Lysosomal Machinery Drives Extracellular Acidification to Direct Non-apoptotic Cell Death

Graphical Abstract

Highlights

- Lysosome-associated genes are required in follicle cells for nurse cell death
- V-ATPases localize to the plasma membrane of follicle cells to acidify nurse cells
- Cathepsin L is released by follicle cells and promotes nurse cell DNA degradation

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In Brief

Mondragon et al. show that V-ATPase proton pumps localize to the plasma membrane of follicle cells and promote extracellular acidification to eliminate adjacent nurse cells in the Drosophila ovary. The follicle cells subsequently release cathepsins by exocytosis into the nurse cells to promote their final degradation.
Lysosomal Machinery Drives Extracellular Acidification to Direct Non-apoptotic Cell Death

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SUMMARY

Cell death is a fundamental aspect of development, homeostasis, and disease; yet, our understanding of non-apoptotic forms of cell death is limited. One such form is phagoptosis, in which one cell utilizes phagocytosis machinery to kill another cell that would otherwise continue living. We have previously identified a non-autonomous requirement of phagocytosis machinery for the developmental programed cell death of germline nurse cells in the Drosophila ovary; however, the precise mechanism of death remained elusive. Here, we show that lysosomal machinery acting in epithelial follicle cells is used to non-autonomously induce the death of nearby germline cells. Stretch follicle cells recruit V-ATPases and chloride channels to their plasma membrane to extracellularly acidify the germline and release cathepsins that destroy the nurse cells. Our results reveal a role for lysosomal machinery acting at the plasma membrane to cause the death of neighboring cells, providing insight into mechanisms driving non-autonomous cell death.

INTRODUCTION

Programmed cell death is essential in the development of an organism for elimination of dangerous cells and to maintain homeostasis (Jacobson et al., 1997). Apoptosis is the most heavily studied type of cell death (Galluzzi et al., 2015); however, it has recently been proposed that apoptosis may not be the most prevalent form of cell death in vertebrate development (Kutscher and Shaham, 2017). Work on non-apoptotic forms of cell death over the last decade has culminated in five proposed classifications of cell death: apoptotic, autophagic, necrotic, non-cell autonomous, and atypical cell death (Martins et al., 2017).

Entosis and phagoptosis are both types of cellular cannibalism that fall within the non-cell autonomous classification of cell death modalities. In entosis, internalized cells form adherent junctions and invade the neighboring cell, bypassing any requirement for conventional phagocytosis machinery (Overholtzer et al., 2007). In contrast, phagoptosis utilizes phagocytosis machinery to drive the death of a nearby cell that would otherwise continue living (Brown and Neher, 2012, 2014). Phagoptosis has been suggested to promote physiological cell deaths, such as neuronal loss associated with stroke (Lana et al., 2017; Neher et al., 2013), Parkinson’s disease (Barcia et al., 2012), amyotrophic lateral sclerosis (ALS) (Liu et al., 2012), and retinitis pigmentosa (Zabel et al., 2016; Zhao et al., 2015).

Developmental germ cell death is a common phenomenon during oogenesis throughout metazoans (Peterson et al., 2015; Tilly, 2001). The Drosophila ovary provides a particularly powerful in vivo model for non-apoptotic germ cell death, given the large size of the cells, genetic tools, and reproducibility of cell death. Each Drosophila ovary is comprised of hundreds of developing egg chambers, each composed of 15 germline nurse cells (NCs) connected to a single oocyte, surrounded by a layer of follicle cells (FCs) (King, 1970). At the end of oogenesis, the NCs dump their cytoplasmic contents into the oocyte, become surrounded by a subset of FCs called stretch follicle cells (SFCs) (Duhart et al., 2017), and are eliminated without the requirement of apoptosis or autophagy genes (Peterson and McCall, 2013). We have previously found that the SFCs require phagocytic machinery to eliminate the NCs, demonstrating that the NCs die by phagoptosis (Santoso et al., 2018; Timmons et al., 2016, 2017); however, our understanding of how NC elimination is carried out remained limited.

Our previous studies revealed a role for lysosomal genes in NC death; however, the exact contribution of lysosomal genes was unknown (Bass et al., 2009; Timmons et al., 2016). Lysosomes have diverse functions: they are responsible for the degradation of materials in endocytosis or autophagy, repairing the plasma membrane through lysosome secretion, and metabolic signaling (Settembre et al., 2013). Lysosomes have also been linked to cell death. For example, in entosis, lysosomes act as the final executor as they fuse with the internalized cell. Lysosomes contain over 50 acid hydrolases that are involved in degradation (Lübke et al., 2009). Of particular interest are cathepsins, which are lysosomal proteases that require acidic conditions to be proteolytically active. Cathepsins can also be secreted by some specialized cells, such as osteoclasts, to degrade bone and by cancer cells to facilitate invasion (Baron et al., 1988; Rozhin et al., 1994).
V-ATPases are a vital component of lysosomes that maintain the acidic pH by pumping protons into the lumen. V-ATPases are composed of a transmembrane complex and a cytoplasmic complex that together hydrolyze ATP to pump protons across a membrane (Cotter et al., 2015). V-ATPases have 14 subunits, encoded by 33 genes in Drosophila, with many of the genes having tissue-specific expression (Allan et al., 2005). V-ATPases are well known for their roles in acidification of lysosomes; however, V-ATPases also play an important role at the plasma membrane in specific cell types in humans, such as osteoclasts for bone resorption (Allan et al., 2005), intercalated cells in the kidney to regulate systemic pH (Brown et al., 2009), clear cells in the testis to maintain acidic luminal fluid (Shum et al., 2009), and cancer cells to acidify the extracellular matrix to facilitate invasion (Stransky et al., 2016).

Here, we report the essential role of lysosome-associated genes in NC death. Specifically, we show a non-autonomous role for V-ATPases and cathepsins in NC acidification and elimination. V-ATPases are enriched and recruited to the plasma membrane of the SFCs to extracellularly acidify the NCs, and cathepsins are released from the SFCs to drive NC degradation in a manner resembling osteoclast degradation of bone. Altogether, this work characterizes a novel role for V-ATPases and cathepsins acting at the plasma membrane to drive the death of a neighboring cell.

RESULTS

Nurse cells Are Acidified Extracellularly

Fifteen NCs are each developing oocyte from the earliest stages of oogenesis through ring canals formed by incomplete cytokinesis (Spradling, 1993). The NCs support the growth and development of the oocyte throughout oogenesis by delivery of organelles, proteins, and RNA to the oocyte. Near the end of oogenesis, the NCs begin to show signs of cell death beginning at stage 10, with distinct changes, including cytoskeletal rearrangements, the leakage of nuclear material, and nuclear remodeling seen by transmission electron microscopy (TEM) (Cooley et al., 1992; Guild et al., 1997). During stage 11, the NCs rapidly transfer their cytoplasm into the oocyte (Spradling, 1993). The nearby SFCs surround the NCs by stage 12 (Duhart et al., 2017) and are required for multiple cell death events in the NCs, including cytoskeletal rearrangements and cytoplasmic transfer (Timmons et al., 2016). During stages 12 and 13, the NCs become acidified and DNA is fragmented (Bass et al., 2009; Foley and Cooley, 1998). The NCs are subsequently degraded by stage 14, leaving the fully intact mature oocyte.

One of the most unusual cell death events of the NCs is their complete acidification (Bass et al., 2009; Timmons et al., 2016). Previously, we had determined that acidic organelles were detected in SFCs prior to the acidification of the NCs, but it was unclear how the NCs became acidified. To investigate how NCs become acidified, we recorded time-lapse images of stage 13 egg chambers, the developmental stage when NC acidification occurs (Figures 1A–1E). SFC membranes were visualized using an SFC-specific GAL4 (see STAR Methods) to drive expression of a membrane-tethered (myristoylated) GFP. To detect acidification of the NCs, egg chambers were labeled with LysoTracker (LT), an acidotropic dye (Timmons et al., 2013). During stage 13, mobile acidic vesicles in the SFCs were observed surrounding NCs prior to their acidification. Fixed tissue staining also demonstrated an accumulation of acidic vesicles in SFCs before the acidification of NCs (Figures 1F–1G’ and S1A–S1E) (Timmons et al., 2016). The acidic vesicles and acidified NCs exhibited a different pattern than lysosomes detected by LAMP1 staining (Figures S1F–S1G). Live imaging over the course of an hour showed the dynamic activity and accumulation of LT vesicles in the SFCs before NC acidification (Video S1).

To further understand the process of NC acidification, we generated flies with a membrane-bound pH detector, pHRed-CAAX (Figure 1H). pHRed is a genetically encoded pH sensor (Tantama et al., 2011) modified with a CAAX motif that localizes pHRed to the cytoplasmic side of the plasma membrane (Hancock et al., 1991). To confirm that pHRed served as an engulfment detector, we expressed it in neurons and NCs and monitored its fluorescence following induction of apoptosis. In both cases, pHRed was not detected in healthy cells but was detected as punctate staining adjacent to dying cells, suggesting that the dying cell material was taken up and acidified within phagosomes (Figures S1H–S1I’). We next characterized pHRed in late-stage NCs to determine how NCs were acidified during developmental cell death. Unlike the labeling from engulfed apoptotic cells, pHRed was first detected along the NC membrane adjacent to SFCs, followed by pHRed labeling of entire NC remnants (Figures 1I–1J’). These two distinct phases of pHRed detection indicate that acidification initiates when the NC membrane is intact and progresses as the NC membrane is broken down and dispersed throughout the cell. The initial acidification of the membrane suggests that the NCs are acidified extracellularly by the SFCs and subsequently degraded.

V-ATPases Are Required for Acidification and Clearance of NCs while Cathepsins Are Only Required for Clearance of NCs

Previously, we discovered that lysosomal trafficking genes were required non-autonomously in the FCs for NC acidification (Timmons et al., 2016). To further understand the role of lysosomes in NC elimination, we screened additional lysosome-associated genes using RNAi to knock down selected genes in the FCs and determine the effect on the NCs. The screen revealed a major requirement for V-ATPases and cathepsins. Although lysosomes are an essential component of phagosome maturation, little is known about the potential requirement of lysosomal machinery for cell death. Given the unusual acidification of NCs, we tested whether these lysosomal components were required for NC acidification and clearance. In normal development, NCs are asynchronously acidified and cleared between stages 13 and 14; quantification revealed that 50.5% of NCs were acidified in stage 13 and 91.9% were cleared by stage 14 (Figures 2A, 2C, 2E, and 2F). Knock down of V-ATPase subunits Vha100-2 or Vha16-1 in FCs significantly reduced the acidification of NCs in stage 13 egg chambers to 6.6% and 5.6% respectively (Figures 2B, 2C, and 2E). In contrast, no acidification defect was observed when CP1, the Drosophila ortholog of cathepsin L (Tryselli and Hultmark, 1997), was knocked down (Figures 2D
and 2E). Knock down of either of the V-ATPase subunits or CP1 resulted in persisting NCs (Figures 2B’–2D’ and 2F). These findings suggest a two-step process where V-ATPase activity is required in the SFCs to first acidify the NCs and, subsequently, the SFCs utilize CP1 for NC degradation.

V-ATPases Are Enriched in Stretch Follicle Cells and Localize to the Plasma Membrane

To visualize V-ATPase expression in the ovary, we examined reporter lines for several of the subunits. We first examined a Vha68-2 (subunit A) enhancer trap that expresses nuclear GFP.
previously shown to correlate with Vha68-2 expression (Zhang et al., 2015). We colabeled egg chambers with an antibody against Eya that is expressed specifically in SFCs (Grammont, 2007). In stage 13 egg chambers, we found that Vha68-2 expression was increased 2.5-fold in SFCs (Figures 3A, 3A’, S2A, and S2B) compared to other FCs. Colabeling with LT revealed that Vha68-2 expression was enriched in FCs adjacent to NCs (Figures 3B and 3B’). These findings demonstrate that V-ATPase expression is enriched in SFCs during stage 13 when NCs are being acidified.

To determine V-ATPase subcellular localization, we examined GFP protein traps for VhaSFD (subunit H), Vha13 (subunit G), and Vha44 (subunit C) (Buszczak et al., 2007; Morin et al., 2001; Nagarkar-Jaiswal et al., 2015). Null mutants of VhaSFD are homozygous lethal (Allan et al., 2005), but VhaSFD-GFP flies are homozygous viable, suggesting that the subunit functions normally with the GFP tag. All three of these V-ATPase protein traps had similar enrichment in SFCs, such as the Vha68-2 enhancer trap, but surprisingly, they localized to the plasma membrane of the SFCs (Figures 3A, 3A’, 3B, and 3B’) rather than lysosomes (Figures 3C and 3C’) rather than lysosomes. Immunohistochemistry with an antibody against ATP6V1B1, the human homolog of Vha55 (subunit B), also demonstrated enrichment at the SFC plasma membrane (Figures 3D and 3D’), and colocalization was observed with a membrane marker (Figure S3). Taken together, these data demonstrate that V-ATPases are enriched in SFCs and localize to the plasma membrane. The localization of V-ATPases is different than either LT or LAMP1 staining (Figures 1 and S1A–S1G’), suggesting that V-ATPases are not delivered to the membrane by lysosome fusion. This localization combined with their requirement for NC acidification suggests that they function by extracellularly acidifying the nearby NCs, similar to V-ATPases acting at the plasma membrane in bone resorption or cancer cell invasion.

In human osteoclasts, V-ATPases are the primary proton pumps driving acidification of bone; however, to prevent a large difference in membrane potential, a chloride pump is also present at the plasma membrane (DiCiccio and Steinberg, 2011). Loss of the chloride channel (CLCN7) in humans leads to inefficient acidification of bone and leads to osteoporosis (Kornak et al., 2001). The Drosophila ortholog of CLCN7 is ClC-b, and it has been previously studied for its role in endolysosomes (Wong et al., 2017). To determine the expression and localization of ClC-b, we utilized a GFP protein trap. In stage 13 egg chambers, we found that ClC-b was enriched specifically in the SFCs and localized to the membrane as the NCs were becoming acidified (Figures 3E and 3E’). Altogether, this suggests that the SFCs may be utilizing the same machinery as osteoclasts for extracellular acidification.

V-ATPases and Cathepsins Are Non-autonomously Required for DNA Fragmentation of Nurse Cells

Previously, we found that disruption of phagocytosis genes in SFCs led to a block in both NC acidification and DNA fragmentation visualized by TUNEL staining (Timmons et al., 2016). To explore the role of acidification in DNA fragmentation, we performed TUNEL staining on V-ATPase and CP1 knockdowns.
control stage 13 egg chambers, 60.9% of NCs labeled positively with TUNEL. Knocking down V-ATPase subunits or CP1 significantly reduced TUNEL-positive NCs to 8.56% (Vha16-1), 18.15% (Vha100-2), and 22.24% (CP1) (Figures 4A–4E). Thus, we conclude that V-ATPase activity and CP1 are both required for DNA fragmentation, a defining step of cell death (Galluzzi et al., 2015).

To visualize FC-derived CP1 during NC death, we expressed a hemagglutinin (HA)-tagged CP1 and a membrane-tethered GFP in the FCs. Initially, CP1 formed aggregates within the FCs; however, we also discovered that CP1 was released from the FCs into some of the NCs in stage 13 (Figures 4F–4F‴). Cathepsins are active only in acidic conditions, so we examined whether the NCs containing FC-derived CP1 were acidified. Consistently, acidified NCs contained CP1 that had been released from the FCs, whereas NCs that had not yet been acidified did not contain CP1 (Figures 4G–4G‴).

To determine whether cathepsins were released from FCs by exocytosis, we knocked down two SNARE proteins associated with exocytosis, namely, Snap24 and Syx6 (Littleton, 2000; Nimmeyer and Schwarz, 2000). We detected CP1 with an antibody that localized similarly to the CP1-HA construct (Figures S4A–S4A‴). Quantification of the amount of CP1 localized in NCs revealed a significant decrease when Snap24 or Syx6 was knocked down in the FCs (Figures S4B–S4E). Additionally, knock down of either of these genes resulted in a significant decrease in NC acidification in stage 13 egg chambers and showed persisting NC nuclei in stage 14 egg chambers (Figures S4F and S4G). However, we found that a V-ATPase subunit localized to the SFC membrane normally in a Snap24 knockdown (Figure S4H), indicating that V-ATPases are targeted to the membrane independent of exocytosis.

Altogether, our findings suggest that CP1 is released from the FCs following NC acidification and both the acidification by V-ATPases and proteolytic activity of CP1 are required for the DNA fragmentation and elimination of the NCs.

**DISCUSSION**

Phagoptosis is defined as a type of cell death that requires phagocytosis machinery (Brown and Neher, 2012). We have previously demonstrated that NC death requires phagocytic machinery, such as Draper and Ced-12 (Santoso et al., 2018; Timmons et al., 2016). In the present study, we identified lysosome-associated genes required by the SFCs that non-autonomously control the elimination of NCs. To our knowledge, this
is the first example of V-ATPases at the plasma membrane driving acidification and subsequent cathepsin release to destroy a nearby cell. These findings suggest that signaling from the phagocytic machinery promotes this use of lysosomal proteins in NC elimination. Whether other examples of phagoposis use the lysosomal machinery in this way remains to be determined.

V-ATPases are enriched at the plasma membrane of several specialized cell types in humans, such as osteoclasts, intercalated cells, clear cells, and some cancer cells. In insects, V-ATPases can be found on the plasma membranes of cells in certain tissues, such as Malpighian tubules (Bertram and Wessing, 1994; Day et al., 2008) and vas deferens (Bebas et al., 2002) to regulate pH or in earlier stages of oogenesis to play a role in bioelectric patterning (Kruger and Bohrmann, 2015). The specific isoforms associated with the plasma membrane V-ATPase holoenzyme have previously been identified (Allan et al., 2005). In this paper, we demonstrated that 7 of these plasma-membrane-associated V-ATPase subunits (Vha16-1, Vha100-2, Vha68-2, VhaSFD, Vha13, Vha44, and Vha55) are either enriched in the SFCs, localize to the SFC membrane, or are required for NC acidification.

**Figure 4. Cathepsin L Is Released by SFCs and Required for NC DNA Degradation**

(A–D) Stage 13 egg chambers of the indicated genotypes labeled with TUNEL (magenta, arrows) and DAPI (cyan).

(E) Quantification of TUNEL-positive NC nuclei in stage 13 egg chambers. ****p ≤ 0.0001.

(F–F'') FC > mCD8-GFP, CP1-HA stage 13 egg chamber labeled with anti-HA (magenta) and DAPI (cyan). CP1-HA (magenta) is produced in the FCs (green, arrowhead) and released into the NCs (arrow).

(G–G'') FC>CP1-HA egg chambers labeled with LT (magenta) and DAPI (cyan). CP1 is detected within LT-positive NCs (arrows) and not LT-negative NCs (arrowhead). Scale bars, 50 μm.
Our previous work also highlighted Draper as being required for both the presence of LT vesicles in SFCs and NC acidification, suggesting that Draper initiates this process (Timmons et al., 2016). Other studies have demonstrated a link between Draper and autophagy genes (Etchegaray et al., 2016; McPhee and Baehrecke, 2010). Further work will need to be done to elucidate the upstream signaling components required to promote the phagoptotic potential of V-ATPases and cathepsins, and the role of SFC LT vesicles preceding NC acidification.

The findings reported here suggest a role for membrane-localized V-ATPases and cathepsin release in promoting cell death by phagoptosis. We have demonstrated that this mechanism is used during developmental NC death, but this mechanism may be used more broadly in other cell deaths that have been found to be non-apoptotic. Developmental germ cell death occurs in many organisms ranging from Hydra (Baum et al., 2005) to humans (Baker, 1963), and it is possible that surrounding somatic cells could contribute to the death of the germ cells in these other species. Our work also brings up the intriguing possibility that the lysosomal machinery can be harnessed to murder neighboring cells in other contexts.

STAR METHODS

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

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2019.03.034.

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AUTHOR CONTRIBUTIONS

Conceptualization, A.A.M., A.Y., J.E., and K.M.; Methodology, A.A.M., A.Y., J.E., W.C., and K.M.; Investigation, A.A.M., A.J.O., Y.Z., and O.N.; Writing – Original Draft, A.A.M.; Writing – Review and Editing, A.A.M. and K.M.; Funding Acquisition, A.A.M., K.M., A.Y., and W.-S.C.; Supervision, A.A.M. and K.M.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Eya                 | DSHB   | Cat# Eya10H6; RRID:AB_528232 |
| ATP6V1B1            | Abagent| Cat# AP11538C-ev; RRID:AB_2797396 |
| Goat-α-mouse Cy3    | Jackson ImmunoResearch | Cat# 115-165-003; RRID:AB_2338680 |
| CP1                 | R&D Systems | Cat# MAB22591; RRID:AB_2087830 |
| LAMP1               | Abcam  | Cat# Ab30687; RRID:AB_775973 |
| Goat-α-rabbit 647   | Jackson ImmunoResearch | Cat# 111-605-144; RRID:AB_2338078 |
| Bacterial and Virus Strains |        |            |
| NEB 5’ 5-alpha F’IQ Competent E. coli | NEB | C2992H |
| Chemicals, Peptides, and Recombinant Proteins |        |            |
| LysoTracker         | Thermo Fisher Scientific | L75283 |
| Vectashield with DAPI | Vector Laboratories | H-1200 |
| Hoechst 33342       | Thermo Fisher Scientific | 62249 |
| In Situ Cell Death Detection Kit, TMR Red | Sigma-Aldrich | 12156792910 |
| Experimental Models: Organisms/Strains |        |            |
|  Stretch follicle cell) PG150-GAL4 | Dr. Ellen LeMosy | N/A |
| (Follicle cell) GR1-GAL4 | Dr. Trudi Schupbach | N/A |
| (Germline) NGT;nanos-GAL4 | Dr. Pernille Rorth | N/A |
| UAS-CP1-HA          | FlyORF  | 780 |
| Vha68-2-GFP         | Dr. Francesca Pignoni | N/A |
| Clc-b-GFP           | Dr. Kartik Venkatachalam | N/A |
| Vha13-GFP           | Bloomington Stock Center | 50828 |
| Vha44-GFP           | Bloomington Stock Center | 63202 |
| VhaSF-D-GFP         | Bloomington Stock Center | 6840 |
| Luciferase RNAi (JF01355) | Bloomington Stock Center | 31603 |
| Vha100-2 RNAi (HMC05732) | Bloomington Stock Center | 64859 |
| Vha16-1 RNAi (HMS02171) | Bloomington Stock Center | 40923 |
| CP1 RNAi (HMS00725) | Bloomington Stock Center | 32932 |
| Snap24 RNAi (JF03146) | Bloomington Stock Center | 28719 |
| Synx6 RNAi (JF03125) | Bloomington Stock Center | 28505 |
| Myr-GFP (pJFRC29- 10XUAS-IVS-myr::GFP-p10) | Dr. Norbert Perrimon | N/A |
| Myr-RFP             | Bloomington Stock Center | 7118 |
| Gal80ts             | Bloomington Stock Center | 7019 |
| Recombinant DNA     |        |            |
| pHRed-cAAX Plasmid  | Dr. Won-Suk Chung | N/A |
| UASP (pPW) Plasmid  | DGRC   | 1130 |

CONTACT FOR REAGENT AND RESOURCE SHARING

Any requests for resources and reagents should be directed to the lead contact, Kim McCall (kmccall@bu.edu).
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Fly Strains
PG150-GAL4 was used to drive expression in SFCs (indicated by SFC > in figures) and was provided by Ellen LeMosy (Dinkins et al., 2008). GR1-GAL4 was used to drive expression in FCs (indicated by FC > in figures) and was provided by Trudi Schüpbach (Goentoro et al., 2006). LuciferaseRNAi was used as a control for all RNAi experiments. UAS-CP1-HA was obtained from FlyORF (Bischof et al., 2013). Vha68-2-GFP was provided by Francesca Pignoni (Zhang et al., 2015). ClC-b-GFP was provided by Kartik Venkatachalam (Wong et al., 2017). UAS-Myr-GFP was provided by Norbert Perrimon (Pfeiffer et al., 2012).

Fly Husbandry
Flies were age matched and dissected before 2 weeks of age. All RNAi crosses were performed in combination with Gal80ts at 18°C and moved to 29°C for 48 hours, except the crosses for Figure S4 which were done without Gal80ts at 25°C and moved to 29°C for 48 hours. All other fly crosses were kept at 25°C. All flies were well fed with yeast paste for 2 days (new yeast paste each day) before dissection to increase egg chamber production.

METHODS DETAILS

Molecular Cloning
pHRed-CAAX was PCR amplified from a plasmid obtained from Dr. Won-Suk Chung and cloned into a Gateway entry vector pENTR using the pENTR/D-TOPO cloning kit (Invitrogen). Gateway LR Clonase (Invitrogen) was used to recombine the construct downstream of the GAL4 responsive promoter UASp (in the plasmid pPW, received from Drosophila Genome Resource Center, Bloomington, IN). The P element vector with pHRed-CAAX was injected into embryos for P-element transformation by BestGene (Chino Hills, CA).

Live Imaging
Live imaging was performed as described (Peters and Berg, 2016). Briefly, stage 13 egg chambers were individually dissected from flies in room temperature Schneider’s medium. Once isolated, egg chambers were transferred to Schneider’s medium (product number 21720-024, Thermo Fisher Scientific) with LysoTracker (LT, LysoTracker Red DND-99 – Invitrogen by Thermo Fisher Scientific L75283 – 1:1000 dilution) and Hoechst 33342 (product number 62249, Thermo Fisher Scientific– 10 μM). Egg chambers in solution were transferred to the imaging chamber which had a glass bottom. An immobilization blanket (small Kimwipe) was used to keep egg chambers in place during imaging. The immobilization blanket was placed in the solution on top of the egg chambers and a brass washer was placed on the blanket to hold it in place. Live imaging was captured on a Nikon C2+Si laser scanning confocal microscope.

Immunohistochemistry
For LT staining, whole ovaries were dissected from flies in Grace’s insect media (product number BW04-457F, Thermo Fisher Scientific) and incubated in LT solution (Thermo Fisher Scientific, 1:50 in PBS) for 6 minutes, rinsed with PBS for 30 min while rotating, fixed in 300 uL Graces, 200 uL Heptane and 4% Paraformaldehyde, washed with PBS + 1% Triton X-100 (PBT) for 15 min, and stained with DAPI. For antibody staining, ovaries were immediately fixed after dissection, washed with PBT for 1 hour, blocked with PBANG (PBT, 0.5% BSA, 5% Normal Goat Serum), and incubated overnight in primary antibody diluted in PBANG. Samples were rinsed with PBT twice, washed with PBT + 0.5% BSA for 2 hours, and incubated in secondary antibody diluted in PBANG for 1 hour, protected from light. Samples were rinsed with PBT twice, washed with PBT + 0.5% BSA for 2 hours, rinsed with PBS, and incubated in 2 drops of Vectashield with DAPI overnight before mounting. Primary antibodies were Eya (Developmental Studies Hybriodoma Bank), a stretch follicle cell specific marker, at 1:300, ATP6V1B1 (Abgent) at 1:10, CP1 (R&D Systems) at 1:200, LAMP1 (abcam) at 1:500. Secondary goat-α-mouse Cy3 (Jackson ImmunoResearch) was used 1:100 and goat-α-rabbit 647 (Jackson ImmunoResearch) was used at 1:100. Imaging of egg chambers was performed on an Olympus Fluoview FV10i laser scanning confocal or a Nikon C2+Si laser scanning confocal microscope.

TUNEL
To assay for DNA fragmentation, we dissected ovaries from flies in 2% paraformaldehyde in PBS with 0.1% Triton X-100. Ovaries were then fixed for 45 min, rinsed with PBT twice, washed in PBT + 0.1% sodium citrate and 0.1% Triton X-100 at 65°C for 30 min. Tissue was then washed 3 times in PBT for 20 min each. Tissue was incubated in 36 uL TUNEL label solution and 4 uL enzyme solution (In Situ Cell Death Detection Kit, TMR Red – Sigma-Aldrich, Cat #12 156 792 910) for 3 hours at 37°C. Samples were washed 4X in PBS for a total time of 1 hour and mounted in Vectashield with DAPI.

Quantification and Statistical Analysis
All data were graphed and analyzed in Graphpad Prism, and an unpaired t test was performed on each set of data compared to control. The mean was graphed ± SEM. Each experiment had at least three biological replicates. At least 10 flies were randomly selected.
for each replicate and egg chambers were dissected and staged using previously described methods (Spradling, 1993) on an Olympus BX60 upright fluorescence microscope. For quantification of Vha68-2 GFP intensity (Figure S2B), the mean GFP intensity of Eya+ follicle cells (SFCs) and Eya− follicle cells was measured by ImageJ after outlining only Eya+ or Eya− nuclei. Measurement of CP1 pixels (Figure S4E) was also performed in ImageJ. NC nuclei (regions of interest) were outlined based on DAPI staining, the CP1 channel was converted to a black and white image, and the pixels in the regions of interest were counted in the CP1 channel.

**Sample Sizes**

Figure 2E - GR1 (FC) > LucRNAi, n = 72 egg chambers; GR1 (FC) > Vha16-1RNAi, n = 28; GR1 (FC) > Vha100-2RNAi, n = 50; GR1 (FC) > CP1RNAi, n = 19.

Figure 2F - GR1 (FC) > LucRNAi, n = 85; GR1 (FC) > Vha16-1RNAi, n = 72; GR1 (FC) > Vha100-2RNAi, n = 118; GR1 (FC) > CP1RNAi, n = 38.

Figure 4E - GR1 (FC) > LucRNAi, n = 42; GR1 (FC) > Vha16-1RNAi, n = 47; GR1 (FC) > Vha100-2RNAi, n = 43; GR1 (FC) > CP1RNAi, n = 50.