound UB chain (Daviet and Colland, 2008; Mevissen and Komander, 2017). DUBs have been the focus of drug discovery efforts due to their pivotal roles in editing the length and topology of UB chains undergoing proteolysis (Komander et al., 2009; Reyes-Turcu et al., 2009). The majority of DUBs are Cys proteases, including UB-specific proteases (USP), ovarian tumor proteases, UB C-terminal hydrolases, and the motif interacting with UB domain-containing novel DUB (MINDY). Others are Zn-dependent metalloproteases of the JAMM family (Nijman et al., 2005; Mevissen and Komander, 2017). DUBs have varying degrees of specificities in cleaving UB chains. For examples, JAMM DUBs are Lys63 specific, and motif interacting with UB domain-containing novel DUB are Lys48 specific (Fig. 1F) (Sato et al., 2008; Abdul Rehman et al., 2016). DUBs have been the focus of drug discovery efforts due to their pivotal roles in editing the length and topology of UB chains on cellular proteins in response to alterations in cell physiology. They are also potentially attractive targets as, unlike RING-type E3s, they have defined catalytic centers (Daviet and Colland, 2008; Nicholson et al., 2014). Understanding the role of a DUB in cell biology demands a detailed understanding of its linkage specificity and the cellular substrates it recognizes. Protein engineering has provided UB probes with tailored functionalities to help reveal the...
molecular basis for recognition of ubiquitinated proteins by DUBs.

5. Ubiquitin-Binding Domains. The message encoded in UB chains attached to cellular proteins needs to be read to be translated into specific outcomes (Husnjak and Dikic, 2012; Sokratous et al., 2014). This is accomplished through binding to specific UBDs (Fig. 1A). Analogous to the diversity in UB chain topology, more than 20 families of UBDs have been identified; they are found throughout the cell in shuttle and adaptor molecules, within subunits of the 19S cap of the 26S proteasome, and are embedded in E2s, E3s, and DUBs. Structures of UBDs vary widely and include, for example, UB-associated domains (UBA) and coupling of UB conjugation to ER degradation domains, which are triple-helical bundles; UB-interacting motifs (UIM), which have a single long α-helix; and zinc finger domains. These readers of signals can recognize various structural features on UB such as the classic hydrophobic patch surrounding Ile44, the TEK box (Thr9-Glu10-Lys11), the loop region of Glu51-Lys63, or the C-terminal Gly of UB (Husnjak and Dikic, 2012). UBDs are natural affinity reagents for UB-modified proteins. UBDs recognize one or more UB moieties, and many demonstrate linkage specificity. For example, a single PH receptor for UB domain of the proteasome receptor Rpn13 is specific for Lys48-linked UB chains (Husnjak et al., 2008). This is similarly the case for the tandem UIMs of the proteasome subunit Rpn10 (S5a) (Fig. 1G) (Zhang et al., 2009). Taking advantage of characterized UBDs, tandum UB-binding entities (TUBEs) have been developed with up to five UBD repeats that can capture UB chains either non-selectively or with a high degree of linkage specificity (Scott et al., 2015; Mattern et al., 2019). TUBEs and other UB chain affinity reagents are powerful tools to follow the dynamics of protein ubiquitination in the cell and interpret the signals encoded in UB chains.

B. Platforms of Protein Engineering

1. Divergent Versus Directed Evolution. Protein engineering in the laboratory aligns well with divergent evolution in nature. Gene duplication is the first step of divergent evolution (Gilbert, 1978; Eigen, 1992). Through this process, an organism acquires more than one copy of a gene. By retaining a copy to carry out its original functions, there is the flexibility for other copies to diversify through mutagenesis or gene recombination. Natural selection favors the survival of the organism with a duplicated copy of the gene that provides a beneficial trait. Divergent evolution could allow the gene pool of an organism to expand when it confers adaptive advantages for the organism in the environment. Similarly, directed evolution starts with a gene that encodes a protein that is likely to acquire certain binding or catalytic properties by protein engineering (Arnold and Volkov, 1999). The target gene is diversified in the laboratory by error-prone polymerase chain reactions or saturated mutagenesis at key positions. A screen or selection procedure directs the gene pool under a selection pressure to enrich the gene with mutations that give rise to desired traits in binding or catalysis. The choice of the gene as the starting point and the design of screens or selection systems to apply evolutionary pressure is the key to the successful outcome of protein engineering by directed evolution.

2. Screen Versus Selection. Once a protein library is generated by expression from a diversified gene pool, it is assayed to reveal clones with desired features. In a screening experiment, each clone within the library is assayed to reveal clones with desired catalytic or binding properties. For screening of individual clones to be efficient and cost-effective, migration from a low-throughput bench approach to medium- or high-throughput approaches employing robotics is desirable and increasingly used. The availability of genetic systems based on yeast two-hybrid or luciferase reporter genes also makes it possible to screen libraries with 10⁵–10⁶ distinct clones. An alternative approach is a selection experiment in which all clones are combined in one pot to compete for the catalytic turnover of a substrate or binding to a target. Clones showing desired catalytic or binding properties are retained on a solid support and affinity-purified, whereas negative clones are eliminated by washing. Because selection combines all the clones in a library, it can have significantly higher throughput than a screen, as 10⁷–10⁸ clones can be assessed in one round of selection. It is relatively easy to design a selection scheme based on binding affinity. It is more challenging to design a selection experiment to enrich clones with desired catalytic activities. Typically, this is done by coupling catalysis with the labeling of the mutant enzymes with an affinity probe such as biotin (Yin et al., 2004; Zhang et al., 2013b). Proteins showing desired catalytic features can then be selected by binding. In this example, the biotin-conjugated proteins would bind to streptavidin. Another challenge for selection reactions is the deconvolution of the selection results. It is often technically challenging or even unfeasible to identify the mutations on the selected protein itself. A coupling of the phenotype (the protein with the desired catalytic or binding properties), with the genotye (the gene encoding the protein), would enable the identification of the selected protein mutants by sequencing the DNA encoding the mutant. Phage display and yeast cell surface display provide high-throughput selection platforms to enrich clones by catalysis or binding, and enable the deconvolution of selected library clones by DNA sequencing (Kay et al., 1996; Barbas et al., 2000; Chao et al., 2006).
3. Phage and Yeast Cell Surface Display. George Smith first demonstrated the expression of peptides or proteins as a fusion to the pIII protein on the filamentous phage (Parmley and Smith, 1988; Smith and Petrenko, 1997). Later a display system using pCom3H phagemid gained popularity as it allowed the monovalent display of protein library fused to pIII on the surface of M13 phage (Barbas et al., 1991). For affinity-based selection, the phage library can be biopanned against a target immobilized on a solid support (Fig. 2A). Catalysis-based selection often takes advantage of a mechanism-based inhibitor that can form a covalent adduct with the enzyme upon its turnover at the enzyme active site and that is also conjugated to an affinity probe such as biotin. Mutant enzymes with higher catalytic activities would have a faster turnover of the inhibitor and would therefore be more readily covalently conjugated to the biotin-linked inhibitor. The corresponding phage particles could then be

Fig. 2. Protein-engineering platforms for manipulating the activities of UB and its transferring cascade. (A) Phage display of a protein library and selection (biopanning) based on affinity binding with a receptor immobilized on a solid support. (B) Yeast cell surface display with a protein library expressed as a fusion with the yeast cell surface protein Aga2 and binding-based selection with a biotinylated receptor. Yeast cells with biotin labeling are labeled with a streptavidin–fluorophore conjugate and selected by fluorescence-activated cell sorting. (C) EPL for adding functional groups to the UB C terminus. UB is expressed as a fusion to the intein and chitin-binding domain (CBD). The fusion protein is purified by binding to chitin resin and cleaved from the resin with a thiol regent such as sodium β-mercaptoethane sulfonate (MESNa). The UB thioester is released from the chitin resin and reacts with an amine-functionalized probe to add electrophiles such as vinylmethylester (VME) to the UB C terminus. Or the UB thioester can ligate with a synthetic peptide to generate precise modifications at the C terminus. (D) UAA can be incorporated into UB to facilitate the synthesis of UB chains of defined linkages or add specific functionalities to the UB molecule. UAA shown are Lys or Tyr derivatives that can be loaded on Amber suppressor tRNA with engineered Pyrro-Lys or Tyr tRNA synthetases. In the genetically encoded orthogonal protection (GOPAL) and activated ligation method for diUB synthesis, Lys residues in UB are post-translationally protected in the form of either Boc-Lys or Alloc-Lys.
preferentially selected by binding to streptavidin (Yin et al., 2004). For either approach, once the phage particles are selected, they can be amplified for subsequent rounds of selection or for sequencing to reveal the amino acid sequence of the selected clones. Yeast cell surface display is another platform that enables the coupling of phenotype and genotype for high-throughput selection (Colby et al., 2004; Chao et al., 2006). Protein libraries are, for example, expressed as fusions with the yeast cell surface protein Aga2p. Selection is based on the binding of the yeast-displayed library to a fluorescently-labeled probe followed by selecting yeast cells displaying high-affinity binders by fluorescence-activated cell sorting (Fig. 2B) (Zhang et al., 2013a,b). Catalysis-based selection may be achieved by reacting the protein library with affinity probes in the presence of enzymes to couple enzyme catalysis with labeling of proteins by affinity probes on the yeast cell surface.

4. Protein Semisynthesis by Expressed Protein Ligation. It has been a challenge to generate precise PTM patterns on a cellular protein to elucidate the effect of a specific modification on the cellular function of the protein. Expressed protein ligation (EPL) fills this gap as it allows for the appending of synthetic peptides with defined modifications to truncated proteins expressed from bacterial cells. EPL can generate proteins with precise PTMs, and, most importantly for this review, it allows modification of the C terminus of proteins with amine functionalities to install reactive warheads (Fig. 2C) (Flavell and Muir, 2009). To do this, the protein fragment for the EPL reaction is expressed as a fusion with the intein protein and the chitin-binding domain so the fusion can be purified by binding to the chitin column. Rearrangement of the intein generates a thioester intermediate between the C terminus of the expressed protein fragment and the N-terminal Cys of the intein. A thioester exchange reaction between the intein Cys of the intein and the N-terminal Cys generates a reactive protein thioester. Synthetic peptides with an N-terminal Cys and a defined PTM pattern can then be ligated with the protein thioester to generate full-length proteins with the desired C termini. Alternatively, the protein thioester can be conjugated to small-molecule probes with an amine functionality to install reactive warheads at the C terminus (Fig. 2C).

5. Site-Specific Incorporation of Unnatural Amino Acids. The development of UAA incorporation with engineered tRNA synthetase has enabled a broad scope of functionalities to be introduced in any position in a protein (Fig. 2D) (Wang and Schultz, 2004). The engineered tRNA synthetase would load an UAA to an amber suppressor tRNA that would carry the UAA to the ribosome and insert the UAA at the site specified by the amber stop codon (UAG) in the expressed transcript. The UAA incorporation method circumvents tedious synthesis of peptides for the incorporation of unnatural functionalities into the target protein by EPL. It also offers the freedom of incorporating an UAA at any position of the target protein, not restricted to the C terminus as in the EPL reaction. A variety of UAAs bearing PTM functionalities has been incorporated, such as phospho-Ser, Thr, and Tyr (Park et al., 2011; Lee et al., 2013; George et al., 2016; Hoppmann et al., 2017; Luo et al., 2017; Zhang et al., 2017a); acetylated Lys (Neumann et al., 2008); and methylated Lys (Nguyen et al., 2009, 2010; Ai et al., 2010; Groff et al., 2010; Wang et al., 2017b). Furthermore, a Lys analog with a δ-thiol entity is also available for incorporation as an UAA, opening the possibility of synthesizing UB chains of various linkages with the combined use of UAA incorporation and EPL (Fig. 2D) (Virdee et al., 2011).

II. Probing the Mechanism of Ubiquitin Transfer by Protein Engineering

A relay of enzymatic transfers of the carboxy terminus of UB involving a cascade of E1, E2, and E3 delivers UB to cellular targets. Protein engineering has been used to capture crucial intermediates in this cascade to reveal, in some cases, dramatic conformational changes in component enzymes that would not be evident from assessment of static structures. The activity of ubiquitinating enzymes, particularly E3 activity toward substrates, is tightly regulated and coupled to dynamic changes of cell physiology. This is also found with DUBs, as exemplified by the regulated expression and phosphorylation of A20, and the phosphorylation of VCIP135 (Dixit et al., 1990; Zhang et al., 2014; Wertz et al., 2015). To probe the activity of both specific UB transfer enzymes and DUBs, protein-engineering tools have been developed to generate UB or UB conjugates equipped with reactive functionalities. These are being employed as activity-based probes to report the writing, editing, and reading of UB chains. As cellular signals are encoded in the diverse linkages of UB chains, fragments of UB chains or UB-substrate conjugates of defined linkages are being made available by protein engineering based on EPL and UAA incorporation. These represent critical tools to interpret the signals encoded in UB chains.

A. Designer Ubiquitin Conjugates To Probe the Catalytic Mechanism of Ubiquitin Transfer Cascade

1. Ubiquitin-E1 and E1-E2 Conjugates To Capture the Conformational Change of E1. The crystal structure of the yeast E1 Uba1 in complex with UB shows UB binding to the adenylation domain of the enzyme with the C-terminal carboxylate of UB approaching ATP to form UB-AMP conjugate (Lee and Schindelin, 2008). It also reveals a large distance between the catalytic Cys residue in the thiolation domain E1 with UB-AMP.
To enable UB transfer, a conformational change of the thiolation domain is expected to close the gap between the catalytic Cys of E1 and the C-terminal carboxylate of UB engaged in the UB-AMP conjugate bound to the adenylation domain of E1. Furthermore, after the formation of UB~E1 thioester conjugate, the E2 enzyme binds to the UFD domain of E1 to position its catalytic Cys residue in the vicinity of UB~E1 thioester bond to enable the transthioesterification reaction. This would require another conformational change of the E1~E2 complex. To capture the conformation of E1 at the moment of thioester formation with UB, EPL was used to conjugate the adenosine nucleoside to the UB C-terminal carboxylate through a vinyl sulfonamide linkage (UB-AVSN) (Fig. 3A) (Lu et al., 2010; Hann et al., 2019). The sulfonamide group is an isostere of the phosphate group in the native UB-AMP conjugate that enables the neighboring vinyl group to function as an electrophile to react with the catalytic Cys residue of E1 when the thiolation domain of E1 is approaching UB-AMP. The crystal structure of UB-AVSN bound to yeast Uba1 was solved, revealing the conformation of the E1 enzyme during the handover of UB from the adenylation domain of E1 to its thiolation domain to form the E1~UB thioester (Hann et al., 2019) (Fig. 1B). The SUMO-AVSN conjugate was also synthesized and coupled with the SUMO E1. The analogous crystal structure of the conjugate similarly captured the conformation of the catalytic Cys residue of E1 approaching SUMO-AMP for thioester bond formation (Olsen et al., 2010). In another study, the catalytic Cys residues of an E1~E2 pair (Uba1-Ubc4) was conjugated through a disulfide linkage to stabilize the complex and force E1 to adopt a conformation simulating its conformation at the moment of UB transfer from E1 to E2 (Olsen and Lima, 2013). The crystal structure of the E1~E2 conjugate reveals that the E2 bends toward the thiolation domain of E1 to enable the catalytic Cys residue of E2 to approach the UB~E1 thioester (Fig. 1C). The engineering of UB-E1 and E1~E2 conjugates has enabled the capture of key moments of UB transfer through the E1 enzyme. It also provides a molecular explanation for the known requirement for dissociation of E2 from canonical binding sites on E3s to be loaded with UB by E1 (Elet et al., 2005).

2. Ubiquitin-E2 Conjugates To Elucidate the Conformation of Ubiquitin Passing through E2. To stabilize the E2~UB conjugates for structural studies by NMR or X-ray crystallography, researchers have mutated the catalytic Cys residues of E2s to Ser or Lys, which can function in nucleophilic attack on the E1~UB thioester, but would form stable oxyster or amide conjugates with UB, respectively (Eddins et al., 2006; Pruneda et al., 2011; Middleton et al., 2014). Alternatively, Gly76 of UB can be mutated to Cys to allow for nonenzymatic formation of a disulfide linkage with the E2 enzyme and also stabilize the conjugate (Serniwicka and Shaw, 2009). Structural and biophysical studies of these engineered E2-UB conjugates reveal that UB has an unrestrained open conformation that adopts a restricted closed conformation with respect to the E2 enzyme when the E2-UB is docked to RING-type E3s for direct UB transfer to the substrate (Pruneda et al., 2012; Metzger et al., 2014) (Fig. 1, E and H). In contrast, when E2-UB conjugates are bound to HECT or RBR E3s, an open conformation of UB is observed (Kamadurai et al., 2009; Lechtenberg et al., 2016), suggesting UB transfer to a Cys residue of a HECT or RBR or to a Lys residue of a substrate would rely on different positioning and/or reactivities of the E2~UB conjugate, which results from distinctive positioning of UB in relation to E2s.

3. Ubiquitin-E3 Conjugates To Elucidate the Conformation of Ubiquitin Passing from E3 to Substrates. To structurally characterize the activated form of the HECT domain, UB has been linked to the catalytic Cys residue of the mammalian HECT E3 NEDD4 through a disulfide bond to generate a relatively stable mimic of a HECT~UB thioester (Maspero et al., 2013). To capture the catalytically-active conformation of the related yeast HECT E3 Rsp5 at the moment of UB transfer to its substrate Sna3, the C-terminal Gly of UB was mutated to Cys and linked to the catalytic Cys of Rsp5 HECT domain through a three-way cross-linker functionalized with two maleimide groups and one alkyn group (Fig. 3B) (Kamadurai et al., 2013). To accomplish this, a peptide corresponding to the C-terminal portion of substrate protein Sna3 was synthesized with the Lys residue at the ubiquitination site replaced by an azide-functionalized sidechain. A click reaction between alkyn and azide (Kolb et al., 2001) generated a three-way complex mimicking the moment of UB handover from the catalytic Cys residue of HECT to the acceptor Lys of Sna3. Although the structure of the HECT-UB-substrate complex did not show the structure of the cross-linker at the core of the complex due to missing electronic densities, it reveals the potential for reorientation of the C-lobe of the HECT domain to facilitate UB transfer to the substrates (Fig. 1D).

B. Activity-Based Probes for Ubiquitin-Transferring Enzymes and Deubiquitinating Enzymes in the Cell

1. Ubiquitin with C-Terminal Electrophiles as Probes of Deubiquitinating Enzyme. The C terminus of UB can be conveniently derivatized by EPL with new functionalities such as electrophiles acting as DUB traps (Hemelaar et al., 2004; Love et al., 2009). Several examples are described below, all of which involve generation of an activated C-terminal carboxylate by EPL with intein (Flavell and Muir, 2009). To prime for EPL, UB is expressed with a C-terminal intein tag fused with a chitin-binding domain. The fusion is purified by affinity chromatography with chitin agarose. Intein-mediated self-cleavage is then triggered by the addition of a thiol reagent such as
Fig. 3. Enzyme-UB conjugates as probes to study the structure and function of UB transfer cascades and DUB. (A) UB-AVSN conjugate for reacting with the catalytic Cys residue of the E1 enzyme to capture the conformational change in E1 during E1–UB thioester formation. (B) Three-way conjugate of UB, Rsp5 HECT, and Sna3 substrate peptide synthesized by click chemistry and thiol–maleimide coupling. (C) UB-vinylmethylsteer probe for reacting with the catalytic Cys residues of DUBs. (D) UB with a C-terminal Dha residue that can be conjugated to the catalytic Cys residues of E1, E2, E3, and DUB enzymes. (E) UB with a C-terminal alkyne for conjugation with the catalytic Cys residue of DUB. (F) UB with a C-terminal disulfide for disulfide exchange with the catalytic Cys residue of DUB. (G) E2-UB conjugates containing a vinylsulfide linker, which is reactive with catalytic Cys residues of E1 and E3 enzymes.
elimination of the catalytic Cys of E2 through an addition–alkyne moiety with a tosyl substitution installs the linker on the C-terminal azide generated by EPL. Click reaction of the UB-E2 conjugate began with UB derivatized with an acrylamide linker as a Cys trap on UB, a thioester-linked fluorophore to report the activity of the E3 enzymes. Although these reagents have great potential for use in cell-free systems to capture catalytic Cys residues of DUBs, but can also migrate through the E1-E2-E3 cascade and be covalently linked to the catalytic Cys residues at each step of UB transfer (Mulder et al., 2016). This functionality is installed at the C terminus of UB by eliminating the thiol from the Gly76Cys mutant of UB (Fig. 3D) (Chalker et al., 2012). A nonconventional electrophile such as a terminal alkyne can also be appended to the UB C terminus to react with the catalytic Cys residue of DUBs (Fig. 3E) (Ekkekbus et al., 2013). Using a similar approach, the C terminus of the SUMO protein has been functionalized with an alkyne probe to covalently capture SUMO-specific proteases (Sommer et al., 2013). A disulfide was also used to functionalize the UB C terminus so it could react with the catalytic Cys residues of DUBs by disulfide exchange (de Jong et al., 2017) (Fig. 3F). UB with C-terminal electrophiles generated by protein engineering provide versatile activities to probe DUB and UB transferring enzymes in the cell lysates. Besides installing Cys traps on UB, a thioester-linked fluorophore has been added to the UB C-terminal carboxylate (Park et al., 2017). This probe, known as UbFluor, can directly load UB onto the catalytic Cys residues of HECT or RBR E3s accompanied by the release of the fluorophore to report the activity of the E3 enzymes. Although these reagents have great potential for use in cell-free systems, the in vitro intein ligation and functionalization make their use in cells problematic.

### 2. Ubiquitin-E2 Conjugates as Probes of E1 and E3 Enzymes

To generate probes targeting specific stages of UB transfer, E2-UB conjugates were designed with an embedded vinylsulfide functionality to trap Cys nucleophiles of E1 or E3 (Pao et al., 2016). The synthesis of the UB-E2 conjugate began with UB derivatized with a C-terminal azide generated by EPL. Click reaction between the UB-azide and alkyne-functionalized acrylate with a tosyl substitution installs the linker on the UB C terminus with an electron-deficient alkyne moiety. The tosyl group in the linker is then replaced with the catalytic Cys of E2 through an addition–elimination mechanism to generate the UB-E2 probe (Fig. 3G). The probe generated can react with the catalytic Cys residues of E1 and E3 enzymes (Stanley et al., 2015). The UbcH7 (UBE2L3) version of the probe could be conjugated to Parkin, a RBR E3, and was used to detect Parkin activities in cell lysates of neuronal cells or fibroblasts from Parkinson’s disease patients (Pao et al., 2016). Besides identifying known HECT and RBR E3s that formed covalent conjugates with the probe, MYCBP2 was strongly labeled by the probe. This protein is an E3 with an annotated RING domain. However, unlike conventional RING E3s that directly mediate UB transfer to substrates or RBR E3s, evidence suggests MYCBP2 transfers UB through a double Cys relay, leading to UB conjugation to the hydroxyl groups on substrates with a preference for transfer to Thr (Pao et al., 2018). Thus, the E3 was named a RING-Cys-relay E3, and it is the first E3 shown to have a preference for Thr ubiquitination. Similar design of E2-UB conjugates with an acrylamide linker as a Cys trap has been used to probe HECT E3 activity in cell lysates (Byrne et al., 2017). E2-UB probes demonstrate the power of protein engineering to generate customized tools to discover new cellular protein ubiquitination activities.

### 3. diUB Probes with Cys Traps To Target Deubiquitinating Enzyme and E3s

To probe the linkage specificity of DUBs, diUB conjugates with Lys-specific isopeptide linkages between the C-terminal carboxylate of one UB (distal UB or donor UB, UB distal) and the Lys sidechain of other UB (proximal UB or acceptor UB, UB proximal) have been synthesized with built-in thiol-reactive linkers to trap Cys residues on DUBs (Ekkekbus et al., 2014; Hewings et al., 2017). Linkage-specific diUB was prepared by total peptide synthesis with a Gly76Cys mutation at the isopeptide linkage of the diUB. The thiol elimination reaction generated a Dha functionality at the linkage that could trap the catalytic Cys residues of DUBs (Fig. 4A) (Kumar et al., 2010; Haj-Yahya et al., 2014). Following a similar strategy, linear diUB with Dha at the linkage was generated by total peptide synthesis with Gly76Cys mutation at the Met1 linkage. Thiol elimination at the Cys residue generated the linear diUB probe with the Dha functionality. The probe was shown to capture OTULIN, a DUB specific for linear UB chains (Weber et al., 2017). diUB probes were also generated by incorporating azido-homoAla (Aha) in place of a Lys in the proximal UB to enable the linkage of the distal UB through click reaction (Fig. 2D). In such a design, Aha was used to replace a specific Lys residue on the proximal UB through site-specific UAA incorporation. Correspondingly, the C terminus of the distal UB was functionalized with an acrylamide–alkyne linker by EPL. Click reaction between the two parts generated a linkage-specific diUB as a DUB trap, although it is not obvious if the triazole ring at the linkage would affect DUB recognition of the diUB probe (Fig. 4B) (McGouran et al., 2013). Other linkers that have been generated include an acrylamide trap that more closely mimics the native isopeptide linkage of a diUB (Li et al., 2014; Mulder et al., 2014). Linkage-specific tri-ubiquitin probes can be assembled by coupling a distal UB functionalized with
a C-terminal α,β-unsaturated ketone linker with a Cys residue replacing a specific Lys on the diUB (Fig. 4C). The tri-ubiquitin probe has been used to investigate the recognition of UB chains by the DUB USP9X (Li et al., 2014; Paudel et al., 2019). Similar to the synthesis of diUB probes, UB-substrate conjugates were prepared with a Dha group at the isopeptide linkage to trap DUBs that are specific for cleaving UB from the substrate proteins (Whedon et al., 2016; Meledin et al., 2018). Through the Dha functionality, a photocross-linking group has also been installed at the junction of the two UB molecules to trap DUB by photoactivation (Tan et al., 2017).

C. Linkage-Specific Ubiquitin Chains Enabled by Solid-Phase Peptide Synthesis and Unnatural Amino Acid Incorporation

1. Total Peptide Synthesis of Ubiquitin Chains. UB chains of defined linkages are critical tools for assaying and differentiating the chain specificities of DUBs and UBDs. Peptides with a partial sequence of the proximal UB including its ubiquitination sites were prepared by peptide synthesis, and their conjugates with the distal UB at a specific Lys were used to mimic diUB conjugates and assay DUB activities (Faesen et al., 2011; Weller et al., 2014). Total synthesis of diUB of defined linkages was achieved by a series of ligation reactions of peptide fragments to generate diUBs with native isopeptide linkages (El Oualid et al., 2010; Kumar et al., 2010). Tetra UB chains having K48 isopeptide linkages have also been assembled by total peptide synthesis (Kumar et al., 2011). This synthesis sets the stage for synthesis of tetra UB chains having other native linkages (Spasser and Brik, 2012). DiUB probes have also been prepared with fluorophores of a Förster resonance energy transfer (FRET) pair specifically labeling the two ends of a diUB conjugate to screen chain specificities of DUB enzymes (Geurink et al., 2016). DiUB from total synthesis were further derivatized at the isopeptide linkages with amide alkylation, α-carbon modification, and linker modifications to probe the conformation of diUB recognized by UBDs (Haj-Yahya et al., 2013).

![Fig. 4.](image-url)
2. Ubiquitin Chain Synthesis Enabled by Unnatural Amino Acid Incorporation. Total peptide synthesis can elegantly generate UB chains with defined Lys linkages. There are downsides, however, that include the equipment and expertise necessary for these syntheses and the limited quantity of products. E2s with strict linkage specificities have been shown to have the capacity to generate specific UB linkages in an E3-independent manner. This has been established for K48 and K63 linkages using E2-25K or Ubc13-Mms2, respectively (Piotrowski et al., 1997; Hofmann and Pickart, 2001), and more recently for K11 linkages using Ube2S (Castañeda et al., 2013). These enzymes represent powerful tools for generating defined diUB conjugates. To produce a homogenous K48 or K63 diUB chain, the proximal UB is blocked at the C terminus with an Asn77 extension, which can later be cleaved off by a DUB after diUB synthesis is complete. On the distal UB, the Lys residue recognized by E2 for further chain elongation is blocked by mutation to a Cys or Arg. These variations force E2-catalyzed coupling to occur only once between the designated proximal and distal UB molecules (Fig. 5A). For K11 diUB chains, Ube2S has been fused to a zinc finger UBD to enhance recruitment of the UB monomer and accelerate K11 chain formation (Bremm and Komander, 2012). Although UB mutants with various combination of Lys to Arg mutations may force the assembly of specific linkages by E2s and E3s, the use of linkage-specific E2s enables the synthesis of native K11, K48, and K63 diUB linkages with nonparticipating Lys residues left intact (Castañeda et al., 2016). For other diUB linkages (i.e., K6, K27, K29, and K33), the incorporation of UAAAs at positions corresponding to the site of linkage formation provides a generally applicable route to generate UB chains of various linkages. Recently, a chemosynthetic method known as genetically encoded orthogonal protection and activated ligation (GOPAL) has been developed and shown to be useful for diUB synthesis (Fig. 5B) (Virdee et al., 2010; Castañeda et al., 2011). GOPAL takes advantage of the pyrro-Lys tRNA synthetase (Pyl-RS) that incorporates a Lys analog with a Boc-protected ε-amino group (BocK) (Fig. 2D). After BocK incorporation in the place of a specific Lys in what will become the acceptor UB, the remaining amino functionalities (the N-terminal amine and six Lys residues) are globally protected with Cbz or Alloc groups (Fig. 2D). Acid treatment selectively removes the Boc group to unmask the reactivity of the designated Lys. The deprotected Lys is then used to couple with the inter-derived thioester of the distal UB that is also globally protected on the amines. After coupling, the protecting groups are removed to afford diUB of a native linkage. Such an approach generates native isopeptide linkages between the two recombinant UBs produced in Escherichia coli. Other strategies for UB chain synthesis have been developed to incorporate derivatized Lys as a UAA that would facilitate EPL at the site of UB conjugation. For example, a pyrro-Lys tRNA synthetase was engineered that could incorporate δ-thiol-Lys into proteins (Fig. 2D) (Virdee et al., 2011). The δ-thiol group attached to a Lys sidechain in what will become the proximal UB undergoes transesterification with the UB thioester, and subsequent S-N acyl transfer would form the precursor for the isopeptide linkage for UB chain synthesis. To complete the synthesis, after ligation, the δ-thiol group is desulfurized to leave a native isopeptide bond between the two UB. UB chain synthesis enabled by UAA incorporation is an extremely valuable approach to generating linkage-defined diUB for decoding chain-encoded UB signals. As an example, diUB probes assembled by UAA-assisted semisynthesis, as well as by total peptide synthesis, have been functionialized by photocross-linking groups to identify UBD proteins as readers of the UB linkage (Liang et al., 2017; Braxton et al., 2019).

3. Engineered Ubiquitin Chains of Non-native Linkages. UB chains of native linkages are ideal choices of reagents when studying the specificity of DUBs for various chain types. However, depending on the purpose of the study, UB chains of non-native linkages are a close mimic of native chains that can be employed to reveal chain topology and identify reader proteins of UB chains. Often UB chains of non-native linkages are relatively easy to assemble, and there is a wider choice of coupling chemistry available. Linkage-specific E2s provide a facile way to assemble tetra UB chains or even longer conjugates with a periodic thioether linkage (Varadan et al., 2002; Pickart and Raasi, 2005; Raasi and Pickart, 2005). In such a method, E2 is initially used for diUB formation with the Lys residue for further chain elongation mutated to a Cys or Arg. These chains have been used to obtain insights into chain structure and function, including the initial determination that K48-linked tetra UB chains are specifically recognized by the proteasome, and to study interactions with UBDs (Thrower et al., 2000; Raasi and Pickart, 2005; Varadan et al., 2005; Eddins et al., 2007). Nonenzymatic reactions such as thiol-ene coupling have also been used to assemble diUB and UB chains (Fig. 5C). To enable the coupling reaction, the C terminus of the distal UB is functionialized with an allyl...
Fig. 5. Synthesis of diUB and tetraUB conjugates of specific linkages. (A) Synthesis of tetraUB of K48 linkages using a K48-specific E2, E2-25K. In the first step, diUB is assembled using UB with an Asp77 extension as the proximal UB and UB with a Lys48Cys mutation as the distal UB. E2-25-catalyzed UB conjugation generates a diUB with a K48 isopeptide linkage. Next, the diUB conjugate is divided into two parts, with one part reacting with ethylenimine to derivatize the Cys48 residue on diUB with an amine functionality so it can be recognized by E2-25K as a Lys residue.
group. On the side of the proximal UB, the Lys residue for UB conjugation is replaced with a Cys by mutagenesis. Taking advantage of the fact that UB has no native Cys residues, a photoinitiator was used to induce radical formation on the thiol group of the Cys side chain (Trang et al., 2012). Thiol-ene coupling between the two UB generates a diUB with a thioether linkage that was one carbon longer than the native isopeptide bond. The coupling strategy can be further expanded to generate linear or branched UB chains. UB chains can also be synthesized by copper-catalyzed click chemistry between alkyne and azide functionalities yielding a triazole linkage that is resistant to DUB cleavage (Valkevich et al., 2012). In one example, the C terminus of the distal UB was functionalized with alkyne and the proximal UB was produced by solid-phase peptide synthesis with the incorporation of azido-ornithine at the site of Lys for UB conjugation. Click reaction generated diUB of specific linkages to identify chain-specific UBDs (Flierman et al., 2016; Zhang et al., 2017c). The click ligand can also be installed by UAA incorporation. For example, alkyne-functionalized propargyl-Lys (PrK) can be used to replace a specific Lys in UB at the site of UB chain formation through UAA incorporation by Pyl-RS (Fig. 2D; Fig. 5D). In the same UB molecule, azido-homoAla (Aha) was used to replace Gly76 at the C terminus of UB by changing the Gly codon to a Met codon, so Aha was added to the UB C terminus as a Met analog. Click reactions with the double mutant of UB generated UB chains of defined linkages through a triazole linker (Eger et al., 2010; Debelouchina et al., 2017). The global protection strategy was applied to the synthesis of human Dishevelled protein with two mono-UB modifications (Madrzak et al., 2015). The UB chains from total synthesis were coupled to substrate proteins via a maleimide linkage with a Cys residue on the substrate proteins replacing a specific Lys at the ubiquitination site or via an oxim linkage with an aldehyde group attached to the Cys residue in the substrate (Hemantha et al., 2014; Singh et al., 2016). UB was linked to a substrate peptide by total synthesis through oxime coupling between an aldehyde-functionalized UB C terminus and a substrate peptide presenting an aminooxy-derivatized Lys residue (Shanmugham et al., 2010). Click chemistry has been used to generate monoubiquitinated PCNA with UAA incorporation to introduce Aha at the UB C terminus and PrK at the ubiquitination site in PCNA (Fig. 2D) (Eger et al., 2011). Recently, Pyl-RS has been engineered to incorporate a Lys derivative with azido-diGly peptide at a specific position of substrate proteins. Reduction of the azido group would generate diGly-conjugated Lys with a free amino end as the substrate of sortase-mediated peptide ligation that would conjugate UB to the specific Lys residue on the substrate (Fig. 5F). The ligation product would have C-terminal mutations in UB to match the specificity of the sortase (Fottner et al., 2019). In another report, wild-type (wt) Pyl-RS was used to incorporate Cys-conjugated Lys into calmodulin (Fig. 2D). EPL reaction between UB thioester with the Cys-conjugated Lys residue on calmodulin generated a UB-CaM conjugate that has a Cys residue replacing the C-terminal Gly of the distal UB at the isopeptide connection (Fig. 5G) (Li et al., 2009).

III. Manipulating Ubiquitin Transfer by Protein Engineering

UB rivals the phosphate in the scope of cellular targets modified and the cellular outcomes of the modification.
Being a protein itself, UB can be further modified by phosphorylation, acetylation, or glycosylation, which adds another layer of complexity to the signals encoded in protein ubiquitination (Swatek and Komander, 2016). Because UB is genetically encoded, it can be plugged into all protein-engineering platforms such as site-directed mutagenesis, phage display, and yeast cell surface display to engineer how it is transferred through the E1-E2E3 cascade, processed by DUBs, and bound by UBDs. By manipulating the reactivity of UB with these writers, editors, and readers, we can potentially control the ubiquitination status of specific proteins in the cell and regulate their degradation, interactions, trafficking, and enzymatic activities.

A. Engineering Ubiquitin To Manipulate Ubiquitin Transfer

1. Ubiquitin and Ubiquitin-Like Protein-Mimicking Peptides. UB C-terminal peptide bearing the LRLRGG sequence (amino acids 71–76) was found to be a surrogate substrate of the UB transfer cascade, although the activity of peptide transfer is orders of magnitude lower than that of the full-length UB (Jonnalagadda et al., 1988; Madden et al., 2008). To define the sequence space of the UB C terminus that could be accommodated by E1 to activate UB transfer, we (Yin and colleagues) randomized UB C-terminal residues covering LRLRG in the context of full-length UB and developed a phage selection scheme to identify UB variants that could be activated by E1 (Zhao et al., 2012a). The UB library was displayed on phage with exposed randomized C termini, and selection was carried out based on thioester bond formation between the UB variants and E1 immobilized on a streptavidin plate. The selection yielded UB variants with alternative C-terminal sequences that could be recognized by E1 for transfer through the cascade. Furthermore, the C-terminal peptides derived from the UB variants from phage selection have hundreds-fold higher activity with E1 compared with the native LRLRG sequence, presumably due to the optimized interaction between the E1 and the peptides from phage selection. The peptides known as UB-mimicking peptides could be transferred through the ubiquitination cascade, where they blocked the transfer of the full-length UB from passing through the cascade by forming thioester conjugates with E1, E2, and HECT E3s (Zhao et al., 2012c; Jin et al., 2018). In this way, these peptides serve as inhibitors of UB transfer through the cascades, although they would not target a specific protein ubiquitination pathway. Using a similar phage selection platform, short C-terminal peptides have been identified that mimic the transfer of UBL proteins NEDD8 and SUMO through their respective enzymatic cascades. In vitro assays showed the inhibition of UB transfer through their cascades by the NEDD8- and SUMO-mimicking peptides (Zhao et al., 2013a,b, 2014).

2. Ubiquitin Variants That Regulate the Activity of Deubiquitinating Enzyme and E3 in the Cell. DUB and E3 enzymes frequently harbor UB-binding sites for substrate recognition or allosteric control of activity (Komander et al., 2009; Maspero et al., 2011). The association of these UB-binding sites with UB would have rapid on and off rates to accommodate the processivity of UB chain formation and the editing capacity of DUBs. By enhancing the binding affinity of UB with these sites on DUB and E3, Zhang and Sidhu (2014) and Gorelik and Sidhu (2017) found that engineered UB variants (Ubv) could function as specific DUB or E3 inhibitors. To do this, they identified a binding interface for the DUB USP21 on UB that comprised ~30 residues. These amino acids were randomized in a UB phage display library and panned against USP21 as well as USP8 and USP2a (Ernst et al., 2013). Ubvs were identified that specifically interact with their cognate DUB in cells and similarly show evidence of selective modulation of function. Thus, Ubvs from phage selection can function as genetically encoded DUB inhibitors. The crystal structures of USP2a and USP21 with Ubvs bound showed the Ubvs adopted the same binding mode as the wt UB at the distal UB-binding site of the DUBs. Interestingly, the crystal structure of USP8-Ubv complex shows a significantly different mode of binding than USP8-UB, suggesting phage selection using the Ubv library could find alternative binding modes with DUBs that maximize binding affinity. Zhang and Sidhu (2014) and Gorelik and Sidhu (2017) also generated Ubvs for binding to DUBs of various families as well as viral DUBs. The E2 enzyme Cdc34, and HECT, RING, and U-box E3s have also been assessed, and Ubvs for these that inhibited UB transfer have been identified. In some cases, this has been found to occur through previously unappreciated E3-UB interactions (Brown et al., 2016; Gorelik et al., 2016, 2018; Leung et al., 2016; Gabrielsen et al., 2017; Zhang et al., 2017b; Ordureau et al., 2018). In another report, the computation-assisted design was used as a guide to generate Ubv libraries, resulting in Ubvs that inhibit the DUB USP7 (Zhang et al., 2013c). Besides their roles as inhibitors of E3s and DUBs, engineered Ubv can also activate E3s (Zhang et al., 2016; Gabrielsen et al., 2017). In the case of the HECT E3, NEDD4L, an Ubv showed enhanced affinity for a UB-binding site in the N-lobe of the HECT domain and thereby allosterically released the self-inhibited state of the HECT domain—activating UB transfer. Ubvs also have revealed novel sites of protein–protein interaction on an E3. The structural characterization of Ubv binding with the Fbw7-Skp1 subcomplex of the E3 SCFBw7 [Skp1-Cullin-F box (SCF) E3 complex with Fbw7 as the F-box component (also known as CRL1Fbw7)] showed the Ubv to be bound to a site normally occupied by Cul1, and, as a result, the Ubv inhibits the association of
Cul1 with Skp1-Fbw7. This led to the structure-based design of a Ubv that functions similarly with Skp1-Fbw11 (Gorelik et al., 2016) and to the selection of Ubv inhibitors of SCF assembly for 17 additional F-box proteins complexed with Skp1 (Gorelik et al., 2018). Thus, phage selection using a Ubv library has led to the identification of an interface that could be targeted by designer protein ligands to interfere with the assembly of SCF complexes. As there are other Cullins that have a similar interaction interface as Cul1, there is the potential for other CRL E3 families to be similarly targeted (Gorelik et al., 2018). Overall, Ubvs have potentially broad applications in probing the catalytic mechanisms of E3s and DUBs by forming high-affinity complexes with enzymes and allowing their structural characterization by X-ray crystallography and electron microscopy. Ubvs could also be used as genetically encoded inhibitors of E3 and DUBs to validate the therapeutic potentials of specific E3 or DUB targets for drug development. Ubvs are recombinant engineered proteins. As potential protein therapeutics, they present challenges related to issues such as stability, cell penetration, and potential immune responses due to introduction of foreign antigenic epitopes on the Ubv. However, with the development of gene and protein...
delivery platforms, Ubvs may have the potential to be further developed to target DUBs and E3s in vivo.

3. Split Ubiquitin. Aside from engineering UB to directly probe components and substrates of the UB system, the stable UB fold has been employed as the basis for sensors to detect protein–protein interactions. Varshavsky’s laboratory demonstrated the capacity of an N-terminal fragment and a C-terminal fragment of UB to self-assemble in yeast. When a mutation was introduced, self-assembly occurred only when two binding partners that were fused to the separate UB fragments interacted. As the system was originally engineered, this assembly resulted in DUB-mediated cleavage of a protein linked to the C terminus of UB. This cleavage therefore served as a proximity sensor (Johnsson and Varshavsky, 1994) (Fig. 6A). This propensity for self-assembly has been adapted for use as sensors in the discovery of novel protein–protein interactions by screening of expression libraries in yeast. This complements yeast two-hybrid systems in that it can be used to screen for transcription factors and is not dependent on the ability of the assembled complex to localize to the nucleus (Laser et al., 2000). In the original iteration of this screening assay, the prey (library to be screened) and N-terminal UB fragment are expressed from a yeast plasmid library as N to C fusions. This was similarly done for the bait (protein for which partners are being screened for) and the C-terminal UB fragment. Critical to the assay is the fusion of a selection marker (initially Ura3) to the other (C-terminal) end of the bait-containing fusion protein. When bait and prey interact, the two halves of UB associate and the C-terminal Ura3 fusion is cleaved by a DUB. Once cleaved, the N-terminal Arg of the engineered Ura3 is exposed. N-terminal Arg is a primary destabilizing amino acid—a degron recognized by the yeast N-end rule E3 (Varshavsky, 1996). Thus, in the yeast cells in which such interaction occurs, Ura3 is ubiquitinated and degraded, allowing for selection by growth in media containing 5-fluoro-orotic acid—expression of Ura3 in yeast cells converts 5-fluoro-orotic acid to 5-fluorouracil, which is toxic to the cell. Plasmid can be recovered from transformed yeast and sequenced, and the identity of the interaction protein(s) determined. The split UB can be fused to other reporters to induce their DUB-mediated cleavage and can be used for screen interactions between specific membrane proteins (Stagljar et al., 1998), for detecting conformational changes in proteins, as well as for other purposes (Müller and Johnsson, 2008). This concept is now being adapted and used in mammalian cells as are other, more recently developed, protein biosensors such as split GFP, β-lactamase, luciferase, and dihydrofolate reductase (DHFR) (Wehr and Rossner, 2016).

B. Engineering E3s To Regulate New Cellular Targets

1. Engineering E3 Adaptors. CRLs are multisubunit complexes that recruit their substrates through adaptor proteins, recognizing specific degron sequences on substrates. The degron sequences on key signaling proteins are frequently mutated, which results in decoupling their regulation by E3s (Hayes and McMahon, 2009). It would be useful to engineer E3s to recognize the mutant degrons and restore substrate regulation through ubiquitination. In CRL3Keap1, the adaptor protein Kelch-like enoyl-CoA hydratase-associated protein (Keap1) binds to the substrate Nrf2. Mutations in the degron sequence allow mutant Nrf2 to evade proteasomal degradation by CRL3Keap1 and promote cancer development (Zhang et al., 2013a). To engineer a Keap1 that recognizes mutant Nrf2, we (Yin and colleagues) generated a library of the Kelch repeat domain of Keap1 displayed on the yeast cell surface with randomized residues at the degron-binding site. The yeast library was allowed to bind biotin-labeled degron peptide-bearing cancer-related mutations (Fig. 2B). Next, yeast cells were incubated with streptavidin–fluorophore conjugates to label cells displaying Kelch domain mutants that show high affinities for the biotinylated degron peptide. Sorting of yeast has identified Kelch-repeat domains with altered specificity for the Nrf2 mutant. It remains to be tested whether the engineered Keap1 could induce degradation of Nrf2 with a mutated degron in cells, but this work suggested that the yeast selection platform is amenable for engineering E3-substrate recognition (Zhang et al., 2013a).

2. E3 Fusions To Redirect the Ubiquitination Targets. E3s determine the specificity of substrate ubiquitination. Conceptually, E3s consist of at least two modular domains: a substrate-binding domain and a UB transfer domain (e.g., HECT, RBR, RING, U-box) (Fig. 1A). An E3 can therefore be targeted to a different substrate by mutating its substrate-binding region. For example, the UB ligase Cbl regulates epidermal growth factor signaling by targeting phosphorylated epidermal growth factor receptor for degradation. When the Src homology 2 (SH2) domain of Cbl is replaced with the SH2 domain from growth factor receptor–binding domain 2, the resulting chimeric E3 is also able to target HER2 for degradation (Li et al., 2007). Based on this idea, designer E3s can be constructed by fusing a substrate-binding domain with a UB transfer domain to target any undesirable cellular protein (Fig. 6B). Indeed, the SH2 domain of growth factor receptor–binding domain 2 fused to either the RING domain of Cbl or U-box domain from carboxyl terminus of Hsc70-interacting protein (CHIP) is sufficient to downregulate EGFR in lung cancer cells (Lee et al., 2014; Zhong et al., 2015). Similarly, a SH2 domain fused to the U-box domain of CHIP downregulates breakpoint cluster region protein–Abelson murine leukemia viral oncogene homolog 1 kinase and inhibits tumor growth of chronic myeloid leukemia cells in mouse xenografts (Ru et al., 2016).

When the protein targeted for degradation is an enzyme, its substrate may be engineered to serve as
the targeting domain for the designer E3. Type A botulinum neurotoxin (BoNT/A) prevents neurotransmission by cleaving soluble NSF attachment protein 25, a SNAP receptor protein in the presynaptic terminal, thereby preventing synaptic vesicle fusion. A designer E3 consisting of a mutant soluble NSF attachment protein 25 resistant to BoNT/A cleavage fused with the HECT domain from E6-AP successfully targets the toxin for proteasomal degradation (Fig. 6B). Besides the HECT domain from E6-AP, the substrate adaptor of Cul2 E3, has been fused to a nanobody directed against GFP (aGFP) to induce the degradation of GFP-tagged proteins (Fulcher et al., 2010). The choice of the UB ligase domain does not appear to be critical, although the optimal length of the linker region connecting the substrate-binding region to the ligase domain may differ for the ligase domains.

Designer E3s can also be constructed from multi-subunit E3 complexes such as members of the modular SCF E3 family of E3s. Among the SCF E3 subunits, the F-box protein serves to recruit the substrate for ubiquitination. A hybrid F-box protein was constructed by fusing β-transducin repeat containing protein with the N-terminal 35-amino-acid region of the human papilloma virus E7 protein to target the retinoblastoma protein for degradation (Fig. 6B) (Zhou et al., 2000). Recently, nanobodies—single-chain antibodies from camelid species—have been used as substrate adaptors due to their specificity and small size. A chimeric protein consisting of a nanobody against BoNT/A light chain fused to the minimal F-box domain of β-transducin repeat containing protein was demonstrated to target BoNT/A toxin for degradation. More importantly, it has been shown to accelerate neuronal recovery following intoxication, supporting the idea that the stability of the toxin accounts for its long-lasting effect (Kuo et al., 2011). In addition to F-box proteins, the von-Hippel-Lindau protein, the substrate adaptor of Cul2 E3, has been fused to a nanobody directed against GFP (aGFP) to induce the degradation of GFP-tagged proteins (Fulcher et al., 2016). The U-box E3 CHIP has also been fused to aGFP to induce the degradation of subunits of K⁺ ion channels bearing a GFP tag (Kanner et al., 2017). Because GFP fusions are commonly used for investigating the subcellular localization of proteins, designer E3s based on aGFP nanobodies will become versatile reagents for regulating the levels of these fusion proteins.

Instead of directly fusing to an E3 or E3 subunit, molecules that bridge the substrate and E3 can be used to target specific substrates for degradation. These proteolysis-targeting chimera (PROTAC) molecules contain a binding domain for the substrate joined by a variable linker to a binding domain for the E3. The first reported PROTAC used ovalicin linked to the phospho-degron peptide of IkB-κ to target MetAP-2 to SCPβ-TRCP (Sakamoto et al., 2001; Lai and Crews, 2017). Subsequent PROTACs have been developed to recruit substrates to the von-Hippel-Lindau E3 complex using a degron derived from HIF1α (Schneekloth et al., 2004). Similarly, thalidomide-like immunomodulatory drugs (IMiDs), have been found to be efficacious in multiple myeloma by binding to Cereblon (CRL4Cereblon) and targeting bound lymphoid-specific transcription factors (IKZF1 and IKZF3) for ubiquitination and proteasomal degradation (Krönke et al., 2014; Lu et al., 2014). These are now being adapted to target Cereblon to other proteins (Lu et al., 2015; Winter et al., 2015; Koduri et al., 2019).

Engineered E3 variants are not suitable for general therapeutic development without parallel development of specialized delivery systems. Nevertheless, they can be useful for studying the effects of degrading specific cellular targets. The recent development of small-molecule PROTACs and the discovery and development of IMiDs and molecular glue with better cell permeability will help realize the therapeutic use of these molecules. Readers are referred to recent reviews such as Lai and Crews (2017), Asatsuma-Okumura et al. (2019), and Pettersson and Crews (2019).

3. Engineered E3 Substrates. A simple way to target a specific protein for degradation is to attach a degron to the protein. For example, a protein can be targeted for degradation by the N-end rule pathway by fusion with UB at its N terminus (Varshavsky, 1996). When the UB is cleaved by cellular DUBs, the first amino acid following the UB determines the half-life on the protein according to the N-end rule. This approach can be used to bypass the normal degradative process in the cell and study the effects of protein degradation in cellular function. Fzo1 is the mitofusin that regulates outer mitochondrial membrane fusion in Saccharomyces cerevisiae (Hermann et al., 1998; Rapaport et al., 1998). Cells lacking Fzo1 have fragmented mitochondria and fail to respire when grown in nonfermentable carbon source (Cohen et al., 2008). Fzo1 is normally degraded by SCFMdm30. Surprisingly, cells lacking Mdm30 also have fragmented mitochondria even though Fzo1 accumulates in these cells. Expression of a Ub-Fzo1 fusion, which can be degraded by the N-end rule pathway independently of Mdm30, bypassed the requirement of Mdm30 for mitochondrial reparation and demonstrated that degradation of Fzo1 is required for outer mitochondrial membrane fusion (Cohen et al., 2011).

Fluorescence proteins fused to UB or a degron peptide could function as reporters of the activity of UB transfer cascades or in cells (Salomons et al., 2005). UB Gly76Val, GFP, in which the N-terminal UB has its C-terminal Gly76 mutated to Val, making it resistant to cleavage by DUBs, is rapidly degraded by the proteasome and can be used to monitor proteasome activity in cells and in vivo (Dantuma et al., 2000). To this end, an unstable GFP (GFP¹) was generated by fusing GFP with the CL1 degron peptide on its C terminus (Bence et al., 2005) (Fig. 6C). CL1 was isolated from a screen in yeast of C-terminal extensions that produced...
domains of both viral and yeast DUBs were generated. The means by which the CL1 fusion is targeted for degradation in human cells had not been known until recent studies suggested that it is degraded at the ER membrane through a UB-mediated ER-associated degradation pathway requiring the RING E3 TRC8, UBE2G2, and an ER-associated degradation accessory protein AUP1 (Stefanovic-Barrett et al., 2018; Leto et al., 2019).

In another approach to monitor degradation, a fusion protein was constructed in the order of following components: Renilla luciferase (RL), K48R mutant of UB, firefly luciferase (FL), and RhoB, a substrate of HECT E3 Smurf1. Both Renilla and firefly luciferase are tagged with multiple FLAG epitopes for affinity purification. When the construct is expressed in cells, DUB cleavage of the fusion generates an equal concentration of FL-UB and RL-RhoB in the cell. Because UB bears the K48R mutation, the FL-UB fusion is less likely to be ubiquitinated and therefore should be relatively stable. FL-RhoB would be readily ubiquitinated by SMURF1 that has been coexpressed in the cell and degraded. The ratio of the activity of RL and FL was measured to report the activity of SMURF1 in the cell (Tian et al., 2019). Such an engineered reporter system was used to screen SMURF1 inhibitors. Substrate proteins of an E3 could also be engineered to couple its stability with the presence of a small-molecule ligand. The split UB system (discussed above) was adapted to regulate substrate stability using the small-molecule rapamycin (Fig. 6D). In the system, the substrate was expressed as a fusion in the following order from amino to carboxyl terminus: degron peptide, FKBP12-rapamycin–binding protein (FRB), C-terminal fragment of UB, and substrate. Upon treatment with rapamycin, the FRB domain in the fusion protein would bind to FK506-binding protein (FKBP) that was fused with the N-terminal fragment of UB. This would bring the two UB fragments together to assemble a UB fold and trigger DUB cleavage to release the substrate protein from the degron-containing fusion. In this way, assembly of a UB fold upon addition of rapamycin serves to stabilize the substrates from ongoing degradation (Pratt et al., 2007).

C. Engineering Deubiquitinating Enzymes

DUBs have been expressed as fusions with either substrates or E3 enzymes to control the ubiquitination status of substrates in the cell (Stringer and Piper, 2011; MacDonald et al., 2017). To probe the role of ubiquitination in endosomal trafficking, fusions between endosomal membrane proteins and the catalytic domains of both viral and yeast DUBs were generated. It was found that the fusion proteins were blocked from sorting into multivesicular bodies and accumulated on the cell surface. This suggests the DUB fused to the target protein can remove UB from the fusion and affect its trafficking in the endocytic pathway (Stringer and Piper, 2011). The same group also generated fusions between all annotated yeast E3 open reading frames (ORFs) and the catalytic domain of the UL36, a DUB derived from Herpes virus. The E3-DUB fusions reversed ubiquitination of substrates of the fused E3s and thereby functioned as dominant-negative mutants of these E3s (Fig. 7A). These "antiligases" serve as specific inhibitors of the fused E3 and can be used to probe the function of these E3s in cells (MacDonald et al., 2017).

DUB can be engineered to transfer UB to a Lys. The native function of the yeast DUB Yuh1 is to cleave C-terminal extensions beyond Gly76 of UB. During the normal catalytic cycle of the DUB, Yuh1 would form an acyl-enzyme intermediate with the UB through a catalytic Cys residue, and, subsequently, the intermediate would hydrolyze to release UB and complete the catalytic cycle. Yeast cell surface display was used to screen a Yuh1 library to enhance its transamination activity so that, instead of hydrolysis, the acyl-enzyme intermediate would react with an active site Lys on the DUB so that UB forms an isopeptide conjugate with the mutant DUB. Such identity switch converts a DUB to an enzyme capable of transferring UB to itself independent of E1 and E2 (Chang and Strieter, 2018).

Recently, a protease targeting UBLs has been repurposed for cleaving UB chains to dissect chain topology. It was noted that leader protease (Lbpro) from the foot-and-mouth disease virus cleaves the UBL protein ISG15 before the C-terminal diGly motif to prevent its conjugation with viral proteins. This serves to disarm host antiviral responses (Swatek et al., 2018). Based on the modeled structure of Lbpro-UB complex, a L102W mutation was introduced to the enzyme-binding site of ISG15 so the engineered enzyme (Lbpro*) could recognize and cleave UB before its C-terminal diGly motif. When Lbpro* is used to process UB chains conjugated to a substrate protein, it would remove UB chains from the substrate, leaving the diGly motifs on Lys residues that served as sites of UB conjugation (Fig. 7B). This is in contrast to cleavage by DUBs, which cleave the isopeptide linkage of a UB chain after the diGly motif to regenerate unmodified Lys residues on UB. After Lbpro* cleavage, UB units derived from a chain in which there were no branch sites (i.e., involving a linkage from one Lys) would bear one diGly motif at the site of Lys linkage, whereas UB units derived from branch points to which distal UBs had been covalently linked to more than one Lys would bear two or more diGly motifs. Trypsin, the protease most commonly used to cleave proteins after positively charged Lys and Arg residues to generate peptide fragments for mapping ubiquitination sites, also cleaves UB chains into peptides with
Fig. 7. Engineering of DUBs and UBDs. (A) DUB fusion as antiligase to inhibit the ubiquitination of the E3 substrates. (B) Viral protease for ISG15 cleavage (Lbpro*) is engineered to recognize UB chains and cleave UB before the C-terminal diGly motif on each UB unit. After cleavage, the diGly peptide remained covalently linked to Lys residues of substrate proteins or of linked UB, providing a footprint of the UB modification. In contrast, native DUBs would disassemble UB chains into UB units with unmodified Lys, and trypsin would digest UB chains into peptides with diGly motifs attached to Lys at the ubiquitination sites. (C) GFP-TUBE fusion to image the polyubiquitinated proteins in the cell. Trypsin-resistant TUBE can protect UB chains from cleavage by proteasome or DUB to facilitate the purification of UB-modified proteins. (D) Bispecific antibody recognizing K11 and K48 linkages is assembled by knobs-into-holes mutations to bind to branched UB chains.
diGly motif conjugated to Lys at the linkage sites; however, the branching pattern of the UB chain is often lost because all UB units are fragmented into peptides (Fig. 7B). A UB-clipping method has been developed using Lbpro to cleave UB chains and deduce the chain topology based on the ratio of UB units bearing various numbers of diGly and the sites of diGly modifications. Analysis of mitochondrial ubiquitination reveals a complex topology of UB chains on depolarized mitochondria orchestrated by the dual action of Pink1 and Parkin during initiation of mitophagy (Swatek et al., 2019).

**D. Engineering Ubiquitin-Binding Domains**

UBDs can be employed in the laboratory as affinity reagents for purifying ubiquitinated proteins. For such purposes, glutathione S-transferase fusions to S5a/Rpn10, Rad23, and Dsk4 UBDs have been used to isolate UB conjugates from cell and tissue lysates (Layfield et al., 2001; Mayor and Deshaies, 2005). Multiple repeats of UBD motifs have been fused to generate affinity reagents for polyubiquitinated proteins (Hjerpe et al., 2009). These TUBEs would typically have four repeats of a UBD. Their polyvalency results in high-avidity reagents. Depending on the UBD, TUBEs can either recognize a variety of UB linkages or be linkage specific (Scott et al., 2015; Gao et al., 2016; Mattern et al., 2019). To facilitate the isolation of polyubiquitinated substrates of an overexpressed E3, constructs encoding FLAG epitope-tagged TUBEs that are resistant to trypsin have been developed for coexpression in cells with E3s. The TUBEs protect UB chains from both DUBs and proteasome activity (Yoshida et al., 2015) (Fig. 7C). After affinity purification from cell lysates, the immunoprecipitated material can then be subject to trypsin digestion and reimmunoprecipitation with an antibody directed against the UB isopeptide linkage (anti–GG-ε-K) (Kim et al., 2011; Xu and Jaffrey, 2011). This results in a highly enriched population of peptides corresponding specifically to substrate-UB linkages, which enhances the ability to detect these linkages by mass spectrometry. Sensors tracking the formation and the location of polyUB chains have also been developed by engineering UBD motifs (Fig. 7C). Triple repeats of the UIM from the UB-binding protein Vps27 (Vx3) were designed and shown to have a high degree of specificity for K63 UB chains. Vx3 was expressed in cells as a fusion with enhanced green fluorescent protein and used to image the location of newly-formed K63 UB chains during nuclear factor κB pathway activation (Sims et al., 2012). In another report, fusions of the UBD domains of both NFκB essential modulator (NEMO) and A20 binding and inhibitor of NFκB (ABIN) with GFP were generated to detect linear UB chains. In the same study, GFP was fused with the UIM of Rap80 to detect K63 chains in cells (van Wijk et al., 2012). The engineered UBD affinity reagents along with linkage-specific antibodies and GG-ε-K antibodies together constitute a powerful suite of tools for characterizing UB chain topology, reading the signals encoded in substrate-linked UB chains, and determining substrates linked to specific UB chains (Newton et al., 2008; Kim et al., 2011; Xu and Jaffrey, 2011).

Antibodies have also been generated as artificial UBDs to differentiate the linkage types of UB chains. To generate linkage-specific antibodies, fragment antigen-binding libraries with randomized complementarity-determining regions were displayed on phage and panned against linear, K11, K48, and K63 UB chains. Fragment antigen binding recognizing various linkages of UB chains were selected and expressed in an IgG format for the identification of cellular proteins modified with various types of UB chains (Newton et al., 2008; Matsumoto et al., 2010, 2012). Recently, K11 and K48 linkage-specific antibodies were assembled into a heterodimer with a knobs-into-holes strategy to generate a bispecific antibody to recognize K11/K48 chains of mixed linkages (Merchant et al., 1998; Yau et al., 2017) (Fig. 7D). To suppress homodimer formation between antibody heavy chains, a sterically bulky Thr to Trp mutation was introduced as a knob into the C_{3H3} region of the heavy chain specific for the K11 UB chain. In contrast, multiple mutations were installed on the C_{3H3} of the heavy chain specific for the K48 UB chain to create a hole and structurally complement the knob mutation. Half antibodies for K11 and K48 UB chains are expressed separately and assembled into bispecific antibodies as an affinity reagent to purify cellular proteins modified with branched UB chains. Enrichment of UB conjugates using the bispecific antibody found mixed K11/K48 chains present on substrates of E3 enzymes critical to cell cycle regulation by anaphase-promoting complex/cyclosome and recognition of misfolded proteins by UB protein ligase E3 component n-recogtin 5 (Yau et al., 2017).

**IV. Identifying E3 Substrates by Protein Engineering**

Dysfunction of specific UB ligases can have profound consequences that contribute to a host of disorders from cancer to neurodegeneration to autoimmune diseases (Rougeulle et al., 1997; Shimura et al., 2000; Nakayama and Nakayama, 2006; Kobashigawa et al., 2011; Lipkowitz and Weissman, 2011). Identifying the range of substrates for specific E3s is therefore of critical importance to understanding the regulation of cell physiology. The large number of E3s and their often transient interactions with substrates presents a significant challenge to the identification of target proteins. Although an oversimplification, methods for E3 substrate identification can be grouped into three categories—affinity binding to the native E3 or fragment of an E3; monitoring changes in protein stability
Fig. 8. Methods for identifying E3 substrates. (A) Identifying E3 substrates based on affinity binding between E3 and substrate proteins. (B) Identifying E3 substrates by monitoring changes in protein stability upon perturbation of E3 activity with a small-molecule inhibitor, short hairpin RNA, or by overexpression of an E3. Alternatively, ubiquitinated proteins are isolated from the cell; trypsin digestion generates peptides with diGly modifications at Lys residues. These branched peptides can be isolated by an anti-GG-e-K antibody, and their levels monitored by mass spectrometry (MS) to correlate with changes in protein ubiquitination upon modulation of E3 activity. (C) For UBAIT, an E3-UBA fusion is used to form conjugates with E3 substrates due to the in cis transfer of UB to substrates bound to E3. (D) An E3-UBA fusion was used to bind to polyubiquitinated proteins synthesized by the same E3 enzymes. (E) To generate a NEDDylator for E3 substrate identification, Ubc12, which is the E2 for the NEDD8 UBL, is...
or ubiquitination upon perturbation of E3 activity; and covalently or noncovalently trapping substrates using engineered E3 fusions (Fig. 8). Affinity-based approaches such as coimmunoprecipitation, yeast two-hybrid screening, and protein microarrays have been used to directly assess E3-substrate interactions (Fig. 8A). These methods have been central to identifying E3-substrate interactions, but are naturally limited by affinities and unrecognized adaptors. Activity-based screening approaches experimentally manipulate E3 activity and correlate this with changes in the stability or ubiquitination of specific proteins. One example is a method known as global protein stability (GPS) profiling, which tracks the stability of thousands of cellular proteins fused to a fluorescence protein (Yen and Elledge, 2008; Yen et al., 2008; Emanuele et al., 2011). The use of TUBEs and anti–GG-G-G-K antibodies, or both together, has also been employed to determine alterations in ubiquitination in response to E3 manipulation (see III. D. Engineering Ubiquitin-Binding Domains above) (Fig. 8B) (Kim et al., 2011; Xu and Jaffrey, 2011; Sarraf et al., 2013). These often highly data-intensive, proteomic approaches can be extremely powerful. Like all approaches, there are pitfalls, both physiologic and technical. One physiologic consideration is that perturbing the activity of one E3 may have downstream direct or indirect effects on the activity of other E3s (Weissman et al., 2011). Readers are referred to recent reviews on various proteomics methods (Kim et al., 2011; Harper and Tan, 2012; Ordureau et al., 2015; Beaudette et al., 2016; Iconomou and Saunders, 2016; Rayner et al., 2019). Protein engineering has added powerful tools for E3-substrate identification by engineering UB transfer enzymes with tailor-made activities to identify the cellular substrates of a specific E3.

A. E3 Fusions To Identify E3 Substrates

1. E3-Ubiquitin Fusions. UB-activated interaction traps (UBAIT) is a protein-engineering technique developed to identify substrates of specific E3s (Fig. 8C) (O’Connor et al., 2015). In UB-activated interaction traps, UB is fused to the C terminus of affinity-tagged E3s. After activation, the fused UB will preferentially interact with the linked E3 and be covalently attached to substrates of the E3-UB fusion. The E3-UB fusion therefore functions as a trap for substrates and can be purified through the affinity tag, followed by proteomic analysis. Using multiple E3s in both yeast and mammalian cells, proof of principle has been provided for this approach utilizing both HECT and RING E3s.

2. E3–Ubiquitin-Associated Domain Fusions. The UB ligase substrate-trapping method was developed to identify SCF E3 substrates (Fig. 8D) (Mark et al., 2014, 2016). In this method, F-box proteins, which are the substrate-specific recruiting subunits of the SCF E3s, are fused to UBA domains that bind to polyUB chains of K48 and K63 linkages. When a substrate that is recruited to a specific F-box undergoes ubiquitination, the proximity of the fused UBA will result in stable association of the substrate and copurification with the F-box fusion. This method has been used to profile the substrate specificities of eight F-box proteins and identified several new SCF substrates.

3. E2-E3 Fusions. The “NEDDylator” method has been developed to redirect a UB E3 to transfer NEDD8, a UBL protein to the E3 substrates (Fig. 8E) (Zhuang et al., 2013). Because the number of NEDDylated proteins is markedly less than the number of ubiquitinated proteins in the cell, NEDDylators would considerably narrow down potential substrates of a specific E3 to those with NEDD8 modifications. The NEDDylator approach was validated by a C-terminal fusion of the NEDD8-specific E2, UbcH12, with a mutant XIAP that has been inactivated by deletion of its RING domain. The N-terminal substrate-binding domain of XIAP in the fusion protein recognizes XIAP substrates and allows for their UbcH12-mediated NEDDylation. Using this approach, NEDDylated proteins could be affinity-purified from the cell lysates and assessed by proteomics. This technique has allowed the identification of PGAM5, a mitochondrial-localized protein phosphatase, as a validated XIAP substrate (Zhuang et al., 2013).

B. Identifying E3 Substrates by a Reporter Gene or Chemical Cross-Linking

1. Split Dihydrofolate Reductase Assays To Screen E3 Substrates in E. coli. The bacterium E. coli lacks an endogenous UB-conjugating system. This has now been taken advantage of to reconstitute highly specific E1- E2-E3 cascades. Such a system can be used to isolate ubiquitination substrates in a high-throughput screening platform. This is carried out by constructing fusions of the N-terminal fragment of DHFR with UB and fusions of the C-terminal fragment of DHFR with a library of potential substrates and expressing these along with ubiquitinating enzymes (Levin-Kravets et al., 2016) (Fig. 8F). Recognition of a substrate by its cognate E3 in E. coli leads to the transfer of UB to the substrate, juxtaposing the N- and C-terminal fragments of DHFR fused to UB and substrate, respectively. The reassembly of DHFR renders the cell resistant to fused to a defective E3 that lacks an intact ligase domain, but still retains the ability to interact with substrates. The presence of Ubc12 in the fusion facilitates transfer of NEDD8 to interacting proteins. This allows for their identification among the limited pool of NEDD8-modified cellular proteins. (F) To identify E3 substrates in an E. coli system, N- and C-terminal fragments of DHFR are fused to UB and a substrate library, respectively, and expressed in E. coli in which ubiquitination has been reconstituted using a specific E1, E2, and E3. Transfer of the DHFR-UB fragment from E1, E2, and E3 to a substrate protein results in reassembly of a functional DHFR. This renders these specific E. coli clones resistant to trimethoprim, allowing for identification of substrates by plasmid isolation and sequencing. (G) 125I-labeled phenylazide probe is conjugated to a Cys residue introduced into the UB chain by mutagenesis. This was used to identify proteasome subunits that come into contact with K48-linked tetraUB.
Fig. 9. OUT cascade. (A) Identifying E3 substrates by OUT. xUB is transferred through the engineered xE1-xE2-xE3 cascade to the substrates of a specific E3 in the cell. xUB-conjugated proteins are purified by tandem affinity purification under denaturing conditions with the 6xHis-biotin tag (HBT) fused to xUB and identified by liquid chromatography with tandem mass spectrometry (LC-MS/MS). (B) Engineering xUB-xE1 pair by phage display. xE1 with mutations in the adenylation domain was labeled with biotin and immobilized on the streptavidin plate. UB library with randomized C terminus was displayed on M13 phage and selected based on the formation of thioester conjugate with xE1 on the plate. (C) Engineering xE1-xE2 pair by phage display. E2 library with randomized N-terminal helix was displayed on the M13 phage and selected based on biotin-UB transfer from the
trimethoprim antibiotics. As a result, clones expressing the substrates of a specific E3 can grow on agar plates supplemented with trimethoprim and be identified. Using this method, new substrates of the yeast HECT E3 Rsp5 have been identified (Levin-Kravets et al., 2016).

2. Chemical Cross-Linking. Chemical cross-linking methods represent promising approaches to identify interactions in the UB system. Early work using chemical cross-linking in the UB system assessed the interaction of tetra-UB with the 19S cap of the mammalian proteasome. In this work, a tetrameric UB chain with K48 linkage was generated by a combination of enzymatic and chemical conjugation with a P37C mutation introduced in the distal UB of the tetramer chain. Using disulfide exchange, an $^{125}$I-labeled photoreactive reducible cross-linker was added to the Cys (Fig. 8G). The mixture of proteasome and $^{125}$I tetra UB was then subject to UV cross-linking. A specifically iodinated species was characterized by two-dimensional isoelectric focusing electrophoresis and peptide sequencing and identified as a 19S ATPase, Rpt5 (S6') (Lam et al., 2002). Another approach, based on chemical cross-linking, has also been employed to capture transient interactions of UB with proteasomes (Archer et al., 2008). In this case, a single UB is tagged with a short peptide that binds to a biarsenical fluorescent probe FlAsH, which is linked to biotin and a dihydroxyphenylalanine moiety. The tagged UB and proteasomes are coinubulated and treated with NaIO$_4$, which oxidizes the catechol ring of dihydroxyphenylalanine to ortho-quinone. This can then cross-link to nucleophilic residues on the proteasome. Using this method and employing yeast proteasomes, a different ATPase subunit of the proteasome was identified. It should be noted that, unlike other 19S subunits, neither of the ATPase subunits identified have known UBDs.

In another study, chemical cross-linkers were introduced into a substrate by UAA incorporation to provide proof of principle to capture transient interaction between a substrate and an E2. The UAA BprY, which is a Tyr analog with an alkylbromide group attached to the phenylhydroxyl residue of Tyr, was introduced into the substrate PCNA (Yang et al., 2017) (Fig. 2D). When PCNA and the E2 were reacted in the test tube or coexpressed in *E. coli*, PCNA was specifically conjugated to the E2 UBE2D3/Ubch5c, but not to other proteins/peptides. This conjugation occurs through S$_2$2 substitution of the bromo group of BprY by the catalytic Cys residue of E2.

C. Orthogonal Ubiquitin Transfer

1. The Concept of Orthogonal Ubiquitin Transfer. We (Yin and colleagues) have developed a method known as orthogonal UB transfer (OUT) to identify the direct substrates of E3s. In OUT, an affinity-tagged UB variant (xUB) is exclusively transferred through an engineered xE1-xE2-xE3 cascade to the substrates of a specific E3 (“x” designates the engineered forms of the enzymes or UB) (Fig. 9A) (Zhao et al., 2012b). By expressing affinity-tagged xUB and the engineered xE1-xE2-xE3 cascade in the cell, we are able to purify xUB-modified proteins and identify them by proteomics. In this way, we can profile ubiquitination targets of a specific E3 and elucidate the associated cell signaling pathways. We have engineered OUT cascades for the HECT E3 E6AP and the U-box E3s E4B and CHIP. Using this approach, we have identified and confirmed their substrate proteins in HEK293 cells. We have established OUT as an efficient platform to identify the direct substrates of an E3. To enable OUT, we need to engineer specific xUB-xE1, xE1-xE2, and xE2-xE3 pairs that have no cross-reactivities (orthogonal) with wt UB and wt E1, E2, and E3 enzymes (Fig. 9A). This will guarantee that xUB is confined on the single track of the OUT cascade on its way to the substrates of the xE3. xUB is expressed with the xE1-xE2-xE3 cascade in the cell with the N terminus of xUB tagged with 6×His and biotin tags (HBT-xUB) (Tagwerker et al., 2006). xUB-conjugated proteins are then purified by tandem affinity chromatography by Ni-NTA and streptavidin columns. Proteins bound to the streptavidin resin are digested by trypsin, and their identities revealed by liquid chromatography with tandem mass spectrometry. xUB-conjugated proteins identified in this way are potential direct substrates of xE3. Because we are only changing the E2-binding site but not the substrate-binding site to generate xE3, it is proper to assume that the xE3 targets identified by the OUT screen are the substrates of wt E3.

We took a bump-and-hole approach to generate the orthogonal enzymatic pairs (Hwang and Miller, 1987; Alaimo et al., 2001). The general protocol is to introduce mutations at the binding interface of the enzymes so that the mutated enzymes would reject the binding of their native partners. We then use phage or yeast cell display to select for complementary mutations that would restore enzyme binding and enable the transfer of xUB. The mutated enzyme pair and the wt pair would have the same binding modes, but they have mutually

xE1 enzyme. Phage displaying xE2-xE3 pairs with xE1 would be loaded with biotin-UB and selected by binding to the streptavidin plate. (D) Engineering xE2-xE3 pairs with HECT E3 by yeast cell sorting. HECT domain library was displayed on the surface of yeast cells as fusions to yeast cell surface protein Aga2. xUB was loaded on the HECT domain by transferring through the xE1-xE2 cascade. Mutant HECT that can connect with the xE1-xE2 pair was loaded with biotin-xUB and labeled with streptavidin conjugated to the PE fluorophore. Yeast cell sorting by fluorescence-activated cell sorting would identify xHECT domain that can pair with xE1-xE2 to assemble the OUT cascade. (E) Engineering xE2-xE3 pairs with U-box E3 by phage display. A U-box library of E4B was displayed on phage surface and selected based on xUB transfer from xE1 and xE2. Phage-displaying U-box mutants that can pair with xE2 would be loaded with biotin-UB and selected by binding to the streptavidin plate.
rejecting binding interfaces. As a result, the wt UB could not be transferred through the engineered OUT cascade, and xUB would not cross over to the native E1-E2-E3 cascade. The advantage of such orthogonality is that 1) xUB is guaranteed to be transferred to the target proteins of xE3 in the OUT cascade for unambiguous identification of E3 substrates; 2) xUB would not compete with wt UB in the cell so the engineered system would assume minimal disturbance on various biologic processes in the cell.

2. Engineering the xUB-xE1 and xE1-xE2 Pairs for the Orthogonal Ubiquitin Transfer Cascade. The first two stages of OUT are mediated by xUB-xE1 and xE1-xE2 pairs that deliver xUB to a designated xE2 (Fig. 9A). We first generated a xUB-xE1 pair with yeast UB and yeast E1 (Uba1) (Zhao et al., 2012b). We generated xUba1 by introducing two sets of mutations to yeast E1, one to the adenylation domain to block binding of wt UB, and one to the UFD to block binding of wt E2s. Using phage selection, we identified xUB with mutations R42E and R72E, and xUbc1, the yeast E2, with mutations in the N-terminal helix that allows it to interact with xUba1, but not the wt E1, and also enables xUB transfer from xUba1 to xUbc1 (Fig. 9, B and C) (Zhao et al., 2012b). To implement OUT in human cells, we transplanted mutations from yeast xUba1 and xUbc1 to human E1 (Uba1) and E2s (UbcH5b and UbcH7), respectively, based on the sequence homology of the E1 and E2 enzymes. We expressed human xUB, xUba1, and xUbcH5b or xUbcH7 in HEK293 cells and tested their orthogonality with wt E1 and E2 enzymes. We confirmed that xUB was exclusively loaded on xE1 or xE2 enzymes and it would not cross over to the wt cascade during the transfer (Wang et al., 2017a; Bhuripanyo et al., 2018). These studies established that we could use mutant transplantation to extend the OUT cascade across species and reach different E2 enzymes.

We were also able to use mutant transplantation to build OUT cascades with two human E1 enzymes, Uba1 and Uba6, to differentiate their functions. We transplanted mutations in the adenylation domain of yeast xUba1 to human Uba1 and Uba6. The E1 mutants, xUba1(A) and xUba6(A), could exclusively activate xUB and transfer it to wt E2 and E3 partners and their downstream targets. We separately expressed xUB-xUba1(A) or xUB-xUba6(A) pairs in the HEK293 cells and found that the two E1s have nonredundant roles in cell regulation because they had a distinct profile of UB transfer targets (Liu et al., 2017a,b). This study added the proof that the two E1s launch UB transfer through distinctive E1-E2-E3 cascades for cell signaling (Jin et al., 2007).

3. Constructing the Orthogonal Ubiquitin Transfer Cascade with HECT E3 E6AP. The R5E and K9E mutations in the N-terminal helix of xUbcH7 interfered with the transfer of xUB to wt HECT E3s (Huang et al., 1999). We used yeast cell surface display to identify a mutant of E6AP HECT domain that can uptake xUB from xUbcH7 (Fig. 9D). We then incorporated the mutant HECT into full-length E6AP and confirmed that it could transfer xUB from the xUba1-xUbcH7 pair to p53 in the presence of E6. We could thus use E6AP mutant as xE6AP to generate the OUT cascade. We expressed the OUT cascade of E6AP in HEK293 and compared the profiles of xUB-conjugated proteins with control cells expressing the xUba1-xUbcH7 pair without xE6AP. Three repeats of the proteomic screen revealed 144 proteins that we found to be consistently dependent on xE6AP for conjugation to xUB. Among these, we confirmed that E6AP ubiquitinated mitogen-activated protein kinase 1 (MAPK1), /beta-catenin, cyclin-dependent kinase 1, cyclin-dependent kinase 4, protein arginine methyltransferase 5 (PRMT5), and ubiquitin-like-domain-containing protein 8 (UbxD8) in cells and induced their degradation. /beta-catenin was confirmed to be an E6AP substrate by two reports published at the same time as ours (Kuslansky et al., 2016; Yi et al., 2017).

4. Constructing the Orthogonal Ubiquitin Transfer Cascades with U-Box E3 E4B and CHIP. xUbcH5b will not pair with either wt E4B or CHIP to mediate xUB transfer due to the mutations in the N-terminal helix of xUbcH5b that interferes with its binding to the U-box domain of the E3s (Benirschke et al., 2010). To identify U-box mutants complementary with xUbcH5b, we randomized residues in loop1 of E4B U-box that interact with K4E and K8E mutations in xUbcH5b (Bhuripanyo et al., 2018). We used phage selection to identify catalytic active mutants of E4B U-box based on xUB transfer (Fig. 9E). One U-box mutant (R1233K, L1236I, D1238H) from the selection restores xUB transfer from xUbcH5b. We also found the mutations from the E4B U-box can be transplanted to the CHIP U-box to activate CHIP for xUB transfer. We incorporated the mutations from phage selection into the full-length E3s to generate xE4B and xCHIP to assemble the OUT cascade. Profiling of their substrates with the OUT platform in HEK293 cells identified 185 and 225 potential substrates of E4B and CHIP, respectively. Among them we have confirmed that PRMT1, MAPK3, protein phosphatase 3 catalytic subunit /alpha, PGAM5, and ovarian tumor protease B1 are ubiquitinated by E4B and that PRMT1, MAPK3, protein phosphatase 3 catalytic subunit /alpha, and /beta-catenin were ubiquitinated by CHIP in HEK293 cells based on in vitro ubiquitination, E3 silencing by short hairpin RNA, and cycloheximide chase assays.

The OUT platform is not automatically adaptable to all E3s, and this is a limit to its general application. To guarantee the exclusive delivery of xUB to designated E3s and its substrates, we need to engineer an OUT cascade unique for each E3. We see potential for transplantation of mutations to expand the OUT cascade within a class of E3s. The E2-binding regions in HECT, U-box, RBR, and RING E3s are homologous.
within each class, and, as proof of principle, we have succeeded in extending the OUT cascade by transplanting mutations from the U-box domain of E4B to CHIP (Bhuripanyo et al., 2018). If transplantation is not efficacious for an E3 of interest, the phage and yeast display methods could be used to engineer individual xE2-xE3 pairs (Wang et al., 2017a; Bhuripanyo et al., 2018). Another limitation of OUT is that coexpression of HBT-xUB and the xE1-xE2-xE3 cascade, although successful in HEK293 cells, could be a challenge to other cell types. The recently developed clustered regularly interspaced short palindromic repeats (CRISPR/Cas9) genome-editing method may provide an opportunity to introduce the OUT cascade into the native genetic background to identify E3 substrates (Cong et al., 2013).

V. Conclusions

In the years since the original characterization of the UB system as a means of targeting proteins for proteasomal degradation, the field has undergone explosive growth. This has been fueled by the development of proteasome inhibitors, the recognition of the various distinct classes of UB ligases and deubiquitinating enzymes, and the realization that ubiquitination can, through UB-binding domains, alter the fate and function of proteins in ways other than targeting to proteasomes. We now understand that the UB system touches on essentially all cellular processes, and its dysfunction is linked to an ever-expanding list of human diseases. It is therefore critical that we achieve an in-depth mechanistic understanding of this system.

In this review, we have surveyed the rapidly increasing number of approaches to dissecting and manipulating the UB system through protein engineering. Depending on one’s perspective, protein engineering can be conceived of as synthetic chemistry on the scale of protein molecules or, alternatively, synthetic biology enabled by laboratory-directed evolution in generating designer proteins. It is particularly fitting that, for a system first discovered by biochemists and initially characterized in vivo through yeast genetics, advances in our understanding would now be guided by this toolbox of approaches.

The use of powerful mutation-based protein screening and selection platforms is being complemented by genetically expressed fusion proteins and structure-based mutations to explore sequence space, interrogate and affect interactions, and dissect and alter catalytic activities. This genetic approach is naturally integrated with advances in chemical biology, allowing for site-specific conjugation, protein ligations, and the introduction of reactive chemical functionalities through unnatural amino acids.

The UB system is a rich source of molecular targets in disease. In the context of emerging genetic and protein delivery technologies, there is great potential for the development of UB system therapeutics, derived from protein engineering, in personalized medicine. As we look optimistically ahead and consider the trajectory of advances, it is reasonable to posit that, in the next decades, protein-based reagents will enter the clinic joining proteasome inhibitors, IMiDs, E1 inhibitors, and other UB system-based small molecules in the treatment of human disease.

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Authorship Contributions

Wrote or contributed to the writing of the manuscript: Zhao, Tsai, Jin, B. Wang, Y. Wang, Zhou, Carpenter, Weissman, Yin.

References


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Deubiquitinating enzymes (DUBs) play a crucial role in regulating the cellular levels of ubiquitin and its conjugates. DUBs work in a site-specific manner and are regulated by their binding partners. The current focus is on understanding the structural, dynamical, and functional properties of these enzymes to discover new DUBs as therapeutic targets. This review provides a comprehensive, integrated overview of the structural basis of DUBs, the mechanism of their enzymatic activity, and the identification of functional domains and residues that are involved in substrate docking and catalytic activity. Understanding the structural and functional properties of DUBs is essential for developing new therapeutic strategies.


Protein Engineering in the Ubiquitin System


