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ATG5 provides host protection acting as a switch in the atg8ylation cascade between autophagy and secretion

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Summary

ATG5 is a part of the E3 ligase directing lipidation of ATG8 proteins, a process central to membrane atg8ylation and canonical autophagy. Loss of Atg5 in myeloid cells causes early mortality in murine models of tuberculosis. This in vivo phenotype is specific to ATG5. Here we show using human cell lines that absence of ATG5 but not of other ATGs directing canonical autophagy promotes lysosomal exocytosis and secretion of extracellular vesicles, and, in murine Atg5^{fl/fl} LysM-Cre neutrophils, their excessive degranulation. This is due to lysosomal disrepair in ATG5 knockout cells and the sequestration by an alternative conjugation complex, ATG12-ATG3, of ESCRT protein ALIX which acts in membrane repair and exosome secretion. These findings reveal a previously undescribed function of ATG5 in its host-protective role in murine experimental models of tuberculosis and emphasize the significance of the branching aspects of the atg8ylation conjugation cascade beyond the canonical autophagy.

eTOC blurb

Competing interests

The authors declare no competing interests.

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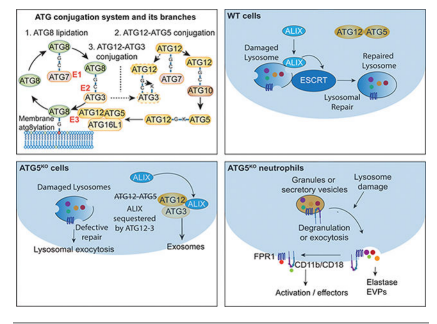
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Wang et al., describe properties of a core autophagy protein ATG5 that fall outside of its canonical autophagy functions. Loss of ATG5 dysregulates lysosomal repair, increases exocytosis and exosome release, and in neutrophils causes hyperactivation and degranulation, partly explaining the function of ATG5 in mouse models of tuberculosis

Graphical Abstract



Introduction

Autophagy in mammalian cells is a complex set of homeostatic processes ¹, including canonical degradative ^{2,3} and noncanonical processes ^{4,5}. During ATG-dependent canonical autophagy, membranes from the constitutive secretory pathway ^{6–16} contribute to the formation of phagophores through fusion with endosomal organelles ^{17,18}. Nascent phagophores enlarge via lipid transfer and close into double membrane autophagosomes ². At the end-stage, canonical autophagosomes fuse with lysosomes to form autolysosomes ¹⁹.

A hallmark of canonical ^{6,20–23} and noncanonical autophagy ^{4,5} is lipidation of mammalian ATG8 proteins (mATG8s), mirroring lipidation of Atg8 in yeast ^{24,25}. Whereas one of the mATG8s, LC3B, is commonly used to identify autophagosomes ²⁶, it is also present on single membranes other than autophagosomes ^{4,27–36}. Among the functions of mATG8s are the enhancement of cargo sequestration into autophagosomes ³⁷, membrane remodeling ³⁸ perturbation ³⁹ and expansion ^{40,41} during autophagosome biogenesis, autophagosome-lysosome fusion ¹⁹, and more general membrane stress responses ⁵.

The modification of autophagic and other membranes by mATG8s, a process referred to as membrane atg8ylation ⁵, depends on conjugation of ubiquitin-like protein ATG12 to ATG5 by way of successive reactions involving E1-like enzyme ATG7 and E2-like enzyme ATG10 ²⁰. The resulting ATG12-ATG5 conjugate in a complex with Atg16L1, forms the Atg12-

Atg5-Atg16L1 E3 ligase which guides lipidation of mATG8s ²⁰. In a parallel cascade, mATG8s are activated at their exposed C-terminal Gly residue by the E1-like ATG7 and the E2-like ATG3 enzymes ²⁰. A cross-over between the ATG12 conjugation and mATG8 conjugation branches has been reported, whereby a noncanonical ATG12-ATG3 conjugate is generated ⁴² with unconventional outputs via the ESCRT protein ALIX action in exosome secretion ⁴³. ALIX also participates in the repair of lysosomal membranes thus maintaining acidification and functionality of lysosomes ^{44–47}, whereas robustness and quality of the lysosomes is a determinant of effective autophagy ¹⁹.

The maturation of autophagosomes into autolysosomes where the captured cargo is degraded has many physiological outputs ⁴⁸. Among such functions are the control of inflammation and direct elimination of intracellular pathogens ⁴⁹. In macrophage infection models, induction of autophagy has been reported as a mechanism of controlling intracellular Mycobacterium tuberculosis (Mtb) ⁵⁰. In murine models of Mtb infection, genetic knockout of Atg5 in myeloid cells caused neutrophilic inflammation and increased mortality 51-53, whereas a loss of other Atg genes had no apparent consequences in short term studies ⁵³. Collectively, a conundrum has emerged of whether canonical autophagy acts here as a pathway and if all ATGs contribute to control of Mtb. In this study, we show that ATG5 has unique properties and functionalities outside of the canonical autophagy pathway. In the absence of ATG5, but not in the absence of other canonical autophagy factors, lysosomes are uncapable of recruiting ALIX and ESCRT repair machinery and are more sensitive to damage. The increased vulnerability of lysosomal membranes in ATG5 knockout cells contributes to a range of exocytic and secretory phenotypes. The lysosomal handicap impacts degranulation and other secretory functions of neutrophils, the inflammatory cells that have been previously linked to Mtb pathogenesis 51-54.

Results

Loss of ATG5 but not of other ATGs renders lysosomes excessively susceptible to damage

The initial reports testing in vivo role of autophagy genes in Mtb infection have shown that myeloid-specific Atg5 knockout, using Atg5^{fl/fl} LysM-Cre transgenic mice, renders mice more susceptible to respiratory Mtb infection ^{51,52}. Follow-up short-term infection studies (80 days) have reported that this Atg5 phenotype was unique among Atg genes and that a similar myeloid knockout in Atg7 (Atg7^{fl/fl} LysM-Cre) transgenic mice was not sensitized to Mtb ⁵³. We assessed these observations and extended them by employing a standard long-term low dose chronic Mtb infection model (aerosol exposure) ^{55–57} by monitoring mice for up to 200 days (Figure 1A). The results affirmed prior short-term observations with statistically significant differences in mortality between Atg5^{fl/fl} LysM-Cre⁺ vs. Atg5^{fl/fl} LysM-Cre⁻ mice but not between Atg7^{fl/fl} LysM-Cre⁺ vs. Atg7^{fl/fl} LysM-Cre⁻) mice to the metagenomic effects since we used only littermates in infection experiments ⁵⁸.

Prior work has also indicated alterations in inflammatory mediators linked to properties of macrophages and neutrophils ^{51–53} that remain to be fully characterized. Our recent studies and studies by others have indicated that mammalian ATG factors participate in the processes of atg8ylation ⁵, ESCRT-dependent repair ⁵⁹, and canonical autophagy ^{2,60},

which among other effects contribute to lysosomal quality control. Lysosomes are important degradative organelles ⁶¹, of particular significance in professional phagocytic cells. Hence, we tested whether there are differences in lysosomal quality between primary bone marrowderived macrophages (BMMs) from Atg5^{fl/fl} LysM-Cre⁺ and Atg7^{fl/fl} LysM-Cre⁺ mice. In these experiments we used LysoTracker Red (LTR) as a measure of the lysosomal status, and quantified LTR⁺ profiles by high content microscopy (HCM). HCM depends on epifluorescence mode of image capture to encompass all cellular profiles in a volume and allows unbiased machine-driven image acquisition and data analysis independently of the operator (each experiment consists of machine-identified 500 valid primary objects/cells per well, 5 wells/sample, in biological replicates of n 3) 18,47,59,62,63 . We treated ex vivo BMMs from Atg5^{fl/fl} LysM-Cre⁺ and Atg7^{fl/fl} LysM-Cre⁺ mice with a standard Cathepsin C-dependent lysosome-damaging agent Leu-Leu-OMe (LLOMe), which enters cells and endomembranous organelles by diffusion and specifically in lysosomes is converted into membrane-damaging polymers via a reverse dipeptidase action of cathepsins ^{47,64–68} (Figure S1A). The Atg5^{fl/fl} LysM-Cre⁺ BMMs showed lysosomal hypersensitivity (LyHYP) phenotype when treated with LLOMe in comparison to BMMs from Atg7^{fl/fl} LvsM-Cre⁺ mice, evidenced by the reduced number of LTR⁺ profiles of LTR⁺ compartments per cell (Figures 1B and 1C). Additionally, we confirmed dependence on ATG5 of the LyHYP phenotype in BMMs, by ex vivo downregulating ATG5 in BMMs from Atg5^{fl/fl} LysM-Cre⁻ by employing an optimized strategy for CRISPR-dependent gene disruption in primary mveloid cells 69. BMMs from Atg5fl/fl LysM-Cre- mice subjected to CRISPR with sgATG5 and Cas9 protein displayed LyHYP phenotype relative to BMMs treated with control guide RNA and Cas9 (Figures 1D and S1B).

To further dissect the lysosomal quality variances between ATG5- and ATG7-deficient cells, we tested whether the LyHYP phenotype was applicable to other cells. We generated ATG5 (Huh7ATG5-KO) and ATG7 (Huh7ATG7-KO) CRISPR knockouts in human hepatocyte-derived carcinoma cell line Huh7 (Figure 1E). We observed that lysosomes in Huh7ATG5-KO cells are more sensitive to damage, i.e. express LyHYP phenotype, relative to their Huh7^{WT} parental cells (Figures 1F, 1G and S1C; note ratios in Figure 1F inset). To ensure that the cellular content of lysosomes did not change in cells displaying lower number of LTR⁺ profiles, we quantified lysosomal content (LAMP2⁺ organelles) did not significantly differ between Huh7ATG5-KO and Huh7WT parental cells treated with LLOMe (Figures 1H and S1D). We next generated ATG5^{KO} in other human cell lines, HeLa, U2OS, and A549 (Figure S1E), and confirmed a LyHYP phenotype in all of the tested ATG5^{KO} lines (Figures 11 and S1O). No differences in lysosomal marker (LAMP1) profiles were observed in HeLaATG5-KO compared to HeLaWT treated with LLOMe (Figures S1F and S1G). LLOMe at concentrations used did not induce cell death or plasma membrane permeabilization as quantified by PI staining, with only 1-3 % of LLOMe-treated adherent cells being PI⁺ whether ATG5 WT or KO (Figures S1H–S1J). We ruled out that LyHYP could be explained by elevated Cathepsin C cellular levels, which were not increased in the absence of ATG5 in all cell types used herein (Figures S1K and S1L). LyHYP could not be explained by changes in endocytosis, as Dextran-TxR-3000 endocytic uptake was not altered in ATG5 KO relative to ATG5 WT cells (Figures S1M and S1N).

When we compared Huh7^{ATG7-KO} with Huh7^{ATG5-KO} isogenic Huh7 cell lines we observed that, like Atg7^{fl/fl} LysM-Cre⁺ vs Atg5^{fl/fl} LysM-Cre⁺ BMMs, Huh7^{ATG7-KO} and Huh7^{ATG5-KO} displayed different sensitivity to lysosomal damage. Huh7^{ATG5-KO} had a more pronounced loss of LTR⁺ profiles relative to Huh7^{ATG7-KO} (Figure 1J). Moreover, another KO mutant affecting atg8ylation conjugation cascade, ATG3 (Figure 1E), tracked with the milder effects of ATG7 loss relative to the highly pronounced effect of ATG5 loss on increased LyHYP (Figure 1J).

We employed another marker of lysosomal damage, galectin 3 (Gal3), a β -galactosidebinding cytosolic lectin that is recruited to lysosomes during their membrane damage by recognizing exposed lumenal glycans ⁶⁴, and can be used for quantitative assessment of lysosomal damage by HCM as previously described ⁴⁷. Upon treatment with LLOMe, all cell lines tested (Huh7, HeLa, A549, and U2OS) showed a quantifiable Gal3 response (Figures 1K and S1P). Further, Gal3 puncta response to LLOMe treatment was higher in ATG5^{KO} derivatives relative to parental ATG5 WT cells (Figures 1K and S1P). Primary macrophages mirrored these properties, as Atg5^{fl/fl} LysM-Cre⁺ BMMs showed stronger Gal3 response to LLOMe treatment than Atg5^{fl/fl} LysM-Cre⁻ BMMs (Figures 1L and S1Q). We generated partial ATG5^{KO} clones of pancreatic PaTu-8902 carcinoma cell line ^{70,71}, and observed that Gal3 response was elevated in CRISPR clones with 20–30% residual ATG12-ATG5 conjugate (Figures 1M and S1R).

The above relationships were observed using lysosomal damaging agents other than LLOMe, including silica crystals ^{47,72–74} and ORF3a^{SARS-CoV-2 75,76}, which elicited elevated Gal3 responses in ATG5^{KO} cells (Figures 1N, 1O, S1S, and S1T). We additionally verified the effects of ATG5 loss on ORF3a^{SARS-CoV-2}-induced lysosomal damage by LysoTracker Green (LTG) staining, and observed LyHYP in A549^{ATG5-KO} relative to A549^{WT} (Figure S1U and S1V).

Further, we generated a series of isogenic cell lines derived from Huh7 cells, adding ATG16L1, ATG13, ATG14 and VPS34 (Figure S1W) to the ATG7 and ATG3 KOs described above (Figure 1E) and assessed their sensitivity to lysosomal damage using Gal3 for quantification of their responses (Figure 1P). Huh7^{ATG7-KO} and Huh7^{ATG5-KO} isogenic cell lines had different Gal3 responses to LLOMe, with Huh7^{ATG5-KO} cells showing more Gal3 puncta than Huh7^{ATG7-KO} (Figure 1P) in keeping with ATG5 having stronger impact on lysosomal status and quality control than ATG7. The KOs of all other ATGs tested showed less pronounced Gal3 response compared to ATG5^{KO} cells (Figure 1P). Thus, ATG5 displays unique properties among all ATGs tested in rendering lysosomes more susceptible to damage.

We next tested whether LyHYP can be complemented by ATG5. We used two alleles of ATG5, ATG5^{WT} and its conjugation-deficient mutant, ATG5^{K130R} (Figure 1Q). We transfected HeLa^{ATG5-KO} cells with mCherry-ATG5^{WT} or mCherry-ATG5^{K130R} and assessed LyHYP upon LLOMe treatment by quantifying Gal3 and LTG responses by gating on transfected cells (Figures 1R, 1S, S1X, and S1Y). Both ATG5^{WT} and ATG5^{K130R} complemented HeLa^{ATG5-KO} cells and rescued lysosomal hypersensitivity phenotype (Figures 1R, 1S, S1X, and S1Y). Whereas it may appear surprising that both WT and

conjugation-deficient ATG5 complemented the ATG5^{KO}, this was in keeping with the observations that conjugation machinery (represented by ATGs such as ATG3, ATG7, and ATG16L1) did not strongly affect LyHYP (Figure 1P). Thus, ATG5 affects lysosomal sensitivity to damage independent of conjugation and canonical autophagy machinery (Figure 1T).

ATG5 is required for recruitment of ESCRTs to repair damaged lysosomes

Lysosomal homeostasis and membrane maintenance involve ESCRT-dependent mechanisms with protein ALIX playing a major role in lysosomal membrane repair ^{44,47,77,78}. Thus, we tested whether ATG5 has an effect on ALIX recruitment to damaged lysosomes. In Huh7^{ATG5-WT} cells, ALIX puncta were induced and ALIX colocalized with lysosomes upon 30 min of LLOMe-exposure, whereas in Huh7^{ATG5-KO} this response was diminished (Figures 2A–2C and S2A–S2C). This was not due to a decrease in ALIX protein levels (Figure S2D). In contrast to ALIX, another ESCRT-I component, TSG101, known to participate in lysosomal damage repair ^{44,77}, was detectable at very low levels as sparse puncta, and it too was affected (a measurable decrease) by the absence of ATG5 (Figures S2E and S2F).

ESCRT-III components, CHMP4B and CHIMP2A, known to act downstream of ALIX in lysosomal repair ^{44,47,77}, were also diminished in response and recruitment to lysosomes in Huh7^{ATG5-KO} cells relative to the parental Huh7^{ATG5-WT} cells (Figures 2D, 2E and S2G–S2L). The reduced ALIX response phenotype (overall puncta and their overlap with lysosomal marker LAMP2) was observed in ATG5^{KO} vs ATG5^{WT} U2OS and A549 cells (Figures S2M–S2T). We also confirmed the loss of ALIX-LAMP1 overlap phenotype associated with ATG5^{KO} in HL-60 cell line which can be differentiated into polymorphonuclear neutrophil (PMN)-like cells. We had to generate fresh ATG5^{KO} cells (which could not be maintained in passages) for each experiment (n=3). Differentiated HL-60s ATG5^{WT} and ATG5^{KO} cells were analyzed as nonadherent cells in an AMNIS imaging flow cytometer. ATG5^{KO} differentiated HL-60 cells displayed reduced Similarity Bright Detail parameter (an indicator of marker colocalization; IDEAS 6.2 software) for ALIX-LAMP1 overlap (Figure S2U).

We next complemented the loss of ALIX response phenotype in ATG5^{KO} cells, by transfecting them with pDEST-mCherry-ATG5 alleles (Figures 2F and S2V). As in the case of LyHYP complementation with both conjugation-competent (ATG5^{WT}) and conjugation-disabled ATG5^{K130R} alleles, the lack of ALIX response phenotype was complemented by both pDEST-mCherry-ATG5^{WT} and pDEST-mCherry-ATG5^{K130R} (Figures 2F and S2V), suggesting that the effects of ATG5 on ALIX recruitment to damaged lysosomes was independent of the conjugation-lipidation cascade (ATG3, ATG7, and ATG16L1), which normally work with ATG5 in the canonical autophagy pathway ²⁰, lacked pronounced effects on ALIX puncta response and overlaps with damaged lysosomes (Figures 2G–2I and S2W). Further, knockouts of other factors involved in canonical autophagy ², ATG13, ATG14, and VPS34, did not display an appreciable ALIX-deficiency phenotype compared to ATG5^{KO} (Figures 2H, 2I, S2W and S2X).

Finally, we confirmed by immunoblotting of purified lysosomes (LysoIP; Figure 2J) ⁷⁹ the defect in ALIX recruitment to lysosomes caused by the absence of ATG5. Purified lysosomes from HeLa^{ATG5-KO} had diminished ALIX and downstream ESCRT-III components (CHMP4B and CHMP2A) relative to HeLa^{ATG5-WT} cells exposed to lysosomal damage by LLOMe (Figures 2K and 2L). We conclude from the collective data presented above that in the absence of ATG5, ALIX and other ESCRTs cannot be efficiently recruited to lysosomes (Figure 2M), despite the presence of molecules that normally recruit ALIX to damaged lysosomes such as Ca²⁺ efflux and Gal3 ^{44,47,77,78}.

Proximity biotinylation proteomics of ATG5 interactors

The observations that ATG5, independently of the conventional conjugation/atg8ylation cascade, affects the ability of the cell to maintain functional lysosomes, prompted us to carry out unbiased proteomic analyses of ATG5 interactors. We employed proximity biotinylation ⁸⁰ and LC/MS/MS quantitative DIA approach ^{47,63,81}. After generating stable Flip-In HeLa cell lines expressing (Tet-On) APEX2 fusions, with the ATG5 alleles ATG5^{WT} and ATG5^{K130R}, we validate functionality of APEX2 fusions by detecting APEX2-ATG5^{WT} in conjugates with ATG12 that were absent in APEX2-ATG5^{K130R} expressing cells (Figure 3A(i)).

The above cells were subjected to lysosomal damage or not, and proximity biotinylation performed followed by LC/MS/MS and quantitative DIA analysis (Table S1, Tabs 1-10). Among the potential ATG5^{WT} proximity interactors with changes upon lysosomal damage (Table S1, Tabs 2,3) an enrichment of proteins associated with exosomes and exocytosis/degranulation (Figure 3B) was observed. In addition to all ATG5 proximity interactors (Figure 3B. Table S1, Tabs 1-3), we next applied an algorithm (Figure 3A(iii)) that discriminates potential ATG5 partners comparing conjugation-competent (ATG5^{WT}) and conjugation-disabled (ATG5K130R) ATG5-interactor subsets. Gene Ontology (GO) biological processes and cellular component global analyses of potential interactors that are in the vicinity of ATG5 independently of ATG5's conjugation capacity (Figure 3C, Table S1) corroborated the general trend observed in the volcano plot (Figure 3B) of the cumulative proteomic data. GO analyses of the data within conjugation-dependent ATG5 interactor group (Figure S3A, Table S1) indicated that they are globally different from conjugation-independent subset and showed no enrichment in secretory components (Figure S3A). Thus, the unique properties of ATG5, i.e. those independent of its participation in conventional conjugation with ATG12, appear to be associated, directly or indirectly, with sets of proteins participating in intracellular trafficking and vectorial transport, exocytosis, degranulation, and exosome secretion.

Lysosomal damage induces release of extracellular vesicles and particles and neutrophil degranulation

We next wanted to address whether ATG5 affects release of extracellular vesicles and particles (EVPs) and other exocytic events based on the following rationale: (i) ATG5 deficiency results in more vulnerable lysosomes; (ii) this effect is independent of canonical conjugation; and (iii) functional enrichment analyses of proteomic data that implicated exocytic processes were independent of ATG5's conjugation capacity. We first established

whether lysosomal damage elicits exocytic processes. Using nanoparticle tracking analysis (NTA; NanoSight), we observed extracellular particles in EVP-enrichment preparations ^{82,83} from supernatants of HeLa cells, which increased upon treatment with LLOMe (Figures 4A and 4B), with some shift in their size (Figure S4A). NTA analysis was also applied to cells subjected to lysosomal damage caused by overexpressing ORF3a^{SARS-CoV-2}, known to cause lysosomal damage ⁷⁶ and recently reported to play a role in secretory autophagy (the type termed SALI) via effects on autophagosomal maturation ⁸⁴, and it showed increased particle release (Figures 4C and 4D). These observations were followed up in imaging-flow cytometry (AMNIS; ImageStream), whereby we detected CD63⁺ extracellular vesicles (EVs) (Figure S4B) released from cells subjected to lysosomal damage either by LLOMe (Figure 4E) or ORF3a^{SARS-CoV-2} (Figure 4F).

Release of CD63⁺ EV was elicited by lysosomal damage in primary cells as well. This included macrophages, i.e. BMMs (Figure 4G), and bone marrow derived neutrophils (BMNs) (Figures 4H, 4I and S4C). Furthermore, we tested levels of CD11b (ITGAM) on Ly6G⁺ BMNs as a measure of neutrophil activation and degranulation ^{85–88}. These membrane proteins become exposed on the cell surface upon granule exocytosis. We found that the percent of CD11b^{Hi} Ly6G⁺ BMNs increased upon LLOMe treatment (Figures 4J–4L). This increase was comparable (Figures 4J–4L) to the effects of the conventional inducer of BMN degranulation, fMLF ^{89,90}. Thus, lysosomal damage elicits exocytosis of EVPs and degranulation in PMNs. The latter also evident from increase in extracellular levels of elastase, a component of neutrophil primary granules ⁹¹ (Figure 4M).

ATG5 knockout enhances EVP release in response to lysosomal damage

Having established that lysosomal damage elicits exocytic processes including EV release, we tested whether ATG5 status affects this phenomenon. ATG5-deficient HeLa cells displayed appreciable increase in release of CD63⁺ EVs and total EVPs relative to HeLa^{WT} cells (Figures 5A and 5B). This ATG5-deficient phenotype was partially complemented in transfections with either mCherry-ATG5^{WT} or its conjugation-incompetent mutant mCherry-ATG5^{K130R} (Figures S5A and S5B). Quantitative immunoblotting analysis of enriched exosomes from the supernatants of cells treated with LLOMe indicated elevated markers/ components of exosomes (ALIX, TSG101, and syntenin-1; ^{92,93}) in ATG5-deficient HeLa cells (Figure 5C). A similar increase in exosomal markers was observed in ATG5-deficient cells expressing ORF3a^{SARS-CoV-2} (Figures 5D–5F).

We next tested primary neutrophils and compared EVPs released by murine BMNs derived from bone marrow of Atg5^{fl/fl} LysM-Cre⁺ and Atg5^{fl/fl} LysM-Cre⁻ transgenic mice. By NTA (NanoSight) analysis of particles released from these BMNs, we observed increased release of EVPs (Figures 5G). This was followed up by imaging-flow cytometry (AMNIS ImageStream) of enriched CD63⁺ exosomes, showing that both neutrophils/BMNs (Figure 5H) and macrophages/BMMs (Figure 5I) lacking ATG5 displayed augmented release of EVPs (including exosomes) in response to lysosomal damage.

ATG5 knockout enhances PMN degranulation and lysosomal exocytosis in response to lysosomal damage

The BMNs derived from Atg5^{fl/fl} LysM-Cre⁺ mice displayed increased parameters of exocytosis of secretory vesicles and granules (degranulation) in response to lysosomal damage. This was manifested as increase in CD11b, a cell surface marker of exocytosis, which was elevated on live Atg5^{fl/fl} LysM-Cre⁺ but not on Atg7^{fl/fl} LysM-Cre⁺ BMNs treated with LLOMe (Figures 6A-6C and S5C). The % of CD11bHi Ly6G⁺ cells were compatible with previously reported % levels with bone marrow derived neutrophils ^{94,95} vis-à-vis the customarily higher % levels found in peripheral blood PMNs. The markers of degranulation/exocytosis were also increased in Atg5fl/fl LysM-Cre+ BMNs upon treatment with the conventional inducer fMLF (Figures 6A and S5C), indicating that the receptor system for detection of fMLF was likely elevated on the surface on Atg5^{fl/fl} LysM-Cre⁺ BMNs, compatible with increased basal release of secretory vesicles where fMLF receptors are principally localized ⁹⁶. This was corroborated by the augmented cell surface FPR1 levels in Atg5^{fl/fl} LysM-Cre⁺ BMNs compared with Atg5^{fl/fl} LysM-Cre⁻ BMNs, reaching significance under LLOMe-treatment conditions (Figures 6D and S5D). Neutrophil elastase (a component of primary/azurophilic granules) was released in response to lysosomal damage in higher amounts from Atg5^{fl/fl} LysM-Cre⁺ BMNs than from Atg5^{fl/fl} LysM-Cre⁻ BMN (Figure 6E). Thus, we conclude that neutrophils lacking ATG5 undergo increased exocytosis and degranulation in response to stimulation by physiological agonists (fMLF) or lysosomal damage.

Lysosomal exocytosis was increased in Atg5^{fl/fl} LysM-Cre⁺ BMNs relative to Atg5^{fl/fl} LysM-Cre⁻ BMNs (Figure 6F). For this we used LAMP2 normally present in lysosomes but translocating to plasma membrane where it is exofacially exposed upon fusion of lysosomes with the cell surface. By comparison to activation of lysosomal exocytosis (proxied by cell surface LAMP2) upon lysosomal damage, fMLF did not elicit this response (Figure 6F). For flow cytometry carried out with live neutrophils, sodium azide was used to block reuptake and endocytosis (Figures 6F and 6G) ^{97–99}. To complement this by HCM with adherent cells, we allowed endocytosis (if any) to take place during staining of live and unpermeabilized Huh7 cells with anti-LAMP2 lumenal domain mAb and quantified LAMP2 puncta. By this method, Huh7^{ATG5-KO} cells exposed to LLOMe showed increased number of LAMP2 puncta relative to untreated Huh7^{ATG5-KO} cells, whereas Huh7^{ATG7-KO} cells did not have that phenotype (Figures 6H and 6I). This was confirmed by staining for LAMP1 using the same procedure with live cells (Figures 6J and 6K) and confocal microscopy (Figures 6L and 6M).

In conclusion, ATG5 defect promotes lysosomal exocytosis and elicits degranulation in PMNs in response to damage and stimulation by physiological ligands (Figure 6N).

Alternative conjugation contributes to lysosomal vulnerability and exocytic processes

Recent studies have indicated the existence of an alternative conjugation pathway whereby ATG12 can form a sidestep conjugate ATG12-ATG3 that contributes to increased exosome secretion ^{42,43}. We detected increased levels of ATG12-ATG3 conjugates in ATG5^{KO} cells (Figures S6A and S6B). Furthermore, we observed that ATG3 puncta are increased in

Huh7^{ATG5-KO} cells vs. Huh7^{WT} irrespective of LLOMe treatment, whereas distinct, large ALIX puncta are elicited by lysosomal damage in Huh7^{WT} cells and do not appear in Huh7^{ATG5-KO} cells (Figures 7A–7E). Moreover, colocalization between ATG3 and ALIX significantly increases in response to LLOMe in the absence of ATG5, i.e., in Huh7^{ATG5-KO} cells, whereas this phenomenon is absent in Huh7^{WT} cells (Figures 7E and S6D). ALIX was increased in protein complexes with ATG3, which included elevated ATG12-ATG3 conjugate (Figure S6C). A knockout of ATG3 alone did not alter the EV release in response to LLOMe (Figures 7F, 7G, and S6E; HeLa^{ATG3-KO} vs. HeLa^{WT}). However, a knockout of ATG3 in HeLa^{ATG5-KO} cells, thus generating double KO cells (HeLa^{ATG3/5-DKO}), abrogated the elevated EVPs release in response to LLOMe treatment seen in single, ATG5^{KO} cell line (Figures 7F, 7G and S6E; HeLa^{ATG5-KO} vs. HeLa^{ATG3/5-DKO}). This was confirmed by measuring CD63⁺ EVs using AMNIS in Huh7 cells (Figures S6F and S6G). Thus, ablation of ATG3 in the absence of ATG5 results in normalization of exocytic processes.

Furthermore, the number of ALIX puncta and ALIX-LAMP1 colocalization following LLOMe treatment, which decreased in Huh7^{ATG5-KO} cells, were partially recovered by ATG3 KO in Huh7^{ATG5-KO} cells (Huh7^{ATG3/5-DKO} cells vs Huh7^{ATG5-KO}; Figures 7H, 7I and S6H, top row). Mirroring the partial recovery of ALIX recruitment in Huh7^{ATG3/5-DKO} relative to Huh7^{ATG5-KO} cells in response to lysosomal damage, Gal3 as a marker of the extent of the damage was partially suppressed in Huh7^{ATG3/5-DKO} cells (Figures 7J and S6H, bottom row). This is indicative of improvement in the ability of cells lacking both ATG5 and ATG3 to repair lysosomes. This paradoxical observation is consistent with a model (Figure 7K) whereby ATG12-ATG3 increases in the absence of ATG5, whereas ATG12 is normally tied up in ATG12-ATG5 conjugates when ATG5 is available at normal levels. In the absence of ATG5, the alternative ATG12-ATG3 conjugate contributes to ALIX's mislocalization and diminishes its function in the lysosome repair.

Discussion

In this study, we uncovered unanticipated aspects of the protein conjugation and lipidation cascade commonly associated with autophagy. Whereas ATG5 has been primarily studied as a part of the E3 ligase leading to mATG8s lipidation, here we show that ATG5 is positioned at a critical intersection with other conjugation branches (Figure S6I) affecting lysosomal quality, exocytosis, and neutrophil degranulation independently of the conventional mATG8 lipidation and canonical autophagy. In the absence of ATG5, ESCRT proteins are not efficiently recruited to the membrane repair sites on damaged lysosomes. Under these conditions, lysosomes, storage granules, secretory vesicles, and EVPs are prone to exocytosis. The dysregulated exocytic processes in neutrophils may contribute to infection outcomes in Mtb murine models exceeding ATG5's role in canonical autophagy.

We find that a loss of ATG5, and the resulting ATG12-ATG3 conjugate formation favored in the absence of ATG5, diminish available ALIX thus compromising lysosomal membrane repair ^{44,47,77}. This results in a plethora of downstream effects such as lysosomal hypersensitivity to damage and includes roles in exosome formation ⁴³ and other processes. These phenomena are different from the recently described eructophagy ¹⁰⁰ since the latter

depends on ATG genes other than ATG5. Other studies propose additional ATG5-specific effects, including measurable metabolic consequences ¹⁰¹.

Reduced levels of ATG5 may occur in physiological and disease states. Lower ATG5 levels are a risk factor in Parkinson's diseases (PD) ^{102–105}. Changes in ATG5 levels have been reported in diabetes ¹⁰⁶ and cancer ^{107–111}. Lysosomal status has been linked specifically to pancreatic cancer cells ¹¹² and lysosomal degradation of plasma membrane MHC-I molecules ¹¹³. ATG5 is implicated in autoimmunity and inflammation ⁴⁹. Loss of ATG5 affects cardiac capacity ¹¹⁴. Thus, changes in ATG5 may affect a variety of diseases.

Our present study uncovers effects of ATG5 knockout on neutrophils, first responders to bacterial infection ^{115,116}. When neutrophils are hyperactivated, they can cause tissue damage ^{117,118}. Neutrophils are activated via receptors ^{119,120} such as FPR1 and CD11b/ CD18 associated with neutrophil degranulation, and were increased in ATG5 knockouts. Granule components are found in neutrophil NETs released during NETosis ¹²¹. Although ATG5 does not alter NETosis ¹²², NETs contain granule components ¹²³ and may be affected by an ATG5 loss. In human populations, active tuberculosis disease is associated with persistent neutrophilic signatures of inflammation ⁵⁴; we postulate that this is linked to the findings in our study concerning neutrophil degranulation, exocytosis, and EVP release.

Limitations of the study

Limitations of our study include a focus on neutrophils and a diversity of exocytic processes altered in ATG5-deficient cells that could not be easily parsed for effects in vivo. It remains to be determined which subsets are responsible for increased susceptibility to Mtb in mice with Atg5-deficient myeloid cells. This is compounded by the duality of phenotypes downstream of the ATG5 deficiency, i.e., lysosomal damage vs. exocytosis and exosomal secretion, and a partial link of both aspects to the sequestration of ALIX. There are additional effects of ATG5 on the lysosomal status beyond the sequestration of ALIX by ATG12-ATG3, and these processes remain to be delineated.

STAR Methods

RESOURCE AVAILABILITY

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Vojo Deretic (vderetic@salud.unm.edu).

Materials availability—Plasmids and cell lines generated in this study are available from the lead contact.

Data and code availability—Raw MS DIA/DDA data have been deposited at the MassIVE proteomics repository MassIVE (MSV000090348) and Proteome Exchange (PXD036850).

Any additional information required to reanalyze the data reported in this work paper is available from the Lead Contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice—Atg5^{fl/fl} LysM-Cre⁻, Atg5^{fl/fl} LysM-Cre⁺, Atg7^{fl/fl} LysM-Cre⁻, and Atg7^{fl/fl} LysM-Cre⁺ mice were previously described ^{51,125}.

Housing and husbandry conditions of experimental animals—All mice were housed in AAALAC-accredited Animal Research Facility (ARF) of the University of New Mexico Health Sciences Center (UNM HSC) and institutionally approved husbandry conditions and approved breeding protocols were followed. *M. tuberculosis*-infected animals were housed in a separate ABSL3 suite within the UNM HSC ARF facility and all staff followed strict ABSL3, BSL3, and animal protocols approved by the UNM HSC Biosafety committee and the Institutional Animal Care and Use Committee. The study was compliant with all relevant ethical guidelines for animal research.

Cells and cell line models—HEK293T, HeLa, A549 and U2OS cells were from ATCC. Huh7 cells were from Rocky Mountain Laboratories. Patu-8902 cells were from Alec Kimmelman. Murine primary cells (Bone marrow derived macrophages and neutrophils; BMMs and BMNs) were used for degranulation, exosome secretion, and lysosomal damage studies. BMMs were isolated from femurs Atg5/7^{fl/fl} LysM-Cre⁻ and Atg5/7^{fl/fl} LysM-Cre⁺ mice and cultured in DMEM supplemented with mouse macrophage colony stimulating factor (mM-CSF, #5228, CST) for 5 to 10 days before use. BMNs were isolated from femurs Atg5/7^{fl/fl} LysM-Cre⁻ and Atg5/7^{fl/fl} LysM-Cre⁺ mice according to ¹²⁶ and gating strategy according to ¹²⁷. Briefly, mouse bone marrow cells were collected, filtered, and centrifuged, followed by 0.2% NaCl lysis red blood cells and gradient centrifugation with Histopaque 1119 and 1077. Cells were then washed and recovered for 1 hour before use. Flp-In cell lines were generated using constructs from Terje Johansen, cell lines for LysoIP were generated using constructs obtained from Addgene, and knockout cell lines were generated by CRISPR/Cas9-mediated knockout system as detailed below.

METHOD DETAILS

Antibodies—Antibodies from Abcam were ATG5 (1:2000 for Western blot (WB), ab108327), ATG7 (1:2000 for WB, ab52472), mCherry (1:1000 for WB, ab183628), Syntenin-1 (1:1000 for WB, ab19903), TSG101 (1:1000 for WB; 1:200 for immunofluorescence (IF), ab83), VPS34 (1:1000 for WB, ab124905).

Antibodies from Biolegend were ALIX (1:400 for IF, #634502), Galectin-3 (1:200 for IF, #125402).

Antibodies from Cell Signaling Technology were ATG13 (1:1000 for WB, #13468), GM130 (1:1000 for WB, #12480), HA-tag (1:1000 for WB, #3724), LAMP1 (1:1000 for WB; 1:1000 for IF, #9091), PDI (1:1000 for WB, #3501).

Antibodies from Proteintech were ALIX (1:500 for WB, 12422-1-AP), CHMP2A (1:500 for WB; 1:500 for IF, 10477-1-AP), and CHMP4B (1:500 for WB; 1:500 for IF, 13683-1-AP). Antibodies from Sigma Aldrich were ATG3 (1:1000 for WB; 1:300 for IF, #A3231) and CD63 (1:400 for IF, SAB4301607).

Other antibodies used in this study were from the following sources: beta-Actin (1:500 for WB, sc-47778), Cathepsin C (1:500 for WB, sc-74590) and GAPDH (1:500 for WB, sc-47724) from Santa Cruz Biotechnology; LAMP2 (1:1000 for IF, H4B4) from DSHB of University of Iowa; FITC-FPR1 (1:400 for IF, FPR1-FITC) from Fabgennix; ATG14 (1:500 for WB, PD026) and ATG16L1 (1:500 for IF, PM040) from MBL International; CD11b (1:500 for IF, NB110-89474) from Novus Biological; Ly-6G (1:500 for IF, #60031) from STEMCELL Technologies; and ATG12 (1:1000 for WB, 366400) from Thermofisher. Alexa Fluor 488, 568, 647 (1:500 for IF) and IgG-HRP (1:10000 for WB) secondary antibodies were from ThermoFisher Scientific. IgG Polyclonal Antibody (IRDye[®] 800CW and IRDye[®] 680RD) (1:10000 for WB) secondary antibodies were from LI-COR Biosciences.

Plasmids, siRNAs, and transfection—Plasmids used in this study, such as ATG5 cloned into pDONR221 using BP cloning, and expression vectors were made utilizing LR cloning (Gateway, ThermoFisher) in appropriate pDEST vectors for immunoprecipitation assay.

ATG5 mutants were generated utilizing the QuikChange site-directed mutagenesis kit (Agilent) and confirmed by sequencing (Genewiz). Plasmid transfections were performed using the ProFection Mammalian Transfection System, FuGENE[®] HD Transfection Reagent (Promega), or Lipofectamine 2000 Transfection Reagent (ThermoFisher Scientific).

Generation of CRISPR mutant cells—Knockout cells (HeLaATG5-KO, HeLaATG3-KO, HeLaATG3/5-DKO, Huh7ATG5-KO, Huh7ATG3-KO, Huh7ATG7-KO, Huh7ATG13-KO, Huh7ATG14-KO, Huh7ATG16L1-KO, Huh7VPS34-KO, Huh7ATG3/5-DKO, U2OSATG5-KO, and A549ATG5-KO) were generated by CRISPR/Cas9-mediated knockout system. The lentiviral vector lentiCRISPRv2 carrying both Cas9 enzyme and a gRNA targeting ATG5 (gRNA-puro: AAGAGTAAGTTATTTGACGT), ATG3 (gRNA-Hygro: GTGAAGGCATACCTACCAAC), ATG7 (gRNA-puro: CTTCCGTGACCGTACCATGC), ATG13 (gRNA-puro: CTGTCCCAACACGAACTGTC), ATG14 (gRNA-puro: CTACTTCGACGGCCGCGACC), ATG16L1 (gRNA-puro: ACCAAATGCAGCGGAAGGAC), VPS34 (gRNA-puro: ACAGACATTTGTAGATCGGT) (VectorBuilder) were transfected into HEK293T cells together with the packaging plasmids psPAX2 and pCMV-VSV-G at the ratio of 5:3:2. Two days after transfection, the supernatant containing lentiviruses was collected. Cells were infected by the lentiviruses with 8-10 μ g/mL polybrene. 36 h after infection, the cells were selected with puromycin (1–10 μ g/mL) for one week in order to select knockout cells. All knockouts were confirmed by western blot. Selection of single clones was performed by dilution in 96-well.

Generation of CRISPR mutant in primary cells—CRISPR knockout of ATG5 in primary murine BMMs was performed as described ⁶⁹. Briefly, monocyte-derived macrophages were cultured in macrophage medium (DMEM high glucose supplemented with 10% FBS [Gibco], 2 mMGlutaMAX, 100U/ml penicillin, 100 µg/ml streptomycin, and M-CSF 100 ng/ml [Peprotech]). Medium was changed every 2–3 d by adding half the volume of medium. gRNAs (IDT) were mixed with Cas9 (IDT Alt-R S.p. Cas9 Nuclease V3) at a molar ratio of 2:1 and incubated at room temperature for 20 min. Macrophages were detached (Detachin, Genlantis), washed, counted and resuspended in 20 µl of P3

primary nucleofection solution (Lonza) per 10⁶ cells. The cells were then added to the Cas9-gRNAs complex and mix by pipetting. The mix was the loaded into the nucleofector cassette strip (Lonza) and nuclefected using the Buffer P3, CM-137 condition in the Lonza 4D-Nucleofector (4D-Nucleofector Core Unit: Lonza, AAF-1002B; 4D-Nucleofector X Unit: AAF-1002X). The cells were then cultured and KO efficiency assayed by Western blot.

Generating Flp-In-ATG5-WT/K130R^{Tet-ON} cell line—Transfect Hela Flp-In host cells with ATG5/K130R reconstructed plasmid and the pOG44 expression plasmid at ration of 9:1. 24 h after transfection, wash the cells and add fresh medium to the cells. 48 h after transfection, split the cells into fresh medium around 25 % confluent. Incubate the cells at 37 °C for 2–3 h until they have attached to the culture dish. Then them medium was removed and added with fresh medium containing hygromycin. Feed the cells with selective medium every 3–4 days until single cell clone can be identified. Pick hygromycin-resistant clones and expand each clone to test. The tested clones incubated in the medium containing 1–2 µg/mL tetracycline for 24 h were determined by western blot for the expressing of ATG5.

LysoIP assay—Lentiviruses constructs for generating stable LysoIP cells were purchased from Addgene. HEK293T cells were transfected with pLJC5-TMEM192–3xHA or pLJC5-TMEM192–2XFLAG constructs in combination with pCMV-VSV-G and psPAX2 packaging plasmids, 60 h after transfection, the supernatant containing lentiviruses was collected and centrifuged to remove cells and then frozen at -80 °C. To establish LysoIP stably expressing cell lines, cells were plated in 10cm dish in DMEM with 10 % FBS and infected with 500µL of virus-containing media overnight, then add puromycin for selection.

Selected cells in 15 cm plates with 90 % confluency were used for each LysoIP. Cells with or without treatment were quickly rinsed twice with PBS and then scraped in1mL of KPBS (136 mM KCl, 10 mM KH₂PO₄, pH7.25 was adjusted with KOH) and centrifuged at 3000 rpm for 2 min at 4 °C. Pelleted cells were resuspended in 950 μ L KPBS and reserved 25 μ L for further processing of the whole-cell lysate. The remaining cells were gently homogenized with 20 strokes of a 2 mL homogenizer. The homogenate was then centrifuged at 3000 rpm for 2 min at 4 °C and the supernatant was incubated with 100 μ L of KPBS prewashed anti-HA magnetic beads (ThermoFisher) on a gentle rotator shaker for 15 min. Immunoprecipitants were then gently washed three times with KPBS and eluted with 2 × Laemmli sample buffer (Bio-Rad) and subjected to immunoblot analysis.

High content microscopy (HCM)—Cells in 96 well plates were fixed in 4 % paraformaldehyde for 5 min. Cells were then permeabilized with 0.1 % saponin in 3 % Bovine serum albumin (BSA) for 30 min followed by incubation with primary antibodies for 2 h and secondary antibodies for 1h. Hoechst 33342 staining was performed for 3 min. High content microscopy with automated image acquisition and quantification was carried out using a Cellomics HCS scanner and iDEV software (ThermoFisher Scientific). Automated epifluorescence image collection was performed for a minimum of 500 cells per well. Epifluorescence images were machine analyzed using preset scanning parameters and object mask definitions. Hoechst 33342 staining was used for autofocus and to automatically define cellular outlines based on background staining of the cytoplasm. Primary objects were cells,

and regions of interest (ROI) or targets were algorithm-defined by shape/segmentation, maximum/minimum average intensity, total area and total intusity, etc., to automatically identify puncta or other profiles within valid primary objects. Each experiment (independent biological repeats; n 3) consists of machine-identified 500 valid primary objects/cells per well, 5 wells/sample. All data collection, processing (object, ROI, and target mask assignments) and analyses were computer driven independently of human operators.

Co-immunoprecipitation assay—Cells transfected with 8–10 μ g of plasmids were lysed in NP-40 buffer (ThermoFisher Scientific) supplemented with protease inhibitor cocktail (Roche, 11697498001) and 1 mM PMSF (Sigma, 93482) for 30 min on ice. Supernatants were incubated with (2–3 μ g) antibodies overnight at 4 °C. The immune complexes were captured with Dynabeads (ThermoFisher Scientific), followed by three times washing with 1X PBS. Proteins bound to Dynabeads were eluted with 2×Laemmli sample buffer (Bio-Rad) and subjected to immunoblot analysis.

Immunofluorescence confocal microscopy and analysis—Cells were plated onto coverslips in 6-well plates. After treatment, cells were fixed in 4% paraformaldehyde for 5 min followed by permeabilization with 0.1% saponin in 3% BSA for 30 min. Cells were then incubated with primary antibodies for 2 h and appropriate secondary antibodies Alexa Fluor 488 or 568 (ThermoFisher Scientific) for 1 h at room temperature. Coverslips were mounted using Prolong Gold Antifade Mountant (ThermoFisher Scientific). Images were acquired using a confocal microscope (META; Carl Zeiss) equipped with a 63 3/1.4 NA oil objective, camera (LSM META; Carl Zeiss), and AIM software (Carl Zeiss).

Flow cytometry—BMNs after treatment were washed with 3% BSA in PBS supplemented with 0.1% of NaN₃ before staining for surface markers. After staining, cells were then resuspended with 3% BSA in PBS supplemented with 0.1% of NaN₃ until acquisition on Amins ImageStreamx MKII (ISx, EMD Millipore, Seattle, WA, USA). For neutrophil degranulation studies, Ly6G⁺ neutrophils were gated for analysis of surface CD11b.

Exosome isolation—Exosomes are isolated using SmartSECTM (SSEC200A-1) and ExoQuick-TC (EXOTC10A-1) according to manufacturer's instructions from supernatant of cells. For mCherry-ATG5^{WT} and mCherry-ATG5^{K130R} transfection experiments, cells were incubated in LLOMe in a capped Eppendorf tube for 1 h in 37°C after transfection for 48 h. For mCherry-ORF3a^{SARS-CoV-2}, supernatant was collected from 6 h after transfection when lipofectamine 2000 was washed away and new medium was added until 48 h after transfection.

ImageStreamx small particle acquisition and analysis—Multispectral imaging flow-cytometric acquisition of EVs and small particle calibrators was performed using Amins ImageStreamx MKII (ISx, EMD Millipore, Seattle, WA, USA) with fluidics set at low speed, sensitivity set to high, magnification at $60\times$, core size 7 µm, and the "Hide Beads" option unchecked prior to every acquisition in order to visualize speed beads in analyses. All parameters are stored in acquisition template except the latter, which requires unchecking prior to each acquisition. The ISx was equipped with the following lasers run at maximal power to ensure maximal sensitivity: 405 nm (120 mW), 488 nm (200 mW), 561

nm (200 mW), and 642 nm (150 mW). Upon each startup, the instrument calibration tool ASSIST[®] was performed to optimize performance and consistency. Each of the two charged couple device (CCD) cameras with which the ISx is equipped have six channels of detection. Two channels (Ch01 and Ch09) were set to brightfield (BF), permitting spatial coordination between cameras. Channel 12 was set to side-scatter (SSC), and further fluorescence channels were used for antibody detection as required. The advanced fluidic control of ISx, coupled with the presence of continuously running speed beads enable cell/particle enumeration using the "objects per mL" feature within the IDEAS[®] data analysis software. To avoid the risk of coincident particle detection, EV samples were not run at concentrations greater than 1010 objects/mL. All samples were acquired using INSPIRE[®] software, with a minimum of 5,000 G1 events collected, or as dictated by the type of analysis to be undertaken. Data analyses were performed, and spectral compensation matrices produced, using ISx Data Exploration and Analysis Software (IDEAS[®]).

Nanoparticle tracking analysis—Isolated EVPs (as described above) from cell culture supernatants were diluted in PBS and used for nanoparticle tracking analysis using a Nanosight NS300 instrument (NanoSight), followed by evaluation using the Nanoparticle Tracking Analysis (NTA) software. Conditions were as follows: camera type, sCMOS; detection threshold, 5; recording for 800 frames at 25 frames per s.

APEX2 labelling and streptavidin enrichment for LC–MS/MS analysis—

HeLaAPEX2-ATG5-WT and HeLaAPEX2-ATG5-K130R cells were incubated in complete medium supplemented with 500 µM biotin-phenol (AdipoGen) with or without 2 mM LLOMe for 30 min. A 1-min pulse with 1 mM H_2O_2 at room temperature was stopped with quenching buffer (10 mM sodium ascorbate, 10 mM sodium azide and 5 mM Trolox in PBS). All samples were washed twice with quenching buffer, and twice with PBS for 1 min. For LC–MS/MS analysis, cell pellets were lysed in 500 µl ice-cold lysis buffer (6 M urea, 0.3 M NaCl, 1 mM EDTA, 1 mM EGTA, 10 mM sodium ascorbate, 10 mM sodium azide, 5 mM Trolox, 1% glycerol and 25 mm Tris-HCl, pH 7.5) for 30 min by gentle pipetting. Lysates were clarified by centrifugation and protein concentrations were determined using Pierce 660 nm protein assay reagent. Streptavidin-coated magnetic beads (Pierce) were washed with lysis buffer. A total of 1 mg of each sample was mixed with 100 µl of streptavidin beads. The suspensions were gently rotated at 4 °C overnight to bind biotinylated proteins. The flow-through after enrichment was removed and the beads were washed in sequence with 1 ml IP buffer (150 mM NaCl, 10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 1 mM EGTA, 1% Triton X-100) twice; 1 ml 1 M KCl; 1 ml of 50 mM Na₂CO₃; 1 ml 2 M urea in 20 mM Tris HCl (pH 8.0); and 1 ml IP buffer. Biotinylated proteins were eluted and processed for mass spectrometry.

Sample preparation for LC-MS/MS—Protein samples on magnetic beads were washed four times with 200 μ l of 50 mM Triethyl ammonium bicarbonate (TEAB) with a twentyminute shake time at 4 °C in between each wash. Roughly 2.5 μ g of trypsin was added to the bead and TEAB mixture and the samples were digested over night at 800 rpm shake speed. After overnight digestion the supernatant was removed and the beads were washed once with enough 50 mM ammonium bicarbonate to cover. After 20 min at a gentle shake

the wash is removed and combined with the initial supernatant. The peptide extracts are reduced in volume by vacuum centrifugation and a small portion of the extract is used for fluorometric peptide quantification (Thermo scientific Pierce). One microgram of sample based on the fluorometric peptide assay was loaded for each LC-MS analysis.

Liquid chromatography-tandem mass spectrometry—Peptides were desalted and trapped on a Thermo PepMap trap and separated on an Easy-spray 100 μ m × 25 cm C18 column using a Dionex Ultimate 3,000 nUPLC at 200 nL/min. Solvent A = 0.1% formic acid, Solvent B = 100% Acetonitrile 0.1% formic acid. Gradient conditions = 2% B to 50% B over 60 min, followed by a 50%–99% B in 6 min and then held for 3 min than 99% B to 2% B in 2 min and total run time of 90 min using Thermo Scientific Fusion Lumos mass spectrometer. The samples were run in DIA mode; mass spectra were acquired using a collision energy of 35, resolution of 30 K, maximum inject time of 54 *ms* and an AGC target of 50K, using staggered isolation windows of 12 Da in the *m/z* range 400–1,000 *m/z*.

DIA Quantification and Statistical Analysis—DIA data was analyzed using Spectronaut 14.10 (Biognosys Schlieren, Switzerland) using the directDIA workflow with the default settings. Briefly, protein sequences were downloaded from Uniprot (Human Proteome UP000005640), ATG5 from Uniprot and common laboratory contaminant sequences from https://thegpm.org/crap/. Trypsin/P specific was set for the enzyme allowing two missed cleavages. Fixed Modifications were set for Carbamidomethyl, and variable modification were set to Acetyl (Protein N-term) and Oxidation. For DIA search identification, PSM and Protein Group FDR was set at 1%. A minimum of 2 peptides per protein group were required for quantification. Proteins known to be endogenously biotinylated were excluded from consideration. Pathway enrichment analysis and gene ontology of Biological Process and Cellular Component of the ATG12-ATG5 conjugationindependent or conjugation-dependent protein subsets were performed using STRING (https://string-db.org/).

Quantification and statistical analysis

Data in this study are presented as means \pm SEM (n 3). Data were analyzed with either analysis of variance (ANOVA) with Tukey's HSD post-hoc test, or a two-tailed Student's *t*-test. For HCM, n 3 includes in each independent experiment: 500 valid primary objects/ cells per well, from 5 wells per plate per sample. Statistical significance was defined as: † (not significant) p 0.05 and *p < 0.05, **p < 0.01.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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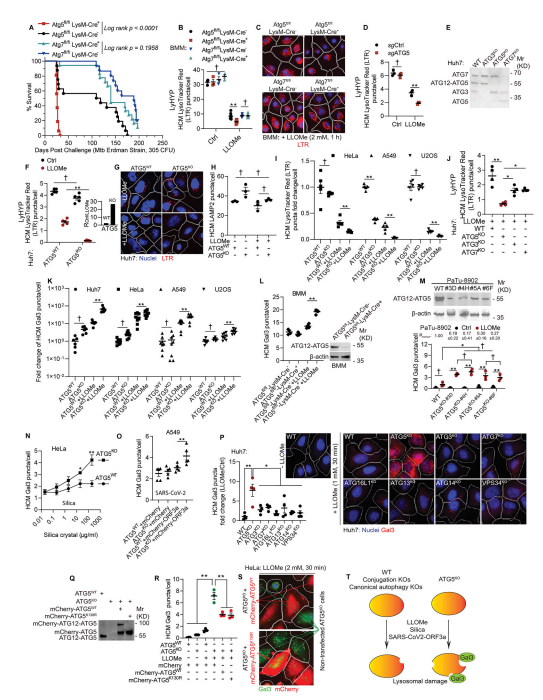
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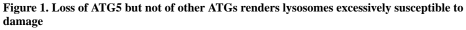
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Highlights

- ATG5 is at a cardinal junction between canonical and noncanonical ATG conjugates
- Loss of ATG5 favors a sidestep conjugate ATG12-ATG3 which sequesters ESCRT ALIX
- Loss of ATG5 causes lysosomal hypersensitivity to damage and increased exocytosis
- The noncanonical role of ATG5 affects neutrophils and tuberculosis pathogenesis





A, Survival curves (mice) post aerosol challenge with Mtb (Erdman; initial lung deposition of 305 CFU). Log-rank (Montel-Cox) test. **B**, Quantification and **C**, representative images of LysoTracker Red (LTR) puncta in bone marrow derived macrophage (BMMs) isolated from Atg5^{fl/fl} LysM-Cre⁻, Atg5^{fl/fl} LysM-Cre⁺, Atg7^{fl/fl} LysM-Cre⁻, Atg7^{fl/fl} LysM-Cre⁺ mice treated with 2 mM of LLOMe for 1 h using High Content Microscopy (HCM). Data, means \pm SE (n 4); unpaired *t*-test. Each data point represents independent biological replicates

with BMMs from different animals. D, HCM quantification of LTR in BMMs subjected to CRISPR inactivation of Atg5 gene (Atg5^{BMM-CRISPR-KO}) compared to Atg5^{BMM-WT} control guide RNA treated cells. Data, means \pm SE (n=3); unpaired *t*-test. **E**, Immunoblots, Huh7 and mutant cells. F, HCM quantification and G, representative image of LTR puncta in Huh7ATG5-WT and Huh7ATG5-KO cells treated with 1 mM LLOMe for 30 min. Data, means ± SE (n=5); two-way ANOVA with Tukey's multiple comparisons. Inset, ratio of LTR puncta in Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells treated:untreated with LLOMe. H, HCM quantification of LAMP2 puncta in Huh7ATG5-WT and Huh7ATG5-KO cells treated with 1 mM LLOMe for 30 min. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. I, HCM quantification of LTR puncta in HeLa (2 mM LLOMe, 30 min), A549 and U2OS (1 mM LLOMe, 30 min) and mutant cells. Data, means \pm SE (n=5); unpaired t-test. J, HCM quantification of LTR puncta in Huh7 and mutant (1 mM LLOMe, 30 min). Data, means \pm SE (n=4); one-way ANOVA with Tukey's multiple comparisons. **K**, HCM quantification of Gal3 puncta in cell lines and under conditions as in panel I. Data, means \pm SE (n=6); two-way ANOVA with Tukey's multiple comparisons. L, HCM quantification of Gal3 puncta in BMMs from Atg5fl/fl LysM-Cre- and Atg5fl/fl LysM-Cre+ mice treated with 2 mM LLOMe for 1 h. Data, means \pm SE (n=4); two-way ANOVA with Tukey's multiple comparisons. Each point represents independent biological replicate from BMMs derived from bone marrows of different animals. M, Immunoblot (top) and HCM quantification (bottom) of Gal3 puncta in PaTu-8902^{WT} and PaTu-8902^{ATG5-KO} cells treated or not with 1.5 mM LLOMe for 30 min. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. N, HCM quantification of Gal3 puncta in HeLa^{ATG5-WT} and HeLa^{ATG5-KO} cells treated with silica. Data, means \pm SE (n=6); unpaired *t*-test. **O**, HCM quantification of Gal3 puncta in A549^{ATG5-WT} and A549^{ATG5-KO} cells overexpressing mCherry or mCherry-ORF3a^{SARS-CoV-2}. Data, means ± SE (n=5); two-way ANOVA with Tukey's multiple comparisons. P, HCM quantification and representative image of Gal3 puncta in Huh7 and mutant cells (1 mM LLOMe, 30 min). Data, means \pm SE (n=4); One-way ANOVA with Tukey's multiple comparisons. **Q**, Immunoblot of Huh7WT and Huh7ATG5-KO cells overexpressing or not overexpressing mCherry-ATG5WT or mCherry-ATG5^{K130R}. **R**, HCM quantification and **S**, representative image of Gal3 puncta in HeLaATG5-WT and HeLaATG5-KO cells control or transfected with mCherry-ATG5WT or mCherry-ATG5^{K130R} (2 mM LLOMe, 30 min). Data, means ± SE (n=3); unpaired *t*-test. T, Summary of the findings in Figure 1. In all HCM graphs, each individual data point represents independent biological replicate, with values based on 49-64 fields/well, >500 primary objects (cells) per well, 5 wells per sample (used only for sampling error) with bars denoting the mean for the biological replicates and standard errors of the mean. Statistical significance symbols in all panels, †p 0.05, *p<0.05, **p<0.01.

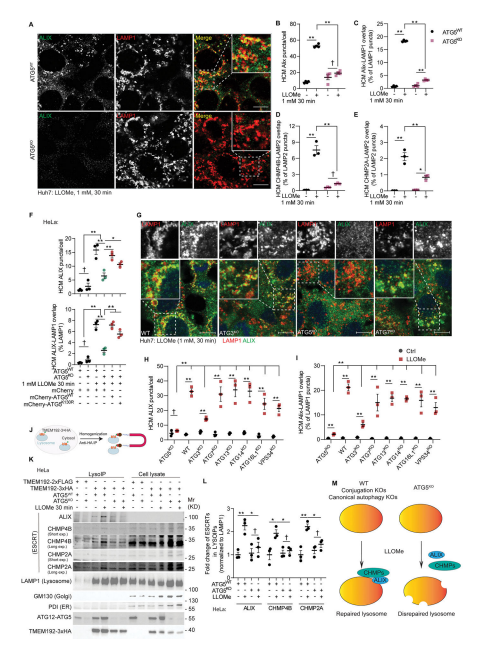


Figure 2. ATG5 is required for recruitment of ESCRTs to repair damaged lysosomes A, Representative confocal images of ALIX and LAMP1 staining in Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells treated with 1 mM LLOMe for 30 min. Scale bars, 10 µm. **B&C**, HCM quantification of ALIX puncta and ALIX-LAMP1 overlap in Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells treated with or without 1 mM LLOMe for 30 min. Data, means \pm SE (n=4); two-way ANOVA with Tukey's multiple comparisons. **D&E**, HCM quantification of CHMP4B-LAMP2 overlap and CHMP4B-LAMP2 overlap in Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells treated as panel B. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **F**, HCM quantification of ALIX puncta and ALIX-LAMP1 overlap in HeLa^{ATG5-WT} and HeLa^{ATG5-KO} cells transfected with mCherry, mCherry-ATG5^{WT}, or mCherry-ATG5^{K130R} followed by treatment with or without 2 mM LLOMe

for 30 min. Data, means \pm SE (n=3); unpaired *t*-test. **G**, Representative confocal images and **H&I**, quantification (HCM) of ALIX puncta and ALIX-LAMP1 overlap in Huh7 and mutant cells treated as in panel B. Data, means \pm SE (n=3); unpaired *t*-test. Scale bars, 10 µm. **J**, Schematic description, **K**, Immunoblot, and **L**, quantification of lysosome ESCRT components by anti-HA immunoprecipitation (LysoIP; TMEM192–3xHA) from HeLa^{ATG5-WT} and HeLa^{ATG5-KO} cells treated as in panel F. Long exp (long exposure) and Short exp (short exposure) correspond to the same blot. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **M**, Summary of the findings in Figure 2. All HCM graphs represent independent biological replicates as in detailed in Figure 1. Statistical significance symbols in all panels, †p 0.05, *p<0.05, **p<0.01.

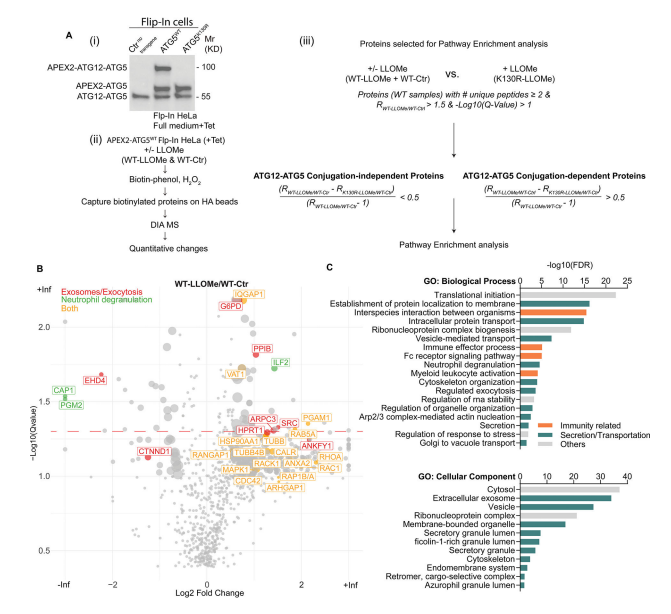


Figure 3. Proximity biotinylation proteomics of ATG5 interactors

• **A**, (i) Stable FlpIn-HeLa^{APEX2-ATG5-WT} or FlpIn-HeLa^{APEX2-ATG5-K130R} cells were (ii) incubated with or without 2 mM LLOMe in full medium supplemented with 2 µg/mL tetracycline for 30 min. Biotinylated proteins were captured by HA beads and subjected to DIA MS. (iii), conjugation-independent or conjugation-dependent interactors of ATG5 were determined by relative fold change of FlpIn-HeLaAPEX2-ATG5-K130R and FlpIn-HeLaAPEX2-ATG5-WT cells treated with LLOMe compared with FlpIn-HeLa^{APEX2-ATG5-WT} cells without LLOMe incubation. **B**, Volcano plot of mass spec result of FlpIn-HeLa^{APEX2-ATG5-WT} incubated with or without LLOMe. **C**, GO terms, pathway enrichment analysis of conjugation-independent proteins.

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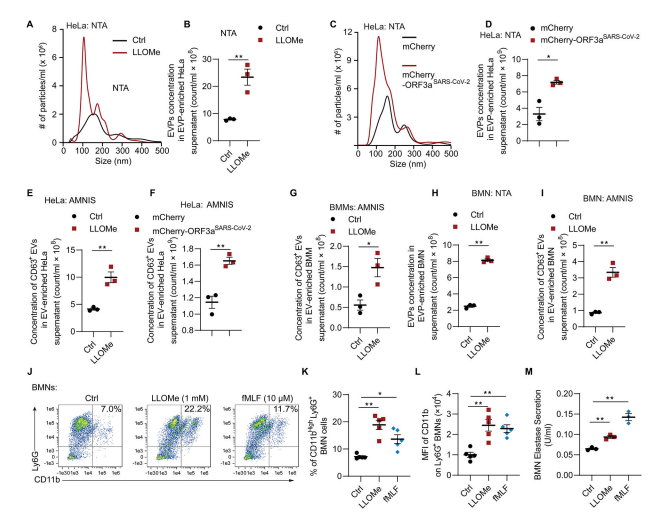


Figure 4. Lysosomal damage induces extracellular vesicle release and neutrophil degranulation A, Representative graph of concentration (y-axis), size distribution (x-axis) and B, quantification of EVPs from supernatant of HeLa cells treated with or without 2 mM LLOMe for 1 h using nanoparticle tracking analysis (NTA). Data, means \pm SE (n=3); unpaired *t*-test. **C**, Representative tracing of concentration, size distribution and **D**, quantification of EVPs from supernatants of HeLa cells expressing mCherry or mCherry-ORF3a (transient transfection for 24 h) using NTA. Data, means \pm SE (n=3); unpaired *t*-test. E, Quantification of CD63⁺ extracellular vesicles from supernatant of HeLa cells treated as in panel A using AMNIS. Data, means \pm SE (n=3); unpaired *t*-test. **F**, Quantification of CD63⁺ extracellular vesicles from supernatant of HeLa cells transfected with mCherry or mCherry-ORF3a for 42 h using AMNIS. Data, means \pm SE (n=3); unpaired *t*-test. G, Quantification of CD63⁺ extracellular vesicles from supernatants of BMMs treated as in panel A using AMNIS. Data, means \pm SE (n=3); unpaired *t*-test. **H**, Quantification of EVPs from supernatant of BMNs (murine bone marrow derived neutrophils) treated with or without 1 mM LLOMe for 1 h using NTA. Data, means \pm SE (n=3); unpaired *t*-test. I, Quantification of CD63⁺ extracellular vesicles extracted from supernatant of BMNs treated as in panel H using AMNIS. Data, means \pm SE (n=3); unpaired *t*-test. J, Representative flow cytometry and K, quantification of % CD11b^{High} Ly6G⁺ BMNs treated with 1 mM

LLOMe for 30 min, or 10µg/ml dihydrocytochalasin B for 5 min followed by 10 µM fMLF for 15 min. Data, means \pm SE (n=5); one-way ANOVA with Tukey's multiple comparisons. **L**, Mean fluorescence intensity (MFI) of CD11b on BMNs treated as in panel J. Data, means \pm SE (n=5); one-way ANOVA with Tukey's multiple comparisons. **M**, Quantification of elastase activity in supernatant of BMNs treated as in panel J. Data, means \pm SE (n=3); one-way ANOVA with Tukey's multiple comparisons. **M**, Quantification of elastase activity in supernatant of BMNs treated as in panel J. Data, means \pm SE (n=3); one-way ANOVA with Tukey's multiple comparisons. In all panels, each data point represents independent biological replicate. For BMM and BMN experiments, each point represents independent biological replicates from different animals. Statistical significance symbols in all panels, †p 0.05, *p<0.05, **p<0.01.

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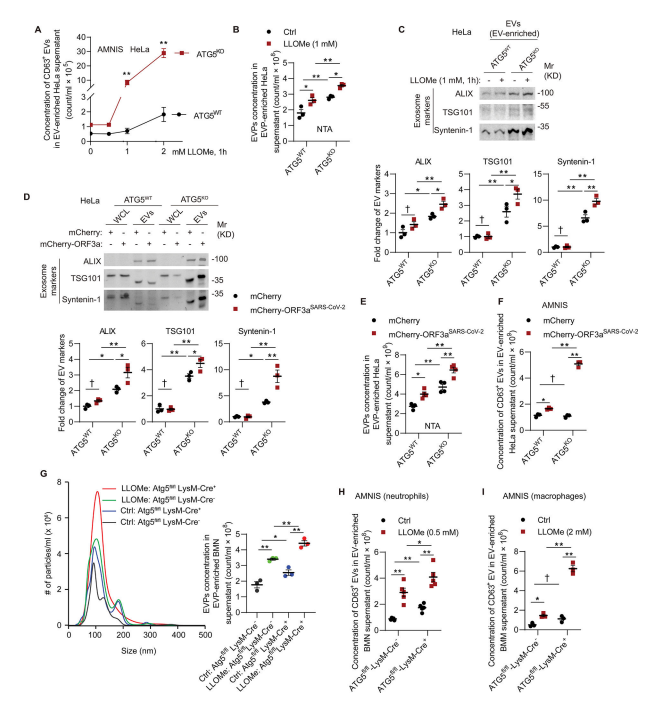


Figure 5. ATG5 knockout enhances EV release in response to lysosomal damage

A, Flow cytometry quantification of CD63⁺ EVs extracted from supernatant of HeLa^{ATG5-WT} and HeLa^{ATG5-KO} cells incubated with indicated concentration of LLOMe for 1 h. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **B**, Quantification of EVPs extracted from supernatant of HeLa^{ATG5-WT} and HeLa^{ATG5-KO} cells treated with or without 2 mM LLOMe for 1 h using NTA. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **C**, Immunoblot and quantification of exosomal markers in EVs isolated from supernatant of HeLa^{ATG5-WT} and HeLa^{ATG5-KO}

cells incubated with 1 mM LLOMe for 1 h. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. D, Immunoblot and quantification of exosomal markers in EVs isolated from supernatant of HeLaATG5-WT and HeLaATG5-KO cells transfected with mCherry or mCherry-ORF3a for 42 h. Data, means ± SE (n=3); two-way ANOVA with Tukey's multiple comparisons. E, Quantification (NTA) of EVPs from supernatant as in panel D. Data, means \pm SE (n=4); two-way ANOVA with Tukey's multiple comparisons. F, Flow cytometry quantification of CD63⁺ EVs extracted from supernatant as in panel D. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **G**, Representative graph of concentration, size distribution and quantification (NTA) of EVPs from supernatant of BMNs from Atg5fl/fl LysM-Cre⁻ and Atg5fl/fl LysM-Cre⁺ mice incubated with or without 0.5 mM LLOMe for 1 h. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. H, Flow cytometry quantification of CD63⁺ EVs from supernatant as in panel G. Data, means \pm SE (n=5); two-way ANOVA with Tukey's multiple comparisons. I, Flow cytometry quantification of CD63⁺ EVs from supernatant of Atg5^{fl/fl} LysM-Cre⁻ and Atg5^{fl/fl} LysM-Cre⁺ BMMs incubated with or without 2 mM LLOMe for 1 h. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. For BMM and BMN experiments, each point represents independent biological replicates from different. Statistical significance symbols in all panels, †p 0.05, *p<0.05, **p<0.01.

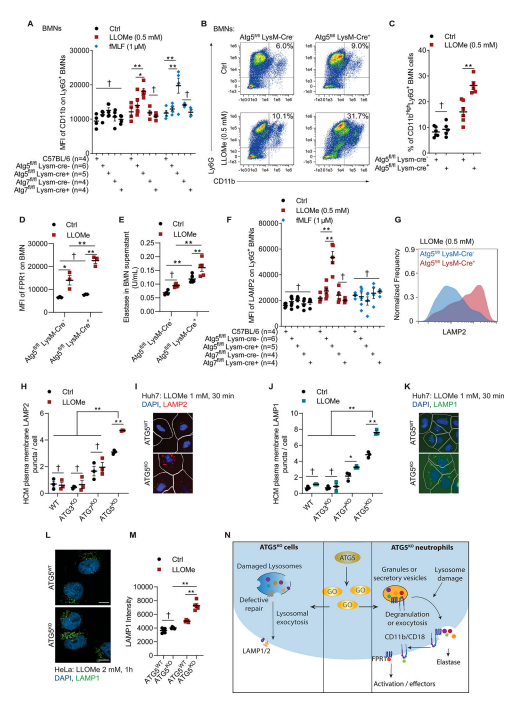


Figure 6. ATG5 knockout enhances PMN degranulation in response to lysosomal damage A, MFI of CD11b on BMNs incubated with 0.5 mM LLOMe for 30 min, or 10µg/ml dihydrocytochalasin B for 5 min followed by 1 µM fMLF for 15 min. Data, means \pm SE (n=4–6); two-way ANOVA with Tukey's multiple comparisons. **B**, Representative flow cytometry of BMNs and **C**, quantification of % CD11b^{High} Ly6G⁺ BMNs from Atg5^{fl/fl} LysM-Cre⁻ and Atg5^{fl/fl} LysM-Cre⁺ mice incubated with or without 0.5 mM LLOMe for 30 min. Data, means \pm SE (n=5–6); two-way ANOVA with Tukey's multiple comparisons. **D**, MFI of FPR1 on BMNs treated as in panel C. Data, means \pm SE (n=3); two-way ANOVA

with Tukey's multiple comparisons. E, Quantification of elastase activity in supernatant of BMNs from Atg5^{fl/fl} LysM-Cre⁻ and Atg5^{fl/fl} LysM-Cre⁺ mice treated as in panel C. Data, means \pm SE (n=4–6); two-way ANOVA with Tukey's multiple comparisons. **F**, MFI of LAMP2 on Atg5^{fl/fl} LysM-Cre⁻ and Atg5^{fl/fl} LysM-Cre⁺ BMNs treated as in panel A. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **G**, Representative flow cytometry of LAMP2 intensity from Atg5^{fl/fl} LysM-Cre⁻ and Atg5^{fl/fl} LysM-Cre⁺ treated as in panel A. H, HCM quantification and I, representative images of plasma membrane LAMP2 puncta on Huh7WT, Huh7ATG3-KO, Huh7ATG7-KO, and Huh7ATG5KO cells treated with or without 1 mM LLOMe for 30 min. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. J, HCM quantification and K, representative images of plasma membrane LAMP1 puncta on Huh7 and mutant cells treated with or without 1 mM LLOMe for 30 min. Data, means \pm SE (n=3); two-way ANOVA with Tukey's multiple comparisons. L, Representative confocal images and M, quantification of plasma membrane LAMP1 puncta on HeLaATG5-WT, and HeLaATG5-KO cells treated with or without 2 mM LLOMe for 30 min. Scale bars, 10 μ m. Data, means \pm SE (n=4); two-way ANOVA with Tukey's multiple comparisons. Fluorescence intensity was calculated based on 20 fields (with a total 100 cells) per biological replicate. N. Summary of data in Figure 6. GO (middle panel), ATG5 interactors identified in proteomic analyses. All HCM graphs represent independent biological replicates as in detailed in Figure 1. For BMMs and BMNs experiments, each point represents independent biological replicates with BMNs from different animals. Statistical significance symbols in all panels, p 0.05, p < 0.05, **p<0.01.

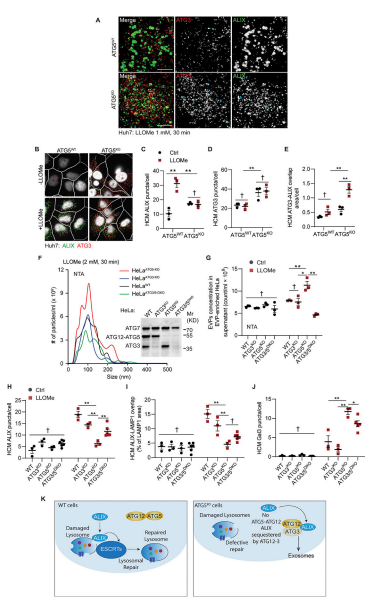


Figure 7. Alternative conjugation contributes to lysosomal vulnerability and exocytic processes A, Representative confocal images of Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells incubated with 1 mM LLOMe for 30 min. Scale bars, 5 μ m. **B-E**, Representative images (**B**) and quantifications of ALIX puncta (**C**), ATG3 puncta (**D**), and ATG3-ALIX overlap area (**E**) in Huh7^{ATG5-WT} and Huh7^{ATG5-KO} cells treated as panel A. Data, means ± SE (n=3); two-way ANOVA with Tukey's multiple comparisons. **F**, Representative graphs and **G**, quantification of EVPs from supernatant of HeLa and mutant cells treated with 2 mM LLOMe for 30 min. Data, means ± SE (n=3); two-way ANOVA with Tukey's multiple comparisons. Inset: Immunoblot of HeLa and mutant cells. **H-J**, HCM quantification of ALIX puncta (**H**), ALIX-LAMP1 overlap area (**I**), and Gal3 puncta (**J**) in Huh7 and mutant cells treated as in panel A. Data, means ± SE (n 3); two-way ANOVA with Tukey's multiple comparisons. **K**, Summary of the mechanistic findings in Figure 7. All HCM graphs represent independent

biological replicates as in detailed in Figure 1. Statistical significance symbols in all panels, †p 0.05, *p<0.05, *p<0.01.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit Anti-APG5L/ATG5 (EPR1755(2))	Abcam	ab108327
Rabbit Anti-ATG7 (EP1759Y)	Abcam	ab52472
Rabbit Anti-mCherry	Abcam	ab183628
Rabbit Anti-Syntenin-1	Abcam	ab19903
Rabbit Anti-TSG101 (4A10)	Abcam	ab83
Rabbit Anti-VPS34 - N-terminal	Abcam	ab124905
Mouse Anti-ALIX	BioLegend	#634502
Mouse Anti-Galectin-3	BioLegend	#125402
Rabbit ATG13(E1Y9V)	Cell Signaling Technology	#13468
Rabbit Gm130 (D6B1)	Cell Signaling Technology	#12480S
Rabbit HA-Tag (C29F4)	Cell Signaling Technology	#3724
Rabbit Lamp1 (D2D11)	Cell Signaling Technology	#9091S
Rabbit Pdi (C81H6)	Cell Signaling Technology	#3501S
Mouse LAMP2	DSHB of University of Iowa	H4B4
Rabbit FPR1 Antibody FITC	Fabgennix	FPR1-FITC
Rabbit Anti-Atg14	MBL International	PD026
Rabbit Anti-ATG16L1	MBL International	PM040
Rabbit CD11b	Novus Biological	NB110-89474
Rabbit Alix	Proteintech	12422-1-AP
Rabbit CHMP2A	Proteintech	10477-1-AP
Rabbit CHMP4B	Proteintech	13683-1-AP
Rabbit beta-Actin (C4)	Santa Cruz Biotechnology	sc-47778
Mouse Anti-cathepsin C (D-6)	Santa Cruz Biotechnology	sc-74590
Mouse Anti-GAPDH (0411)	Santa Cruz Biotechnology	sc-47724
Rabbit Anti-ATG3	Sigma Aldrich	#A3231
Rabbit Anti-CD63	Sigma Aldrich	SAB4301607
Rat Ly-6G (Clone 1A8)	STEMCELL Technologies	#60031
Mouse ATG12	ThermoFisher	366400
Alexa Fluor 488 Goat anti-Rabbit secondary antibody	ThermoFisher	#A-11034
Alexa Fluor 488 Goat anti-Mouse secondary antibody	ThermoFisher	#A-11029
Alexa Fluor 568 Goat anti-Rabbit secondary antibody	ThermoFisher	#A-11011
Alexa Fluor 568 Goat anti-Mouse secondary antibody	ThermoFisher	#A-11004
Alexa Fluor 647 Goat anti-Rabbit secondary antibody	ThermoFisher	#A27040
Alexa Fluor 647 Goat anti-Rat secondary antibody	ThermoFisher	#A-21247
Goat anti-rabbit IgG-HRP secondary antibody	ThermoFisher	#31460
Goat anti-mouse IgG-HRP secondary antibody	ThermoFisher	#31430

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Goat Anti-Rabbit IgG Polyclonal Antibody (IRDye® 800CW)	LI-COR Biosciences	926-32211
Goat Anti-Mouse IgG Polyclonal Antibody (IRDye® 680RD),	LI-COR Biosciences	926-68070
Bacterial and virus strains		
Mycobacterium tuberculosis Erdman	Manzanillo et al. 124	PMID: 22607800
NEB 5-alpha Competent <i>E.coli</i> (High Efficiency)	New England Biolabs	C2987
One Shot Mach1 Phage-Resistant Competent E.coli	ThermoFisher	C862003
Chemicals, peptides, and recombinant proteins		
Leu-Leu-methyl ester hydrobromide (LLOMe)	Sigma Aldrich	L7393
Pierce [™] Streptavidin Magnetic Beads	ThermoFisher	88816
Tetracycline hydrochloride	Sigma Aldrich	T3383
Puromycin dihydrochloride	Sigma Aldrich	P9620
Hygromycin B	Sigma Aldrich	H0654
Silica crystal	US Silica	MIN-U-SIL-15
mouse macrophage colony stimulating factor (mM-CSF)	Cell Signaling Technology	5228
Hoechst 33342	ThermoFisher	H3570
DAPI	Sigma Aldrich	D9542
Prolong Gold Antifade Mountant with DAPI	ThermoFisher	P36931
Polybrene	Sigma Aldrich	TR-1003-G
LysoTracker [™] Red DND-99	ThermoFisher	L7528
Biotinyl tyramide	AdiopoGen Life Sciences	CDX-B0270-M500
X-tremeGENE [™] HP DNA Transfection Reagent	Sigma Aldrich	6366236001
Histopaque [®] -1119	Sigma Aldrich	11191
Histopaque [®] -1077	Sigma Aldrich	10771
N-Formyl-Met-Leu-Phe	Sigma Aldrich	47729-10MG-F
LR Clonase Plus Enzyme Mix	ThermoFisher	11791100
BP Clonase Plus Enzyme Mix	ThermoFisher	11789100
Nuclease-Free Duplex Buffer	Integrated DNA Technologies	11-05-01-03
Alt-R [®] S.p. Cas9 Nuclease V3	Integrated DNA Technologies	1081059
Detachin	Genlantis	T00100
Sodium ascorbate	Sigma Aldrich	A7631
Sodium azide	Sigma Aldrich	S2002
Trolox	Sigma Aldrich	238813
Dextran, Texas Red [™] , 3000 MW, Lysine Fixable	ThermoFisher	D3328
Critical commercial assays		
QuikChange II Site-Directed Mutagenesis Kit	Agilent	200523
TidyBlot	BioRad	STAR209
P3 Primary Cell 4D-Nucleofector® X Kit L	Lonza	V4XP-3024
SmartSEC [™] Single for EV Isolation	SBI System Biosciences	SSEC200A-1

REAGENT or RESOURCE	SOURCE	IDENTIFIER
ExoQuick-TC	SBI System Biosciences	EXOTC10A-1
Gateway Vector Conversion System	ThermoFisher	11828-029
Lipofectamine 2000 Transfection Reagent	ThermoFisher	12566014
Molecular Probes EnzChek Elastase Assay Kit	ThermoFisher	E12056
Deposited data		
Raw MS DIA data	https://massive.ucsd.edu	MSV000090348
Raw MS DIA data	http://www.proteomexchange.org	Proteome Exchange = PXD03685
Experimental models: Cell lines		
HeLa WT	ATCC	CCL-2
HeLa ATG5 ^{KO}	This paper	N/A
HeLa ATG3 ^{KO}	This paper	N/A
HeLa ATG3/5 ^{DKO}	This paper	N/A
Huh7 WT	Rocky Mountain laboratories	N/A
Huh7 ATG5 ^{KO}	This paper	N/A
Huh7 ATG3 ^{KO}	This paper	N/A
Huh7 ATG3/5 ^{DKO}	This paper	N/A
Huh7 ATG7 ^{KO}	This paper	N/A
Huh7 ATG13 ^{KO}	This paper	N/A
Huh7 ATG14 ^{KO}	This paper	N/A
Huh7 ATG16L1 ^{KO}	This paper	N/A
Huh7 VPS34 ^{KO}	This paper	N/A
A549 WT	ATCC	CCL-185
A549 ATG5 ^{KO}	This paper	N/A
U2OS WT	ATCC	HTB-96
U2OS ATG5 ^{KO}	This paper	N/A
HEK293T	ATCC	CRL-3216
HeLa WT-TMEM192-3XHA	This paper	N/A
HeLa ATG5 ^{KO} -TMEM192-3XHA	This paper	N/A
HeLa WT-TMEM192-2XFLAG	This paper	N/A
HeLa ATG5 ^{KO} -TMEM192-2XFLAG	This paper	N/A
Flp-In-HeLa APEX2-ATG5 ^{WT}	This paper	N/A
Flp-In-HeLa APEX2-ATG5 ^{K130R}	This paper	N/A
HL-60 WT	This paper	N/A
HL-60 ATG5 ^{KO}	This paper	N/A
Patu-8902 ^{WT}	Alec Kimmelman	N/A
Patu-8902 ^{ATG5-KO}	This paper	N/A
Experimental models: Organisms/strains		
C57BL/6	The Jackson Laboratory	Strain #000664

REAGENT or RESOURCE	SOURCE	IDENTIFIER
ATG5 ^{fl/fl} LysM-Cre-negative mice	Castillo et al. ⁵¹	PMID: 23040807
ATG5 ^{fl/fl} LysM-Cre-positive mice	Castillo et al. ⁵¹	PMID: 23040807
ATG7 ^{fl/fl} LysM-Cre-negative mice	Komatsu et al. ¹²⁵	PMID: 15866887
ATG7 ^{fl/fl} LysM-Cre-positive mice	Komatsu et al. ¹²⁵	PMID: 15866887
Oligonucleotides		
ATG5 CRISPR gRNA: AAGAGTAAGTTATTTGACGT	VectorBuilder	gRNA#1544
ATG3 CRISPR gRNA: GTGAAGGCATACCTACCAAC	VectorBuilder	gRNA#1119
ATG7 CRISPR gRNA: CTTCCGTGACCGTACCATGC	VectorBuilder	gRNA#6556
ATG13 CRISPR gRNA: CTGTCCCAACACGAACTGTC	VectorBuilder	gRNA#4570
ATG14 CRISPR gRNA: CTACTTCGACGGCCGCGACC	VectorBuilder	gRNA#58
ATG16L1 CRISPR gRNA: ACCAAATGCAGCGGAAGGAC	VectorBuilder	gRNA#1540
VPS34 CRISPR gRNA: ACAGACATTTGTAGATCGGT	VectorBuilder	gRNA#5287
ATG5 primers for gateway cloning: Forward: GGGGACAAGTTTGTACAAAAAAGCAGGCTTCACAGA TGACAAAGATGTGCTTCGAG Reverse: GGGG-AC-CAC-TTT-GTA-CAA-GAA-AGC- TGG-GTC-TCAATCTGTTGGCTGTGGGATG	Integrated DNA Technologies	N/A
ATG5 primers for K130R mutation: Sense: 5'- GCAATTGAAGCTCATTTTATGTCATGTATGAGAGAAG CTGATGCTTTA-3' Antisense: 5'- TAAAGCATCAGCTTCTCTCATACATGACATAAAATGAG CTTCAATTGC-3'	Integrated DNA Technologies	N/A
Alt-R [®] CRISPR-Cas9 guide RNA for ATG5 sgRNA1: TCCATCCAAGGATGCGGTTG sgRNA2: AAGAGTCAGCTATTTGACGT	Integrated DNA Technologies	Mm.Cas9.ATG5.1.AA Mm.Cas9.ATG5.1.AB
Recombinant DNA		
pDEST-Flp-In-APEX2-ATG5 ^{WT}	This paper	N/A
pDEST-Flp-In-APEX2-ATG5 ^{K130R}	This paper	N/A
pDEST-mCherry-ATG5 ^{WT}	This paper	N/A
pDEST-mCherry-ATG5K130R	This paper	N/A
pDEST-mCherry-SARS-CoV-2 ^{ORF3a}	This paper	N/A
pCMV-VSV-G	Addgene	#8454
psPAX2	Addgene	#12260
pOG44	This paper	N/A
pLJC5-TMEM192-3xHA	Addgene	#102930
pLJC5-TMEM192-2xFLAG	Addgene	#102929
Software and algorithms		
iDEV software	ThermoFisher	N/A
Zen 3.4	Carl Zeiss	N/A
Spectronaut software	Biognosys Inc	N/A
Prism 8	GraphPad	https://www.graphpad.com/
Image Lab	BIO-RAD	N/A

REAGENT or RESOURCE	SOURCE	IDENTIFIER
IDEAS 6.2	Luminex	N/A
ImageJ	NIH	https://imagej.nih.gov/ij/
NTA 3.3	Malvern Panalytical	N/A
R (v4.1.2)	The R Foundation	https://www.r-project.org
Other		
NP40 Cell Lysis Buffer	ThermoFisher	FNN0021
Protease Inhibitor Cocktail Tablets	Roche	11697498001
PMSF	Sigma Aldrich	93482
Anti-HA Magnetic Beads	ThermoFisher	88836
Dynabeads Protein G	ThermoFisher	10003D