Minireview

Chemically Induced Cellular Proteolysis: An Emerging Therapeutic Strategy for Undruggable Targets

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Traditionally, small-molecule or antibody-based therapies against human diseases have been designed to inhibit the enzymatic activity or compete for the ligand binding sites of pathological target proteins. Despite its demonstrated effectiveness, such as in cancer treatment, this approach is often limited by recurring drug resistance. More importantly, not all molecular targets are enzymes or receptors with druggable ‘hot spots’ that can be directly occupied by active site-directed inhibitors. Recently, a promising new paradigm has been created, in which small-molecule chemicals harness the naturally occurring protein quality control machinery of the ubiquitin-proteasome system to specifically eradicate disease-causing proteins in cells. Such ‘chemically induced protein degradation’ may provide unprecedented opportunities for targeting proteins that are inherently undruggable, such as structural scaffolds and other non-enzymatic molecules, for therapeutic purposes. This review focuses on surveying recent progress in developing E3-guided proteolysis-targeting chimeras (PROTACs) and small-molecule chemical modulators of deubiquitinating enzymes upstream of or on the proteasome.

Keywords: deubiquitinating enzyme, induced proteolysis, PROTAC, small-molecules, ubiquitin-proteasome system, undruggable target

INTRODUCTION

Cellular proteolysis can be achieved through two major pathways: the ubiquitin-proteasome system (UPS) and the lysosomal degradation pathway (Ciechanover, 2005). Ubiquitination, a type of protein post-translational modification, is covalent conjugation of the small, stable protein, ubiquitin through the formation of an isopeptide bond between a Gly of ubiquitin and a Lys on the substrate. This reaction occurs through an enzymatic cascade of ubiquitin-modifying enzymes comprising E1 (ubiquitin activating enzyme), E2 (ubiquitin conjugating enzyme), and E3 (ubiquitin ligase) (Finley, 2009; Komander and Rape, 2012). Although ubiquitin conjugation may serve non-proteolytic functions, its most recognized function is targeted protein degradation via the 26S proteasome, a multi-protein protease complex (Fig. 1) (Finley and Chau, 1991). Although ubiquitin conjugation may serve non-proteolytic functions, its most recognized function is targeted protein degradation via the 26S proteasome, a multi-protein protease complex (Fig. 1) (Finley, 2009; Komander and Rape, 2012). Ubiquitination process is tightly controlled, reflecting the critical importance of protein turnover rates for cellular function. This requirement strongly suggests that the ubiquitination reaction must be remarkably versatile. Notable in this context, more than 600 E3 ligases are known, helping to explain the observed substrate specificity (Deshaies and Joazeiro, 2009); moreover, deubiquitinating enzymes (DUBs) can reverse this reaction by break-
ing down ubiquitin polymers (Wilkinson, 1997). E3 ligases transfer ubiquitin from a ubiquitin-charged E2 complex to the substrate and this reaction can be repeated, producing a polyubiquitinated substrate that is recognized by the proteasome. The specificity of E3 ligases suggests that E3 inhibitors could produce more targeted effects and thus might be less harsh than E1- or E2-targeting drugs. In fact, several studies have shown that specifically targeting E3 ligase may be therapeutically effective against certain cancers (Chan et al., 2013; Shangary et al., 2008; Vassilev et al., 2004). In addition to E3 ligases, the human proteome contains approximately 100 DUBs belonging to at least six subfamilies (Komander et al., 2009), which play key roles in ubiquitin-mediated proteolytic pathways as well as other biological processes, including (a) rescue of protein substrates from degradation by ubiquitin removal, (b) editing of ubiquitin chains to regulate the function or half-lives of protein substrates, and (c) ubiquitin recycling into free chains or ubiquitin monomers.

Since the initial discovery of the proteolytic machinery, a plethora of evidence has shown that the UPS is closely associated with various diseases, such as cancers and neurodegenerative diseases (Popovic et al., 2014). These findings have garnered immense attention in treating such diseases by targeting the UPS. For example, following a path similar to that for kinase inhibitors, bortezomib and its progeny have been successfully developed as FDA-approved anticancer drugs for the treatment of multiple myeloma (MM) and other blood cancers, and in fact, these agents are proteasome inhibitors that block the proteolytic reaction (Manasanch and Orlowski, 2017; Richardson et al., 2005; Stewart et al., 2015). Moreover, a number of E3 ligase inhibitors and DUB inhibitors have been tested in preclinical or clinical phase studies, representing novel strategies for target-directed therapies (Goepinath et al., 2016; Farshi et al., 2015; Weathington and Mallampalli, 2014).

Traditional active site-directed therapies, which generally employ small-molecule inhibitors and antibodies, are designed to inhibit the activity of target proteins. Small-molecule chemicals have proven effective for inhibiting several oncogenic enzymes, including protein kinases (e.g., imatinib and erlotinib), histone deacetylases (e.g., belinostat), and other oncogenic enzymes, including protein kinases (e.g., olaparib) (Manasanch and Orlowski, 2017; Richardson et al., 2005; Stewart et al., 2015). In addition to E3 ligases, the human proteome contains approximately 100 DUBs belonging to at least six subfamilies (Komander et al., 2009), which play key roles in ubiquitin-mediated proteolytic pathways as well as other biological processes, including (a) rescue of protein substrates from degradation by ubiquitin removal, (b) editing of ubiquitin chains to regulate the function or half-lives of protein substrates, and (c) ubiquitin recycling into free chains or ubiquitin monomers.

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An emerging new paradigm of disease treatment—"targeted protein degradation (TPD)"—promises to help overcoming the limitations of typical active site-directed therapies (Zhou, 2005). TPD aims to induce degradation of target proteins, especially those considered “undruggable”, by harnessing the endogenous protein quality control machinery—the UPS. One elaborate approach is to induce ubiquitin tagging and subsequent target protein degradation through E3-guided proteolysis-targeting chimeras (PROTACs). Small-molecule DUB inhibitors that act upstream of or on the proteasome can also be utilized for antagonizing substrate deubiquitination. This review will focus on recent progress in the development of TPD, specifically PROTACs and DUB inhibitors, and their therapeutic potential. Figure 1 highlights PROTAC and DUB inhibitors as chemical proteolytic inducers in the UPS pathway.

**INDUCED PROTEOLYSIS BY PROTACS**

PROTAC, an E3-guided proteolysis-targeting chimera, is a bifunctional molecule composed of three parts: an E3 ligase binding ligand, a substrate binding ligand, and a linker between the two. E3 ligase generally requires sequential recognition events through multi-step E1/E2/substrate reactions (Deshaies and Joazeiro, 2009). PROTACs can simplify this process—they transiently recruit target substrate and E3 ligase in close proximity and facilitate the ubiquitination of the substrate (Coleman and Crews, 2018). As a result, target proteins become more efficiently polyubiquitinated and undergo degradation by the 26S proteasome (Fig. 1). This idea originated with the elegant work of Sakamoto and colleagues, in which the researchers developed a chimera linking the E3 ligase SCF-ß-TRCP (Skp1/Cullin/F box-ß-TRCP) to methionine aminopeptidase 2 (MetAP2) (Sakamoto et al., 2001). This chimeric molecule, PROTAC-1, contains a 10-amino acid IKeR phosphopeptide that can be recognized by β-TRCP at one end, while the other side is an MetAP1 inhibitor, ovalicin. PROTAC-1 indeed promotes degradation of MetAP1 in a ubiquitination-dependent manner. Since first-generation PROTACs had low cell permeability because of the peptidic sequence, next-generations adopted von Hippel-Lindau (VHL) as an E3 ligase instead of SCF-ß-TRCP (Schneekloth et al., 2004). This variant, developed by Crews and colleagues, exploited a short hydroxyproline-containing peptide derived from the VHL substrate, hypoxia-inducible factor 1-α (HIF1α), and they applied the resulting chimeras for selective degradation of androgen receptor (AR) and FK506-binding protein 12 (FKBP12) in intact cells. Although the hydroxyproline-containing degron displayed improved
Fig. 1. Mechanisms of induced protein degradation by PROTACs and DUB inhibitors. (Left) General scheme of ubiquitin-proteasome system. An enzymatic cascade of E1-E2-E3 transfers ubiquitin to Lys residues on the substrate. As a consequence, the polyubiquitinated substrate is recognized by the 26S proteasome and undergoes degradation. (Right) Induced proteolysis by PROTAC and/or DUB inhibitor. PROTAC links E3 and the target protein, enhances E3-mediated ubiquitination, and promotes degradation of the target molecule. By contrast, DUBs acting upstream of the proteasome (violet) or on the proteasome (olive drab) can inhibit degradation through substrate deubiquitination. Therefore, DUB inhibitors may facilitate the proteasomal degradation pathway by antagonizing deubiquitination.

Table 1. Representative examples of PROTACs

| E3 ligase/HyT | E3 binder/HyT (type) | Target binder (type) | Target | Reference |
|--------------|----------------------|----------------------|--------|-----------|
| β-TRCP      | lβx phosphopeptide (P) | Ovalicin (S) | MetAP2 | Sakamoto et al., 2001 |
| cIAP        | MeB5 (S) | ATRA (S) | CRABP-I, II | Itoh et al., 2010 |
|             | BE04 (S) | Ch55 / Estrone / DHT (S) | RAR / ER / AR | Itoh et al., 2011 |
|             | BE04 (S) | KHS108 (S) | TACC3 | Ohoka et al., 2014 |
|             | BE04 (S) | Alkyl chloride analog (S) | HaloTag | Tomoshig et al., 2015 |
| CRBN        | Thalidomide (S) | JQ1 / SLF (S) | BET family / FKB12 | Winter et al., 2015 |
|             | Pomalidomide (S) | Bosutinib, Dasatinib (S) | BCR-ABL | Lai et al., 2016 |
|             | Thalidomide (S) | Aminopyrazole analog (S) | CDK9 | Robb et al., 2017 |
|             | Pomalidomide (S) | Centinib (S) | ALK | Zhang et al., 2018 |
|             | Lenalidomide (S) | HIB97 (S) | BET family | Zhou et al., 2018 |
|             | Pomalidomide (S) | Brustubulin derivative (S) | BTK (Bruton’s Ty kinase) | Buhimschi et al., 2018 |
| Pomalidomide (S) | HDAC inhibiting aldehydes (S) | HDAC6 | Yang et al., 2018 |
| Pomalidomide (S) | Thalidomide (S) | HDAC6 | Yang et al., 2018 |
|             | Thalidomide (S) | HDAC inhibiting aldehydes (S) | HDAC6 | Yang et al., 2018 |
|             | Thalidomide (S) | SiruIn Rearranging Ligand (SirReal) (S) | Siruin 2 (Sirt2) | Schiedel et al., 2018 |
| Keap1       | Keap1 binding motif (P) | Tau binding motif (P) | Tau | Lu et al., 2018 |
| MDM2        | Nutlin-3 (S) | SARM (S) | AR | Schneekloth et al., 2008 |
|             | HiFα degron (S) | DHT / AP21998 (S) | AR / FKB12 | Schneekloth et al., 2004 |
|             | HiFα degron (P) | DHT / Estradiol (S) | AR / ER | Rodriguez-Gonzalez et al., 2008 |
|             | HiFα degron (P) | TrkA degron / ErbB3 degron (P) | FRS2α / PI3K | Hines et al., 2013 |
| HiFα derivative (S) | Vandetanib / Phenoxy TZD analog (S) | RIKP2 / ERRα | Bondeson et al., 2015 |
|             | HiFα derivative (S) | JQ1 (S) | BRD4 | Zengerle et al., 2015 |
|             | HiFα derivative (S) | Chloroalkane analog (S) | Halotag | Buckle et al., 2015 |
|             | HiFα derivative (S) | Triazolo-diazepine acetalide (S) | BET family | Raina et al., 2016 |
|             | HiFα degron (P) | CPP-tri_a (S) | AKT | Henning et al., 2016 |
|             | HiFα derivative (S) | Dasatinib (S) | c-ABL | Lai et al., 2016 |
|             | HiFα degron (P) | EN300-72284 (S) | Smad3 | Wang et al., 2016a |
|             | HiFα degron (P) | tau binding motif (P) | Tau | Chu et al., 2016 |
| HiFα derivative (S) | TBK1 ligand (S) | TBK1 (TANK-Binding Kinase 1) | Crow et al., 2018 |
| HiFα degron (P) | PD-PEM (P) | ERα | Jiang et al., 2018 |
| HiFα derivative (S) | Lapatinib, Gefitinib, Aftatinib (S) | EGFR, HER2, c-Met | Burslem et al., 2018 |
| HyT         | Adamantane (S) | Haloalkane reactive linker (S) | HaloTag | Neklesa et al., 2011 |
|             | BocArg (S) | Trimethoprim / EA, Thiobenzofurazan (S) | DHFR / GST | Long et al., 2012 |
|             | Adamantane (S) | TX1-85-1 (S) | HER3 | Xie et al., 2014 |
|             | Adamantane (S) | RUS59063 (S) | AR | Gustafson et al., 2015 |

a: HyT, Hydrophobic tagging; b: P, peptide, S, small molecule
cell applicability, this VHL-recognized peptide still lacked ideal drug-like properties. Recent efforts have further validated MDM2/Nutlins, Cereblon (CRBN)/thalidomide, cIAP ligands, and optimized VHL ligands as E3-recruiting moieties for target protein degradation in PROTAC technology (Table 1). Notably, landmark papers from Bradner and Crews groups demonstrated that small-molecule VHL- and CRBN-directed PROTACs are highly effective in degrading estrogen-related receptor α (ERRα) and Bromodomain-containing protein 4 (BRD4) in animal cancer models (Bondeson et al., 2015; Winter et al., 2015). PROTACs can be designed to behave like traditional small-molecules, yet their distinctive mode of action provides innovative therapeutic opportunities. There has also been considerable effort devoted to PROTACs into commercial drugs, exemplified by the recent foundation of Arvinas and C4 Therapeutics (Neklesa et al., 2017).

PROTACs have several advantages over active site-directed therapies, ideally overcoming the drawbacks of small-molecule inhibitors and therapeutic antibodies. The most prominent feature of PROTAC is that induced protein degradation, in principle, renders conventionally undruggable targets druggable. As previously noted, typical active site-directed drug targets constitute only about 13% of the thousands of disease-related proteins, mostly enzymes or membrane receptors. This clearly indicates that transcription factors, scaffolding proteins, and toxic protein aggregates have remained therapeutically intractable. PROTACs, by linking ligands and appropriate E3 ligases, can induce the proteolysis of traditionally undruggable proteins, including transcriptional regulators (e.g., Myc, Gli, β-catenin, and STAT family) and signaling scaffolds (e.g., ERBB3, KSR, Gab family, β-arrestin, BCL10, and AKAPs) (Coleman and Crews, 2018; Neklesa et al., 2017). Protein aggregates are particularly interesting targets because they are common features of several neurodegenerative diseases, such as tauopathies and amyotrophic lateral sclerosis (ALS) (Ross and Poier, 2004). Recent studies have shown that the low efficacy problem of early PROTACs can be substantially resolved, as evidenced by the apparent nanomolar range of EC50s of more recently developed ones (Lai and Crews, 2017). Analogous to E3-guided PROTAC, the ‘hydrophobic tagging (HyT)’ technology has also been developed to destabilize target proteins and/or recruit chaperones for induced protein degradation (Huang and Dixit, 2016; Lai and Crews, 2017). The novel repertoire of E3 ligases (e.g., Keap1), HyT binders, and target ligands has rapidly expanded, strongly suggesting that PROTAC may serve as a platform technology (Chu et al., 2016; Lu et al., 2018). Table 1 summarizes representative examples of PROTACs.

### SMALL-MOLECULE INHIBITORS TARGETING DEUBIQUITINATING ENZYMES UPSTREAM OF OR ON THE PROTEASOME

Reversing the ubiquitination process is exclusively fulfilled by deubiquitinating enzymes (DUBs), numbering nearly 100 in the human genome (Komander et al., 2009). To date, six subfamilies of DUBs have been identified: (1) ubiquitin carboxyl-terminal hydrolases (UCHs), (2) ubiquitin specific proteases (USPs), (3) ovarian tumor like proteases (OTUs), (4) JAMM/MPN metallocarboxypeptidases, (5) Machado-Jacobs disease proteases (MJDs), and (6) motif interacting with Ub-containing novel DUB family (MINDY) (Abdul Rehman et al., 2016; Clague et al., 2013). DUBs should represent promising drug targets because they become increasingly implicated in various human diseases (Harrigan et al., 2018). For example, a growing number of DUBs (e.g., USP28, JOSD1, UCHL1, CSN5, USP9x, USP10, USP11, USP22, and USP48) were found to be overexpressed in diverse cancer types. Moreover, genetic alterations of DUBs (e.g., USP6/Tre2 and USP28) can be truly oncogenic in certain cancers (Fraile et al., 2012; Sacco et al., 2010). The isopeptidase activity of DUBs can be selectively targeted by inhibiting the catalytic sites with drug-like compounds, and in this sense, they are typical targets of active site-directed inhibitors. In an interesting twist, however, most cellular proteins are degraded by the ubiquitin-mediated proteasome pathway; thus, DUBs in principle should antagonize the destruction of these proteins (Fig. 1). Indeed, deubiquitination can occur on histones, transcription factors, and other regulatory proteins, so the pool of target substrates must far exceed the number of conventionally druggable targets. Therefore, similar to PROTACs, small-molecule DUB inhibitors may also induce the degradation of previously intractable targets. While of great therapeutic interest, only a handful of DUB inhibitors have been reported, and none has reached in advanced clinical trials, providing a valuable opportunity for developing first-in-class therapeutic DUB inhibitors (Farshi et al., 2015; Table 2).

Analogous to the cell cycle, deubiquitination reactions at two points-upstream of and on the proteasome—may serve as critical checkpoints for ubiquitin-mediated degradation pathways (Fig. 1). DUBs acting upstream of the proteasome may act as a “rescue crew”, salvaging specific substrates by deubiquitination before they are engaged in degradation. Accordingly, chemical blockers of the DUB may accelerate the turnover of the specific substrate for proteasome-mediated degradation. For instance, USP7/HAUSP is the most well-recognized drug target because of its cellular role as a regulator of the tumor suppressor p53. USP7 deubiquitinates MDM2, an E3 ligase of p53, and in turn destabilizes p53 (Colland et al., 2009). Thus, by stabilizing p53, a USP7 inhibitor may be beneficial for the treatment of certain cancers. The USP7 inhibitors, HBO 19,818 and P5091, were previously reported to selectively inhibit USP7 in vitro and in vivo, and importantly, P5091 showed cytotoxicity in relapsed MM cells derived from cancer patients (Chauhan et al., 2012; Reverdy et al., 2012). More recently, a series of USP7 inhibitors have been discovered and among them, a non-covalent inhibitor FT671 and a covalent inhibitor FT827 were identified by molecular-based screening (Turnbull et al., 2017). These compounds were found to be highly selective for USP7 in counter-screening against a panel of DUBs including USP10 and USP47, which are also inhibited by P5091. In vivo experiments in cancer cell lines and MM.1s xenograft animal models demonstrated that FT671 induces p53 stabilization and MDM2 degradation, leading to anti-tumor activity via USP7 blocking. Another NMR and structure-based screening study identified the USP7 inhibitors,
### Table 2. Representative examples of deubiquitinating enzyme inhibitors

| Target DUB | DUB inhibitor |
|------------|---------------|
| UCH-L1     | LDN-57444     |
|            | Liu et al., 2003 |
| USP1/UAF1  | Pimozide      |
|            | Chen et al., 2011 |
|            | GW7647        |
|            | ML323         |
| USP7       | HBX 41108     |
| (USP47\(a\)) | Colland et al., 2009 |
|            | HBX 19818     |
|            | Reverdy et al., 2012 |
|            | HBX 28258     |
|            | Tian et al., 2011 |
|            | Chauhan et al., 2012 |
|            | P22077        |
|            | P5091         |
|            | P5091 analogue\(a\) |
|            | FT671         |
|            | Turnbull et al., 2017 |
|            | FT827         |
|            | GNE-6640      |
|            | Kategaya et al., 2017 |
|            | XL188         |
|            | Lamberto et al., 2017 |

(continued)
| Target DUB | DUB inhibitor |
|------------|---------------|
| USP7 (USP47) | ![Compound 4](Gavory et al., 2018) |
| USP9x/USP5/USP14/UCH37 | WP1130 (Kapuria et al., 2010) |
| USP14 (UCH37) | IU1 (Lee et al., 2010) 
| | CuPt (Liu et al., 2014a) 
| | b-AP15 (D’Arcy et al., 2011) |
| USP30 | 15-oxospiramilactone (Yue et al., 2014) 
| | MF-094 (Kluge et al., 2018) 
| | MF-095 |
| RPN11 | ![Capzimin](Li et al., 2017) |
| Broad DUB inhibitor | PR-619 (Tian et al., 2011) |

a: reported to inhibit both USP7 and USP47; b: reported to inhibit both USP14 and UCH37

GNE-6640 and GNE-6776 (Kategaya et al., 2017). These compounds may selectively interfere with K48 linkage-directed ubiquitin chain cleavage mediated by USP7, suggesting that K48-linked substrates such as MDM2 could be susceptible. More recently, an elegant fragment-based screen combined with structure-guided medicinal chemistry identified a highly potent and selective USP7 inhibitor, “compound 4” (IC$_{50}$ = 6 nM). This allosteric inhibitor showed
strong anti-proliferative effects against several cancer cell lines with equal or even greater efficacy compared to known clinical MDM2 antagonists (Gavory et al., 2018). A mitochondria-localized DUB, USP30 may also represent a promising therapeutic target due to its involvement in mitophagy-related Parkinson’s disease as well as cancers. USP30 antagonizes Parkin-mediated ubiquitination on multiple mitochondrial substrates (Bingol et al., 2014; Liang et al., 2015). Recently, a potent USP30 inhibitor MF-O94 was developed through high-throughput screening and subsequent structure-activity relationship (SAR) studies of acyl benzensulphonamide derivatives, and this compound showed the increased mitophagy in C2C12 cells (Kluge et al., 2018).

Targeting DUBs on the proteasome may also offer an exciting strategy for induced protein degradation. There are three major and distinctive DUBs on human proteasome: USP14, UCH37, and RPN11 (de Poot et al., 2017; Finley, 2009). USP14 and UCH37 may rescue substrates from degradation prior to the proteasome’s commitment step, whereas RPN11 is coupled to degradation. Finley and colleagues have screened out highly selective USP14 inhibitors, IU1 and its derivatives, and showed that their treatment promotes the degradation of proteopathic substrates in neurodegenerative disease models (Boselli et al., 2017; Lee et al., 2010; 2016). USP14 inhibitors may uncheck and bypass the deubiquitination-mediated proteolytic checkpoint on the proteasome under certain conditions of proteotoxic stress. By contrast, the proteasome 19S DUB inhibitors, b-AP15 and VLX1570, were reported to suppress tumor progression by inhibiting both USP14 and UCH37 activities (D’Arcy et al., 2011; Wang et al., 2015; 2016b). b-AP15 treatment leads to accumulation of polyubiquitinated conjugates and inhibition of protein degradation. Recently, capzimin was identified as a potent and specific RPN11 inhibitor (Li et al., 2017). Capzimin, a quinoline-8-thiol (8-TQ) derivative, induced the stabilization of proteasome substrates and inhibited cancer cell proliferation probably through the unfolded protein response (UPR). Unlike IU1, the anti-tumor effects of b-AP15 and capzimin might rely on ‘restrained protein degradation’ rather than induced proteolysis.

**FUTURE PERSPECTIVES**

Here we described PROTACs and DUB inhibitors-two emerging strategies of chemically induced proteolysis that utilize the endogenous ubiquitin-proteasome system to inhibit previously undruggable targets. While certainly bearing tremendous promise for new therapeutic applications, these approaches could also face several challenges. For example, current PROTACs are orally unavailable, probably due to its relatively large size, typically 700-1000 Da. Their pharmacokinetic properties also need to be improved for better drug metabolism. Besides, only a few E3 ligases have been exploited, and not all E3 ligases might be co-expressed with target proteins in specific tissues, which makes diagnostics arduous (Huang and Dixit, 2016). PROTAC optimization-E3 ligase selection, ligand availability, and linker design—is another challenging issue. In this context, ligand screening can be performed by advanced screening tools, such as computeraided drug design and DNA-encoded small molecule libraries, which can be accomplished on the order of $\sim10^3$ compounds in a single vial (Chan et al., 2015). Although DUB inhibitors might be more orally bioavailable, their specificity and utility still remain to be explored. Given the smaller pool of DUB members compared to over 600 E3 ligases, DUB inhibitors may target only a subset of substrates with limited specificity. Nevertheless, one can envisage that ‘degradation inducers’ might pioneer the valuable therapeutic strategies and provide more advanced platform technologies, leading to a new era of UPS-related drug development. It will be also interesting to investigate potential combinatorial treatment with E3-guided and DUB-guided degradation inducers or to design new classes of chemically induced proteolysis chimeras recruiting substrates, E3s, DUBs, chaperones, or proteasomes.

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