The Drosophila effector caspase Dcp-1 regulates mitochondrial dynamics and autophagic flux via SesB

Lindsay DeVorkin,1,2 Nancy Erro Go,1 Ying-Chen Claire Hou,1 Annie Moradian,3 Gregg B. Morin,1,3 and Sharon M. Gorski1,2

1The Genome Sciences Centre, British Columbia Cancer Agency, Vancouver, British Columbia V5Z 1L3, Canada
2Department of Molecular Biology and Biochemistry, Simon Fraser University, Burnaby, British Columbia V5A 1S6, Canada
3Department of Medical Genetics, University of British Columbia, Vancouver, British Columbia V5Z 1L3, Canada

Increasing evidence reveals that a subset of proteins participates in both the autophagy and apoptosis pathways, and this intersection is important in normal physiological contexts and in pathological settings. In this paper, we show that the Drosophila effector caspase, Drosophila caspase 1 (Dcp-1), localizes within mitochondria and regulates mitochondrial morphology and autophagic flux. Loss of Dcp-1 led to mitochondrial elongation, increased levels of the mitochondrial adenine nucleotide translocase stress-sensitive B (SesB), increased adenosine triphosphate (ATP), and a reduction in autophagic flux. Moreover, we find that SesB suppresses autophagic flux during midoogenesis, identifying a novel negative regulator of autophagy. Reduced SesB activity or depletion of ATP by oligomycin A could rescue the autophagic defect in Dcp-1 loss-of-function flies, demonstrating that Dcp-1 promotes autophagy by negatively regulating SesB and ATP levels. Furthermore, we find that pro-Dcp-1 interacts with SesB in a nonproteolytic manner to regulate its stability. These data reveal a new mitochondrial-associated molecular link between nonapoptotic caspase function and autophagy regulation in vivo.

Introduction

Autophagy is a cellular self-digestion process in which long-lived proteins and organelles are engulfed in double-membrane vesicles called autophagosomes. After fusion with lysosomes to form autolysosomes, the cellular contents are degraded, and the resulting breakdown products, including constituents for ATP production and protein synthesis, are recycled back into the cell (Rabinowitz and White, 2010). Autophagic flux, which refers to the complete process of autophagy, occurs at basal levels and is also up-regulated in response to various cellular stresses, including nutrient deprivation (Filkins, 1970; Scott et al., 2004), chemotherapies (Bursch et al., 1996; Kanzawa et al., 2003; Høy-Hansen and Jäättelä, 2008), and reactive oxygen species (Scherz-Shouval et al., 2007; Chen et al., 2009). As such, autophagy acts to remove toxic metabolites, harmful protein aggregates, and damaged organelles and provides a mechanism for energy production in instances of metabolic stress to promote cell survival. It is therefore not surprising that defects in autophagic flux have been implicated in several human diseases, including cancer (Kondo et al., 2005; Kimmelman, 2011) and neurodegeneration (Wong and Cuervo, 2010).

The relationship between apoptosis and autophagy is complex, as core machinery components and signaling molecules of the pathways are interconnected (Eisenberg-Lerner et al., 2009). However, the molecular mechanisms governing these interactions are largely unknown. Several studies have begun to examine the multifunctional signaling molecules that take part in each pathway. For example, antiapoptotic Bcl-2 and Bcl-xL bind Beclin1, a core autophagy component of the class III phosphoinositide 3-kinase complex, and inhibit autophagy (Pattingre et al., 2005; Maituri et al., 2007). Furthermore, the core autophagy protein Atg12 enhances mitochondrial apoptosis by binding to and inactivating antiapoptotic family members, including Bcl-2 and Mcl-1 (Rubinstein et al., 2011). Several studies have also revealed that core autophagy proteins undergo proteolytic processing,
and in some cases, the newly generated fragment gains a novel proapoptotic role (Yousefi et al., 2006; Cho et al., 2009; Wirawan et al., 2010; Zhu et al., 2010).

Studies using *Drosophila melanogaster* have also examined the complex relationship between autophagy and apoptosis. Overexpression of the autophagy gene *Atg1* in the larval fat body, a nutrient storage organ analogous to the mammalian liver, induces apoptosis in a caspase-dependent manner (Scott et al., 2007). Degradation of larval tissues, including the salivary gland (Berry and Baehrecke, 2007) and the midgut (Denton et al., 2009), during development requires autophagy. Although these studies show that autophagy can lead to the induction of death as well as contribute to death-related processes, autophagy functions primarily as a cell survival mechanism in *Drosophila* in response to cellular stress. For example, JNK signaling in the intestinal epithelium and fat body stimulates autophagy gene transcription to promote cell survival during oxidative stress (Wu et al., 2009). Furthermore, autophagy is induced to high levels in the larval fat body (Scott et al., 2004) and the midgut (Wu et al., 2009) to remobilize nutrients and promote cell survival after starvation.

The *Drosophila* ovary is also sensitive to nutritional cues. The ovary is made up of 15–20 ovarioles, each containing a series of developing egg chambers that arise from the germarium and progress through 14 well-defined stages. Each egg chamber is made up of 15 germline nurse cells and one oocyte surrounded by a layer of somatically derived follicle cells. Cell death in midstage egg chambers occurs sporadically and in response to environmental cues, including nutrient deprivation (Drumond-Barbosa and Spradling, 2001). Midstage egg chambers undergoing cell death are characterized by nurse cell nuclei condensation and fragmentation, engulfment of the nurse cell cytoplasm by the surrounding follicle cells and follicle cell death (Giorgi and Deri, 1976). Autophagy has been observed in degenerating midstage egg chambers (Hou et al., 2008; Nezis et al., 2009), and we previously showed that both cell death and autophagy are dependent on the effector caspase *Drosophila* caspase-1 (Dcp-1; Song et al., 1997; Laundrie et al., 2003; Hou et al., 2008). However, whether autophagic flux occurs in response to starvation in nondegenerating and degenerating midstage egg chambers and the mechanism of Dcp-1–induced autophagy remain to be identified.

In this study, we demonstrate that autophagic flux occurs in healthy and degenerating midstage egg chambers in response to starvation in a Dcp-1–dependent manner. We find that Dcp-1 localizes to the mitochondria under nutrient-rich conditions in which it functions to regulate mitochondrial network morphology and ATP levels. The proform of Dcp-1 interacts with the adenine nucleotide translocase (ANT) stress-sensitive B (SesB), a mitochondrial protein that functions to regulate ATP levels, in a nonproteolytic manner. SesB mutant analysis reveals a new role for SesB as a negative regulator of autophagic flux in *Drosophila* midstage egg chambers. Depletion of ATP or SesB loss-of-function flies can rescue the autophagy defect in *Dcp-1* loss-of-function flies, demonstrating that SesB acts downstream of Dcp-1 in the regulation of autophagy. These data uncover a novel mechanism of caspase-mediated regulation of autophagy in vivo.

### Results

**Starvation-induced autophagic flux occurs during *Drosophila* midoogenesis and is regulated by Dcp-1**

Using GFP-LC3 as a marker of autophagosomes, we previously showed that loss of Dcp-1 results in a reduction of starvation-induced autophagosome formation in vitro in *Drosophila* *l(2)mbn* cells and in vivo in degenerating midstage egg chambers (Hou et al., 2008). However, whether Dcp-1 affects autophagic flux, including the degradative completion of autophagy in the lysosome, remains to be identified. To test this, we first examined transgenic flies expressing a UASp-GFP-mCherry-DrAtg8a transgene specifically in the germline using the nosGAL4 driver. Under nonautophagy-inducing conditions, the dual-tagged Atg8a protein is diffuse throughout the cytoplasm and appears yellow (overlap of green and red). When autophagy is induced, Atg8a becomes lipidated and associates with the autophagosomal membrane where it fluoresces as yellow puncta. Once autophagosomes fuse with lysosomes, GFP fluorescence is quenched by acidic hydrolases, and the resulting autolysosomes fluoresces red, signifying that autophagic flux has occurred (Nezis et al., 2010). Healthy egg chambers from flies conditioned on yeast paste (fed) showed diffuse yellow GFP-mCherry-DrAtg8a staining throughout the nurse cells (Fig. 1 A). After amino acid deprivation (starvation), we observed an increase in the percentage of autolysosomes (68% vs. 7% in fed conditions) in healthy nondegenerating midstage egg chambers (Fig. 1 B and Fig. S1 C), indicating that autophagic flux occurs in otherwise healthy egg chambers. In response to a cell death signal, nurse cell nuclei condense and fragment, follicle cells take up portions of the nurse cell cytoplasm, and this is followed by follicle cell death (Giorgi and Deri, 1976). In addition to the cell death phenotype, degenerating midstage egg chambers taking up nurse cell cytoplasm (Fig. 1 C) and those later in the degeneration process (Fig. 1 D) contained both autophagosomes and autolysosomes. In late degenerating midstage egg chambers, the GFP signal was lost, and the remaining follicle cells fluoresced red (Fig. 1 E), indicating that there may be a late stage acidification of the dying follicle cells. All together, these data show that autophagic flux occurs in the germline in response to starvation.

We next examined whether Dcp-1 regulates autophagic flux in the germline. Well-fed *Dcp-1* loss-of-function flies (*Dcp-1Prev*) expressing GFP-mCherry-DrAtg8a in the germline had diffuse yellow staining (Fig. 1, F and J). After starvation, degrading midstage egg chambers from *Dcp-1Prev* flies, which are characterized by a premature loss of follicle cells and persisting nurse cell nuclei (Laundrie et al., 2003), contained a reduction in the percentage of autolysosomes compared with the control (Fig. 1, G [compare with D] and K). Nondegenerating midstage egg chambers from starved *Dcp-1Prev* flies also showed reduced autophagic flux (Fig. S1), indicating that the reduction in autophagy in *Dcp-1Prev* flies is not simply caused by a lack of engulfment of dying nurse cells by follicle cells. To determine whether Dcp-1 is sufficient to induce autophagic flux, GFP-mCherry-Atg8a and an active form of Dcp-1 lacking its prodomain (tDcp-1) were expressed in the germline. We observed an increase in the percentage of autolysosomes in both nondegenerating (Fig. 1, H and J)
Figure 1. **Dcp-1 is necessary for autophagic flux during midoogenesis.** GFP-mCherry-DrAtg8a was expressed in the germline using the nosGAL4 driver. Staining shows DNA, GFP, and mCherry. (A) UASp-GFP-mCherry-DrAtg8a/+;nosGAL4/+ flies conditioned on yeast paste had diffuse GFP-mCherry-DrAtg8a staining in midstage egg chambers. (B) Nondegenerating midstage egg chambers from starved flies contained autophagosomes (yellow) and autolysosomes (red). (C and D) Egg chambers early in the degeneration process showed follicle cells that take up portions of the nurse cell cytoplasm (C) followed by condensation and fragmentation of the nurse cell nuclei and further uptake of the nurse cell cytoplasm into follicle cells (D). (E) Late stage degenerating egg chambers lose all GFP staining and fluoresce red. (F) Fed Dcp-1^{Prev1}/Dcp-1^{Prev1};UASp-GFP-mCherry-DrAtg8a/nosGAL4 flies showed diffuse GFP-mCherry-DrAtg8a staining in the germline. (G) Starved Dcp-1^{Prev1}/Dcp-1^{Prev1};UASp-GFP-mCherry-DrAtg8a/nosGAL4 flies showed reduced autolysosomes in degenerating midstage egg chambers. (H and I) Fed flies overexpressing truncated Dcp-1 (tDcp-1) in the germline showed increased autophagosomes and autolysosomes in nondegenerating midstage egg chambers (H) and also contained degenerating midstage egg chambers that lose all GFP fluorescence and fluoresce red (I). Bars: (main images) 25 μm; (insets) 12.5 μm. Insets in A–I show diffuse cytoplasmic staining of Atg8a or autophagosomes and autolysosomes. (J and K) The percentages of autolysosomes (% Autolysosomes) were manually calculated in at least eight egg chambers for each genotype as indicated. Error bars represent the means ± SD. Statistical testing was performed using one-way ANOVA with a Dunnet post test (****, P < 0.0001) or a two-tailed Student’s t test (**, P < 0.005).
Fig. S2, C and D), indicating that although Dcp-1 overexpression can enhance the autophagic response, Dcp-1 is not required for starvation-induced autophagy in the fat body. Dcp-1 is required for the degradation of Ref(2)P after starvation. Ref(2)P, the Drosophila homologue of p62 (Nezis et al., 2008), is a substrate of autophagy and was shown to be a marker of autophagic activity (Nezis et al., 2010; Bartlett et al., 2011). Therefore, we used Ref(2)P analyses to confirm that Dcp-1 is a positive regulator of autophagic flux. Compared with fed wild-type flies (Fig. 3 A), nutrient-deprived flies had reduced Ref(2)P in follicle cells and nurse cells (Fig. 3 B), and Western blot analysis of wild-type ovaries revealed that the level of Ref(2)P was reduced after starvation (Fig. 3 C).

Ovaries from fed or starved Atg7d77/d14 mutant flies showed an accumulation of Ref(2)P (Fig. 3, C and D). It was reported that starvation itself leads to transcriptional activation of Ref(2)P (Érdi et al., 2012), and this is likely why we see an even further increase in Ref(2)P levels after starvation in Atg7 flies. Compared with starved wild-type flies, degenerating mid-stage egg chambers from starved Dcp-1Prev1 flies contained increased levels of Ref(2)P (Fig. 3 E), and Western blot analysis of whole ovaries confirmed that Ref(2)P failed to be completely

and degenerating (Fig. 1 I) midstage egg chambers even in the absence of starvation.

To confirm our in vivo findings, we examined the role of Dcp-1 in autophagic flux in vitro using a Drosophila S2 cell line stably expressing GFP-RFP-Atg8a. To validate the cell line, we first starved cells and observed a significant increase in autolysosomes that could be inhibited by addition of the late stage autophagy inhibitor Bafilomycin A1 (Fig. 2, A and B), as expected. Next, we used RNAi to knock down Rheb, a negative regulator of autophagy, or Atg7, a positive regulator of autophagy, and detected an increase and reduction in autolysosomes, respectively (Fig. 2, C and D). Dcp-1 RNAi resulted in a significant reduction in autolysosomes (Fig. 2, C and D), consistent with the Dcp-1Prev1 phenotype in midstage egg chambers, confirming a role for Dcp-1 as a positive regulator of autophagic flux.

We also investigated whether Dcp-1 regulates starvation-induced autophagy in the larval fat body using LysoTracker red (LTR), an acidotropic dye that labels both autolysosomes and lysosomes. After 1 h of starvation, fat bodies from larvae overexpressing Dcp-1 contained increased LTR and Atg8a puncta, whereas control larvae contained low LTR levels and diffuse Atg8a staining (Fig. S2, A and B). Fat bodies from 4-h-starved wild-type flies and Dcp-1Prev1 flies had high LTR and Atg8a puncta (Fig. S2, C and D), indicating that although Dcp-1 overexpression can enhance the autophagic response, Dcp-1 is not required for starvation-induced autophagy in the fat body.

Dcp-1 is required for the degradation of Ref(2)P after starvation

Ref(2)P, the Drosophila homologue of p62 (Nezis et al., 2008), is a substrate of autophagy and was shown to be a marker of autophagic activity (Nezis et al., 2010; Bartlett et al., 2011). Therefore, we used Ref(2)P analyses to confirm that Dcp-1 is a positive regulator of autophagic flux. Compared with fed wild-type flies (Fig. 3 A), nutrient-deprived flies had reduced Ref(2)P in follicle cells and nurse cells (Fig. 3 B), and Western blot analysis of wild-type ovaries revealed that the level of Ref(2)P was reduced after starvation (Fig. 3 C). Ovaries from fed or starved Atg77714 mutant flies showed an accumulation of Ref(2)P (Fig. 3, C and D). It was reported that starvation itself leads to transcriptional activation of Ref(2)P (Érdi et al., 2012), and this is likely why we see an even further increase in Ref(2)P levels after starvation in Atg7 flies. Compared with starved wild-type flies, degenerating mid-stage egg chambers from starved Dcp-1Prev1 flies contained increased levels of Ref(2)P (Fig. 3 E), and Western blot analysis of whole ovaries confirmed that Ref(2)P failed to be completely

Figure 2. Dcp-1 regulates autophagic flux in vitro. (A) Drosophila S2 cells stably expressing GFP-RFP-Atg8a showed an increase in the percentage of cells containing two or more autolysosomes after starvation, which was blocked after Bafilomycin A1 (BafA1) treatment. At least 50 cells were manually quantitated in three independent experiments (n = 3). Statistical significance was determined using one-way ANOVA with a Dunnett post test (**, P < 0.01). (B) Representative images of S2-GFP-RFP-Atg8a cells after the indicated treatments. (C) S2-GFP-RFP-Atg8a cells were treated with dsRNAs and subjected to fed or starvation conditions as indicated. At least 50 cells were manually quantitated in three independent experiments (n = 3). Statistical significance was determined using one-way ANOVA with a Dunnett post test (**, P < 0.01; ***, P < 0.001; ****, P < 0.0001). (D) Representative images of S2-GFP-RFP-Atg8a cells after the indicated RNAi treatments. Error bars represent the means ± SD. Bars, 10 µm.
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To elucidate the mechanism by which Dcp-1 mediates autophagic flux, we first determined the subcellular location of Dcp-1 in Drosophila l(2)mbn cells by immunostaining with an antibody to Dcp-1 (Fig. S3; Tenev et al., 2005). We observed colocalization between Dcp-1 and the mitochondrial markers ATP synthase-α (ATPsyn-α; Fig. 4 A) and MitoTracker red (Fig. 4 B) but did not observe colocalization between Dcp-1 and markers for the endoplasmic reticulum or cis-Golgi (Fig. S3, A and B). Dcp-1 was also found to colocalize with β-Tubulin (Fig. S3 C).

Dcp-1 localization to the mitochondria

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Figure 3. Ref(2)P analyses confirm that Dcp-1 is required for autophagic flux in ovaries. Staining shows DNA, Ref(2)P, and Armadillo. Zoomed insets show Ref(2)P staining. (A) Nondegenerating midstage egg chambers from fed w1118 flies showed Ref(2)P staining in follicle cells and nurse cells. (B) Starved w1118 flies contained degenerating midstage egg chambers with reduced Ref(2)P. (C) A representative Western blot of ovaries from w1118 and Atg7077/d14 flies subjected to fed or starvation conditions for 4 d. Ref(2)P was detected by immunoblotting. Actin served as a loading control. Densitometry was performed to quantify Ref(2)P protein levels relative to actin. Graph represents ± SD from five independent experiments (n = 5). Statistical significance was determined using a two-tailed Student’s t test. *, P = 0.004 for fed samples; **, P = 0.008 for starved samples. (D) Starved Atg7 flies showed an accumulation of Ref(2)P in the follicle cells and nurse cells of degenerating midstage egg chambers. (E) Starved Dcp-1Prev1 flies contained increased levels of Ref(2)P in degenerating midstage egg chambers. (F) A representative Western blot of ovaries from w1118 and Dcp-1Prev1 flies that were subjected to fed or starvation conditions for 4 d. Ref(2)P was detected by immunoblotting. Actin served as a loading control. Densitometry was performed to quantify Ref(2)P protein levels relative to actin. Graph represents ± SD from five independent experiments (n = 5). Statistical significance was determined using a two-tailed Student’s t test. *, P = 0.03. Bars: (main images) 25 µm; (zoomed images) 10 µm.

degraded after starvation (Fig. 3 F). The effect of Dcp-1 on Ref(2)P levels is modest compared with Atg7; we found that Ref(2)P accumulates only in degenerating midstage egg chambers of Dcp-1Prev1 flies after starvation. The variability in the levels of Ref(2)P, as shown in Fig. 3 F, is most likely caused by the extent of the persisting or “undead” egg chambers present in the samples. In contrast to Atg7 flies, Dcp-1 does not affect basal autophagy. Altogether, these data show that Dcp-1 is a positive regulator of starvation-induced autophagic flux during Drosophila mideogenesis.
found that the proform (zymogen) of Dcp-1 was located only in the membrane-enriched fraction in nutrient-rich and starvation conditions, whereas the processed p10 subunit of Dcp-1 was found in both the membrane and cytosolic fractions (Fig. 4 C). Similar results were obtained after subcellular fractionation of ovaries (Fig. 4 D). To determine whether Dcp-1 is located inside the mitochondria or is associated with the surface of the outer mitochondrial membrane, a proteinase K protection assay was performed with or without hypotonic disruption of the mitochondria. Mitochondrial proteins within the mitochondria are resistant to proteinase K treatment, whereas proteins located on the surface of the outer mitochondrial membrane, but not embedded within the lipid bilayer of mitochondria, are sensitive to proteinase K (Smith et al., 1994; Setoguchi et al., 2006). VDAC, an embedded outer mitochondrial membrane protein, and ATPsyn-α, a mitochondrial matrix protein, remained intact upon proteinase K treatment as expected but were digested by proteinase K after hypotonic disruption of the mitochondria (Fig. 4 E). In addition, Pink1, a mitochondrial protein associated with the outer mitochondrial membrane, was degraded after proteinase K treatment as expected (Fig. 4 F). Examination of Dcp-1 showed that it was digested only after hypotonic disruption of the mitochondria,
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of control cells in nutrient-rich media contained mitochondria with a normal morphology, and starvation led to an increase in the percentage of cells with elongated mitochondria (Fig. 5 B). Treatment with Dcp-1 RNAi resulted in a significant increase in cells that contained elongated mitochondria in both nutrient-rich and starvation media (Fig. 5 B). Given that the changes observed in mitochondrial morphology after Dcp-1 RNAi were not associated with changes in mitochondrial mass under the conditions tested (Fig. 4 G) suggests that Dcp-1 may play a role in maintaining the mitochondrial network under fed and starvation conditions.

To determine whether Dcp-1 alters mitochondrial morphology in vivo, we expressed mitochondrial targeting GFP (mitoGFP) in the germ line of Dcp-1 flies. All fed control flies contained short, tubular mitochondria that were dispersed throughout the nurse cell (Fig. 5 C) and were scored as healthy (Fig. 5 D, abbreviated as H). Starvation induces a series of mitochondrial events in degenerating midstage egg chambers, including mitochondrial remodeling and clustering, uptake by the follicle cells, and finally, degradation within the follicle cells (Fig. 4, E and F). We speculate that the proform of Dcp-1 translocates into the mitochondria where a fraction of it is processed into its active form, and this active form either remains in the mitochondria or translocates back into the cytosol. Together, these data demonstrate that pro–Dcp-1 localizes within mitochondria.

Figure 5. Loss of Dcp-1 promotes mitochondrial elongation. (A) l(2)mbn cells were labeled with the ATPsyn-α, and mitochondrial morphology was scored as fragmented, normal, or elongated. (B) Cells were treated with control or Dcp-1 dsRNA and subjected to nutrient-rich media or 1 h of starvation. Quantifications represent the percentage of cells with elongated mitochondria divided by the total number of cells examined. At least 100 cells were examined manually in three independent experiments (n = 3). Error bars represent the mean ± SD. Statistical significance was determined using one-way ANOVA with a Bonferroni post test (*, P < 0.05; **, P < 0.01). (C) Mitochondrial targeted GFP (mitoGFP) was expressed in the germline using the nosGAL4 driver. Staining shows mitoGFP, Armadillo, and DNA. (D) Mitochondria were scored as healthy (H), clustered (C), or elongated and overly connected (E). All of the mitochondria from fed UASp-mitoGFP/+;nosGAL4/+ flies were scored as healthy. n = 15 egg chambers manually scored. (E) mitoGFP was expressed in Dcp-1 flies using the nosGAL4 driver. (F) 54% of egg chambers from UASp-mitoGFP/+;Dcp-1/marm,Dcp-1/marm;nosGAL4/+ flies contained elongated mitochondria, 39% contained mitochondria that were scored as healthy, and 7% contained clustered mitochondria. n = 28 egg chambers manually scored. Bars: (A) 5 µm; (C and E, main images) 25 µm; (C and E, zoomed images) 10 µm.

indicating that Dcp-1 is internalized within the mitochondria (Fig. 4, E and F). We speculate that the proform of Dcp-1 translocates into the mitochondria where a fraction of it is processed into its active form, and this active form either remains in the mitochondria or translocates back into the cytosol. Together, these data demonstrate that pro–Dcp-1 localizes within mitochondria.

Loss of Dcp-1 alters morphology of the mitochondrial network

To examine a potential role of Dcp-1 at the mitochondria, we first analyzed whether loss of Dcp-1 alters mitochondrial mass. No changes in mitochondrial mass as detected by 10-N-nonyl acridine orange (NAO), a fluorescent dye that binds mitochondrial cardiolipin independently of mitochondrial membrane potential (Maftah et al., 1989), were detected in nutrient-rich or starvation conditions (Fig. 4 G). Next, we determined whether loss of Dcp-1 alters mitochondrial morphology. Mitochondria were labeled with ATPsyn-α and were scored as having a fragmented, a normal (containing both short and elongated mitochondria), or an elongated morphology (Fig. 5 A). The majority of control cells in nutrient-rich media contained mitochondria with a normal morphology, and starvation led to an increase in the percentage of cells with elongated mitochondria (Fig. 5 B). Treatment with Dcp-1 RNAi resulted in a significant increase in cells that contained elongated mitochondria in both nutrient-rich and starvation media (Fig. 5 B). Given that the changes observed in mitochondrial morphology after Dcp-1 RNAi were not associated with changes in mitochondrial mass under the conditions tested (Fig. 4 G) suggests that Dcp-1 may play a role in maintaining the mitochondrial network under fed and starvation conditions.

To determine whether Dcp-1 alters mitochondrial morphology in vivo, we expressed mitochondrial targeting GFP (mitoGFP) in the germ line of Dcp-1 flies. All fed control flies contained short, tubular mitochondria that were dispersed throughout the entire nurse cell (Fig. 5 C) and were scored as healthy (Fig. 5 D, abbreviated as H). Starvation induces a series of mitochondrial events in degenerating midstage egg chambers, including mitochondrial remodeling and clustering, uptake by the follicle cells, and finally, degradation within the follicle cells.
morphology even in the absence of a starvation signal, underscoring a novel nonapoptotic role of an effector caspase in regulating mitochondrial dynamics under both fed and starvation conditions.

High ATP levels in Dcp-1 flies suppress autophagic flux

Loss of Dcp-1 promotes elongation of the mitochondrial network, and mitochondrial elongation has been shown to sustain ATP levels after starvation to promote cell survival (Gomes et al., 2011). In addition, the cellular energy sensor AMP-activated protein kinase is activated when the ratio of ATP/AMP falls, for example, during periods of starvation (Salt et al., 1998; Gleason et al., 2007), thus stimulating energy producing pathways such as autophagy (Kim et al., 2011). Therefore, we determined whether Dcp-1 also alters ATP levels as a mechanism to induce autophagic flux. Cells treated with Dcp-1 RNAi showed a significant increase in ATP levels within 4 h of starvation (Fig. 6 A), suggesting (Tanner et al., 2011). Fed Dcp-1 flies expressing mitoGFP in the germline contained mitochondria with an altered morphology even in the absence of starvation. 54% of midstage egg chambers from Dcp-1 flies contained mitochondria that were elongated and overly connected (Fig. 5, E and F, abbreviated as E), whereas only 39% of midstage egg chambers contained mitochondria that were healthy (Fig. 5 F, abbreviated as H). To control for the different genetic backgrounds between wild-type and Dcp-1 flies with respect to the mitochondrial phenotypes, Dcp-1 was overexpressed in the germline of Dcp-1 flies to determine whether the mitochondrial phenotype could be rescued. Indeed, 94% of midstage egg chambers from Dcp-1 flies overexpressing wild-type Dcp-1 showed healthy mitochondria compared with only 33% of midstage egg chambers from Dcp-1 flies (Fig. S4, A–C), indicating that the mitochondrial phenotype is caused by the Dcp-1 mutation. These data demonstrate that Dcp-1 plays a role in the regulation of mitochondrial network morphology even in the absence of a starvation signal, underscoring a novel nonapoptotic role of an effector caspase in regulating mitochondrial dynamics under both fed and starvation conditions.

Figure 6. Dcp-1 alters ATP levels that in turn regulate autophagy. (A) Total cellular ATP levels were measured in l(2)mbn cells treated with control or Dcp-1 dsRNA. Data represent ± SEM of three independent experiments (n = 3). Statistical significance was determined using a two-tailed Student’s t test (*, P = 0.02). (B) Total cellular ATP levels were measured in ovaries from fed or starved w1118 and Dcp-1 flies. Data represent ± SEM of five independent experiments (n = 5). Statistical significance was determined using a two-tailed Student’s t test (*, P = 0.014; ***, P < 0.001). (C) Fed or starved w1118 and Dcp-1 flies were treated with DMSO or 25 µg/ml oligomycin A, and Ref(2)P levels were assessed by immunoblot analysis. Actin served as a loading control. Densitometry was performed to quantify protein levels relative to actin. Graph represents ± SD from three independent experiments (n = 3). Statistical significance was determined using a two-tailed Student’s t test (*, P = 0.02). (D and E) Control UASP-mCherry-GFP-DrAtg8a/+;nosGAL4/+ flies (D) and Dcp-1/Dcp-1;UASP-mCherry-GFP-DrAtg8a/nosGAL4/+ flies (E) were subjected to starvation conditions supplemented with DMSO or 25 µg/ml oligomycin A. Bars, 25 µm. Quantifications show percentage of autolysosomes (autolysosomes/total autophagic structures). At least eight egg chambers were manually quantitated per genotype per condition (n = 8). Statistical testing was determined using a two-tailed Student’s t test. ***, P = 0.0002.
that at least under starvation in vitro, the elongated mitochondrial phenotype in Dcp-1 RNAi-treated cells is associated with increased ATP levels. To confirm this in vivo, we examined ovaries from wild-type and Dcp-1<sup>Prev1</sup> flies subjected to fed or starvation conditions and found ATP levels to be significantly increased in ovaries of Dcp-1<sup>Prev1</sup> flies under both conditions tested (Fig. 6 B). We reasoned that the increased ATP levels in Dcp-1<sup>Prev1</sup> flies may inhibit autophagic flux after starvation. Reduction of ATP with oligomycin A, an inhibitor of the mitochondrial ATP synthase, induces autophagy in the IPLB-LdFB insect cell line (Tettamanti et al., 2006), and so, we tested whether oligomycin A would induce autophagy in Drosophila ovaries. Although oligomycin A did not alter Ref(2)P levels under fed conditions, ovaries from starved Dcp-1<sup>Prev1</sup> flies treated with oligomycin A showed a significant reduction in Ref(2)P protein compared with those treated with DMSO (Fig. 6 C). In addition, starved oligomycin A–treated Dcp-1<sup>Prev1</sup> flies expressing GFP-mCherry-DrAtg8a in the germ-line showed an increase in the percentage of autolysosomes in degenerating midstage egg chambers compared with those treated with DMSO (red puncta; Fig. 6 E), whereas we observed no additional increase in the percentage of autolysosomes in starved control flies treated with oligomycin A (Fig. 6 D). This suggests that high ATP levels are sufficient to block autophagic flux in Dcp-1<sup>Prev1</sup> flies after starvation. Collectively, our data demonstrate that Dcp-1 controls autophagic flux by a mechanism involving the regulation of the mitochondrial network and maintenance of ATP levels.

**Dcp-1 negatively regulates the levels of the ANT SesB**

We next determined the molecular mechanism by which Dcp-1 could regulate ATP levels to induce autophagy. *Drosophila SesB* encodes an ANT that functions to exchange ATP for ADP across the mitochondrial inner membrane (Terhzaz et al., 2010). RNAi of SesB in malpighian tubules significantly decreased ATP levels and altered mitochondrial morphology from a threadlike appearance to a short and globular morphology (Terhzaz et al., 2010). Treatment of l(2)mbn cells with Dcp-1 RNAi resulted in a significant increase in the levels of SesB protein under both fed and starvation conditions (Fig. 7 A). This observed increase in SesB protein was not caused by an increase in mitochondrial mass as measured by NAO (Fig. 4 G) nor could it be fully explained by alterations in SesB transcript levels (Fig. S5 A), which were unaltered in fed conditions, increased 12.4-fold after 2 h of starvation, and returned to levels comparable with fed conditions by 4 h of starvation. To determine whether the effect on SesB protein was also observed in vivo, we examined ovaries from fed and nutrient-deprived wild-type and Dcp-1<sup>Prev1</sup> flies. We observed decreased SesB levels after starvation in wild-type flies (Fig. 7 B), whereas ovaries from fed or nutrient-deprived Dcp-1<sup>Prev1</sup> flies contained increased SesB levels, indicating that Dcp-1 may be required to negatively regulate the levels of SesB and, thus, ATP. We also observed a significant increase in ATPsyn-α levels under fed and 2 h of starvation in vitro (Fig. 7 A) but did not detect a significant change in ATPsyn-α levels in vivo (Fig. 7 B). Given the consistent changes observed in SesB, we next tested whether SesB is a direct target of Dcp-1’s proteolytic activity using in vitro cleavage assays. Catalytically active Dcp-1 (Dcp-1<sup>FL</sup>) and inactive Dcp-1 (Dcp-1<sup>CαCα</sup>) were expressed in l(2)mbn cells and purified by Ni<sup>2+</sup>-nitrilotriacetic acid (NTA) columns. Only Dcp-1<sup>FL</sup> was able to cleave in vitro translated Drice (Fig. 7 C), another *Drosophila* effector caspase and known substrate of Dcp-1 (Song et al., 2000). Treatment of in vitro translated SesB with Dcp-1<sup>FL</sup> or Dcp-1<sup>CαCα</sup> failed to induce cleavage of SesB (Fig. 7 C), indicating that Dcp-1 does not directly cleave SesB but rather affects its stability. To test whether Dcp-1 directly interacts with SesB as a mechanism to regulate its stability, an immunopurification (immunoprecipitation [IP]) and tandem mass spectrometry (MS; MS/MS) fragmentation assay were performed. To prevent the proteolytic cleavage of Dcp-1’s substrates and allow for their identification, we overexpressed catalytically inactive V5-tagged Dcp-1<sup>CαCα</sup> in l(2)mbn cells and subjected cells to nutrient-full medium or 2 h of starvation. Dcp-1 was immunoprecipitated with an anti-V5 antibody (Fig. 7 D), and immunoprecipitates were analyzed by liquid chromatography–MS/MS for protein detection and identification. Four unique peptides corresponding to SesB were identified in the Dcp-1 immunoprecipitate, but not in the vector control, in two independent replicates, indicating that SesB is a binding partner of Dcp-1 (Table S2). To confirm the interaction between Dcp-1 and SesB, we expressed FLAG-tagged SesB in l(2)mbn cells and found that FLAG-SesB immunoprecipitated endogenous pro–Dcp-1 (Fig. 7 E). Notably, no processed Dcp-1 was detected (Fig. 7 E), and we found that FLAG-SesB was mitochondrial localized (Fig. S5 B), further indicating that the interaction between Dcp-1 and SesB takes place in the mitochondria and occurs in a nonproteolytic manner. All together, these data show that pro–Dcp-1 interacts with SesB and affects its stability in a nonproteolytic manner.

**SesB is a negative regulator of autophagic flux and interacts genetically with Dcp-1**

The loss of SesB after starvation suggests that SesB may negatively regulate autophagy. To test this, we examined ovaries from SesB hypomorphic flies (SesB<sup>Org</sup>) that were shown to have reduced ANT activity (Rikhy et al., 2003). Consistent with this previous study, ovaries from SesB<sup>Org</sup> flies had reduced ATP levels compared with ovaries from wild-type flies (Fig. S5 C). Although fed wild-type flies rarely contain degenerating midstage egg chambers (Table S1), fed SesB<sup>Org</sup> flies contained an increase in degenerating midstage egg chambers (Table S1) that had increased LTR staining (Fig. S5 F) and reduced Ref(2)P (Fig. 8, A and B). Loss of SesB resulted in a similar phenotype in the larval fat body (Fig. S2). In addition, overexpression of GFP-mCherry-DrAtg8a in the germ-line of fed SesB<sup>Org</sup> flies revealed an increase in the percentage of autolysosomes in both nondegenerating (Fig. 8, C [left] and D) and degenerating (Fig. 8, C, right) midstage egg chambers, confirming that there is increased autophagic flux in SesB<sup>Org</sup> flies. Degenerating midstage egg chambers from SesB<sup>Org</sup> flies also stained positively for TUNEL (Fig. 8 E), suggesting that SesB is normally required to suppress autophagic flux and DNA fragmentation during *Drosophila* oogenesis.

To determine the epistatic relationship between Dcp-1 and SesB, we examined fed and nutrient-deprived SesB<sup>Org</sup>; Dcp-1<sup>Prev1</sup> double mutant flies. Because of the high levels of SesB in
Our results reveal that starvation-induced autophagic flux occurs in both midstage egg chambers that have not entered the degeneration process as well as in those that are undergoing cell death. Furthermore, we find that the effector caspase Dcp-1 is required for autophagic flux in degenerating midstage egg chambers in addition to its role in cell death. We find that one mechanism of Dcp-1–induced autophagic flux is mediated through SesB. In humans, there are four mitochondrial ANT isoforms, each with a tissue-specific distribution and different roles in apoptosis. ANT1 and ANT3 were proposed to be proapoptotic, whereas ANT2 and ANT4 were shown to be antiapoptotic (Brenner et al., 2011). However, the roles of mammalian ANT proteins in autophagy have yet to be characterized. Our data show that reduced Dcp-1

Discussion

Our results reveal that starvation-induced autophagic flux occurs in both midstage egg chambers that have not entered the degeneration process as well as in those that are undergoing cell death. Furthermore, we find that the effector caspase Dcp-1 is required for autophagic flux in degenerating midstage egg chambers in addition to its role in cell death. We find that one mechanism of Dcp-1–induced autophagic flux is mediated through SesB. In humans, there are four mitochondrial ANT isoforms, each with a tissue-specific distribution and different roles in apoptosis. ANT1 and ANT3 were proposed to be proapoptotic, whereas ANT2 and ANT4 were shown to be antiapoptotic (Brenner et al., 2011). However, the roles of mammalian ANT proteins in autophagy have yet to be characterized. Our data show that reduced
Figure 8. *SesB* hypomorphic flies have an enhanced autophagy phenotype, and epistasis analysis indicates that *SesB* acts downstream of Dcp-1. (A) Ovaries from fed *SesB*<sup>Org</sup> flies show reduced Ref(2)P staining in degenerating midstage egg chamber (arrow). (B) Ovaries from fed *w<sup>1118</sup>* and *SesB*<sup>Org</sup> were analyzed by Western blotting using Ref(2)P. Actin served as a loading control. Results are representative of three independent experiments (n = 3). (C) *SesB*<sup>Org</sup> flies expressing GFP-mCherry-DrAtg8a in the germline showed increased autophagosomes and autolysosomes in nondegenerating (left) and degenerating (right) midstage egg chambers. Insets show autophagosomes and autolysosomes (left) and autolysosomes (right) in *SesB*<sup>Org</sup> midstage egg chambers. (D) Quantitation of percentages of autolysosomes (autolysosomes/total autophagic structures) in nondegenerating ovaries from fed *w<sup>1118</sup>* and *SesB*<sup>Org</sup> flies. 10 egg chambers were manually examined per genotype. Error bars represent the means ± SD. Statistical significance was determined using a two-tailed Student’s *t* test (***, *P* < 0.0001). (E) Degenerating midstage egg chambers (arrows) from fed *SesB*<sup>Org</sup> flies are TUNEL positive. (F) Degenerating midstage egg chambers from starved control *SesB*<sup>Org</sup>;*Dcp-1<sup>Prev1</sup>* flies contained persisting nurse cell nuclei and increased Ref(2)P. n = 117 ovarioles examined. Box represents zoomed image. (G) Degenerating midstage egg chambers from fed *SesB*<sup>Org</sup>;*Dcp-1<sup>Prev1</sup>* flies contained condensed and fragmented nurse cell nuclei and low levels of Ref(2)P. n = 156 ovarioles examined. (H) In response to starvation, Dcp-1 acts to negatively regulate the levels of SesB. SesB is a negative regulator of autophagy, and reduced SesB levels mediated by Dcp-1 lead to decreased ATP and induction of autophagic flux. Bars: (A, C [main images], E, F [main images], and G [main images]) 25 µm; (C, insets) 12.5 µm; (F and G [zoomed images]) 10 µm.
Dcp-1 leads to increased levels of SesB protein in fed and starvation conditions during *Drosophila* oogenesis and in *Drosophila* cultured cells. We detected no significant change in SesB transcript levels in fed conditions or after 4 h of starvation, but a significant increase was observed in cells after 2 h of starvation. This finding suggests that a transcription-related mechanism may play some role in the observed cellular response but is not sufficient to account for all of the observed changes in protein levels. Although Dcp-1 does not cleave SesB, the proform of Dcp-1 interacts with SesB, and we predict that this interaction regulates the stability of SesB. We find that SesB is required to suppress autophagic flux during midoogenesis even under nutrient-rich conditions, and reduction of SesB in *Dcp-1*^prov^ flies rescues the autophagic defect after starvation. This is the first study to our knowledge showing that an ANT functions as a negative regulator of autophagy.

The *Drosophila* genome encodes seven caspases, and to date, only the initiator caspase Dronc and the effector caspase Drice have been shown to localize to the mitochondria (Dorstyn et al., 2002). In mammalian cells, caspases have been detected at the mitochondria during apoptosis (Krajewski et al., 1999; Susin et al., 1999; Chandra and Tang, 2003); however, the role of caspases at the mitochondria, especially under nonapoptotic conditions, is poorly understood. Our results demonstrate that Dcp-1 localizes to the mitochondria where it functions to maintain the mitochondrial network morphology. Under nutrient-rich conditions, nondegenerating midstage egg chambers from *Dcp-1*^prov^ flies contained mitochondria that appeared elongated and overly connected, and ovaries contained increased ATP levels, indicating that Dcp-1 normally functions to negatively regulate mitochondrial dynamics and ATP levels. Consistent with our findings, overexpression of the caspase inhibitor p35 in the amnioserosa suppressed the transition of mitochondria from a tubular to a fragmented state during delamination (Muliyyil et al., 2011), further suggesting that inhibition of caspases hinders normal mitochondrial dynamics.

Dcp-1 acts to finely tune the apoptotic process, and cell death only occurs when caspase activity reaches a certain apoptotic threshold (Florentin and Arama, 2012). Effector caspases involved in nonapoptotic processes may be restricted in time or space to regulate caspase activity (Kaplan et al., 2010; Florentin and Arama, 2012). As Dcp-1 functions not only in autophagy and apoptosis but also at the mitochondria to regulate mitochondrial morphology and ATP levels, one question that remains is how the activity of Dcp-1 is regulated. As Dcp-1 has autocatalytic activity (Song et al., 1997), perhaps Dcp-1 is sequestered in mitochondria to prevent its full activation. Mitochondrial localized mammalian pro–Caspase 3 and 9 are S-nitrosylated in mitochondria to prevent its full activation. Mitochondrial localized mammalian pro–Caspase 3 and 9 are S-nitrosylated in their catalytic active site, leading to the inhibition of their activity (Mannick et al., 2001). Perhaps mitochondrial Dcp-1 is also S-nitrosylated, serving to limit Dcp-1’s activity. In addition, mammalian Hsp60 and Hsp10 were shown to interact with mitochondrial localized pro–Caspase 3 in which they function to accelerate pro–Caspase 3 activation after the induction of apoptosis (Samali et al., 1999). Perhaps Dcp-1 associates with *Drosophila* Hsp60 or Hsp10 in the mitochondria to regulate its mitochondrial related functions. However, further studies are required to identify upstream regulators of Dcp-1 that regulate its mitochondrial, autophagic, and apoptotic functions.

Effector caspases are the main executioners of apoptotic cell death; however, it is becoming increasingly evident that caspases have nonapoptotic functions in differentiation, proliferation, cytokine production, and cell survival (Kuranova and Miura, 2007; Galluzzi et al., 2012). For example, Caspase 3 was shown to regulate tumor cell repopulation in vitro and in vivo (Huang et al., 2011), and it was also shown to be required for skeletal muscle (Fernando et al., 2002) and macrophage differentiation (Sordet et al., 2002). In *Drosophila*, the initiator caspase Dronc maintains neural stem cell homeostasis by binding to Numb in a noncatalytic, nonapoptotic manner to regulate its activity (Ouyang et al., 2011). In addition, Dcp-1 is required for neuro-muscular degeneration in a nonapoptotic manner (Keller et al., 2011). Our results show that Dcp-1 also has a nonapoptotic role during oogenesis, in which it is required to maintain mitochondrial physiology under basal conditions. Loss of Dcp-1 alters this physiology, leading to increased SesB and ATP levels that in part prevent the induction of autophagic flux after starvation. These data support the notion that caspases play a much more diverse role than previously known and that the underlying mechanisms should be better understood to appreciate the full impact of apoptosis pathway modulation for treatment in human pathologies.

**Materials and methods**

**Fly strains**

*Prev1*^w^ was used at the wild-type control strain in this study. Other fly strains used are as follows: *nos-GAL4::VP16* (abbreviated as *nosGAL4*), a driver containing the Gal4/VP16 transactivator under the control of the *nos* promoter; Reth, 1998), *UASp-mitoGFP* (*expresses* the COX VIII mitochondrial targeting sequence fused to EGFP under the *UASp* promoter; Cox and Spradling, 2003), *Dcp-1*^prov^ (contains a 40-bp partial P element insertion in the coding region of *Dcp-1*, resulting in a frame shift in *Dcp-1* and an in-frame stop codon within the 40-bp insertion; Laundrie et al., 2003), and *UASp-FL-Dcp-1* (drives expression of full-length Dcp-1 under the *UASp* promoter; Peterson et al., 2003) were gifts from K. McColl (Boston University, Boston, MA). *Atg7*^mut^ (*P* element excision removing *CG3335* and exons 5 and 6 and most of exon 4 from *Atg7*) and *Atg7*^mut^ (*P* element excision removing the translation and transcription start sites and the majority of the *Atg7* coding region) were gifts from T. Neufeld (University of Minnesota, Minneapolis MN; Juhász et al., 2007). *UASp-GFP-mCherry-DrAtg8a* (drives expression of GFP-mCherry-DrAtg8a under the *UASp* promoter) was a gift from T.E. Rusten (Centre for Cancer Biomedicine, Oslo University Hospital, Montebello, Oslo, Norway; Nezis et al., 2010), and *SesB* multiphosphorylated flies, which contain a glutamate to lysine change at amino acid 266 that disrupts normal SesB activity, were obtained from the Bloomington Stock Center (stock 21590).

**Cell culture conditions**

*Drosophila* l(2)mbn cells were grown in Schneider’s medium [Invitrogen] supplemented with 10% FBS in 25-cm² suspension cell flasks (Sarstedt) at 25°C. *Drosophila* S2-RFP-GFP-Atg8a cells were grown in ESF921 medium (Expression Systems) in the presence of 50 μg/ml Zeocin. All experiments were performed 3–4 d after passage.

**Construction of RFP-GFP-ATG8a plasmid and creation of S2-RFP-GFP-Atg8a stable cell line**

ATG8a was amplified using the primers 5’-TAGCCTAGCTAGATCTACTAAGTCCAAATACAAAGGAGGA-3’ and 5’-TGGCCTAGCTGGAATGTTATGTTTGGGCCCAGGCC-3’. Atg8a was cloned into the BglII–EcoRI sites of mRFP-GFP-LC3 (Kimura et al., 2007). The cytomegalovirus promoter from *Drosophila S2-RFP-GFP-Atg8a* were grown in SF921 medium (Expression Systems) in the presence of 50 μg/ml Zeocin. All experiments were performed 3–4 d after passage.

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Center) using the primers 5′-TGACAGATGCGATATACGAGTAACTTC-TATATGTC-3′ and 5′-TGACAGATGCGCTAGTGATAGATTTGTC-3′. The P<sub>τ</sub>-RFP-GFP-ATG6a and P<sub>τ</sub>-RFP-GFP-ATG8a were added to 10 µl Celfi invented (Invitrogen) plus 100 µl Grace’s medium (Invitrogen) and incubated for ≥30 min. 3.75 × 10<sup>5</sup> S2 cells in 400 µl Grace’s media were incubated with transfection medium overnight. 1 ml ES9F21 medium (Expression Systems) was added back to the cells, and the cells were incubated for an additional 3 d before adding Zeocin at a concentration of 0.6–0.8 mg/ml. The media were replaced every 4 d until the negative control showed 0% viability (~3 wk). Transfected cells were maintained in ES9F21 + 0.1 mg/ml Zeocin.

Double-stranded RNA (dsRNA) synthesis

Each PCR primer for RTPCR was designed to contain a 5′ T7 RNA polymerase binding site (5′-TAATACGACTCACTATAGGG-3′) followed by sequences specific for the target gene. The ampicillin resistance gene was used as a control dsRNA and was amplified using the primers 5′-TAATACGACTCACTATAGGG-3′ and 5′-TAATACGACTCACTATAGGG-3′. The PCR products were generated by RTPCR using SuperScript One-Step RTPCR with Platinum Taq (Invitrogen). RTPCR products were ethanol precipitated and used as a template for in vitro transcription reactions using T7 Ribomax Express RNAi System (Promega). Quality of the RNA was analyzed by gel electrophoresis. dsRNA was quantitated using PicoGreen and adjusted to 200–400 ng/µl with nuclelease-free water.

RNAi

For RNAi, cells were washed and resuspended in ES9F21 medium to a concentration of 2 × 10<sup>6</sup> cells/ml. 333 µl of adjusted cells was plated in each well of a 24-well plate. 5–10 µg dsRNA was added per well and incubated at 25°C for 1 h. After incubation, 667 µl Schneider’s + 10% FBS was added back to each well and was incubated for an additional 72 h at 25°C.

Immunofluorescence experiments

Flies were conditioned on wet yeast paste for 2 d (fed) and then transferred to a vial containing 10% sucrose for 4 d for amino acid starvation. Ovaries were dissected in PBS and fixed with 4% paraformaldehyde. Ovaries were washed with PBS (PBS + 0.3% Triton X-100), permeabilized with 0.5% Triton X-100, and blocked with 2% BSA in PBS. For in vitro experiments, 200 µl cells were plated into an 8-well CC2-coated chamber slide and incubated for ≥30 min to let the cells adhere. For starvation treatments, media were replaced with 2 mg/ml glucose/PBS. For MitoTracker experiments, 500 µl MitoTracker red (Invitrogen) was added to each well and was incubated for 30 min in the dark. Media were removed and fixed with 4% paraformaldehyde for 20 min. Cells were washed with PBS/T and fixed with 0.5% Triton X-100.

Primary antibodies included rabbit anti-Ref(2)P [1:10,000], mouse anti-actin [1:500; JLA20; Developmental Studies Hybridoma Bank], mouse anti-actin (1:1,000; E7), mouse anti–ATP synthase (1:1,000), mouse anti–Porin (1:1,000; MitoSciences), guinea pig anti–Dcp-1 (1:500; Tenev et al., 2005), guinea pig anti–Dcp-1 (1:500; Tenev et al., 2005), rabbit anti-Porin (1:1,000; MitoSciences), mouse anti-ANT (1:500; MitoSciences), rabbit anti-Dcp-1 [1:500; Laundrie et al., 2003], guinea pig anti–Dcp-1 [1:500; Tenev et al., 2005], rabbit anti-Porin (1:1,000; EB), rabbit anti-Doc-1 [1:1,000; Barth et al., 2011]. Membranes were incubated with HRP-conjugated secondary antibodies or infrared-labeled secondary antibodies and were detected using the ECL Enhanced Western Blotting System (GE Healthcare) or the Odyssey Western Blotting System (LI-COR Biosciences). Densitometry was performed using ImageQuant 5.2 software (GE Healthcare).

Isolation of crude mitochondrial and cytosolic fractions

Approximately 5 × 10<sup>6</sup> cells were collected by centrifugation at 800 rpm for 10 min, resuspended in cold SEM-10 [10 mM MOPS, pH 7.5, 320 mM sucrose, and 1 mM EDTA with complete protease inhibitor cocktail (Roche)], and ground using a dounce homogenizer. Lysates were centrifuged twice at 3,000 rpm to remove cell debris, and supernatants were collected. The membrane fraction (pellet) was separated from the cytosolic fraction (supernatant) by centrifugation at 12,000 rpm. The pellets were washed once with 500 µl SEMP and finally resuspended in 50 µl SEMP. Protein concentrations were determined using the Bicinchoninic Acid Protein Assay. Aliquots of 50 µg protein were either spun down (membrane fraction) or TCA precipitated (cytosolic fraction) and analyzed by Western blotting.

Proteinase K protection assay

For mitochondrial lysis, mitochondria were resuspended in the appropriate amount of swelling buffer (50 mM Tris, pH 7.5) for 30 min on ice with vortexing. Intact and lysed mitochondria were incubated in 20 µg/ml proteinase K (Roche). Proteinase K was inactivated by adding PMSF (Fluka) to a final concentration of 1 mM. Intact mitochondria were spun down at 4°C, and 12,000 rpm for 20 min while lysed samples were TCA precipitated.

Oligomycin A treatment

3–5-d-old flies were transferred to a vial containing wet yeast paste supplemented with 200 µl of 25-g/ml Oligomycin A or DSMO added directly to the top of the yeast paste. After 2 d of treatment, flies were transferred to a vial containing a Kimwipes soaked with 10% sucrose supplemented with 25 µg/ml Oligomycin A or DSMO for 4 d.
Mitochondrial mass analysis and determination of ATP levels
RNA-treated cells were transferred from Schneider + 10% FBS to 2 mg/ml glucose/PBS starvation media plus 10 μg dsRNA for 1–24 h. For mitochondrial mass measurements, cells were incubated with 10 μM NAO for 10 min at 25°C in the dark. Cells were resuspended and put on ice to be analyzed by flow cytometry (FACSCalibur; BD). A minimum of 30,000 cells was acquired for triplicate samples per experiment. The mean fluorescence was analyzed using FlowJo software version 7.2.6 (Tree Star, Inc.).

ATP levels from I2/mim cells or Drosohila ovaries were measured using the ATP Determination Kit (Invitrogen) according to the manufacturer’s recommendations. Luminescence was measured using a plate reader (Wallac 1420 Victor; PerkinElmer).

Generation of constructs and in vitro synthesis
Drice (Tenev et al., 2005) was PCR amplified using primers containing Arfβ1 and Arfβ2 sequences. SesB was amplified from a full-length cDNA construct (Drosophila Genomics Resource Center) using primers containing Arfβ1 and Arfβ2 sequences. PCR products were cloned into the pDONR221 Gateway Entry Vector (Life Technologies) and were sequenced. The entry clones were then shuttled into the pEX2-DEST and pEX1-DEST expression vectors, respectively, for in vitro translation experiments. Sequences used for plasmid construction can be found in Table S1. SesB and Drice were synthesized using the Expressway Mini Expression Module (Invitrogen) according to the manufacturer’s instructions. 1 μg of template DNA was used for every 100 μl of translation reaction in a 4 h reaction. PCR primers were used as follows: Drice Arfβ1, 5′-GGGACAAGTTTGTACAAAAAAGC-3′; Drice Arfβ2, 5′-GGGGACCACTTTGTACAAGAAAGCTGGGTCTCAAACCCGTCCG-3′; SesB Arfβ1, 5′-GGGACAAGTTTGTACAAAAAAGCCTGTTGACAAAGGTCGCTTCTTTCCG-3′; and SesB Arfβ2, 5′-GGGGACCACTTTGTACAAAAAAGCCTGTTGACAAAGGTCGCTTCTTTCCG-3′.

Transfection and purification of Dcp-1 and in vitro cleavage assays
For transfection experiments, 0.5 μg His-V5–Dcp-1His or His-V5–Dcp-1Caa plasmid DNA (Tenev et al., 2005) was added to 10 μl Cellfectin plus 100 μl Grace’s medium and incubated for ≥30 min. 3.75 × 10⁴ cells in 400 μl Grace’s medium were incubated with transfection medium overnight before receiving 1 ml Schneider’s medium + 10% FBS. Cells were incubated for an additional 3 d before Ni-NTA purification. Purification of His-V5–Dcp-1His or His-V5–Dcp-1Caa was performed using Ni-NTA spin columns (HisPur; Thermo Fisher Scientific). Cells were resuspended in 400 μl equilibration buffer (PBS and 10 mM imidazole) with 1% Triton X-100, incubated in a rotary shaker at 4°C for 10 min, and centrifuged at 15,000 rpm at 4°C for 15 min to remove insoluble material. Subsequent steps were performed according to the manufacturer’s protocol. Eluted Dcp-1His and Dcp-1Caa were checked for caspase activity using the Caspase-Glo 3/7 Assay kit (Promega) according to the manufacturer’s instructions. The solutions were immediately used for in vitro cleavage assays. The caspase reaction buffer used in this experiment was as previously described (Tenev et al., 2005) and consisted of 20 mM Tris pH 7.5, 150 mM NaCl, 2 mM DTT, and 0.1% Triton, X-100. A 100 μl reaction volume was set up consisting of increasing volumes of Ni-NTA-purified Dcp-1His or Dcp-1Caa [10, 20, and 40 μl] and 5 μl of in vitro translated SesB or Drice. The reaction mixture was incubated at 25°C overnight and precipitated with 400 μl acetone for Western blot analysis.

Quantitative RT-PCR (QRT-PCR) analysis
RNA-treated cells were subjected to nutrient-full or starvation conditions for 2 or 4 h and were collected and pelleted at 850 rpm for 5 min. Cell pellets were lysed in 1 ml TRIZOL (Invitrogen), and total RNA was extracted according to the manufacturer’s instructions. The elutions were immediately used for in vitro translation experiments. The reaction mixture was incubated at 25°C overnight and precipitated with 400 μl acetone for Western blot analysis.

Statistics
In each graph, data represent ± SEM or SD of n independent experiments. As indicated in the legends, statistical significance was calculated by analysis of variance (ANOVA) plus a Dunnett or Bonferroni post test, or a two-tailed Student’s t-test between the indicated samples was used. P-values are shown in the legends.

Online supplemental material
Fig. S1 shows that Dcp-1His filae have reduced autolyososomes in nondegenerating midstage egg chambers after starvation. Fig. S2 shows LTR and Atg8a antibody staining in fat bodies of Dcp-1His/C, GAL4A/+; UASdcp-1His/C, SesB or control, and control flies. Fig. S3 shows a lack of colocalization between Dcp-1 and the ER or Golgi but does show localization between Dcp-1 and βTubulin. Fig. S4 shows that overexpression of Dcp-1 can rescue the Dcp-1null mitochondrial phenotype. Fig. S5 shows SesB QRT-PCR analysis of the LTR. FLAG-SesB localization to mitochondria, and that ovaries from SesB+ flies have reduced ATP and contain increased LTR staining in degrading midstage egg chambers. Table S1 shows quantification of TUNEL-positive germaria and midstage egg chambers. Table S2 shows identification of SesB by MS. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201303144/DC1.

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