OTULIN Antagonizes LUBAC Signaling by Specifically Hydrolyzing Met1-Linked Polyubiquitin

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SUMMARY

The linear ubiquitin (Ub) chain assembly complex (LUBAC) is an E3 ligase that specifically assembles Met1-linked (also known as linear) Ub chains that regulate nuclear factor κB (NF-κB) signaling. Deubiquitinases (DUBs) are key regulators of Ub signaling, but a dedicated DUB for Met1 linkages has not been identified. Here, we reveal a previously unannotated human DUB, OTULIN (also known as FAM105B), which is exquisitely specific for Met1 linkages. Crystal structures of the OTULIN catalytic domain in complex with diubiquitin reveal Met1-specific Ub-binding sites and a mechanism of substrate-assisted catalysis in which the proximal Ub activates the catalytic triad of the protease. Mutation of Ub Glu16 inhibits OTULIN activity by reducing $k_{cat}$ 240-fold. OTULIN overexpression or knockdown affects NF-κB responses to LUBAC, TNFα, and poly(I:C) and sensitizes cells to TNFα-induced cell death. We show that OTULIN binds LUBAC and that overexpression of OTULIN prevents TNFα-induced NEMO association with ubiquitinated RIPK1. Our data suggest that OTULIN regulates Met1-polyUb signaling.

INTRODUCTION

Ubiquitination is an important posttranslational modification that regulates diverse processes, including protein degradation, intracellular trafficking, transcription, kinase activation, and the DNA damage response (Hershko and Ciechanover, 1998; Komander and Rape, 2012). This variety of functions is mediated by eight different types of polyubiquitin (polyUb) linkages, and the roles of Lys48- and Lys63-linked polyUb have been studied in great detail, much less is known about the remaining “atypical” Ub chains (Behrends and Harper, 2011; Kulathu and Komander, 2012).

Met1-linked polyUb (Met1-polyUb) is the source of the cellular Ub pool, given that Ub is translated as a polyprotein (Ozkaynak et al., 1984) and posttranslationally processed by dedicated DUBs, such as USP5 (also known as IsoT) (Amerik AYu et al., 1997). This chain type can also be assembled by the linear Ub chain assembly complex (LUBAC), a multisubunit E3 ligase consisting of HOIP, HOIL-1L, and SHARPIN (Gerlach et al., 2011; Ikeda et al., 2011; Kirisako et al., 2006; Tokunaga et al., 2011). LUBAC has roles in NF-κB activation (Haas et al., 2009; Tokunaga and Iwai, 2012; Tokunaga et al., 2009; Walczak et al., 2012) and is required for full activation of the inhibitor of κB (IκB) kinase (IKK) complex. IKK activation leads to the phosphorylation and degradation of IκB and the activation of the NF-κB transcription factor (Karin and Ben-Neriah, 2000). It is not fully understood how Met1-polyUb regulates this process, but it involves the binding and modification of the IKK subunit NEMO with Met1-linked chains. NEMO harbors a Met1-specific Ub-binding domain (UBD) that is important for NF-κB signaling (Komander et al., 2009; Rahighi et al., 2009).

Much less is known about DUBs that regulate Met1-polyUb chains, and a specific DUB for Met1-linkages has not been identified. Of the roughly 80 active DUBs in the human genome, many show weak or no activity toward Met1-linked chains (Faesen et al., 2011; Komander et al., 2009). A potential reason is the distinct chemistry of a peptide versus an isopeptide linkage (Figure 1A).

Ovarian tumor (OTU) domain DUBs regulate important cell-signaling pathways. A20 regulates NF-κB signaling (Hymowitz and Wertz, 2010), OTUD5 (also known as DUBA) regulates IRF3 signaling (Kayagaki et al., 2007), and OTUB1 regulates the DNA damage response (Nakada et al., 2010). OTU DUBs can be linkage specific. Structural work has revealed the basis for OTUB1 Lys48 specificity (Juang et al., 2012; Wiener et al.,
Figure 1. Identification and Specificity of OTULIN
(A) Chemical difference between an isopeptide (left) and Met1-peptide linkage (right) in diUb. The distal Ub (top) is linked via its C-terminal Gly75-Gly76 sequence to a Lys side chain ε-amino group in another Ub or on a substrate, generating a branched peptide. In a Met1-linked chain, the C-terminal Gly76 is connected to Met1 of the distal Ub in a standard peptide linkage. The Met1 and Gln2 side chains, as well as the Met1 carbonyl (red), represent steric differences in comparison to an isopeptide linkage. A green arrow indicates the scissile bond in a DUB reaction.

(B) Sequence alignment of FAM105B/OTULIN with OTUB1. Sequence identity is 18% for the catalytic domain. Secondary structure elements are shown for OTUB1. The OTU domain is indicated in blue, and catalytic residues are labeled with yellow stars.

(C) The domain structure of OTULIN colored as in (B).

(D) Linkage specificity of OTULIN. diUb (1 mM) of all possible linkage types is hydrolyzed over a time course by 10 nM OTULIN and visualized on silver-stained 4%–12% gradient SDS-PAGE gels. See Figure S1 D for the assay at 1 mM OTULIN concentration.

(E) Cleavage of tetraUb chains, as in (D). The last substrate is a Met1-tetraUb with G76S mutation in all Ub moieties.

(F) Hydrolysis of Met1-diUb by OTULIN wild-type (WT) and catalytic mutants as indicated.

(G) The OTU domain of OTULIN encodes Met1-linked Ub specificity. diUb specificity analysis as in (D) with OTULIN 80–352 at a 10 nM concentration.

(H) Affinity measurements by fluorescence anisotropy with OTULIN (1–352) C129A or OTULIN (80–352) C129A and FlAsH-tagged Met1-diUb, as described in the Extended Experimental Procedures. Error bars represent SD from the mean of measurements performed in triplicate.
A
OTULIN
helical arm (S1)

B

C
Asp336
His339 (cat) 64%
Cys129 (cat) 28%

D
Ala336
Cys129 (cat) 51%
His339 (cat) 45%

E

F

G

H

I

J

(legend on next page)
RESULTS

FAM105B/OTULIN, a Met1-Linkage-Specific OTU DUB

Given the high sequence divergence of OTU domains, we set out to identify unstudied OTU enzymes using a bioinformatical screen based on generalized profile analysis (Bucher et al., 1996). Iterative profile refinement, starting from a multiple-sequence alignment of experimentally validated OTUs, indicated an OTU domain with a complete catalytic triad in the uncharacterized human protein FAM105B (Figure 1B). FAM105B comprises 352 amino acids (aa), and the OTU domain spans the majority of the protein (aa 80–352) and an N-terminal region with predicted helical content (Figure 1B, 1C). The catalytic domain is highly conserved between species (Figure S1A available online). Bacterially expressed full-length FAM105B did not hydrolyze common fluorescent substrates such as Ub-AMC (Figure S1B). Ub-based suicide inhibitors that comprise an electrophilic group at the Ub C terminus (Borodovsky et al., 2002) covalently modify most OTU domain DUBs but showed no reactivity against FAM105B (Figure S1C). However, DUB assays against diubiquitin (diUb) of all eight linkage types revealed that FAM105B exclusively hydrolyzed Met1-diUb (Figure 1D). The enzyme was active at 10 nM concentration (Figure 1D) and remained Met1 linkage specific at a 1 μM concentration (Figure S1D). Specificity is maintained when longer Ub chains are used as substrates (Figure 1E) but depended on an intact Ub Gly76-Met1 linkage sequence between Ub moieties, given that mutant tetraUb with Ser76-Met1 linkages was not hydrolyzed (Figures 1E and S1E). Catalytic mutants of FAM105B (C129A, H339A, and N341A) did not hydrolyze Met1-diUb (Figure 1F). Having established FAM105B as a Met1-linkage-specific OTU DUB, we named the enzyme OTULIN (OTU DUB with linear linkage specificity). OTULIN is unique, given that the 14 annotated human OTU DUBs cannot hydrolyze Met1-diUb (Mevisser et al., 2013).

Molecular Basis for OTULIN Specificity

Structural studies revealed how OTULIN achieved its unique specificity for Met1 linkages. The catalytic domain of OTULIN (OTULINcat, aa 80–352) is sufficient for linkage specificity (Figure 1G), and OTULINcat C129A bound Met1-diUb with a similarly high affinity as full-length OTULIN C129A, as revealed by fluorescence anisotropy measurements (K_D 196 versus 178 nM, Figure 1H).

OTULINcat was crystallized, and its structure was determined to 1.3 Å resolution with SeMet phasing (Figure 2A and Table S1). OTULINcat adopts an OTU fold most similar to OTUB1 (root-mean-square deviation [rmsd] 2.1 Å, DALI Z score 8.7) (Figure 2B). Interestingly, catalytic triad residues His339 and Cys129 display two alternate conformations. In the “active” conformation (occupancy ~30%), the catalytic triad is formed by interactions between Asn341, His339, and Cys129 (Figure 2C); e.g., as observed in OTUB1 in complex with Ub suicide inhibitor (Wiener et al., 2012) (Figures S2A–S2C). In the “inhibited” conformation (occupancy ~70%), Asp336 pulls His339 away from its catalytic position (Figure 2C), and Cys129 flips to an inactive rotamer. Next, we determined the structure of OTULIN D336A to 1.35 Å resolution (Figures 2D and S2A and Table S1). There were no global structural perturbations (Figures S2B and S2F), but His339 was now in the active rotamer, and Cys129 showed increased occupancy of the active rotamer (Figure 2D). Consistently, Ub suicide inhibitors that did not modify wild-type (WT) OTULIN modified OTULIN D336A and also OTULIN N341D, in which the catalytic Asn341 was

See Figure S1C and the Extended Experimental Procedures.
Figure 3. Substrate-Assisted Catalysis in OTULIN
(A) The OTULIN-diUb complex is shown with OTULIN under a blue surface and Met1-diUb colored as red and orange for the distal and proximal moieties, respectively. The catalytic center and Lys63-binding pocket is shown boxed.

(B) A close-up view of (A) showing all Ub Lys residues on the proximal Ub (yellow). Lys63 and Met1 are spatially close.

(C) Affinity measurements of OTULIN C129A with K48-, K63, and Met1-diUb linkages performed with fluorescence anisotropy (Ye et al., 2011). Error bars represent SD from the mean of measurements performed in triplicate.

(D–F) A close-up image of the OTULIN catalytic center showing key residues. Dotted lines indicate hydrogen bonds.

(D) Unliganded OTULIN, shown as in Figure 2 C.

(K) Summary table of kinetic parameters for OTULIN variants.

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changed to a negatively charged Asp (Figure 2E). Both mutants stabilize His339 in the active conformation, generating a more reactive enzyme.

To understand how OTULIN acted on Met1-polyUb specifically, we determined the structure of OTULINcat C129A bound to Met1-diUb to 1.9 Å resolution (Figures 2F and S2 and Table S1). The distal and proximal Ub moieties occupy extensive S1 and S1’ Ub-binding sites on OTULIN, respectively (Figures 2F–2H). Residues mediating Ub binding are highly conserved in OTULIN orthologs (Figure S2I). The binding interface with the distal Ub covers 1,045 Å² and involves the Ile44 patch that interacts with a helical arm (aa 254–264) conserved in all OTU domains (Figures 2F–2H). However, compared to the OTUB1–Ub structure (Wiener et al., 2012) (Figure S2J), the distal Ub rotates by ~18° in the S1 binding site (Figure S2K). The proximal Ub binds with an interface of 880 Å² to an S1’ Ub-binding site formed by helices α1 and α2 of OTULINcat via an unusual binding surface on Ub involving the Ub helix and the Phe4 patch (Figure 2H). Point mutations in the S1 (L259E and E314R) or S1’ (W96A and TQK100-102AAA)Ub-binding sites reduced OTULIN activity toward Met1-diUb (Figure 2I) by decreasing Met1-diUb affinity (Figure 2J).

OTULIN Specificity: Selective diUb Binding

The extensive S1’ Ub-binding site is likely to be important for OTULIN specificity in that it orients the proximal Ub such that only Met1 points toward the catalytic center (Figure 3A). In this orientation of the proximal Ub, all Lys residues are remote from the catalytic center, except for Lys63, which is spatially close to Met1 (Figure 3B). Importantly, OTULIN wedges these linkage points apart by two loops (aa 125–127 and 282–284) that fix Lys63 in a dedicated binding pocket (Figure 3B). Nonetheless, a differently linked Ub chain including a Lys63-linked diUb would rotate the proximal Ub moiety by several degrees, and such orientation would likely not fit the OTULIN S1’ binding site. Indeed, Lys63-linked diUb bound with 100-fold reduced affinity (<112 nM for Met1-linked diUb versus 12 μM for Lys63-linked diUb), and no binding was detected for Lys48-linked diUb (Figure 3C). This shows that the Ub-binding sites of OTULIN already distinguish between structurally similar Met1 and Lys63 chain types by two orders of magnitude.

OTULIN Specificity: Substrate-Assisted Catalysis

The complex structure revealed that the proximal Ub directly participates in the organization of the catalytic center. Autoinhibition of the catalytic triad in the absence of substrate (Figure 3D) is released by the binding of the Met1-linked proximal Ub (Figures 3E and 3F). The carbonyl group of Met1 sterically interferes with the inhibited conformation of His339, pushing it into an active conformation (Figure 3F). Lys-linked Ub chains lack a structural equivalent of this carbonyl moiety in the linkage (Figure 1A). More significantly, Glu16 of the proximal Ub is inserted into the catalytic center, displacing the inhibitory OTULIN residue Asp336, further restricting the mobility of His339 (Figures 3E and 3F). In addition, Glu16 coordinates the third residue in the catalytic triad Asn341, aligning it toward His339 (Figure 3E).

Importantly, Met1-diUb with mutations of Glu16 in the proximal Ub was hydrolyzed with significantly lower activity in qualitative gel-based assays, whereas mutation of nearby Gin2 (which also interacts with OTULIN, Figure 3E) had no effect (Figures 3G and S3A). All mutants were hydrolyzed by the nonspecific DUB USP21 (Ye et al., 2011) (Figure S3B) and bound to OTULINcat C129A in analytical gel filtration (Figure S3C), and Met1-diUb E16Aprox affinity toward OTULIN C129A was only slightly decreased (612 versus <112 nM, Figure S3D).

We used a quantitative fluorescent kinetic assay for diUb cleavage (Virdee et al., 2010) to examine whether Glu16 on a proximal Ub had a direct role in catalysis (Figures 3H and 3I). Met1-diUb E16Aprox decreased kcat 240-fold and enzyme efficiency (kcat/KM) 74-fold (because of a 3.5-fold lower KM for the mutant) in comparison to Met1-diUb (Figures 3H and 3I). Interestingly, the negative charge on Ub is important for OTULIN activity, given that Met1-diUb E16Qprox still showed a 35-fold lower kcat for WT OTULIN. Next, we tested whether the more reactive OTULIN N341D or D336A mutants (Figure 2E) showed improved diUb hydrolysis activity. WT Met1-diUb is a poor substrate for OTULIN N341D, most likely because of the repulsion of negative charges (Ub Glu16 and OTULIN Asp341) in the catalytic center (Figures 3I and K). Indeed, OTULIN N341D showed improved kinetics when Ub Glu16 was mutated, and both kcat and KM improved, although WT activity was not regained. The OTULIN N341D mutant worked best with Met1-diUb E16Qprox, suggesting that the requirement for a negative charge was partially compensated (Figures 3J and 3K). The coordination of Asn341 is a key event in OTULIN activation, as was confirmed when the only OTULIN residue coordinating the Asn341 side chain, Tyr91, was mutated to Phe, resulting in 20-fold reduction of kcat while not affecting KM (Figures 2I, 2J, 3K, and S3F). Strong effects of Glu16 mutation were also observed in the OTULIN D336A mutant (Figures 3J and 3K). Altogether, this showed that the coordination of the catalytic triad through Ub interaction is important for OTULIN activation (Figure 3L).

Hence, we reveal a mechanism of substrate-assisted catalysis in which Glu16 of the proximal Ub activates the catalytic triad of OTULIN by both restricting the movement of the catalytic His339 and introducing a negative charge, presumably for the proper
coordinated of Asn341 for catalysis, OTULIN’s usage of a catalytic Asn improves interaction with Glu16 containing Met1-linked chains.

It appears that OTULIN has evolved Met1-linkage-specific Ub-binding sites to specifically interact with linear chains. Additionally, to further distinguish chain types, OTULIN invokes a mechanism of substrate-assisted catalysis in order to ensure that only Met1 linkages are hydrolyzed. OTULIN’s remarkable specificity suggests that Met1-linked polyUb have to be tightly regulated independently of other ubiquitination events in cells.

**Cellular OTULIN Antagonizes LUBAC Signaling**

The identification of OTULIN as a Met1-linkage-specific DUB prompted us to study its role in cells. The human FAM105B gene is ubiquitously expressed (http://biogps.org/#/3d3Cgo73nereport%26id%3D90268). A polyclonal antiserum detected OTULIN in human embryonic kidney (HEK) 293ET and other cell lines (Figure S4A–S4C). C-terminal GFP-tagged OTULIN is cytoplasmic, active, and Met1 specific (Figure S4D–S4F). OTULIN is evolutionarily restricted to vertebrates and selected invertebrate lineages but is not detected in D. melanogaster and C. elegans. Interestingly, all OTULIN-comprising taxa also contain genes for components of the Met1-polyUb chain assembly machinery LUBAC.

Expression of the LUBAC components HOIP, HOIL-1L, and SHARPIN induced Met1-polyUb, which was removed when OTULIN was coexpressed (Figure 4A, lanes 2 and 3). Inactive OTULIN C129A led to the significant enrichment of Met1-polyUb in cells, and OTULIN Ub chain-binding mutants W96A and L259E (Figures 2I and 2J) increased Met1 linkages in cells, albeit not in D. melanogaster and C. elegans. Interestingly, all OTULIN-comprising taxa also contain genes for components of the Met1-polyUb chain assembly machinery LUBAC.

Ubiquitin Glu16 Is Important for Met1-polyUb Signaling

Next, with a cellular readout for OTULIN overexpression at hand, we set out to test OTULIN’s mechanism of substrate-assisted catalysis in cells by characterizing the effects of an Ub E16A mutant. Control experiments assessing whether this mutant is still assembled in Met1-polyUb by HOIP gave the surprising result that a minimal HOIP ligase construct (Smit et al., 2012; Stiegitz et al., 2012) was impaired in assembling Met1-linked chains from Ub E16A (Figure 5A). Furthermore, fluorescent Met1-diUb E16A protein bound to the NEMO UBAN domain with reduced affinity (Figure 5B), which is consistent with the known interaction between the NEMO UBAN domain and Ub Glu16 (Figure 5C). Despite this, co-overexpression of Ub E16A and LUBAC in HEK 293ET cells still activated NF-κB, and this could not be completely reversed by OTULIN (Figure 5D), consistent with OTULIN’s inability to efficiently hydrolyze Ub E16A-containing polymers (Figure 3). These results suggested that Ub Glu16 is important in multiple aspects of Met1-polyUb signaling.

OTULIN Affects LUBAC-Mediated Cytokine Responses

Transient overexpression of OTULIN or the well-studied NF-κB inhibitor A20 (Hymowitz and Wertz, 2010) blocks poly(I:C) induced NF-κB activity (Figure 6A). In comparison, OTULIN inhibits TNFα-mediated NF-κB by ~50%, which correlates with effects observed by LUBAC downregulation (Haas et al., 2009; Tokunaga et al., 2009) (Figure 6A). In OTULIN-expressing stable cell lines, transcription of NF-κB target genes was reduced in response to 10 ng/ml TNFα (Figure 6B), and, although IκBα was rapidly phosphorylated and almost completely degraded after 15 min of TNFα stimulation in control cell lines, IκBα phosphorylation, degradation, and NF-κB activation was delayed in
OTULIN-overexpressing cells (Figure 6C). Moreover, stable overexpression of OTULIN sensitized cells to TNFα-induced cell death (Figures 6D and 6E), consistent with observations in cpdm mice that lack Sharpin (Gerlach et al., 2011; Ikeda et al., 2011; Tokunaga et al., 2011; Walczak et al., 2012) or in humans with mutations in HOIL1 (Boisson et al., 2012). OTULIN overexpression promoted enhanced and persistent JNK activation and c-Jun phosphorylation at later time points, leading to the cleavage of PARP and caspase-3, two key players in the regulation of cell-death induction (Figure 6F). As for IkxBα phosphorylation, OTULIN overexpression also resulted in a delay of JNK activation kinetics (Figures 6C and 6E).

Small interfering RNA (siRNA) against OTULIN resulted in an increase in LUBAC-induced NF-kB activation in HEK 293ET and U2OS cells and also in T-REx 293 cells stably expressing a doxycycline-inducible OTULIN-targeting microRNA (miRNA) (Figures 7A and S6A–S6C). LUBAC-dependent induction of NF-kB could be prevented by co-overexpression of...
active, but not inactive, OTULIN (Figure 7A). Interestingly, western blotting against the overexpressed LUBAC components revealed that HOIP was ubiquitinated in OTULIN miRNA cell lines (Figure 7B), and coexpression of OTULIN removed these chains, showing that they are Met1 linked (Figure 7B). This suggested that one role of OTULIN in the LUBAC complex (Figure 4E) is to prevent the autoubiquitination of HOIP. However, LUBAC recruitment to the TNF-RSC was unchanged in OTULIN knockdown cell lines, and Met1-ubiquitination were enriched in the TNF-RSC, implying that OTULIN also regulates Met1-Ub on other components (Figure 7C). Accordingly, we observed a slight increase in RIPK1 ubiquitination enriched with a Met1-linkage-specific Ub-binding matrix in OTULIN-depleted U2OS cells (Figure 7D), suggesting that RIPK1 Met1 ubiquitination in response to TNFα is targeted by OTULIN (Figure 4F).
Figure 6. OTULIN Overexpression Inhibits TNFα Signaling
(A) Luciferase assays performed as in Figure 4B in HEK 293ET cells transfected with OTULIN, A20, or both OTULIN and A20 and stimulated with TNFα (100 ng/ml) or poly(I:C) (10 μg/ml). Western blots below show transfected protein levels. Error bars represent SD from the mean for experiments performed in triplicate. p values are given to indicate significance. *, the control set to 1.

(B–F) Stable doxycycline-inducible control and OTULIN-overexpressing T-REx 293 cells after treatment with 1 μg/ml doxycyclin (Dox) for 24 hr (see the Extended Experimental Procedures).

(B) Quantitative PCR (qPCR) analysis of selected NF-κB target genes in control (gray) and OTULIN-overexpressing cells (black) after treatment with 10 ng/ml TNFα for the indicated times.

(C) Analysis of TNFα-stimulated NF-κB activation and IκBα phosphorylation upon TNFα treatment (100 ng/ml) over the indicated time course by western blotting with indicated antibodies and EMSA. See (F) for tubulin control.

(D) Clonogenic survival of indicated cell lines ± Dox and ± TNFα (50 ng/ml) for 24 hr (see the Extended Experimental Procedures).

(E) Cell viability counts of cells treated as in (D). Error bars represent SD from the mean for experiments performed in triplicate.

(F) Analysis of TNFα-stimulated signaling cascades upon TNFα treatment (100 ng/ml) over the indicated time course by western blotting.
Figure 7. OTULIN Depletion Amplifies LUBAC Signaling

(A–C, E–F) Experiments in stable T-REx 293 cell lines inducibly expressing control or OTULIN-targeting miRNA (see Experimental Procedures and Figure S6).

(A) NF-κB luciferase activity shown as in Figure 4B in cells transfected with LUBAC and indicated OTULIN variants. Error bars represent SD from the mean of experiments performed in duplicate. *p values are given to indicate significance. *, the control set to 1; n.s., nonsignificant.

(B) Western blotting analysis of lysates from (A) with indicated antibodies. *, nonspecific band.

(C) Immunoprecipitation of the TNF-RSC by FLAG-TNFα (100 ng/ml) from control and OTULIN-depleted cell lines at indicated time points western blotted against Met1-polyUb, HOIP, and TNFR1.

(D) Met1-Ub-specific GST-M1-SUB or general GST-TUBE1 Ub pulldown from U2OS cells stimulated with TNFα (10 ng/ml) after transfection of OTULIN siRNA or a nontargeting (NT) control siRNA.

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Stable OTULIN knockdown affected cytokine responses and led to enhanced NF-κB activation after poly(I:C) and TNFα (Figure 7E), and, even though the initial induction of NF-κB target genes was not severely altered after OTULIN knockdown, expression of A20, IκBα, IL-8, and TNFα was sustained in response to TNFα (Figure 7F). Initial phosphorylation and degradation of IκBα after TNFα stimulation was only slightly enhanced in OTULIN knockdown cells (Figure S6D). However, JNK and c-Jun phosphorylation was sustained at late time points, again leading to cell death (Figures S6D–S6F). The fact that both overexpression and knockdown of OTULIN led to cell death was surprising and requires further investigation.

OTULIN knockdown cell lines were deregulated in their NF-κB response, but we could not observe a strong effect on canonical NF-κB signaling. Jurkat cells express higher amounts of OTULIN (Figure S4C), and OTULIN knockdown by siRNA led to an increase in phospho-IκBα and subsequent enhancement of NF-κB DNA binding (Figure 7G). This was due to increased IKK activation (Figure 7H), consistent with current models of LUBAC function (Tokunaga and Iwai, 2012; Walczak et al., 2012). Altogether, our data are consistent with a role of OTULIN in LUBAC-mediated Met1-polyUb signaling.

**DISCUSSION**

Here, we identify FAM105B/OTULIN as a human OTU DUB specific for Met1-linked Ub chains. OTULIN is a missing piece in Met1-Ub chain biology, for which no linkage-specific DUB has been described to date. Our data suggest that OTULIN is an antagonist of LUBAC in vitro and in cells. We reveal the molecular basis for the remarkable Met1 linkage specificity by structural and biophysical studies and give insights into potential roles of OTULIN as a LUBAC interactor and antagonist in cells. Consistent with the biochemical findings, overexpression of OTULIN removes Met1-polyUb and dampens LUBAC-mediated NF-κB signaling, whereas knockdown of OTULIN leads to increased Met1-polyUb, IKK activation, and NF-κB signaling after TNFα stimulation. However, both overexpression and knockdown of OTULIN sensitzed cells to TNFα-induced cell death, suggesting more complex roles for Met1 linkages in balancing signaling pathways downstream of the TNFα receptor.

**Structural Insights into Ub Chain Linkage Specificity in DUBs**

Linkage specificity in DUBs is not well understood, and the structure of apo OTULIN did not explain its Met1 specificity, because of its structural similarity to the Lys48-specific OTUB1. The key insight into linkage specificity for these two enzymes came from the complex structures of the enzyme with two Ub molecules bound across the active site. A high-affinity Ub-binding site in OTULIN allows it to select Met1-linked chains with 100-fold preference over Lys63 linkages and disallows the binding of Lys48-linked and presumably other chains. This mechanism is shared by many DUBs; e.g., TRABID (Licchesi et al., 2012) and OTUB1 (Juang et al., 2012; Wiener et al., 2012). However, in a physiological situation where multiple Ub chain linkages are often mixed, substrate-targeted DUBs may cleave additional Ub chain types. OTULIN prevents such promiscuity by requiring a properly positioned Ub residue, Glu16, for catalytic efficiency. Even in Lys63-linked chains, the required rotation of the proximal Ub would displace Ub Glu16, preventing activity. Therefore, OTULIN is the first DUB for which substrate-assisted catalysis has been demonstrated to achieve linkage specificity. It is possible that substrate-assisted catalysis explains the linkage (or substrate) specificity of other DUBs. It will also be interesting to see whether DUB activators exist that complement the catalytic triad in an analogous manner as that observed in OTULIN.

Ub-mediated substrate-assisted catalysis was recently shown to be crucial for the assembly of Lys11-linked polyUb, where Ub Glu34 complements the active site of the E2 Ub-conjugating enzyme UBE2S (Wickiffe et al., 2011). The fact that Glu16 also impairs HOIP activity also suggests the functionality of this side chain in assembly reactions. Altogether, this shows that Ub is more than just a binding partner for other proteins and that it can actively participate in enzymatic reactions.

**Cellular Role of OTULIN in Counteracting LUBAC**

In transfection experiments, we demonstrated that OTULIN could counteract cellular LUBAC responses, suggesting that OTULIN helps to balance the amount of Met1-polyUb in cells. Furthermore, the mechanism of substrate-assisted catalysis suggests that OTULIN may not target individual proteins but, instead, target Met1-linked Ub chains, regardless of where they are attached. Indeed, we identified three proteins that change in their ubiquitination status when OTULIN levels are modulated. NEMO and RIPK1 are among the few reported targets of LUBAC (Gerlach et al., 2011; Tokunaga et al., 2009).

We provide evidence that OTULIN directly interacts with LUBAC. The observation that depletion of OTULIN leads to the modification of HOIP with Met1-polyUb suggests that HOIP, like many other E3 ligases, undergoes autoubiquitination and that OTULIN has the ability to prevent this. This most likely explains why the overexpression of OTULIN chain-binding mutants lead to increased Met1-polyUb in cells (Figure 4A). Moreover, this interaction most likely recruits OTULIN to LUBAC targets. However, the functional consequences of a LUBAC-OTULIN interaction are unclear, given that HOIP stability, recruitment, and activity appear to be unaffected, and this requires further investigation.

(E) NF-κB luciferase activity in control and OTULIN-depleted cell lines after treatment with TNFα (10 ng/ml) or poly(I:C) (1 μg/ml). Error bars represent SD from the mean of experiments performed in duplicate. *p values are given to indicate significance. **, the control set to 1; n.s., nonsignificant.
(F) Analysis of selected NF-κB target genes by qPCR in control and OTULIN-depleted cell lines over a time course of TNFα stimulation (10 ng/ml).
(G) Jurkat cells transfected with nontargeting (NT) control siRNA or three different OTULIN-specific siRNAs were treated with TNFα (25 ng/ml) as indicated. EMSA signals were quantified by densitometry. SP, smart pool.
(H) OTULIN or control siRNA-transfected Jurkat cells were stimulated for 15 and 30 min with TNFα (25 ng/ml). After NEMO IP, kinase assays were performed with GST-IκBα (aa 1–53) as a substrate and quantified by densitometry.
Regarding the functional requirement for Met1-linked chains in NF-κB responses, key questions remain to be answered. First, in contrast to Lys63-linked chains, for which roles in NF-κB signaling have long been verified, for example, by elegant replacement strategies (Xu et al., 2009), Met1-linked polymers have a very low abundance, and are hard to detect in cell lysates. This may change with the discovery that catalytically inactive OTULIN (Figure 4A) or OTULIN knockdown (Figure 7C) leads to the stabilization of Met1 linkages. A second, more important issue is the lack of proteins modified with Met1-polyUb in vivo after a physiological stimulus. We believe that OTULIN can also be instrumental here, given that its specificity can be exploited as a Met1-specific Ub-binding protein when inactivated or in mass spectrometry methods. We anticipate that the identification of LUBAC and OTULIN targets may reveal a surprising variety of cellular proteins not restricted to NF-κB signaling.

**Role of OTULIN in Cytokine Responses**

Our data identify roles of OTULIN in TNFα signaling, which is in agreement with reported roles of Met1-polyUb in this pathway (Haas et al., 2009; Tokunaga et al., 2009; Walczak et al., 2012). However, in TNFα signaling, the putative redundant or nonredundant involvement of many different Ub chain types leads to a complicated interplay of Ub signaling (Kulathu and Komander, 2012). The functional importance of Met1-linked chains in the mix is supported by genetic evidence from cpdm mice lacking SHARPIN (Gerlach et al., 2011; Ikeda et al., 2011; Tokunaga et al., 2011) and data from human patients with mutations in HOIL1 (Boisson et al., 2012). So far, there is no unifying molecular mechanism that explains why so many different forms of polyUb are seemingly important for the activation of identical kinase cascades upon TNFα stimulation. Interestingly, the effects of OTULIN overexpression or knockdown seem more pronounced in poly(l:C) versus TNFα signaling (Figures 6A and 7E), and it will be important to study roles of OTULIN in other pathways that depend on Met1-linked chains, such as NOD2 signaling (Damgaard et al., 2012).

Our data support a role of Met1 linkages in providing an important scaffold for productive complex formation, given that the loss of Met1-linked chains induced by OTULIN overexpression prevents the association of NEMO with ubiquitinated RIPK1. Ubiquitinated RIPK1 is targeted also by the DUBs A20 (Hymowitz and Wertz, 2010) and CYLD (Sun, 2010). CYLD hydrolyzes Met1 linkages (Komander et al., 2009) and, hence, could also contribute to antagonize LUBAC signaling. However, although A20 and CYLD are induced by TNFα for the establishment of a negative feedback loop, neither OTULIN (Figure 7F) nor LUBAC (Haas et al., 2009) are induced by TNFα, again suggesting that they function as a differentially regulated signaling module. OTULIN is subject to phosphorylation, acetylation, and ubiquitination in cells, and it will be interesting to study whether these modifications regulate its activity or function.

Overall, our data suggest that the identified DUB OTULIN is an antagonist of the LUBAC E3 ligase complex. Consistent with reported roles for Met1-polyUb, changing OTULIN expression affects LUBAC-mediated processes, including NF-κB signaling. Detailed genetic analysis will be necessary in order to understand how LUBAC and OTULIN balance Met1-polyUb chains in vivo. This might confirm some reported roles and may reveal new cellular roles for this atypical Ub chain type.

**EXPERIMENTAL PROCEDURES**

**Identification of OTULIN**

FAM105B/OUTLIN was identified bioinformatically with generalized profile analysis, as described in the Extended Experimental Procedures.

**Cloning, Expression, and Purification**

FAM105B was cloned from brain complementary DNA (Invitrogen), expressed in E. coli from pPOPIN-F vector, and purified by anion exchange and gel filtration.

**Crystallography and Structure Determination**

Apo, SeMet-substituted, and mutant OTULIN were crystalized from 100 mM MES and imidazole, 30 mM MgCl₂, 30 mM CaCl₂, 10% (w/v) PEG 4k, and 20% glycerol (pH 6.5). The structure of apo OTULIN was determined by single anomalous dispersion, and subsequent structures were determined by molecular replacement.

**Qualitative DUB Linkage Specificity Assay**

Qualitative deubiquitination assays were performed as previously described (Komander et al., 2009).

**Binding Studies and DUB Kinetics**

Binding studies were performed as in Ye et al. (2011), and kinetic studies were performed as in Virdee et al. (2010) with FIAsh-tagged Met1-dUb variants.

**Cellular Studies of OTULIN**

OTULIN and A20 were expressed from pOPIN-F or pcDNA4/TO/MRGS6H, HOIP, SHARPIN and HOIL-1L from pcDNA3.1, and GST-tagged NEMO were expressed from pcDNA4/TO/MRGS6H-OTULIN or a pDEST30-EmGFP construct with an miRNA targeting the 3’ untranslated region of FAM105B were generated according to the manufacturer’s protocol (Invitrogen). Knockdown analysis was performed in miRNA cell lines or with eight different siRNA sequences, as listed in the Extended Experimental Procedures.

**NF-κB Activity Analysis**

NF-κB activity was assessed by luciferase assays, for which cells were cotransfected with MSP sin rev NF-κB firefly and pRL-TK Renilla (Promega) luciferase plasmids by immunofluorescence with anti-p65 staining or by quantitative PCR, as described in the Extended Experimental Procedures. TNFα signaling was analyzed by western blotting with the antibodies listed in the Extended Experimental Procedures.

**ACCESSION NUMBERS**

Coordinates and structure factors have been deposited in the Protein Data Bank and can be found at accession numbers 3ZNV (WT OTULIN apo), 3ZNX (OTULIN apo D336A), and 3ZNZ (OTULIN-Met1-diUb).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, six figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.05.014.

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