Autophagy induction in atrophic muscle cells requires ULK1 activation by TRIM32 through unanchored K63-linked polyubiquitin chains


Optimal autophagic activity is crucial to maintain muscle integrity, with either reduced or excessive levels leading to specific myopathies. LGMD2H is a muscle dystrophy caused by mutations in the ubiquitin ligase TRIM32, whose function in muscles remains not fully understood. Here, we show that TRIM32 is required for the induction of muscle autophagy in atrophic conditions using both in vitro and in vivo mouse models. Trim32 inhibition results in a defective autophagic response to muscle atrophy, associated with increased ROS and MfRF1 levels. The pro-autophagic function of TRIM32 relies on its ability to bind the autophagy proteins AMBRA1 and ULK1 and stimulate ULK1 activity via unanchored K63-linked polyubiquitin. LGMD2H-causative mutations impair TRIM32’s ability to bind ULK1 and induce autophagy. Collectively, our study revealed a role for TRIM32 in the regulation of muscle autophagy in response to atrophic stimuli, uncovering a previously unidentified mechanism by which ubiquitin ligases activate autophagy regulators.

INTRODUCTION

Autophagy is a catabolic process that ensures the removal of excess or damaged cellular components in physiological and pathological conditions and provides metabolic supplies when extracellular nutrients are scarce (1, 2). By keeping the intracellular environment free of harmful material, autophagy plays a key protective role in several human diseases (3). Hereditary myopathies are among the best-documented examples of the close relationship between dysregulated autophagy and human disorders (4). Defective autophagy has been observed in collagen VI–related myopathies, Duchenne muscular dystrophy, and Emery-Dreifuss muscular dystrophy, while impaired recognition of autophagy cargos is present in the limb-girdle muscular dystrophy (LGMD) 1D and in myofibrillar myopathies (4, 5). Conversely, excessive autophagy activation characterizes the merosin-deficient congenital muscular dystrophy 1A (6).

Cellular alterations caused by autophagy dysregulation in muscles include the presence of damaged mitochondria and enlarged endoplasmic reticulum, the impaired turnover of sarcomeric proteins, and the increased susceptibility to cell death (5). Restoration of appropriate autophagy levels by pharmacological or dietary approaches ameliorates the myopathic defects in mouse models for collagen VI–related and Duchenne muscular dystrophies, providing preclinical evidence of the relevance of dysfunctional autophagy in these diseases (7–9). These studies have also uncovered the existence of signaling pathways that link autophagy to the proper functioning of muscle cell compartments, such as the extracellular matrix (through collagen VI and laminin-2), cytoskeleton–to–plasma membrane interaction (dystrophin), and nuclear envelope (lamin A/C) (4). The mechanisms by which these signals converge to the autophagy machinery to modulate its activity, however, remain unknown.

Tripartite motif-containing 32 (TRIM32) is a member of the TRIM protein family, a large group of E3 ubiquitin ligases characterized by the presence of a RING finger domain, a B-box domain, a coiled-coil region, and a variable C-terminal region (10). NHL domains characterized the C terminus of TRIM32, which are involved in protein dimerization and substrate recognitions (11). Mutations in the NHL domains of the TRIM32 are causative of LGMD2H and sarcotubular myopathy, which are mild and severe manifestations of the same disorder (12). Both trim32 knock-out (KO) and knock-in mice carrying a disease-associated mutation have confirmed the myopathic phenotype as a consequence of TRIM32 dysfunction (13, 14) and highlighted the presence of neurological defects that may also contribute to the muscular disorder (13). A missense mutation in the B-box domain of TRIM32 causes the Bardet-Biedl syndrome type 11, a disease characterized by obesity, retinal degeneration, genito-urinary tract malformations, and cognitive impairment, but not showing muscle alterations (15). In keeping with this observation, TRIM32 also plays a role in the regulation of processes not directly related to muscle function, such as immunity, neural differentiation, and cancer (16).

How TRIM32 mutations cause muscular dystrophy has not been entirely clarified. A pro-atrophy role of TRIM32 has been initially postulated on the basis of its ability to ubiquitinate actin, tropomyosin, troponins, α-actinin, and desmin (17, 18). TRIM32 also inhibits the pro-survival phosphatidylinositol 3-kinase/Akt pathway through the
degradation of the desmosome component plakoglobin (19). Studies on Trim32 KO mice have shown that TRIM32 is not necessary to trigger muscle atrophy, but it plays a key role in muscle regrowth after atrophy (20). This finding is in agreement with the observation that patients with LGMD2H often manifest loss of motility after prolonged immobilization (21). Defective muscle regeneration was also observed in Trim32 KO mice upon damage induced by cardiotoxin treatment (22).

Muscle regrowth failure after atrophy has been ascribed to the impaired degradation of PIAS4, a SUMO (small ubiquitin-like modifier) ligase that induces premature senescence (20), and of the proliferation inhibitor NDRG2 (23). Whether TRIM32 also plays a protective role in differentiated muscle cells during or after atrophy induction remains to be elucidated.

Recently, several members of the TRIM protein family have been demonstrated to promote autophagy induction by interacting with the upstream regulators ULK1 (Unc-51 like autophagy activating kinase 1) and BECLIN 1 (24–27). In addition, TRIM proteins act as cargo receptors for selective autophagy (27–29).

AMBRA1 (activating molecule in BECN1-regulated autophagy protein 1) is a positive regulator of autophagy that binds and regulates BECLIN 1 and ULK1 activity (30–33) by favoring their non-degradative ubiquitination. In addition, AMBRA1 interacts with the E3 ubiquitin ligases CULLIN4 and CULLIN5 (34) to regulate the temporal dynamics of autophagy response and, with PARKIN and HUWE1, to promote mitophagy (35, 36). Evidence of a role of AMBRA1 in muscle homeostasis have been recently reported (37). Ablation of AMBRA1 in zebrafish leads to a severe myopathy characterized by disorganized myofibers and aberrant mitochondria morphology (37). Abnormal muscle structure organization was also observed in AMBRA1 mutant mouse embryos (37). Here, we show that AMBRA1 interacts with TRIM32 and mediates autophagy induction in muscle cells under atrophic conditions by stimulating ULK1 activity via unanchored K63-linked polyubiquitin.

RESULTS

AMBRA1 interacts with the E3 ubiquitin ligase TRIM32

AMBRA1 was found in association with TRIM32 in a mass spectrometry-based protein interaction screening (34). The binding between AMBRA1 and TRIM32 was confirmed by coimmunoprecipitation in 293 T cells expressing MYC-AMBRA1 and FLAG-TRIM32 proteins (Fig. 1A) and in C2.7 myoblasts at endogenous levels (Fig. 1B). The domains of AMBRA1 and TRIM32 responsible for their interaction were mapped by means of deletion mutants. Coinmunoprecipitation experiments in 293 T cells transfected with vectors encoding the N-terminal, central, or C-terminal region of AMBRA1 showed that TRIM32 preferentially associates with the C-terminal part of AMBRA1 (Fig. 1C). On the other hand, coimmunoprecipitation experiments in TRIM32 KO 293 T cells transfected with TRIM32 mutants encoding the catalytic domain (RING/B-box), the coiled-coil domain, or the NHL repeats showed that the catalytic domain of TRIM32 is responsible for the binding to Ambra1 (Fig. 1D).

TRIM32 is required for the induction of autophagy by atrophic stimuli

The interaction of TRIM32 with AMBRA1 prompted us to analyze the role of TRIM32 in the regulation of autophagy in muscle cells. We performed experiments in a murine myoblast cell line (C2.7 cells), which is able to differentiate into myotubes upon serum withdrawal. At first, we asked whether AMBRA1 and TRIM32 were required for sustaining basal autophagy in undifferentiated and differentiated cells. We measured autophagy flux in cells in which AMBRA1 or TRIM32 expression was down-regulated by using specific lentiviral short hairpin RNAs (shRNA; shAmbra1 and shTrim32). Analysis of LC3-II levels in the presence or absence of lysosome inhibitors revealed that basal autophagy flux is defective when AMBRA1 expression is silenced both in myoblasts and myotubes (fig. S1A). We also observed a partial inhibition of the myosin heavy chain expression, according to the role of basal autophagy in supporting the differentiation of C2 myoblasts (38). In contrast, we observed no significant alterations of autophagy flux in either undifferentiated or differentiated cells in which TRIM32 expression was down-regulated, indicating that TRIM32 is dispensable for sustaining basal autophagy (fig. S1B).

TRIM32 was reported to play an important role in muscle fiber recovery from atrophic conditions (20). Since autophagy is up-regulated during muscle atrophy (5), we decided to investigate the role of TRIM32 in the induction of autophagy in response to atrophic stimuli. To set up the experimental conditions to induce muscle atrophy in vitro, we treated the differentiated C2.7 cells with dexamethasone, a synthetic analog of glucocorticoids, for 24 hours and verified atrophy induction by analyzing the expression level of the atrophic gene MuRF1 (fig. S2A). In parallel, we evaluated autophagy flux after 4 hours of treatment by LC3 immunoblotting analysis, which confirmed that autophagy is induced by dexamethasone (fig. S2B).

Similar to that observed in basal autophagy, shAmbra1 C2.7 cells treated with dexamethasone showed lower LC3-II levels (fig. S2C). Notably, a significant impairment of autophagy induction was also observed when TRIM32 knock-down or KO myotubes were exposed to dexamethasone (Fig. 2A and fig. S2, D and E). Autophagy impairment upon TRIM32 down-regulation was not restricted to C2.7 cells, as a similar defect was also observed in rat L6E9 myoblasts upon dexamethasone treatment (fig. S2F). We confirmed failure of Trim32-silenced C2.7 cells to increase autophagy flux by measuring the lysosomal degradation rate of the autophagy cargo receptor NBR1 (next to BRCA1 gene 1 protein) at a later time point of dexamethasone treatment (fig. S3A), as well as by ultrastructural analysis. In particular, transmission electron microscopy showed that the number of degradative compartments (amphisomes, lysosomes, and autolysosomes) per cell section, which reflects autophagic activity (39), was similar in untreated shControl and shTrim32 C2.7 cells (fig. S3B), corroborating that TRIM32 is not involved in basal autophagy. Atrophy stimulation in control cells led to an increased number of degradative compartments per cell section, confirming autophagy induction under these conditions (Fig. 2B and fig. S3C, top). In contrast, no increase was observed in Trim32-depleted cells exposed to dexamethasone, underlying that TRIM32 is required for autophagy in atrophic myotubes (Fig. 2B and fig. S3C, bottom).

To substantiate the involvement of TRIM32 in the regulation of autophagy in atrophic conditions, we analyzed autophagy flux upon nutrient deprivation as an alternative atrophic stimulus. Also, in this case, a reduced induction of autophagy was observed in Trim32-silenced cells when LC3-II levels were analyzed (fig. S4A). We confirmed defective autophagy by measuring the lysosomal degradation rate of the autophagy cargo receptor p62 in shTrim32-silenced cells at a later time point of nutrient deprivation (fig. S4B).

We then assessed whether TRIM32 is required for an efficient autophagy response to atrophic stimuli in vivo using a mouse model of Trim32 deficiency (22). We treated Trim32 wild-type (WT) and
KO mice with dexamethasone for 8 hours and analyzed LC3, NBR1, and p62 levels in quadriceps muscles by immunoblotting. As shown in Fig. 2C, a significant impairment of autophagy is observed in Trim32 KO mice upon dexamethasone treatment when compared to the WT counterpart. NBR1 and p62 RNA levels were also measured by quantitative polymerase chain reaction (qPCR) to rule out a transcriptional contribution to the observed changes (fig. S5, A and B). Defective autophagy response was confirmed by confocal microscopy as well. Quadriceps muscles from WT mice treated with dexamethasone showed a robust increase of LC3-positive vesicles, which mostly colocalize with the cargo receptor p62 (fig. S5, C and D). Instead, the accumulation of LC3-positive vesicles was reduced...
Fig. 2. Trim32 is required for autophagy induction by atrophic stimuli in muscle cells. (A) shCTR and shTrim32 C2.7 cells differentiated for 3 days were treated with dexamethasone (dexa) for 4 hours (h) or left untreated. One hour before lysis, cells were incubated with the lysosome inhibitors E64d and pepstatin A (E64d/PepA), as indicated. LC3-II and TRIM32 levels were analyzed by immunoblotting (left). GAPDH (glyceraldehyde-3-phosphate dehydrogenase) was included as a loading control. The graph (right) reports means ± SD of LC3-II/GAPDH values from three independent experiments; *P < 0.05. (B) Differentiated shCTR and shTrim32 C2.7 cells were treated with dexamethasone for 4 hours and processed for EM. ER, endoplasmic reticulum; N, nucleus. Yellow asterisks highlight degradative compartments (amphisomes, lysosomes, and autolysosomes). Scale bars, 1 μm. The graph (right) reports means ± SEM of degradative vacuoles per cell per section. Images of untreated cells are reported in fig. S3B. (C) WT (Trim32+/+) and KO (Trim32−/−) mice were injected with dexamethasone (5 mg/kg) or saline solution (0.9% NaCl) as a negative control. Mice were sacrificed 8 hours after injection, and protein extracts were prepared from quadricep muscles. LC3-II, NBR1, and p62 levels were analyzed by immunoblotting (left). Arrows point to TRIM32- and NBR1-specific bands; the number sign (#) indicates nonspecific signals. The graph (right) reports means ± SD of LC3-II/GAPDH, NBR1/GAPDH, and p62/GAPDH values from at least three independent experiments; *P < 0.05.
in Trim32 KO mice with the residual ones, showing diminished colocalization with p62 (fig. S5, E and F).

Autophagy inhibition is known to exacerbate muscle damage in response to atrophic stimuli, resulting in the accumulation of reactive oxygen species (ROS) from dysfunctional mitochondria and an increased induction of atrophic genes (40, 41). Therefore, we assessed whether these alterations were associated with the autophagy impairment observed upon TRIM32 inhibition. TRIM32-silenced myotubes and the relative controls were treated with dexamethasone, and the production of ROS was analyzed after 24 hours (fig. S6A), while the expression of the atrophic gene Murf1 was monitored at 8 and 24 hours (fig. S6, B and C). We also treated control cells with the autophagy inhibitor 3-methyladenosine (3MA), as a positive control of autophagy inhibition (fig. S6D). Concomitantly, we analyzed LC3-II levels at 24 hours after dexamethasone treatment to confirm that autophagy impairment was still present at the time points where ROS and Murf1 levels were measured (fig. S6E). Results showed that both atrophy-related parameters were worsened in TRIM32-silenced cells, similar to those observed in control cells exposed to 3MA. Together, these results point to a role of TRIM32 in the regulation of autophagy in atrophic conditions.

TRIM32 stimulates ULK1 activity in an AMBRA1-dependent manner through unanchored K63-linked polyubiquitin

TRIM proteins were reported to control autophagy by modulating the activity of BECLIN 1 and ULK1 (27). We therefore asked whether TRIM32 could regulate autophagy by interacting with these autophagy regulators. Coimmunoprecipitation experiments performed in 293 T cells transfected with vectors encoding TRIM32, ULK1, and BECLIN 1 showed that TRIM32 efficiently interacts with ULK1 (Fig. 3A). This interaction is specific for ULK1, since the binding with BECLIN 1 was not observed in the same experimental settings (Fig. 3A).

We mapped the domain of TRIM32 responsible for ULK1 interaction by immunoprecipitation of a series of TRIM32 deletion mutants, which showed that TRIM32 binds ULK1 through its catalytic domain (Fig. 3B). On the basis of the ability of AMBRA1 to associate with both ULK1 and TRIM32, we asked whether AMBRA1 plays a role in the interaction between these proteins. Coimmunoprecipitation analysis showed that the association between TRIM32 and ULK1 is strongly affected when AMBRA1 expression is down-regulated (Fig. 3C), indicating that AMBRA1 is required to assemble TRIM32 and ULK1 in a complex.

Since TRIM32 is an E3 ubiquitin ligase, we assessed whether its interaction with ULK1 results in an increase of ULK1 ubiquitination. To this aim, we analyzed degradative and nondegradative ubiquitination of endogenous ULK1 in TRIM32-transfected 293 T cells using K48- and K63-linked polyubiquitin chain antibodies, respectively. As shown in Fig. 4A, TRIM32 triggers K63-linked ubiquitination of ULK1, while K48-linked ubiquitination remains unaltered (fig. S7A). The increase in ULK1 ubiquitination depends on the ligase activity of TRIM32, since a catalytic-inactive mutant is not able to trigger ULK1 ubiquitination (fig. S7B). We also confirmed the ability of TRIM32 to ubiquitinate ULK1 by in vitro assays using either recombinant or immunopurified ULK1 (Fig. 4B and fig. S7C). In keeping with its role in TRIM32–ULK1 complex formation, AMBRA1 down-regulation leads to a significant decrease of TRIM32-mediated ubiquitination of ULK1 (fig. 4C).

Since TRIM32 was reported to produce both substrate-linked and free polyubiquitin chains (23, 42), we analyzed whether TRIM32-induced modification of ULK1 is due to either covalent ubiquitination or

Fig. 3. TRIM32 associates with ULK1 in an AMBRA1-dependent manner. (A) Protein extracts from 293 T cells transfected with vectors encoding HA-TRIM32, FLAG–BECLIN 1, and MYC–ULK1, as indicated, were subjected to immunoprecipitation using an anti-HA antibody. Immunopurified complexes were analyzed by immunoblotting using anti-FLAG, anti-MYC, and anti-HA antibodies. (B) TRIM32 KO 293 T cells were cotransfected with vectors encoding MYC–ULK1 and the following FLAG-TRIM32 constructs: full length, catalytic domain (RING/B-box, amino acids 1 to 133), central region containing the coiled-coil domain (amino acids 134 to 198), and NHL repeats (amino acids 199 to 325). Protein extracts were immunopurified using an anti-FLAG antibody. Immunopurified complexes were analyzed by immunoblotting using anti-FLAG and anti-MYC antibodies. A scheme of the TRIM32 domain architecture is shown; the red bar indicates the ULK1-interacting domain. (C) Control shRNA and shAmbra1 293 T cells were transfected with a vector encoding MYC–ULK1, alone or in combination with FLAG-TRIM32. Cells were lysed, and protein extracts were subjected to immunoprecipitation using an anti-FLAG antibody. Immunopurified complexes were analyzed by immunoblotting using anti-MYC and anti-FLAG antibodies. Total extracts were also probed with an anti-AMBRA1 antibody to verify AMBRA1 silencing.
the binding to unanchored polyubiquitin. To this aim, we subject-
ed ubiquitinated ULK1, immunopurified from either TRIM32-
transfected cells or an in vitro assay, to heat denaturation to disrupt
noncovalent interactions. Immunoblotting analysis showed that
the ubiquitination signal is lost following denaturation [Fig. 4, B and D
(see lanes labeled as Re-IP)], indicating that ULK1 binds nonco-
vallently to K63-linked polyubiquitin chains produced by TRIM32. More-
over, we observed that the interaction with the polyubiquitin chains
is mediated by the C-terminal domain of ULK1, as shown by co-
transfecting TRIM32 with ULK1 deletion mutants (fig. S7D).

Since K63-linked ubiquitination of ULK1 stimulates its kinase ac-
tivity (31), we asked whether TRIM32 is able to promote ULK1 pro-
autophagic function. A phosphorylation analysis of the autophagic
proteins VPS34 and BECLIN 1, two well-characterized targets of ULK1
(43, 44), showed that ULK1 kinase activity is potentiated by TRIM32
(Fig. 4, E and F). In line with these results, we also observed that
overexpression of WT TRIM32, but not of the catalytic mutant C39S,
results in an increased autophagy flux, as revealed by LC3 immuno-
blotting (fig. S7E). Together, these data indicate that TRIM32 can in-
duce autophagy by interacting with ULK1 in an AMBRA1-dependent
manner and promoting its activity through unanchored polyubiquitin.

**TRIM32 is required for ULK1 activation through K63-linked polyubiquitin in atrophic conditions**

The ability of TRIM32 to bind and activate ULK1 through unanchored
polyubiquitin prompted us to analyze the functional relation be-
 tween TRIM32 and ULK1 in the induction of autophagy by atrophic
stimuli. First, by coimmunoprecipitation assays performed in un-
treated and dexamethasone-treated C2.7 cells, we observed that
atrophy induction stimulates the interaction of ULK1 with TRIM32
(Fig. 5A), as well as its association with K63-linked polyubiquitin
(Fig. 5B). Then, we investigated whether TRIM32 is required for the in-
creased association of ULK1 to K63-linked polyubiquitin upon
atrophy stimulation. Down-regulation of TRIM32 expression in
C2.7 cells by RNA interference markedly reduced the amount of K63-
linked polyubiquitin that coimmunoprecipitated with ULK1 upon
dexamethasone treatment (Fig. 5C). Consistently, ULK1 activity was
decreased upon TRIM32 down-regulation, as shown by analyzing the
phosphorylation status of BECLIN 1 and ATG14 (AuTopaHy
related 14), two ULK1 substrates (Fig. 5, D and E). Together, these
results indicate that autophagy impairment in TRIM32-deficient
muscle cells is associated with a defective activation of ULK1 medi-
ad by K63-linked polyubiquitin.

We also confirmed the key role of ULK1 in the response to atrophic
stimuli by inhibiting the expression of NEDD4L, an E3 ubiquitin
ligase that targets ULK1 to proteasome-mediated degradation (45).
As shown in fig. S8 (A and B), NEDD4L silencing in C2.7 myoblasts
resulted in ULK1 protein stabilization, leading to a higher autophagy
induction by nutrient starvation.

**TRIM32 pathogenic mutants are defective for ULK1 binding and autophagy induction**

Specific mutations in TRIM32 are causative of the muscular disease
LGMD2H (12). Since autophagy dysregulation plays an important role
in various muscular dystrophies (4), we asked whether the disease-
associated TRIM32 mutants have an impaired proautophagic activity.
First, TRIM32 mutants were tested for the ability to bind ULK1 in a
coinmunoprecipitation experiment. As shown in Fig. 6A, the bind-
ing of TRIM32 to ULK1 is severely affected by all tested pathogenic
mutations. Since the ULK1-interacting domain of TRIM32 is differ-
ent from the NHL repeats (see Fig. 3B), where the disease-associated
mutations are located, we decided to better elucidate the role of this
domain in the binding to ULK1 by generating a TRIM32 mutant
lacking the entire NHL repeats. When tested in a coimmunoprecip-
ation assay, the ΔNHL TRIM32 mutant also failed to interact with
ULK1 (fig. S8C), indicating that, although not sufficient for the
binding, the NHL repeats need to be present to allow the interaction
of TRIM32 with ULK1 in the context of the entire protein.

We also tested whether pathogenic mutations affect the interac-
tion between TRIM32 and AMBRA1. We observed that AMBRA1 is
still able to interact with all mutants tested, suggesting that pathogenic
mutations specifically impair the interaction with ULK1 (fig. S8D).

To verify whether the defective binding of TRIM32 pathogenic
mutants with ULK1 affects their ability to stimulate ULK1 proauto-
phagic activity, we further characterized two of them: Trim32D487N
and Trim32R394H. First, we analyzed the levels of ULK1-associated
polyubiquitin and of ULK1-mediated BECLIN 1 phosphorylation
upon TRIM32 WT or mutant protein overexpression in 293 T cells.
We found that both pathogenic TRIM32 mutants showed a reduced
ability to promote ULK1 association to K63-linked polyubiquitin
(fig. S8E) and BECLIN 1 phosphorylation on Ser15 (fig. S8F), con-
firming their defective ability to induce ULK1 activity.

Then, we analyzed whether the pathogenic TRIM32 mutants have
an impaired ability to induce autophagy in response to atrophy in-
duction in myoblast cells. To this aim, we complemented Trim32 KO
C2.7 cells with retroviral vectors expressing TRIM32 WT, pathogenic
mutants Trim32D487N and Trim32R394H, or the catalytic mutant
Trim32C398S. We treated cells with dexamethasone and analyzed auto-
phagy flux by measuring LC3-II (Fig. 6B and fig. S9, A and B) and
NBR1 (fig. S9C) levels in the presence or absence of lysosome inhibi-
tors. Immunoblot analysis showed that TRIM32 mutants have an
impaired autophagic induction. Also, in this case, the inability of
TRIM32 mutants to trigger autophagy was associated with a defective
activity of ULK1, as shown by analyzing phosphorylation of BECLIN
1 and ATG14 on Ser15 and Ser29, respectively (fig. S10, A and B).

Last, we evaluated whether cells obtained from a patient with
LGMD2H show a defective autophagy response to dexamethasone-
duced atrophy. To this aim, fibroblasts from a healthy donor (HD) and
a patient with LGMD2H, carrying a complete gene deletion in
one allele and a nonsense c.1837 C>T (R613X) mutation in the NHL
domain of the other allele (46), were transfected in myoblasts
by MYOD ectopic expression (fig. S10C). We treated cells with dexa-
methasone and analyzed LC3-II levels in the presence or absence of
lysosome inhibitors to measure autophagic flux. Notably, autophagy
induction by dexamethasone treatment was impaired in myotube
cells carrying a TRIM32 mutation (Fig. 6C and fig. S10D). More-
over, in line with that observed in Trim32-silenced cells, autophagy
induction was exacerbated in LGMD2H patient cells, as indicated
by analyzing MuRF1 levels (fig. S10E). Together, these results show
that disease-associated mutations affect the proautophagic function
of TRIM32 (fig. S10F).

**DISCUSSION**

Autophagy is critical for muscle adaptation to sublethal cellular stress
(47). Exercise increases autophagy levels to meet energetic demands
and to eliminate dysfunctional cell constituents, such as mitochondria
and sarcomere proteins, which may accumulate during contraction
Fig. 4. TRIM32 stimulates ULK1 activity in an AMBRA1-dependent manner through unanchored K63-linked polyubiquitin. (A) Protein extracts from FLAG-TRIM32–transfected 293 T cells were subjected to immunoprecipitation using an anti-ULK1 antibody. Immunopurified complexes were analyzed by immunoblotting to detect ULK1 and K63-linked ubiquitin. Total extracts were also probed with an anti-FLAG antibody to verify TRIM32 transfection. (B) In vitro ULK1 ubiquitination assay using immunopurified Flag TRIM32 and recombinant ULK1, HA ubiquitin, E1 Ub–activating enzyme (UBE1), and E2 Ub–conjugating enzyme (UBE2N). Reactions were also performed in the absence of UBE1 to verify ubiquitination reaction specificity. At the end of the reaction, ULK1 was immunopurified and ubiquitination evaluated by immunoblotting using an anti-HA antibody (lanes 1 and 2). To verify whether ubiquitin was covalently linked, immunopurified ULK1 was boiled in 1% SDS after the ubiquitination reaction, reimmunoprecipitated (Re-IP), and analyzed by immunoblotting using anti–HA and anti–ULK1 antibodies (lanes 3 and 4). (C) shCtr or shAmbra1 293 T cells were transfected with a vector encoding FLAG-TRIM32 or an empty vector (−). Protein extracts were immunoprecipitated using an anti-ULK1 antibody. Immunopurified complexes were analyzed by immunoblotting to detect ULK1 and K63-linked ubiquitin. Total extracts were also probed with FLAG and AMBRA1 antibodies to verify TRIM32 transfection and AMBRA1 silencing, respectively. (D) 293 T cells were transfected with vectors encoding HA-tagged ubiquitin, FLAG-ULK1, and TRIM32, as indicated. Protein extracts were subjected to immunoprecipitation using an anti-FLAG antibody. Immunopurified complexes were analyzed by immunoblotting to detect K63-linked ubiquitin and ULK1 using specific antibodies. TRIM32 expression was also analyzed in total extracts. To verify whether ubiquitin was covalently linked, immunopurified FLAG-ULK1 was boiled in 1% SDS, reimmunoprecipitated, and analyzed by immunoblotting using anti–K63-linked ubiquitin and anti-ULK1 antibodies. (E) 293 T cells were transfected with vectors encoding FLAG-VPS34, alone or in combination with HA-TRIM32, TRIM32, VPS34, and phospho-VSP34 [pVPS34; Ser^{49} (S249)] were analyzed by immunoblotting. GAPDH was included as a loading control. (F) 293 T cells were transfected with vectors encoding FLAG–BECLIN 1, alone or in combination with HA-TRIM32. The levels of TRIM32, BECLIN 1, and phospho–BECLIN 1 [pBECLIN 1; Ser^{15} (S15)] were analyzed by immunoblotting. GAPDH was included as a loading control.
Autophagy is also induced during prolonged inactivity or other atrophic stimuli, mainly to ensure the removal of surplus of organelles, such as mitochondria and sarcoplasmic reticulum, during muscle fiber remodeling. Although it contributes to muscle dismantling, autophagy is considered a protective mechanism by preventing the accumulation of harmful signals, such as ROS, generated by inoperative compartments.

How autophagy is regulated in muscle cells has been extensively investigated at a transcriptional level, highlighting the important role of FoxO, glucocorticoid receptors, nuclear factor-κB (NF-κB), ...
Fig. 6. TRIM32 pathological mutants are unable to interact with ULK1 and promote autophagy. (A) 293T cells were transfected with a vector encoding for MYC-ULK1, alone or in combination with HA-TRIM32 WT, or the indicated pathogenic TRIM32 mutants. Protein extracts were subjected to immunoprecipitation using an anti-HA antibody. Immunopurified complexes were analyzed by immunoblotting to detect the presence of TRIM32 and ULK1. (B) TRIM32 KO C2.7 cells were infected with retroviruses encoding WT FLAG-TRIM32 or the indicated pathogenic TRIM32 mutants. EMPTY, TRIM32 KO C2.7 cells infected with noncoding retroviruses. Upon 3 days in differentiation medium, cells were treated with dexamethasone for 4 hours and incubated with the lysosome inhibitors E64d and pepstatin A for 1 hour before lysis, as indicated. LC3-II and TRIM32 levels were analyzed by immunoblotting using specific antibodies. GAPDH was included as a loading control. (C) Fibroblasts from a WT donor and a patient with LGMD2H were transdifferentiated to myoblasts by MYOD (myogenic differentiation 1) ectopic expression. Cells differentiated for 3 days were treated, or not, with dexamethasone for 2 and 4 hours. One hour before lysis, cells were incubated with the lysosome inhibitors E64d and pepstatin A, as indicated. LC3-II and myosin heavy chain (MYOSIN HC) levels were analyzed by immunoblotting (bottom panel). GAPDH was included as a loading control.
and SMAD transcription factors (51). In contrast, the mechanisms by which stresses related to muscle inactivity are transduced to the autophagy machinery for the activation of the process remain less characterized.

Here, we report that the E3 ubiquitin ligase TRIM32 is required for the induction of autophagy in muscle cells by atrophic stimuli using both in vitro and in vivo models. The proautophagic activity of TRIM32 resides in its ability to bind to AMBRA1 and ULK1 and to synthesize K63-linked polyubiquitin chains that bind noncovalently to ULK1 to promote its activity (fig. S10F).

Unanchored polyubiquitin chains are known to play a key role in the regulation of kinase activity (52). Most evidence come from studies on innate immune pathways, showing that transforming growth factor β–activated kinase 1 (TAK1), inhibitor of nuclear factor κB kinase ε (IKKe), and NF-kappa-B essential modulator (NEMO) can be activated by the noncovalent interaction with polyubiquitin chains, which result in changes of the structural conformation and/or the interaction partners of these kinases (53–55). In this context, TRIM32 E3 ligases are emerging as key players (56), as shown for TRIM5, TRIM6, TRIM25, and TRIM32, with the latter adopting this strategy to activate NEMO in response to cytosolic double-stranded DNA (42). Here, we show that the unanchored polyubiquitin chains are also involved in the regulation of the autophagic process. TRIM32 synthesized K63-linked ubiquitin chains associated with ULK1 via its C-terminal domain, resulting in an augmented phosphorylation of its substrates BECLIN1 and VPS34. In contrast to that reported for other TRIMs, we did not observe an increased interaction between ULK1 and BECLIN1 complexes upon TRIM32 activation, suggesting that the binding with the polyubiquitin chains stimulates ULK1 activity rather than favoring protein complex interactions.

Our data also highlighted the central role of AMBRA1 in conveying multiple ubiquitin signals to the autophagy machinery for its activation in specific cellular contexts. In atrophic muscle cells, AMBRA1 acts as an essential cofactor, being required for both the interaction and the ubiquitination of ULK1 by TRIM32. AMBRA1 was previously shown to regulate ULK1 activity by mediating its covalent ubiquitination by the E3 ligase TNF receptor associated factor 6 (TRAF6) (31). Although the function of TRAF6 and TRIM32 may appear redundant in autophagy regulation, these proteins play opposite roles in the activation of the atrophy program (20, 57, 58). TRAF6 inactivation significantly impairs the induction of MuRF expression under atrophic conditions due to its role in the activation of the NF-κB pathway (57). Conversely, TRIM32 down-regulation leads to higher MuRF1 induction in myoblasts treated with dexamethasone, consistent with that observed when autophagy genes are deleted (40).

An important role of TRIM32 in muscle autophagy has been attributed to its ability to inhibit premature senescence of satellite cells responsible for muscle regrowth in vivo (20). Our results showing a defective muscle autophagy in atrophic conditions suggest that, in addition to regulating stem cells, TRIM32 may also contribute to the preservation of the function of differentiated muscle cells by reducing ROS accumulation and MuRF1 expression.

Evidence that the alteration of TRIM32-dependent autophagy may have an important implication in LGDM2H is underlined by two sets of data. First, we detected defective autophagy in myoblasts derived from fibroblasts of a patient with LGDM2H. Second, we found that disease-associated mutants of TRIM32 are unable to interact with ULK1 (their binding to AMBRA1, however, is unaffected) and to induce autophagy in muscle cells upon dexamethasone treatment, which is accompanied with accumulation of the autophagy cargo receptors NBR1 and p62, higher induction of MuRF1, and increased ROS production (fig. S10F). Further studies are required to clarify the molecular basis of this alteration, since the NHL repeats, where pathological mutations are located, do not mediate the interaction of TRIM32 with ULK1, which occurs through the RING/B-box region. We found that a TRIM32 mutant lacking the NHL repeats is also unable to bind ULK1, suggesting that this domain is essential to making the RING/B-box domain accessible to the targets in the presence of the coiled-coil region.

In conclusion, we have identified TRIM32 as an E3 ubiquitin ligase that regulates ULK1 activity in muscles under atrophy conditions, highlighting a novel autophagy pathway with potential relevance in human disorders. In light of these results, it is predictable that the proautophagic activity of TRIM32 could be relevant in other TRIM32-regulated processes, such as neuronal development, tumorigenesis, and immune response (16), in which its partner AMBRA1 is also playing important roles (30, 59, 60).

MATERIALS AND METHODS

Cell culture

293 T cells (American Type Culture Collection) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) (Sigma-Aldrich, D6546), supplemented with 10% fetal bovine serum (Gibco, 10270), 2 mM l-glutamine, and 1% penicillin/streptomycin solution (Sigma-Aldrich, G7513; P0781) at 37°C under 5% CO2. Murine myoblast C2.7 cells (61), rat myoblast L6E9 cells (62), and human fibroblast–derived myoblast cells from an HD and a patient with LGMD2H were cultured in DMEM, supplemented with 20% fetal bovine serum, 2 mM l-glutamine, and 1% penicillin/streptomycin solution at 37°C under 5% CO2. Myoblast differentiation was induced by culturing cells in DMEM supplemented with 2% horse serum (Life Technologies, 16050122), 2 mM l-glutamine, and 1% penicillin/streptomycin for an indicated amount of time. Fibroblasts from an HD and a patient with LGMD2H were provided by the Telethon Network of Genetic Biobanks project no. GTB12001. To prepare fibroblasts, a patient’s skin biopsy was obtained after informed consent and approval of the Ethics Committee of the University of Ferrara. No cell lines used in this study were found in the database of commonly misidentified cell lines maintained by the International Cell Line Authentication Committee and National Center for Biotechnology Information biosample. Cells were screened for mycoplasma contamination by PCR (ABMgood, G238).

To evaluate autophagy, C2.7 cells and HD/LGMD2H-differentiated myoblasts were treated with 400 μM dexamethasone (Calbiochem, 265005) in the presence or absence of E64d/pepstatin A (5 μg/ml) (Santa Cruz Biotechnology, sc-201280A and sc-45036) or 5 nM Bafilomycin A1 (Sigma-Aldrich, B1793) or incubated with nutrient-deprived medium Earle’s balanced salt solution (Sigma-Aldrich, E2888) for the indicated time, according to guidelines. The autophagy inhibitor 3MA (Sigma-Aldrich, M9281) was used at 5 mM.

Myoblast transdifferentiation

Myoblast preparation was obtained by transducing human fibroblasts from healthy or Trim32 mutant patients with the lentivirus cytomegalovirus (CMV) MyoD-ER(T) (Addgene) with a multiplicity of infection of 20 and treated with 1 μM 4OH-tamoxifen (Sigma-Aldrich, H7904)
to activate MyoD-ER(T). The transdifferentiation was confirmed by PCR analysis of MyoD and myogenin expression. Autophagy analysis was performed in the absence of 4OH-tamoxifen.

**Animal studies**

Trim32 KO mice were previously described (22). TRIM32 KO and WT mice were maintained and treated according to approved protocols and in accordance with the institutional and national guidelines and regulations (IP00001489) approved by the Oregon Health and Science University. To analyze the autophagy response to dexamethasone, 3-month-old mice (n = 4 for each genotype) were injected subcutaneously with dexamethasone sodium salt (5 mg/kg; Sigma-Aldrich, D-1756) and prepared in a saline solution (vehicle, 0.9% NaCl). The control group received an injection of saline solution (vehicle, 0.9% NaCl). Eight hours after injection, mice were sacrificed, and quadriceps muscles were harvested and deep-frozen in liquid nitrogen before storing at −80°C for subsequent protein extraction. For immunoblotting analysis, frozen muscles were crushed into a fine powder with a hammer, resuspended in the extraction buffer (Coimmunoprecipitation Kit, Thermo Fisher Scientific, 14321D), supplemented with protease, phosphatase, and deubiquitinase inhibitors as described above, homogenized using a Dounce homogenizer, incubated at 4°C for 30 min, and centrifuged at 13,000 rpm for 10 min to remove debris. Same amounts of total protein (30 μg per well) were loaded on SDS–polyacrylamide gel electrophoresis (PAGE) for immunoblotting analysis.

**Transfection and viral infection**

293 T cells were transiently transfected with expression vectors using the calcium phosphate method, as previously described (32). For retroviral production, packaging cells [293 gp/bsr (gag-pol/blaC10)–targeting gene] were cotransfected with 15 μg of retroviral vectors and 5 μg of pCMV-VSV-G using the calcium phosphate method. For lentiviral production, 293 T cells were cotransfected with 10 μg of lentiviral vectors, 2.5 μg of pCMV-VSV-G, and 7.5 μg of pspAX2 plasmid by using the calcium phosphate method. After 48 hours, the supernatant containing the retroviral or lentiviral particles was recovered, ultracentrifuged at 19,800 rpm on an SW28 rotor for 2 hours, and resuspended in phosphate-buffered saline (PBS) (500 μl for 20 ml of supernatant). Cells were infected with 80 μl of viral suspension in a medium supplemented with polybrene (4 μg/ml) for 8 hours. Two consecutive rounds of infections were performed to improve efficiency.

To establish Trim32 KO cells for mutant complementation assays, C2.7 cells were transiently transfected with CRISPR-Cas9 vectors by Lipofectamine LTX and Plus Reagent (Invitrogen, 15338-100), as indicated by the supplier. Transduced cells were incubated with puromycin dihydrochloride (2.5 μg/ml; Santa Cruz Biotechnology, sc-108071) for only 48 hours to select transfected cells, where inactivation of Trim32 CRISPR-Cas9 has occurred, and then cultured in the absence of puromycin to avoid selecting cells with a stably integrated CRISPR-Cas9 vector.

**Plasmids**

pCDNA3 hemagglutinin (HA)–Trim32 has been previously described (10). The HA-Trim32 mutants that correspond to human complementary DNA (cDNA) pathogenic Trim32 mutations were engineered with the QuickChange Site-Directed Mutagenesis Kit (StrateGene).

All retroviral constructs were based on a modified version of a pLPCX vector (Clontech) (32). pLPCX plasmids encoding FLAG- and MYC-WT AMBRA1, MYC-AMBRA1 mutants (amino acids 1 to 532, 533 to 751, and 761 to 1269), and FLAG–BECLIN 1 were previously described (30–32). pLPCX FLAG–TRIM32 and pLPCX TRIM32 were obtained by subcloning from pCDNA3 HA–TRIM32. pLPCX FLAG–TRIM32 deletion mutant RING/B-box (amino acids 1 to 136), coiled-coil domain (amino acids 136 to 326), NHL repeats (amino acids 327 to 653), and ΔNHL (amino acids 1 to 325) were created using the appropriate oligonucleotides and amplification, followed by in-frame insertion into the pLPCX FLAG vector. pCDNA3 HA–Trim32 C39S was generated by site-specific mutagenesis; pLPCX FLAG–TRIM32 C39S was obtained by subcloning from pCDNA3 HA–TRIM32 C39S. pLPCX FLAG–ULK1 and pLPCX MYC–ULK1 was obtained by subcloning from pCDNA3 Myc-tag ULK1 (31). pLPCX FLAG–ULK1 deletion mutants (amino acids 1 to 828 and 829 to 1050) were created using appropriate oligonucleotides and amplification, followed by in-frame insertion into the pLPCX FLAG vector. pRK5 HA–UBIQUITIN was obtained from Addgene (#17608). A CRISPR-Cas9 All-in-One lentiviral vector specific for human TRIM32 and murine TRIM32 [hTRIM32 single-guide RNA (sgRNA), K2465605; mTRIM32 sgRNA, K3452705] and control sgRNA (scramble sgRNA, K010) were purchased from ABMgood.

For stable murine Trim32 mRNA interference, two lentiviral Trim32 mRNA–targeting pLKO.1 plasmids were used (TRCN0000040831 and TRCN0000048032; Sigma-Aldrich). For stable murine Ambra1 mRNA interference, two lentiviral Ambra1 mRNA–targeting pLKO.1 plasmids were used (TRCN0000189905 and TRCN0000189940; Sigma-Aldrich). For stable human AMBRA1 mRNA interference, a lentiviral AMBRA1 mRNA–targeting plasmid was used (TRCN0000168652; Sigma-Aldrich). For stable murine Nedd4L mRNA interference, a lentiviral Nedd4L mRNA–targeting plasmid was used (TRCN0000086870; Sigma-Aldrich). A pLKO.1 containing a nonmammalian shRNA was used as a negative control (Sigma-Aldrich).

**Antibodies**

The primary antibodies used in this study were rabbit anti–HA antibody (Sigma-Aldrich, H6908), rabbit anti–FLAG antibody (Sigma-Aldrich, F7425), mouse anti–MYC antibody [Santa Cruz Biotechnology, sc-40 (9E10)], rabbit anti–MYC antibody (Millipore, 06-549), rabbit anti–BECLIN 1 [Santa Cruz Biotechnology, sc-11427 (H-300)], goat anti–BECLIN 1 [Santa Cruz Biotechnology, sc-10086 (D-18)], rabbit phospho–BECLIN 1 (Ser15) [Cell Signaling Technology, 138255 and 84966 (D4B7R)], rabbit anti–ATG14 [Cell Signaling Technology, 96752 (D1A1)], rabbit phospho–ATG14 [Cell Signaling Technology, 13155 (S29)], rabbit anti–NBR1 (Novus Biologicals, NBP1-71703), LC3 (Cell Signaling Technology, 27755), mouse anti–human AMBRA1 [Santa Cruz Biotechnology, sc-398204 (G6)], rabbit anti–human AMBRA1 (Novus Biologicals, 2619002), rabbit anti–mouse Ambra1 (Millipore, ABC131), mouse anti–multi ubiquitin (MBL International, ST1200), rabbit anti–K63-linked ubiquitin (Millipore, 05-1308), rabbit anti–K48-linked ubiquitin (Millipore, 05-1307), rabbit anti–ULK1 [Santa Cruz Biotechnology, sc-33183 (H240)], rabbit anti–ULK1 [Cell Signaling Technology, 80545 (D8H5)], rabbit anti–TRIM32 (Thermo Fisher Scientific, PA5-22316), rabbit anti–phospho–VP534 [Ser249] (Cell Signaling Technology, 138575), rabbit anti–VP534 (Life Technologies, 382100), mouse
fluoride (Sigma-Aldrich, S-7920), 0.5 mM sodium orthovanadate was complemented with protease and phosphatase inhibitors [Pro -
secondary antibodies [anti-goat 705-036-147, anti-rabbit 711-036-
Detection was achieved using horseradish peroxidase–conjugated
or polyvinylidene difluoride (Millipore, IPVH20200) membranes. 
Technologies, 4 to 12% EA0378BOX and 3 to 8% NW04120BOX)
and electroblotted onto nitrocellulose (Whatman Amersham, 10600041) and 2 mM MgCl2 (Sigma-Aldrich, M8266]) was used to analyze ubiquitination levels of immu-
noprecipitated proteins and for immunoblotting assays. Lysis buffer was complemented with protease and phosphatase inhibitors [Pro-
tease Inhibitor Cocktail plus (Sigma-Aldrich, P8340), 5 mM sodium 
fluoride (Sigma-Aldrich, S-7920), 0.5 mM sodium orthovanadate (Sigma-Aldrich, S6508), 1 mM sodium molybdate (Sigma-Aldrich, S-6646), 50 mM 2-chloroacetamide (Sigma-Aldrich, C0267), 2 mM 1,10-phenanthroline monohydrate (Sigma-Aldrich, 320056), and 0.5 mM phenylmethylsulfonyl fluoride (Sigma-Aldrich, P7626). Co-
immunoprecipitation was performed with 1 mg of protein extracts from transfected cells or 3 mg in the case of endogenous proteins assays. For endogenous protein immunoprecipitation, protein extracts were incubated overnight with 2 μg of antibody, and immu-
nocomplexes were recovered using 25 μl of Protein A Sepharose (GE Healthcare, GE 17-1279-01). For immunoprecipitation of over-
expressed tagged proteins, protein extracts were incubated with 25 μl of anti-FLAG, anti-HA, or anti-MYC antibodies conjugated to agarose 
beads (Sigma-Aldrich: A22220, A2095, and A7470, respectively) for 
2 hours. To test the covalent binding of polyubiquitin chains to ULK1, 
the second-round immunoprecipitation assay, the immunoprecipi-
tates were denatured by boiling for 5 min at 95°C in the lysis buffer 
containing 1% SDS. The eluates were then diluted 1:10 with lysis buffer 
and reimmunoprecipitated with the anti-FLAG antibody conjugated 
to agarose beads for 2 hours. To test the covalent binding of polyubiquitin chains to ULK1, the first-round immunoprecipitation assay, cell extracts were pre-
apared as described above, and then, lysates were incubated with the anti-FLAG antibody conjugated to agarose beads for 2 hours. 
Immunoprecipitates were washed three times with RIPA buffer. Before the second-round immunoprecipitation assay, the immunoprecipi-
tates were denatured by boiling for 5 min at 95°C in the lysis buffer containing 1% SDS. The eluates were then diluted 1:10 with lysis buffer 
and reimmunoprecipitated with the anti-FLAG antibody conjugated to 
agarose beads for 2 hours.
Ubiquitination assay
293 T cells were independently transfected with plasmids encoding 
MYC-ULK1 or FLAG-TRIM32. Forty-eight hours later, cells were 
lysed in RIPA buffer for ULK1 purification and tris buffer for TRIM32 
immunoprecipitation, both containing protease inhibitors, as pre-
vioously indicated. Lysates were cleared by centrifugation and sub-
jected to immunoprecipitation for 2 hours using agarose-coupled 
antibodies against MYC or FLAG tags. TRIM32 protein was eluted 
for 1 hour by means of FLAG peptide (Sigma-Aldrich, F3290) at 
400 ng/μl in the 1X Ubiquitin Conjugation Reaction Buffer (Boston 
Biochem, B-70). The ubiquitination assays were performed in 100 μl 
of reaction volume, combining 20 μl of immunopurified MYC-ULK1 
bound to agarose-beaded Myc antibody, 20 μl of eluted TRIM32, 
and the following recombinant components: 100 nM E1 Ub-
activating enzyme (Ube1; Boston Biochem, E-305), 1 μM E2 Ub-
conjugating enzyme (Ube2N; Boston Biochem, E2-664), and 50 μM 
HA-ubiquitin (Boston Biochem, U-110) resuspended in Ubiquitin 
Reconstitution Buffer (Boston Biochem, B-90). In some in vitro ubiquitination experiments, recombinant ULK1 protein (Signal-
Chem, U01-11G) was used instead of the immunopurified one. 
The reaction was performed in 1X Ubiquitin Conjugation Reaction 
Buffer, supplemented with Mg2+-adenosine triphosphate (Boston 
Biochem, B-20) at 2 mM and incubated at 30°C for 2 hours in a rock-
ring platform. When indicated, a denaturation step was added as pre-
viously described, followed by a reimmunoprecipitation of ULK1 
using the rabbit anti-ULK1 H240 antibody (Santa Cruz Biotech-
nology, sc-33183). The incorporation of ubiquitin was analyzed by 
immunoblotting using rabbit anti-HA antibody (Sigma-Aldrich, 
H6908) to detect HA-ubiquitin.
Real-time PCR
Real-time PCR was performed, as previously described (63). Briefly, 
RNA was extracted by using a TRIzol reagent (Invitrogen, 15596-018). 
cDNA synthesis was generated using a reverse transcription kit 
(Promega, A3500), according to the manufacturer’s recommenda-
tions. qPCR reactions were performed with the Rotor-Gene 6000 
(Corbett Research Ltd.) thermocycler. The Maxima SYBR Green/
ROX qPCR Master Mix (Thermo Fisher Scientific, K0253) was used to produce fluorescently labeled PCR products during repetitive cy-
cling of the amplification reaction, and the melting curve protocol 
was used to check for probe specificity, as described previously (30). 
The following primer sets for all amplicons were designed using the 
Primer-Express 1.0 Software System (Roche): mouse MRF1 forward 
(5′-CCAAGGAAAGAGCAGTATG-3′) and reverse (5′-GCAGG-
CTCCTCTGGTATTTG-3′), mouse p62 forward (5′-TGAAACAT-
GGACACTTTGCTGGC-3′) and reverse (5′-AATTTGGACCT-
TCTGTTGAGGACA-3′), mouse NBR1 forward (5′-GAGATT-
AGAGGGAGGAGATT-3′) and reverse (5′-CTTCAGAGGAGG-
CAGAAGAC-3′), mouse GAPDH forward (5′-TTTACAGGCG-
CACTCAAG-3′) and reverse (5′-CCAGTAGACTCCACAGA-3′), 
human MYOD forward (5′-CACAGCGAGCAGTATG-3′) and 
MYOD reverse (5′-GTGCTCTTGGGTTTCAG-3′), human 
MYOG forward (5′-GGTGTAGTGAGGTTGTAAG-3′) and 
MYOG reverse (5′-GCCATCTACCTCCTGCTTAC-3′) and reverse 
(5′-CCGTTGACTCCGACCTTCAC-3′).
Flow cytometry
Staining for mitochondria was performed by incubating differenti-
ated C2.7 cells with 5 μM CellROX Deep Red Reagent (Thermo 
Fisher Scientific, C10422), following the manufacturer’s protocols, 
and directly analyzed without fixing. Cell analysis was performed 
using FACScan (Becton-Dickinson).
Muscle tissue analysis by fluorescent microscopy
To analyze the autophagy response to dexamethasone, 6-month-old mice (n = 4 for each genotype) were injected subcutaneously with dexamethasone sodium salt (5 mg/kg; Sigma-Aldrich, D-1756) and prepared in a saline solution (vehicle, 0.9% NaCl). The control group received an injection of saline solution (vehicle, 0.9% NaCl). Twenty-four hours after injection, mice were sacrificed, and quadriceps were fixed with 4% formaldehyde for 24 hours at room temperature, and after dehydration with a series of alcohol-xylene dilutions, the tissue was embedded in paraffin and cut into 7-μm sections. Sections were then dewaxed and boiled for 8 min in preheated 10 mM citric acid retrieval buffer (pH 6.0). Sections were subsequently blocked in PBS containing 1% goat serum and 0.4% Triton X-100 for 1 hour before the primary antibody incubation in PBS containing 0.1% Tween 20 overnight at 4°C. Nuclei were stained with Hoechst 33342 (Sigma-Aldrich). The incubation with the secondary antibody in PBS containing 0.1% Tween at room temperature for 1 hour. Fluorescent microscopy was performed at room temperature using the DeltaVision RT fluorescence microscope (Applied Precision, Issaquah, WA) equipped with a CoolSNAP HQ camera (Photometrix, Kew, Australia). Images were generated by collecting stacks of images with focal planes 0.30 μm apart and subsequently deconvolved using the SoftWoRx software (Applied Precision). The following antibodies were used: rabbit anti-LC3 (PM036; MBL International, Woburn, MA), guinea pig anti-p62 (GP62-C; Progen, Darra, Australia), Alexa Fluor 568–conjugated goat anti–guinea pig immunoglobulin G (IgG; Thermo Fisher Scientific, A-11075), Alexa Fluor 488–conjugated goat anti-rabbit IgG (H + L) (Thermo Fisher Scientific, A27034).

Electron microscopy
Differentiated C2.7 cells carrying control or TRIM32-targeting shRNA were incubated with 400 μM dexamethasone for 4 hours. Cells were fixed before and after dexamethasone treatment by direct addition of 5% glutaraldehyde (Merck Millipore, 1042390250) and 4% paraformaldehyde (Sigma-Aldrich, 441244) in 0.1 M cacodylate buffer (pH 7.4; Sigma-Aldrich, 20840-100G-F) to the culture medium. After a 20-min incubation at room temperature, the fixation solution was replaced by 2.5% glutaraldehyde and 2% paraformaldehyde in 0.1 M cacodylate buffer (pH 7.4), and fixation was prolonged overnight. Cells were then embedded in EPON resin [for 25 ml, 12 g of glycic ether (SERVA, 21045.02), 8 g of 2-dodecensuccinic anhydride (SERVA, 20755.02), 5 g of methylindic anhydride (SERVA, 29452.02), and 560 μl of N-benzylidimethylamine (Electron Microscopy Sciences, 11400-25)]. Subsequently, 55-nm sections were cut and stained with uranyl acetate and lead citrate. Cell sections were examined using an 80-kV CM100 transmission electron microscope (Phillips). Three different grids with sections obtained from the same preparations were statistically evaluated. For every grid, the average number of degradative compartments (amphisomes, autolysosomes, and lysosomes) per cell section was determined by counting 25 randomly selected cell profiles.

Statistical analysis
Statistical analysis of electron microscopy data was analyzed using the Mann-Whitney test (independent samples and two-sided; GraphPad). Statistical analysis of immunoblotting, PCR, and fluorescence-activated cell sorting (FACS) data were performed using unpaired, two-tailed Student’s t test (Excel software). Values are shown as means ± SD of at least three independent experiments. P values <0.05 were marked by an asterisk. Densitometric analysis of immunoblots was performed using the Adobe Photoshop software. The control ratio was arbitrarily defined as 1.00. Normal distribution was assumed on the basis of the appearance of the data, since n < 5. No statistical method was used to predetermine sample size for animal studies. The animal experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment. No exclusion criteria were applied to exclude samples or animals from analysis.

SUPPLEMENTARY MATERIALS
Supplemental material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/eaau8857/DC1
Fig. S1. AMBRA1, but not TRIM32, is required for basal autophagy in myoblast cells.
Fig. S2. Role of TRIM32 and AMBRA1 in autophagy induction by atrophic stimuli in muscle cells.
Fig. S3. Analysis of autophagy defects in TRIM32 silenced myoblasts.
Fig. S4. Role of Trim32 in autophagy induction by nutrient starvation in muscle cells.
Fig. S5. Analysis of autophagy defects in Trim32−/− mice.
Fig. S6. Analysis of MuRF1 and ROS levels in dexamethasone-treated muscle cells.
Fig. S7. Characterization of the proautophagic properties of TRIM32.
Fig. S8. Regulation of ULK1 activity by NEDD4L and TRIM32.
Fig. S9. Characterization of autophagy properties of C2.7 cells expressing TRIM32 pathological mutants.
Fig. S10. Defective autophagy induction in human and murine myoblast cells expressing TRIM32 pathological mutants.

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Autophagy induction in atrophic muscle cells requires ULK1 activation by TRIM32 through unanchored K63-linked polyubiquitin chains


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