DUB esterase activity further decodes ubiquitin’s enigma

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Posttranslational modifications of proteins are often key to understanding their biological function, localization, and fate. In particular, the covalent attachment of ubiquitin, a small 76-amino acid polypeptide, to substrates has attracted recent attention and is being exploited to generate novel drugs capable of removing pathogenic targets in a selective fashion (proteolysis-targeting chimeras) (1). Conjugation of ubiquitin can be reversed by deubiquitinating enzymes (DUBs), reflecting additional regulation (2). Control for reversing protein ubiquitylation has been the subject of Satpal Virdee and coworkers (3), who have developed a DUB amino acid profiling assay that led to the discovery of a class of ubiquitin esterases, classically assigned as ubiquitin specific proteases. As reported in PNAS, the authors show, for a subset of DUBs from the Machado–Josephin Domain (MJD) family, previously with unknown function, their ability to cleave ubiquitin not only from lysine-based “classical” isopeptide bonds but also from ubiquitin moieties linked to serine and threonine side chains via ester linkages (3).

Protein ubiquination seems predominantly restricted to the attachment of ubiquitin C termini to lysine (Lys) residues, forming “classical” isopeptide and, in the case of linear ubiquitin chains, peptide bonds (Fig. 1A). This is, in part, due to the unique length of the hydrocarbon chain of Lys that appears to be most optimal for E2/E3 enzymes’ reaction and transition of ubiquitin molecules to substrates (4). At the same time, there has been a precedent for ubiquitin covalent linkages other than to lysine epsilon amine side chains representing “canonical ubiquitylation.” For instance, in the context of ubiquitin conjugation to substrate proteins, the ubiquitin C terminus is transiently reacting with cysteine (Cys) thiol groups to form ubiquitin thioesters, in particular, with E2 and E3 enzymes (Fig. 1B) (5). At present, there is little evidence of DUBs reversing this reaction, although ubiquitin thioester formation may regulate DUB function itself, as proposed for USP4 and USP15 (6). Evidence for noncanonical ubiquitylation on alternative amino acid residues was suggested by several observations. First, the discovery of enzymatically catalyzed ubiquitin adenosine 5′-diphosphate (ADP) ribosylation was made, leading to the attachment to protein substrate’s serine (Ser) residues via phosphodiester bonds (ubiquitin attachment via arginine through the ribosyl group), mediated by bacterial pathogen-derived effectors (7).

Ubiquitin ADP ribosyl Ser phosphodiester can be hydrolyzed by the esterase activity of SidJ (8). Second, protein ubiquitination via esterification of Thr and Ser residues was demonstrated to occur via the E2 enzyme Ube2j2 (9) and the E3 ligase HOIL-1 (10). Also, Satpal Virdee’s team initially discovered MYCCBP2, an E3 ubiquitin ligase of the RING-Cys relay family, capable of performing this reaction (11, 12) (Fig. 1C). However, it was previously unknown whether mammalian DUBs might target these unusual sites for deubiquitination. To address this, the same group has now extended this line of research and developed representative model substrates and screen 53 DUBs for nonlysine activity, thereby providing important insights into DUB function (3). Satpal Virdee’s team found that, generally, ubiquitin-specific protease (USP) and ubiquitin C-terminal hydrolase (UCH) class DUBs exert both isopeptidase and esterase activity with comparable kinetics. In contrast, ovarian tumor domain (OTU) DUBs had little esterase activity, with the exception of TRABID and virally encoded vOTU. Notably, Satpal Virdee’s team (3) uncovered that a poorly studied class of DUBs of the MJD family (13) has potent and highly selective threonine esterase activity (Fig. 1C). These findings suggest that nonlysine ubiquitination appears to be common and coexisting with its conventional counterpart, possibly serving distinct biochemical purposes (3).

These intriguing observations made by the Virdee team open up a plethora of routes for ubiquitin biology, but there remain unanswered questions. For instance, from a thermodynamic point of view, thioester bonds are less stable as compared to (iso)-peptide bonds under aqueous conditions, such as aqueous, neutral pH and physiological temperature (37 °C) (14, 15). This is,

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at least in part, a possible reason why analytical detection of non-
lysine ubiquitylation on proteins has been a challenge, as their
chemically labile nature may lead to premature hydrolysis during
sample preparation and/or measurement in biochemical or mass
spectrometry-based assays. On the other hand, such modifications
make biological sense, for example, as short-lived reaction interme-
diates, which is the case for ubiquitin thioesters during transthioy-
lation and trans(iso)-peptide reactions, until formation of the final
product, typically a Lys-Ub protein substrate, which is chemically
more stable. Considering all these variations of ubiquitin-linked
chemistry, perhaps it does not completely come as a surprise that
nonlysine-linked ubiquitin also exists beyond Cys residues, such as
ubiquitin esters with Ser/Thr residues of targeted proteins as de-
scribed by Virginia De Cesare et al. (3).

It remains to be determined, in future studies, what exactly the
stability and lifetime of protein adducts are that contain intracellular
esters that are expected to be less stable as compared to thioesters
and, in particular, (iso)-peptide bonds. The authors argue that, under
certain circumstances, this may not necessarily be the case (3). As
stated above, ester bonds are much more prone to hydrolysis than
are peptide bonds, which suggests that ubiquitin chains based on ester
linkages, if proven to widely exist in vivo, may play a more transient role
than ubiquitin chains based on conventional (iso)-peptide linkages.

This is relevant, as the entire study has been performed
predominantly with recombinant enzymes and artificial substrates,
so the biological significance as well as how widespread ubiquitin
esterification of protein substrates is remains to be demonstrated
(3). There is precedence in the literature describing how ubiquiti-
nation of Ser, Thr, or Lys residues on the cytoplasmic tail can in-
duce ERAD of MHC-I by viral E3 ligase mK3 (16). Also, Ser-Thr
ubiquitination mediates down-regulation of BST-2/tetherin and
relief of restricted virion release by HIV-1 Vpu (17), and ester
bonds were observed between ubiquitin and components of
the MYD88 Myddosome (18). However, the role of DUBs—in par-
ticular, how selective OTUs and the MJDs, such as JosD1, with
apparent selective esterase activity, may act as ubiquitin-specific
carboxylic ester hydrolases (CEHs) in this cellular context—is not
yet fully understood (19). According to the De Cesare study, many

Fig. 1. DUB esterases as part of the ubiquitin code arsenal. Protein substrate (de)-ubiquitylation on different amino acid side chains. (A) Classical ubiquitin (Ub) conjugation and deubiquitylation of lysine “iso peptide bonds.” (B) Cys-ubiquitylation yielding Ub-thioesters, mostly prevalent in transthioylation reactions involving E1, E2, and E3 enzymes. (C) Ser/Thr ubiquitylation and cleavage via DUBs (in red) from the USP, UCH, MJD, and OTU subfamilies. Scissors indicate DUB esterase activities (in red).
DUBs have apparent cysteine protease and esterase activities, and structural features discriminating between these two enzymatic activities are at the beginning of being unraveled. For instance, CEHs are predominantly serine hydrolases and classified as clans A-D (20), whereas a large subset of DUB cysteine proteases belong to the papain-like family of the clan CA (21). Dissecting this will open up another layer of regulation that controls ubiquitin-driven biological processes.

In summary, the discovery made in the study by Virginia De Cesare et al. (3) expands the biochemical concept of protein ubiquitylation, suggesting Ser/Thr ubiquitin conjugates may be widespread and intertwined with classical ubiquitin modifications. This has potentially wide implications for our general understanding of the ubiquitin system and associated biology in the normal, but also a pathophysiological, context.

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