ATG9A regulates proteostasis through reticulophagy receptors FAM134B and SEC62 and folding chaperones CALR and HSPB1.
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SUMMARY
The acetylation of ATG9A within the endoplasmic reticulum (ER) lumen regulates the induction of reticulophagy. ER acetylation is ensured by AT-1/SLC33A1, a membrane transporter that maintains the cytosol-to-ER flux of acetyl-CoA. Defective AT-1 activity, as caused by heterozygous/homozygous mutations and gene duplication events, results in severe disease phenotypes. Here, we show that although the acetylation of ATG9A occurs in the ER lumen, the induction of reticulophagy requires ATG9A to engage FAM134B and SEC62 on the cytosolic side of the ER. To address this conundrum, we resolved the ATG9A interactome in two mouse models of AT-1 dysregulation: AT-1 sTg, a model of systemic AT-1 overexpression with hyperacetylation of ATG9A, and AT-1S113R/+, a model of AT-1 haploinsufficiency with hypoacetylation of ATG9A. We identified CALR and HSPB1 as two ATG9A partners that regulate the induction of reticulophagy as a function of ATG9A acetylation and discovered that ATG9A associates with several proteins that maintain ER proteostasis.

INTRODUCTION
Ne-lysine acetylation in the endoplasmic reticulum (ER) maintains protein homeostasis (proteostasis) within the organelle and secretory pathway (reviewed in Farrugia and Puglielli, 2018). In particular, the acetylation status of the autophagy protein ATG9A regulates the induction of ER-specific autophagy (also referred to as reticulophagy or ER-phagy) and the disposal of toxic protein aggregates that form in the lumen of the organelle (Pehar et al., 2012; Peng et al., 2014, 2016, 2018). ER acetylation is ensured by AT-1 (also referred to as SLC33A1), a membrane transporter that translocates acetyl-CoA from the cytosol to the ER lumen, and ATase1 (also referred to as NAT8B) and ATase2 (also referred to as NAT8), two ER-bound type II membrane proteins that acetylate ER-resident and -transiting proteins (Jonas et al., 2010; Ko and Puglielli, 2009).

In humans, defective ER acetylation, as caused by loss-of-function mutations or gene duplication events of AT-1/SLC33A1, is linked to hereditary spastic paraplegia (heterozygous mutation), developmental delay and progeria-like dysmorphism (gene duplication) (Chipulkar et al., 2016; Huppke et al., 2012a; Lin et al., 2008; Prasad et al., 2012; Sanders et al., 2011). These disease phenotypes are efficiently mimicked by mouse models of reduced or increased AT-1 activity (Hullinger et al., 2016; Peng et al., 2014, 2018). Importantly, reduced ER acetylation in AT-1s113r/− hypomorphic mice causes increased (aberrant) induction of reticulophagy, whereas increased ER acetylation in AT-1-overexpressing (AT-1 sTg) mice has the opposite effect (Hullinger et al., 2016; Peng et al., 2014, 2018). Furthermore, inhibition of ATase1 and ATase2, downstream of AT-1, can restore reticulophagy and disposal of protein aggregates from the ER lumen in AT-1 overexpressors (Peng et al., 2016, 2018).

We recently reported that the acetylation status of ATG9A regulates its ability to interact with FAM134B (also referred to as RETREG1) and SEC62 (Peng et al., 2018), two ER membrane-bound proteins that can act as receptors for LC3B through their LC3B-interacting region (LIR) (Fumagalli et al., 2016; Khaminets et al., 2015; Lennemann and Coyne, 2017; Mochida et al., 2015). Importantly, the acetylation of ATG9A occurs on two lysine residues (K359 and K363) that face the ER lumen, whereas the LIR of FAM134B and SEC62 is on the C terminus of both proteins, exposed to the cytosolic side of the ER membrane (Fumagalli et al., 2016; Khaminets et al., 2015; Lennemann and Coyne, 2017; Mochida et al., 2015; Pehar et al., 2012). This
topological conundrum is further complicated by the fact that FAM134B appears to assemble within the ER membrane with a reticulon-like conformation and has no available domain exposed to the ER lumen that could engage the acetylated loop of ATG9A (Bhaskara et al., 2019).

In this study, we dissected the structural aspects of the ATG9A-FAM134B and ATG9A-SEC62 interaction, resolved the ATG9A interactome, and took advantage of AT-1S113R/+ and AT-1 sTg mice to discover two ATG9A-interacting proteins—calreticulin (CALR) and heat shock protein beta 1 (HSPB1)—which engage from the lumenal and cytosolic sides of the ER membrane, respectively, to regulate the induction of reticulophagy. Collectively, our results disclose a previously unidentified aspect of the complex mechanisms that allow the ER acetylation machinery to maintain proteostasis within the ER and secretory pathway. Our findings also disclose pathogenic events underlying the disease phenotypes caused by dysfunctional ER acetylation.

RESULTS
FAM134B and SEC62 engage ATG9A on the ER membrane with their cytosolic exposed C terminus

Membrane modeling suggests that FAM134B assembles with a reticulon-like conformation within the ER membrane to force curvature of the lipid bilayer (Figure 1A) (Bhaskara et al., 2019). However, different topology and disorder-prediction platforms suggest the presence of an ER lumen-exposed loop between amino acid (aa) 140 and 208 with two disordered segments of about 10 aa each (aa156–169 and aa187–198). Therefore, it is theoretically possible that disordered segments of FAM134B could move out of the ER membrane to interact with ATG9A within the ER lumen. To test this possibility, we introduced the Asn-Gly-Thr consensus sequence for N-glycosylation within the aa150–200 region of FAM134B. We reasoned that, if these segments were indeed exposed to the ER lumen, we would observe a migration shift...
caused by N-glycosylation of the Asn residue. Specifically, we introduced the Asn-Gly-Thr sequence (Dempski and Imperiali, 2002; Petrescu et al., 2004) at six independent locations. We also mutated N177 to neutralize a potential N-glycosylation motif within the aa140–208 segment of FAM134B. We only observed a migration shift of FAM134B when the Asn-Gly-Thr sequence was placed at aa200 (Figures 1B and 1C). Furthermore, no migration shift occurred with the N177D substitution (Figure 1B). Collectively, these results indicate that the aa140–208 region of FAM134B has limited movement and, therefore, is unlikely to engage ATG9A from the luminal side of the ER. These results also support the “hairpin” topology model proposed by Bhaskara et al. (2019).

FAM134B is solely found in the ER, whereas ATG9A can be found on different cellular locations, which highlights its complex biological functions (Imai et al., 2016; Lee and Finkel, 2009; Nishimura et al., 2017; Ohashi and Munro, 2010; Tamura et al., 2010; Webber et al., 2007; Young et al., 2006). As ATG9A is acetylated by ATase1 and ATase2 within the ER lumen (Pehar et al., 2012; Peng et al., 2018), it is reasonable to assume that only the immature and ER-based form of ATG9A can engage FAM134B. To test this hypothesis, we transfected HEK293 cells with both ATG9A-GFP and FAM134B-DDK. Immunoprecipitation of FAM134B only pulled down the ER-based and immature form of ATG9A, thus supporting our hypothesis (Figures 1D and 1E). This was further confirmed by the visualization of ATG9A-FAM134B interaction on the ER surface with high-definition structured illumination microscopy (SIM) (Figure 1F).

The C terminus of FAM134B has intrinsic disordered features, which make it a good candidate for protein-protein interactions. To dissect the mechanism that allows FAM134B to engage ATG9A, we generated truncated versions of FAM134B missing the large C-terminal region (aa235–495), as well as the predicted “lumen loop” (aa141–207). The results show that, indeed, truncation of the entire C-tail greatly reduced the interaction with ATG9A, whereas deletion of the aa141–207 loop did not (Figures 2A and 2B). Iterative Threading ASSEmbly Refinement (i-TASSER) modeling of aa235–495 domain predicts a long arm with a globular end (aa310–495) uniquely positioned to engage in protein-protein interactions (Figure 2C; the arm is shown in blue). This globular domain is predicted to assemble as three very similar sub-domains, each assembled as two opposing α-helices connected by a coiled region (aa310–380, aa380–440, and aa440–495; see Figure 2D). Deletion of the arm (aa228–310) or sub-domains aa310–380 and aa380–440 increased, whereas deletion of the aa440–495 sub-domain reduced the FAM134B-ATG9A interaction (Figures 2E–2H). Interestingly, the LIR, which is necessary for FAM134B to recruit LC3B, is situated within the aa440–495 domain as a connecting loop between the two small α-helices of the sub-domain (Figure 2I; the LIR is in yellow). However, deletion of the LIR did not affect the ATG9A-FAM134B interaction, indicating that, although necessary for the engagement of LC3B, it is not required for interacting with ATG9A (Figure 2J).

Next, we decided to target the ATG9A-SEC62 interaction. The topology of SEC62 is shown in Figure 3A. As with FAM134B, SEC62 was also found to interact with the ER-based immature form of ATG9A following co-immunoprecipitation (Figure 3B; see inset) as well as SIM-mediated visualization (Figure 3C). Truncation of the cytosolic C terminus of SEC62 greatly reduced the ATG9A-SEC62 interaction (Figures 3D and 3E). The C terminus of SEC62 is highly disordered and displays several flexible areas; as a result, we were unable to generate reliable structural models. However, even with SEC62, we observed that truncation of two different segments of the C terminus elicited opposing results, with the aa260–289 deletion stimulating the ATG9A-SEC62 interaction and the aa291–395 deletion (which included the LIR) reducing the ATG9A-SEC62 interaction (Figures 3F–3I).

The aforementioned results suggest that FAM134B and SEC62 behave similarly in multiple aspects. First, they both engage ATG9A at the ER with their cytosolic C terminus. Second, they both engage ATG9A with the amino acid region that encompasses the LIR. Finally, in both cases, the interaction appears to require specific structural features rather than a simple amino acid sequence, thus suggesting that intrinsic dynamic events within the C terminus of FAM134B and SEC62 are necessary for functional engagement of ATG9A.

Engagement of ATG9A by FAM134B and SEC62 is accompanied by induction of reticulophagy

To determine the functional relevance of the above-mentioned findings, we generated HEK293 cells with stable expression of the ER autophagy tandem reporter (EATR) (Liang et al., 2018). The probe includes a
tandem mCherry-eGFP reporter fused to RAMP4 (Figure S1A). As RAMP4 is an integral ER-membrane resident protein, EATR detects induction of reticulophagy (Liang et al., 2018). We transfected the aforementioned HEK293EATR cells with two opposing mutants of FAM134B and SEC62. The results show that deletion mutants that displayed reduced (Δ235–495 for FAM134B and Δ291–395 for SEC62) engagement of ATG9A were unable to stimulate reticulophagy, whereas deletion mutants that increased (Δ310–380 for FAM134B and Δ260–289 for SEC62) the engagement of ATG9A displayed increased induction of reticulophagy (Figures 4A and 4B).

I-TASSER modeling of FAM134B predicts two opposing interacting Ca++-binding helix-loop-helix EF hands (Gifford et al., 2007; Kawasaki and Kretsinger, 2017). One of them (EF2) is situated within the aa440–495 loop that is required for binding to ATG9A, whereas the other (EF1) is situated on the opposing arm (Figures 4C and 4D; see also Figures 2C and 2D). The opposing and interacting arrangement of the two EF hands would suggest that intrinsic Ca++ dynamics can influence engagement of LC3B and induction of

Figure 2. The globular C terminus of FAM134B is required for ATG9A interaction
(A and B) Deletion of aa235–495 of FAM134B prevented functional interaction with ATG9A. Representative western blot is shown in (A), whereas quantitative changes are shown in (B). Data are represented as mean ± SD. *p < 0.0005.
(C and D) I-TASSER model of the cytosolic C terminus of FAM134B. The entire C-terminus (C) and the globular domain (D) are shown. Different colors highlight the domains targeted in follow-up experiments.
(E–H) Deletion of different domains within the cytosolic C terminus of FAM134B affected the ATG9A-FAM134B interaction differently. Representative western blots are shown in (E and G), whereas quantitative changes are shown in (F and H). Data are represented as mean ± SD. **p < 0.005; #p < 0.0005.
(I) I-TASSER model of the aa440–495 region of FAM134B with the LIR domain (shown in yellow).
(J) Deletion of the LIR domain of FAM134B did not prevent functional interaction with ATG9A.
reticulophagy. To test whether this was indeed the case, we mutated the Ca++-stabilizing residues within each EF hand to Ala, specifically E270, D272, and E274 within EF1 and D464 and E467 within EF2 (Figures 4C and 4D). When transfected into HEK293EATR cells, the EF1 mutant increased, whereas the EF2 mutant decreased the induction of reticulophagy (Figures 4E and 4F). Importantly, the EF1 mutant version of FAM134B displayed a sustained reticulophagy flux that appeared independent of starvation (Figure 4F). In contrast, the EF2 mutant version of FAM134B was unable to respond at all to starvation (Figure 4F). Therefore, it is plausible to assume that Ca++ dynamics between EF1 and EF2 are key for the progression of reticulophagy, perhaps through an “opening” mechanism that allows the aa440–495 loop to engage LC3B (see Figure 4G).

Sequence alignment suggests the presence of two EF hands on the C terminus of SEC62 that follow the same pattern of FAM134B (Figures 4C and 4D). However, secondary structure prediction as well as I-TASSER failed to support this scenario. The inconsistency might be because the C terminus of SEC62 is highly dynamic. Indeed, previous studies have confirmed the Ca++-binding properties of one of the EF hands (labeled as EF1 in Figure 4C) (Linxweiler et al., 2013). Therefore, we decided to mutate predicted...
Figure 4. FAM134B and SEC62 mutants that engage ATG9A can induce reticulophagy

(A and B) Reticulophagy flux induced by different FAM134B (A) or SEC62 (B) deletion mutants. Data are represented as mean ± SD. *p < 0.005.

(C and D) Sequence alignment of presumed EF hands in FAM134B and SEC62. Both EF1 (C) and EF2 (D) are shown. The Ca++-stabilizing amino acids that were mutated in the following experiments are indicated by arrowheads. The LIR domain is highlighted.

(E and F) EF1 and EF2 of FAM134B differentially affect reticulophagy flux. (E) Flux at 2hr and 4hr. (F) Time-course. Data are represented as mean ± SD. *p < 0.05; **p < 0.005; #p < 0.0005.

(G) I-TASSER model of the opposing EF1 and EF2 of FAM134B showing predicted interaction.

(H and I) EF1 and EF2 of SEC62 have a modest effect on the induction of reticulophagy. (H) Flux at 2hr and 4hr. (I) Time-course. Data are represented as mean ± SD. *p < 0.0005.
Ca**-stabilizing residues within each EF hand of SEC62 (Figures 4C and 4D) and test their functional role. In contrast to FAM134B, only the EF2 mutant version of SEC62 was able to significantly affect the induction of reticulophagy upon starvation (Figure 4H). However, this effect was not as dramatic and immediate as with FAM134B (Figure 4I). Therefore, the biological significance of the EF structure and Ca** dynamics with SEC62 remains unclear.

To validate the results obtained with the EATR system, we also used live imaging to visualize mCherry puncta as an indication of successful transition from the ER membrane to the lysosomes. The results were consistent with EATR for both FAM134B and SEC62 (Figures S1B and S1C). Indeed, we observed increased puncta with the Δ310–380 (for FAM134B) and Δ260–289 (for SEC62) deletion mutants, as well as with the EF1 mutants (Figures S1B and S1C). Next, we generated HEK293 cells with stable expression of an mCherry-RAMP4 fusion system (referred to as mCherry cleavage from ER [CCER]) (Liang et al., 2018). In this system, the mCherry is cleaved from RAMP4 during progression of reticulophagy, thus causing a migration shift on immunoblotting (Figure S1D) (Liang et al., 2018). Again, we observed concordance with EATR- and imaging-based approaches (Figures S1E and S1F).

When taken together, the array of modeling, structural biochemistry, and targeted mutagenesis indicates that engagement of ATG9A by the C terminus of FAM134B and SEC62 is accompanied by the induction of reticulophagy. Furthermore, at least with FAM134B, this process appears to depend on Ca** dynamics that might influence the engagement of LC3B. The possible regulatory functions of Ca** are supported by the fact that calcium-modulating agents, thapsigargin and BAPTA, were able to influence the induction of reticulophagy in vivo (Figure S2).

**CALR and HSPB1 are ATG9A partners**

Collectively, these results reveal previously unidentified mechanistic aspects of the ATG9A-FAM134B and ATG9A-SEC62 interaction. However, they do not resolve the topological conundrum that we discussed earlier. Indeed, the acetylation of ATG9A occurs on the luminal side of the ER, whereas the engagement of FAM134B or SEC62 occurs on the cytosolic side. A possible scenario is that additional ATG9A-interacting proteins might be required to connect a luminal (acetylation) to a cytosolic (FAM134B or SEC62 engagement) event. To address this, we overexpressed wild-type (WT) ATG9A with a myc tag at the C terminus in HEK293 cells and generated stable clones. Next, we purified ATG9A by affinity chromatography following cell permeabilization with either Triton X-100 or DDM, and analyzed the migration profile of ATG9A under native conditions. Depending on the different detergent used, we resolved two large complexes of approximately 720 and 900 kDa (Figure 5A) suggesting that ATG9A exists within a large supercomplex. Importantly, similar high-molecular-mass species have already been described in yeast, indicating that this is an intrinsic aspect of ATG9A biology (He et al., 2008).

Next, we prepared ER-enriched fractions and purified ATG9A-interacting proteins following Triton X-100 or DDM permeabilization and affinity chromatography. This ATG9A supercomplex was then resolved by mass spectrometry (MS). MS identified several proteins, which were broadly classified as cytosolic and ER associated based on our own interest in ER-specific events. The 35 ER-associated proteins included ER luminal proteins, integral ER membrane proteins, and proteins that can associate with the cytosolic side of the ER membrane (Figure 5B and Table S1). KEGG pathway and STRING analysis broadly defined three groups of proteins. (1) Folding/chaperones, which included a subset of 16 proteins responsible for folding of nascent polypeptides. Of note are PDIA3 and PDIA6, two members of the large protein disulfide isomerase family, which recognize free sulfhydryl groups and catalyze the formation of disulfide bonds of nascent glycoproteins. Also of note are UDP-glucose-glycoprotein glucosyltransferase (UGGT1), which is responsible for reglucosylation of unfolded glycoproteins as part of the calreticulin/calnexin cycle, and calreticulin (CALR) and calnexin (CANX) themselves, which are Ca**-binding chaperones responsible for quality control of nascent glycoproteins. (2) ERES/COP, which included nine proteins involved in the assembly of ER exit sites (ERES) as well as formation and trafficking of vesicles between the ER and the Golgi apparatus. Of note are SEC22B, SEC23B, SEC24C, and SEC31A, which are essential for the organization of the ERES and the packaging of correctly folded polypeptides for transport to the Golgi apparatus. Also of note are COPA, COPB1, COPB2, COPG1, and COPE, all members of the large coatamer superfamily. They are cytosolic proteins that are recruited on the surface of the ER and Golgi apparatus to mediate anterograde and retrograde ER-Golgi transport of cargo proteins. (3) Cytosolic chaperones, which included a group of six cytosolic members of the heat shock protein superfamily that can be recruited on the ER membrane as part of their chaperone-like activity.
Figure 5. Identification and characterization of the ER-based ATG9A interactome
(A) Migration profile of the ATG9A supercomplex.
(B) ER-based ATG9A-interacting proteins are shown in clusters based on KEGG and STRING classification.
(C) Endogenous Hspb1 and Calr preferentially associate with Atg9a in AT-1 sTg mice. Co-immunoprecipitation was performed with intact native ER vesicles.
(D) Schematic view of the experimental conditions used in (C) and (E–G). Atg9a is shown in blue.
To support the MS findings, we isolated ER membranes, immunoprecipitated ER-bound endogenous Atg9a, and probed for all target proteins identified in Figure 5B. We were able to confirm in vivo association with 32 of the proteins identified by MS (see Figures 5C and S3A). Only exceptions were a subset of three proteins (Sec24c, Sec31a, and Cdc42), which could not be detected in a consistent fashion. As mentioned above, we launched this MS study to identify possible ATG9A-interacting proteins that specifically regulate the ability of ATG9A to engage FAM134B and SEC62 as a function of ATG9A acetylation. Therefore, we took advantage of AT-1S113R/+ and AT-1 sTg mice, which display reduced and increased acetylation of Atg9a, respectively (Peng et al., 2014, 2018). We reasoned that ATG9A-interacting proteins that are relevant for our study should display different association with ATG9A as a function of ATG9A’s own acetylation status. Among the 35 ER-associated proteins that were identified in the ATG9A interactome, only six displayed preferential association with endogenous Atg9a in one of our models: heat shock protein beta-1 (Hspb1; also referred to as Hsp27), Calr, protein disulfide-isomerase A3 (Pdia3), peptidyl-prolyl cis-trans isomerase B (Ppib), coatomer subunit alpha (Copa), and coatomer subunit gamma 1 (Copg1). Specifically, Hspb1, Calr, Ppib, and Copg1 appeared to preferentially associate with Atg9a in AT-1 sTg mice, which exhibit hyperacetylated Atg9a and reduced reticulophagy (Dieterich et al., 2019; Peng et al., 2018), whereas Copa appeared to preferentially associate with Atg9a in AT-1S113R/+ mice, which exhibit hypoacetylation of Atg9a and increased reticulophagy (Peng et al., 2014, 2016) (Figures 5C and S3A). This initial co-immunoprecipitation experiment was done by pulling down endogenous Atg9a from intact native ER membranes (Figure 5D). A limitation of this approach is that the ER vesicles are intact and, therefore, can include proteins that incorporate or associate with the vesicles but are not specifically bound to Atg9a (Figure 5D). To resolve this limitation, we permeabilized ER vesicles with 0.5% Triton X-100 before immunoprecipitation of endogenous Atg9a in one of our models: heat shock protein beta-1 (Hspb1; also referred to as Hsp27), Calr, protein disulfide-isomerase A3 (Pdia3), peptidyl-prolyl cis-trans isomerase B (Ppib), coatomer subunit alpha (Copa), and coatomer subunit gamma 1 (Copg1). Specifically, Hspb1, Calr, Ppib, and Copg1 appeared to preferentially associate with Atg9a in AT-1 sTg mice, which exhibit hyperacetylated Atg9a and reduced reticulophagy (Dieterich et al., 2019; Peng et al., 2018), whereas Copa appeared to preferentially associate with Atg9a in AT-1S113R/+ mice, which exhibit hypoacetylation of Atg9a and increased reticulophagy (Peng et al., 2014, 2016) (Figures 5C and S3A). This initial co-immunoprecipitation experiment was done by pulling down endogenous Atg9a from intact native ER membranes (Figure 5D). A limitation of this approach is that the ER vesicles are intact and, therefore, can include proteins that incorporate or associate with the vesicles but are not specifically bound to Atg9a (Figure 5D). To resolve this limitation, we permeabilized ER vesicles with 0.5% Triton X-100 before immunoprecipitation of endogenous Atg9a (Figure 5D). We found a striking divergence between AT-1S113R/+ and AT-1 sTg mice with Hspb1 and Calr, but not with Pdia3, Ppib, Copa, or Copg1 (Figures 5E and S3B). Interestingly, both Hspb1 and Calr were found to preferentially associate with ER-bound Atg9a of AT-1 sTg mice (Figures 5E–5G). Finally, we co-transfected human ATG9A-GFP with either HSPB1-DDK or CALR-DDK in HEK293 cells. Co-immunoprecipitation experiments confirmed that both HSPB1 and CALR can interact with ATG9A (Figures 5H–5K). Importantly, immunoprecipitation of CALR, an ER luminal protein, only pulled down the immature and ER-based form of ATG9A (Figure 5J), whereas immunoprecipitation of HSPB1, a cytosolic protein, pulled down both the immature and mature versions of ATG9A (Figure 5H). These results provide further significance to our findings. Finally, high-definition SIM confirmed that Atg9a-Hspb1 (Figure 5L) and Atg9a-Calr (Figure 5M) interaction can occur at the ER.

Collectively, the aforementioned results identified HSPB1 and CALR as previously unidentified ATG9A-interacting proteins. They also revealed that both HSPB1 and CALR preferentially associate with acetylated ATG9A, which is unable to engage FAM134B or SEC62 and trigger reticulophagy (Peng et al., 2018). Our MS-based studies also revealed a surprising association of ER-bound ATG9A with a series of proteins that are involved with essential aspects of ER biology (discussed later).

**CALR and HSPB1 play different roles in the regulation of reticulophagy**

To dissect the role of CALR in ATG9A-mediated induction of reticulophagy, we overexpressed human CALR in HEK293 cells and then determined levels of acetylated ATG9A. Surprisingly, we found a marked increase in the levels of acetylated ATG9A (Figures 6A and 6B), suggesting that CALR can either acetylate ATG9A or stabilize already acetylated ATG9A. To differentiate between these two possibilities, we analyzed the acetylation profile of the ER following overexpression of CALR, but did not observe significant differences (Figure 6C). We also determined acetyltransferase activity of affinity-purified CALR, but were not able to detect significant acetyl-CoA:lysine acetyltransferase activity in vitro (data not shown).
Figure 6. CALR and HSPB1 differentially regulate the induction of reticulophagy

(A and B) Overexpression of CALR-stabilized acetylated ATG9A. Representative western blots are shown in (A), whereas quantitative changes are shown in (B). Data are represented as mean ± SD. *p < 0.05.

(C) Overexpression of CALR did not affect the acetylation profile of the ER.

(D) Overexpression of CALR did not affect the induction of reticulophagy. Data are represented as mean ± SD.

(E and F) siRNA-mediated downregulation of CALR stimulated the induction of reticulophagy. (E) Reticulophagy flux was measured at 2 and 4 h. (F) Western blot to document the downregulation of CALR. siNEG were used as negative siRNA controls. Data are represented as mean ± SD. #p < 0.0005.

(G–H) I-TASSER structure of dimeric (G) and monomeric (H) forms of HSPB1 with mutations and deletions targeted in (I–O).

(I) Overexpression of HSPB1 stimulated the induction of reticulophagy, whereas disease-associated mutants or downregulation of HSPB1 did not. Reticulophagy flux was measured at 2 and 4 h. Data are represented as mean ± SD. *p < 0.0005.

(J) Western blot to document the downregulation of HSPB1. siNEG were used as negative siRNA controls.
Next, we used the EATR system to determine whether CALR itself could influence the induction of reticulophagy. Interestingly, overexpression of CALR did not elicit any effect (Figure 6D), whereas small interfering RNA (siRNA)-mediated downregulation of CALR caused a significant increase in induction and progression of reticulophagy (Figures 6E and 6F).

Overall, the aforementioned results are consistent with a model where CALR recognizes—and perhaps stabilizes—already acetylated ATG9A, thus preventing the engagement of FAM134B and/or SEC62 by ATG9A (discussed later). The limiting element of this model would be the acetylation status of ATG9A. As a result, overexpression of CALR would not affect the induction of autophagy because the acetylated pool of endogenous ATG9A is already “sequestered” by endogenous CALR. However, downregulation of endogenous CALR would prevent “sequestration” of ATG9A, thus favoring engagement of FAM134B and/or SEC62 and induction/progression of reticulophagy.

To dissect the role of HSPB1, we took advantage of the already reported disease association with axonal Charcot-Marie-Tooth type 2F (CMT2F) and distal hereditary motor neuropathy 2B (HMN2B). Specifically, we targeted the R136W single mutation and the R140G/K141Q double mutation, as well as the N terminus (Almeida-Souza et al., 2010; Echaniz-Laguna et al., 2017; Ikeda et al., 2009; Muranova et al., 2015; Nefedova et al., 2013a, 2013b). It is worth mentioning that HSPB1 exists in equilibrium between different assembly states: monomers, dimers, and oligomers (Alderson et al., 2019; Arrigo, 2017). The R136W and R140G/K141Q mutations preclude the dimerization of HSPB1, whereas deletion of the N terminus prevents the formation of HSPB1 oligomers (Alderson et al., 2019; Arrigo, 2017). Therefore, the mutations and deletion that we chose provide biological as well as disease-relevant information (see Figures 6G and 6H).

Expression of WT HSPB1 stimulated the induction and progression of reticulophagy as assessed by EATR. However, all the point mutations and terminal deletions that we used in this study appeared to be defective and unable to stimulate EATR flux at the same level of the WT protein (Figure 6I). Interestingly, siRNA-mediated downregulation of HSPB1 did not elicit any effect (Figures 6I and 6J), suggesting that although WT HSPB1 can stimulate the induction of reticulophagy, it is not essential for its normal progression.

When taken together, the aforementioned results suggest that CALR and HSPB1 are ATG9A-interacting proteins; they play opposing roles and act on the opposing sides of the ER membrane to regulate the ATG9A-mediated induction of reticulophagy.

DISCUSSION

The ER acetylation machinery includes three essential components: an ER membrane transporter, AT-1, which translocates acetyl-CoA from the cytosol to the ER lumen, and two ER-bound acetyl-CoA:lysine acetyltransferases, ATase1 and ATase2, which acetylate ER-resident and -transiting proteins (Farrugia and Puglielli, 2018; Jonas et al., 2010; Ko and Puglielli, 2009). Loss-of-function mutations and gene duplication events that disrupt ER acetylation are associated with severe human diseases (Chiplunkar et al., 2016; Huppke et al., 2012a; Lin et al., 2008; Prasad et al., 2012; Sanders et al., 2011). Furthermore, mouse models of reduced or increased AT-1 activity develop severe phenotypes that mimic the associated human diseases (Hullinger et al., 2016; Peng et al., 2014, 2018). A combination of cell- and animal-based studies indicate that the ER acetylation machinery is necessary to maintain proteostasis within the ER and secretory pathway (Farrugia and Puglielli, 2018; Pehar et al., 2012; Peng et al., 2014, 2016, 2018).

The removal of toxic protein aggregates through reticulophagy is an essential aspect of the proteostatic functions of ER acetylation. Reduced ER acetylation in AT-1S113R/+ hypomorphic mice causes increased (aberrant) induction of reticulophagy, whereas increased ER acetylation in AT-1-overexpressing mice has the opposite effect (Hullinger et al., 2016; Peng et al., 2014, 2018). Furthermore, inhibition of ATase1 and ATase2, downstream of AT-1, can restore reticulophagy in AT-1 overexpressors (Peng et al., 2016, 2018). Mechanistically, the regulation of reticulophagy involves acetylation of ATG9A on two lysine residues (K359 and K363) that face the lumen of the ER (Pehar et al., 2012). The acetylation status of ATG9A, in turn, regulates the ability of ATG9A to interact with FAM134B and SEC62, which can then engage cytosolic LC3B through the LIR on their C terminus (Peng et al., 2018).

Here, we report mechanistic aspects of the ATG9A-FAM134B and ATG9A-SEC62 interactions. Both FAM134B and SEC62 engage ATG9A on the cytosolic side of the ER through specific structural features.
that appear to involve changes in Ca\(^{++}\) dynamics. The modeling data obtained with FAM134B are particularly intriguing, as they support a scenario where interaction with ATG9A and changes in Ca\(^{++}\) binding through two opposing EF hands might allow structural rearrangement of the globular end of FAM134B that is necessary for the LIR to “open” and engage LC3B. The typical helix-loop-helix structure of the EF hand is stabilized by Ca\(^{++}\), which is engaged by the side chains of Glu and Asp residues. This provides geometric and spatial constraint to the otherwise very flexible turn-loop structure (Gifford et al., 2007; Kawasaki and Kretsinger, 2017). EF hands tend to occur in pairs, and cooperativity is often ensured by their opposing placement on the folded protein that generates a defined 11 Å distance between bound Ca\(^{++}\) ions (Biekofsky and Feeney, 1998; Biekofsky et al., 1998). However, the opposing EF hands tend to be slightly different. This difference causes asymmetric Ca\(^{++}\) binding, which ultimately ensures dynamic fluctuation of specific regions within a folded polypeptide (Kawasaki and Kretsinger, 2017; Lopez et al., 2002; Shaw et al., 1991). Consistent with the above, inactivation of the opposing EF1 and EF2 hands on the globular domain of FAM134B elicited opposing effects. Inactivation of EF1 increased the ability of FAM134B to induce reticulophagy, whereas inactivation of EF2 decreased the ability of FAM134B to induce reticulophagy, suggesting that structural fluctuations are indeed necessary for FAM134B to engage LC3B. This is somewhat reminiscent of the assembly-disassembly fluctuations described with circular dichroism and NMR studies of both typical and atypical EF hands (Gifford et al., 2007; Kawasaki and Kretsinger, 2017). The fact that selective deletions of opposing domains within the globular end of FAM134B positively and negatively influenced ATG9A-FAM134B interaction provides further structural support to the model delineated above.

A challenging aspect of our findings is that the acetylation of ATG9A occurs in the lumen of the ER, whereas the engagement of FAM134B or SEC62 occurs on the cytosolic side of the organelle. To resolve this, we used unbiased MS to resolve a crude “ATG9A interactome.” The analysis identified several proteins broadly divided into “cytosolic” and “ER-associated.” The former included cytoskeletal and cytoskeletal-associated proteins. The latter included ER-integral as well as ER-associated proteins. It is worth stressing that, although translated in the ER, ATG9A normally translocates out of the ER toward the Golgi apparatus, the plasma membrane, and LC3B-positive autophagosomes during the induction of autophagy (Bejarano et al., 2014; Ohashi and Munro, 2010; Pun et al., 2013; Saitoh et al., 2009; Tamura et al., 2010; Young et al., 2006). Also important is the fact that ATG9A can interact with ERES and COP structures (Graef et al., 2013; Kakuta et al., 2012; Lynch-Day et al., 2010; Suzuki et al., 2013), and that cross talk between the autophagy machinery and the ERES/COP assembly machinery has been described (Sanchez-Wandelmer et al., 2015). Therefore, the ATG9A interactome that we report here appears to reflect essential aspects of ATG9A biology. It is also worth mentioning that both FAM134B and SEC62 are found on the rough ER, where the bulk of protein biosynthesis occurs (Fumagalli et al., 2016; Khaminets et al., 2015; Mochida et al., 2015; Rubinstein, 2015; Schuck, 2016). Furthermore, SEC62 is part of the translocon, which allows insertion of newly synthesized proteins within the ER (Gemmer and Forster, 2020). It has been proposed that FAM134B and SEC62 couple the machinery that allows biosynthesis and insertion of newly synthesized proteins into the ER with the machinery that controls the disposal of unfolded/misfolded polypeptides, and maintains the size of the ER (Nakatogawa and Mochida, 2015; Schuck, 2016). The presence of integral components of the ER quality control machinery as well as ERES and COP II structural components within the ATG9A interactome seems to support the idea of vicinity and cross talk of the essential machineries that maintain proteostasis within the ER.

By using mouse models of decreased (AT-1^5113R/+), and increased (AT-1 sTg) AT-1 activity, we were able to identify two ATG9A-interacting proteins, CALR and HSPB1, that preferentially bind acetylated ATG9A. Importantly, CALR is a non-classical chaperone involved in the quality control of newly synthesized glycoproteins (Hebert and Molinari, 2007; Trombetta and Parodi, 2003); being an ER luminal protein, it can only engage ATG9A within the lumen of the organelle. Our findings suggest that CALR can recognize—and perhaps stabilize—acetylated ATG9A; therefore, CALR is the first confirmed ER luminal ATG9A-interacting protein. Calnexin (CANX), the membrane-bound structural homolog of CALR, has recently emerged as a possible “facilitator” of ER-phagy/reticulophagy through its interaction with FAM134B (Fregno et al., 2018). Therefore the CANX/CALR system may collectively regulate the induction of ER-phagy/reticulophagy through FAM134B and ATG9A, respectively. Interestingly, in addition to CALR, the MS analysis of the ATG9A interactome also identified CANX, PDIA3, PDIA6, and UGGT1; these are all part of the classical CALR/CANX cycle that retrieves unfolded glycoproteins as part of the normal quality control activities of the ER. These findings further support the existence of a close interaction (and spatial vicinity) between the machinery that ensures quality control and the one that ensures reticulophagy within the ER.
In addition to CALR, the MS analysis of the ATG9A interactome identified the small heat shock protein HSPB1 as a cytosolic ATG9A-interacting protein. HSPB1 mutations have been linked to axonal forms of CMT disease and HMN (Almeida-Souza et al., 2010; Capponi et al., 2011; Echaniz-Laguna et al., 2017; Evgrafov et al., 2004; Holmgren et al., 2013; Ikeda et al., 2009; Irobi et al., 2010; Lin et al., 2011; Muranova et al., 2015; Nefedova et al., 2013a, 2013b), which belong to the general family of hereditary sensory autonomic and motor neuropathies (HSANs/HMNs). How these mutations affect HSPB1 activity is unknown, although studies suggest altered substrate-binding activity (Almeida-Souza et al., 2010; Capponi et al., 2011; Echaniz-Laguna et al., 2017; Evgrafov et al., 2004; Holmgren et al., 2013; Ikeda et al., 2009; Irobi et al., 2010; Lin et al., 2011; Muranova et al., 2015; Nefedova et al., 2013a, 2013b). However, the real substrate(s) is/are unknown. Our data indicate that CMT/HMN-associated mutant versions of HSPB1 inactivate the ability of HSPB1 to stimulate reticulophagy. Interestingly, AT-1 or FAM134B mutations are also associated with related forms of HSANs/HMNs (Almeida-Souza et al., 2010; Capponi et al., 2011; Davidson et al., 2012; Echaniz-Laguna et al., 2017; Evgrafov et al., 2004; Holmgren et al., 2013; Ikeda et al., 2009; Irobi et al., 2010; Lin et al., 2011; Muranova et al., 2015; Nefedova et al., 2013a, 2013b). When taken together, these data support the possibility that defects in induction and progression of reticulophagy might underlie disease-associated mutations affecting AT-1, FAM134B, and HSPB1.

Based on our findings, we can envision a working model where increased influx of acetyl-CoA into the ER lumen caused by increased expression and activity of AT-1 in AT-1 sTg and AT-1 nTg mice leads to hyper-acetylation of the ER pool of ATG9A, perhaps caused by allosteric activation of ATase1 (Rigby et al., 2020) (Figure 7A). Conversely, reduced influx of acetyl-CoA into the ER lumen caused by haploinsufficiency of AT-1 in AT-1 sTg and AT-1 nTg mice leads to hyper-acetylation of the ER pool of ATG9A (Figure 7B). When acetylated, ATG9A is recognized—and perhaps sequestered—by CALR, thus preventing interaction with FAM134B or SEC62, engagement of LC3B, and induction of reticulophagy (Figure 7A). Under this condition, the cytosolic-exposed domain(s) of ATG9A is occupied by HSPB1, probably in the monomeric form. In contrast, when non-acetylated, ATG9A is not recognized by CALR and is able to interact with FAM134B or SEC62. This might force displacement or rearrangement of HSPB1. The ATG9A-FAM134B and ATG9A-SEC62 interaction allows engagement of LC3B and induction of reticulophagy.
reticulophagy (Figure 7B). The structural arrangement of HSPB1 seems to be an important component, because disease-associated mutations that prevent oligomerization of HSPB1 appear to act as negative effectors. Although several aspects of this model are supported by the present study and by already published work (Hullinger et al., 2016; Pehar et al., 2012; Peng et al., 2014, 2016, 2018; Peng and Puglielli, 2016; Rigby et al., 2020), there are different components that remain to be fully dissected and, as such, we consider this only a working and incomplete model.

The specific role of K359/K359Ac and K363/K363Ac and how they influence ATG9A activity remains an unresolved aspect of our findings. All ATG9 proteins, including human ATG9A, are predicted to contain six transmembrane (TM) helices with three loops facing the luminal side. While this manuscript was under preparation and editorial process, four different ATG9 cryoelectron microscopic (cryo-EM) structures were reported (Guardia et al., 2020; Lai et al., 2020; Maeda et al., 2020; Matoba et al., 2020). Three of them (from yeast and human ATG9A) yielded a model with four spanning TM helices and two reentrant helices (Guardia et al., 2020; Maeda et al., 2020; Matoba et al., 2020), whereas one (from Arabidopsis thaliana) yielded a model with six spanning TM helices (Lai et al., 2020). In all cases, ATG9 appeared to organize as a trimer with a central open pore. ATG9A is acetylated on two lysine residues, K359 and K363, both residing in the second of the three intraluminal loop within the six TM predicted model (Pehar et al., 2012). The ER-based ATases, which acetylate ATG9A, are both type-II membrane proteins with the catalytic domain facing the ER lumen (Farrugia and Puglielli, 2018; Ko and Puglielli, 2009). Therefore, the predicted intraluminal location of K359 and K363 is consistent with the topology of the ATases. The six TM helices cryo-EM model of A. thaliana ATG9 would be consistent with the intraluminal K359/K363 acetylation model. In contrast, the four fully spanning and two reentrant helices cryo-EM model of human ATG9A would place K359 and K363 in the central pore of the trimeric complex. Although the central pore of the trimeric structure appears to be necessary for the autophagy-related functions of ATG9A, the dynamics of the trimeric assembly itself are currently unknown (Maeda et al., 2020; Matoba et al., 2020). Specifically, it is unknown when and how the ATG9A monomers interact to build the trimeric structure. As the acetylation status of ATG9A regulates its autophagy-related functions (Farrugia and Puglielli, 2018; Pehar et al., 2012; Peng et al., 2014, 2016, 2018), we can speculate that the acetylation/deacetylation of K359/K363 might play, either directly or indirectly, some role in the trimerization process of ATG9A or in the proposed scramblase/flipase activity of the open pore. However, this remains to be studied. The in vivo implication of the structural models also remains to be fully studied. However, ongoing studies with Atase1−/− and Atase2−/− mice in our laboratory might reveal additional mechanistic aspects related to ATG9A assembly and, perhaps, further clarify pathogenic events underlying the disease phenotypes caused by dysfunctional ER acetylation.

Limitations of the study

1. The fact that ATG9A might work at the intersection between engagement of the secretory pathway and induction of reticulophagy is an intriguing new area of investigation. Two independent studies (one recently published [Dietenich et al., 2021] and one in press) from our laboratory using mouse models of increased (AT-1 sTg) and reduced (AT-1S113R/+ , Atase1−/−/C0 and Atase2−/−/C0) ER acetylation confirm this dual function of the ER acetylation machinery. However, how this balance is achieved remains to be dissected.

2. This study revealed a previously unknown regulatory function of Ca++. Ca++ regulates the ATG9A-FAM134B and ATG9A-SEC62 interaction on the cytosolic side of the ER. Ca++ also regulates CALR activity within the ER lumen. Importantly, CALR itself normally functions as a high-affinity “storage system” for ER luminal Ca++. The precise mechanism(s) underlying the role of Ca++ release and—perhaps—transport across the ER membrane remains to be dissected.

3. The assembly state of HSPB1 is crucial to the activity of the protein and seems to be important for the pathophysiology of HSPB1-associated HSANs/HMNs. However, why and how remains to be determined.

Resource availability

Lead contact
Further information, requests, and enquiries should be directed to and will be fulfilled by the lead contact and corresponding author, Prof. Luigi Puglielli (lp1@medicine.wisc.edu).

Materials availability
All unique/stable reagents generated in this study are available on request from the lead contact.
Data and code availability
The published article includes all data generated or analyzed during this study.

METHODS
All methods can be found in the accompanying Transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102315.

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AUTHOR CONTRIBUTIONS
B.K.S., N.S.O., Y.P., and S.L.S. performed the experiments and analyzed the data. L.P. designed the overall study and wrote the paper with input from all authors.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental information

ATG9A regulates proteostasis through reticulophagy
receptors FAM134B and SEC62 and folding
chaperones CALR and HSPB1

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Figure S1. FAM134B and SEC62 mutants that engage ATG9A can induce reticulophagy, Related to Figure 4.

(A) Schematic view of the EATR assay design.

(B and C) Live imaging showing induction of reticulophagy as caused by different FAM134B (B) or SEC62 (C) mutants.

(D) Schematic view of the CCER assay design.

(E and F) Western blotting showing cleavage of mCherry as caused by different FAM134B (E) or SEC62 (F) mutants.
Figure S2. Calcium modifying drugs, Thapsigargin (Tg) and BAPTA (Bt), differentially affected the induction of reticulophagy, Related to Figure 4.

Reticulophagy flux as regulated by Tg and BAPTA. Data are represented as mean ± SD. #p < 0.0005.
Figure S3. Identification and characterization of the ER-based ATG9A interactome, 
Related to Figure 5.

(A) Western blotting showing that Atg9a interacting proteins co-immunoprecipitate with ER-based endogenous Atg9a. Co-immunoprecipitation was performed with intact native ER vesicles.

(B) Functional co-immunoprecipitation of selected proteins from (A; see arrows) was also performed with permeabilized native ER vesicles.
Figure S4. Uncropped blots included in the main manuscript, Related to Figures 1-2.
Figure S5. Uncropped blots included in the main manuscript, Related to Figure 3.
Figure S6. Uncropped blots included in the main manuscript, Related to Figure 5.
Figure S7. Uncropped blots included in the main manuscript, Related to Figure 6.
TRANSPARENT METHODS

Animals

All mouse studies were carried out in accordance with the NIH Guide for the Care and Use of Laboratory Animals, and received ethical approval by the Institutional Animal Care and Use Committee of the University of Wisconsin-Madison. Generation of AT-1S113R/+ and AT-1sTg mice is described elsewhere (Peng et al., 2014; Peng et al., 2018). Mice were studied at ~2-3 months of age. Age-matched wild-type (WT) littermates were used as controls. Both males and females were used for the experiments reported in this study.

Cell cultures

H4 (human neuroglioma; RRID:CVCL_1239; ATCC:HTB-148) and HEK293 (human embryonic kidney; RRID: CVCL_0045; ATCC:CRL-1573) cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Corning #10–013-CV) supplemented with 10% Fetal Bovine Serum (Corning #35–011-CV) and 1% penicillin/streptomycin (Gibco™ #10378016), thereafter referred to as DMEM-10. Cells were maintained at 37°C in a humidified atmosphere with 5% CO2. Cells were obtained and authenticated by American Type Culture Collection, and are not listed as a commonly misidentified cell line by the International Cell Line Authentication Committee (ICLAC; Version 9). Unless otherwise specified, cells were harvested by scraping in ice-cold PBS and pelleting by centrifugation at 6,000g for 5 minutes at 4°C.

Structural modeling

Structural modeling was performed using the Iterative Threading ASSEmbly Refinement (I-TASSER) platform, which analyzes PDB structural templates to select high-scoring 3D
models (Roy et al., 2010; Yang et al., 2015). PDB analysis employed the following threading platforms: CEthreader, FFAS3D, HHpred, HHsearch, MUSTER, Neff-MUSTER, PPAS, PRC, PROSPECT2, SP3, and SparksX. Models were further validated by combining secondary structure and solvent accessibility analysis. The top five scoring templates were built using MODELLER; they were then assigned a confidence score based on the significance of threading template alignments and the convergence parameters of the structure assembly simulations. The structure that gave the highest confidence score was used for our analysis. In the case of FAM134B, the following top structure templates were identified from the PDB library by LOMETS: 6gmhM; 2pff; 6fltX; 2pff; 5gaoE; 5yfpD; 5yz0C; 5ly9A; 4jioA. In the case of HSPB1, the following top structure templates were identified from the PDB library by LOMETS: 6dv5B; 6dv5A; 4ydzA; 6dv5; 2ygd; 6t1rA; 6dv5; 6dv5A; 6dv5B; 2bolA. The dimeric structure of the α-crystallin domain of HSPB1 (PDB:2n3j) was also used to generate our final HSPB1 model (Rajagopal et al., 2015). Secondary structure prediction was performed with the Jpred 4, PredictProtein, and NetSurfP-2.0 platforms. Ligand-protein binding interactions were assessed through the BioLip database. Disorder prediction was performed with DisEMBL, PredictProtein and NetSurfP-2.0 platforms. Images were prepared with Pymol (Schrödinger, Inc).

Cell transfection

H4 and HEK293 cells were transiently transfected with Lipofectamine 3000 (Invitrogen #100022234) according to the manufacturer's instructions. Original plasmids used: human pCMV6-ATG9A-tGFP (OriGene #RG222568); human pCMV6-ATG9A-DDK (OriGene #RC222513); human pCMV6-CALR-DDK (OriGene #RC203222); human pCMV6-HSPB1-
DDK (OriGene #RC201800); human pCMV6-FAM134B-DDK (OriGene #RC208619); human pCMV6-SEC62-DDK (OriGene #RC204452); pCMV6-Entry empty plasmid (OriGene #PS100001). Co-transfections were performed with 300fmol of DDK-tagged protein and 600fmol ATG9A-tGFP, unless otherwise noted. Cells were harvested 48 hours after transfection.

Site-directed mutagenesis

Plasmids were obtained from OriGene and are listed in the “Cell transfection” section. Molecular constructs were prepared by Integrated DNA Technologies. Primers are listed in Table S2. Site-directed mutagenesis was performed using the QuikChange Lightning Site-Directed Mutagenesis kit per manufacturer’s instructions (Agilent Technologies, 210518 and 210519). All constructs were prorogated from a single colony. Plasmid DNA was extracted using either QIAprep Spin Miniprep Kit or QIAGEN Plasmid Maxi Kit (Qiagen, 27106 and 12165). All constructs were confirmed by DNA sequencing.

Generation of EATR, CCER, and mCherry-ER-3 stable lines

To generate the EATR stable line, HEK293 cells were transfected with SspI digested TetOn-mCherry-eGFP-RAMP4 vector (gift from Jacob Corn; Addgene plasmid #109014) and selected in DMEM-10 with 400µg/mL G418 (Invitrogen #10131027). To generate the CCER stable line, HEK293 cells were transfected with MluI digested pLenti-X1-hygro-mCherry-RAMP4 vector (gift from Jacob Corn; Addgene plasmid #118391) and selected in DMEM-10 + 100µg/mL Hygromycin B (Invitrogen #10687010). To generate the mCherry-ER-3 stable line, HEK293 cells were transfected with undigested mCherry-ER-3 vector (gift from Michael Davidson; Addgene plasmid #55041) and selected in DMEM-10 with 400µg/mL G418.
Polyclonal resistant cells were expanded and cultured in G418 or Hygromycin B until plating for experiments.

**EATR assay**

HEK293 cells stably expressing TetOn-mCherry-eGFP-RAMP4 were plated on 12-well plates two days before transfection. On the day of transfection, media was changed to DMEM-10 with 400ug/mL G418 and 16ug/mL Doxycycline to enhance expression of the tandem reporter. Twenty-four hours before flow cytometry analysis, cells were transfected with 200fmol of plasmid using Lipofectamine 3000 according to manufacturer's instructions. To stimulate autophagy, cells were incubated for 2 or 4 hours in 50% DMEM-10 and 50% EBSS then trypsinized, centrifuged, and resuspended in PBS containing 2% FBS, 10mM EDTA, and 1ug/mL DAPI on ice. For time course experiments, autophagy was not stimulated and cells were trypsinized, centrifuged, and resuspended as above, but were placed in a 37°C water bath at time 0 and sampled every 10 minutes. Flow cytometry was performed on Attune NxT flow cytometers (Thermo Fisher Scientific) configured with 405, 488, 561, and 637nm lasers. Single, live, and mCherry-expressing gates were set using the entire experiment population. The “elevated flux” gate was defined as events one standard deviation above the least-squares regression line of the eGFP vs mCherry plot of non-transfected cells incubated in 100% DMEM-10 and 16ug/mL Doxycycline under equivalent experimental conditions. Data analysis was performed using Attune NxT Software version 4.2 and in-house Python scripts.

For calcium modulating experiments, cells were prepared equivalently to other EATR assay experiments except media was changed to 100% EBSS 2 hours before analysis instead of a 50% DMEM-10 and 50% EBSS solution to force induction of reticulophagy. Immediately after
media change, cells were treated with a 10x solution containing additives and either calcium modulating agent or DMSO. The final solution contained 1mM Probenecid (Sigma, P8761) and 0.05% Pluronic F-127 (Sigma, P2443) as well as either 1μM Thapsigargin (Sigma, T9033), 1μM BAPTA-AM (Thermo Fisher Scientific, B6769), or 0.1% DMSO (vehicle). Probenecid and Pluronic F-127 were added to improve loading of BAPTA-AM into cells.

**EATR Live Imaging**

HEK293 cells stably expressing mCherry-eGFP-RAMP4 were transfected as for the EATR assay. Cells were trypsinized and replated 24 hours post-transfection on a poly-D-lysine-coated, glass-bottomed, 96-well plate (Cellvis P96-1.5H-N) and incubated for 48hrs to allow for maximum construct expression. Single 1024x1024, 0.208um/pixel images were acquired on a Nikon A1R-SI microscope equipped with 405, 488, 561, and 640 nm lasers and a CFI Plan Apochromat Lambda 60X Oil 1.4 NA objective at a pinhole size of 219.67um. The acquisition software was NIS-Elements AR 5.11.01.

**CCER Assay**

HEK293 cells stably expressing mCherry-RAMP4 were plated on 12-well plates two days before transfection. On the day of transfection, media was changed to DMEM-10 and cells were transfected with 200fmol of the indicated plasmids using Lipofectamine 3000 according to manufacturer's instructions. After 24 hours, cells were harvested via trypsin.
HEK293 cells stably expressing mCherry-ER-3 were plated on poly-D-lysine (Sigma P6407) coated glass coverslips and allowed to adhere overnight. Cells were then starved with EBSS for 6hrs to stimulate reticulophagy and fixed in 4% PFA for 15min. Cells were washed in PBS for 5min and permeabilized in 0.5% Saponin (Sigma 47036) in PBS for 20min. Saponin was used to preserve ER membrane structure. Fixed and permeabilized cells were then washed for 5min in PBS to remove excess detergent. Antibody staining was done sequentially to avoid cross-reaction, starting with ATG9A then FAM134B, SEC62, CALR, or HSPB1. Cells were blocked for 1hr in 5% goat serum and 0.1% Saponin in PBS, then incubated in primary antibody in 5% goat serum and 0.1% Saponin in PBS overnight at 4°C. Cells were washed 3x5min in PBS with 0.1% Tween-20, incubated in secondary antibody in 5% goat serum and 0.1% Saponin in PBS for 1 hour, and again washed 3x5min in PBS with 0.1% Tween-20. After sequential stainings, cells were washed one more time in PBS to remove excess detergent and mounted onto slides using ProLong™ Diamond Antifade Mountant (Invitrogen P36961).

**SIM and Image Reconstruction**

Z-stack 1024x1024, 0.032um/pixel images were acquired at 0.2um intervals on a Nikon-Structured Illumination Microscope (Eclipse Ti-E) equipped with 405, 488, 561, and 647 nm lasers, a CFI Apochromat TIRF 100XC Oil 1.49 NA objective, and iXon 3 EMCCD (Andor Technologies). The acquisition software was NIS-Elements AR 4.60.00. Reconstructions were generated using Imaris 9.6.0.
**ER enrichment**

Total ER membranes were isolated from 8-week-old WT, AT-1$^{S113R/+}$, and AT-1 sTg mice using the ER Enrichment Kit (Novus Biologicals, NBP2-29482) according to the manufacturer’s instructions.

**LC-MS/MS**

**ATG9A complex isolation.** Enriched ER fractions prepared as described above were extracted in either 1% Triton X-100 or 1% DDM. The ATG9A-myc complex was then purified by affinity chromatography using AminoLink immobilized anti-c-Myc monoclonal antibodies (Pierce ProFound, VWR-PI23620) (Jonas et al., 2010; Ko and Puglielli, 2009). Eluted ATG9A-associated proteins were analyzed by high-resolution high-accuracy MS.

**Enzymatic “In Liquid” Digestion.** TCA/acetone precipitated protein pellets were resolubilized and denatured in 7.5μl of 8M Urea / 50mM NH$_4$HCO$_3$ (pH8.5) / 1mM TrisHCl. Subsequently diluted to 30μl for reduction step with: 1.25μl of 25mM DTT, 2.5μl MeOH, and 18.75μl 25mM NH$_4$HCO$_3$ (pH8.5). Incubated at 52°C for 15 minutes, cooled on ice to room temperature then 1.5μl of 55mM IAA was added for alkylation and incubated in darkness at room temperature for 15 minutes. Reaction was quenched by adding 4μl of 25mM DTT. Subsequently 1μl of Trypsin/LysC solution [100ng/μl of 1:1 Trypsin (Promega) and LysC (FujiFilm) mix in 25mM NH$_4$HCO$_3$] and 13.5μl of 25mM NH$_4$HCO$_3$ (pH8.5) was added to 50μl final volume. Digestion was conducted for 2 hour at 42°C then additional 0.5μl of Trypsin/LysC solution was added and digestion proceeded for 4hrs at 37°C. Reaction was terminated by acidification with 2.5% TFA [Trifluoroacetic Acid] to 0.3% final.
NanoLC-MS/MS. Digests were cleaned up using OMIX C18 SPE cartridges (Agilent, Palo Alto, CA) per manufacturer protocol and eluted in 20µl of 60/40/0.1% ACN/H2O/TFA, dried to completion in the speed-vac and finally reconstituted in 12µl of 0.1% formic acid. Peptides were analyzed by nanoLC-MS/MS using the Agilent 1100 nanoflow system (Agilent) connected to hybrid linear ion trap-orbitrap mass spectrometer (LTQ-Orbitrap Elite™, Thermo Fisher Scientific) equipped with an EASY-Spray™ electrospray source. Chromatography of peptides prior to mass spectral analysis was accomplished using capillary emitter column (PepMap® C18, 3µM, 100Å, 150x0.075mm, Thermo Fisher Scientific) onto which 3µl of extracted peptides was automatically loaded. NanoHPLC system delivered solvents A: 0.1% (v/v) formic acid, and B: 99.9% (v/v) acetonitrile, 0.1% (v/v) formic acid at 0.50 µL/min to load the peptides (over a 30 minute period) and 0.3µl/min to elute peptides directly into the nano-electrospray with gradual gradient from 0% (v/v) B to 30% (v/v) B over 78 minutes and concluded with 5 minute fast gradient from 30% (v/v) B to 50% (v/v) B at which time a 5 minute flash-out from 50-95% (v/v) B took place. As peptides eluted from the HPLC-column/electrospray source survey MS scans were acquired in the Orbitrap with a resolution of 120,000 followed by MS2 fragmentation of 20 most intense peptides detected in the MS1 scan from 350 to 1800 m/z; redundancy was limited by dynamic exclusion.

MS data analysis. Raw MS/MS data were converted to mgf file format using MSConvert (ProteoWizard: Open Source Software for Rapid Proteomics Tools Development). Resulting mgf files were used to search against Uniprot Homo sapiens proteome database (UP000005640) with a decoy reverse entries and a list of common contaminants (134,183 total entries) using in-house Mascot search engine 2.2.07 [Matrix Science] with variable Methionine oxidation, Asparagine or Glutamine deamidation plus Lysine and protein N-terminus acetylation. Peptide mass tolerance
was set at 15 ppm and fragment mass at 0.6 Da. Protein annotations, significance of identification and spectral based quantification was done with Scaffold software (version 4.3.2, Proteome Software Inc., Portland, OR). Protein identifications were accepted if they could be established at greater than 97.0% probability to achieve an FDR less than 1.0% and contained at least 2 identified peptides. Protein probabilities were assigned by the Protein Prophet algorithm (Nesvizhskii, AI Anal Chem. 2003 Sep 1;75(17):4646-58). Proteins that contained similar peptides and could not be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony. MS parameters as well as the final output of the MS analysis is reported in Table S1. Identified proteins were further analyzed using the KEGG (Kanehisa, 2019; Kanehisa et al., 2020) and STRING (Szklarczyk et al., 2019) platforms.

**siRNA Mediated Gene Silencing**

*Silencer Select* siRNAs (siNeg: *Silencer™* Select Negative Control No. 1, siCALR 1: s115, siCALR 2: s116; siHSPB1: s194537 and s194538) were purchased from Thermo Fisher Scientific. siRNAs were resuspended in RNase-free water to a concentration of 50μM for long term storage at -20°C. siRNAs were diluted to a 2μM working solution for short term use. Lipofectamine 3000 was used for all siRNA transfections per manufacturer’s instructions. For a 12-well plate, 15μL of 2μM siRNA was suspended in mixture containing 100μL Opti-MEM (Invitrogen, 31985-070) and 3μL Lipofectamine 3000. In the case of siHSPB1, 7.5μL 2μM siRNA s194537 and siRNA s194538 working solutions were added. HEK293 cells or HEK293 cells stably expressing TetOn-mCherry-eGFP-RAMP4 were plated on 12-well plates and immediately transfected with siRNA. After 24 hours, cells were again transfected with siRNA. After 48 hours, media was changed and cells were transfected a third time with siRNA. The next
day, cells were either harvested via trypsin for Western blot, or were prepared for the EATR assay as described above. DMEM-10 without G418 was used during these experiments in order to minimize toxic effect of gene silencing.

**Western blotting**

Western blotting was performed on a 4–12% Bis-Tris SDS-PAGE system (NuPAGE, Invitrogen) (Costantini et al., 2006; Jonas et al., 2010; Peng et al., 2014; Peng et al., 2018). The primary antibodies used in this study are listed in Table S3. Appropriate secondary antibodies were diluted 1:15000 or 1:20000 in blocking solution at room temperature for 1 hour. Membranes were washed with 1X Tris-Buffered Saline with 0.1% Tween® 20 Detergent (Thermo Fisher Scientific 28360), and protein bands were developed using the LI-COR Odyssey infrared imaging system (LI-COR Biosciences). The original uncropped Western blot images included in the main manuscript can be found in Figures S4-S7.

**Co-immunoprecipitation**

Protein extracts were generated using GTIP buffer (10 mM Tris, pH 7.6, 2 mM EDTA, 0.15 M NaCl) supplemented with 1% Triton X-100 (Roche Applied Science 11332481001), 0.25% Nonidet P-40 (Sigma N-6507), complete protein inhibitor mixture (Roche Applied Science 11836170001), and phosphatase inhibitors (mixture set I and set II; Calbiochem). Extracts were pre-cleared with protein A magnetic beads (Polysciences 84600, Bio-Rad 1614013). For immunoprecipitation, protein A magnetic beads were washed with cold GTIP buffer solution and pre-incubated for 4hr at 4°C with rotation in the presence of the specific antibody. The lysate was then added to washed beads and incubated for 12hr at 4°C.
Elution of protein-coupled beads was carried out in 20-μl LDS sample buffer 4X with heating at 95°C for 5 minutes.

**Statistics**

Data are expressed as mean ± standard deviation of separate experiments. Data analysis was performed using GraphPad Prism Version 7.0.5. Comparison of the means was performed using Student's t test, or one- or two-way ANOVA. Differences were declared statistically significant if p<0.05, and the following statistical significance indicators are used: *p<0.05; **p<0.005; #p<0.0005.

**References to Transparent Methods**

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