Expanding the ubiquitin code through post-translational modification

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Abstract

Ubiquitylation is among the most prevalent post-translational modifications (PTMs) and regulates numerous cellular functions. Interestingly, ubiquitin (Ub) can be itself modified by other PTMs, including acetylation and phosphorylation. Acetylation of Ub on K6 and K48 represses the formation and elongation of Ub chains. Phosphorylation of Ub happens on multiple sites, S57 and S65 being the most frequently modified in yeast and mammalian cells, respectively. In mammals, the PINK1 kinase activates ubiquitin ligase Parkin by phosphorylating S65 of Ub and of the Parkin Ubl domain, which in turn promotes the amplification of autophagy signals necessary for the removal of damaged mitochondria. Similarly, TBK1 phosphorylates the autophagy receptors OPTN and p62 to initiate feedback and feedforward programs for Ub-dependent removal of protein aggregates, mitochondria and pathogens (such as Salmonella and Mycobacterium tuberculosis). The impact of PINK1-mediated phosphorylation of Ub and TBK1-dependent phosphorylation of autophagy receptors (OPTN and p62) has been recently linked to the development of Parkinson’s disease and amyotrophic lateral sclerosis, respectively. Hence, the post-translational modification of Ub and its receptors can efficiently expand the Ub code and modulate its functions in health and disease.

Keywords mitophagy; phosphorylation; post-translational modification; ubiquitin

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See the Glossary for abbreviations used in this article.

Ubiquitin—a versatile post-translational modifier

The cellular functions of most proteins are regulated by post-translational modifications (PTMs). As these PTMs are inducible and reversible, they allow eukaryotic cells to dynamically respond to external stimuli by modulating intracellular signal transduction pathways. Among the best-studied PTMs are ubiquitylation, phosphorylation and acetylation [1].

Ubiquitylation is a post-translational modification in which ubiquitin (Ub), a 76 amino acid protein, is attached to target proteins through the sequential actions of an E1 Ub-activating enzyme, an E2 Ub-conjugating enzyme and an E3 Ub ligase [2–4]. This cascade requires the initial ATP-dependent activation of Ub by the E1, which links the C-terminal glycine residue of Ub via a thioester bond to a cysteine residue within the E1 active site [4]. The activated Ub intermediate is then transferred to the catalytic cysteine residue of an E2 enzyme. The E3 Ub ligase conjugates the C-terminal glycine of Ub via an isopeptide bond to the ε-amino group of the target lysine (K) of the substrate [5]. E3 ligases are divided into families, based on their catalytic domain structure and mode of catalysis. HECT E3 ligases temporarily accept activated Ub, whereas RING E3 ligases catalyse the direct transfer of Ub from the E2 to the substrate. Parkin belongs to the RBR family, which is formed by RING-HECT hybrid E3 ligases [6].

An important feature of ubiquitylation is that more Ub molecules can be added onto one of its own seven lysine residues or onto the ε-amino terminus of the first Ub. Depending on which lysine residue within Ub is utilized to anchor the subsequent Ub molecule, chains of different types and lengths are formed. Linkages can occur on M1, K6, K11, K27, K29, K33, K48 or K63 of Ub [7,8]. Branched chains and mixed linkage types also occur [9,10]. These different linkage types function as signals that are specifically recognized by Ub-binding proteins, which relay the signal to ultimately determine the cellular response or regulate the modified protein by changing its enzymatic activity, localization or stability [11]. The removal of Ub(s) attached to target proteins (known as deubiquitylation) is catalysed by deubiquitylating enzymes (DUBs) and represents an important regulatory mechanism of Ub signals in vivo [12].

Interplay between PTMs

Post-translational modifications can engage in extensive crosstalk, with the potential to either positively or negatively modulate signalling networks [13] (Fig 1). For example, acetylation can compete with ubiquitylation at lysine residues and thus enhance the stability of target proteins by suppression of ubiquitylation [14,15]. On the other hand, acetylation of target proteins can trigger their subsequent Ub-mediated degradation by promoting ubiquitylation of another lysine within the substrate protein [16].
In particular, ubiquitylation and phosphorylation are closely interlinked and mutually affect each other’s functions. For example, phosphorylation can stimulate or inhibit the activity of E3 Ub ligases [17–19] or DUBs [20]. It can also promote or prevent interactions with their target proteins by controlling their intracellular localization [21]. Additionally, phosphorylation can mark proteins for recognition by E3 Ub ligases [22] and DUBs [23], or can inhibit substrate recognition [24]. Similarly, the phosphorylation of UBDs can alter the ability of a protein to recognize specific Ub modifications, thus affecting signal propagation or protein fate [25–27]. Two prominent examples of the interplay between phosphorylation and ubiquitylation are cell cycle control and NF-κB activation. During cell cycle progression, the cyclin-dependent kinases (CDKs) and E3 ligases APC and SCF coordinate the controlled degradation of proteins [28]. In the NF-κB pathway, multiple E3 ligases initiate the proximal ubiquitylation cascade—including TRAFs, IAPs, Pellino and LUBAC—that can be further regulated through phosphorylation events, modifying downstream substrate ubiquitylation. The ubiquitylation of substrates is recognized by the UBAN domain of IKK component NEMO, resulting in NF-κB activation [29–31].

This regulatory and functional interplay also holds true for the reverse situation, as ubiquitylation can activate or inactivate kinase activity by non-degradative [32,33] and degradative mechanisms [34]. For example, E3 ligases of the IAP family (inhibitor of apoptosis protein) have been recently identified as key regulators of ERK5 activation pathway, via physical and functional disassembly of the ERK5-MAPK module by non-degradative ubiquitylation [35]. Recent data indicate that the phosphorylation of Ub itself has an enormous impact on the ubiquitylation cascade, as it can promote polyUb chain formation, but also inhibit certain E2s, E3s and DUBs, as discussed in detail below. This review summarizes the current understanding of the regulation of Ub by the well-studied PTM removal of PTMs determines substrate fate.

**Localization and stoichiometry of Ub phosphorylation**

Ub can be phosphorylated at multiple sites in yeast and mammalian cells (Fig 2). Phosphoproteomic analyses have shown that Ub may be phosphorylated at T7 [36–38], T12 [36,39], T14 [39–41], S20 [42–44], S57 [38–40,45–50], Y59 [51–54], S65 [49,55–59] and T66 [39] (Fig 2). However, little is known about the responsible kinases/phosphatases and stoichiometries of these Ub phosphorylation sites, and the functional relevance of most of them remains to be identified.

Multiple factors likely impact on the function of Ub phosphorylation within cells, including its intracellular localization and the stoichiometry of Ub or substrate phosphorylation. For example, a subpopulation of Ub can be modified on a specific site by a locally restricted kinase, indicating that only a small but locally enriched population of modified Ub is functionally relevant and thus detected by proteomic analyses. For instance, the stoichiometry of p-S65-Ub is below 0.5% of total Ub in exponentially growing yeast and ~1.5%

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**Glossary**

AMSH  | associated molecule with the SH3 domain of STAM  
APC   | anaphase promoting complex  
AR-JP | autosomal recessive juvenile parkinsonism  
Cif   | cycle inhibiting factor  
CHBP  | cycle inhibiting factor homolog in Burkholderia pseudomallei  
CK2   | casein kinase 2  
DUB   | deubiquitylating enzyme  
ERK   | extracellular-signal-regulated kinase  
HAT   | histone acetyltransferase  
HDAC  | histone deacetylase  
IAP   | inhibitor of apoptosis  
IKK   | IkappaB kinase  
Pellino | LUBAC  
MAPK  | mitogen activated protein kinase  
MS    | mass spectrometry  
MUL1  | mitochondrial E3 ubiquitin protein ligase 1  
MAPL  | mitochondrial-anchored protein ligase  
MULAN | mitochondrial ubiquitin ligase activator of NF-κB  
NEDDD8| neural precursor cell expressed, developmentally down-regulated 8  
NF-κB | nuclear factor kappa-light-chain-enhancer of activated B cells  
PGAMS | phosphoglycerate mutase 5  
PINK1 | PTEN-induced putative kinase 1  
RING | really interesting new gene  
RBR  | RING in between RING  
SCF  | Skp, Cullin, F-box containing complex  
SNARE | soluble N-ethylmaleimide-sensitive factor attachment receptor  
TBK-1 | TANK-binding kinase 1  
TRAF | TNF receptor-associated factor  
UBA  | ubiquitin-associated domain  
UBAN | ubiquitin binding in Abin and NEMO  
UBD  | ubiquitin-binding domain  
UBZ  | ubiquitin-binding zinc finger domain  
Ubi  | ubiquitin-like  
USP  | ubiquitin-specific protease  

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**Figure 1. Generation and control of PTMs.**

External stimuli result in the generation of different PTMs on target proteins. Phosphorylation, acetylation and ubiquitylation can influence each other and are attached to and removed from substrate proteins by different enzymes. Phosphorylation occurs mainly on serine, threonine and tyrosine residues, whereas acetylation and ubiquitylation target lysine residues. The attachment or removal of PTMs determines substrate fate.

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**Table 1.**

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Box 1: Acetylation of ubiquitin

Reversible lysine acetylation regulates several cellular processes—such as gene expression, cell cycle and cellular metabolism—and can impair phosphorylation-dependent protein–protein interactions [11,112]. The surface of Ub is theoretically amenable to most PTMs and acetylation of Ub was recently detected in cells [41]. The acetylation of Ub at K6 and K48 is dynamically regulated by all three classes of histone deacetylases (HDACs) and blocks the synthesis of Ub chains. K6- or K48-acetylated Ub can be activated by the E1, transferred to the E2, but cannot be used for E2-mediated K11-, K48-, or K63-linked Ub chain assembly (see schematic illustration). Ub acetylation, which neutralizes the positive charges of lysine residues, was proposed to affect the non-covalent interaction of Ub with E2 enzymes [41]. Through overexpression of acetyl-mimetic Ub (K6Q), K6 acetylation was shown to repress Ub chain elongation on substrates. Ideally, the conclusions of this study would need to be confirmed with K6-acetylated Ub rather than acetyl mimetic Ub, as acetyl-lysine and glutamine are chemically different. In addition, only a small proportion of Ub is acetylated in vivo. Using mass spectrometry, acetylation on K6 and K48 was found on only 0.03 and 0.01% of total Ub in cells [42]. Interestingly, the acetylation of Ub occurs on two lysine residues that are frequently used for chain elongation. How the competition of ubiquitylation and acetylation is regulated on K6 and K48 residues of Ub is not yet known, but it might have a significant impact on the regulation of mono- versus polyubiquitylation of target proteins, and hence drastically impact the fate of these proteins. However, histone 2B is the only known endogenous substrate of acetylated Ub in cells [41]. UBOs recognize different Ub chain linkage types [11], and there is evidence that S65-phosphorylation of Ub may modulate the binding affinity to certain Ub-binding domains present in autophagic adaptor proteins [81]. As K6 and K48 of Ub are close to the hydrophobic patch, which is the major binding site for Ub-binding motifs, acetylation or phosphorylation might have a direct impact on the interaction of Ub receptors with Ub. The identification of Ub-binding motifs specific for acetyl- or phospho-Ub would allow the identification of additional pathways regulated by Ub PTMs.

Phosphorylation of ubiquitin modulates the ubiquitylation cycle

Since the ubiquitylation cascade involves a variety of different players [5], it can be assumed that the phosphorylation of Ub influences components of the ubiquitylation machinery as well as Ub-binding proteins. Current studies point to significant phosphorylation-mediated structural and biophysical changes of Ub and Ub chains, which might have a wide spectrum of consequences on Ub biology [49, 59]. In yeast, Ub S65E is preferentially incorporated into K6- and K11-chains, while K27-linkages are disfavoured (K63 could not be assessed by MS due to the S/E mutation) [49]. Moreover, proteins modified by S65E-Ub are more ubiquitylated, suggesting that either the disassembly of chains containing Ub S65E is impeded [49, 59], or Ub S65E increases the conjugation activity of certain E2-E3 pairs, or both [49]. Indeed, Ub chain generation in vitro is hindered through the phosphorylation of S65 on Ub, as it impairs the activity of several E2 and E3 enzymes in mammalian cells [59, 61] and yeast [49]. E1-mediated E2 charging with p-S65-Ub is not affected, whereas a subset of E2-conjugating enzymes (mammalian: UBE2R1, UBE3E1, UBE2T, UBE2K, UBE2N/UBE2V1, and yeast: Ubc13/Mms2s, Ubc1) were unable to efficiently discharge phosphorylated Ub to form Ub chains, even though S65 does not contact the active site or interact with any other part of the enzyme (with the exception of Ubc13/Mms2) [49, 59, 61]. Although one could speculate that the close proximity of p-S65 to K63 might disturb K63-linked chain formation, several of the tested K63-specific E2-E3 pairs were not inhibited in chain formation [55, 59] and K63-linked p-S65-Ub chains could be detected by mass spectometry [55, 62]. Also the mammalian E3 ligases TRAF6 and HOIP (RFN31) were unable to use phosphorylated Ub for Ub chain synthesis, whereas phosphorylated Ub had no negative impact on mammalian Nedd4L, cIAP1- or yeast Rsp5-mediated chain assembly [49, 59]. Nevertheless, there is currently no evidence that p-S65 exists as monoUb in cells or that it is used for chain synthesis. Interestingly, Parkin E3 ligase activity
Box 2: Deamidation of ubiquitin, NEDD8 and Pup

Deamidation is a ubiquitous protein modification in which an amide is converted into an acid. This irreversible conversion of glutamine and asparagine into glutamic acid and aspartic acid, respectively, can occur non-enzymatically without any reactant species or catalyst. Enzymatic in vivo deamidation of glutamine, in contrast to asparagine, has also been reported. Deamidation generally regulates protein turnover in vivo; however, several bacterial virulence factors have also evolved to use enzymatic glutamine deamidation to modify the functions of host proteins. At physiological pH, deamidation introduces a negative charge at the deamidation site and can thus alter the protein structure [113,114]. Deamidation can be detected in most proteins, including Ub and the Ub-like modifiers NEDD8 and Pup (see schematic illustration). The bacterial effectors CHBP from *Burkholderia pseudomallei* and Cif from *Enteropathogenic E. coli* (EPEC) use a novel mechanism for the inhibition of Ub chain elongation in host cells [115]. Cif/CHBP harbour a papain-like hydrolytic fold and deamidate Ub at Q40 to generate E40 in *vivo* and in infected cells. Ub chain synthesis catalysed *in vitro* by several different E2/E3s is abolished in the presence of catalytically active CHBP. Covalently modified Ub E40 does not affect the formation of E1 or E2 Ub thioester intermediates, but blocks the discharge of Ub from E2 to E3. How a single charge substitution in Ub (Q40E) could have such a dramatic effect on Ub chain elongation remains unclear. Crystallographic analysis of Ub Q40E could determine whether Ub deamidation significantly influences its 3D structure, thereby impeding chain formation, or whether the E2-binding motif is negatively affected. In infected cells, the production of Ub E40 impairs TNFα-induced signalling, thereby preventing downstream NF-κB-dependent transcription. Hence, CHBP-mediated deamidation of Ub in *Burkholderia*-infected cells is an effective virulence mechanism that undermines the host Ub system [115]. Notably, CHBP and Cif also target the Ub-like protein NEDD8, in which Q40 is conserved [115,116]. NEDD8 is conjugated to cullins—which are part of cullin-RING Ub E3 ligases (CRLs)—and stimulates E3 ligase activity [117,118]. However, this effect is abolished if deamidated NEDD8 is attached to cullins, resulting in impaired degradation of CRL substrates, such as p27, Nrf2, HIF-1α and RhoA [116]. CHBP-mediated NEDD8 deamidation kills macrophages (while other cells remain viable), thus constituting an additional virulence mechanism for *Burkholderia* and EPEC to counteract macrophage-mediated host defence. CHBP/Cif-mediated NEDD8 deamidation is obviously detrimental during infection, but could be exploited to serve therapeutic purposes, as the neddylation-mediated activation of CRL has a well-established role in cell proliferation, cancer progression and several other diseases [119,120]. Modification by deamidation is not only relevant for Ub and NEDD8, but also for Pup, a bacterial Ub-like modifier. In analogy to Ub in eukaryotes, Pup is attached to lysine residues of proteins destined for proteasomal degradation. However, unlike Ub, Pup requires deamidation of its C-terminal glutamine to glutamate before its attachment [121].

is activated by p-S65-Ub, but also hindered by concentrations exceeding 20% of total Ub in *in vitro* ubiquitylation assays [55,59]. Taken together, S65 phosphorylation can impact the activity of certain E2-E3 combinations to assemble polyUb chains, although the structural and mechanistic basis for this phenomenon remains unclear.

Phosphorylation of Ub chains also affects the activity of DUBs [49,59,61]. The phosphorylation of S65 on Ub impedes efficient deubiquitylation by mammalian USP8, USP15, USP30, ataxin-3, USP2, AMSH and USP5 [49,59]. Interestingly, the site of phosphorylation in diUb (whether distal or proximal) is highly relevant: USP2 (a promiscuous DUB) and AMSH (a K63-linkage-specific DUB) are inactive against the doubly phosphorylated diUb [61], whereas AMSH is inhibited by phosphorylation of the proximal Ub and USP2 is inhibited by phosphorylation at the distal site but not affected by the modification of the proximal Ub [61].

S65 phosphorylation of ubiquitin activates Parkin

The functional interplay between phosphorylation and ubiquitylation has been extensively studied in the context of mitochondrial autophagy (mitophagy), mediated by PINK1 and the E3 Ub ligase Parkin [63,64] (Fig 3A). This process involves the selective engulfment of damaged mitochondrial material by a specialized double membrane that forms the autophagosome and delivers its cargo to the lysosome for degradation [63,64]. Disturbances in this pathway, for example by mutations affecting the activities of PINK1 or Parkin, are associated with neurodegenerative diseases such as Parkinson’s disease [65–67].

Mechanistically, mitophagy is induced through stabilization and autophosphorylation of PINK1 at the mitochondrial outer membrane (MOM) upon mitochondrial depolarization [68,69]. PINK1 activity is required to recruit Parkin to mitochondria and to activate its E3 ligase activity (Fig 3A and B). Parkin then promotes the ubiquitylation of a variety of MOM proteins [70,71], thereby priming the organelle for recognition by autophagic adaptor proteins that link it to the autophagosomal membrane [63]. However, how PINK1 activates and recruits Parkin to mitochondria has been unclear until recently. Crucial insights into this process were gained through quantitative proteomic approaches and novel enrichment strategies for ubiquitylated and phosphorylated proteins. PINK1 was shown to have two critical phosphorylation targets required for optimal activation of Parkin, mitochondrial ubiquitylation and mitophagy: serine 65 of the UbI domain of Parkin [72,73], and serine 65 of either Ub [55–58] or Ub chains assembled on mitochondria [55,62] (Fig 3A).

Before mitochondrial depolarization, interactions between the Parkin N-terminal UbI domain, its C-terminus [74] and an inhibitory interface involving its catalytically important RBR domains [75–77] render cytoplasmic Parkin inactive. This native inactive state is partially released upon phosphorylation of serine 65 in the UbI domain; however, to be fully active p-S65-Parkin also requires the presence of S65-phosphorylated Ub generated by PINK1. Phosphorylated Ub is hardly used by Parkin for conjugation *in vitro*, but it non-covalently binds to Parkin’s RBR region [57,58] and accelerates the discharge of Ub loaded on the E2 UBCH7 [57,58,60]. p-S65-Ub (and not p-S20-Ub or p-S57-Ub) binds Parkin to allosterically trigger its E3 ligase activity [58,60,78].
Recent structural studies have provided crucial atomic insights into the role of p-S65-Ub binding to Parkin, which initiates the multistep process required for its activation (Fig 3B) [78–81]. Parkin is inhibited at multiple levels and disruption of the autoinhibited conformation primes Parkin for activation. Binding of the Ubl domain to the RING1 interface keeps the molecule in a closed conformation [74], the REP element blocks access for the E2-Ub loading [75] and the catalytic cysteine—which is C431 in the RING2— is occluded by the RING0 domain, thus blocking catalysis [75–77]. Binding of p-S65-Ub to Parkin leads to a conformational reorganization in its RING1 domain, and displacement of the Ubl domain and REP element, opening a binding surface to engage Ub conjugated with E2 [78–81]. Minimal conformational changes in the RING2 domain upon binding of p-Ub reveal that the thiol group of the active site C431 is capable of accepting ubiquitin from the E2–Ub conjugate [80]. Thus, the conformational change that enables access of the E2-Ub conjugate in the proximity of the active site seems to be an essential step for the catalytic activation of Parkin [80]. The p-Ub-binding site is located on the opposite side of the Ubl-binding patch, on a hinge region at RING0/RING1, indicating that antagonistic binding of the Ubl and pUb is mediated through negative allostery [80,81]. Mutations in the p-S65-Ub-binding interface of Parkin (K151, H302, R305, A320 and G284) impede Parkin activation and its recruitment to depolarized mitochondria [78–81]. Interestingly, the G284R mutation in Parkin is associated with AR-JP and the AR-JP mutation L283P and cancer-associated H279P mutations might also interfere with p-Ub binding [79]. Parkinson’s disease missense mutations in the Parkin IBR-RING1 interface (R275W, Q311H, G328E) could interfere with sequential p-Ub binding and Ubl release [80]. Further AR-JP mutations that occur in the Ubl domain (R42P, A46P and R33Q) [74] could result in loosening of the Ubl from the RING1 and thus in enhanced E3 ligase activity [81]. The structural model nicely accounts for the two required steps in Parkin activation: binding of p-S65-Ub to Parkin and PINK1-mediated phosphorylation of S65-Ubl of Parkin (Fig 3A and B). Yet, the situation in vivo might be more complex, as Ub-replacement experiments in cells indicate that Parkin can efficiently ubiquitylate a subset of mitochondrial substrates in cells expressing non-phosphorylatable Ub S65A (approximately 90% of total Ub) [60].
Figure 3. PINK1-mediated phosphorylation of ubiquitin and Parkin.
(A) Upon mitochondrial damage, PINK1 is recruited and activated at the mitochondrial surface resulting in the sequential phosphorylation of both Ub and the Parkin Ubl domain on their respective Ser65 sites. This leads to Parkin activation and ubiquitylation of multiple substrates on the mitochondrial outer membrane (MOM), which in turn become favourable substrates for PINK1. In such a way, PINK1 and Parkin generate high-density p-S65-Ub chains on MOM proteins. This increases the binding and retention time of Parkin on mitochondria, leading to the amplification of Ub signals. Phosphorylation of Ub chains also serves as a commitment step, as p-Ub chains are resistant to deubiquitylation by the majority of DUBs. The multiple ubiquitylation signals then commit mitochondria for degradation by attracting multiple autophagy receptors, such as OPTN and NDP52. (B) Conformational domain rearrangements for Parkin activation. (a) Inactive PARKIN: Closed auto-inhibited structure of full-length PARKIN (PDB: 4k95). The catalytic cysteine 431 of the RING2 domain (in red) is blocked by the UPD. The I44 patch of the Ubl domain interacts with the RING1 helix and the IBR interacts with Ubl, thereby covering the S65 phosphorylation site of the Ubl domain. Additionally, the C-terminal Ubl and REP block the E2 access to the RING1 domain. (b) Dynamic intermediate Parkin (PDB: 5caw; 143–461 a.a.) in complex with p-Ub. The interaction between p-Ub and Parkin straightens the kinked pUBH helix, opens the IBR and releases the Ubl and its phosphorylation site from the Parkin RBR core. (c) Model of the active form of Parkin. The RING domain of a Cbl-UBCH7 crystal structure (PDB: 1fbv; magenta) conjugated to ubiquitin (PDB: 4q5e; yellow) is superposed to PARKIN RING1 (PDB: 5caw). The Ubl and REP domains are released, making the RING1 domain accessible to the E2–Ub. The interaction of Parkin with the E2–Ub could induce conformational changes in that active site of the RING2 domain and fully activate Parkin.
The stoichiometry of non-phosphorylated vs. phosphorylated Ub on different substrates, or within Ub chains, might be critical in controlling Parkin activity and in the recruitment of specific effector proteins to p-Ub-decorated mitochondria. For example, whether Parkin can be activated by a single p-Ub on a substrate or a p-Ub moiety within a larger non-phosphorylated Ub chain is still unclear. Notably, unphosphorylated Ub does not induce Parkin activation [57], but has to be present in the reaction for chain building [55,59].

The stoichiometry of Ub phosphorylation will also depend on phosphatases that can reverse the process. The PINK1-mediated phosphorylation of Ub is reversible and declines with cessation of mitochondrial stress [82]. A mitochondrial phosphatase PGAM5 has been shown to interact with PINK1 [83,84] and could be a potential candidate to dephosphorylate p-S65 on Ub. It will be intriguing to identify phosphatases that specifically remove the phosphate group from free Ub or from Ub chains attached to substrates, thereby regulating not only Parkin activation but more broadly the biological responses mediated by binding of UBD-containing effector proteins.

Phosphorylation of Ub expands the repertoire of Ub-binding proteins

The unique conjugation machinery of ubiquitin enables the assembly of diverse and complex Ub modifications on substrate proteins. More than 20 different types of UBDs implicated in decoding the multiple functions of specific Ub modifications have been characterized to date [11]. The large majority of them bind with a relatively low affinity to a hydrophobic patch centred on isoleucine 44 of Ub. Phosphorylation of Ub changes its surface properties, inducing a modest conformational change adjacent to the hydrophobic patch [59]. Therefore, the recognition of phosphorylated monoUb and polyUb chains by Ub-binding proteins is likely altered upon phosphorylation. Recent work has indicated that the Ub-binding proteins NDP52 and OPTN are the primary autophagic receptors for PINK1/Parkin-mediated mitophagy [85]. The UBAN domain of OPTN and the UBZ domain of NDP52 are essential domains required for the removal of both cytosolic Salmonella [86,87] and mitochondria [85,88]. In the case of mitophagy, Youle and colleagues have shown that PINK1-mediated phosphorylation of Ub generates a specific signal required for recruiting OPTN and NDP52 to damaged mitochondria thereby activating mitophagy [85]. Interestingly, OPTN and NDP52 were not only shown to act as receptors for LC3-positive autophagosomal membranes, but also to recruit additional key components of the autophagy machinery including ULK1, DFCP1 and WIPI1 in order to promote in situ growth of the autophagosome and efficient mitophagy [85]. The kinase activity of PINK1 and the ability of OPTN and NDP52 to bind to Ub are essential for this process, whereas Parkin-mediated ubiquitylation amplifies it (Fig 3A). Yet, the basis for the specific interaction of OPTN and NDP52 with pS65-Ub is not understood. Both autophagy receptors preferentially bind to p-S65-Ub and to phosphomimetic S65D-Ub in vitro [85]. However, in vitro OPTN, NDP52 and p62 interacted less well with phosphorylated K63-Ub chains [60]. Similarly, none of the tested UBD-containing proteins that bind to the conserved hydrophobic patch around I44 on Ub showed increased affinities to phosphomimetic monoUbS65E [49] or phosphorylated Ub chains [60] in vitro. Thus, the cellular milieu favours the binding of autophagy receptors to phosphorylated Ub through a mechanism that cannot be recapitulated in vitro using purified proteins. Harper and colleagues showed that around 20% of mitochondrial Ub is phosphorylated [55]. Ub chains consisting of both phosphorylated and non-phosphorylated Ub molecules could control the recruitment of autophagic receptors and LC3 in vivo. In addition, phosphorylation of Ub at multiple sites may generate a new set of signalling cues that are recognized by autophagy receptor complexes (harbouring known or novel UBDs).

**TBK1-mediated phosphorylation controls ubiquitin-dependent autophagy**

In addition to PINK1, TBK1 was found to play a key role in engaging selective autophagy pathways for efficient removal of pathogens, protein aggregates and mitochondria [25,27,87,89]. OPTN, as well as TANK, Sintbad and Nap1 act as adaptor proteins that direct TBK1 to distinct complexes within cells [90]. For example, in the case of bacterial invaders, TBK1 is activated and recruited via OPTN and NDP52 to Ub-decorated cytosolic bacteria [86,87]. Analogously, TBK1 acts in concert with OPTN to mediate mitophagy of depolarized mitochondria [85]. Notably, direct binding of OPTN to Ub chains on cargoes has been proposed to promote TBK1 oligomerization and activation [91]. Locally accumulated TBK1 in turn phosphorylates the autophagy receptors OPTN and p62 on multiple sites, including the UBA domain of p62 and the UBAN domain of OPTN [25,27,87] (Richter and Dikic, unpublished observation) (Fig 4). This feedback phosphorylation increases the binding affinity of the autophagy receptors to Ub chains, thus strengthening their commitment to selected cargoes. This occurs, for example, on TBK1-mediated phosphorylation of S403 in the UBA domain of p62, which strongly enhances its binding to K63- and K48-linked polyUb chains [25] and has been implicated in the autophagic elimination of Mycobacterium tuberculosis [27] and polyubiquitylated mitochondria during PINK1/Parkin-mediated mitophagy [25]. Phosphorylation of S403-p62 by CK2 can also promote targeting of polyubiquitylated proteins into sequestosomes and their subsequent clearance by aggrephagy [26]. One interesting observation is that TBK1 may phosphorylate OPTN and p62 that are localized to different mitochondrial surface microdomains, with likely distinct functional consequences in the mitophagy pathway [85,88]. This is reminiscent of microdomain structures found on the Salmonella surface in the early phase of autophagy [87,92].

TBK1 can also phosphorylate S177 adjacent to the LIR motif of OPTN, resulting in an increase in LC3 binding that provides a feed-forward mechanism for the clearance of cytosolic Salmonella and restriction of intracellular bacterial proliferation [87]. Moreover, S332 in the LIR motif of p62 is also subjected to phosphorylation [26], indicating that, in analogy to OPTN, the LIR/LC3 interactions of p62 might be regulated through phosphorylation. However, further studies are required to assign a functional role for S332 phosphorylation of p62.
Thus, both the Ub/UBD interface that couples autophagy receptors to cargo, as well as the LIR/LC3 interface that links cargo to autophagic membranes, undergo phosphorylation (Fig 4). In this context, TBK1-mediated phosphorylation of autophagy receptors effectively prolongs their residence time on the surface of cargoes and also promotes directionality and the delivery of cargoes for autophagosomal degradation.

**Integrative model for phosphorylation-based regulation of mitophagy**

The modulation of critical binding interfaces by phosphorylation is emerging as a general regulatory concept in selective autophagy pathways as nicely demonstrated for the PINK1- and TBK1-dependent amplification loop in the control of mitophagy (Fig 5). PINK1-mediated phosphorylation of Ub and Parkin Ubl on their conserved serine 65 residues acts as an early event leading to the recruitment and activation of Parkin on mitochondria. Parkin ensures its own continuance on mitochondria through assembling Ub chains on MOM proteins. Simultaneously, autophagy receptors (such as OPTN and p62) are recruited to the sites of mitochondrial damage. Through TBK1-mediated phosphorylation on their UBDs and LIRs, autophagy receptors facilitate the removal of damaged mitochondria by bridging the p-ubiquitylated cargo to LC3-coated phagophores (Fig 5). Taken together, PINK1-mediated phosphorylation of Ub and Parkin and TBK1-mediated phosphorylation of autophagy receptors establish a robust mechanism that drives the selective labelling of damaged mitochondria to ensure their delivery for lysosomal degradation.

**Phosphorylation of Ub and UBDs in neurodegenerative disease pathogenesis**

Deregulation of the Ub system is often associated with severe phenotypes and diseases, including a variety of neurodegenerative disorders [93]. The potential physiological relevance of the PINK1- and TBK1-mediated phosphorylation loops is emphasized by genetic and biochemical studies showing that mutations in autophagy receptors that affect either their Ub-binding or their LC3-binding ability are linked to neurodegenerative disorders, in particular frontotemporal lobar degeneration (FTLD), amyotrophic lateral sclerosis (ALS) and Parkinson’s disease (PD) [88,94–96]. Most recently, genetic alterations impairing TBK1 activity or its binding to OPTN have been linked to FTLD and ALS [97–99]. Likewise, defects in the PINK1/Parkin pathway that disrupt signal amplification cause Parkinson’s disease [67,100,101]. Given the crucial role of p-S65-Ub in mitophagy, anti-p-S65-Ub antibodies may prove to be a useful tool for future studies of Parkinson’s disease. Indeed, p-S65-Ub can be detected under endogenous conditions in stressed neurons and accumulates in human brain during ageing and Parkinson’s disease, whereas it is largely absent in material obtained from Parkinson patients carrying PINK1 mutations [82]. Moreover, proteomics could potentially be developed as a diagnostic tool, as its quantitative nature would be crucial for
the discrimination between basal mitophagy and pathological conditions.

A caveat when studying the effects of Parkin mutations on the development of Parkinsonism in mouse models is that Parkin knockout mice do not display a neurodegenerative phenotype. In order to overcome this issue and study Parkin function in vivo, the Youle laboratory created a mouse model with accelerated accumulation of mitochondrial DNA mutations [102]. The loss of Parkin paired with the accumulation of dysfunctional mitochondria caused a loss of dopaminergic neurons, but neurons from other neuroanatomical regions appeared unaffected. The levels of p-S65-Ub measured by quantitative proteomics increase in the brains of these mice, indicating that in vivo mitochondrial damage leads to the accumulation of a specific p-Ub autophagic signal [102].

Conclusions

Initial work on the crosstalk of ubiquitylation with other PTMs revealed interesting bidirectional regulatory mechanisms. PTMs on the Ub moiety fine-tune signalling and increase the repertoire and selectivity of downstream interacting proteins. Recent work on PINK1/Parkin and TBK1/OPTN, showing interdependency of ubiquitylation and phosphorylation in selective mitophagy pathways, has provided new mechanistic insight and generated great excitement in the field (Figs 3–5).

There is likely a much broader spectrum of phosphorylation-dependent Ub functions in different pathways. Moreover, the identification of other mammalian kinases, as well as the yeast kinase(s) that can phosphorylate Ub on S65, S57 and other residues, is likely to reveal many of the pleiotropic effects of the expanded Ub code in cellular signalling. One interesting avenue for future research is to further explore the crosstalk between PINK1 and TBK1, which modify the Ub code and Ub receptors, respectively, thereby influencing their binding properties and their physiological roles. In addition, the role of other Ub ligases that act together or in parallel with Parkin should be further investigated. There is evidence that MUL1/MAPL/MULAN, another RING E3 ligase associated with mitochondria, can act as an alternative E3 ligase to Parkin to mediate mitophagy [103,104]. Furthermore, Parkin has been shown to maintain mitochondrial integrity by increasing linear ubiquitylation of NEMO, and this
autophagy-independent pathway involves Parkin-mediated activation of LUBAC [105].

Phosphomimetic Ub mutants—which are commonly used in research—provide a pragmatic experimental tool as the entire population of the studied protein is changed. Nevertheless, caution should be exerted when using these approaches regarding the physiological relevance of the conclusions drawn—even though they could be correct. Diverting results may partly be ascribed to the varying stoichiometry of phosphorylated substrates, such as Ub or Ub receptors. Moreover, the effect of serine to glutamic/aspartic acid mutations only partially mimics the altered surface or the conformational change induced by phosphorylation and could also account for a non-physiological consequence of the mutation. For example, MEK that is phosphorylated at S217/S221 by Raf has a 7,000-fold higher activity than the dephosphorylated enzyme, but the phosphomimetic mutations S217E/S221E of MEK only induce a 180-fold higher activity [106]. Similarly, due to the fast off-rates, a physiological binding partner of phosphorylated Ub is unlikely to be trapped in a complex with a Ub species that harbours a phosphomimetic residue (E/D). Likewise, phosphomimetic Ub or UbParkin proteins do not mimic their stoichiometrically phosphorylated forms, as phosphomimetics lack activity during Ub chain synthesis, whereas phosphorylated proteins are active [60].

The tRNA-based method to directly incorporate O-phosphoserine at desired positions and to stoichiometrically generate milligram quantities of serine-phosphorylated proteins in *E. coli* has already been used for S65-phosphorylated Ub (as well as other serine sites) [60,107,108]. Another recombinant approach to generate phosphorylated Ub molecules even included threonine- and tyrosine-phosphorylated forms of Ub [109]. These new techniques could lead to improved analysis, aiding the definition of genuine interactors of phosphorylated Ub.

Future studies will reveal whether there are even more PTMs that control the function of Ub and it will be important to identify the modifying enzymes that catalyse these PTMs. Moreover, using powerful quantitative proteomics will enable the identification, quantification and tracking of new PTMs on Ub with great sensitivity. Advanced proteomic tools to identify the flux and stoichiometry of phosphorylation and ubiquitylation are already available [110]; the technology would have to be further advanced for less studied PTMs.

Taken together, the modification of Ub and its receptors through phosphorylation, acetylation (Box 1) or deamidation (Box 2) can efficiently diversify the Ub code, thus adding another layer of complexity to Ub biology with important medical implications.

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**Conflict of interest**

The authors declare that they have no conflict of interest.

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