Protein tyrosine kinase regulation by ubiquitination: Critical roles of Cbl-family ubiquitin ligases

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Abstract

Protein tyrosine kinases (PTKs) coordinate a broad spectrum of cellular responses to extracellular stimuli and cell-cell interactions during development, tissue homeostasis, and responses to environmental challenges. Thus, an understanding of the regulatory mechanisms that ensure physiological PTK function and potential aberrations of these regulatory processes during diseases such as cancer are of broad interest in biology and medicine. Aside from the expected role of phospho-tyrosine phosphatases, recent studies have revealed a critical role of covalent modification of activated PTKs with ubiquitin as a critical mechanism of their negative regulation. Members of the Cbl protein family (Cbl, Cbl-b and Cbl-c in mammals) have emerged as dominant "activated PTK-selective" ubiquitin ligases. Structural, biochemical and cell biological studies have established that Cbl protein-dependent ubiquitination targets activated PTKs for degradation either by facilitating their endocytic sorting into lysosomes or by promoting their proteasomal degradation. This mechanism also targets PTK signaling intermediates that become associated with Cbl proteins in a PTK activation-dependent manner. Cellular and animal studies have established that the relatively broadly expressed mammalian Cbl family members Cbl and Cbl-b play key physiological roles, including their critical functions to prevent the transition of normal immune responses into autoimmune disease and as tumor suppressors; the latter function has received validation from human studies linking mutations in Cbl proteins to human leukemia. These newer insights together with embryonic lethality seen in mice with a combined deletion of Cbl and Cbl-b genes suggest an unappreciated role of the Cbl family proteins, and by implication the ubiquitin-dependent control of activated PTKs, in stem/progenitor cell maintenance. Future studies of existing and emerging animal models and their various cell lineages should help test the broader implications of the evolutionarily-conserved Cbl family protein-mediated, ubiquitin-dependent, negative regulation of activated PTKs in physiology and disease.

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1. Introduction

Phosphorylation of proteins on tyrosine residues is an evolutionarily conserved mechanism used by higher eukaryotes to alter the activity of cellular pathways in response to changes in extracellular environment. This fundamentally simple biochemical process, mediated by protein tyrosine kinases (PTKs), serves as an initial trigger in cellular activation processes whose downstream components include other major signal transduction enzyme cascades such as serine/threonine kinases, lipid kinases, phosphatases, phospholipases, small GTPases and other pathways. PTK-mediated signaling ensures orderly cell-environment and cell-cell interactions during development, tissue homeostasis, tissue repair and remodeling, as well as protection against pathogens. It is therefore not surprising that deregulation of PTK-initiated signal transduction is a major cause of human diseases such as cancer [1]. Given their critical importance in signal transduction cascades that regulate normal development and homeostasis in higher organisms, as well as the causal linkage of their aberrant activities to human diseases,
understanding the basis of PTK regulation is therefore of fundamental interest in biology and medicine.

2. Protein tyrosine kinases as key mediators of trans-membrane signaling

2.1. Both receptor and non-receptor protein tyrosine kinases mediate trans-membrane signaling

58 of the 90 PTKs in the human genome are trans-membrane receptors (referred to as receptor tyrosine kinases or RTKs) and are therefore directly poised for signal transduction in response to extracellular cues. Classical members of this family include growth factor receptors such as platelet-derived growth factor receptor (PDGFR) and epidermal growth factor receptor (EGFR). However, RTKs are now known to play crucial roles in essentially all developmental and homeostatic signaling pathways triggered by interactions with soluble or cell-surface associated ligands [1]. Nearly all of the remaining PTKs, the so-called non-receptor PTKs, are also involved in membrane receptor-initiated signaling through their constitutive or inducible association with cell surface receptors. For example, members of the Src-family of PTKs associate with antigen receptors on immune cells and members of the JAK-family of PTKs with cytokine receptors; in each case, non-covalently associated PTKs are essential for cellular activation upon receptor engagement, as demonstrated by naturally-occurring or engineered mutations of genes encoding these PTKs [2,3]. Thus, PTKs serve as essential membrane-proximal receptor signal transducers. Importantly, cell surface receptors with intrinsic or associated PTK activity commonly serve as biochemical or functional partners of other cell surface receptors such as G-protein coupled receptors (GPCRs), integrins and others, and are also activated by altered metabolic states of cells, such as oxidative stress. These collaborative interactions of PTKs further expand their critical roles to essentially every physiological response in higher eukaryotes [1].

2.2. Tyrosine phosphorylation-dependent signaling

Similar to serine/threonine phosphorylation, tyrosine phosphorylation can directly regulate enzymatic activity as exemplified by the phosphorylation of activation loops of PTKs. Importantly, phosphotyrosine (pY)-containing peptide sequences generated on PTKs themselves or on their interacting partners serve as docking sites that are recognized by Src-Homology 2 (SH2) and phosphotyrosine-binding (PTB) domains on signaling proteins. These interactions promote the recruitment of key regulatory and/or enzymatic components of signaling pathways such as Ras-Erk, PI3-kinase and PLCγ to membrane-associated activated receptors resulting in the activation of these downstream pathways [1]. Given the primary role of phosphorylation-dependent regulation of signaling protein complex formation and subsequent activation of enzymatic cascades, dephosphorylation by phosphotyrosine phosphatases (PTPs) is thought to provide one mechanism to negatively regulate PTK-dependent cellular activation. Notably, however, only in a limited number of cases have PTPs been categorically established as negative regulators of PTK signaling and paradoxically a number of PTPs mediate positive roles in PTK-dependent signaling cascades likely through dephosphorylation of sites that serve in negative regulation of PTK signaling [4]. Consistent with this dual role, recent cancer genome sequencing analyses have revealed that PTPs may function in both negatively (as putative tumor suppressors) and positively (as oncogenes) regulating oncogenesis [5].

2.3. Endocytic traffic of protein tyrosine kinase-associated receptors as a determinant of duration and diversification of signaling

Similar to ligand-binding cell surface receptors in general, PTK-associated receptors invariably undergo ligand-induced internalization through the endocytic pathway (Fig. 1). The rapid relocation of activated receptors from the cell surface into endocytic compartments has emerged as a fundamental mechanism to control signaling downstream of PTK-associated cell surface receptors. Studies over the last quarter century have elegantly demonstrated that a major consequence of the internalization of activated receptors is their trafficking to lysosomes where they undergo degradation [6,7]. This provides a mechanism of signal termination that appears to be essential for regulating signaling outputs from activated PTK-associated receptors under physiological conditions. Indeed, endocytosis-impaired receptors, such as EGFR with a deletion of its C-terminal sequences, become hyperactive and transforming [8]. Similarly, higher signaling potency of ErbB2, compared to its family members such as EGFR, has been attributed to its being internalization (and lysosomal degradation)-impaired [9,10].

Further studies have demonstrated that lysosomal degradation is not an invariant, or the only, consequence of the endocytosis of activated PTK-associated receptors. For example, while studies of EGFR using its prototypical ligand EGF have demonstrated lysosomal degradation as a major consequence, TGFα stimulation is associated with less lysosomal degradation of EGFR and a higher degree of its recycling back to the cell surface [11,12]. More recent analyses of all available EGF ligands have further established the distinct itineraries of internalized EGFR depending on the ligand [13]. In other studies, it has been established that concentrations of ligands can also dictate distinct itineraries of internalized receptors. For example, lower concentrations of EGF sufficient for induction of cell proliferation were associated with internalization of EGFR but not its degradation while higher concentrations of EGF promoted EGFR degradation [14]. These findings, together with studies of additional PTK-associated as well as non-PTK-associated receptors have contributed to the idea of endosomal signaling [15,16].

According to this newer concept, internalization of activated receptors by itself is not a signal attenuation mechanism. Instead, internalized receptors continue to signal within endosomal compartments until they are degraded in the lysosome or inactivated by other mechanisms and returned to the cell surface. The extent of receptor signaling within endosomes may indeed surpass that at the cell surface. In all cell models, the life of an activated RTK on the cell surface is very short (1–3 min) yet substantial degradation of internalized, activated receptor typically takes ten times longer: thus, activated RTKs spend nearly 90% of their lifespan within endosomes [6,7]. Studies in many cell systems have now shown that internalized RTKs continue to signal [15,17,18]. For example, signaling proteins such as Grb2 and Shc remain physically associated with activated EGFR and NGF receptors even after they traverse to endosomes [17–19]. Signaling from endosomes is indeed required for Trk receptor-mediated gene regulation in the neuronal cell body in response to stimulation at distant nerve endings [16].

Continued signaling ability of internalized PTK-associated receptors therefore provides a potential mechanism to regulate the duration of signal output depending on cellular needs. Thus, ligands (or concentrations of ligands) that promote lysosomal degradation can be anticipated to induce a relatively un-sustained signal output. In contrast, ligand conditions that do not promote lysosomal degradation can be anticipated to provide more sustained signaling outputs [6,7]. Studies of growth factor receptors in model cell systems have demonstrated that the transient vs. sustained activation of a single signaling axis can have dramatically different consequences for cell fate decisions. For instance, elegant studies using PC12 neuroblastoma cells have shown that transient Erk activation by EGF leads to a proliferative outcome while sustained Erk activation in response to NGF is associated with neuronal differentiation [20]. In a more recent example, maintenance of primary mammary epithelial organoid cultures in EGF versus amphiregulin determined the dominance of basal vs. luminal cell populations based on the extent of Erk activation [21].

In addition to regulating the duration and intensity of signaling outputs, evidence also supports the role of endosomal signaling by
PTK-associated receptors in signal diversification. For example, inhibition of EGFR endocytosis with a dynamin-K44A mutant reduced the level of phosphorylation of EGFR, the p85 subunit of PI3-kinase and Erk1, but increased the phosphorylation of PLCγ1 and Shc [22]. Recent studies indicate that RTK internalization is required for full MAPK activation, apparently due to the localization of a critical MEK-MAPK tethering protein within endosomes [23,24]. Additional studies have uncovered endocytic pathways that deliver active receptor signaling complexes to ER and Golgi for Ras activation [25]. Indeed, the requirement of endocytosis to activate signaling pathways extends to other receptors, such as G-protein-coupled receptors and TGFβ receptor [26–28]. Notably, endocytosis of EGF-EGFR complexes in the presence of a washable tyrosine kinase inhibitor (TKI), followed by inhibitor washout, demonstrated that initiation of signaling directly from endosomes was sufficient to promote cell survival and cell-cycle entry [29,30]. The new paradigm of endosomal signaling by receptors activated by their ligands at the cell surface therefore provides a unique mechanism for spatio-temporal regulation to achieve optimal duration and diversification of signaling outputs.

Given the vastly distinct functional consequences of endosomal sorting of internalized PTK-associated receptors into alternative lysosomal vs. recycling fates, the regulatory mechanisms involved have been of great interest. Early studies demonstrated that EGF-dependent lysosomal targeting of EGFR was a saturable process, suggesting the likelihood that certain key regulatory proteins were rate-limiting [31]. Recent studies have established that this postulated mechanism for lysosomal sorting of activated PTK-associated cell surface receptors is mediated by members of the Cbl family of ubiquitin ligases. The next sections therefore focuses on a brief overview of the ubiquitin pathway followed by a discussion of how the Cbl family of proteins controls lysosomal sorting of activated PTK-associated receptors to provide a fundamental mechanism of negative regulation of signaling. We will then discuss the physiological implications of this mechanism and pathological consequences of its aberrations.

3. Ubiquitin machinery and the control of cellular signaling by protein tyrosine kinase-coupled cell surface receptors

3.1. Components of the ubiquitin pathway

The pathway of post-translational modification of proteins with ubiquitin involves a highly conserved enzymatic cascade [32–34]. A ubiquitin-activating or E1 enzyme initiates the cascade in which a series of enzymatic reactions lead to an ATP-dependent thioester bond formation with the C-terminus of a ubiquitin molecule. This activated E1-ubiquitin conjugate binds with high affinity to a ubiquitin-conjugating enzyme (Ubc) and transfers active ubiquitin to E2, again linked via a thioester bond. The E2 enzymes stably associate with ubiquitin ligase enzymes (E3s) of which the two main classes are the catalytic variety of HECT-domain containing E3s and the non-catalytic category of RING finger domain (and related domains)-containing E3s. The E2 enzymes bound to HECT domain-containing E3s transfer the ubiquitin moiety to E3 (via a thioester bond), which in turn transfers ubiquitin to the ε-amino groups of lysine residues in target polypeptides through a stable iso-peptide linkage. The E3s of the RING finger domain category function as scaffolds, with the E3-bound E2 directly transferring ubiquitin to the target.

![Diagram of the ubiquitin pathway](image-url)
The E1–E2–E3 cascade helps establish the wide range of targets that can be subjected to ubiquitin modification relatively specifically [32–34]. Only two ubiquitin-directed functional E1s exist in mammals and it is likely that they function in distinct cellular processes, although their redundant roles remain to be fully explored. There are nearly three dozen E2s, a majority of them dedicated to the ubiquitin pathway. In contrast, several hundred ubiquitin pathway E3 proteins are encoded by the mammalian genome. The large number of E3s broadens the range of target proteins that are regulated by ubiquitin modification. E3s typically use distinct domains/motifs (and often several of them) to specifically interact with their substrates, with many of the interactions involving recognition of motifs generated by post-translational modifications (such as phosphorylation; this will be further illustrated with the interactions of Cbl proteins below). These specific interactions and their modulation under defined physiological conditions provide a mechanistic basis for ubiquitin modification of a limited group of substrates during specific cellular processes.

3.2. Types of ubiquitin modification and their consequences

In general, the major categories of ubiquitin modification of target proteins involve mono-ubiquitination and poly-ubiquitination with the latter involving chains linked through various lysine residues in ubiquitin (e.g., commonly observed K48- or K63-linked chains) or mixed chains [32–34]. While the precise regulatory mechanisms that result in a particular type of ubiquitin modification are not well understood, the type of ubiquitin modification dictates distinct functional consequences for the modified proteins. The common form of K48-linked poly-ubiquitin chains serves as a proteasomal targeting signal leading to degradation in the proteasome. In contrast, K63-linked poly-ubiquitin chains do not appear to function in proteasomal targeting. Both K63-linked chains as well as mono-ubiquitin modification, which is frequently seen in signal transduction, transcriptional and other regulatory pathways, dictate protein–protein interactions that do not lead to proteasomal degradation.

In the context of PTK-associated cell surface receptors, which is the main focus of this review, the receptors themselves or their associated adaptor proteins undergo K63-linked poly-ubiquitin and/or mono-ubiquitin modifications [35]. These modifications are recognized by a network of proteins with ubiquitin binding domains. These interactions provide a basis for recognition of ubiquitin-modified PTK-associated receptors by a series of multi-protein complexes referred to as the Endosomal Sorting Complex Required for Transport (ESCRT). The serial handover by ESCRT 0-III complexes is critical for sorting ubiquitin-modified receptors on outer limiting membrane of the multi-vesicular body (MVB) compartment of the endocytic pathway for delivery to inner vesicles of this organelle. This step is critical for further transport of receptors to the lysosome where they are degraded by lysosomal hydrolases (Fig. 1). While a greater structural and functional complexity is seen with the mammalian ESCRT machinery, the components and the architecture of ESCRT complexes is conserved from yeast to mammals. The ESCRT machinery has been thoroughly reviewed elsewhere [35–39], and will not be discussed in further detail here.

3.3. The role of the de-ubiquitination machinery

Over 100 proteases in the mammalian genome are members of several subfamilies of de-ubiquitinases (DUBs). These enzymes play multiple roles in regulating the ubiquitination process. Members of this group are essential for generating ubiquitin from its precursor covalent fusions with itself or other polypeptides as well as in regenerating ubiquitin from ubiquitin-modified proteins. Importantly, members of this enzyme family are involved in proteasomal degradation of ubiquitinated proteins as they play essential roles in processing and removing ubiquitin chains to facilitate the entry of ubiquitin-targeted polypeptides into proteasomal core, which harbors the catalytic proteases. Significantly, emerging evidence indicates key, and seemingly opposite, roles of DUBs in regulating endocytic traffic of PTK-associated receptors. Studies in yeast identified an essential role of the de-ubiquitinase Doa4/Ubp4 in vacuolar sorting of ubiquitinated receptors. Studies of the mammalian homolog of this DUB, Usp8/UBPY, in the context of RTKs have produced somewhat conflicting results. One set of studies indicates that Usp8/UBPY-mediated de-ubiquitination reduces the lysosomal degradation of RTKs such as EGFR [40]. Other studies have suggested an opposite role of facilitating RTK degradation [41]. More in depth analyses have begun to shed light on this complexity and apparently contradictory findings from different studies. It has now become clear that Usp8/UBPY also regulates the level of ubiquitination of other endocytic pathway regulators; such as the components of the receptor internalization and ESCRT machinery and alterations in Usp8/UBPY are associated with marked changes in the structure of endocytic compartments [42]. Similarly, endosomal deubiquitinase AMSH (Associated Molecule with the SH3-domain of STAM) has been shown to antagonize the ubiquitin-dependent lysosomal degradation of EGFR by trimming the K63-linked ubiquitin chains [43] while other studies have shown this DUB to be required for MVB-dependent EGFR degradation [43.44]. Furthermore, recent studies have shown that Usp8/UBPY and AMSH may regulate each other [45] as well as E3s such as the ErbB3-directed Nrdp1 [46]; therefore, their direct versus indirect effects require further elucidation. Notably, Usp8/UBPY is transcriptionally up-regulated by PTK-coupled cytokine receptor activation and its inducible upregulation was shown to alter colony stimulating factor 3 receptor traffic [47]. Thus while initial studies support a role of the Usp8/UBPY and AMSH deubiquitinases in providing a layer of regulation in the ubiquitin-dependent lysosomal sorting and degradation of activated PTK-associated receptors, the generality of this concept needs to be established in the context of additional RTKs and the cellular and/or molecular contexts that lead to distinct outcomes of their DUB function in RTK traffic need to be delineated [48].

Further pointing to this complexity, ubiquitin specific protease 2a (USP2a), a DUB that can remove both K48- and K63-linked poly-ubiquitination chains from substrates, localizes to the early endosome and exhibits oncogenic transforming activity, de-ubiquitinates and stabilizes internalized EGFR and promotes its recycling back to the cell surface [49]. Recently, the ovarian tumor protease DUB family member Cezanne-1 (also known as A20) was demonstrated to directly interact with EGFR, induce de-ubiquitination of the receptor and curtail its ligand-induced degradation; notably, Cezanne-1 was overexpressed in breast cancer patients and its mRNA overexpression predicted aggressive disease [50]. It is pertinent that Cezanne-1/A20 is an established regulator of NF-κB signaling and other pathways [51], more in depth analyses into its RTK regulatory function are warranted.

As the relative substrate selectivity and subcellular localization of DUBs during dynamic cellular processes remain relatively poorly characterized, further studies of their roles that reflect the regulation of the endocytic itinerary of PTK-associated cell surface receptors versus other functions are needed. The roles of DUBs demonstrated by these isolated initial examples also highlight the need for more unbiased studies of the larger DUB family. Such analyses should help uncover the potential roles of DUBs in the endocytic traffic of the wider group of PTK-associated receptors with attention to the tissue-specific and developmental patterns of expression of DUBs as these become more clearly defined, and may help elucidate novel mechanisms of PTK mediated oncogenesis that can be targeted for new therapeutic interventions.

3.4. Direct targeting of PTKs by the ubiquitin machinery

Following the well-established antagonistic relationship of phosphorylation and dephosphorylation as physiological switches, early studies of PTK regulation naturally focused on phosphorytrosine phosphatases (PTPs). As indicated above, PTPs provide one mechanism for negative regulation of PTK signaling although the vast majority of PTPs appear to have complex roles, including positive roles in PTK

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signaling. Early studies of the first PTK, Src, and subsequent studies of other cellular PTKs have also shown these cellular signaling switches to be potent proto-oncogenes whose oncogenic activity can be unleashed by overexpression, mutational or heterologous fusion-dependent activation, and other mechanisms [52]. Thus, it is not surprising that ubiquitination, an ancient pathway of regulating protein function, has evolved as an additional biochemical mechanism of PTK regulation.

Although ligand-induced ubiquitination of a RTK, PDGFR, was observed in early studies [53] the potential for PTK ubiquitination as a regulatory mechanism became clearly established in the early to mid-1990s through the convergence of a number of studies on the proto-oncogene Cbl, which is the focus of this review and is discussed in detail below.

The idea that direct interaction with Cbl or its family members (below), targets PTKs for ubiquitination and that this has regulatory consequences has led to considerable interest in other E3s with potentially similar roles under distinct biological scenarios [32–34]. Numerous examples of E3s that may facilitate ubiquitin-dependent regulation of PTKs are emerging and the list is likely to grow as more comprehensive studies of the ubiquitome are undertaken. A feature that sets the E3s of the Cbl family apart from all other E3s linked to PTK regulation is that they exclusively target PTKs and selected PTK signaling components in a “PTK activation-dependent” manner. The basis of this activated PTK-dependence involves mechanisms that mediate the binding interactions of Cbl-family E3s as well as the regulation of their activity by PTKs, as will be discussed below. Thus, Cbl-family E3s serve a unique role as activation-induced feedback negative regulators of PTKs. The discussion below on the biological consequences of genetic ablation of Cbl-family E3s together with emerging evidence that their mutations lead to human disease provide functional evidence that ubiquitination-dependent negative feedback provides a layer of regulation of activated PTKs that is distinct from the roles of PTPs and other mechanisms, and yet is essential for physiological control of PTKs.

4. Cbl-family ubiquitin ligases

4.1. General structure of Cbl-family E3s

Cbl (Casitas B-lineage Lymphoma) protein is the founding member of an evolutionarily-conserved family that includes three distinct gene products (Cbl or c-Cbl; Cbl-b; and Cbl-c, Cbl-3 or Cbl-SL) in mammals (Fig. 2). Cbl was initially isolated as the transforming protein...
oncogene (viral Cbl or v-Cbl) of a murine leukemia virus [54]. Subsequent isolation of the cellular homologue of mouse Cbl demonstrated that v-Cbl represents the truncated N-terminal part of Cbl fused to viral Gag sequences; the Gag sequences, however, proved dispensable for transformation of rodent fibroblasts by v-Cbl [55].

Mammalian Cbl proteins fall into two distinct categories (Fig. 2): the longer Cbl and Cbl-b gene products are highly related in their primary structure as well as their domain architecture. In contrast, Cbl-c represents a shorter version that lacks some of the key C-terminal domains and motifs. All three members of the Cbl family of proteins share a highly homologous N-terminal region that serves as the structural platform for direct binding to specific pY-containing peptide motifs in activated PTKs (Fig. 3) and is accordingly referred to as the Tyrosine Kinase-Binding (TKB) domain; this domain is in fact a unique assembly of a four-helical (4H) bundle, an EF hand domain and a variant SH2 domain, a tandem domain combination only found in Cbl-family proteins [56]. The TKB domain is followed by a highly conserved helical linker and a RING (Really Interesting New Gene) finger domain; these two regions together form the structural platform for binding to a ubiquitin conjugating enzyme (E2) [57] and this interaction is essential for the E3 activity of Cbl proteins. The three N-terminal domains (TKB, linker and RING finger) form the basic E3 module of Cbl proteins (Fig. 4) and these domains are necessary as well as sufficient for activated PTK-directed E3 activity of Cbl [58,59].

Cbl and Cbl-b proteins contain additional C-terminal motifs/domains known or presumed to mediate protein-protein interactions (Fig. 5) [60–65]. These include: a proline-rich region that mediates interactions of Cbl proteins with SH3 domain-containing proteins such as Src-family kinases, Grb2, CIN-85 and others; several tyrosine residues whose phosphorylation generates binding motifs for interaction with SH2 domain-containing signaling proteins that form key intermediates in PTK-dependent signaling networks, including Vav-family of guanine nucleotide exchange factors (GEFs) for Rho-family GTPases (bind to human Cbl Y700), p85 subunit of PI 3-kinase (binds to human Cbl Y731) and Crk-family of adaptor proteins that link Cbl proteins to C3G, a GEF for Ras-related small GTPase Rap1 (binds to human Cbl Y774). The sequences near the C-terminus of Cbl and Cbl-b encode a leucine zipper (LZ)-like motif/Ubiquitin-Associated (UBA) domain that is known to facilitate dimerization under certain experimental conditions [66]; this region also functions as a ubiquitin-associated (UBA) domain capable of binding to ubiquitin chains [67]. However, the functional roles of the LZ/UBA domain remain unclear although its ability to mediate ubiquitin-dependent dimerization has been suggested to provide a potential mechanism of activation of Cbl proteins [68].

In contrast to Cbl and Cbl-b, Cbl-c lacks most of the C-terminal regions except for a short proline-rich region for potential interactions with SH3 domain-containing proteins. Cbl-c also lacks a LZ/UBA domain as well as the region that harbors the tyrosine phosphorylation sites discussed above.

### 4.2. Evolutionary conservation of Cbl-family

Cbl proteins homologous to mammalian Cbl proteins have been identified in non-mammalian species including model genetic organisms

<table>
<thead>
<tr>
<th>Cbl-TKB domain binding target Protein</th>
<th>Motif sequence relative to consensus</th>
<th>Residues</th>
<th>Reference</th>
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<td>[82, 90]</td>
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<td>[84]</td>
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Fig. 3. Phosphotyrosine-containing sequence motifs that mediate Cbl TKB domain binding to its targets. The consensus motifs and individual target protein sequence motifs with critical amino acids that contribute to binding affinity are indicated [208–210]. Φ indicates hydrophobic residue. The motifs and sequences are adapted from [106].
such as C. elegans and Drosophila and even simple organisms such as Dictyostelium (Fig. 2) [60–65]. These Cbl orthologs exhibit a domain structure similar to that of mammalian Cbl-family proteins, with striking conservation of the domain architecture as well as the primary structure strongly suggesting a conservation of the biological functions as well as the biochemical mechanisms by which Cbl proteins exert their functional effects, as discussed below.

4.3. Cbl-family proteins as "activated PTK-selective" E3s—evolution of the concept

A number of key early observations contributed to the current paradigm that Cbl-family E3s function as selective regulators of activated PTKs in higher organisms. Identification of the v-Cbl oncogene [54,72] implied that the N-terminal region of the c-Cbl (which is incorporated in the oncogene) might interact with critical cellular proteins to dominantly deregulate their function and unleash an oncogenic program. However, the nature of cellular machinery targeted by Cbl remained unknown until multiple lines of evidence helped direct attention to PTKs. Several independent studies in the mid-1990s, investigating either Cbl or various PTKs, quickly converged to establish that Cbl is a direct binding partner and substrate of mammalian non-receptor and receptor type PTKs. Notably, Cbl was found to be tyrosine phosphorylated in cells oncogenically transformed with v-Ab1 indicating that Cbl is a substrate for a deregulated PTK [73]. This led to further analyses in which Cbl was found to be tyrosine phosphorylated in response to T-cell receptor engagement in cell lines, indicating that Cbl was a substrate of physiological PTK machinery [74]. Independent studies identified a novel 120 kD cellular protein (p120) that directly interacted with SH3 domains of T-cell receptor-associated Src-family kinases Fyn and Lck and was rapidly tyrosine phosphorylated upon T-cell receptor stimulation [75]. Subsequent purification of p120 identified it as Cbl [76]. Additional studies showed that stimulation through T-cell receptor as well as related B cell antigen receptor led to tyrosine phosphorylation of Cbl and its complex formation with downstream signaling proteins including Grb2, PI3-kinase p85 subunit, and Crk adaptor proteins [60,77–79]. Other studies quickly established tyrosine phosphorylation as well as signaling protein complexes of Cbl upon stimulation of EGFR in multiple cell systems [73–75,80–83]. In parallel, genetic screens in C. elegans identified the worm ortholog of Cbl (SLI-1) as a negative regulator of EGFR (LET23)-mediated vulva development.
and showed that the loss-of-function mutation in SLI-1 was a single amino acid missense mutation (G to E; equivalent to human Cbl G306E) within the evolutionarily-conserved region preserved in v-Cbl [84]. Collectively, these advances suggested a critical role of the conserved N-terminal region of Cbl in its function with the possibility that this region may be related to the interaction of Cbl with PTKs.

Direct binding assays using purified recombinant Cbl fragments incorporating the N-terminal region of Cbl, corresponding to v-Cbl sequences (Cbl amino acids 1–357; also referred to as Cbl-N), allowed a test of this hypothesis. GST-Cbl-N selectively and directly bound to phosphorylated ZAP70 tyrosine kinase in an activated T cell line and to related PTK Syk in an activated B lymphocyte line [85,86]. From screening of oriented phosphopeptide libraries and mutagenesis, it was established that Cbl-N bound to the negative regulatory pY-containing peptide motifs within the linker region of ZAP70 and Syk [87] (Fig. 3). Importantly, a point mutation (human Cbl-G306E) corresponding to the loss of function allele of C. elegans SLI-1 completely abrogated the ability of the TKB domain to interact with pY-PTK-peptide motifs on activated PTKs thereby linking the genetically identified loss of Cbl’s function as a negative regulator in C. elegans with PTK-binding activity in mammalian cells. Similar pY-containing motifs that allow direct binding on activated PTKs have been identified on other PTKs [88] (hence the designation Tyrosine Kinase-Binding or TKB domain) [60,85].

The idea of a TKB domain-mediated regulatory interaction of Cbl with activated PTKs was further supported by observations in fibroblasts which showed that the overexpression of oncogenic mutants of Cbl hyper-activated while the overexpression of wild-type Cbl reduced the level of PDGFR signaling in a manner that required an intact TKB domain [82,88]. Studies of fibroblasts overexpressing human EGFR showed that transforming Cbl mutant expression was associated with increased EGF phosphorylation [89]. Biochemical studies in fibroblasts with or without Cbl overexpression demonstrated that Cbl-dependent negative regulation of PDGFR receptor was a consequence of its Cbl-dependent ubiquitination which was associated with an enhancement of the ligand-induced down-regulation of the receptor from the cell surface [88,90]. Parallel studies demonstrated the facilitation by Cbl of EGF-induced down-regulation of EGFR and further demonstrated this to be due to enhanced lysosomal traffic of the internalized EGFR [59].

These findings led to the idea that Cbl-dependent ubiquitination of RTKs, such as PDGFR and EGFR, facilitates their internalization and/or lysosomal sorting akin to a model based on genetic and biochemical studies of yeast G-protein coupled pheromone receptors [91,92]. Indeed, these analyses were extended to hepatocyte growth factor receptor c-MET and mutation of Cbl binding site of c-MET led to its acquiring a highly oncogenic trait [93]. Furthermore, ubiquitination and downregulation of cell surface M-CSF receptor on bone marrow macrophages was reduced in Cbl-null mice [94]. Further studies with a number of non-receptor PTKs such as Fyn, Lck, Src, ZAP-70 and Syk that are associated with other cell surface receptors [58,96–99] helped establish the generality of Cbl-dependent ubiquitination of activated PTKs.

The ability of Cbl to induce PTK ubiquitination and the presence of a highly conserved RING finger domain (shown in other proteins to bind to E2) adjacent to the TKB domain prompted in vitro biochemical analyses that established that Cbl functions as an E3 towards an activated PDGFR receptor; the TKB domain was needed to contact the negatively charged tyrosine residues of the PDGFR-α [81,129]. While the requirement of the kinase activity of the target PTKs was found to be necessary for Cbl-dependent ubiquitination [115,116], the requirement of the kinase activity could reflect the need to generate a Cbl TKB domain-binding motif on RTKs or adaptor proteins, we showed that the Cbl TKB domain-mediated binding was dispensable yet the kinase activity was required for ubiquitination of the Src-family kinase Fyn [73,116]. Further studies revealed that target PTKs promote the phosphorylation of a conserved tyrosine residue in the helical linker of Cbl (human Cbl Y371) whose Y–F mutation leads to loss of E3 function [117].

Current crystal structural analyses of the TKB, helical and C terminus of un-phosphorylated versus Y371 phosphorylated states provided evidence that phosphorylation of Y371 reorients the helical region resulting in increased E2 binding and positioning of E2 closer to the phospho-peptide binding pocket of the TKB domain [105,113,114]. Thus, binding to target PTKs is thought to promote a linker phosphorylation-dependent activation of Cbl E3 activity. While the recent structural studies investigated this in the context of a TKB domain-bound PTK target, it remains unclear if a similar mechanism might account for requirement of the kinase activity for ubiquitination of targets recruited via domains others than the TKB domain (as with SFKs and SH2 domain-containing proteins; see below). It is also not clear if Y371 in Cbl is directly phosphorylated by the TKB domain-associated PTKs or by secondarily-activated PTKs such as Src-family kinases. In this regard, it is notable that Cbl-mediated ubiquitination and down-regulation of PDGFR-α was impaired if Src-association sites on PDGFR were mutated or in cells lacking SFKs [118].

Once bound to target PTKs, Cbl and Cbl-b proteins also undergo phosphorylation on several tyrosine residues located within the
sequences C-terminal to the RING finger domain and these generate binding sites for SH2 domain-containing signaling proteins [discussed above]. It is noteworthy that the signaling proteins that form complexes with Cbl proteins through SH2–phosphopeptide interactions (hence PTK activity-dependent associations) also undergo Cbl-family E3-dependent ubiquitination; this has been clearly demonstrated for PI3-kinase, whose p85 subunit interacts with pY731 of Cbl [119], for Vav and Vav2 [120,121] which interacts with Cbl pY700, and Crk/CrkL-associated C3G protein, a guanine nucleotide exchanger for Rap GTPases that interacts with Cbl pY774 [122]. Since structural studies have not included the C-terminal region of Cbl or Cbl-b, it is at present unknown what mechanisms might juxtapose signaling proteins bound to these sequences with the RING finger domain-bound E2. The C-terminal sequences and their phosphorylation (and interactions with SH2 domain-containing proteins) however do not appear to be required for ubiquitination of TKB domain-bound targets as suggested by the functionality of mammalian Cbl-c and that of C. elegans SLI-1 and the shorter isoform of Cbl in Drosophila as well as studies with truncated Cbl proteins [discussed above].

4.5. Nature and consequences of Cbl E3-dependent ubiquitin modification of activated PTKs

In general, Cbl E3s facilitate mono- or poly-ubiquitination of the associated targets [112,123–128]. Cbl-dependent mono-ubiquitination and K63-linked poly-ubiquitination appears to be a feature of activated RTKs [112,124–128] while other targets, such as ZAP70/Syk and Src-family kinases, appear to undergo poly-ubiquitination [50,57–99]. Notably, in most cases, these distinct modifications dictate degradation of ubiquitinated targets of Cbl proteins but through distinct pathways: mono- or K63-linked poly-ubiquitination of activated cell surface receptors facilitates their sorting to the lysosomes, based on recognition of the ubiquitin chains by the Endosomal Sorting Complex for Transport (ESCRT) machinery which facilitates the entry of ubiquitin-tagged receptors into inner vesicles of the multi-vesicular body (MVB) compartment (see above) (Fig. 1); in contrast, many poly-ubiquitinated targets of Cbl proteins appear to undergo proteasomal degradation [35–39]. However, in some instances, cytoplasmic signaling proteins associated with activated RTK complexes may indeed be sorted together with RTKs into the MVB and thereby undergo lysosomal degradation [126,129]. Interestingly, degradation-independent negative regulation by Cbl proteins has been suggested in selected cases. For example, negative regulation of PI3-kinase in T cells by Cbl-b was associated with ubiquitination but not degradation, leading to the authors’ suggestion that Cbl-family E3 dependent ubiquitination may mediate negative regulation of PI3-kinase and potentially other targets by regulating subcellular localization [119].

5. Functional roles of Cbl-family proteins

5.1. Functional roles of Cbl-family proteins based on evidence from in vitro model systems

A large body of literature now documents the functional consequences of the negative regulatory role of Cbl proteins using cellular systems. As these have been reviewed in detail over the past several years [63,77,78,130,131], this section will provide an overview of the message that has emerged out of these studies.

As PTKs generally serve as membrane proximal switches to convert extracellular signals into intracellular functional programs, the vast majority of in vitro studies have naturally focused on the role of Cbl proteins at the level of receptors themselves and at the level of proximal signaling events downstream of PTK-associated receptors. As stated above, the E3 activity of Cbl proteins plays a key role in the ubiquitination of RTKs as well as receptors (such as antigen receptors on immune cells) that are non-covalently linked to PTKs. There is general agreement that the overall consequence of Cbl-family protein-dependent ubiquitin modification of cell surface receptors is an enhancement of their lysosomal degradation via ESCRT protein-dependent facilitation of sorting at the MVB compartment [76,129,132]. Another proposed mechanism of the regulation of endocytic traffic of PTK-associated surface receptors by Cbl proteins, however, remains controversial. A series of investigations in the context of RTKs such as EGFR and c-MET, have indicated that Cbl proteins facilitate RTK internalization from the cell surface through a mechanism that may be independent of their E3 activity [133,134]. The model emanating from these studies suggests that Cbl proteins associate with activated RTKs via their TKB domains and recruit the Crb proline-rich region-associated SH3 domain-containing proteins of the CIN85 family. CIN85 forms a complex with endophilin A, which is known to be important in receptor internalization. Thus, Cbl-family proteins are proposed to function as adapters to promote the internalization of activated RTKs from the cell surface into early endocytic vesicles (Fig. 1). Studies using mutant proteins in transfection studies support this model [133]. Studies in other systems, however, have suggested that Cbl proteins are dispensable for initial internalization of activated RTKs and are essential only for the later step of lysosomal sorting. A rather physiological approach using mouse embryonic fibroblasts with complete deletion of Cbl showed that initial internalization of activated EGFR was not materially impaired [135] and our initial findings using Cbl plus Cbl-b double knockout cells further support these conclusions (unpublished observations). Consistent with these findings, EGFR was shown to become internalized in a Cbl-independent manner in response to oxidative stress resulting in EGFR recycling rather than degradation [136]. In addition, EGFR mutants defective in Cbl binding as well as ubiquitin modification are still internalized [137,138]. Thus, while it is possible that Cbl proteins can facilitate receptor internalization this function may be easily compensated for in the absence of Cbl protein recruitment. However, Cbl proteins are essential for sorting of the internalized PTK-associated receptors to the lysosome (Fig. 1). Further analyses are needed to fully elucidate the relative importance of Cbl proteins working through an adapter mechanism versus E3 activity, at two distinct steps in the endocytic traffic of PTK-associated cell surface receptors.

The second aspect of the functional roles of Cbl-family proteins that has been elucidated through cellular studies relates to their importance in regulating early biochemical pathways activated by PTK-associated receptor signaling. Thus, in a range of cellular models ranging from transfection analyses, knockdown analyses or cells derived from mice with deletion of Cbl or Cbl-b studies have shown that Cbl proteins negatively regulate the signaling outputs from activated PTKs into major downstream pathways such as the Ras-Erk, PI3-kinase-AKT and STAT pathways [139–143]. In general, loss of Cbl protein-dependent negative regulation kinetically prolongs the activation of these signaling pathways upon receptor stimulation. These effects are likely to be a result of the overall down-regulation of activated receptors through lysosomal degradation (which terminates their ability to continue to signal), targeting of associated PTKs (such as Src-family kinases, ZAP70/Syk) for proteasomal degradation, and the ubiquitination of signaling proteins themselves if these are able to interact with Cbl proteins directly. The latter mechanism appears to contribute to the negative regulation of PI3-kinase-AKT (via p85 subunit interaction with pY731 of Cbl), Erk (via interaction of Rap-GEF C3G with Cbl via Crk adapters; possibly also via interactions with sprotty proteins), and Rho GTPase family (via interaction with GEFs of the Vav family) [97,99,115,120,121,140,144–147]. Other signaling pathways are often impacted in specific cell types.

How the regulation of PTK-associated receptor traffic and immediate biochemical pathways translate into specific biological responses is however poorly understood. Perhaps the best links in this regard have been established in specific immune system cellular studies. It is clear, for example, that Cbl proteins regulate the actin cytoskeletal remodeling in T-cells that in turn is required for the stability of the immunological synapse; as a result, Cbl-b deficient T cells have longer lived
immunological synapses [148]. Similarly, Cbl-b deficient T cells are resistant to induction of anergy [149]. Loss of Cbl-b or Cbl plus Cbl-b in mice renders T-cells hypersensitive to T cell receptor activation for cell proliferation and cytokine secretion and removes the requirement for co-stimulatory signals normally delivered through CD28 or related proteins [150]. However, the linkage of these functional effects with specific biochemical pathways and regulation of T cell receptor endocytic trafficking remains poorly explored. How receptor proximal events regulated by Cbl proteins translate into distal functional consequences of PTK-associated receptor activation in other cell systems is even less clear. For example, even though RTKs such as EGF, PDGF and c-MET have been extensively analyzed with respect to receptor trafficking events and early biochemical responses regulated by Cbl proteins, how the physiological responses mediated by these receptors are regulated by Cbl proteins remains to be fully examined. One trait often altered in cells lacking Cbl proteins is cell migration response to growth factors such as EGF or HGF; for example, EGF or HGF-induced migration was increased in human mammary epithelial cells with Cbl plus Cbl-b knockdown yet their EGF-driven proliferation was unaffected [121]. In contrast, cytokine-induced proliferation of Cbl-null or Cbl plus Cbl-b-null hematopoietic stem and progenitor cells is markedly enhanced [151–153]. Little is known about how Cbl proteins might influence distal transcriptional events, cell proliferation and differentiation responses. These analyses are of fundamental importance to our understanding of the basis of physiological and pathological roles of Cbl proteins emerging from animal models and human diseases (see below).

While the focus of the current review is on the role(s) of Cbl-family E3s, it will be important for future studies to consider their roles in the context of other negative regulatory pathways that cells utilize to attenuate PTK-coupled receptor signaling. Apart from Cbl-family E3s other ubiquitin ligases have emerged as negative regulators of PTK-coupled receptor by impinging on pathways similar to those controlled by Cbl-family proteins: for example, Nedd proteins, ITCH, Nrdp1, CHIP, Cullin-5 and SIAH proteins have been shown to play a role in the ubiquitination and downregulation of PTK-coupled receptors [63,154–156]. Consistent with findings from earlier genetic studies in model organisms, recent analyses in mammalian systems have also revealed an important additional layer of negative feedback regulators that are induced upon receptor activation: examples of such inducible feedback inhibitors include LRIG1 (leucine rich repeats and immunoglobulin-like domain 1), suppressor of cytokine signaling 4 and 5 (SOCS4 and SOCS5) and receptor-associated late transducer and immunoglobulin-like domain 1), suppressor of cytokine signaling regulators that are induced upon receptor activation: examples of have also revealed an important additional layer of negative feedback regulation in model organisms, recent analyses in mammalian systems studies in model organisms, recent analyses in mammalian systems have shown that Cbl-Sprouty association sequesters Cbl from activated RTKs resulting in their inefficient ubiquitination and downregulation with a consequent increase in down-stream signaling. Thus, this complex interaction appears to utilize a known negative regulator of RTKs (Sprouty) to abrogate the effects of another negative regulator (Cbl) effectively converting Sprouty into a positive regulator of RTK signaling [145]. Further analyses indicate that Cbl-TKB-Sprouty 2 pY55 interaction is required for the negative regulatory function of Sprouty 2 towards Ras-ERK1/2 signaling [145,166]. In addition, this interaction has been shown to promote the ubiquitination and degra-dation of Sprouty 2 [169]. Finally, other studies have noted a biphasic regulation with positive as well as negative regulatory roles of Cbl-Sprouty complex that may influence RTK signaling in a spatio-temporal manner [170,171]. Notably, induction of Sprouty 1 upon activation of CD4+ T cells resulted in inhibition or enhancement of receptor signals depending on the differentiation state of cells, suggesting that Cbl and Sprouty proteins may function as a context-dependent regulatory complex for more precise and versatile regulation of receptor signals during development and tissue homeostasis [172]. Importantly, further studies, including in vivo analyses, of the relative importance of Cbl proteins versus other negative regulators and their cross-talk should provide fruitful avenues to elucidate the regulatory mechanisms that control biological processes mediated by PTK-associated receptor signaling.

5.2. Functional roles of Cbl-family proteins based on evidence from in vivo studies

5.2.1. Studies in model organisms

An early indication of a negative regulatory role of a Cbl-family protein for a PTK-associated receptor came from studies in C. elegans when a genetically identified negative regulator of the EGFR homolog LET23 was identified as the worm Cbl (SLI-1) [84] (Table 1). Drosophila Cbl homolog has similarly emerged as a negative regulator of EGFR signaling during photoreceptor development [173]. Drosophila Cbl is also required for dorso-ventral patterning and germ cell migration and functions by regulating RTK endocytic traffic [174]. More recently, a Cbl ortholog in zebrafish was shown to negatively regulate FGFR signaling during embryonic development [175]. The mechanism and consequences of the regulatory role of Cbl identified in these studies are quite interesting with potentially important implications for
**Table 1**

In vivo roles of Cbl family genes revealed by genetic mutation and deletion approaches.

<table>
<thead>
<tr>
<th>Organism (Gene)</th>
<th>Mutation /Knockout (KO)</th>
<th>Phenotype</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. elegans (Slt-1)</td>
<td>Loss of function mutation</td>
<td>No phenotype in wildtype worms; Restoration of vulva development defects in worms expressing a hypomorphic EGFR (let-23) allele</td>
<td>[84]</td>
</tr>
<tr>
<td>Drosophila (D-Cbl)</td>
<td>Transgene overexpression</td>
<td>No phenotype in wildtype flies; expression in a sensitized background (expressing a disabled Sevenless kinase, SveF) eliminated photoreceptor R7 differentiation.</td>
<td>[173]</td>
</tr>
<tr>
<td>Zebrafish (Cbl)</td>
<td>Loss of function mutation of D-Cbl</td>
<td>Defective dorso-ventral patterning during oogenesis, ovarian development and embryonic development. Disruption of the wing development.</td>
<td>[174]</td>
</tr>
<tr>
<td>Zebrafish (Cbl)</td>
<td>Ring finger mutant of D-Cbl (mutation of Cys-369, corresponds to Cys-381 in human Cbl); affects both D-Cbl and D-CblS</td>
<td>Abrogated border cell migration during oogenesis.</td>
<td>[211]</td>
</tr>
<tr>
<td>Mouse (Cbl)</td>
<td>Deletion mutation (identical to loss of function mutant of Schupbach group) and Ring finger mutation of D-Cbl (mutation at Arg-406); affects both D-Cbl and D-CblS</td>
<td>Abnormal eye development, increased omatidial spacing and larger head.</td>
<td>[212]</td>
</tr>
<tr>
<td>Mouse (Cbl)</td>
<td>T-&gt;F mutation in linker region Tyr-352 and 355 (corresponding to Tyr-368 and 381 of human Cbl)</td>
<td>Mutant fish show patterning defects during embryonic development due to alterations of fgf8 (fibroblast growth factor 8) morphogen gradient.</td>
<td>[175]</td>
</tr>
<tr>
<td>Mouse (Cbl)</td>
<td>Complete (germline) Knockout (Cbl&lt;sup&gt;m112abr&lt;/sup&gt;)</td>
<td>Hyperplasia of lymphoid organs, splenomegaly, increased mammary gland branching.</td>
<td>[176]</td>
</tr>
<tr>
<td>Mouse (Cbl)</td>
<td>Complete (germline) knockout (Cbl&lt;sup&gt;m111hor&lt;/sup&gt;)</td>
<td>CHI&lt;sup&gt;−/−&lt;/sup&gt; mice exhibited upregulated glucose metabolism, lean muscles and low fat deposition in the body despite hyperphagia, increased whole-body insulin action and energy expenditure.</td>
<td>[179]</td>
</tr>
<tr>
<td>Mouse (Cbl)</td>
<td>Complete (germline) Knockout (Cbl&lt;sup&gt;−/−&lt;/sup&gt;)</td>
<td>Mice are born healthy and fertile. Mild splenomegaly due to erythroid hyperplasia. Loss of Cbl in these mice altered thymic positive selection by enhancing CD4 lineage Tcells. Cbl knockout male mice are hypofertile. Absence of Cbl in the testicular gonadal cells results in failure to induce apoptosis during spermatogenesis.</td>
<td>[177]</td>
</tr>
<tr>
<td>Mouse Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;−/−&lt;/sup&gt;)</td>
<td>Cbl-deficient mice exhibit expansion of hematopoietic stem/progenitor (HSC) pool in the bone marrow. Born at expected Mendelian ratios and fertile. At the age of 6 month display huge mass of salivary gland due to accumulation of T cells and B-cells, and lymphoid neorganogenesis in most vital organs of the body. Generalized spontaneous autoimmune disorder around 6 months of age due to hyperactivation of autoreactive T and B-cells. Older mice developed progressive changes in facial features and enhanced osteoclastogenesis, CD40-induced B-cell proliferation is enhanced in mutant mice. Furthermore, Cbl-b&lt;sup&gt;−/−&lt;/sup&gt; mice display enhanced thymus-dependent antibody responses and germinal center formation.</td>
<td>[180], [213]</td>
</tr>
<tr>
<td>Mouse Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;−/−&lt;/sup&gt;)</td>
<td>Mutant mice are fertile and grossly normal levels of T-, B- and Natural Killer cells in peripheral and central lymphoid organs. Mice more susceptible to induced experimental autoimmune encephalomyelitis. Deficiency of Cbl-b leads to loss of requirement for CD28 co-stimulation for T-cell activation. Mice showed symptoms of Type-2 diabetes by manifesting glucose intolerance and insulin resistance around 20 weeks of age. Infiltration of activated macrophages seen in white adipose tissue. Cbl-b-deficient mice are resistant to unloading-induced muscle atrophy and loss of muscle function as a result of reduced IRS-1 (insulin receptor substrate-1) degradation. No gross or micro-anatomic phenotype.</td>
<td>[181], [214], [182]</td>
</tr>
<tr>
<td>Mouse Cbl-c</td>
<td>Knockout (Cbl&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
<td>Embryonic lethal by day E10.</td>
<td>[150]</td>
</tr>
<tr>
<td>Mouse Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
<td>Severe vascular lesions, lymphadenopathy, splenomegaly, extramedullary hematopoesis and symptoms of autoimmune diseases. Impaired T-cell receptor down modulation. Mutant mouse exhibited disease sign after 3 weeks. Females became moribund at 12 weeks, male mice by 16 weeks.</td>
<td>[150]</td>
</tr>
<tr>
<td>Mouse Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
<td>Mice develop spontaneous SLE-like autoimmune diseases. Impaired B-cell intrinsic tolerance induction, Bcell and induction of B-cell anergy. Half of the mutant mice became moribund by 10 months of age.</td>
<td>[185]</td>
</tr>
<tr>
<td>Mouse Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
<td>Mutant mice born at sub-Mendelian ratios. Mice develop severe myeloproliferative disorders (MPD) by 5 weeks of age with a median survival of 67 days.</td>
<td>[153]</td>
</tr>
<tr>
<td>Mouse Cbl-c and Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
<td>Embryonic lethal by day E10.</td>
<td>[150]</td>
</tr>
<tr>
<td>Mouse Cbl-c and Cbl-b</td>
<td>Knockout (Cblb&lt;sup&gt;c−/−&lt;/sup&gt;)</td>
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<td>[153]</td>
</tr>
</tbody>
</table>
other systems. FGF8 morphogen gradient is critical for shaping events during embryogenesis in Zebrafish. While loss of Cbl did not alter the morphogen gradient, Cbl-deficient tissues showed an expansion of the range of FGF8 target genes that were expressed apparently corre-
lating with reduced lysosomal targeting of active FGFR1 receptor sig-
ning complexes[175]. Thus, Cbl appears to regulate how a critical
developmental morphogen gradient is interpreted by target tissues
and this could have a bearing for the role of mammalian Cbl proteins
as well. It is notable that combined deletion of Cbl and Cbl-b in mice
results in early embryonic lethality through unknown mechanisms
[150]. The elegant genetic studies in model organisms provide strong
evidence that Cbl proteins play a critical role in physiological func-
tions mediated by RTKs and provide a rationale for more directed
studies in mammals.

5.2.2. Studies in mammalian systems

Complementing the evidence from biochemical and mammalian cell
biological analyses, mouse models (Table 1) have provided a number of
novel insights into physiological pathways where negative regulation
provided by Cbl proteins is critical. Germ-line deletion of Cbl proteins is critical. Germ-line deletion of
Germ-line deletion of Cbl proteins is critical. Germ-line deletion of

Table 1 (continued)

<table>
<thead>
<tr>
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</tr>
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<tbody>
<tr>
<td>Mouse Cbl-c</td>
<td>MMTV-Transgenic overexpression of Cbl-c in mammary epithelium.</td>
<td>Retardation in mammary ductal branching.</td>
<td>[215]</td>
</tr>
<tr>
<td>Mouse Cbl (G304E)</td>
<td>Knock in Cbl(^{G304E})</td>
<td>No gross abnormality in organs and tissues except for mild splenomegaly and dilated ureterine horns and fallopian tubes in mice of 6 to 9 weeks old.</td>
<td>[216]</td>
</tr>
<tr>
<td>Mouse Cbl-c (C379A)</td>
<td>Knock in Cbl(^{C379A})</td>
<td>Homozygous mutant mice died in utero or during first 24 h after birth. However, the Cbl-c(^{C379A}) mice exhibited additional defects, notably progressive loss of thymus.</td>
<td>[217]</td>
</tr>
<tr>
<td>Mouse Cbl-c (Y737)</td>
<td>Knock in Cbl(^{Y737})</td>
<td>Mice expressing mutant Cbl with inability to interact with the SH2 domain of phosphatidylinositol (PI) 3-Kinase p85 subunit exhibit increased bone mass and volume, and decreased osteoclast function.</td>
<td>[218]</td>
</tr>
<tr>
<td>Mouse Cbl-b (C373A)</td>
<td>Knock in Cbl(^{C373A})</td>
<td>RING finger domain mutant mice are healthy and produce homozygous offspring with expected frequency. Enhanced high-affinity IgE receptor (FcεRI) signaling in mutant mice is independent of E3 ligase activity.</td>
<td>[219]</td>
</tr>
</tbody>
</table>

The targets of Cbl proteins that mediate their embryonic developmental roles remain to be identified.

Given the apparently redundant but essential roles of Cbl and Cbl-b during embryonic development, it is of substantial interest to
dissect out to what extent these two family members (with their clos-
er biochemical similarity compared to Cbl-c) might function redund-
antly or in a specific manner. One mechanism for relatively specific
functions might be provided by their relative expression levels. At
the mRNA level, Cbl and Cbl-b appear to be widely expressed but
with one or the other family member predominating in specific tis-
ues, which appears to be concordant with the phenotypes of individ-
ual knockouts. For example, higher Cbl expression in the testes and
thymus are consistent with male infertility and T cell development
al phenotypes of Cbl null mice while hypersensitivity of peripheral T
cells in Cbl-b null mice coincides with higher Cbl-b expression
relative to Cbl [150,178,184]. However, the overlapping expression
patterns in most tissues suggest that either specific cell types within
these tissues rely on one family member versus the other or that
Cbl and Cbl-b function predominantly in a redundant manner in
many tissues.

Mice with a conditional (floxed) allele of Cbl on a Cbl-b-null back-
ground have allowed tissue-selective KO of Cbl plus Cbl-b to assess
the question of redundancy and to reveal the full functional impact
of negative regulation by Cbl proteins. Combined deletion of Cbl
plus Cbl-b in peripheral T lymphocytes (using Lck-Cre) indeed led
to a substantially more severe and spontaneous systemic autoim-
mune/inflammatory disease and dramatic hyper-responsiveness of T
cells to antigen stimulation [150]. Similarly, Cbl/Cbl-b double deletion
in the B lymphoid compartment of mice led to loss of peripheral toler-
ance [185]. Finally, deletion of Cbl and Cbl-b in hematopoietic stem
cells led to a myeloproliferative disease that was fatal within 8 weeks
of birth [153]. The latter phenotype is replicated by a RING finger do-
main missense mutant knock-in on a Cbl-null background [152].

Collectively, the KO animal models suggest that Cbl and Cbl-b proteins serve selective physiological roles likely due to their relative
levels of expression in various tissues. More importantly, these
models have emphasized that the two Cbl-family members Cbl and
Cbl-b function redundantly as essential regulators of both the embry-
onic development and adult tissue homeostasis in mammals. Given
the success of conditional induction of Cbl deficiency in a Cbl-b-null
background, further studies using this approach should help elucidate
the roles of these widely expressed Cbl family members in mammals.
In this regard, engineering of a Cbl-b floxed mouse model (ongoing in our laboratory) should provide a more facile and definitive tool to study functional impact of Cbl plus Cbl-b as redundant but critical negative regulators at various steps in development and in adult function. Notably, deletion of the predominantly epithelial tissue-expressed Cbl-c gene is without a phenotype [183]; however, combined deletion of Cbl-c with Cbl or Cbl-b has not been examined; these studies are currently being carried out in our laboratory and should be of considerable general interest.

5.2.3. Evidence from human diseases

Given the critical roles of PTK signaling in animal development, homeostasis and diseases such as cancer, together with emerging evidence of the role of Cbl family proteins in animal models, there have been strong efforts to link aberrations of Cbl family proteins to human disease. Recent results have indeed linked Cbl mutations to a set of human malignancies providing unequivocal evidence of the critical role of Cbl family protein-dependent negative regulation of PTKs and establishing Cbl proteins as tumor suppressors. Since 2007, a number of investigations have identified mutations clustered in the helical linker and RING finger domain of Cbl (and occasionally of Cbl-b) in about 5% of myeloid leukemia patients with myelo-dysplastic syndromes/myeloproliferative disorders: these include the juvenile myelo-monocytic leukemia (JMML), chronic myelo-monocytic leukemia (CMML) and atypical chronic myeloid leukemia (aCML) subtypes [186–194]. The cancer-associated Cbl mutations predominantly involve residues that have been shown by mutational analyses and/or structural studies to be essential for the ubiquitin ligase activity of Cbl, those mutants that were examined in vitro were found to lack the E3 ligase activity [186,195,196]. Furthermore, a cancer-associated mutant introduced into mouse hematopoietic cells produced mastocytosis [197] and as stated above a RING finger knock-in mutant of Cbl in mice phenocopied the myeloproliferative disorder seen in human patients as also did the deficiency of Cbl and Cbl-b [152,153]. Findings of a direct link between Cbl-family E3 mutations and human leukemia strongly support the critical role of Cbl protein-dependent negative regulation of PTK signaling. Identification of relevant target PTKs and a better understanding of the mechanisms of how mutant Cbl proteins promote oncogenesis represent exciting areas of ongoing and future research. Several recent reviews have summarized progress and emerging trends in this area [62,198–201]. It is somewhat intriguing that mutations of the linker and RING finger domains have thus far not surfaced in solid tumors. Whether this is a reflection of a lower frequency of mutations and need for larger studies or biological differences in the functional consequences of Cbl protein-dependent negative regulation in hematopoietic versus other tissues remains to be determined. Notably, recent studies of Cbl in lung cancer patients identified several mutations that are located in N- or C-terminal regions but not in the linker region and the RING finger domain [202]. As these mutations appear to moderately inhibit Cbl function in cell models, further studies with other solid tumors are warranted.

Given the growing evidence that human leukemias arise from hematopoietic stem/progenitor cells, linkage of Cbl mutations to leukemia also links these proteins to stem/progenitor cell homeostasis. Animal models with myeloproliferative disease as a result of the loss of Cbl protein function by gene deletion or knock-out of mutant Cbl [152,153] further support this role. In this regard, Cbl has also been shown to negatively regulate the PDGF-dependent survival of neural progenitor cells, and Cbl and Cbl-b were shown to regulate the asymmetric distribution of EGFR in neural stem cells as part of their self-renewal mechanisms [203,204]. Further studies should help clarify the role (general or limited) of Cbl proteins in stem cell maintenance and the potential target PTKs that they regulate in stem cell contexts. In this context, further analyses of the embryonic lethality of Cbl/Cbl-b double knockout mice should be quite revealing. Prominent autoimmune phenotypes in mice with Cbl-b (or Cbl plus Cbl-b) deletion have prompted analyses of polymorphisms/mutations in human patients with autoimmune disease. While several smaller studies have failed to detect any associations, some recent studies suggest an association between particular polymorphisms of Cbl-b with systemic lupus erythematosus [205]. Studies in the Komeda diabetes prone rat model identified Cbl-b as a susceptibility gene but the mechanism by which the truncation mutation in a region past the RING finger (and hence unlikely to affect the E3 activity) might contribute to disease remains unclear [206,207]. Regardless of the paucity of evidence in this area, future studies are warranted especially in view of the clear evidence from mouse models that aberrant Cbl protein function can contribute to inflammatory and autoimmune phenomena.

6. Conclusions and future directions

Structural, biochemical and cell biological studies of Cbl proteins have established that members of the Cbl protein family function as activated PTK-selective E3 ubiquitin ligases. Cbl family protein-dependent ubiquitination targets activated PTKs for degradation either by facilitating their entry into lysosomes or by promoting their proteasomal degradation. Signaling intermediates associated with PTK signaling and/or associated with Cbl proteins also appear to be subjected to ubiquitination and negative regulation although the role of degradation needs to be clarified. Cellular and animal studies have established that at least two members (Cbl and Cbl-b) of the Cbl protein family play key physiological roles: prominent examples of these roles include protection against autoimmune disease and tumor suppressor function, the latter receiving validation from human studies that have linked mutations in Cbl to human leukemia. Cbl mutations in human leukemia and insights from animal models together support an important role of Cbl and Cbl-b in stem/progenitor cell maintenance. Thus, future studies of existing and emerging animal models and cellular models derived from these, together with studies of additional human pathologies should help fully realize the range of physiological functions regulated by the evolutionarily-conserved Cbl protein family.

While functions of the relatively widely-expressed Cbl-family members Cbl and Cbl-b have begun to be elucidated, the roles of the structurally distant family member Cbl-c remain unclear. As Cbl-c is selectively expressed in epithelial tissues, future studies of this protein by incorporating its deficiency in animals that lack Cbl, Cbl-b or both proteins should help clarify whether this protein functions redundantly with its family members or by itself, perhaps under specific physiological conditions. Concurrently, a better linkage of the roles of Cbl proteins in the early biochemical and cell biological events associated with activated PTKs and the cellular and organismic roles deciphered from genetic studies should help underscore the importance of the Cbl protein family and the need for its evolution into a multi-gene family in mammals. In this regard, systematic examination of protein–protein interaction networks of all Cbl proteins and their impact on transcriptional pathways and biological outputs of PTK signaling under different physiological conditions of cells and tissues should add to our understanding of how Cbl proteins perform their diverse functional roles.

Studies directed at understanding how Cbl mutants associated with hematological disorders promote oncogenesis should provide further avenues to explore the physiological functions of Cbl proteins as well as provide potential avenues to develop agents that might prove useful in mutant Cbl-driven human cancer.

Abbreviations

TKB Tyrosine kinase binding
PTK Protein tyrosine kinase
EGFR Epidermal growth factor receptor
ErbB2 v-erb B2 erythroblastic leukemia viral oncogene 2 or EGFR-2
E1 Ubiquitin activating enzyme


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