**Retrofusion of intralumenal MVB membranes parallels viral infection and coexists with exosome release**

### Highlights

- MVBs are complex organelles with intraluminal vesicles bound by the limiting membrane
- Intraluminal membranes are in a dynamic equilibrium with the limiting membrane
- Retrofusion of internal vesicles is controlled by processes used for viral fusion
- Exosomes arise from internal MVB vesicles not participating in retrofusion

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### In brief

Multivesicular bodies (MVBs) are complex organelles harboring internal vesicles. Using a chemically controlled system, Perrin et al. visualize a part of these internal vesicles fusing back to the limiting membrane. This process of retrofusion exists in equilibrium with lysosomal degradation and exosome release and is inhibited by antiviral proteins.

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Retrofusion of intraluminal MVB membranes parallels viral infection and coexists with exosome release

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SUMMARY

The endosomal system constitutes a highly dynamic vesicle network used to relay materials and signals between the cell and its environment.1 Once internalized, endosomes gradually mature into late acidic compartments and acquire a multivesicular body (MVB) organization through invagination of the limiting membrane (LM) to form intraluminal vesicles (ILVs).2 Cargoes sequestered into ILVs can either be delivered to lysosomes for degradation or secreted following fusion of the MVB with the plasma membrane.3 It has been speculated that commitment to ILVs is not a terminal event, and that a return pathway exists, allowing “back-fusion” or “retrofusion” of intraluminal membranes to the LM.4 The existence of retrofusion as a way to support membrane equilibrium within the MVB has been widely speculated in various cell biological contexts, including exosome uptake5 and major histocompatibility complex class II (MHC class II) antigen presentation.6–9 Given the small physical scale, retrofusion of ILVs cannot be measured with conventional techniques. To circumvent this, we designed a chemically tunable cell-based system to monitor retrofusion in real time. Using this system, we demonstrate that retrofusion occurs as part of the natural MVB lifestyle, with attributes parallel to those of viral infection. Furthermore, we find that retrofusion and exocytosis coexist in an equilibrium, implying that ILVs inert to retrofusion comprise a significant fraction of exosomes destined for secretion. MVBs thus contain three types of ILVs: those committed to lysosomal degradation, those retrofusing ILVs, and those subject to secretion in the form of exosomes.

RESULTS AND DISCUSSION

To address the challenge of observing retrofusion in real time, we designed a chemically tunable reporter system to visualize and monitor the occurrence of this process in living cells. To follow relocation of membrane components from the intraluminal vesicles (ILVs) back to the limiting membrane (LM), we furnished the canonical multivesicular body (MVB) marker tetraspanin membrane protein CD63 with a GFP tag harboring an N-terminal nuclear localization signal (NLS) and a C-terminal tobacco etch virus (TEV) protease-specific cleavage site (TCS), resulting in NLS-GFP-TCS-CD63 stably expressed in MelJuSo cells, henceforth referred to as GFP-CD63 (Figures 1A and 1B). The TEV protease was introduced into the same cells as two inactive parts, coexpressed along with NLS-DsRED from a single polycistronic vector (Figures 1A and 1B), which can be brought together on demand by a rapamycin analog termed “dimerizer.”10 Upon addition of dimerizer, protease activity would be reconstituted and result in cleavage of available GFP-CD63 (i.e., GFP exposed to the cytosol, unlike the GFP in ILVs). As a consequence, CD63 at the LM of MVBs would lose its GFP fluorescence, and newly liberated NLS-GFP would be targeted to the nucleus, averting accumulation of cytosolic background (Figures 1A and 1B).

In the absence of dimerizer, GFP-CD63 localized to both ILVs and the LM of MVBs at a 2:1 ratio (Figures 1C and 1D), mirroring the distribution of endogenous CD63,11 whereas the split protease remained inactive (Figure 1B), and no nuclear GFP was observed (Figure 2A; t = 0). GFP-CD63 was also present at the cell surface (Figure 2A). Upon addition of dimerizer to the cells, and consequent reconstitution of the TEV protease (Figure 1B), GFP fluorescence in acidic vesicles of the endocytic tract marked by SiR-lysosome began to diminish over time, accompanied by a concomitant rise of GFP signal in the nucleus (Figure 2A; Video S1). Because GFP in the nucleus can arise from cleaved CD63 molecules in endosomes as well as from the cell surface, we directly measured the decay of GFP fluorescence in the late endosomes. To correct for background fluorescence, cytoplasmic GFP was subtracted from the SiR-lysosome-positive GFP signal,
yielding normalized fluorescence intensity associated with endolysosomal organelles over time (Figures 2B and S1A; Video S2). Following the initial equilibration phase (Figures 2B and S1A; t = 0–90 min), a near-linear decay of GFP signal from SiR-lysosome-positive compartments was observed (t = 90–360 min) and its slope A was assessed (Figures 2A, 2B, and 2C; Video S1). This decay was contingent on TEV protease activity, as evidenced by no appreciable loss of endolysosomal GFP (or rise in nuclear GFP) in the absence of dimerizer treatment (Figures 2A, 2B, and 2C; Video S3).

Because only GFP exposed on the cytosolic side of the LM can be accessible for TEV cleavage, we expected that, in the absence of retrofusion, dimerizer-induced decay of endolysosomal GFP fluorescence would plateau at roughly 66%–70%, corresponding to the amount of intraluminal GFP-CD63 (Figure 1D). Further loss of this GFP signal would necessitate repopulation of the LM by GFP-CD63 derived from retrofusing ILVs. Inspection of the GFP signal decay following the system’s engagement with dimerizer (Figures 2B and S1A) revealed that 41% ± 5% of the starting GFP signal associated with endolysosomes remained at 6 h post treatment (Figures 2B and 2D), implying that a proportion of GFP signal lost from endocytic compartments depends on the occurrence of retrofusion. Indeed, electron microscopy (EM) analysis following 6-h incubation with dimerizer and immunogold labeling against both GFP and CD63 demonstrated a decrease in GFP staining relative...
Figure 2. Monitoring retrofusion of GFP-CD63-positive ILVs in real time

(A) Representative confocal fluorescence stills from a time-lapse experiment with GFP-CD63 cells (white) imaged in the presence (+) or absence (−) of dimerizer along with color overlays of GFP (green) with SiR-lysosome (red) marking ELs are shown. DsRED-positive (magenta) nuclei indicate expression of the split TEV protease at t = 0. Scale bars, 10 μm.

(B) Representative plot of EL-associated median GFP fluorescence intensity over time (min), corrected by cytoplasmic GFP background subtraction; (+), dimerizer (gray); (−), dimerizer (red) treatment. Slope A characterizes the near-linear GFP decay. Only loss of GFP signal in MVBs is quantified in our experiments as nuclear NLS-GFP is also provided by GFP-CD63 molecules at the cell surface.

(C) Quantification of slope A relative to (+) dimerizer. Shown is mean ± SD from n = 7 independent experiments for (−) dimerizer and n = 35 for (+) dimerizer.

(D) Quantification of GFP fluorescence remaining in late endosomes at 6 h after dimerizer addition, expressed as % GFP fluorescence at t = 0. Shown is mean ± SD from n = 23 independent experiments.

(legend continued on next page)
to that of CD63 on both the LM and intraluminal membranes (ILM) (Figures 2E and 2F), confirming intraluminal GFP recovery. Of note, dimerizer treatment did not affect the distribution of CD63 (ILM versus LM) in MVBs (Figure S1B). Past the 6-h time point, the decay of endolysosomal GFP began to plateau (Figures 2B and S1A) and, by 9 h post dimerizer addition, 20% of the initial fluorescence intensity still remained in MVBs. This suggests that a significant fraction of GFP-CD63-positive ILVs does not participate in retrofusion and is likely subject to other pathways.

Because MVBs continuously receive input of new membranes and sort cargoes toward diverse fates, notably including lysosomal proteolysis, we tested whether biosynthesis or degradation of GFP-CD63 contributes to the GFP decay measured during the time frame of our microscopy assay. A pulse-chase experiment with 1-deoxymannosidase I (DMM), an inhibitor of mannosidase I,12 was used to assess the relative proportion of old GFP-CD63 bearing long polyglycosylated glycans versus newly synthesized GFP-CD63 carrying high-mannose N-linked glycans (migrating at a lower molecular weight on the gel) over time (Figure 2G), yielding a half-life longer than 24 h (Figure 2H), which was expected because CD63 is known to be highly stable.13 These data indicate that the natural turnover of GFP-CD63 does not significantly contribute to the decay of endolysosomal GFP fluorescence following dimerizer addition.

Because the readout of our system is contingent on the performance of the TEV protease, we examined whether the decay of GFP-CD63 is limited by the rate of TEV-mediated cleavage. To this end, the cytoplasmic tail of the lysosomal membrane protein LAMP1, which localizes almost exclusively to the LM of MVBs,14 was fused to a TEV cleavage site followed by mTurquoise2-NLS, and the resulting LAMP1-TEV-TQ (turquoise) construct was stably coexpressed with the TEV protease and GFP-CD63. Following addition of dimerizer, the mTurquoise2 signal associated with MVBs decreased more rapidly and reached a lower end value than its GFP-CD63 counterpart expressed in the same cells (Figures 2I and S1C), indicating that inaccessibility of ILV-localized GFP-CD63 to the TEV protease, rather than TEV cleavage efficacy, is the limiting factor in our measurements.

MVBs are acidic organelles, which is fundamental to their proteolytic function and also, intriguingly, crucial to fusion of many viruses with the LM.15 To test whether retrofusion of ILVs also benefits from the acidic environment of the MVB, endolysosomal pH was elevated using bafilomycin A, a V-ATPase inhibitor pretreated with MVBs decreased more rapidly and reached a lower end value than its GFP-CD63 counterpart expressed in the same cells (Figures 2I and S1C), indicating that inaccessibility of ILV-localized GFP-CD63 to the TEV protease, rather than TEV cleavage efficacy, is the limiting factor in our measurements. retrofusion. Once again, no significant effect on the decomposition of GFP-CD63 in MVBs, nor the intraluminal content, was observed in the presence of thioperamide, implying that the resulting retrofusion attenuation cannot be attributed to lack of intraluminal GFP-CD63 (Figures 1C, 1D, 3D, 3E, and S2C). In the case of U18666A, a slight increase in intraluminal GFP-CD63 was observed, which may be attributed to attenuation of retrofusion, although lipid alterations might also affect the formation of ILVs. Based on this evidence, we propose that lipid composition of ILVs is crucial for recycling of the ILM to the LM. Because LBPA and cholesterol have both been implicated in viral infection,22–24 these results again hint at fundamental parallels between endosomal equilibrium and viral escape, explored below.

The first experimental suggestions regarding the existence of retrofusion were made, albeit indirectly, in the context of viral infection.24 and it has been proposed that some viruses hijack the retrofusion pathway to escape lysosomal degradation and

(E) Electron micrograph featuring immunogold labeling with GFP10 nm and CD6315 nm gold antibodies of GFP-CD63-expressing cells in the absence or 6 h in the presence of dimerizer. Scale bars, 100 nm.
(F) Quantification of GFP-CD63 and untagged CD63 abundance on ILVs and the LM in the presence or absence of dimerizer, expressed as the ratio of GFP/CD63 (GFP and CD63, respectively, representing the sum of GFP and CD63 in all MVBs) assessed by immunogold labeling. Shown is mean ± SD from over 25 MVBs from n = 4 independent experiments.
(G) Immunoblot analysis of GFP-CD63 turnover. The mannosidase I inhibitor DMM is added to the cells at t = 0 h and cells are grown for the times indicated followed by western blot (WB) analyses and staining with GFP antibody. DMM yields CD63 molecules with high-mannose glycans (HM) that run at a lower position than mature CD63, as indicated. The position of marker proteins is indicated.
(H) Quantification of mature GFP-CD63 signal normalized to total GFP (sum of mature and HM GFP-CD63 signal) and normalized to t=0 h at each time point following DMM addition. Shown is mean ± SD from n = 4 independent experiments.
(I) Quantification of slope A of GFP-CD63 relative to TQ-Lamp1 decay following dimerizer addition. Shown is mean ± SD for n = 7 independent experiments. For primary data and cell images, see Figures S1A and S1B.

Statistical differences between the groups were assessed using unpaired Student’s t test (C and I) or multiple t test (F) (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001). See also Figure S1 and Videos S1, S2, and S3.
Figure 3. Alterations in MVB pH and lipid repertoire hamper ILV retrofusion

(A) Scheme summarizing the different treatments used to modulate membrane composition and dynamics within the MVB: (1) bafilomycin A (Baf A) blocks acidification; and (2) thioperamide and U18666A lead to accumulation of LBPA and cholesterol in MVBs, respectively.

(B) Effect(s) of Baf A and thioperamide treatments on the rate of ILV retrofusion. Representative confocal images of GFP-CD63 (white) distribution before (t = 0 h) and after (t = 6 h) treatment with dimerizer are shown, along with color overlays of GFP (green) with SiR-lysosome (red) in retrofusion-monitoring cells. DsRED-positive (magenta) nuclei indicate expression of the split TEV protease at t = 0. Scale bars, 10 μm.

(C) Quantification of slope A (paired t test) and the percentage of remaining fluorescence intensity at 6 h for the indicated conditions normalized to control. Shown is mean ± SD; n = 5 independent experiments for Baf A, n = 4 for thioperamide, and n = 6 for U18666A. See also Figure S2.

(D) Quantification of GFP-CD63 as detected by immunogold labeling on ILVs relative to the LM (expressed as ratio per MVB) following incubation with U18666A, thioperamide, or Baf A. Shown are the relative results from n ≥ 98 MVBs from n = 2 independent experiments.

(E) Quantification of MVB structure (normal, lightly swollen, or swollen) as expressed as a percentage of MVBs for each condition, U18666A, thioperamide, or Baf A. Shown are the relative results from n = 98 MVBs from n = 2 independent experiments.

Statistical differences between the groups were assessed using paired or unpaired Student’s t test (C), or Mann-Whitney test (D) (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, ns = not significant). See also Figure S2.
access the host’s cytoplasm. Furthermore, type I interferons, induced upon viral infection, can stimulate expression of interferon-induced transmembrane proteins (IFITMs) that block viral escape from endosomes. Specifically, IFITM3 expression has been shown to promote accumulation of cholesterol in MVBs of infected cells, hence impeding retrofusion of vesicular stomatitis virus (VSV)-positive ILVs. We therefore considered whether stable introduction of TQ-IFITM1–3 (Figure 4A), all of which colocalized with GFP-CD63 in endolysosomal compartments (Figures 3 and 4), would affect the rate of constitutive retrofusion. GFP fluorescence decay was reduced by ~50% in the presence of TQ-IFITM3 (Figures 4C–4E) but was affected to a lesser extent by IFITM1 and IFITM2 (Figures 4E, S3A, and S3C). This difference is consistent with previous findings showing that IFITM3 expression restricts viral access to the host’s cytoplasm more efficiently than expression of IFITM1 or 2. It has been proposed that IFITM3 blocks viral membrane hemifusion by affecting membrane curvature required during viral fusion. Our data suggest that IFITM3 may hamper ILV retrofusion through a similar mechanism. Taken together with the effects of endosomal pH and ILV lipid content on retrofusion (Figure 3), the above observations reveal mechanistic parallels between ILV retrofusion and viral fusion with endosomal membranes and lend experimental support to the notion that some viruses exploit the former pathway to further their infection program, whereas certain antiviral host proteins can inhibit it at the cost of attenuated retrofusion.

Depending on their cargoes, ILVs can be expelled into extracellular space following fusion of the LM with the plasma membrane. These extracellular vesicles, or exosomes, can transfer ILV contents between cells; exosome transmission has been implicated in the pathogenesis of various diseases, including tumor metastasis and dissemination of prion-like proteins; implicated in the pathogenesis of various diseases, including tumor metastasis and dissemination of prion-like proteins; and retrofusion has been proposed as a key step in exosome up-regulation of treatment with dimerizer (Figure 4F), implying that at least a third of exosomes come from dynamic ILVs (which have lost their associated GFP), whereas the majority derive from inert ILVs (harboring intact GFP-CD63 not accessible to the cytosolic activated TEV protease during the harvesting period). To explore the existence of an equilibrium between these populations, we tested whether inhibiting retrofusion would promote exocytosis of affected ILVs. To this end, exosomes were isolated from the medium of control cells versus those ectopically expressing TQ-IFITM3 treated either in the presence or absence of dimerizer. In the case of the TQ-IFITM3-expressing cells, the abundance of GFP-CD63 in

Figure 4. Activation of antiviral response genes attenuates retrofusion of ILVs and renders a fraction of dynamic exosomes inert

(A) Schematic representation of the fusion of viruses with the LM of late endosomes and the potential function of IFITM proteins in restricting fusion and thus infection of host cells.

(B) Confocal images showing the localization of TQ-IFITM3 to late endocytic compartments. Fluorescence overlays of TQ-IFITM3 (blue) with GFP-CD63 (green) are shown, as indicated. Scale bar, 10 μm.

(C) Effect(s) of IFITM3 overexpression on the rate of ILV retrofusion. Representative confocal images of GFP-CD63 (white) distribution before (t = 0 h) and after (t = 6 h) treatment with dimerizer are shown, along with color overlays of GFP (green) with SiR-lysosome (red) in retrofusion-monitoring GFP-CD63 cells. DsRED-positive (magenta) nuclei indicate expression of the split TEV protease at t = 0. Scale bars, 10 μm.

(D) Representative plots of normalized late-endosome-associated median GFP fluorescence intensity over time (min) following dimerizer addition in control cells (gray line) versus those overexpressing TQ-IFITM3 (red line).

(E) Quantification of slope A for the indicated conditions normalized to control cells in the same experiment. Shown is mean ± SD from n = 6 independent experiments for TQ-IFITM3, n = 6 for TQ-IFITM2, and n = 5 for TQ-IFITM1. Statistical differences between each condition and its control were assessed using paired Student’s t test and those between different conditions using unpaired Student’s t test. See also Figure S3.

(F) Workflow for exosome isolation from retrofusion-monitoring cells.

(G) Quantification of the ratio of GFP over total CD63 signal in exosome fractions (left graph) and WCL (right graph) prepared following incubation of control cells or cells overexpressing TQ-IFITM3 and cultured in the presence (+) or absence (−) of dimerizer for 9 h. Shown is mean ± SD normalized to control from n > 5 independent experiments.

(H) Quantification of the percentage of GFP-CD63 secreted in exosomes relative to the total cell lysate where GFP-CD63 is expressed in MVBs and the plasma membrane. The relative contribution of GFP-CD63 fluorescence in MVBs was determined by 3D reconstitution of GFP fluorescence in cells (see Video S4). GFP-CD63 in exosomes was compared in a dilution series of GFP labeling as detected by WB analyses and the ratio of GFP fluorescence in the total cell versus MVB was determined by microscopy. Shown is the result of n = 3 independent exosome versus total lysate isolations assessed by WB. Shown is mean ± SD. Statistical differences between groups were assessed using unpaired Student’s t test unless otherwise indicated (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, ns = not significant). See also Figures S3 and S4 and Video S4.
exosome isolates was normalized to both total (untagged) CD63 and TG-IFITM3 in the same samples. Only a 10% decrease in the ratio of GFP-positive over CD63/IFITM3-positive exosomes was noted upon dimerizer treatment (Figure 4G and S4C), suggesting that IFITM3-mediated block of retrofusion renders at least a fraction of dynamic exosomes inert. Because our findings indicate that most exosomes are inert to retrofusion prior to exocytosis, it begs the questions of how retrofusion is controlled and what determines the fate of ILVs. Cytosolic factors, such as the endosomal sorting complexes required for transport (ESCRT)-associated protein Alix,36 may contribute to intra-endosomal membrane dynamics to some extent, but cargo could also play an important role in dictating the ultimate ILV destination. Although retrofusion and exosome formation have some relationship, the exact molecular details still need to be determined in future studies.

Finally, we sought to exploit our chemically inducible cell-based system to assess the relative contribution of different fates experienced by ILVs at steady state (retrofusion, secretion, degradation). The numbers, however, represent a rough approximation as we follow the bulk of endosomes whereas particular endosomes may be more specialized in a specific pathway. Exosomes released over a 6-h period were isolated from cells and the amount of GFP-CD63 in exosomes was related to GFP-CD63 in total cell lysates (6%; Figures 4H and S4D–S4F), and further corrected for the distribution of GFP-CD63 in endosomes versus the plasma membrane (62% in MVB; Video S4). Around 10% of GFP-CD63 ended up in exosomes (Figure 4H). Along with pulse-chase experiment results (Figures 2G and 2H), we calculated the percentage of GFP-CD63 (from whole-cell lysate [WCL] or MVBs) that is either degraded in lysosomes or secreted in exosomes (Figure S4G). Considering these numbers, at least 11% of ILVs may participate in retrofusion in the 6 h of detection (Figure S4H). This number is likely higher because the TEV protease is not modifying every GFP-CD63 exposed to the cytosol. How individual ILVs differ to end up in one of the three different pathways is unclear, but is likely influenced by the type of substrate involved (e.g., epidermal growth factor receptor [EGFR] degradation upon EGF binding).37

In conclusion, the development and application of the first chemically tunable system to visualize and quantify the rate of ILV retrofusion in living cells have enabled us to show that constitutive retrofusion is a dynamic process occurring as part of the normal MVB lifecycle in unperturbed cells. However, only a part of the ILV population is able to fuse back with the limiting membrane of the MVB. In other words, MVBs harbor different pools of ILVs existing in equilibrium. One pool contributes to dynamics within MVBs and allows intraluminal proteins to return to the LM. The other, more inert, pool encompasses the bulk of secreted exosomes, which can transfer information to other cells, and the rest accounts for ILV cargo destined for lysosomal degradation. What distinguishes these different ILV pools is unclear at this point. Furthermore, we have observed a number of parallels between ILV retrofusion and viral entry, including the influence of pH and lipids, as well as the attenuating effects of the host antiviral protein IFITM3. Collectively, these findings support the notion that viruses employ the dynamic nature of MVBs to complete their infection cycle and expose a new twist on the emerging paradigm of pathogen-instigated subversion of host processes.

STAR Methods

Detailed methods are provided in the online version of this paper and include the following:

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A video abstract is available at https://doi.org/10.1016/j.cub.2021.06.022#mmc7.

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Author Contributions

J.N. and L.J. developed the concept; P.P. and J.N. designed the experiments; P.P. performed most of the experiments supported by constructs and cell lines generated by L.J.; H.J. and D.v.E. performed immunogold EM; L.M.V. and B.v.d.B. performed most of the experiments supported by constructs and cell lines generated by L.J.; H.J. and D.v.E. performed immunogold EM; L.M.V. and B.v.d.B. designed the macro for time-lapse analysis; J.N., P.P., and I.B. drafted the manuscript with input from all authors.

Declaration of Interests

The authors declare no competing interests.

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45. Calafat, J., Janssen, H., Stähle-Backdahl, M., Zuurbier, A.E., Knol, E.F., and Egesten, A. (1997). Human monocytes and neutrophils store transforming growth factor-alpha in a subpopulation of cytoplasmic granules. Blood 90, 1255–1266.
## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Mouse anti-CD63 NKI-C3 | NKI | N/A        |
| rabbit anti-GFP     | NKI    | N/A        |
| HRP-goat anti-Mouse IgG (H+L) | ThermoFisher Scientific | Cat# G21040; RRID:AB_2536527 |
| HRP-goat anti-rabbit IgG (H+L) | ThermoFisher Scientific | Cat# G21234; RRID:AB_2536530 |
| Mouse monoclonal Lamp2 antibody | SantaCruz | Cat# sc-18822; RRID:AB_626858 |
| Rabbit monoclonal calnexin antibody | Cell Signaling Technology | Cat#2679; RRID:AB_2228381 |
| Mouse monoclonal transferrin receptor antibody | ThermoFisher Scientific | Cat#13-6800; RRID:AB_2533029 |
| Mouse monoclonal HLA-DR antibody | NKI     | N/A        |
| Mouse monoclonal Tag101 antibody | ThermoFisher Scientific | MA1-23296; RRID:AB_561859 |
| **Chemicals, peptides, and recombinant proteins** | | |
| 1-Deoxymannojirimycin hydrochloride | Sigma-Aldrich | 73465-43-7 |
| Thioperoxidase maleate | Cayman Chemical | 30397-1-2 |
| U18666A | Cayman Chemical | 148440-81-7 |
| Bafilomycin A1 | Tebu-Bio | 88899-55-2 |
| Heterodimerizer | Takara | |
| SiR-Lysosome | Tebu-Bio | SC012 |
| **Experimental models: Cell lines** | | |
| Melanoma cell line MelJuso | Johnson et al. [38] | database of the DSMZ MEL-JUSO |
| **Oligonucleotides** | | |
| NLS mGFP fwd Nhe1 | This paper | N/A |
| AGGCACCACATGGTGAAACGACC AGCAAGACAAAGAAGACAGGAC AAGCAAGAGAAAGAGGATGTTG AGCAAGGGCAAGGAG AGCAAGGGCAAGGAG | This paper | N/A |
| GFP ENLYFQS rev bglII | This paper | N/A |
| CTACTCTGGAATACAGATTTTCCC CGCCCTCGTACAGCTGTCCTCAT | This paper | N/A |
| C-TEV 119-242 fwd Nhe1 | This paper | N/A |
| CCCAGCTAGCAAGAGCATGTCTAGCATGTTG | This paper | N/A |
| C-TEV 119-242 rev BamH1 | This paper | N/A |
| CCGAGGTCTCTAGTGGAGTACACCAATT | This paper | N/A |
| N-TEV 1-118 fwd SpeI | This paper | N/A |
| CCCAATAGTGGAGAAGCTTGTGGTAAAAGGG | This paper | N/A |
| N-TEV 1-118 rev BamH1 | This paper | N/A |
| CCGAGGTCTCTTGAGTGGAAGTGGTGGT | This paper | N/A |
| F2A Fwd 1 | This paper | N/A |
| GATTCACGCGGCAAGAGCCAATTCCAAC CCTCTTTTGGGCA | This paper | N/A |
| F2A Fwd 2 | This paper | N/A |
| TGACGCACTATCGGGATCAGGA GGGCCGATCAACACAGTAAACTTCGAC CTITTTGAAGGTCAGGGATTGTCAGTC GAACCCCTGGAACAG | This paper | N/A |
| F2A Rev 1 | This paper | N/A |
| GATCTCCTGAGGGTCTGACT CGACATCCCTCTGCTAGCTAAAGGTCTG AAGTTTGAATGTCG | This paper | N/A |
| F2A Rev 2 | This paper | N/A |
| TTTCACGGCGCTCCTGATCCCG ATGTATCTGCTGAGGCCAAGAGGTTTGAATTGCTTCCGGGCTTGG | This paper | N/A |
| IFITM1 fwd HinDIII | This paper | N/A |
| CCCAAGCTTCTAGTGGAGAAGGAG | This paper | N/A |

(Continued on next page)
**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Jacques Neefjes (j.j.c.neefjes@lumc.nl).

**Materials availability**
All unique/stable reagents generated in this study are available from the Lead Contact without restriction.

**Data and code availability**
- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Cell lines and culturing**
Cell Line Authentication was performed by Eurofins Genomics. MelJuso (human melanoma) cells were cultured in IMDM ( Gibco) supplemented with 7.5% fetal calf serum (FCS, Greiner) at 37°C. For exosome isolation, cells were cultured in IMDM ( Gibco) supplemented with 9% exosome-depleted fetal bovine serum (ThermoFisher Scientific). The cell-based system was constructed in the following way. First, NLS-GFP-TCS-CD63 was transfected into MelJuSo cells (Effectene, QIAGEN). Following selection on G418 for stable expression, cells were retrovirally transduced with the split snipper TEV viral supernatant (protocol from Retroviral systems; Nolan lab) and selected on Puromycin. After single-cell sorting of double positive cells (GFP/DsRed), cells were screened for activation of the TEV protease in response to dimerizer. Clones demonstrating cleavage of GFP upon dimerizer addition were selected for the study. Where appropriate, Turq-IFITM1-3 was introduced by transfection (Effectene, QIAGEN) and, following selection on Hygromycin, triple positive (Turq/GFP/DsRED) cells were sorted and expanded for further experiments.

**METHOD DETAILS**

**Antibodies and reagents**
Mouse anti-CD63 NKI-C341 and rabbit anti-GFP,42 followed respectively by HRP-goat anti-Mouse and HRP-goat anti-rabbit IgG (H+L) secondary antibodies (ThermoFisher Scientific) were used for detection of endogenous or overexpressed proteins by SDS-PAGE and western blot. Mouse monoclonal Lamp2 antibody was purchased from SantaCruz, rabbit monoclonal calnexin antibody was purchased from Cell Signaling Technology, mouse monoclonal transferrin receptor antibody and mouse monoclonal Tsg101

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**REAGENT or RESOURCE** | **SOURCE** | **IDENTIFIER**
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IFITM1 rev BamHI CCCAGGATCCCTAGTAAACCGGTT | This paper | N/A
IFITM2 fwd HinDIII CCCAAAGCTTTGATGAGACCATGGGGAAAC | This paper | N/A
IFITM2 rev BamHI CCCAGGATCCCTGCTGCTGGCCTGGAC | This paper | N/A
IFITM3_HindIII_fwd CCCAAAGCTTTGATGAACTACGTGCTGGAC | This paper | N/A
IFITM3-BamHI_rev CCCAGGATCCCTATCCATAGGCTGGGAGATC | This paper | N/A
Recombinant DNA
pCD63-EGFP-bos | Blott et al. | N/A
FKBP N-TEV and FRB C-TEV | Gray et al. | N/A
Software and algorithms
LAS X | Leica Microsystems | https://www.leica-microsystems.com/
Fiji 1.52p | National Institutes of Health | https://imagej.net/Fiji/Downloads
Microsoft Excel | Microsoft Inc. | N/A
Adobe Illustrator CC 2018 | Adobe Inc. | N/A
GraphPad Prism 8.4.2 | GraphPad | https://www.graphpad.com/
antibody were purchased from ThermoFisher Scientific, mouse monoclonal HLA-DR antibody (1B5) was obtained from the NKI, Amsterdam NL, SIR-Lysosome was purchased from Tebu-Bio (used at 50 μM for live cell imaging). Heterodimerizer (i.e., dimerizer) was purchased from Takara (0.35μM for live cell imaging, 0.5μM for exosome isolation). 1-Deoxymannojirimycin hydrochloride (DMM) and thymidine were purchased from Sigma-Aldrich. Thioperamide maleate and U18666A were purchased from Cayman Chemical (10 μM for live cell imaging, and 3 μg/mL respectively). Bafilomycin A1 was purchased from Tebu-Bio (100 nM for live cell imaging).

**Constructs**
The NLS-GFP-TCS-CD63 construct was generated by PCR adding a NLS (MVKRPAATKKAGQAKKKK) at the 5’ end of GFP and a TEV cleavage (ENLYFQS) moiety at 3’ end and this replaced the GFP part in the original GFP-CD63 construct. The split sniper TEV was build up by dividing the FKBP N-TEV and FRB C-TEV by a F2A motif allowing both parts to be expressed separately and at relative equal amounts under the same promoter. This box was cloned into a pMX Puro IRES2 NLS DsRed2 plasmid. The templates for IFITM1-3 were isolated from the Gateway pDONR223 (Entry ORF Library) library. cDNA’s were first cloned into the mTurq2-C1 vector before being subcloned into pcDNA3.1 Hygro (Invitrogen). All constructs were sequence verified.

**Construction of the system**
The cell-based system was constructed in the following way. First, NLS-GFP-TCS-CD63 was transfected into MelJuSo cells (Efectene, Qiagen). Following selection on G418 for stable expression, cells were retrovirally transduced with the split sniper TEV viral supernatant (protocol from Retroviral systems Nolan lab) and selected on Puromycin. After single-cell sorting of double positive cells (GFP/DsRed), cells were screened on activation of the TEV protease in response to dimerizer. Clones demonstrating cleavage of GFP upon dimerizer addition were used in the study. Where appropriate, Turq-IFITM1-3 was introduced by transfection (Efectene, Qiagen) and, following selection on Hygromycin, triple positive (Turq/GFP/DsRED) cells were sorted and expanded for further experiments.

**Confocal microscopy**
Live cells were incubated with SIR-Lysosome (50 μM) for at least 30 min. The medium was removed and medium at a pH of 6.3 (adjusted with acetic acid 10mM in water) was added to the cells. Samples were imaged using a Leica SP8 WLL confocal microscope, HC PL APO 63x/1.40 oil immersion objective and HyD detectors. For Z stack imaging, a Leica SP8 with Andor Dragonfly spinning disc module was used, and 0.2 μM Z stacks were acquired with a 63x oil immersion objective. The microscope was equipped with a humidified climate control system at 37˚C supplemented with 5% CO2. Images were collected using a digital zoom of 1.0 in 512 by 512 scanning format with line averaging [4x], at a rate of 180 s per frame for a period of at least 6 h. In the case of thioperamide or U18666A treatment, the cells were incubated with the compounds for 19h and 24h respectively before imaging. Cells were treated with Bafilomycin A1 for 6h before imaging.

**Quantification of ILV retrofusison**
Post-collection image processing and analysis were performed using Fiji. A macro was programmed to analyze fluorescence time-lapses (the different steps are shown in Video S2). Briefly, the macro was designed such that the user first sets an intensity threshold to define the area of endolysosomes (EL) based on SIR-lysosome fluorescence in the first and last frame, and the threshold for the other frames is automatically interpolated to allow for a possible slight fluctuation in fluorescence intensity. This allows quantification of the GFP signal in EL. The segmentation of nuclei (N) is achieved by thresholding the DSRED fluorescence, this allows quantification of the GFP signal in nuclei. Next, to define the area of the plasma membrane, a mask is created that segments the area outside of the nuclei and EL. Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL). Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL). Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL). Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL). Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL). Within this mask, the user can set a threshold on the GFP signal in the first and last frame to define the area of endolysosomes (EL).

**DMM pulse-chase**
Cells were synchronized by a double thymidine block and incubated with 1mM 1-deoxymannojirimycin (DMM) for the times indicated and then lysed in lysis buffer containing 0.5% NP-40, 150 mM NaCl, 50 mM Tris-HCl pH 7.6 and 5 mM MgCl2 for 30 min before addition of Laemmli Sample Buffer (containing 80 mM DTT) followed by 10 min incubation at 95˚C.

**Electron microscopy (EM)**
MelJuSo cells were fixed for 2 hours in 2% paraformaldehyde + 0.2% glutaraldehyde in 0.1 M PHEM buffer (60 mM PIPES, 25 mM HEPES, 2 mM MgCl2, 10 mM EGTA, pH 6.9) and then processed for ultrathin cryosectioning, as previously described. Briefly, 50 nm cryosections were cut at – 120˚C using diamond knives in a cryoultramicrotome (Leica Aktiengesellschaft) and transferred with a mixture of sucrose and methylcellulose onto formvar-coated copper grids. The grids were placed on 35-mm Petri dishes containing 2% gelatine. Ultrathin frozen sections were incubated at room temperature with primary antibody and then incubated with 10-nm...
protein A-conjugated colloidal gold (Klumperman Lab, Utrecht University), as described. After washing, the sections were fixed for 10 minutes in 1% glutaraldehyde, blocked and incubated with the second primary antibody and the second label protein A/15 nm gold. The sections were embedded in a mixture of methylcellulose and uranyl acetate and examined with a Philips CM10 electron microscope (FEI).

In case of exosome imaging, exosome-containing pellets were adsorbed onto formvar and carbon-coated copper grids after which they were negatively stained and embedded in a mixture of methylcellulose and uranyl acetate. Exosomes were imaged with a Tecnai 12 transmission electron microscope (FEI) at 120 kV acceleration voltage.

**Exosome isolation**
Cells were cultured in presence or absence of dimerizer for 2h, then incubated in exosome-depleted medium in the presence (or absence) of dimerizer for 5h. The supernatant (enriched in exosomes) was retrieved and subjected to differential centrifugation at 4°C: 1000 g for 10 min, then 2000 g for 20 min, 10000 g for 30 min and 100300 g for 2h. The exosome pellets were lysed in lysis buffer containing 0.5% NP-40, 150 mM NaCl, 50 mM Tris-HCl pH 7.6 and 5 mM MgCl2 for 30 min before addition of Laemmli Sample Buffer (containing 80 mM DTT) followed by 10 min incubation at 95°C. After collection of the medium for exosome isolation, the cells were lysed as described above.

**Western blotting**
Samples were separated using 8% acrylamide gels and transferred to a PVDF membrane (Immobilon-P, 0.45 μm, Millipore) at 100V for 3h. The membranes were blocked in PBS/5% Skim Milk (Oxoid) and incubated with a primary antibody for 1h diluted in PBS/0.1% Tween20 (Sigma-Aldrich)/5% Milk, washed three times for 10min in PBS/0.1% Tween and incubated with the secondary antibody for 1h diluted in PBS/0.1% Tween/5% Milk and washed three times again in PBS/0.1% Tween. Signals were detected on the Chemidoc XRS+ imager (Bio-Rad) using ECL (SuperSignal West Dura Extended Duration Substrate, Thermo Scientific). Intensity of bands was quantified using ImageJ.

**QUANTIFICATION AND STATISTICAL ANALYSIS**
Statistical analyses were performed with GraphPad Prism 7. All tests are mentioned in the corresponding figure. All experiments were performed independently at least 3 times, as indicated. All graphical data plots were produced using GraphPad Prism 7 and fonts were adjusted with Adobe Illustrator CC.
Supplemental Information

Retrofusion of intralumenal MVB membranes parallels viral infection and coexists with exosome release

Priscillia Perrin, Lennert Janssen, Hans Janssen, Bram van den Broek, Lennard M. Voortman, Daphne van Elsland, Ilana Berlin, and Jacques Neefjes
Addition of dimerizer

**A**

**B**

| % GFP-CD63 or total CD63 on ILVs (per MVB) | Dimerizer | -   | +   |
|--------------------------------------------|-----------|-----|-----|
| GFP-CD63                                   | ***       | ns  |     |
| CD63                                       |           |     |     |

**C**

Addition of dimerizer

| t = 0h | DsRed | GFP | TQ |
|--------|-------|-----|----|
| SIR-lysosome | Overlay GFP-SIR-lysosome | Overlay TQ-SIR-lysosome |

| t = 6h | GFP | TQ |
|--------|-----|----|
| SIR-lysosome | Overlay GFP-SIR-lysosome | Overlay TQ-SIR-lysosome |
Figure S1. Analysis of retrofusion measurement and TEV protease efficiency. Related to Figures 1 and 2.

(A) Representative plot of intracellular median GFP fluorescence intensity from GFP-CD63 cells either (+) or not (-) cultured with dimerizer for the time indicated (min) N: nucleus; EL: late endosomes as labelled by SiR-lysosome; Cyto: cytoplasm. EL: endolysosome; N: nucleus; PM: plasma membrane; Cyto: cytoplasm.

(B) Quantification of GFP-CD63 as detected by immunogold labeling on ILVs relative to LM (expressed as ratio per MVB) following incubation with dimerizer (6h). Shown is median +/- IQR from over 98 MVBs from 2 independent experiments.

(C) Analysis of Lamp1-TEV-TQ expression in GFP-CD63 retrofusion-monitoring cells stained with SiR-lysosome. Confocal fluorescence images of GFP-CD63 and Lamp1-TEV-TQ (white) distribution before (t=0h) and after (t=5h) treatment with dimerizer are shown, along with color overlays of GFP (green) or TQ (blue) with SiR-Lysosome (red). NLS-DsRED (magenta) in the nucleus indicate expression of the split TEV protease at t=0. Bar: 10 µm. TQ : mTurquoise2.
**A**

Addition of dimerizer

|       | DSRED          | GFP            |
|-------|----------------|----------------|
| Ctrl  |                |                |
| t = 0h|                |                |
| t = 6h|                |                |

SiR-lysosome Overlay GFP-SiR-lysosome

**B**

Normalized median fluorescence intensity

- Control
- Bafilomycin A
- Thioperamide
- U18666A

**C**

Control

Bafilomycin A

U18666A

Thioperamide
Figure S2. Effects of lipid and pH manipulation on the rate of ILV retrofusion. Related to Figure 3.

(A) Confocal images of control (left panel; Ctrl) and U18666A treated (right panel) GFP-CD63 cells as used for monitoring retrofusion. Confocal fluorescence images of GFP-CD63 \textit{(white)} distribution before (t=0h) and after (t=6h) treatment with dimerizer are shown, along with color overlays of GFP \textit{(green)} with SiR-Lysosome \textit{(red)}. NLS-DsRED \textit{(magenta)} in the nucleus indicate expression of the split TEV protease at t=0. All scale bars, 10 µm.

(B) Representative plots of normalized median GFP fluorescence intensity in endolysosomes \textit{(as stained with SiR-lysosome)} over time (min) following dimerizer addition in control cells \textit{(gray)} versus those treated with Bafilomycin A, thioperamide or U18666A \textit{(red)}.

(C) Electron micrograph featuring immunogold labeling with GFP antibody of control cells following incubation with U18666A, thioperamide or Bafilomycin A. Scale bar, 200 nm.
A

Addition of dimerizer

| Time (h) | DSRED | GFP |
|----------|-------|-----|
| t = 0h   | ![Image](image1.png) | ![Image](image2.png) |
| t = 6h   | ![Image](image3.png) | ![Image](image4.png) |

Ctrl

SIR-lysosome Overlay GFP-SIR-lysosome

Addition of dimerizer

| Time (h) | DSRED | GFP |
|----------|-------|-----|
| t = 0h   | ![Image](image5.png) | ![Image](image6.png) |
| t = 6h   | ![Image](image7.png) | ![Image](image8.png) |

+ TQ-IFITM1

TQ Overlay GFP-SIR-lysosome

B

Addition of dimerizer

| Time (h) | DSRED | GFP |
|----------|-------|-----|
| t = 0h   | ![Image](image9.png) | ![Image](image10.png) |
| t = 6h   | ![Image](image11.png) | ![Image](image12.png) |

TQ-IFITM1 TQ-IFITM2 TQ-IFITM3

Lysotracker Overlay TQ-lysotracker

C

Normalized median fluorescence intensity

| Time (h) | Control | TQ-IFITM2 |
|----------|---------|-----------|
| 0        | 1.0     | 1.0       |
| 1        | 0.8     | 0.8       |
| 2        | 0.6     | 0.6       |
| 3        | 0.4     | 0.4       |
| 4        | 0.2     | 0.2       |
| 5        | 0.0     | 0.0       |

| Time (h) | Control | TQ-IFITM1 |
|----------|---------|-----------|
| 0        | 1.0     | 1.0       |
| 1        | 0.8     | 0.8       |
| 2        | 0.6     | 0.6       |
| 3        | 0.4     | 0.4       |
| 4        | 0.2     | 0.2       |
| 5        | 0.0     | 0.0       |
Figure S3. Effects of overexpression of the IFITM family members on the rate of ILV retrofusion. Related to Figure 4.

(A) Confocal images showing overexpression of IFITM1 or 2 in GFP-CD63 retrofusion-monitoring cells, as used for the retrofusion experiments. Confocal fluorescence images of GFP-CD63 (white) distribution before (t=0h) and after (t=6h) treatment with dimerizer are shown, along with color overlays of GFP (green) with SiR-Lysosome (red). NLS-DsRED in the nucleus (magenta) and TQ-IFITM1 and 2 (blue) respectively indicate expression of the split TEV protease and the IFITM proteins at t=0. Scale bars, 10 µm

(B) Localization of TQ-IFITM1, 2 and 3 proteins to late endocytic compartments as marked by SiR-Lysosome. Confocal fluorescence overlays of TQ-IFITM1-3 (green) with SiR-Lysosome (red) are shown in the bottom panel.

(C) Representative plots of normalized median GFP fluorescence intensity in endolysosomes (as marked by SiR-Lysosomes) over time (min) following dimerizer addition in control cells (gray) versus those overexpressing TQ-IFITM1 or 2 (red).
Figure S4. Retrofusion and exosomes. Related to Figure 4.

(A) Electron micrographs of exosome isolates prepared as described in Figure 4F. Scale bar, 200 nm.

(B) Biochemical profile of exosome isolates (secreted in absence of dimerizer) was assessed by immunoblot against the indicated markers. TfR: transferrin receptor, recycling endosomes; HLA-DR: MHC class II, LM and ILVs of late endosomes/lysosomes; Lamp2: LM of late endosomes/lysosomes; Golgin97: trans-Golgi network; Calnexin: endoplasmic reticulum; Tsg101: exosome marker. WCL: diluted whole-cell lysate (to prevent overloading control signal). The position of marker proteins is indicated.

(C) Immunoblot analysis of GFP-CD63 (detected with anti-GFP antibodies) versus untagged CD63 in exosome isolates and a fraction of whole cell lysate (WCL) from control cells or cells overexpressing TQ-IFITM3 treated in the presence (+) or absence (-) of dimerizer. The different incubations as well as the position of the marker proteins are indicated.

(D) Immunoblot analysis of CD63 in exosome isolates and WCL from control cells. Exosomes were isolated from 10x10^6 cells. For the WCL, 1.3x10^6 cells were loaded on the same gel. The results from a duplicate exosome isolation were loaded on two lanes as is the corresponding WCL. Position of marker proteins is indicated.

(E) Quantification of GFP-CD63 signal in exosome isolates versus the signal in a fraction of WCL from control cells. Exosomes were isolated from 10x10^6 cells and 1.3x10^6 cells were loaded on the same gel. Shown is mean +/- SD from three independent experiments.

(F) Quantification of the percentage of GFP-CD63 secreted in exosomes relative to the fraction cell lysate (1.3x10^6 cells) including MVBs. This is not corrected for the number of cells providing exosomes. The corrected values are shown in Figure 4H.