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A PI3K-WIPI2 positive feedback loop allosterically activates LC3 lipidation in autophagy

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Autophagy degrades cytoplasmic cargo by its delivery to lysosomes within double membrane autophagosomes. Synthesis of the phosphoinositide PI(3)P by the autophagic class III phosphatidylinositol-3 kinase complex I (PI3KC3-C1) and conjugation of ATG8/LC3 proteins to phagophore membranes by the ATG12–ATG5–ATG16L1 (E3) complex are two critical steps in autophagosome biogenesis, connected by WIPI2. Here, we present a complete reconstitution of these events. On giant unilamellar vesicles (GUVs), LC3 lipidation is strictly dependent on the recruitment of WIPI2 that in turn depends on PI(3)P. Ectopically targeting E3 to membranes in the absence of WIPI2 is insufficient to support LC3 lipidation, demonstrating that WIPI2 allosterically activates the E3 complex. PI3KC3-C1 and WIPI2 mutually promote the recruitment of each other in a positive feedback loop. When both PI 3-kinase and LC3 lipidation reactions were performed simultaneously, positive feedback between PI3KC3-C1 and WIPI2 led to rapid LC3 lipidation with kinetics similar to that seen in cellular autophagosome formation.

Introduction

Macroautophagy (hereafter autophagy) is a conserved intracellular degradation process that ensures cellular homeostasis by the degradation of harmful material such as damaged organelles, misfolded proteins, and bacterial pathogens. It also ensures survival of cells during starvation (Anding and Baehrecke, 2017; Gomes and Dikic, 2014; Kirkin and Rogov, 2019; Wen and Klionsky, 2016; Zaffagnini and Martens, 2016). These functions are achieved by the sequestration of the cargo material within double membrane vesicles, the autophagosomes, which form de novo around the cargo. The precursors of autophagosomes, called phagophores (or isolation membranes), emerge in the cytoplasm as small membrane structures, which capture the cargo as they grow. Upon closure of the phagophores, the structures become autophagosomes, which deliver their cargo for degradation upon fusion with lysosomes (Lamb et al., 2013; Mercer et al., 2018).

Approximately 40 autophagy (ATG) proteins are required for most forms of autophagy. In particular, these are the ULK1/Atg1 kinase complex, the ATG9 vesicles, the ATG14-containing class III phosphatidylinositol-3 kinase complex I (PI3KC3-C1), the β-propellers that bind polyphosphoinositides (PROPPINs) and ATG2, as well as the ATG12 and LC3 conjugation machineries (Bento et al., 2016; Hurley and Young, 2017; Mizushima, 2009; Matsunaga et al., 2009; Obara et al., 2006). PI(3)P-enriched membranes recruit downstream sensor proteins called PROPPINs or WD-repeat protein interacting with phosphoinositides (WIPIs; Proikas-Cezanne et al., 2015). WIPIs have a seven-bladed β-propeller fold and are a platform for the binding of downstream effector proteins. WIPIs bind PI(3)P and phosphatidylinositol 3,5-bisphosphate at two sites through their Phe-Arg-Arg-Gly (FRRG) motifs and increase membrane affinity further using a loop that inserts into the target membrane (Baskaran et al., 2012; Krick et al., 2012; Watanabe et al., 2012). While Saccaromyces cerevisiae has three members of this family, Atg18, Atg21, and Hsv2, humans have four orthologues, namely, WIPI1-4. Additionally, several isoforms derived from alternative splicing exist for the WIPIs. For example, there are six isoforms for WIPI2, including WIPI2b and d.

WIPIs promote phagophore expansion by the recruitment of the LC3 conjugation machinery (Dooley et al., 2014). In yeast, Atg21 interacts with the coiled-coil domain of Atg16 to recruit the Atg12–Atg5–Atg16 complex to the site of autophagosome formation (Juris et al., 2015). In humans, WIPI2b recruits ATG16L1 via a direct interaction with a motif downstream of the coiled-coil domain of ATG16L1 (Dooley et al., 2014). Because the ATG12–ATG5–ATG16L1 complex behaves much like a...
ubiquitin E3 ligase, we refer to it hereafter as the E3 complex. The E3 complex subsequently acts to promote the attachment of LC3 proteins to the membrane lipid phosphatidylethanolamine (PE) in a manner analogous to the action of E3 ligases in ubiquitin conjugation reactions (Hanada et al., 2007). The attachment of LC3 to PE, referred to as lipidation, requires the E1-like ATG7 as well as the E2-like ATG3 enzymes and mediates efficient phosphophore elongation and serves to recruit cargo material into autophagosomes (Ichimura et al., 2000; Zaffagnini and Martens, 2016). The E3 complex itself is the product of a ubiquitin-like conjugation machinery wherein the ATG7 and ATG10 proteins conjugate the ubiquitin-like ATG12 to a lysine residue in ATG5 (Mizushima et al., 1998). The ATG12–ATG5 conjugate subsequently noncovalently binds the ATG16L1 protein (Kuma et al., 2002; Mizushima et al., 1999, 2003).

A number of the individual steps in autophagy have been studied by in vitro reconstitution (Brier et al., 2016), which permits a level of experimental control not possible in cell culture. This minimalist approach can reveal the inherent activities of individual components stripped of the complexity of the cellular context. More sophisticated reconstitutions including multiple purified components and reactions can restore some of the physiological context and complexity while maintaining maximal experimental control. As part of a larger effort to reconstitute mammalian autophagosome biogenesis from start to finish using giant unilamellar vesicles (GUVs) as a model, we studied by in vitro reconstitution (Brier et al., 2016), which permits a level of experimental control not possible in cell culture. This minimalist approach can reveal the inherent activities of individual components stripped of the complexity of the cellular context. More sophisticated reconstitutions including multiple purified components and reactions can restore some of the physiological context and complexity while maintaining maximal experimental control. As part of a larger effort to reconstitute mammalian autophagosome biogenesis from start to finish using giant unilamellar vesicles (GUVs) as a model, we

Results

Reconstitution of PI(3)P- and WIPI-dependent LC3 lipidation

We expressed and purified the recombinant human E3 complex by coexpressing its three subunits ATG12, ATG5, and ATG16L1 (isoform β) together with ATG7 and ATG10 in insect cells (Fig. S1 A). We assessed the oligomeric state of the purified complex in solution by static light scattering (SLS) and detected a monodisperse population of dimeric complexes, where the experimentally determined molecular weight corresponds to 2.90 ± 105 Da ± 0.8%, essentially equal to the computed mass of two 1.48 × 105 Da monomers (Fig. S1 B). Thus, the purified E3 complex exists as a dimer composed of two copies of each subunit ATG5, ATG12, and ATG16L1 (Fig. S1 C). We assessed the ability of the recombinant E3 complex to catalyze LC3 lipidation. We co-incubated PE-containing small unilamellar vesicles (SUVs) with ATG7, ATG3, and LC3BΔ5C, with the latter lacking the five C-terminal residues in order to expose the Gly required for lipidation (LC3B-I; see all purified components of the machinery in Fig. S1 D). Consistent with the observation by Lystad et al. (2019), we found that the presence of the E3 complex promoted LC3B lipidation (LC3B-II; Fig. 1 A). We also found that the presence of ATG16L1 was required for the promotion of LC3B-II formation (Fig. 1 A). Lipidation was far more efficient with di-oleoyl (DO) lipids compared with palmitoyl-oleoyl (PO) lipids (Fig. S1, E and F). LC3-II conversion was >50% complete with DO lipids after 30 min, whereas a comparable degree of conversion required overnight (o.n.) incubation with PO lipids. This effect cannot be explained by preferential membrane binding by the E3 complex (Dudley et al., 2019; Lystad et al., 2019), as PO and DO lipids are bound equally (Fig. S1, G and H). These data, which are consistent with a high proportion of unsaturated lipids in autophagosomal membranes (Schrütter et al., 2020), suggest that the presence of two rather than one unsaturated tail in the lipids of the membrane substrates increases the flexibility of the membrane, facilitating structural rearrangements necessary for LC3 lipidation.

In cells, efficient membrane recruitment of the E3 complex during autophagosome formation depends on the presence of WIPI2 (Dooley et al., 2014). To recapitulate this step, we used GUVs instead of SUVs because they are less curved, rendering membrane binding more stringent. In the absence of WIPI2d, we observed no E3 recruitment to GUVs bearing the same lipid composition as the DO-SUVs used in the previous assays (compare Fig. S1 H and Fig. 1 B). The addition of ATG7, ATG3, mcherry-LC3BΔ5C, and ATP did not result in any detectable mcherry-LC3B lipidation on GUVs (Fig. 1 C).

To test the ability of WIPIs to activate LC3 lipidation, we expressed and purified the three human WIPI proteins: WIPI2d, WIPI3, and WIPI4 isoform 1. We assessed their ability to interact with the E3 complex in a microscopy-based bead interaction assay (Fig. 2 A and Fig. S2 A). We assessed their ability to interact with the E3 complex in a microscopy-based bead interaction assay (Fig. 2 A and Fig. S2 A). mcherry-WIPI2d was specifically recruited to beads coated with E3-GFP (Fig. 2 A). We also observed that the E3-GFP was robustly recruited to beads coated with GST-mcherry-WIPI3, but not GST-mcherry-WIPI4 (Fig. S2 A). Thus, the E3 complex directly binds to WIPI2d and WIPI3. We then tested the ability of WIPI2d to recruit the E3 to GUVs. We added the E3-GFP and mcherry-WIPI2d to PI(3)P-containing GUVs and compared the GFP signal intensity to conditions lacking mcherry-WIPI2d (Fig. 2 B) or PI(3)P (Fig. S2 B). The E3 was robustly and specifically recruited to PI(3)P-containing GUVs in a WIPI2d-dependent manner (Fig. 2 B). When 15% PI was used instead of 5% PI(3)P, no WIPI2d and E3 recruitment was observed, despite the equivalent negative charge of the membrane (Fig. S2 B).

We assessed whether the recruitment of the E3 to PI(3)P-containing GUVs by WIPI2d elicited LC3B lipidation. mcherry-LC3B was efficiently lipidated to these GUVs in an ATP-, WIPI2d-, and PI(3)P-dependent manner (Fig. 2 C and Fig. S2 C). To further confirm that the mcherry signal observed on these GUVs was due to covalent attachment of LC3B to membranes, we performed fluorescence recovery after photo-bleaching (FRAP) experiments (Fig. 2 D; Fracchiolla et al., 2016). The mcherry-LC3B signal did not recover over the time period imaged, suggesting that mcherry-LC3B is covalently linked to the membrane. Moreover, the addition of the ATG4B protease,
which removed LC3B from the membrane, resulted in loss of membrane-associated fluorescence, further corroborating that LC3B is lipidated on the GUV membrane (Fig. 2E). Thus, the GUV system recapitulates the strict dependence of E3 recruitment and LC3B lipidation on PI(3)P and WIPI2 observed in cells (Dooley et al., 2014).

WIPI2d both recruits and allosterically activates E3 for LC3 lipidation  

Given the dramatic stimulatory effect of WIPI2d on LC3B lipidation in the GUV system (Fig. 2C), we asked whether WIPI2d could activate the E3 complex beyond merely recruiting it to the membrane. We observed that the lipidation occurred more efficiently in the presence of WIPI2d (Fig. 2, F and G). Importantly, in the SUV system, WIPI2d did not significantly increase membrane binding by the E3 as assessed by sedimentation assays (Fig. S3 A) and bead-based recruitment assays (Fig. S3, B and C). We therefore conclude that WIPI2d must have an effect on the LC3B lipidation machinery that goes beyond simply recruiting E3 to the membrane.

In yeast, the WIPI orthologue Atg21 recruits Atg16 to the phagophore assembly site (Juris et al., 2015). Atg21 binds to a DE motif (D101, E102) within the coiled coil of Atg16, while WIPI2b binds to an EE-containing motif C terminal to the extended coiled-coil domain (E226, E230; Fig. 3, A and B; Dooley et al., 2014). Fortuitously, the intra-coiled-coil motif of yeast Atg16 is also present in human ATG16L1 and corresponds to residues D164 and E165. We found that yeast Atg21 can directly bind the human E3 complex (Fig. 3 D). This led us to ask whether E3 activation was unique to human WIPI2 or common to any ATG16- and PI(3)P-binding WIPI orthologue. Atg21 was added to LC3 lipidation reactions using SUVs. Atg21 did not significantly enhance LC3 lipidation (Fig. 3, E and F). Thus, the stimulatory effect of WIPI2d as opposed to Atg21 cannot be attributed to enhanced membrane binding of the E3 in the presence of WIPI2d because the membrane binding by

Figure 1. LC3 lipidation machinery is active on SUVs, but not on GUVs. (A) In vitro LC3B lipidation assay on DO-SUVs. ATG7, ATG3, and LC3B were mixed with SUVs (65% PC:15% liver PI:20% PE), in the absence (left) or presence of the E3 complex (1 µM; middle) or the ATG12–ATG5 conjugate (1 µM; right) and incubated at 37°C in the presence of MgCl2/ATP. Samples taken at the indicated time points (minutes to hours) were loaded on a 4–15% SDS-polyacrylamide gel. MW, molecular weight. (B) The E3-GFP complex (0.5 µM) was added to DO-GUVs (65% PC:15% liver PI:20% PE). (C) The E3-GFP (0.5 µM), ATG7, ATG3, and mCherry-LC3B were added to GUVs (65% PC:15% liver PI:20% PE), in the presence of MgCl2/ATP. Representative confocal micrographs are shown.

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Figure 2. WIPI2d both recruits and allosterically activates E3 for LC3 lipidation. (A) Quantification of the mCherry-WIPI2d signal intensity (red bars) measured on GFP-Trap beads coated with either GFP or GFP-tagged E3 (means ± SD; n = 94 [E3-GFP] or 80 [GFP]). P values were calculated using Student’s t test: not significant (ns), P > 0.05; *, 0.01 < P < 0.05; **, 0.001 < P < 0.01; AU, arbitrary units. (B) E3-GFP (0.5 µM) was added to DO-GUVs containing 75% PC:5% PI(3)P:20% PE in the absence or presence of mCherry-WIPI2d (0.5 µM). (C) E3-GFP (0.1 µM) was coincubated with WIPI2d (0.5 µM), mCherry-LC3B, and the lipidation machinery on DO-GUVs [75% PC:5% PI(3)P:20% PE] in the presence of MgCl2/ATP. (D) FRAP experiment on GUVs after lipidation in the presence of ATP as conducted in (C). A quantification is shown (means ± SD; n FRAP = 3, n no FRAP = 2), together with representative images of the two conditions, at times 0 and 25 min after the photobleaching. (E) De-lipidation reactions on GUVs treated as in (C), in the presence of ATP. CIP/ATG4B (left) or buffer (right) was
the E3 was not significantly different under the conditions tested (Fig. S3, A–C).

We took advantage of the previously characterized ATG16L1 binding site in WIPI2b identified in Dooley et al. (2014). Sequence alignment of the WIPI2b and WIPI2d isoforms showed that the positively charged residues R108 and R125 comprising the binding site are present in both isoforms (Fig. 3 C). The R108,125E mutant was defective in binding to the E3 complex on GUVs (Fig. S3 D) and lost the promoting effect on LC3b lipidation (Fig. 3 G and H; and Fig. S3 E), although still retaining its ability to bind PI(3)P membranes (Fig. S3 F). We conclude that WIPI2d has a potent activating effect on LC3b lipidation, depending on its ability to recruit the E3 to the membrane via the ATG16L1 subunit. Beyond this, it further activates LC3b lipidation, likely due to an allosteric effect on the E3.

**PI3KC3-C1 supports LC3b lipidation on unsaturated flat membranes**

During autophagosome nucleation, the PI3KC3-C1 complex translocates to the ER to generate PI(3)P to recruit downstream factors (Axe et al., 2008; Matsunaga et al., 2010). It was previously reported that PI3KC3-C1 is active only on high curvature membranes and has no measurable activity on GUV membranes (Rostislavleva et al., 2015). The ER membrane contains both flat and highly curved membranes, and much of the omegasome domain of the ER involved in autophagy is flat on a molecular scale. The membrane of the ER is less densely packed than post-Golgi compartments (Bigay and Antonny, 2012; Vanni et al., 2014), and we reasoned that highly unsaturated lipid mixtures, also found in autophagosomal membranes (Schütter et al., 2020), might therefore support PI3KC3-C1 activity on the flat membrane of GUVs.

We tested membrane binding and the enzymatic activity of PI3KC3-C1 on GUVs composed of lipids with a headgroup composition resembling the ER (65% phosphatidylcholine [PC]:20% PE:5% phosphatidylserine [PS]:10% PI) but with different hydrophobic tails. PI3KC3-C1 failed to bind to GUV membranes made with the brain lipid extract previously used (Fig. S4, A and B; Rostislavleva et al., 2015). Using a mCherry-Fab1, YOTB, Vac1, and EEA1 (FYVE) domain as a probe for PI(3)P, minimal activity of PI3KC3-C1 was observed on these GUVs (Fig. S4, A and C). By contrast, PI3KC3-C1 strongly bound to and robustly produced PI(3)P on GUVs composed of lipids with two unsaturated tails (DO), but not of only one unsaturated (PO) tail (Fig. 4, A and B; and Fig. S4). Little FYVE domain recruitment was seen on GUVs composed of DO lipids in the absence of PI3KC3-C1 (Fig. S4, A and C). These data show that PI3KC3-C1 is active on flat membranes with unsaturated lipids, suggesting that ER-like membranes with loose lipid packing are good substrates for PI3KC3-C1 even when they are flat.

We tested whether PI3KC3-C1 could support LC3b lipidation on GUV membranes with DO lipids via the WIPI2d–E3 axis. Upon addition of WIPI2d, the E3, and the LC3 conjugation machinery (ATG7, ATG3, mCherry-LC3bΔ5C), PI3KC3-C1 robustly triggered membrane recruitment of the E3-GFP complex and subsequent mCherry-LC3b lipidation (Fig. 4, C and D). Consistent with expectation, the E3 was not detectably recruited to the GUV membrane in the absence of WIPI2d (Fig. 4, C and D). Mutation of the conserved PI(3)P binding FRRG motif significantly reduced E3 membrane recruitment and LC3 lipidation (Fig. 4, C and D). The effect of WIPI2d on the PI3KC3-C1–induced E3 membrane association and LC3b lipidation was dose dependent (Fig. 4 E). We analyzed the effect of WIPI3 on LC3b lipidation in the in vitro system and found that WIPI3 also mediated LC3b lipidation on GUV membranes (Fig. S5), suggesting a potential role of WIPI3 in autophagosome formation in cells. In combination, the data obtained using the reconstituted system showed that PI3KC3-C1 stimulates LC3 lipidation on GUV membranes in a WIPI-dependent manner.

**Positive feedback between PI3KC3-C1 and WIPI2d promotes LC3b lipidation**

Because PI3KC3-C1 strongly promotes WIPI2–dependent LC3b lipidation (Fig. 4, C and D), we asked whether it was solely attributable to its PI 3-kinase enzyme activity or whether the PI3KC3-C1 also had additional roles. To elucidate the role of PI3KC3-C1 more precisely, we assayed how much PI(3)P was produced by PI3KC3-C1 on GUVs by comparing FYVE domain recruitment to GUVs with 10% (mol fraction) PI(3)P in the absence of CI or GUVs with 10% PI in the presence of CI, respectively. We found that the PI3KC3-C1 produced ~2% PI(3)P in 30 min (Fig. 5, A and B). We then measured WIPI2d recruitment to these two GUVs and surprisingly found that WIPI2d was recruited much faster than the FYVE domain to the 10% PI GUVs in the presence of PI3KC3-C1, despite showing lower affinity than the FYVE domain to 10% PI(3)P GUVs (Fig. 5, A and B). The WIPI2d FRRG mutant was not detectably recruited to the either of these two GUVs (Fig. 5, A and B).

To determine whether the enzymatic activity of the PI3KC3-C1 was required for the enhanced recruitment of WIPI2d compared with the FYVE domain, we generated GUVs containing 2% PI(3)P, but no PI. We first analyzed the binding of PI3KC3-C1 itself to these GUV membranes, and only trace binding of PI3KC3-C1 to membrane was observed (Fig. 5, C and D). However, in the presence of WIPI2d, but not the WIPI2d FRRG mutant or FYVE domain, an increased amount of PI3KC3-C1 was recruited to the membrane (Fig. 5, C and D). We observed that more WIPI2d was recruited to these GUVs in the presence of PI3KC3-C1 (Fig. 5, C and D), which suggests that PI3KC3-C1 and

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Figure 3. **WIPI2d specifically promotes LC3 lipidation.** (A) Scheme showing the domain organization of *S. cerevisiae* Atg16 and *H. sapiens* ATG16L1 (isoform β) and their interactors. (B) Alignment of *S. cerevisiae* Atg16 and *H. sapiens* ATG16L1 (isoform β) protein sequences spanning a region around the D101, E102 of Atg16 (Atg21 binding site) and residues D164, E165 in ATG16L1. (C) Alignment of the *H. sapiens* WIPI2 isoforms b and d spanning the region surrounding the ATG16L1 binding site including R108 and R125. (D) Microscopy-based bead protein interaction assay with RFP-Trap beads coated with mCherry-WIPI2d, mCherry-Atg21, or mCherry as baits and incubated with 5 µM E3-GFP as prey. Representative confocal micrographs are shown. (E and F) DO-SUVs [75% PC:5% PI(3)P:20% PE] were incubated with ATG7, ATG3, E3-GFP (0.5 µM), and LC3B in the presence of no PROPPIN, WIPI2d or Atg21 (both 2.5 µM). Samples taken at
PI2d cooperatively bind to membranes. We further observed that GFP-tagged PI3KC3-C1 was specifically recruited to beads coated with WIP12d, but not the FYVE domain (Fig. 5, E and F). These data imply that the two might form a stable physical complex on membranes, which requires PI(3)P to anchor WIPI2. Both PI3KC3-C1 and WIP12d can additionally insert parts of themselves into the membrane (Baskaran et al., 2012; Fan et al., 2011; Rostislavleva et al., 2015; Chang et al., 2019); thus, these two factors could modulate local membrane properties to enhance binding.

The inherent membrane-binding activity of the human E3 (Dudley et al., 2019; Lystad et al., 2019) is insufficient to recruit it to flat membranes. Here, we confirmed previous observations (Dooley et al., 2014) that WIPI2 is essential for functional E3 recruitment onto membranes resembling flat portions of the ER membrane. Moreover, we discovered that WIP12d does not merely target E3 to membranes but also potently activates the LC3 conjugation reaction catalyzed by E3. This effect would be most readily explained by allosteric communication between the WIP12d binding site on ATG16L1 and the ATG3 binding site on the active ATG12–ATG5 unit (Metlagel et al., 2013; Otomo et al., 2013) of the complex. Alternatively, or in addition, WIP12d might reorient the E3 on the membrane to promote LC3 conjugation (Fig. 6). It has been suggested that the long coiled coil of ATG16L1 could span the gap between the omegasome and the phagophore, such that ATG16L1 anchored by omegasome-localized WIP12 could conjugate LC3 to the phagophore in trans (Wilson et al., 2014). This is an attractive model, but our data show that it is at least possible for WIP12 to promote efficient LC3 lipiddation in cis on the same membrane. Structural studies will be needed to reveal these mechanisms in more detail.

When these two novel principles, the WIP12–PI3KC3-C1 positive feedback and the WIP12 allosteric activation, are combined in a single reaction, a massive dose-dependent acceleration of LC3 conjugation is observed. LC3 lipiddation nears saturation in ∼10 min, which is comparable to what is seen in cells (Axe et al., 2008; Kageyama et al., 2011; Karanasios et al., 2013; Koyama-Honda et al., 2013; Zachari et al., 2019). This is emphasized by the data shown in Fig. 4 E. These data explain the observation of a PI3KC3-C1 requirement for efficient LC3 lipiddation in cells (Dooley et al., 2014; Axe et al., 2008; Itakura et al., 2008; Itakura and Mizushima, 2010; Matsunaga et al., 2009; Zhong et al., 2009) and of ER–Golgi intermediate compartment–derived membranes in a cell-free system (Brier et al., 2019). They also explain the strict requirement for WIP12 for the lipiddation reaction in canonical autophagy (Dooley et al., 2014) and even in STING-induced LC3 lipiddation, which bypasses PI3KC3-C1 but is dependent on WIP12 and the E3 (Gui et al., 2019). These observations place WIP12 at a truly central position in autophagy initiation and autophagosome biogenesis. These observations also highlight how unexpected
Figure 4. PI3KC3-C1 supports LC3B lipidation on unsaturated flat membranes. The schematic drawing illustrates the reaction setting. Colors indicate fluorescent protein–fused components. Components in gray are not labeled but are present in the reaction mix. (A) Representative confocal images of GUVs showing the membrane binding of the PI3KC3-C1 and FYVE domain. GFP-tagged PI3KC3-C1 (200 nM) and mCherry-tagged FYVE (1 µM) were incubated with DO-GUVs (64.8% PC:20% PE:5% PS:10% POPl:0.2% Atto647 DOPE) in the presence or absence of ATP/Mn2+ (50 µM/1 mM) at RT. Images were taken after 30-min incubation. (B) Quantification of the relative intensities of PI3KC3-C1 (green bars) and FYVE domain (red bars) on GUV membranes in A (means ± SD; n = 50). P values were calculated using Student’s t test: ****, P < 0.0001. AU, arbitrary units. (C) Representative confocal images of GUVs showing E3 binding and LC3B lipidation. mCherry-tagged LC3B was incubated with GUVs in the presence of PI3KC3-C1 (0.1 µM), WIPI2d (0.4 µM or none) or WIPI2d FRRG mutant (0.4 µM), E3-GFP, ATG7, ATG3, and ATP/Mn2+ (50 µM/1 mM). Images taken at indicated time points are shown. (D) Quantitation of the kinetics of E3 recruitment and LC3B lipidation on the membrane from individual GUV tracing in C (means ± SD; n = 53 [wt], 45 [FRRG], 52 [-]). AU, arbitrary units; wt, wild type. (E) Quantitation of the kinetics of E3 recruitment and LC3B lipidation on the membrane from individual GUV tracing in C (means ± SD; n = 26 [25 nM], 40 [50 nM], 37 [100 nM], and 32 [400 nM]). mCherry-LC3B was incubated with PI3KC3-C1 (0.1 µM), E3-GFP, ATG7, and ATG3 in the presence of WIPI2d with different concentration. AU, arbitrary units.
Figure 5. Positive feedback between PI3KC3-C1 and WIPI2d promotes LC3B lipidation. (A) Quantitation of the kinetics of FYVE domain or WIPI2d recruitment to the membrane from individual GUV tracing (means ± SD; n = 45 [FYVE], 54 [WIPI2dwt], and 40 [WIPI2dFRRG]). mCherry-FYVE, mCherry-WIPI2d, or mCherry-WIPI2d FRRG mutant (0.5 µM) was incubated with 10% PI(3)P DO-GUVs (64.8% PC:20% PE:5% PS:10% PI(3)P:0.2% Atto647 DOPE). AU, arbitrary units; wt, wild type. (B) Quantitation of the kinetics of FYVE domain or WIPI2d recruitment to the membrane from individual GUV tracing (means ± SD; n = 53 [FYVE], 64 [WIPI2dwt], and 66 [WIPI2dFRRG]). mCherry-FYVE, mCherry-WIPI2d, or mCherry-WIPI2d FRRG mutant (0.5 µM) and PI3KC3-C1 (0.1 µM) were incubated with 10% PI DO-GUVs (64.8% PC:20% PE:5% PS:10% PI(3)P:0.2% Atto647 DOPE) in the presence of ATP/Mn2+ (50 µM/1 mM). AU, arbitrary units; wt, wild type. (C) Representative confocal images showing the membrane binding of the PI3KC3-C1 complex, WIPI2d, WIPI2d FRRG mutant, or FYVE domain. GFP-tagged PI3KC3-C1 (0.1 µM) was incubated with 2% PI(3)P GUVs (72.8% PC:20% PE:5% PS:10% POPI:0.2% Atto647 DOPE) in the absence or presence of 250 nM mCherry-WIPI2d, WIPI2d FRRG mutant, or FYVE domain, respectively (top two panels). 250 nM WIPI2d, WIPI2d FRRG mutant, or FYVE domain was incubated with 2% PI(3)P GUVs in the absence of PI3KC3-C1 (bottom). wt, wild type. (D) Quantitation of the kinetics of PI3KC3-C1, WIPI2d, WIPI2d FRRG, and E3-LC3B recruitment to the membrane from individual GUV tracing (means ± SD; n = 30 [PI3KC3-C1], 33 [WIPI2dwt], 33 [WIPI2dFRRG], and 33 [E3-LC3B]). mCherry-tagged PI3KC3-C1, mCherry-WIPI2d, mCherry-WIPI2d FRRG mutant, or GFP-tagged E3-LC3B (0.1 µM) was incubated with 10% PI(3)P DO-GUVs (64.8% PC:20% PE:5% PS:10% PI(3)P:0.2% Atto647 DOPE) in the presence of ATP/Mn2+ (50 µM/1 mM). AU, arbitrary units; wt, wild type.
properties emerge when multiple steps in autophagy are combined in simultaneous reactions, properties whose quantitative nuances are clarified in the setting of in vitro reconstitution. In addition, the data show how combining reactions that each have their own amplification characteristics can drive the overall process with great increased efficiency, helping to explain how autophagosomes are formed de novo in cells in a matter of minutes.

**Materials and methods**

Sequence data are deposited in the following GenBank accession numbers: ATG3: NP_071933.2; ATG4: NM_013325.

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**Protein expression and purification**

Genes coding for protein sequences of human ATG5, ATG12, ATG16L1 (isoform β), ATG7, and ATG10 were codon optimized for the SF9 insect cell expression system, and synthetic genes were purchased from GenScript. The (10xHis-TEVcs-)ATG16L1-GFP(-TEVs-StrepII) coding sequence was assembled via Gibson strategy using the insect codon-optimized ATG16L1 gene sequence and monomeric GFP gene sequence. All the ORFs and their tags (Table 1) were inserted into pLIB or pBIG library vectors (Weissmann et al., 2016) via classical restriction cloning. Human ATG12, (10xHis-TEVcs-)ATG5 and human ATG7, ATG10 polycystronic constructs were assembled via biGBac system approach (Weissmann et al., 2016) using Gibson assembly strategy. The human ATG12, ATG5, 10xHis-TEVs-ATG16L1(±GFP)-TEVs-StrepII, ATG7, ATG10 polycystronic gene constructs were cloned via Golden Gate approach by the Vienna BioCenter Core Facilities (VBCF) Protech Facility.

2.5 µg of bacmid DNA per construct obtained from amplification in DH10BacY cells was used to transfect 1 million SF9 cells per construct using FuGENE transfection reagent (FuGENE HD; Promega). Virus at P0 was harvested and used to produce a stock Virus P1 solution to further infect a 1-liter culture of Sf9 cells at 0.8 million/ml in SF921 medium containing penicillin-streptomycin. Cultures were harvested when cells reached a viability of maximum 95–98%. They were pelleted down and further washed in 1× PBS at 4,000 rpm for 10 min at 4°C. Pellets were flash frozen in liquid nitrogen and stored at −80°C until purification.

For expression of (10xHis-TEVcs-)ATG5-ATG12 conjugate, Sf9 cells were coinfected with Virus stocks P1 of the two polycystronic constructs coding for ATG12, (10xHis-TEVcs-)ATG5 and ATG7, ATG10, respectively. For purification, pellets were thawed and resuspended in ice cold buffer containing 50 mM Heps, pH 7.5, 300 mM NaCl, 10 mM imidazole, 2 mM MgCl2, 2 mM β-mercaptoethanol, complete protease inhibitors (Roche), Protease Inhibitor Cocktail (Sigma), and Benzonase Nuclease (Sigma). Cells were lysed on ice by extrusion in a tissue homogenizer, and lysates were cleared by ultracentrifugation at 25,000 rpm for 45 min at 4°C using a Beckman Ti45 rotor.
Supernatant was applied to a 5-ml nickel-nitrilotriacetic acid (Ni-NTA) column (GE Healthcare) and eluted via a stepwise imidazole gradient (50, 75, 100, 150, 200, and 300 mM). Protein eluted in fractions containing 150 mM imidazole. These fractions were pooled, concentrated down, applied onto a Superdex 200 Increase 10/300 GL (GE Healthcare), and eluted in a buffer containing 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT. Fractions containing pure 10xHis-TEVcs-ATG5-ATG12 were pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

**Table 1.** List of constructs used in this work, together with specifications about internal identification number (when it applies), vector, expression system, protein encoded, and reference study.

| Identification no.| Vector | Expression system | Encoding | Reference/source |
|------------------|--------|-------------------|----------|------------------|
| SMC1178          | pBiGlB | Sf9               | (10xHis-TEVcs-)ATG5, ATG12 (synthetic gene) | This study |
| SMC1179          | pBiGlA | Sf9               | ATG7, ATG10 (synthetic gene) | This study |
| SMC1099          | pGBdest| Sf9               | ATG12, (10xHis-TEVcs-)ATG5, 10xHis-TEVcs-ATG16L1-TEVcs-StrepII, ATG7, ATG10 (synthetic genes) | This study |
| SMC1100          | pGBdest| Sf9               | ATG12, (10xHis-TEVcs-ATG5), 10xHis-TEVcs-ATG16L1-GFP-TEVcs-StrepII, ATG7, ATG10 (synthetic genes) | This study |
| SMC911           | pFast  | Mouse             | 6xHis-TEVcs-ATG7 | Noor Gammoh laboratory |
| SMC861           | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-ATG3 | This study |
| SMC893           | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-LC3B-Gly(Δ5C) | This study |
| SMC948           | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-mCherry-LC3B-Gly(Δ5C) | Zaffagnini et al., 2018 |
| SMC1199          | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-mCherry-WIPI2d | This study |
| SMC1200          | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-mCherry-WIPI2d | This study |
| SMC872           | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-Atp21 | This study |
| SMC929           | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-mCherry-Atp21 | This study |
| SMC1397          | pET Duet-1 | E. coli Rosetta pLysS | 6xHis-TEVcs-WIPI2d R108,125E | This study |
| SMC1392          | pGEX4T-1 | E. coli Rosetta pLysS | GST-Thrcs-ATG4B | This study |
| Addgene 99329    | pLEXm   | HEKGNti           | GST-TEV-ATG14 (synthetic gene) | Baskaran et al., 2014 |
| Addgene 99328    | pCAG    | HEKGNti           | OSF-TEV-BECN1 (synthetic gene) | Baskaran et al., 2014 |
| Addgene 99327    | pCAG    | HEKGNti           | OSF-TEV-VPS34 (synthetic gene) | Baskaran et al., 2014 |
| N/A              | pCAG    | HEKGNti           | VPS15 (synthetic gene) | Stjepanovic et al., 2017 |
| N/A              | pCAG    | HEKGNti           | GST-TEV-GFP-ATG14 (synthetic gene) | This study |
| N/A              | pCAG    | HEKGNti           | GST-TEV-mCherry-WIPI3 (synthetic gene) | This study |
| N/A              | pCAG    | HEKGNti           | GST-TEV-WIPI3 (synthetic gene) | This study |
| N/A              | pCAG    | HEKGNti           | WIPI2d-TEV-Strep | This study |
| N/A              | pCAG    | HEKGNti           | WIPI2d-TEV-Strep (FRRG to AAAA) | This study |
| N/A              | pCAG    | HEKGNti           | mCherry-WIPI2d-TEV-Strep | This study |
| N/A              | pCAG    | HEKGNti           | mCherry-WIPI2d-TEV-Strep (FRRG to AAAA) | This study |
| N/A              | pLEXm   | HEKGNti           | GST-TEV-mCherry-WIPI4 (synthetic gene) | This study |
| N/A              | pGST2   | E. coli BL21 DE3  | GST-TEV-mCherry-FYVE | This study |

*N/A, not applicable.*

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Reconstituting WIPI2 activation of LC3 lipiddation
For expression of ATG12-(10xHis-TEV-)ATG5-(10xHis-TEVcs-)ATG16L1-(6xHis)-TEVcs-StrepII complexes, S9 cells were infected with a single Virus stock P1 corresponding to the poli-cystronic construct coding for GFP-tagged (or not) E3 complex. Cell pellets were thawed and resuspended in ice-cold buffer containing 50 mM Hepes, pH 7.5, 300 mM NaCl, 2 mM MgCl₂, 1 mM DTT, complete protease inhibitors, Protease Inhibitor Cocktail, and Benzonase Nuclease. Cells were lysed on ice by extrusion in a tissue homogenizer, and lysates were cleared by ultracentrifugation at 25,000 rpm for 45 min at 4°C in a Ti45 rotor. Supernatant was applied to a 5-ml StreptTactin column (GE Healthcare) and eluted with 2.5 mM desthiobiotin in 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT. Fractions containing the protein complex were pooled, concentrated down, applied onto a Superdex 6 column (Increase 10/300; GE Healthcare), and eluted with a buffer containing 25 mM Hepes, pH 7.5, 300 mM NaCl, and 1 mM DTT. Fractions containing pure ATG12-(10xHis-TEV-)ATG5-(10xHis-TEVcs-)ATG16L1-(6xHis)-TEVcs-StrepII complexes were pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

Mouse (6xHis-TEVs-)ATG3 was expressed in Escherichia coli Rosetta pLysS cells. Cells were grown in Luria Bertani (LB) medium at 37°C until an OD₆₀₀ of 0.4. The culture was then brought to 18°C and grown to an OD₆₀₀ of 0.8. Protein expression was induced with 100 µM IPTG and grown for a further 16 h at 18°C. Cells were pelleted and resuspended in a buffer containing 50 mM Hepes, pH 7.5, 300 mM NaCl, 10 mM imidazole, 2 mM MgCl₂, 2 mM β-mercaptoethanol, 1 mM Pefablock, complete protease inhibitors, and DNase (Sigma). Cells were lysed by freeze thawing and 2 × 30-s sonication. Lysates were cleared by ultracentrifugation (40,000 rpm for 30 min at 4°C in a Ti45 rotor). Supernatant was filtered (0.45 µm) applied to a 5-ml Ni-NTA column and eluted via a stepwise imidazole gradient (50, 75, 100, 150, 200, and 300 mM). Fractions containing the protein of interest were pooled, concentrated down, and applied onto a Superdex 75 column (16/60 prep grade; GE Healthcare), and eluted with a buffer containing 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT. Fractions containing pure 6xHis-TEV-LC3B Δ5C or 6xHis-TEV-mCherry-LC3B Δ5C proteins were pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

WIPITd gene coding sequence was amplified from a HEpG2 cDNA library and cloned via classical restriction cloning in a PET-Duet vector, with and without a N-terminal mCherry tag, in frame with an N-terminal 6xHis tag. The (6xHis-TEV-)WIPITd R108E,125E mutant construct was obtained by consecutively inserting the two point mutations in the sequence of the WT gene sequence using Round-The-Horn PCR strategy. Primers for mutation of R108E were forward (fw) 5′-GAGACAGG CTGATAAGATG-3′, reverse (rv) 5′-CTTGAGCTTCACAGCCG-3′; primers for mutation of R125E were fw 5′-GAAGACATGAA AGTGCAGCATC-3′, rev 5′-AAATGTTGGATGCAGGGACTC-3′. Primers for mutation of FRGG motif were fw 5′-GGACAAACTC TTTGAGGCTGGCCAGCTGATGAAAGAGGTGTGAGCCG-3′, rev 5′-GCT CACCGACCTTTTACTGTCGCCAGCGAGTCTTCGTGGGAGCC-3′. (6xHis-TEV-)WIPITd WT and R108E,125E mutant and (6xHis-TEV-)mCherry-WIPITd were expressed in E. coli Rosetta pLysS cells. Cells were grown in Terrific Broth medium at 37°C until an OD₆₀₀ of 0.4. Next, the culture was brought to 18°C and grown to an OD₆₀₀ of 0.8. Protein expression was induced with 100 µM IPTG and grown for a further 16 h at 18°C. Cells were pelleted and resuspended in a buffer containing 50 mM Hepes, pH7.5, 300 mM NaCl, 10 mM imidazole, 2 mM MgCl₂, 2 mM β-mercaptoethanol, complete protease inhibitors, and DNase. Cells were lysed by freeze thawing and 2 × 30-s sonication. Lysates were cleared by ultracentrifugation (25,000 rpm for 30 min at 4°C using a Ti45 rotor). Supernatant was filtered (0.45 µm) applied to a 5-ml Ni-NTA column and eluted via a stepwise imidazole gradient (50, 75, 100, 150, 200, and 300 mM). Fractions at 100–150 mM imidazole containing the proteins of interest were pooled, concentrated down, applied onto a Superdex 200 column (16/60 prep grade), and eluted with a buffer containing 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT. Fractions containing pure protein are pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

The S. cerevisiae Atg21 gene coding sequence was amplified from an S. cerevisiae cDNA library and cloned via classical restriction cloning in a pET-Duetl vector, with a N-terminal 6xHis tag followed by a TEV cleavage (and mCherry tag where it applies) site in frame with the protein coding sequence. (6xHis-TEV-)Atg21 and (6xHis-TEV-)mCherry-Atg21 were expressed in E. coli Rosetta pLysS cells. Cells were grown in Terrific Broth medium (for Atg21) or LB medium (for mCherry-Atg21) at 37°C until an OD₆₀₀ of 0.4. Next, the culture was brought to 18°C and grown to an OD₆₀₀ of 0.8. Protein expression was induced with
100 µM IPTG and grown for a further 16 h at 18°C. Cells were pelleted and resuspended in a buffer containing 50 mM Hepes, pH 7.5, 300 mM NaCl, 10 mM imidazole, 2 mM MgCl₂, 2 mM β-mercaptoethanol, complete protease inhibitors, and DNase. Cells were lysed by freeze thawing and 2 × 30-s sonication. Lysates were cleared by ultracentrifugation (40,000 rpm for 30 min at 4°C in a Ti45 rotor). Supernatant was filtered (0.45 µm) applied to a 5-ml Ni-NTA column and eluted via a stepwise imidazole gradient (50, 75, 100, 150, 200, and 300 mM). Fractions at 100–150 mM imidazole containing the proteins of interest were pooled. For mCherry-Atg21, the 6xHis tag was removed with o.n. cut at 4°C using TEV protease. Consequently, pooled fractions were concentrated down and applied onto a Superdex 200 column (16/60 prep grade) and eluted with a buffer containing 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT. Fractions containing pure (±mCherry) Atg21 protein were pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

The Homo sapiens ATG4B gene coding sequence was amplified from a HeLa cDNA library and cloned via classical restriction digestion at pGEX4T1 vector, with an N-terminal GST tag followed by a thrombin cleavage site in frame with the protein cloning in a pGEX4T1 vector, with an N-terminal GST tag followed by a thrombin cleavage site in frame with the protein

The PI3KC3-C1 complex was expressed and purified from HEK293 GrT1 cells as described previously (Chang et al., 2019). ATG14, VPS34, VPS15, and BECN1 constructs were transfected to cells using polyethylenimine (Polysciences). After 60 h of expression, cells were harvested and lysed with lysis buffer (50 mM Hepes, pH 7.4, 1 mM Triton X-100, 200 mM NaCl, 1 mM MgCl₂, 10% glycerol, and 1 mM tris(2-carboxyethyl)phosphine (TCEP)) supplemented with EDTA-free protease inhibitors (Roche). The lysate was clarified by centrifugation (15,000 rpm for 1 h at 4°C) and incubated with glutathione Sepharose 4B (GE Healthcare) for 4 h at 4°C, applied to a gravity column, and washed extensively with wash buffer (50 mM Hepes, pH 8.0, 200 mM NaCl, 1 mM MgCl₂, and 1 mM TCEP). The protein complexes were eluted with wash buffer containing 50 mM reduced glutathione and treated with TEV protease at 4°C o.n. TEV-treated complexes were loaded on a StrepTactin Sepharose gravity flow column (IBA, Gmbh). The complexes were eluted with wash buffer containing 10 mM desthiobiotin (Sigma) and applied to Superose 6 16/50 column equilibrated with gel filtration buffer (20 mM Hepes, pH 8.0, 200 mM NaCl, 1 mM MgCl₂, and 1 mM TCEP). Peak fractions were collected and used immediately for subsequent assays.

GST-mCherry-WIPI3, GST-mCherry-WIPI4, WIPI2d-Strep, WIPI2d FRG mutant-Strep, mCherry-WIPI2d-Strep, and mCherry-WIPI2d FRG mutant-Strep were purified from HEK293 GrT1 cells by a similar protocol used for PI3KC3-C1 purification. The TEV-treated proteins were directly applied to a Superdex 200 column (16/60 prep grade) equilibrated with gel filtration buffer (20 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM TCEP). Fractions containing pure proteins were pooled, concentrated, snap frozen in liquid nitrogen, and stored at −80°C.

SLS
A sample of 90 µl at 10 µM (1.5 mg/ml) of purified ATG12–ATG5–ATG16L1-GFP was applied to a Superose 6 Increase 10/300 GL column in 25 mM Hepes, pH 7.5, 300 mM NaCl, and 1 mM DTT. The column was coupled to a TRESOS II instrument (Wyatt). Data were analyzed using the ASTRA V software (Wyatt).

Preparation of GUVs
GUVs were prepared by electroformation (Romanov et al., 2012) or hydrogel-assisted swelling (Weinberger et al., 2013), as described previously. To prepare GUVs by electroformation, lipids were desiccated for 5 h, and the electroformation was conducted at 30°C in a 309 mOsm sucrose solution. GUVs were used for experiments directly afterward. To prepare GUVs by gel-assisted swelling, polyvinyl alcohol (PVA) with a molecular weight of 145,000 (Millipore) was used as hydrogel substrate. PVA was mixed with water to obtain a 5% (wt/wt) PVA solution and stirred on a heat plate at −90°C until the solution is clear. 300 µl of 5% PVA was then spun coat for 30 s at a speed of 1,200 rpm on a plasma-cleaned cover glass with a 25-mm diameter. To dry the PVA film, the coated glass was placed for 30 min in a heating incubator at 60°C. 10–15 µl of lipids with different compositions at 1 mg/ml stock were deposited uniformly on the PVA film. Details about different lipid compositions are indicated in figure legends. The lipid-coated cover glass was put under vacuum o.n. to evaporate the solvent from the dissolved lipid mixture. The coated cover glass was transferred into a 30-mm dish, and 300 µl of 400 mOsm sucrose solution was added on top of the glass. After swelling for ~1 h at RT, the vesicles were harvested and then stored at RT and used immediately.

Atto647N dioleoylphosphatidylethanol-amine (DOPE; 1 mg/ml, AD 647N-161; Atto TEC) was used as a GUV membrane dye. All the other lipids for GUVs preparation are from Avanti Polar Lipids. GUVs with DO lipids contained dioleoylphosphatidylcholine (DOPC; 850375C, 10 mg/ml), dioleoylphosphatidylethanol-amine (DOPE; 850725C, 10 mg/ml), dioleoylphosphatidylserine (DOPS; 840035C, 10 mg/ml), and palmitoyl oleoylphosphatidyl-nositol (POPI; 850142P, 1 mg/ml). GUVs with PO lipids contained POPC (850457C, 10 mg/ml), POPE (850757C, 10 mg/ml), and dipalmitoylphosphatidylcholine (DPPC; 850726C, 10 mg/ml)
palmitoyloleoylphosphatidylserine (POPS; 840034C, 10 mg/ml), and POPI (850142P, 1 mg/ml). GUVs with brain lipids contained brain PC (840053C, 10 mg/ml), brain PE (840022C, 10 mg/ml), brain PS (840032C, 10 mg/ml), and liver PI (840042C, 10 mg/ml). PI(3)P (850150P, 1 mg/ml) was used for GUVs containing PI(3)P.

Preparation of SUVs
For preparation of SUVs, lipids were mixed to homogeneity in chloroform, dried under a stream of argon, and further desiccated for 1 h under vacuum. The lipid film was rehydrated in 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT. The lipid film was resuspended by gentle mixing at RT and sonicated for 2 min in a bath sonicator. The resuspended SUVs were then extruded 21 times through a 0.4-µm membrane followed by 21 times through a 0.1-µm membrane (Whatman) using the Mini Extruder from Avanti Polar Lipids. The final SUV suspension has a concentration of 1 mg lipids/ml.

Lipids used for SUVs preparation were from Avanti Polar Lipids: DOPC (850375C, 10 mg/ml) and DOPE (850725C, 10 mg/ml). Alternatively, when PO-fatty acid chains were used, SUVs contained POPC (850457C, 10 mg/ml) and POPE (850757C, 10 mg/ml). Other lipids, ATTO390-PE (ATTO-TEC, 1 mg/ml) and PI(3)P (850150P, 1 mg/ml), prepared as in Fracchiolla et al. (2016), and 15% liver PI (840042C, 10 mg/ml).

Liposome sedimentation assay
SUVs prepared as described above were mixed with 1–2-µg protein mixes at a ratio of 1:1 in buffer containing 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT. Reactions were incubated with 0.5 mg/ml liposomes for 30 min at RT (22°C). Next, the reactions were centrifuged at 180,000 × g at 22°C, supernatants and pellets were separated, and equal amounts were run on 4–15% SDS-polyacrylamide gels and stained with Coomassie Brilliant Blue.

SUVs membrane recruitment assay on beads
For experiments shown in Fig. S3, B and C, GFP-Trap (Fig. S3 B) or RFP-Trap (Fig. S3 C) beads (Chromotek) were first coated with the ATG12–ATG5–ATG16L1–GFP protein (Fig. S3 B) or mCherry or mCherry-WIPI2d or mCherry-Atg21 proteins (Fig. S3 C), and where it applies, upon washings with buffer containing 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT, were further incubated with mCherry-tagged Proppins (Fig. S3 B). After washings, 1–2 µl of beads per sample was pipetted into the wells of a 384-well glass-bottomed microplate (Greiner Bio-One) prefilled with a prep of ATTO390-SUVs in buffer containing 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT at a concentration of 0.2 mg/ml.

Membrane protein recruitment: GUV assay
For experiment shown in Fig. 1 B, 15 µl of the ATG12–ATG5–ATG16L1–GFP protein was added to 15 µl of GUVs prepipetted into the wells of a 384-well glass-bottomed microplate precoated with a 5-mg/ml BSA solution in buffer containing 25 mM Hepes, pH 7.5, and 150 mM NaCl for a final concentration of 500 nM mCherry-LC3B

buffer containing 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT. For experiment shown in Fig. 2 B, 15 µl of the ATG12–ATG5–ATG16L1–GFP protein at 500 nM and the mCherry-WIPI2d at 500 nM were added to 15 µl of GUVs. Concentrations are calculated for a final volume of 30 µl. The images were acquired after 30 min of incubation at RT (22°C) in the dark using an LSM700 confocal microscope (Zeiss) with a 20×/0.8 Plan Apochromat objective (lasers: 488 nm, 10 mW; 555 nm, 10 mW), controlled by Zeiss ZEN 2012 software and processed with ImageJ software. Identical laser power and gain settings were used during the acquisition of all conditions. For experiment shown in Fig. S3 D, 15 µl of the ATG12–ATG5–ATG16L1–GFP at 100 nM and WIPI2d at 200 nM (or buffer) were added to 15 µl of GUVs. Concentrations are calculated for a final volume of 30 µl. The images were acquired after at least 30 min of incubation at RT (22°C) in the dark using a spinning disc microscope (Visiport) with a 63×/1.4 Oil differential interference contrast (DIC) objective and processed with ImageJ software. Identical laser power and gain settings were used during the acquisition of all conditions.

In vitro bulk LC3B lipidation assay
Reactions were set up by mixing 1:1 volume of SUVs and protein mix. Protein mix was prepared in buffer containing 25 mM Hepes, pH 7.5, 137 mM NaCl, 2.7 mM KCl, and 0.1 mM DTT at a final concentration of 1 µM mouse ATG7, 1 µM human ATG3, 5 µM LC3BΔSC, and 1 µM E3-like ligase or ATG12–ATG5 conjugate (Fig. 1 A). 0.1 µM E3-like ligase was used in experiments shown in Fig. 2 F together with 0.5 µM WIPI2d. 0.5 µM E3-like ligase was used in experiments shown in Fig. 3, E and H, together with 2.5 µM Atg21 or WIPI2d WT or mutant. Reactions were conducted on a thermoblock heating element at 37°C in the presence of 5 mM ATP and 1 mM MgCl2. 15 µl of reaction was sampled at each time point, mixed with 3 µl of 6x Protein Loading dye, boiled at 60°C for 10 min, and loaded on SDS-polyacrylamide gels followed by staining with Coomassie Brilliant Blue.

In vitro reconstitution of LC3B lipidation on GUVs
For experiments shown in Fig. 1 C, proteins were mixed in 15 µl and added to 15 µl of GUVs prepipetted into the wells of a 384-well glass-bottomed microplate, for a final concentration of 100 nM mouse ATG7, 100 nM human ATG3, 500 nM E3-GFP, and 500 nM mCherry-LC3BΔSC, together with 0.5 mM ATP and 0.5 mM MgCl2. The reaction buffer contains 25 mM Hepes, pH 7.5, 150 mM NaCl, and 0.1 mM DTT.

For experiments shown in Fig. 2 C, proteins were mixed in 15 µl and added to 15 µl of GUVs prepipetted into the wells of a 384-well glass-bottomed microplate, for a final concentration of 100 nM mouse ATG7, 100 nM human ATG3, 100 nM E3-GFP, and 500 nM mCherry-LC3BΔSC, together with 0.5 mM ATP and 0.5 mM MgCl2. Concentrations of proteins and cofactors used are calculated for a final volume of 30 µl of reaction. The images were acquired after 30 min of incubation at RT (22°C) in the dark (Fig. 2 C) using an LSM700 confocal microscope with a 20×/0.8 Plan Apochromat objective (lasers: 488 nm, 10 mW; 555 nm, 10 mW),
controlled by ZEN 2012 software and processed with ImageJ software.

For de-lipidation experiments, 10 U of calf intestinal phosphatase (CIP) was added to the well together with ATG4B at a final concentration of 1 µM, or buffer in the negative control and GUVs imaged under a spinning disc microscope equipped with a 63x/1.4 Oil DIC objective and an electron multiplying charge-coupled device camera.

For experiments with PI3KC3-C1, the reactions were performed at RT in an observation chamber (Lab Tek) coated with 5 mg/ml β casein. 15-20 µl of GUVs was added at last to initiate the reaction with a final volume of 150 µl. The reaction buffer contained 20 mM Hepes, pH 8.0, 190 mM NaCl, and 1 mM TCEP. A final concentration of 100 nM E3-GFP, 100 nM ATG7, 100 nM ATG3, and 500 nM mCherry-LC3BΔ5C was used for all lipiddation reactions with PI3KC3-C1. The final concentrations of PI3KC3-C1, WIP12d, or FYVE domain used in different experiments are indicated in figure legends. After 5-min incubation, during which we picked random views for imaging, time-lapse images were acquired in multitracking mode on a Nikon A1 confocal microscope with a 63× Plan Apochromat 1.4 NA objective. Identical laser power and gain settings were used during the course of all conditions.

**Microscopy-based bead protein–protein interaction assay**

mCherry, mCherry-Atg21, and mCherry-WIP12d bait proteins are incubated at a concentration of 40 µM each with 10 µl of RFP-Trap beads in buffer containing 25 mM Hepes, pH 7.5, 150 mM NaCl, and 1 mM DTT (Fig. 3 D) or GFP-Trap beads (Chromotek) when E3-GFP was used (Fig. 2 A). Beads are incubated with proteins for at least 30 min at 4°C on a rotating wheel and washed three times in 100 µl of buffer. 1 µl of beads taken from a 1:1-volume beads/buffer mix was pipetted onto the well of a 384-well glass-bottomed microplate precoated with a 5-mg/ml BSA solution in buffer containing 25 mM Hepes, pH 7.5, and 150 mM NaCl, containing 15 µl of the ATG12–ATG5–ATG16L1-GFP prey protein at 5 µM. Beads were incubated for at least 30 min at RT (22°C) and imaged using an LSM700 confocal microscope with an A1 confocal microscope using a 63x/1.4 Oil DIC objective with a 100% laser intensity and 10 s/pixel. Quantifications and statistical analysis

For quantification shown in Fig. S3 D, multiple lines were manually drawn across the GUV membranes identified in the bright-field channel. GFP signal intensity was measured at these positions as the maximum intensity signal along the line. Averages and SDs were calculated from multiple datasets, and the average and SD were calculated and are reported.

For FRAP experiments in Fig. 2 D, a line was manually drawn across the GUV membrane identified in the GFP channel at time 0. The position of the line was checked throughout the time course to cross the GUV membrane. GFP and mCherry signal intensities were simultaneously measured at these positions as the maximum intensity signal along the line. A GUV-free area at time point 0 was chosen to measure the background value for the mCherry channel as mean value. This was subtracted from the mCherry intensities measured for all the GUVs of the field of view along the time course. mCherry and GFP signal at time 0 were set to 100%, and the corresponding values of the time course were calculated relative to this. Data are plotted in a scatter graph as a function of time.

For FRAP experiments in Fig. 2 D, a line was manually drawn across the GUV membrane within the area subjected to photo-bleaching. mCherry maximum intensity signal along the line
was measured. The value corresponding to the prebleached time point was set as 100%, and all the values measured after photobleaching were related to this and are plotted in a scatter graph as function of time. The P values were calculated using an unpaired two-tailed Student’s t test. P values were considered as follows: not significant (NS), P ≥ 0.05; *, 0.01 < P < 0.05; **, 0.001 < P < 0.01; ***, 0.0001 < P < 0.001; and ****, P < 0.0001. Data distribution was always assumed to be normal, but this was not formally tested.

**Bead images quantification**

ImageJ software was used for bead images quantifications. Multiple lines were manually drawn across the beads. These lines were saved as regions of interest. mCherry and GFP signal intensity were simultaneously measured at these regions of interest as the maximum intensity signal along the line. Averages and SDs were calculated among the measured values per each condition and are plotted in a bar graph.

**SUV sedimentation**

The gels of three independent experiments were quantified using the Analyze Gel tool of ImageJ software. The amounts of pelleted protein per each condition were measured as the area below the corresponding peak. The measured amount of pelleted protein in the absence of SUVs was subtracted from the measured amount of pelleted protein in the presence of SUVs. These values were plotted as relative or absolute pelleted protein amounts in bar graphs.

**In vitro LC3B lipidation on SUVs**

The gels of three independent experiments were quantified using the Analyze Gel tool of ImageJ software. The amounts of LC3B-I and LC3B-II at each time point were measured as the area below the corresponding peak. The sum of the values corresponding to LC3B-I and LC3B-II peaks was then calculated at each time point, and the fraction of LC3B-II was calculated. Finally, the relative LC3B-II fraction amounts were calculated at each time point and are plotted as a function of time.

When a contaminant band of WIPI2d batch with a running behavior similar to that of LC3B-II was present and overlapped with the protein of interest (Fig. 3 D), the value corresponding to the contaminant protein peak present at time point 0 was subtracted from each value measured for the LC3B-II band in the following time points. Instead, in Fig. 3 F, the value assigned to LC3B-II was arbitrarily set to 0, as both WT and mutant proteins had the same contaminant band, not present at the following time point. The analysis followed as described above.

**Online supplemental material**

Fig. S1 shows biochemical characterization of the E3 and the LC3 lipidation machinery. Fig. S2 shows PI(3)P- and WIPI2d-dependent E3 recruitment to GUVs and LC3 lipidation. Fig. S3 shows the role of WIPI2d binding to the E3 complex goes beyond its membrane recruitment. Fig. S4 shows PI3KC3-C1 is active on unsaturated flat membranes. Fig. S5 shows WIPI3 mediates PI3KC3-C1-triggered LC3 lipidation on GUV membranes.

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Reconstituting WIPI2 activation of LC3 lipidation

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Figure S1. **Biochemical characterization of the E3 and the LC3 lipidation machinery.** (A) Purified human E3-GFP complex resolved on a 10% SDS-polyacrylamide gel and stained with Coomassie Brilliant Blue. MW, molecular weight. (B) SLS plot of recombinant E3-GFP. The protein was applied onto a Superose 6 Increase 10/300 GL column coupled with a TREOS II instrument. BSA was used for calibration. (C) Schematic representation of the dimeric E3-like ligase holo-complex containing two copies of each subunit of ATG12, ATG5, and ATG16L1(+monoGFP). (D) Recombinant mouse ATG7, human ATG3, and LC3B resolved on a 10% SDS-polyacrylamide gel and stained with Coomassie Brilliant Blue. MW, molecular weight. (E) In vitro LC3B lipidation assay using PO-SUVs (65% PC:15% liver PI:20% PE), ATG7, ATG3, E3-GFP (1 µM), and LC3B incubated at 37°C in the presence of MgCl2/ATP. Samples taken at the indicated time points (o.n.) were loaded on a 4–15% SDS-polyacrylamide gel. Time points corresponding to o.n. incubation indicated with an asterisk (*) were swapped during loading. MW, molecular weight. (F) ATG7, ATG3, E3-GFP (1 µM), and LC3B were incubated with DO-SUVs (65% PC:15% PI:20% PE) at 37°C with MgCl2 and ATP. Samples at the indicated time points (o.n.) were loaded on a 4–15% SDS-polyacrylamide gel. MW, molecular weight. (G) Co-sedimentation assay of ATG12–ATG5 with DO-lipid (left) or PO-lipid (right) SUVs (65% PC:35% PI:20% PE). MW, molecular weight. (H) Co-sedimentation assay of the E3 with DO-lipid (left) or PO-lipid (right) SUVs with the indicated lipid composition. MW, molecular weight.
Figure S2. PI(3)P- and WIPI2d-dependent E3 recruitment to GUVs and LC3 lipiddation. (A) GST-mCherry–tagged WIPI3 or WIPI4 was incubated with GFP-tagged E3 and glutathione-coated polystyrene beads. Representative confocal images taken after 30-min incubation are shown. (B) E3 was coincubated with mCherry-WIPI2d and DO-GUVs containing either 65% PC:20% PE:15% PI (bearing the same net negative charge as GUVs in Fig. 2 B) or GUVs containing 80% PC:20% PE (zero net charge). (C) mCherry-LC3B was coincubated with ATG7 and ATG3 in the presence or absence of E3-GFP or WIPI2d with GUVs containing either 75% PC:20% PE:5% PI(3)P (as GUVs in Fig. 2 C) or GUVs containing 80% PC:20% PE (zero net charge).
Figure S3. Role of WIPI2d binding to the E3 complex goes beyond its membrane recruitment. (A) Co-sedimentation assay using DO-SUVs [75% PC:20% PE:5% PI(3)P] and the E3-GFP complex in the presence or absence of WIPI2d or Atg21. Quantification of three independent experiments (means ± SD; n = 3) is shown on the right. P, pellet; MW, molecular weight; ns, not significant; S, supernatant. (B) GFP-Trap beads coated with E3-GFP and mCherry-WIPI2d or mCherry-Atg21 or no PROPPINs were incubated with DO-SUVs containing 72% PC:20% PE:5% PI(3)P:3% ATTO390-PE. Quantification of three independent experiments (means ± SD; n = 3) is shown. AU, arbitrary units; n.s., not significant. (C) RFP-Trap beads coated with mCherry, mCherry-WIPI2d, or mCherry-Atg21 were incubated with DO-SUVs containing 72% PC:20% PE:5% PI(3)P:3% ATTO390-PE. Quantification of three independent experiments (means ± SD; n = 3) is shown. P values were calculated using Student’s t test: **, 0.001 < P < 0.01. AU, arbitrary units. (D) E3-GFP (0.1 µM) recruitment to DO-GUVs containing 75% PC:20% PE:5% PI(3)P in the presence of wild-type (wt) WIPI2d (0.2 µM) or R108,125E mutant WIPI2d (0.2 µM) or no PROPPINs (−). The E3-GFP signal on GUVs was quantified and plotted (means ± SD; n wt = 107, n mut = 197, n no PROPPINs = 170). A blot probed for WIPI2 shows the protein input. AU, arbitrary units. (E) Coomassie-stained gel showing equal amounts of wild-type (wt) and R108,125E mutant proteins used in the bulk lipidation assays of Fig. 3 G and H. MW, molecular weight. (F) Co-sedimentation assay using DO-SUVs [75% PC:5% PI(3)P:20% PE] and the wild-type (wt) WIPI2d or R108,125E WIPI2d. MW, molecular weight.
Figure S4. **PI3KC3-C1 is active on unsaturated flat membranes.** (A) Representative confocal images of GUVs showing the binding of the PI3KC3-C1 complex and FYVE domain on different membranes. mCherry-FYVE domain (1 µM) was incubated with GUVs with brain lipids (64.8% brain PC:20% brain PE:5% brain PS:10% liver Pl:0.2% Atto647 DOPE), PO lipids (64.8% POPC:20% POPE:5% POPS:10% POPI:0.2% Atto647 DOPE), or DO lipids (64.8% DOPC:20% DOPE:5% DOPS:10% POPI:0.2% Atto647 DOPE) in the absence or presence of GFP-tagged PI3KC3-C1 (200 nM) for 30 min. (B) Quantification of the relative intensities of PI3KC3-C1 on different GUV membranes (means ± SD; n = 50). P values were calculated using Student’s t test: ****, P < 0.0001. AU, arbitrary units. (C) Quantification of the relative intensities of FYVE domain on different GUV membranes (means ± SD; n = 50). P values were calculated using Student’s t test: ****, P < 0.0001. AU, arbitrary units.
Figure S5. **WIPI3 mediates PI3KC3-C1 trigged LC3 lipidation on GUV membranes.** (A) Representative confocal images of GUVs showing E3 membrane recruitment and LC3B lipidation. mCherry-tagged LC3B was incubated with PI3KC3-C1 (100 nM), WIPI3 (400 nM), E3-GFP, ATG7, and ATG3 in the presence or absence of ATP. (B) Quantitation of the kinetics of E3 recruitment and LC3B lipidation on the membrane from individual GUV tracing in (A) (means ± SD; n = 20; 15). AU, arbitrary units.