**ARTICLE**

**TBK1 phosphorylation activates LIR-dependent degradation of the inflammation repressor TNIP1**

Jianwen Zhou¹β, Nikoline Lander Rasmussen²*, Hallvard Lauritz Olsvik², Vyacheslav Akimov³, Zehan Hu¹, Gry Evjen⁴, Stéphanie Kaeser-Pebernard⁵, Devanarayanan Siva Sankar⁵, Carole Roubat⁵, Pauline Verhac⁴, Nicole van de Beck⁶, Fulvio Reggiori⁴,⁵,⁶, Yakubu Princely Abudu⁷, Blagoy Blagoev⁵, Trond Lamark⁵, Terje Johansen⁵, and Jörn Dengjel⁵

Limitation of excessive inflammation due to selective degradation of pro-inflammatory proteins is one of the cytoprotective functions attributed to autophagy. In the current study, we highlight that selective autophagy also plays a vital role in promoting the establishment of a robust inflammatory response. Under inflammatory conditions, here TLR3-activation by poly(I:C) treatment, the inflammation repressor TNIP1 (TNFAIP3 interacting protein 1) is phosphorylated by Tank-binding kinase 1 (TBK1) activating an LIR motif that leads to the selective autophagy-dependent degradation of TNIP1, supporting the expression of pro-inflammatory genes and proteins. This selective autophagy efficiently reduces TNIP1 protein levels early (0–4 h) upon poly(I:C) treatment to allow efficient initiation of the inflammatory response. At 6 h, TNIP1 levels are restored due to increased transcription avoiding sustained inflammation. Thus, similarly as in cancer, autophagy may play a dual role in controlling inflammation depending on the exact state and timing of the inflammatory response.

**Introduction**

Macroautophagy, hereafter referred to as autophagy, is a primary cytoprotective process that leads to the removal and lysosomal degradation of non-functional and/or superfluous cytoplasmic material. Autophagy is a constitutive process but may also be triggered by various forms of cell stresses (Mizushima and Levine, 2020). It involves the formation of a double membrane vesicle known as autophagosome that enwraps portions of the cytoplasm, including organelles. The content of the autophagosome is then targeted for degradation through the lysosome. Constitutive autophagy is regarded as a non-selective, bulk process, whereas stress-induced autophagy is often selective, aiming at removing the causes of stress, e.g., depolarized mitochondria in oxidative stress (Youle, 2019). Selective autophagy is carried out either by direct interactions between cargo and lipidated human ATG8 family members (MAP1LC3-A, -B, -C, GABARAP, GABARAPL1, GABARAPL2, commonly referred to as LC3s) that are anchored to autophagosomal membranes or by indirect interactions in which so-called selective autophagy receptors (SARs) tether cargo to LC3s (Johansen and Lamark, 2020). Cargo and receptors are then both degraded within lysosomes (Morishita and Mizushima, 2019), p62/SQSTM1, which is considered the founding member of the protein class of p62/SQSTM1-like receptors (SLRs), recognizes poly-ubiquitinated proteins/organelles destined for lysosomal degradation through its ubiquitin-associated (UBA) domain and interacts with lipidated LC3s through its LC3-interacting region (LIR) motif (Johansen and Lamark, 2020; Pankiv et al., 2007). SLRs are involved in the selective, autophagy-dependent degradation of a highly diverse set of substrates (Zellner et al., 2021).

The limitation of deleterious inflammatory responses is one of the cytoprotective functions attributed to autophagy (Deretic and Levine, 2018). The link between autophagy and tissue inflammation became obvious by genome-wide association studies that identified ATG16L1 as susceptibility locus for Crohn’s disease (Hampe et al., 2007). Since then several autophagy loci have been linked to inflammatory and autoimmune disorders (Mizushima and Levine, 2020). The selective degradation of inflammasome components, e.g., by p62/SQSTM1, is among the best-understood functions of autophagy in limiting tissue inflammation (Deretic and Levine, 2018; Samie et al., 2018). However, autophagy was also shown to support unconventional secretion of the pro-inflammatory cytokine IL1B (Dupont et al., 2011; Zhang et al., 2015), which argues for a fine-tuned and
balanced role of autophagy in regulating inflammatory responses.

TNIP1 (also known as ABIN-1, Nafl, and VAN) is a ubiquitin-binding adaptor protein that has been implicated as a negative regulator of inflammatory signaling and cytokine-induced cell death (Dziedzic et al., 2018; Gao et al., 2011; Oshima et al., 2009; Su et al., 2019; Zhou et al., 2011). Interestingly, recent studies linked TNIP1, either as cargo or as receptor, to autophagy-dependent protein degradation (Shinkawa et al., 2022). Selective autophagy-dependent degradation of TNIP1 through interaction with the SLR optineurin (OPTN) was reported, supporting senescence-associated inflammation (Lee et al., 2021). While TNIP1 itself shows no catalytic activity, its ability to bind linear polyubiquitin chains through its Ub-binding domain in ABIN proteins and NEMO (UBAN) is important for its anti-inflammatory function (Nanda et al., 2011; Wagner et al., 2008). A number of studies suggest that TNIP1 exerts its negative function by recruiting the ubiquititin-editing enzyme TNFAIP3 (also known as A20) to polyubiquitinated targets (Dziedzic et al., 2018; Gao et al., 2011; Mauro et al., 2006). However, the exact mechanism behind this negative regulation is not completely understood. Furthermore, TNIP1 shows activity independent of TNFAIP3 (Kattah et al., 2018; Oshima et al., 2009). The importance of TNIP1 as an anti-inflammatory signal transducer is highlighted by numerous studies implicating TNPI dysregulation in autoimmune disorders (Shamilov and Aneskievich, 2018). Uncovering the molecular function and dynamics of TNIP1 could, therefore, be valuable in understanding the mechanisms behind such complex disorders.

In the current study, we characterize TNIP1 as an autophagy substrate, which is selectively degraded at an early stage (0–4 h) under inflammatory conditions. We highlight that TNIP1 fulfills the structural characteristics of an autophagy receptor with an LIR motif. Upon TLR3 activation, TNIP1 is phosphorylated by TBK1 on LIR proximal serine residues to increase binding to LC3s. Hence, TNIP1 is selectively degraded by autophagy in order to promote a competent initiation of pro-inflammatory signaling.

Results

TNIP1 is ubiquitinated and degraded within lysosomes

To identify new proteins potentially involved in autophagy regulation or autophagosome targeting, we screened ubiquitination dynamics by quantitative mass spectrometry (MS)-based proteomics. U2OS and HeLa cells were differentially labeled by stable isotope labeling by amino acids in cell culture (SILAC), and autophagy was induced by inhibiting mTORC1 by rapamycin. TNIP1 is ubiquitinated and degraded within lysosomes

One protein that caught our attention was TNIP1/ABIN-1, which was identified as significantly ubiquitinated on the amino acid residue Lys389 (Fig. 1 C). TNIP1 is a key repressor of inflammatory signaling (Shamilov and Aneskievich, 2018), and posttranslational mechanisms regulating its protein abundance are largely unknown. In a reverse affinity purification (AP), we used U2OS-StUbEx cells indubitably expressing 6xHis-FLAG-tagged ubiquitin (Akimov et al., 2014), treated cells with rapamycin, or starved for amino acids with and without ConA and used Ni-NTA beads to enrich ubiquitinated proteins. Anti-TNIP1 immunoblotting validated the MS findings and characterized TNIP1 as increasingly ubiquitinated in cells in which lysosomal degradation was inhibited (Fig. 1 D and Fig. S1 A).

As only a fraction of TNIP1 appeared to be ubiquitinated, we investigated whether ubiquitination of Lys389 is necessary for lysosomal targeting of TNIP1. We performed site-directed mutagenesis and analyzed the stability and ubiquitination of wild-type (TNIP1WT) and respective TNIP1 variants. Next to Lys389, we also mutated the neighboring residue Lys371, which we identified in two out of the three SILAC experiments. The blockage of lysosomal degradation led to the accumulation of modified and non-modified variants of TNIP1, indicating that ubiquitination is not decisive for lysosomal degradation (Fig. S1 B). In addition, arginine-variants did neither exhibit alterations in their global ubiquitination pattern nor in their stability, indicating the presence of additional ubiquitination sites, which agrees with database entries (Fig. S1, B–F). Together, these results indicate that non-ubiquitinated and multi-ubiquitinated variants of TNIP1 accumulate upon the blockage of lysosomal acidification and that TNIP1 may be an autophagy substrate. Due to its importance in inflammation, we decided to study the regulation of TNIP1 protein abundance in more detail.

TNIP1 is degraded by autophagy

Because the blockage of lysosomal degradation by ConA led to an accumulation of non- and ubiquitinated TNIP1 variants, we examined whether TNIP1 is degraded by autophagy and whether pro teaseosomal degradation also contributes to regulating TNIP1 protein abundance under basal conditions. TNIP1 behaved similarly as p62/SQSTM1 under basal and mTORC1 inhibited conditions using Torin-1, while the inhibition of lysosomal acidification by ConA led to a significant accumulation of TNIP1 protein, inhibition of the proteasome by MG132 did not (Fig. 2, A and B). Furthermore, confocal immunofluorescent imaging

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showed endogenous TNIP1 accumulation upon Bafilomycin A1 (BafA1), another V-type ATPase inhibitor, treatment (Fig. S2). Importantly, under these conditions TNIP1 colocalized with LC3-positive structures, which also strongly overlapped with p62/SQSTM1. Using airyscan super-resolution confocal microscopy, we found that TNIP1 was located inside LAMP1- and LC3-positive structures in untreated cells, and that TNIP1 is accumulated within these structures upon BafA1 treatment (Fig. 2 C). Another way to test for lysosomal degradation is using a tandem tag autophagy flux reporter system (Pankiv et al., 2007), in which TNIP1 is fused to a tandem mCherry-EYFP tag. While EYFP is sensitive to low pH and therefore quickly loses its fluorescence in acidic lysosomes, the mCherry tag is rather stable under these conditions. In neutral cytosol, both tags of mCherry-EYFP-TNIP1 will be visible, while in lysosomes, only the mCherry-tag will fluoresce and appear as red-only dots. Transient transfection of mCherry-EYFP-TNIP1 did indeed lead to the formation of many red-only dots per cell, supporting the notion that TNIP1 ends up in lysosomal structures (Fig. 2 D). Taken together, this suggests that TNIP1 is degraded by autophagy under basal conditions as...
Figure 2. **TNIP1 is degraded by autophagy.** (A) U2OS cells were treated with 1 μM Torin-1 for 4 h; proteasomal or lysosomal degradation were inhibited by 10 μM MG132 or 2 nM ConA, respectively. Under fed conditions (DMSO) and in Torin-1 treated cells blockage of lysosomal acidification led to a significant increase of TNIP1 protein abundance. Shown are representative blots of three biological replicates. (B) Quantification of blots shown in A (n = 3). * = P < 0.05, ** = P < 0.01, *** = P < 0.001 unpaired, two-sided t test compared to DMSO treated samples. Error bars indicate SEM. (C) Confocal images showing...
colocalization between endogenous TNIP1, LAMPI and LC3 in U2OS cells treated for 5 h with either BafA1 or vehicle DMSO. Cells were immunostained against endogenous TNIP1 (yellow), LAMPI (cyan), and LC3 (purple) and imaged by Airyscan using the Zeiss LSM880 confocal microscope. Inserts highlight TNIP1 localized in LAMPI- and LC3-positive structures. Due to BafA1 treatment leading to accumulation of the immunostained proteins, signal intensities in the DMSO image have been increased relative to the BafA1 treated image during post-processing. Scale bars are 5 µm for the airyscan images and 1 µm for the inserts. (d) U2OS cells were transiently transfected with either mCherry-EYFP or mCherry-EYFP-TNPI2 24 h after transfection, cells were either left untreated or treated with BafA1 for 4 h. BF = bright field. Scale bars, 20 µm. Quantification of red-only TNPI1 dots over total TNPI1 dots was done using Velocity software (PerkinElmer), with intensity cut-offs based on BafA1 intensity of red and green dots (n = 3). Around 2,000–3,000 dots were counted for each condition within each replicate. * P < 0.05, unpaired, two-sided t test. Error bars indicate SD. In B and D, data distribution was assumed to be normal, but this was not formally tested. Source data are available for this figure: SourceData F2.

well as upon Torin-1 treatment, with proteasomal protein degradation playing a negligible role.

Next, we aimed to characterize the molecular events that led to autophagosomal recruitment of TNPI1. We immunoprecipitated HA-tagged TNPI1 and performed an MS analysis of its interactome. This screen identified a number of autophagy-related proteins as possible TNPI1 interactors, including p62/SQSTM1, TAXIBPI, OPTN, and TBK1 (Fig. 3, A and B and Table S2). The former two, p62/SQSTM1 and TAXIBPI, were also identified as MS enriched in IPs of endogenous TNPI1 under basal conditions (Fig. S3 A). The interaction with p62/SQSTM1 was also observed by immunoprecipitation followed by Western blotting (Fig. 3 C). In ATG101 KO, FIP200 KO, and pentaKO HeLa cells (which do not express the SLRs p62/SQSTM1, NBRI, NDP52, TAXIBPI and OPTN [Sarraf et al., 2020]) TNPI1 no longer showed stabilization upon lysosomal blockage (Fig. 3, D and E). This was also true for single KO’s of p62 and OPTN (but not TAXIBPI), which also showed a reduction in TNPI1 abundance upon lysosomal blockage (Fig. S3, B and C). Taken together, this suggests that more than one SLR is mediating the basal autophagic turnover of TNPI1. The staining of endogenous TNPI1 in U2OS cells showed an increase in the number of TNPI1 puncta (Fig. 3, F and G) and increased colocalization between TNPI1 and the SLRs p62/SQSTM1, TAXIBPI, and NDP52 upon BafA1 treatment (Fig. 3 H). Thus, under basal conditions, TNPI1 interacts with SLRs, which target TNPI1 to autophagosomes for lysosomal degradation. To further investigate the basal turnover, we analyzed the autophagic flux using mCherry-EYFP-TNPI1 in U2OS cells KO for FIP200, ATG9, ATG7, and ATG16L1. As quantified in Fig. 3 I, FIP200 and ATG9 KO cells showed little to no red-only dots. This is consistent with the importance of these proteins in both ATG7-dependent and -independent autophagy (Goodwin et al., 2017). ATG7 and ATG16L1 KO’s showed reduced formation of red-only dots, but not a complete loss, indicating that there may also be ATG7-independent degradation of TNPI1 under basal conditions. Altogether, these results suggest that autophagic degradation of TNPI1 under basal conditions is aided by SLRs, and that this turnover may be both ATG7-dependent and -independent.

TNPI1 interacts with human ATG8 family proteins through LIR motifs

Having established that TNPI1 colocalizes with several SLRs and is degraded by autophagy, we aimed to examine whether TNPI1 may also function independently of SLRs in autophagy. TNPI1 contains coiled-coil domains for oligomerization and a ubiquitin-binding UBAN domain similar to that of OPTN, which preferentially binds K63- and M1-linked polyubiquitin chains (Fig. 4 A; Herhaus et al., 2019; Wagner et al., 2008). TNPI1 only needs an LIR motif interacting with ATG8 proteins to be able to function as a SAR itself. Hence, we performed a peptide array screen for potential LIR motifs in human TNPI1. The peptide array, containing overlapping 20-mer peptides of TNPI1 moved by increments of three amino acids to cover the 636 amino acids full-length sequence, was probed with GST-GABARAP and revealed two potential LIRs in TNPI1, with core sequences 83-FDPL-86 and 125-FEVV-128, respectively (Fig. 4 B). The N-terminal region harboring these LIR motifs is missing in the two other TNPI family members, TNPI2 and -3. Of these two candidates, only LIR2 is conserved in the evolution of vertebrates down to cartilaginous fishes, while LIR1 is only conserved down to marsupials and is not found in platypus (Fig. 4 C). LIR2 also has acidic- and phosphorylatable residues flanking the core LIR motif, making it a very strong candidate for a functional LIR that could be positively regulated by phosphorylation (Johansen and Lamark, 2020; Wirth et al., 2019). LIR1 has a proline within the core motif which is usually inhibitory to binding to the ATG8s (Alemu et al., 2012; Johansen and Lamark, 2020). We then further tested the interaction between TNPI1 and the six human ATG8 family proteins by GST-pulldown assays. GST and GST-tagged human LC3 and GABARAP proteins were used to pull down in vitro translated wild-type TNPI1. Here, we observed that TNPI1 bound very well to several of the human ATG8s, with the strongest interaction being with LC3A, LC3B, GABARAP, GABARAPL1, and weak binding to LC3C and GABARAPL2 (Fig. 4, D and E). To test whether any of the two potential LIRs identified in the peptide array scan were responsible for this interaction, we mutated the conserved aromatic- and hydrophobic residues in each core LIR sequence to alanine, namely, F83A/L86A and F125A/V128A (Fig. 4 C). GST-pulldown assays with these mutants revealed that the F125A/V128A mutations in LIR2 strongly reduced the interaction between TNPI1 and the GST-ATG8s, while the F83A/L86A mutations in LIR1 had neither or a very minor effect compared to mutating LIR2 alone (Fig. 4, D and E). We also tested the influence of LIRs for in vivo interactions of TNPI1 with LC3s using HeLa cells expressing HA-TNPI1 variants by performing anti-HA affinity purifications (APs) followed by anti-LC3A/B Western blots (Fig. 4, F and G). In agreement with the in vitro observations, TNPI1 with both LIR motifs mutated (mLIR1+2) interacted significantly weaker with LC3A/B in vivo. Thus, TNPI1 fulfills all structural characteristics of being an SLR, with LIR2 being the motif mainly responsible for binding to ATG8s.
Figure 3. **TNIP1 localizes to p62 bodies.** (A) HA-TNIP1 affinity purification (AP)-MS highlights its interaction with autophagy receptors. HeLa cells expressing HA-TNIP1 and vector control cells (ctrl.) were differentially SILAC-labeled and anti-HA APs were performed under basal conditions, followed by quantitative MS.
TNIP1 does not affect basal autophagy flux

We next asked whether TNIP1 was capable of regulating autophagy flux. For this, we compared the levels of several known SLRs as well as the ratio of LC3-I and II under basal and starvation conditions in WT HeLa and two TNIP1 KO clones. The ratio of LC3-I and LC3-II was not affected by TNIP1 KO, and neither was the conversion upon starvation by HBSS, suggesting that TNIP1 does not affect autophagic flux under the tested manner. Shown are representative blots of three biological replicates. HeLa WT cells, ATG101 KO cells, FIP200 KO cells and PentaKO cells were treated with 10 μM MG132 or 2 mM ConA for 4 h. TNIP1 abundance was significantly increased in ConA-treated HeLa WT cells, while treatment had no effect in HeLa ATG101 KO, FIP200 KO, and pentaKO cells. E shows quantifications of D (n = 3). * = P < 0.05, unpaired, two-sided t test. Error bars indicate SD.

Pro-inflammatory signaling induces LIR-dependent, autophagic degradation of TNIP1

So far, we characterized TNIP1 as a constitutive autophagic cargo as we did not observe changes in autophagosomal recruitment and lysosomal degradation based on the metabolic status of cells. To test whether inflammatory signaling could lead to specific effects, we treated cells with the double-stranded RNA mimic poly(I:C), which mimics viral infection and elicits a TLR3 signaling response (Glavan and Pavletic, 2014). After 4 h of poly(I:C) treatment, we observed a significant decrease in endogenous TNIP1 levels, followed by an increase after 6 h (Fig. 6, A and B). Parallel to the observed decrease of TNIP1, an increase in pro-inflammatory proteins like ISG15 and the chemokine CCL5 was observed (Fig. 6 A and Fig. S4 A). The decrease of TNIP1 was dependent on canonical autophagy, as loss of ATG7, ATG101, and FIP200 inhibited the response, but independent of SLRs, as loss of the SLRs p62/SQSTM1, NBR1, NDP52, TAX1BP1, and OPTN in a pentaKO cell line did not inhibit the response (Fig. 6, A and B). In agreement, lysosomal inhibition by ConA could block this decrease in WT and pentaKO cells but had no effect on the autophagy incompetent cell lines (Fig. S4 B). Proteasomal inhibition by MG132 did not stabilize TNIP1 levels in any of the cell lines (Fig. S4 B). The observed increase at 6 h of treatment is due to transcriptional upregulation of TNIP1 (Fig. 6 C). Hence, TNIP1 is itself specifically targeted to autophagosomes under poly(I:C) treatment. Confocal imaging showed that upon poly(I:C) treatment in WT cells, endogenous TNIP1 changes from being mostly diffuse to forming dots in the cytoplasm (Fig. 6, D and E). The formation of TNIP1 dots in response to poly(I:C) was also observed in ATG7- and pentaKO cells (Fig. 6, D and E). In ATG7 KO cells, there was an increased amount of TNIP1 dots already at the basal level, which increased even further upon poly(I:C) treatment. In pentaKO cells, we observed an increased diffuse TNIP1 staining of cells at the basal
Figure 4. TNIP1 interacts with human ATG8 family proteins through a LIR motif. (A) Schematic drawing of the domain architecture of TNIP1, showing possible LIRs. (B) Peptide array of 20-mer peptides covering full length TNIP1 was used to probe for possible LIRs, using GST-GABARAP. (C) Amino-acid sequence alignment showing conservation of the core consensus LIRs in TNIP1 across species. TNIP1 amino acid sequences were collected from UniProt, and multiple sequence alignment performed with Clustal Omega. Asterisk (*) indicates fully conserved residues; colon (:) indicates conservation between groups of strongly similar properties (>0.5 Gonnet PAM 250 matrix); and a period (.) indicates conservation between groups of weakly similar properties (0–0.5 Gonnet PAM 250 matrix). Mutated residues for the LIR mutants (mLIR1 and mLIR2) are shown in red. (D) In vitro GST-pulldown assay using 35S-labeled myc-TNIP1, myc-TNIP1-F83A/L86A (mLIR1), myc-TNIP1-F125A/V128A (mLIR2) and myc-TNIP1-F83A/L86A/F125A/V128A (mLIR1+2) against recombinant GST and GST-Zhou et al. Journal of Cell Biology 8 of 22 Degradation of TNIP1 by selective autophagy https://doi.org/10.1083/jcb.202108144

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level likely reflecting an increase in basal protein levels. Upon 4 h stimulation with poly(I:C) TNIP1 puncta increased also in pentaKO cells similarly as in ATG7 KO cells. Taken together, our data suggest that neither ATG7 nor SLRs are required for poly(I:C)-induced TNIP1 aggregation into dots. However, ATG7, but not SLRs, is necessary for TNIP1 degradation upon poly(I:C) stimulation.

Because our data suggested that TNIP1 is degraded specifically by autophagy upon poly(I:C) exposure, we investigated whether this was LIR-dependent. To this end, we observed that in cells reconstituted with HA-TNIP1-WT, 6 h of poly(I:C) treatment led to a significant decrease in TNIP1 levels, while we did not observe this decrease in cells expressing HA-TNIP1-mLIR1+2 (Fig. 6, F and G; and Fig. S4 C). ISG15 levels anti-correlated with HA-TNIP1 levels confirming the inhibitory role of TNIP1 in regulating ISG15 expression (Fig. 6, F and G; and Fig. S4 A). Importantly, the relative increase of ISG15 was significantly higher in HA-TNIP1-WT compared to HA-TNIP1-mLIR1+2 expressing cells, supporting the interpretation that LIR-dependent degradation of TNIP1 is critical for a stimulus- and time-dependent expression of pro-inflammatory genes. Note that the time-dependent increase of endogenous and exogenous TNIP1 after prolonged poly(I:C) treatment differed, indicating a long-term transcriptional and/or translational regulation next to the observed short-term autophagy-dependent effects (Fig. 6, F and G; and Fig. S4 D). TNIP1 expression has previously been shown to be transcriptionally regulated by NF-κB, making it likely that this translates in a characteristic modulation of TNIP1 expression. To further test the LIR-dependent recruitment of TNIP1 to autophagosomes under poly(I:C) treatment, we performed anti-GFP-LC3B IPs followed by anti-TNIP1 Western blotting. In these IP experiments, we observed poly(I:C)-dependent increase in interaction of TNIP1 with LC3B in TNIP1 KO cells reconstituted with WT TNIP1 which was significantly reduced in KO cells reconstituted with TNIP1-mLIR1+2 (Fig. 6 H). Thus, pro-inflammatory signaling as exemplified by poly(I:C) treatment appears to lead to the specific, LIR-dependent degradation of TNIP1 by autophagy.

TBK1 positively regulates LIR-dependent interaction of OPTN with LC3B by phosphorylating the LIR motif of OPTN (Wild et al., 2011). In addition, TBK1 is an important mediator of TLR3-induced antiviral signaling (Louis et al., 2018). Since we identified TBK1 as a possible interaction partner of TNIP1 (Fig. 3 A), we tested whether TBK1 also plays a role in selective TNIP1 degradation. Indeed, confocal imaging showed that upon poly(I:C) treatment in WT cells, several of the observed endogenous TNIP1 dots colocalized with phosphorylated (Ser172), i.e., active, TBK1 (Fig. 7 A). Poly(I:C) treatment also led to a time-dependent activation of TBK1 as indicated by phosphorylation of Ser172 (Fig. 7 B). Interestingly, loss of TNIP1 led to an increased activation of TBK1 indicating a feedback regulatory mechanism.

With the TBK1-mediated regulation of the OPTN LIR in mind, we used MS-based phosphoproteomics to study whether TBK1 can phosphorylate serine and threonine residues located N-terminal to the core LIR sequence FEVV of LIR2 in TNIP1 (Fig. 4 C). Due to the surrounding amino acid sequence, we could not follow the standard workflow of bottom-up proteomics experiments using trypsin to generate respective peptides. Instead, we performed a multi-protease digestion protocol using Elastase or ProAlanase to generate optimal sequence coverage (Eisenhardt et al., 2016; Samodova et al., 2020). We were able to identify and quantify two phosphopeptide species which were phosphorylated within the TSS motif just in front of LIR2 (Fig. 7 C). Comparing cells treated or not with poly(I:C) and/or the TBK1 inhibitor MRT67307 clearly indicated that TBK1 phospho-lyates TNIP1 in a stimulus-dependent manner (Fig. 7 C). To determine whether TNIP1 is a direct target of TBK1, we performed in vitro kinase assays and could recapitulate the in vivo observations characterizing TNIP1 as a bona fide TBK1 substrate (Fig. 7 D). To test the effect of phosphorylations of the evolutionary conserved S122 and S123 residues N-terminal to the core LIR motif (Fig. 4 C), we analyzed the binding of the phosphomimicking S122E/S123E TNIP1 mutant to ATG8 family proteins by GST pulldown. Strikingly, the phosphomimicking mutant displayed strongly increased binding to all ATG8 proteins. The binding increase was particularly evident for LC3B (3.4-fold), LC3C (4.4-fold) and GABARAPL2 (5.8-fold; Fig. 7 E). Finally, we tested the effects of poly(I:C) treatment and TBK1 inhibition on TNIP1-LC3B interaction by GFP-LC3B IP. Inhibition of TBK1 led to a decreased interaction of TNIP1-WT with GFP-LC3B indicating that TBK1-dependent phosphorylation of the TNIP1 LIR motif was responsible for the observed increased interaction between LC3 and TNIP1 (Fig. 7 F). In agreement, pharmacological and genetic inhibition of TBK1 led to a stabilization of TNIP1 under poly(I:C) treatment, i.e., interfered with its stimulus-dependent degradation (Fig. 7, G and H). Thus, whereas SLRs appear to support autophagy-dependent, basal turnover of TNIP1, activation of its LIR motif by TBK1 induces its selective, autophagy-dependent removal, supporting the mounting of a transcriptional program to induce a robust inflammation response (Fig. 8).

Discussion

Autophagy is largely regarded as a cytoprotective, anti-inflammatory response ensuring cell and organismal homeostasis (Deretic, 2021; Deretic and Levine, 2018). Several autophagy loci have been linked to genetic predispositions for chronic..
Figure 5. **Loss of TNIP1 leads to an increase in inflammatory proteins.** (A) Loss of TNIP1 does not alter autophagy flux under basal and starvation conditions. TNIP1 knock out HeLa cells generated by CRISPR/Cas9 were used to study its effect on autophagy. Wild-type HeLa cells and the two TNIP1

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knockout clones denoted KO1 and KO2 were kept in either fed or starved (HBSS) conditions and treated with either vehicle (DMSO) or BafA1 for 8 h. Blots were probed for several known SLRs as well as LC3. (B) Quantification of blots shown in A (n = 3). Error bars indicate SD. (C) Loss of TNIP1 leads to an increased transcription of inflammatory genes. Fold changes of RNA and protein intensities of TNPI1 KO and WT cells were compared. Shown are average values of two KO clones compared to WT cells (n = 3 per cell type for RNAseq; n = 5 per cell type for proteomics). Genes that were significantly regulated on RNA and protein level are highlighted in red (P < 0.01). Genes linked to immune effector processes and interferon-stimulated genes are annotated. (D) Protein-protein interactions of significantly regulated proteins. Proteins highlighted in red in C were analyzed on known interactions using STRING DB (Schelter et al., 2019). Interactions between 17 proteins were identified, of which 13 are linked to stress response (marked in red). Thickness of edges indicate confidence of interactions. (E and F) Gene set enrichment analysis of significantly dysregulated mRNAs and proteins identifies an increased transcription and translation of genes involved in inflammation. NES denotes normalized enrichment score. (G–I) TNIP1 represses translation of pro-inflammatory gene products. Whereas knockout of TNPI1 led to an increased abundance of indicated inflammatory proteins (G), which is likely due to transcriptional changes (H, n = 3, error bars indicate SEM), re-expression of TNPI1 WT or TNPI1 mLIR1+2 blunted this phenotype (I). Source data are available for this figure: Source Data F5.

Inflammatory and autoimmune diseases. This indicates that the lack of removal of damaged organelles may indirectly promote tissue destabilizing pro-inflammatory signaling. Autophagy may also contribute on a direct molecular level to limit excessive inflammatory signaling, e.g., by the selective degradation of inflammasome components by SARs, such as p62/SQSTM1 (Shi et al., 2012). Furthermore, by removing other endogenous pro-inflammatory sources including damaged organelles and components from viral and bacterial infections, SARs play an important role in anti-inflammatory responses (Deretic, 2021). The pro-inflammatory functions of autophagy were so far limited to its contribution to interleukin secretion and to senescence-associated inflammation (Dupont et al., 2011; Lee et al., 2021; Zhang et al., 2015). In the current article, we highlight that the role of autophagy in inflammation appears to be more complex than anticipated and that both processes appear to be more intimately intertwined on a molecular level.

As TNIP1 has no reported enzymatic activity itself, it was believed that its functions in inflammation were directly linked to its interaction with TNFAIP3 (also known as A20), a ubiquitin-editing enzyme that contains both ubiquitin ligase and deubiquitinase activities and that was shown to negatively interfere with NF-κB signaling (Shamilov and Aneskievich, 2018; Song et al., 1996). Several recent studies, however, indicate that TNPI1 itself may function as a key repressor of inflammatory signaling, its dysregulation being linked to hyperinflammatory diseases like psoriasis (Nair et al., 2009), systemic lupus erythematosus (Gaveta et al., 2009), systemic sclerosis (Allanore et al., 2011), and senescence (Lee et al., 2021). TNPI1 is thus a potential target for the design of anti-inflammatory therapeutics. It has also been suggested that TNPI1 can function by out-competing other pro-inflammatory mediators for polyubiquitin binding, thereby negatively affecting inflammatory signaling (Shamilov and Aneskievich, 2018).

Whereas single nucleotide polymorphisms were shown to alter TNPI1 expression and microRNAs to decrease TNPI1 mRNA levels (Shamilov and Aneskievich, 2018), posttranslational mechanisms regulating TNPI1 protein abundance are largely unknown. We and others localized TNPI1 to autophagosomes, but did so far not address underlying mechanisms and phenotypical consequences (Meijvang et al., 2018; Shinkawa et al., 2022; Zellner et al., 2021). In the current study, we corroborate observations that TNPI1 can be regarded as a substrate for constitutive autophagosomal degradation. We did not identify changes in autophagy-dependent lysosomal degradation under fed and starved conditions, as well as under blocked mTORC1 signaling. The association of TNPI1 with SLRs and its colocalization with p62 bodies and LC3 puncta further supports this interpretation, indicating that ubiquitination of TNPI1 and direct protein-protein interactions lead to SLR-dependent lysosomal degradation (Fig. 8). Indeed, we could show that TNPI1 was still ubiquitinated even after mutating the two ubiquitination sites identified in this study. This is in agreement with www.phosphosite.org, which lists 11 ubiquitination sites of TNPI1, Lys889 being one of them (Akimov et al., 2018). In addition, also non-modified TNPI1 could be stabilized by lysosomal inhibition. Thus, TNPI1 appears to be constitutively degraded by autophagy, the role of ubiquitination being not entirely clear.

Interestingly, this changes under inflammatory conditions. Poly(I:C) treatment led to SLR-independent, selective removal of TNPI1 by autophagy dependent on its LIR motif. This suggests the existence of distinct TNPI1 pools within cells: non-modified TNPI1, phosphorylated and/or ubiquitinated TNPI1 (Fig. 8). This interpretation is supported by the observation that TNFAIP3 appeared to not be degraded by autophagy in a poly(I:C)-dependent manner, indicating that the pool of TNPI1 which interacts with TNFAIP3 under these conditions is also spared from degradation (see Fig. S4 A). The time-dependent regulation of TNPI1 protein levels indicates that autophagy-dependent degradation within the first 4 h of poly(I:C) treatment contributes to the establishment of a robust inflammatory response. At 6 h, TNPI1 levels rise again due to transcriptional upregulation starting already at 4 h. This regulation occurs to prevent excessive inflammatory signaling, which may lead to cell and tissue damage. In light of the model suggesting that TNPI1 can compete with pro-inflammatory proteins for ubiquitin binding (Shamilov and Aneskievich, 2018), it is possible that TNPI1 is degraded during the early stages of TLR3-activation to prevent this competition. The subsequent increase in TNPI1 at later stages of the signaling response could then be to outcompete other pro-inflammatory mediators in order to limit excessive signaling. In a recent study, OPTN was characterized as a direct, ubiquitination-independent binding partner of TNPI1 contributing to the specific, autophagy-dependent degradation of TNPI1 in senescent cells (Lee et al., 2021). The pathway characterized by us is different as TNPI1 degradation under inflammation conditions appears as SLR-independent (Fig. 8). OPTN and TNPI1 appear to function in parallel and to be positively linked on the transcriptional level since TNPI1 KO led to an upregulation of OPTN mRNA and protein levels, potentially in a compensatory fashion.
Figure 6. Poly(I:C) stimulation induces LIR-dependent, specific degradation of TNIP1 by autophagy. (A) Poly(I:C) treatment leads to time-dependent changes in TNIP1 abundance. Poly(I:C) stimulation leads to an autophagy dependent and SLR independent decrease of TNIP1 abundance within the first 4 h as
Our results strongly suggest that TBK1 is responsible for phosphorylation of the LIR2 proximal TSS motif. TBK1 is an important kinase in response to innate antiviral signaling. Upon TLR3-activation, TBK1 gets activated through TRIF and TRAF3 leading to the phosphorylation and activation of IRF3 and induction of type I interferons (Louis et al., 2018). TRIF itself may undergo autophagy-dependent degradation (Gentle et al., 2017; Inomata et al., 2012; Lim et al., 2019; Samie et al., 2018). How this anti-inflammatory effect of autophagy is coordinated with the described pro-inflammatory acting degradation of TNIP1 will be an interesting question to address in the future studies. TBK1 is also implicated in the regulation of autophagy and is known to phosphorylate several autophagy proteins (Oakes et al., 2017). It is possible that TBK1 promotes TNIP1 degradation in order to ensure the efficient activation of downstream interferon signaling. Our results indicate that TNIP1 may rely on the other SLRs for its basal turnover in unstimulated cells, but upon TBK1 activation in response to innate immune stimuli, the selective degradation of TNIP1 is promoted (Fig. 8). Interestingly, TNIP1 fulfills all characteristics of a SAR, having a UBD, LIR motif, and coiled-coil regions supporting multimerization. TLR1/2-induced degradation of TNIP1 was shown to mediate the degradation of MYD88 and IRAK1 (Shinkawa et al., 2022). Whether TLR3-activation leads to selective TNIP1-dependent degradation of specific cargo proteins will have to be addressed in future studies. The knockout of TNIP1 resulted in a basal increase in the expression and protein levels of several pro-inflammatory genes, even in the absence of pro-inflammatory signaling. This suggests that TNIP1 may have a role in preventing the induction of an inflammatory response even in untreated cells. Whether this is caused by TNIP1-mediated selective degradation of pro-inflammatory mediators or -complexes will also be important to address in future studies.

Taken together, we identified TBK1-dependent phosphorylation sites immediately N-terminal to the core LIR motif of TNIP1 that lead to its selective degradation by autophagy under inflammatory conditions. Autophagy may contribute to the establishment of a potent inflammatory response before it limits excessive cytotoxic inflammatory signaling. Thus in addition to cancer, inflammation is another condition in which autophagy may have a dual role either supporting or inhibiting underlying processes depending on the exact cell state and timing.
Figure 7. Poly(I:C) stimulation induces TBK1-dependent, specific degradation of TNIP1 by autophagy. (A) Immunofluorescence images showing colocalization between TNIP1 and pTBK1 upon poly(I:C) treatment. Cells were either left untreated or treated with 5 µg/ml poly(I:C) for 4 h, and subsequently...
stained for endogenous TNIP1 and pTBK1. Colocalization between TNIP1 and pTBK1 is indicated by arrowheads. Quantification of TNIP1 dots colocalizing with pTBK1 was done using Volocity software (PerkinElmer). Around 160–220 cells were counted for each condition in each independent experiment (n = 3). ∗ = P < 0.05, unpaired two-sided t test. Error bars indicate SD. Scale bar in overview image is 10 µm, and scale bar in insert is 2 µm. (B) Time-course effect of poly(I:C) treatment on TBK1 activation and TNIP1. Representative blot and the corresponding quantification of the relative pTBK1 over total TBK1 levels are shown. (C and D) TBK1 phosphorylates TNIP1 N-terminal of LIR2. (C) In vivo phosphoproteomics using Elastase or ProAlanase as proteolytic enzymes identified indicated phosphopeptides. The single phosphorylation site could not be unambiguously localized to one of the three amino acid residues highlighted in red. Inhibition of TBK1 blocked the respective phosphorylation event (n > 3). (D) In vitro kinase assay using purified TBK1 and TNIP1 coupled to phosphoproteomics indicates that TBK1 directly phosphorylates TNIP1 on one of the amino acid residues highlighted in red. ∗ = P < 0.05, unpaired two-sided t test. Error bars indicate SEM. (E) In vitro GST-pulldown assay using ^35S-labeled myc-TNIP1 and myc-TNIP1-S122E/S123E against recombinant GST and GST-tagged human ATG8s. Bound myc-TNIP1 WT and S122E/S123E was detected using autoradiography (AR). Quantification and fold change of n = 3, ∗ = P < 0.05, ∗∗ = P < 0.01, ∗∗∗ = P < 0.001; unpaired two-sided t test. Error bars indicate SD. (F) The interaction between TNIP1 and LC3B is regulated by TBK1. GFP-LC3B is purified using GFP trap beads. Bound TNIP1 is detected by Western blot. Inhibition of TBK1 by MRT67307 negatively regulates the poly(I:C)-dependent interaction of TNIP1 with LC3. KO1 cells were used for reconstitution. (G) Inhibition of TBK1 negatively interferes with poly(I:C)-dependent degradation of TNIP1. Western blots of whole cell lysate indicate TNIP1 stabilization by TBK1 inhibition. Actin was used as loading control (n = 3). Error bars indicate SEM. ** = P < 0.01, unpaired, two-sided t test. (H) TBK1 KO negatively interferes with poly(I:C)-dependent degradation of TNIP1. Western blots of whole cell lysate indicate TNIP1 stabilization by TBK1 KO in two independent cell lines. Tubulin was used as loading control (n = 3). Error bars indicate SD. ** = P < 0.01, unpaired, two-sided t test. In A, C, D, E, G, and H, data distribution was assumed to be normal, but this was not formally tested. Source data are available for this figure: SourceData F7.

Materials and methods

The reagents, antibodies, and plasmids used in this study are listed in Tables S4, S5, and S6, respectively.

Cell culture and cell treatments

HeLa cells and HEK293T cells were obtained from ATCC. U2OS cells were obtained from ECACC. StUbEx U2OS cells were a gift from Blagoy Blagoev (Department of Biochemistry and Molecular Biology, University of Southern Denmark, Odense, Denmark; Akimov et al., 2014). HeLa CCL2.2, HeLa CCL2.2 ATG7 KO cells, and HeLa CCL2.2 pentaKO cells were a gift from Richard J. Youle (National Institutes of Health, Bethesda, MD; Sarraf et al., 2020). HeLa p62 KOs have been described previously (Abudu et al., 2021). All cells were maintained in Dulbecco’s

![Figure 8](https://example.com/figure8.jpg)

**Figure 8.** Model of TNIP1 regulation. Under basal, unstimulated conditions (left panel), TNIP1 functions as a negative regulator of inflammatory signaling and is subject to constitutive autophagic degradation through interaction with autophagy receptors such as p62/SQSTM1. Upon poly(I:C)-induced activation of TLR3 (right panel), activated TBK1 phosphorylates TNIP1 in the vicinity of its LIR, increasing TNIP1 affinity for human LC3 and GABARAP proteins. This, in turn, leads to a LIR-dependent increase in TNIP1 degradation through selective autophagy. The removal of TNIP1 relieves the negative effect on inflammatory signaling, allowing the establishment of a robust inflammatory response upon antiviral signaling.
modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum, 2 mM L-glutamine, and 100 U/ml penicillin/streptomycin. For starvation experiments, cells were incubated with Hanks’ Balanced Salt solution (HBSS) for the indicated times. Cells were treated with 100 nM rapamycin, 200 nM bafilomycin A1, 5 µg/ml Concanamycin A, 20 µg/ml cycloheximide, or 10 µM MG132, for the indicated time periods. For SILAC, cells were grown for >14 d in SILAC DMEM supplemented with dialyzed fetal calf serum, 2 mM L-glutamine, 100 U/ml penicillin, and “light” (13C615N4-Arginine) stable isotope-labeled amino acids. For transient DNA transfection, subconfluent cells were transfected using TransIT LT1 transfection reagent according to the manufacturer’s instructions.

**Plasmid constructs**

Plasmids used in this study are listed in Table S4. Cloning into pDest-vectors was performed using the Gateway cloning system (Invitrogen). QuickChange site-directed mutagenesis kit (Stratagene) was used to create desired point mutations, which were verified by DNA sequencing (BigDye Sequencing kits, Applied Biosystems). Oligonucleotides for mutagenesis and sequencing were from invitrogen. TNIP1 cDNA was obtained from Genscript (NM_001252390). TNIP1 LIR mutation fragment was synthesized by IDT gBlocks Fragments (Integrated DNA Technologies). TNIP1 and TNIP1 mLIR were cloned into pDONR201 and pLenti CMV Blast DEST (706-1) by Gateway recombination cloning according to manufacturer’s instructions. All plasmid constructs were verified by sequencing (BigDye; Applied Biosystems) and/or restriction digestion.

**Generation of knockout (KO) and stable cell lines**

The TNIP1 KO cells were generated by CRISPR/Cas9-mediated genome-editing tools provided by the F. Zhang laboratory (Broad Institute, MIT, Boston, MA). A target sequence in exon twenty of human TNIP1 was selected (sgTNIP1-1, 5’-CGTACCGATGCTAGCACCC-3’; sgTNIP1-2, 5’-GGGCCCAGGTAGTTCCAACGGAC-3’; sgTNIP1-3, 5’-CAGCCGAAGCGTGCTGAC-3’). The sgRNA oligos were cloned into the Bbs1 site of hSpCas9 plasmid. Lipofectamine LTX and Plus reagent (15338100; Invitrogen) were used to transfect hSpCas9-sgRNA into HeLa cells according to the manufacturer’s instructions. Cells were selected with 2 µg/ml puromycin for 2 d and single cells were isolated by serial dilutions. TNIP1 deficiency was screened by immunoblotting.

CRISPR knockouts of FIP200 and ATG101 in HeLa cells was performed using sgRNAs cloned into the plasmid pX330-U6-Chimeric_BB-CBh-hSpCas9 (#42230; Addgene): ATG101, 5’- ACC AGAGAAAGAAGTCTCGC-3’ and 5’- GTTATCCACCTCGACTGTG-3’; FIP200, 5’- GTAGGTTTTTAGGAATAGGAGG-3’. CRISPR plasmids and pEGFP-puromycin plasmids were co-transfected in HeLa cells using lipofectamine LTX reagent (15338030; Thermo Fisher Scientific), according to manufacturer’s instructions. 24-h post-transfection, co-transfected cells were selected in culture medium supplemented with 3 µg/ml puromycin (Invivogen, ant-pr-1). Efficiently transfected cells were isolated to generate clonal lines by single-cell cloning in 96-well plates; colonies were all evaluated for KO efficiency by Western blotting against the targeted gene. The sgRNAs were designed using CHOP-CHOP CRISPR/gRNA algorithm (Labun et al., 2019).

Generation of HeLa CCL2.2 cells KO for OPTN and TAX1BP1 and HeLa KO for TBK1 was carried out as described previously (Abudu et al., 2021). The following gRNAs were used: sgOPTN: 5’-CCACGAGAACGCTCCAC-3’, sgTAX1BP1: 5’-AGACCTGCAAGGACGCTGCTGAC-3’. For generation of stable cell lines, lentiviral vectors were co-transfected with packaging (psPax2) and envelope plasmids (pMD2.G) in HEK293T cells using jetPrime reagent (Polyplus transfection). Medium was changed to fresh DMEM 15 h after transfection. After 24 h incubation, the supernatant was harvested by filtration through 0.22-µm filter. Polybrene (8 µg/ml) was added before infection of recipient cells. To obtain stable cell lines, infected cells were cultured in the presence of 5 µg/ml blasticidin for 1 wk and monitored by immunoblotting.

**Enrichment of ubiquitinated peptides**

Ubiquitinated peptides were purified according to Akimov et al. (2018). Triple SILAC labeled U2OS and HeLa cells were treated with DMSO, Rapa, and Rapa + ConA. Cells were washed twice with cold PBS and lysed with 12 ml lysis buffer (8 M guanidine-HCl, 25 mM ammonium bicarbonate [ABC], pH 8.5). The lysates were sonicated to reduce viscosity and cleared by centrifugation at 15,000 RCF for 30 min. Protein concentration was determined by BCA protein assay kit (23225 and 23227 Pierce), and the same amount of proteins of each label were mixed. Proteins were reduced with 2 mM DTT for 30 min at room temperature and alkylated with 11 mM chloro acet amide in the dark for 30 min at room temperature. The concentration of guanidine-HCl was diluted to 2 M with 25 mM ammonium bicarbonate and filtered with low binding 0.45-µm PVDF filters (Millipore). The Lys-C endopeptidase was added at a 1:100 enzyme-to-protein ratio to digest proteins at room temperature overnight. The peptide mixture was cleaned and purified using C18 cartridges (Waters) and lyophilized for 24–36 h. The lyophilized peptides were dissolved in 12 ml of IAP buffer (50 M MOPS, pH 7.2, 10 mM sodium phosphate, 50 mM NaCl, pH 7.5–8.0) plus 0.1% Triton X-100. Dissolved peptides were spun down and passed through low binging 0.45-µm PVDF filters (Millipore). The Lys-C endopeptidase was added at a 1:100 enzyme-to-protein ratio to digest proteins at room temperature overnight. The peptide mixture was cleaned and purified using C18 cartridges (Waters) and lyophilized for 24–36 h. The lyophilized peptides were dissolved in 12 ml of IAP buffer (50 M MOPS, pH 7.2, 10 mM sodium phosphate, 50 mM NaCl, pH 7.5–8.0) plus 0.1% Triton X-100. Dissolved peptides were spun down and passed through low binging 0.45-µm PVDF filters and incubated with 500 µl of Ubisite conjugated matrix for 5 h at 4°C. The Ubisite matrix was washed three times with IAP buffer without detergent and three times with 150 mM NaCl. After the third wash, the matrix was transferred to a small Poly-Prep column (Bio-RAD). Enriched peptides were eluted with 250 µl 0.1% TFA three times, and each time incubated for 5 min. The eluted peptides were pooled and neutralized with 1 M ABC buffer to a final concentration at 25 mM, followed by trypsin digestion overnight at 37°C. The fractionation of trypptic peptides was performed as described previously (Akimov et al., 2018). The resulting peptides were lyophilized and cleaned by STAGE-tips.

**Enrichment of ubiquitinated proteins by StUbEx**

StUbEx U2OS cells were induced with doxycycline for 48 h and treated with Rapa and lysosomal degradation was blocked with Anti-TNIP1 (Zhou et al., 2016).

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**Zhou et al.**

Degradation of TNIP1 by selective autophagy

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ConA. The cells were washed with cold PBS and lysed with binding buffer (6 M Guanidinium-HCl, 50 mM Na₂HPO₄/NaH₂PO₄, pH 8.0, 500 mM NaCl, 5 mM imidazole). Sample viscosity was reduced by sonication followed by centrifugation at 11,000 RCF for 30 min at room temperature. Protein concentration was determined by BCA assay. Ubiquitin conjugates were purified by cComplete His-tag purification beads (Roth) for 4 h at room temperature, followed by washes with binding buffer and washing buffers 1-3 (WB1—8 M Urea, 50 mM Na₂HPO₄/NaH₂PO₄, pH 8.0, 500 mM NaCl, 0.1% Triton-X-100; WB2—same as WB1 but with 0.2% Triton X-100; WB3—8 M Urea, 25 mM Tris, pH 8.0, 150 mM NaCl). The ubiquitin conjugates were finally eluted with three times elution buffer (150 mM NaCl, 50 mM Tris, pH 8.0, and 300 mM imidazole).

**Immunoblotting**

For immunoblot analysis, treated cells were harvested and lysed in modified RIPA buffer supplemented with 2% SDS and 0.1% benzonase Nuclease or 1× SDS buffer (50 mM Tris, pH 6.8, 2% SDS, and 10% glycerol). For cells harvested in modified RIPA buffer, lysates were centrifuged for 15 min at 11,000 RCF. For cells harvested in 1× SDS buffer, lysates were incubated at 100°C for 10 min. Protein concentration was measured using Pierce BCA Protein Assay Kit. Quantified cell lysates or AF eluates were either reduced with 2 mM DTT at 75°C for 10 min or 100 mM DTT at 100°C for 10 min and resolved on SDS-PAGE gels. Proteins were transferred to a PVDF or nitrocellulose membrane, and subsequently prepared for either fluorescent or chemiluminescent detection. For fluorescent detection, membranes were blocked in Intersect (PBS or TBS) Blocking Buffer, followed by overnight incubation at 4°C with primary antibody. Membranes were subsequently washed 4× in PBS or TBS containing 0.1% Tween (PBS/T/TBS-T), followed by incubation with secondary antibody diluted in Intersect Blocking Buffer. After 4× wash in PBS/T/TBS-T, a final wash was carried out in PBS/TBS without Tween. Fluorescent signal was detected using LiCOR Odyssey CLx imaging system. Chemiluminescent detection: membranes were blocked in either 5% milk or 5% bovine serum albumin in PBS-T or TBS-T, followed by incubation with primary antibody. Membranes were washed 3–4× in PBS/T-TBS/TBS-T followed by incubation with secondary antibody. After 3–4× washes in PBS/T-TBS/TBS-T, membranes were developed using SuperSignal West Pico Chemiluminescent Substrate on GE Fujiﬁlm LAS4000 Luminescent image analyzer or Odyssey Fc reader (LI-COR Biosciences-GmbH). The densitometry of immunoblotting was performed by either ImageJ (National Institutes of Health) or ImageStudio software (LI-COR Biosciences-GmbH).

**Immunofluorescence staining and confocal fluorescence microscopy**

For imaging, cells were grown on 1.5 cm round 12-mm coverslips (#631-0150; VWR) for immunofluorescence staining or 8-well Lab-Tek chamber coverglass for double-tag analysis (#155411; Thermo Fisher Scientific), and fixed with 4% formaldehyde in PBS for 30 min. For immunofluorescence staining, cells were washed 3× with PBS and then permeabilized with either 0.1% Triton X-100 in PBS for 10 min, or ice-cold methanol for 10 min. Next, cells were washed 5× with PBS or TBS, and blocked for 1 h in 5% BSA in PBS or TBS. Subsequently, cells were incubated with primary antibodies diluted in 1% BSA in PBS or TBS for 1-2 h at room temperature. After 5× wash in PBS or TBS, cells were incubated with Alexa Fluor secondary antibodies diluted in 1% BSA in PBS or TBS for 1 h at room temperature. After 5× wash in PBS or TBS, cell nuclei were stained with 1 µg/ml DAPI diluted in PBS for 5 min, followed by 2× final washes in PBS or TBS. Coverslips were mounted using ProLong Gold or ProLong Glass Antifade Mountant. Cells were imaged using Zeiss LSM800 or LSM880 (Carl Zeiss Microscopy) using a 63 × NA1.4 oil immersion lens for coverslips or a 40 × NA1.2 water immersion lens for chambered coverglass. Images were collected in ZEN software (Zeiss). For Airyscan super-resolution images, optimal pixel size and z spacing as suggested by ZEN was used. Optimal excitation and emission settings were determined using the Smart Setup function. All fluorescence channels were recorded at non-saturating levels, and settings were kept identical between all samples within replicates used for comparisons or quantifications.

**Image analysis**

For quantification of red-only dots of mCherry-EYFP-TNIP1, transiently transfected cells with low expression of mCherry-EYFP-TNIP1 were visually selected and imaged. Cells with high levels of overexpression were excluded, as this resulted in the formation of large aggregates. All images within each replicate were taken using identical settings. For quantification of relative amounts of red-only dots versus total dots, mCherry and EYFP dot detection was performed using a custom-made measurement protocol using intensity thresholding, size exclusion, and noise filtering, based on signal intensity of the BafA1 control in Volocity software (PerkinElmer) ver. 6.3. The number of mCherry-only positive dots was counted by subtracting total EYFP dots from total mCherry dots for each experiment. For quantification of the number of cells positive for red-only mCherry-EYFP-TNIP1 dots, the images were analyzed manually by a subject that was blind to the analyzed conditions.

To quantify the number of TNIP1 dots and the colocalization between TNIP1 and SLRs/pTBK1, populations of objects representing fluorescent puncta in each channel were segmented using a custom-made protocol in Volocity ver. 6.3 (PerkinElmer). Detection of TNIP1, SLRs, and pTBK1 puncta was performed by intensity thresholding, size exclusion, and noise reduction. Overlap between TNIP1 and pTBK1 was identified by excluding TNIP1 objects not touching SLRs/pTBK1. The percentage of TNIP1 dots colocalizing with SLRs/pTBK1 with the total number of TNIP1 dots. For line-profiles, the line-profile tool in Zen Blue software (Zeiss) was used to measure signal intensity across indicated lines.

**SPOT synthesis and peptide array**

TNIP1 peptide array was synthesized on cellulose membranes using MultiPrep peptide synthesizer (INTAVIS Bioanalytical Instruments AG). Membranes were blocked with 5% non-fat milk in TBS-T and peptide interactions were tested using GST-
GABARAP by overlaying the membrane with 1 µg/ml recombinant protein for 2 h at room temperature. Membranes were washed three times in TBS-T. Bound GST-GABARAP was visualized with HRP-conjugated anti-GST antibody (Johansen et al., 2017). Putative LIR motifs in 20, 3 arrays (20-mer peptides moved a window of 3 residues along the protein sequence) were identified as 4–6 consecutive strong spots containing the core LIR consensus [W/F/Y]XX(L/I/V).

**Recombinant protein production and GST-pulldown analysis**

GST and GST-fusion proteins were expressed in *Escherichia coli* strain SoluBL21 (DE3; #C700200; Genlantis). Protein expression was induced by adding 50 µg/ml isopropyl β-D-1-thiogalactopyranoside (IPTG). The bacterial cells were sonicated in lysis buffer (20 mM Tris-HCl, pH 7.5, 10 mM EDTA, 5 mM EGTA, 150 mM NaCl) and GST-fused proteins were immobilized on glutathione Sepharose 4 Fast Flow beads (#17-5132-01; GE Healthcare) by incubating in a rotator at 4°C for 1 h. The beads containing GST-fusion proteins were subsequently used for pulling down in vitro translated proteins or cell lysates. For in vitro translated proteins, a pDest-myc-vector containing the protein of interest and a T7 promoter was used. In vitro translation was performed using the TNT T7 Reticulocyte Lysate System (Promega Corp.), in the presence of radioactive 35S-methionine. In vitro translated protein was subsequently used for pulling down in vitro translated proteins or cell lysates. For in vitro translated proteins, a pDest-myc-vector containing the protein of interest and a T7 promoter was used. In vitro translation was performed using the TNT T7 Reticulocyte Lysate System (Promega Corp.), in the presence of radioactive 35S-methionine. In vitro translated protein was then precleared by incubation with empty Glutathione Sepharose beads in 100 µ1 of NETN buffer (50 mM Tris, pH 8.0, 150 mM NaCl, 1 mM EDTA, 0.5% NP-40) supplemented with complete Mini EDTA-free protease inhibitor (Merck) for 30 min at 4°C. Precleared lysates were then incubated with GST-fusion protein bound beads for 1–2 h on a rotator at 4°C. Beads were then washed 5× with NETN-buffer, and resuspended in 2 × SDS-PAGE gel loading buffer (125 mM Tris, pH 7.5, 4% SDS, 0.04% bromophenol blue, 20% glycerol, 100 mM dithiothreitol), boiled for 10 min, and resolved by SDS-PAGE. Gels were stained with Coomassie Brilliant Blue for protein visualization and then vacuum-dried. The radioactive signal was then detected on imaging plates with FujiFilm BAS-5000 (FujiFilm). Signals from 35S-labeled proteins were measured in terms of unit of photostimulated luminescent (PSL) and quantified in comparison with 10% of the in vitro translated lysate (input) using the Image Gauge software (Fuji; Johansen et al., 2017).

**Affinity purification**

Cells were lysed in ice-cold modified RIPA buffer (25 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA, 1% NP-40, 0.1% Sodium deoxycholate) containing complete protease inhibitor cocktail and phosphatase inhibitor cocktail. Lysates were centrifuged for 15 min at 13,000 RCF and protein concentration was determined by BCA assay. GFP-tagged proteins were affinity purified by GFP-trap (ChromoTek) according to the manufacturer’s instructions. HA-TNIP1 was affinity purified by anti-HA magnetic beads (Pierce). TNIP1 polyclonal antibody-coupled protein G dynabeads were used for affinity purification of endogenous TNIP1. For denaturing purification, the lysate was supplemented with 2% SDS and incubated for 30 min at room temperature to break protein–protein interaction, then diluted to 0.5% SDS with lysis buffer and incubated with HA or anti-TNIP1 antibody for affinity purification.

For TNIP1 interactome analysis by MS, HA-TNIP1 cells and empty vector cells were cultured in “heavy” and “light” SILAC medium, respectively. After 2 wk, cells were harvested and lysed in modified RIPA buffer containing complete protease inhibitor cocktail and phosphatase inhibitor cocktail. Protein concentration was determined by BCA assay, and protein amount was adjusted to equal concentration with lysis buffer, followed by affinity purification with anti-HA magnetic beads. The eluate of “heavy” and “light” samples was combined and fractionated by SDS-PAGE. Proteins were in-gel digested by trypsin and peptides were desalted by STAGE-tips prior to LC-MS/MS analysis.

**Whole proteome analysis**

For whole proteome analysis by MS, HeLa wild-type cells and TNIP1 KO cells were cultured in SILAC medium for 2 wk. Cells were harvested and lysed in modified RIPA buffer containing 2% SDS and 0.1% benzozene Nuclease. Protein concentration was determined by BCA assay and equal amounts of proteins in each label were mixed. Proteins were reduced with DTT and alkylated with IAA, followed with SDS-PAGE fractionation and trypsin in-gel digestion. Tryptic peptides were desalted by STAGE-tips prior LC-MS/MS analysis.

**TNIP1 phosphorylation analysis**

HA-TNIP1 cells were treated with poly(I:C) with or without TBK1 inhibitor MRT67307. Cells were harvested and lysed in modified RIPA buffer containing complete protease inhibitor cocktail and phosphatase inhibitor cocktail, 2% SDS was supplemented to break protein–protein interaction. The samples were diluted to 0.5% SDS with lysis buffer and subjected to affinity purification with anti-HA magnetic beads. After purification, the samples were transferred onto 10 kDa MW cut-off filter with 400 µl 8 M urea and 1 mM DTT. Proteins were digested by elastase or ProAlanase using the FASP protocol (Wisniewski et al., 2009). Digested peptides were eluted twice with 200 µl 50 mM ammonium bicarbonate into fresh tubes and acidified with TFA to a final concentration of 1%. Peptides were frozen and lyophilized prior to phosphopeptide enrichment. The lyophilized peptides were resuspended in 200 µl of 80% ACN, 0.1% TFA. Phosphopeptides enrichment was performed on Agilent AssayMAP Bravo platform according to manufacturer’s instructions. The Agilent AssayMAP phosphopeptide enrichment v2.0 App was used for automated phosphopeptide enrichment using Fe (III)-NTA cartridges (Basel, Switzerland). Cartridges were primed with 100 µl of 50% ACN, 0.1% TFA using a high flow rate of 300 µl/min and equilibrated using 80% ACN containing 0.1% TFA. Samples were loaded onto the cartridge using a low flow rate of 3 µl/min. The cartridges were washed twice with 200 µl 80% ACN containing 0.1% TFA and eluted with 50 µl of 1% ammonium hydroxide (pH 11) and 50 µl of 1% ammonium hydroxide in 80% ACN to a low-binding PCR tube containing 5 µl FA. The eluted peptides were lyophilized and resuspended in 20 µl of 0.1% FA for LC-MS/MS analysis.

**In vitro kinase assay**

HA-TNIP1 cells were harvested and lysed in ice-cold modified RIPA buffer containing complete protease inhibitor cocktail.
TNIP1 protein was enriched by anti-HA magnetic beads and dephosphorylated by lambda phosphatase with 1 mM MnCl2 for 30 min at 30°C. Beads were washed with 2 × 10 ml of kinase buffer (50 mM Tris–HCl, pH 7.6, 10 mM MgCl2, 150 mM NaCl and 1× PhosSTOP) to remove lambda phosphatase. Kinase buffer was added to a final volume of 800 μl with 1 μg TBK1 protein, for control sample, 10 μM TBK1 inhibitor MRT67307 was added. Kinase assays were performed on a rotor at 37°C for 1 h. Finally, reactions were quenched by addition of 8 M urea and 1 mM DTT.

The beads were transferred onto 10 kD MW cut-off filter for the digestion with ProAlanase, followed with phosphopeptide enrichment.

**LC-MS/MS analyses**

Solubilized peptides were injected into a 20 cm fused silica column with an inner diameter of 75 μm and in-house packed with C18 (ReproSil-Pur 120 C18-AQ, 1.9 μm, Dr. Maisch) for reverse phase fractionation by EasyLC 1200 nanoflow-HPLC system (Thermo Fisher Scientific). Peptides were loaded with solvent A (0.1% FA in water) at a max. pressure of 800 Bar and eluted with a step gradient of solvent B (0.1% FA in 80% ACN) from 2 to 25% within 85 min, from 25 to 60% within 5 min, followed by increasing to 100% in 2 min at a flow rate of 250 nl/min. MS/MS analysis was performed on a nano-electrospray ion source equipped QExactive HF-X mass spectrometer (Thermo Fisher Scientific). The spray voltage was set to 2.3 kV with a capillary temperature of 250°C. Mass spectrometer was operated in positive polarity mode and MS data were acquired in the data-dependent mode. The automatic gain control (AGC) target was set to 3 × 10^6; the resolution was set to 120,000, and the ion injection time was set to 15 ms for the full scan at a mass range of m/z = 370–1750. 12 precursors were fragmented using normalized collisional energy (NCE) of 28 by higher-energy collisional dissociation (HCD). MS/MS scans were acquired with a resolution of 30,000, AGC target of 100,000 maximum IT of 54 ms, isolation window of 1.6 m/z, and dynamic exclusion window of 30 s. MS raw files were analyzed using MaxQuant software version 1.6.2.110 (Cox and Mann, 2008), data were searched against UniProt full-length homo sapiens database (21,033 entries, released March, 2016). Cysteine carbamidomethylation was set as a fixed modification, protein amineterminal acetylation and methionine oxidation were set as variable modifications. Ubiquitination of lysine was set as a variable modification and digestion was specific to trypsin/P for Ubisite experiment. Serine-, threonine- and tyrosine-phosphorylation were set as a variable modification and digestion was specific to trypsin/P for Ubisite experiment. The analysis was carried out with “match-between-run” with a time window of 0.7 min. MaxQuant results were analyzed using Perseus (Tyanova et al., 2016).

**Total RNA extraction and library construction**

Four replicates of HeLa WT cells and TNIP1 KO cells (clone 1 and clone 2) were cultured in 10 cm dishes. The total RNA was extracted using RNeasy Mini Kit (Qiagen) according to the instructions of the manufacturer. Cells were lysed with 350 μl RLT buffer and homogenized by passing 5 times through a blunt 20-gauge needle. One volume of 70% ethanol was supplemented, the lysates were transferred to RNeasy spin column and centrifuged at 8,000 × g to collect RNA. The RNA was washed once with 700 μl RW3 buffer and twice with 500 μl RPE buffer. After completely removing the RPE buffer, the RNA was eluted with 50 μl RNase-free water. The quality of RNA samples was analyzed by RNA screen tape (Agilent). The complementary DNA (cDNA) libraries were barcoded using Illumina primers and sequenced on one lane of an Illumina Novaseq 6000 instrument with 2 × 50 bp paired-end sequencing cycles. The sequence data were deposited at the European Nucleotide Archive (accession number: PRJEB45902). The output reads from multiple lanes were combined into single files and quality control was performed with FastQC v0.11.7 and cleaned with fastsp v0.19.5 to remove pol yg tails and keep only full-length reads (Chen et al., 2018). The human genome GRCh38.p13 (ENSEMBL) was used to remap the reads using STAR v2.5.3a (Dobin et al., 2013). The differential expression between WT and KO was analyzed by R software (R Core Team, 2014 http://www.R-project.org/) using the DESeq2 package (Love et al., 2014).

**qRT-PCR**

The total RNA were purified as above, 1 μg RNA was subjected to reverse transcription by Quantitect RT kit (Qiagen). Quantitative PCR was performed using the KAPA SYBR Fast (Universal) qPCR kit (Merck Millipore) according to the manufacturer’s recommendations. The qRTPCR reaction was performed with a Rotor-GeneQ (Qiagen) and the same thermal profile conditions were set as 95°C for 10 min; then 40 cycles were performed of 10 s at 95°C, 20 s 60°C and 20 s 7°C. The following primer pairs were used: TNIP1 forward 5’-CTAGTGTCAGCGGAGTAAAGG-3’, TNIP1 reverse 5’-GCTGTTCCATGACCAGGAA-3’; Actin forward 5’-GGACTTGCAAGAGATGG-3’, Actin reverse 5’-AGCAGTGTTGGCCGTACAG-3’; HPRT1 forward 5’-TGACACTGGCAAACAAATCGA-3’, HPRT1 reverse 5’-GGTTCTTTTACCACGCAAGCT-3’; 18s rRNA forward 5’-CGCGAGAACCATTCCAAGAC-3’, 18s rRNA reverse 5’-GAATCGAACCCTGATTCGCCGTC-3’.

**Gene set enrichment analyses**

Gene set enrichment analyses (GSEA) was performed using default setting weighted enrichment statics and signalNoise metric for ranking genes (Subramanian et al., 2005). Significantly regulated genes in transcriptome data and proteins in whole proteome data were listed in Table S3.

**Quantification and statistical analyses**

Significantly regulated ubiquitination sites were determined by two samples paired t test, FDR < 0.05, S0 = 0.1. In all other cases unpaired, two-sided t tests were used unless stated otherwise. In general, parametric test were used, and data distribution was assumed to be normal, but this was not formally tested. The interacome network was generated by the String database. The statistical analyses including standard deviations, error bars, and P values were performed using Excel (Microsoft), all the details were indicated in figure legends.

**Online supplemental material**

Fig S1 shows TNIP1 gets ubiquitinated and degraded in the lysosome. Fig. S2 shows TNIP1 localizes to autophagosomes. Fig. S3...
shows endogenous TNIP1 interacts with TAXIBP1 and p62/SQSTM1 under basal conditions. Fig. S4 shows regulation of TNIP1 protein abundance. Table S1 shows UbiSite-proteomics approach to identify ubiquitination sites on proteins being potentially involved in autophagy. Table S2 shows HA-TNIP1 interactome. Table S3 shows RNA-Proteome correlation comparing WT and TNIP1 KO HeLa clones. Table S4 shows chemical reagents used in this study. Table S5 shows antibodies used in this study. Table S6 lists plasmids used in this study.

**Data availability**
The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium through the PRIDE (Perez-Riverol et al., 2019) partner repository with the dataset identifier PXD027163.

The RNA sequence data were deposited at the European Nucleotide Archive (accession no. PRJEB45902).

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Degradation of TNIP1 by selective autophagy

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Figure S1. **TNIP1 gets ubiquitinated and degraded in the lysosome.** (A) U2-OS-StUbEx cells inducibly expressing His-FLAG-tagged ubiquitin at endogenous levels were used to enrich ubiquitinated proteins (Akimov et al., 2014). Under control conditions as well as under starvation treatment (HBSS), TNIP1 gets ubiquitinated as shown by anti-TNIP1 immunoblots. Ubiquitinated TNIP1 was stabilized by the addition of concanamycin A (ConA) indicating its lysosomal degradation in treated and nontreated cells. Actin was used as loading control. (B and C) Mutations of identified TNIP1 ubiquitination sites do not lead to reduced lysosomal degradation as indicated by stabilized protein amounts by ConA treatment. This is the case for fed control conditions (DMEM) as well as under active autophagy (Rapa and HBSS treatment). C shows quantification of blots exemplified in B (n = 3, error bars indicate SD). (D and E) Mutated TNIP1K371/389R is still getting ubiquitinated as indicated by anti-TNIP1 IP followed by anti-ubiquitin (D) and anti-TNIP1 (E) Western blot. The addition of ConA leads in all cases to a stabilization of non-ubiquitinated and polyubiquitinated protein variants. (F) Identified ubiquitination sites according to PhosphoSitePlus database and this study. Gray bar depicts the amino acid sequence of TNIP1. Sections in green mark tryptic peptides identified in this study, i.e., sequence coverage of TNIP1. Amino acids marked in blue highlight published ubiquitination sites, number of references shown on y-axis. Amino acids marked in red were identified in this study. Source data are available for this figure: SourceData FS1.

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Figure S2. **TNIP1 localizes to autophagosomes.** U2OS cells kept in either fed or starved (HBSS) conditions and treated with either vehicle (DMSO) or BafA1 for 8 h. Cells were fixed and stained with antibodies against endogenous TNIP1 (green), p62 (orange), and LC3 (purple) and imaged using the Zeiss LSM800 confocal microscope. Line-profile co-localization plots were made using the line-profile quantification tool in the Zen blue imaging software (Zeiss). Vertical axis represents measurements of fluorescent intensity and the horizontal axis the drawn distances. Scale bar = 10 µm.
Figure S3.  **Endogenous TNIP1 interacts with TAX1BP1 and p62/SQSTM1 under basal conditions.** (A) SILAC-based, IP-MS analyses of anti-TNIP1 immunoprecipitations identified TAX1BP1 and p62/SQSTM1 as enriched compared to negative control IPs using beads only. GAPDH is shown as negative control. Shown are average values of three biological replicates (n = 3). Error bars: SD, **P < 0.01, ***P < 0.001, unpaired, two-sided t test. (B and C) Lysosomal turnover of TNIP1 is mediated by several SLRs. HeLa WT and HeLa p62, OPTN, and TAX1BP1 KO cells were either left untreated or treated with 200 nM BafA1 for 12 h. Shown are average values of three biological replicates. Error bars: SD, *P < 0.05, unpaired, two-sided t test. Source data are available for this figure: SourceData FS3.
Regulation of TNIP1 protein abundance. (A) Reduction of TNIP1 correlates with an increase of ISG15 and TNFAIP3/A20 in HeLa cells. The increase of the TNIP1 interaction partner TNFAIP3 under poly(I:C) treatment indicates the existence of distinct TNIP1 pools, i.e., free and bound to TNFAIP3. Arrow marks A20 band. (B) Poly(I:C) treatment leads to an autophagy-dependent and SLR-independent lysosomal degradation of TNIP1. Whereas lysosomal inhibition by ConA treatment leads to a significant block of TNIP1 degradation in WT and pentaKO cells, proteasomal inhibition by MG132 treatment has no effect. In ATG101 KO, FIP200 KO and ATG7 KO, autophagy incompetent cell lines poly(I:C) does not lead to TNIP1 degradation. TLR3 and SQSTM1 are monitored as positive controls, actin as loading control. Shown are representative blots of three biological replicates each. Bar diagram shows quantification, error bars: SEM. * = P < 0.05, ** = P < 0.01, *** = P < 0.001, unpaired, two-sided t test. (C) TNIP1 KO cells transfected with an empty control vector (EV) do also respond to poly(I:C) treatment by an upregulation of ISG15 and CCL5 similar to KO cells transfected with a TNIP1 expression construct. This indicates that ISG15 and CCL5 abundances are not only regulated by TNIP1. Shown are representative blots of three biological replicates. Bar diagram shows quantification. Error bars represent SEM. * = P < 0.05, ** = P < 0.01, *** = P < 0.001; unpaired, two-sided t test. (D) Blockage of protein translation by cycloheximide (CHX) treatment reduced the time- and poly(I:C)-dependent increase of TNIP1 after 6 h of treatment indicating a regulation on translational level. Source data are available for this figure: SourceData FS4.

Figure S4.
shows RNA-Proteome correlation comparing WT and TNIP1 KO HeLa clones. Table S4 shows chemical reagents used in this study. Table S5 shows antibodies used in this study. Table S6 lists plasmids used in this study.