

# Acinus integrates AKT1 and subapoptotic caspase activities to regulate basal autophagy

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How cellular stresses up-regulate autophagy is not fully understood. One potential regulator is the *Drosophila melanogaster* protein Acinus (Acn), which is necessary for autophagy induction and triggers excess autophagy when overexpressed. We show that cell type-specific regulation of Acn depends on proteolysis by the caspase Dcp-1. Basal Dcp-1 activity in developing photoreceptors is sufficient for this cleavage without a need for apoptosis to elevate caspase activity. On the other hand, Acn was stabilized by loss of Dcp-1 function or by the presence of a mutation in Acn that eliminates its

conserved caspase cleavage site. Acn stability also was regulated by AKT1-mediated phosphorylation. Flies that expressed stabilized forms of Acn, either the phosphomimetic Acn<sup>S641,731D</sup> or the caspase-resistant Acn<sup>D527A</sup>, exhibited enhanced basal autophagy. Physiologically, these flies showed improvements in processes known to be autophagy dependent, including increased starvation resistance, reduced Huntingtin-induced neurodegeneration, and prolonged life span. These data indicate that AKT1 and caspase-dependent regulation of Acn stability adjusts basal autophagy levels.

## Introduction

Two modes of macroautophagy, referred to as autophagy from here on, support cell survival by alleviating distinct problems. A first predominantly catabolic mode of autophagy enables acutely starving cells to nonspecifically degrade cellular content and reclaim necessary building blocks. The lysosome-localized mammalian target of rapamycin (TOR; mTOR) signaling complex plays a major role in the regulation of this mode of autophagy (Settembre et al., 2012; Yan and Lamb, 2012; Efeyan et al., 2013; Jewell et al., 2013). A second mode, sometimes called quality control autophagy, appears to operate at low levels in most cells to eliminate deleterious cellular content and debris, ranging in size from invading pathogens and dysfunctional mitochondria to misfolded proteins (Kroemer et al., 2010). Both of these modes use the core autophagy proteins to initiate the generation of isolation membranes (also known as phagophores), promote their growth to autophagosomes, and finally promote their fusion with lysosomes to initiate degradation of the captured content (Mizushima et al., 2011).

Studies of starvation-induced autophagy in yeast and in the context of tumorigenesis led to the discovery of most of the core autophagy proteins (Klionsky and Ohsumi, 1999; Liang

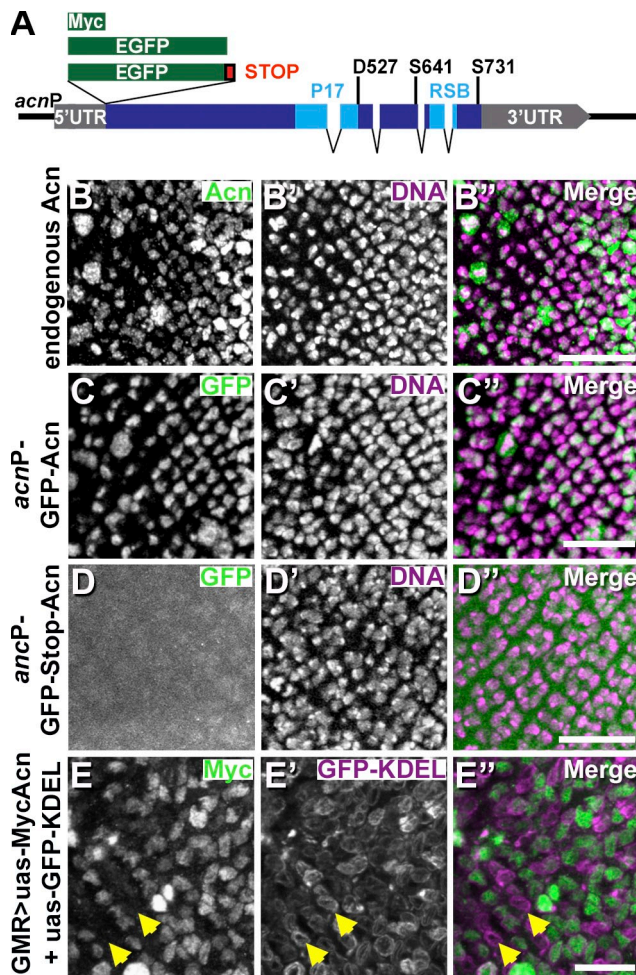
et al., 1999). Since then, significant progress has been made in understanding the homeostatic regulation of autophagy in response to nutrient deprivation (Yang and Klionsky, 2010; White, 2012). Less well understood, however, is the regulation of basal autophagy levels. The importance of the quality control function of basal autophagy was highlighted by autophagy-deficient mice and flies, which undergo neurodegeneration (Hara et al., 2006; Komatsu et al., 2006; Juhász and Neufeld, 2008) and cardiac hypertrophy (Nakai et al., 2007). Clearance of aggregation-prone proteins is a key aspect of autophagy in preventing disease and extending life span (Meléndez et al., 2003; Vellai et al., 2009; Demontis and Perrimon, 2010; Kroemer et al., 2010).

Basal autophagy and life span can be enhanced by increased expression of autophagy genes (Simonsen et al., 2008). For their acute up-regulation, lysosome-bound TFEB (transcription factor EB) plays an important role in transcriptional responses to metabolic changes through its mTOR-regulated nuclear relocalization (Settembre et al., 2012). Long-term adjustment of basal autophagy relies on multiple factors regulating transcriptional and posttranscriptional processes (Füllgrabe et al., 2014).

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Abbreviations used in this paper: Acn, Acinus; ASAP, apoptosis- and splicing-associated protein; Da, Daughterless; ERG, electroretinogram; GMR, glass multimer reporter; mTOR, mammalian TOR; SEM, scanning EM; TEM, transmission EM; TOR, target of rapamycin; UAS, upstream activating sequence.

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**Figure 1. Acn is regulated on the protein level.** (A) Schematic of *acn* locus and genomic transgenes all inserted at 96F3 for this study. RSB is the conserved RNPS1 and SAP18 binding site (Martelli et al., 2012). (B–E'') Projections of confocal sections of eye discs showing early stages of photoreceptor development. Regions of eye discs shown are just posterior to the furrow as in the white frame of Fig. 2 D. Images show wild-type discs stained for endogenous Acn (B) and DNA (B') or discs with indicated transgenes stained for GFP (C and D) and DNA (C' and D') or for Myc (E) and GFP (E'). Arrows in E–E'' point to cells expressing GFP-KDEL but lacking Myc-Acn. Merged images are to the right. Bars, 20  $\mu$ m. Detailed genotypes are given in Table S2.

One such regulator of autophagy is encoded by the *acinus* (*acn*) gene (Haberman et al., 2010). Acn proteins are highly conserved between flies and mammals, but their function is incompletely understood. Acn was originally discovered as a protein that induces DNA condensation or fragmentation after its activation by Caspase-3 in apoptotic cells (Sahara et al., 1999; Joselin et al., 2006). Acn is, however, not required for apoptosis, which progresses unimpeded in *Drosophila melanogaster* and mammalian Acn loss-of-function cells (Joselin et al., 2006; Haberman et al., 2010). Acn, RNPS1, and SAP18 form the nuclear apoptosis- and splicing-associated protein (ASAP) complex (Schwerk et al., 2003; Joselin et al., 2006; Murachelli et al., 2012), which modulates RNA metabolism through interactions with the exon junction complex, spliceosomes, and messenger ribonucleoprotein particles (Tange et al., 2005; Joselin et al., 2006; Singh et al., 2010; Hayashi et al., 2014; Malone et al., 2014).

*Drosophila acn* alleles were identified in a screen for mutants affecting endolysosomal trafficking (Haberman et al., 2010). In the same study, a role of Acn in regulating autophagy was revealed by loss- and gain-of-function experiments in *Drosophila* larval fat bodies, a tissue poised for robust up-regulation of autophagy in response to environmental or developmental cues (Rusten et al., 2004; Scott et al., 2004). In fat body cells lacking *acn* function, early autophagosomes stall and fail to fuse with lysosomes. Conversely, systemic Acn overexpression causes excess autophagy that is lethal to flies. Genetic epistasis experiments placed the requirement of Acn for regulating autophagy downstream or in a pathway parallel to the TOR complex: up-regulation of autophagy in response to dominant-negative TOR is suppressed in *acn* mutants, but enhanced autophagy as a result of Acn gain of function is not repressed by activated TOR (Haberman et al., 2010). A high-content RNAi screen for genes involved in viral autophagy in human cells revealed a conserved role for Acn in autophagy (Orvedahl et al., 2011).

Here, we explore the regulation of Acn function as part of a pathway regulating basal levels of autophagy. We find that Acn integrates multiple inputs as its activity is inhibited by Caspase-mediated cleavage and enhanced by AKT1-dependent phosphorylation. Genetically stabilized forms of Acn increase the level of basal autophagy sufficiently to prolong life span and partially suppress Huntingtin-induced neurodegeneration.

## Results

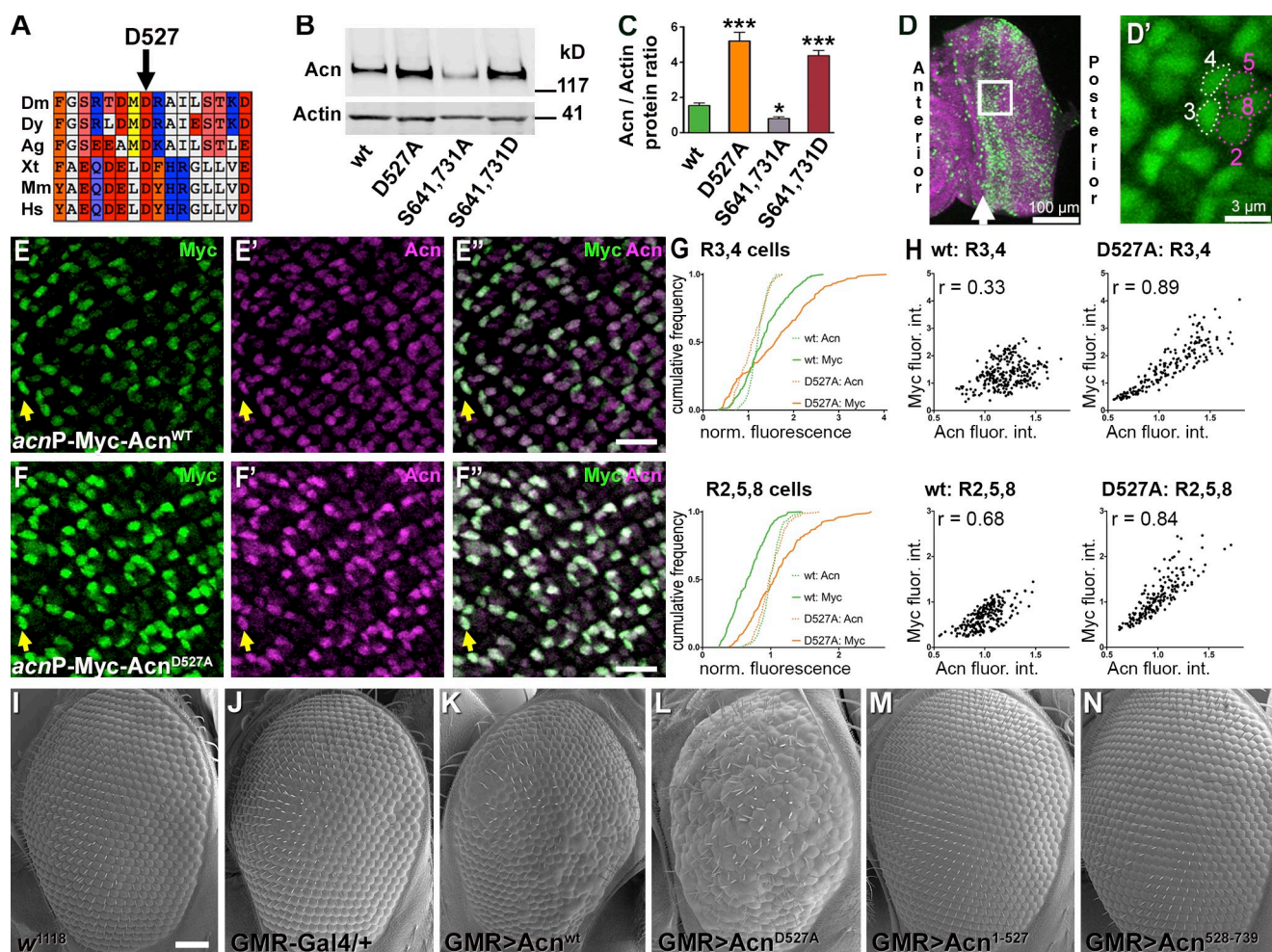
### Acn is dynamically regulated at the protein level

Acn levels in developing photoreceptor cells are dynamically regulated in a cell type-specific manner (Haberman et al., 2010). To investigate the mechanisms that regulate Acn in eye discs, we expressed two GFP transgenes under control of the endogenous 4-kb *acn* genomic region (Fig. 1 A), which is sufficient to rescue all *acn* phenotypes (Fig. S1; Haberman et al., 2010). A transgene in which GFP was fused to the N terminus of Acn mimicked the expression pattern of endogenous Acn (Fig. 1, B and C) and was sufficient to rescue *acn* lethality. In contrast, when GFP was expressed from the same control region but separated from Acn by a stop codon, cell type-specific regulation was lost (Fig. 1 D). This argues against a major contribution of transcriptional control to dynamic changes of Acn expression in developing photoreceptors. To directly test whether Acn, when separated from its genomic control regions and 5' and 3' UTRs, is regulated at the protein level, we used the eye-specific glass multimer reporter (GMR)–Gal4 driver to coexpress Myc-tagged Acn and an endoplasmic reticulum-targeted GFP-KDEL from upstream activating sequence (UAS) transgenes with identical UTRs (Fig. 1 E). As expected, GFP-KDEL was expressed in all cells posterior to the furrow, but Acn levels varied greatly: some cells displayed strong expression, whereas others were devoid of it (e.g., Fig. 1 E, arrowheads). These results indicate that Acn is dynamically regulated at the protein level.

### Acn is cleaved at the conserved D527 site

Acn is highly conserved; *Drosophila* and human Acn share 53% identity in their P17 domains (Fig. 1 A). This includes D527





**Figure 2. Acn is cleaved and inactivated at aspartate D527.** (A) Conserved aspartate D527 of *Drosophila melanogaster* (Dm) Acn is shown in alignment with *Drosophila yakuba* (Dy), *Anopheles gambiae* (Ag), *Xenopus tropicalis* (Xt), mouse (*Mus musculus* [Mm]), and human (*Homo sapiens* [Hs]) proteins. Color code for amino acid groups: brown for aromatic, white for hydrophobic, salmon for hydroxyls, red for acidic, blue for basic, cyan for amido, and yellow for sulfur. (B) Western blot of lysates from larvae probed for Acn and Actin. (C) Quantification of blots for Acn levels relative to Actin ( $n = 3$ ). Error bars show means  $\pm$  SD. \*,  $P < 0.05$ ; \*\*\*,  $P < 0.001$ . (D) Micrograph of eye disc stained for DNA (magenta) and Acn (green). The frame just posterior to the furrow (arrow) indicates the region shown in high magnification images (E and F). (D') Micrograph of an eye disc expressing nuclear GFP. Dotted lines encircle R3/4 and R2/5/8 photoreceptors just posterior to the furrow. (E–F') Dynamic changes in Acn expression in photoreceptors visualized in projections of confocal sections of eye discs from *acnP-Myc-Acn*<sup>wt</sup> (E) and *acnP-Myc-Acn*<sup>D527A</sup> (F) larvae double labeled for Myc and C-terminal Acn epitopes of Myc-Acn (merged images are shown in E' and F'). Arrows point to R3/4 photoreceptor pairs just posterior to the furrow. Bars, 10  $\mu$ m. (G and H) Quantification of Acn and Myc levels in early R3/4 and R2/5/8 cells measured from  $\geq 180$  ommatidia from four eye discs, each for *Acn*<sup>wt</sup> and *Acn*<sup>D527A</sup>. (G) Cumulative frequency histograms of normalized fluorescence intensities for Myc and Acn immunoreactivity. (H) Intensity blots of normalized integrated intensities of Acn and Myc immunoreactivity for individual groups of R3/4 and R2/5/8 cells in *Myc-Acn*<sup>wt</sup> and *Myc-Acn*<sup>D527A</sup> discs and calculated Pearson correlation coefficients  $r$ . fluor. int., fluorescence intensities. (I–N) SEM images of wild-type control (I) and eyes expressing under GMR-Gal4 control the indicated UAS transgenes: none (J), UAS-Acn<sup>wt</sup> (K), UAS-Acn<sup>D527A</sup> (L), UAS-Acn<sup>1-527</sup> (M), and UAS-Acn<sup>528-739</sup> (N). Bar, 50  $\mu$ m. Detailed genotypes and roughness quantification are given in Tables S1 and S2. wt, wild type.

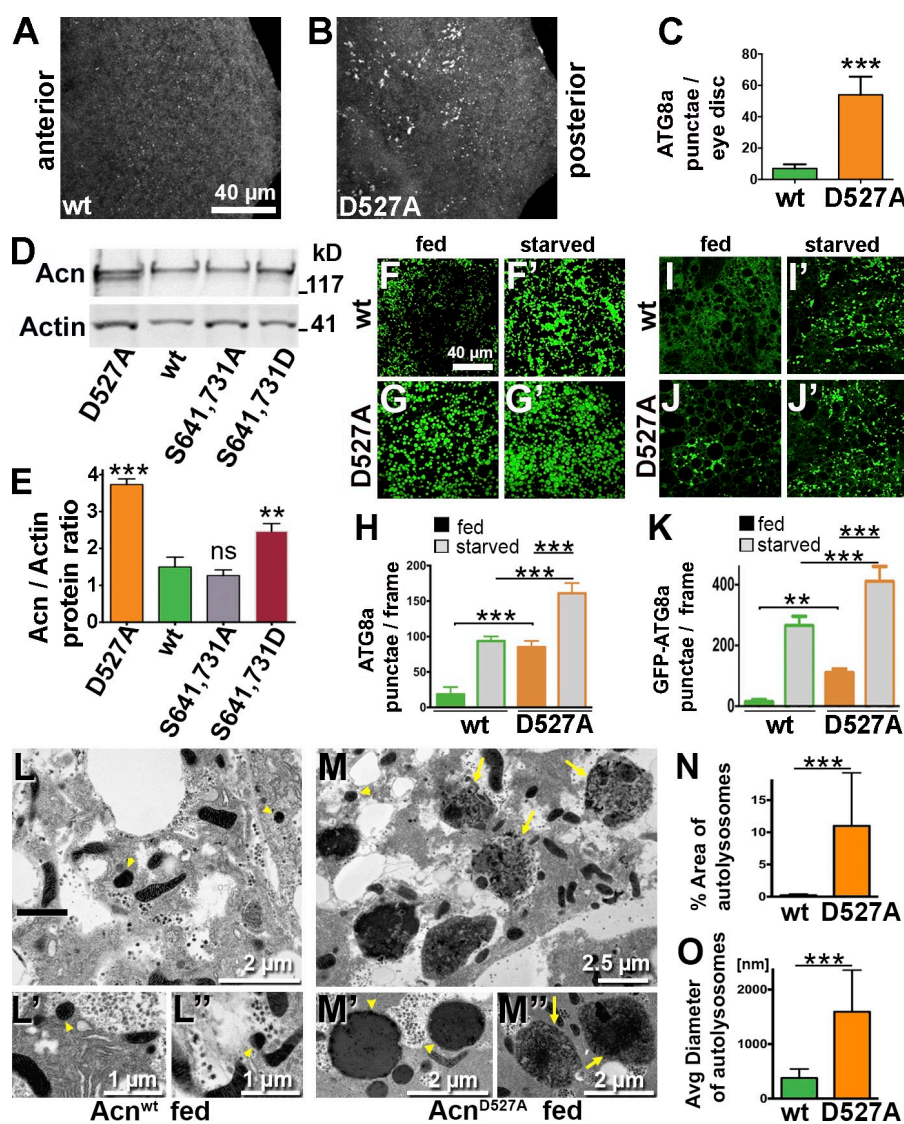
(Fig. 2 A), which aligns with the aspartate residue cleaved by Caspase-3 in apoptotic mammalian cells (Sahara et al., 1999).

To test whether *Drosophila* Acn is proteolytically cleaved at this site in vivo, we generated flies in which endogenous Acn was replaced by Myc-tagged Acn<sup>D527A</sup> or Acn<sup>wt</sup> as a control. Expression was under control of the endogenous *acn* promoter and enhancers within a 4-kb *acn* genomic region (Fig. 1 A). All genomic *acn* transgenes were inserted at 96F3 using the site-specific unidirectional PhiC31 recombinase to avoid insertion site-specific differences in expression levels (Groth et al., 2004; Venken et al., 2006). When assessed by quantitative RT-PCR, the level of expression was indistinguishable for both transgenes, and both rescued lethality and endocytic trafficking

defects of *acn*-null alleles (Fig. S1). We will refer to these rescued flies that, in an *acn*-null background, express exclusively transgenic forms of Acn as Acn<sup>wt</sup> or Acn<sup>D527A</sup> flies from here on. This system allows us to test in vivo the properties of various Acn mutants without interference from the endogenous Acn protein. In such rescued larvae, levels of Acn<sup>D527A</sup> were about three times higher when compared with Acn<sup>wt</sup> (Fig. 2, B and C), indicating that the larval Acn protein was stabilized by loss of the caspase cleavage site.

Acn stabilization was even more obvious in eye discs (Fig. 2, D–H) double stained with antibodies against the N-terminal Myc epitope and an Acn epitope close to the C terminus (Haberman et al., 2010). In eye discs expressing Myc-Acn<sup>wt</sup>, the ratio of Myc

**Figure 3. Stabilized *Acn*<sup>D527A</sup> enhances basal autophagy.** (A and B) Micrographs of fed *Acn*<sup>wt</sup> or *Acn*<sup>D527A</sup> larval eye discs stained for Atg8a. (C) Quantification of Atg8a punctae in eye discs from five larvae. (D) Western blot of fat body lysates from 96-h fed larvae probed for *Acn* and Actin. (E) Quantification of blots for *Acn* levels relative to Actin ( $n = 3$ ). (F–G' and I–J') Micrographs of *Acn*<sup>wt</sup> or *Acn*<sup>D527A</sup> fat bodies encompassing six to eight cells from 96-h fed or starved size-matched larvae. (F–G') Fat bodies stained with antibodies against Atg8a. (H) Quantification of Atg8a punctae averaged from five larvae from one representative experiment out of three repeats. (I–J') GFP-Atg8a fluorescence detected in live fat bodies. (K) Quantification of GFP-Atg8a punctae averaged from five larvae from one representative experiment out of three repeats. (L–M') TEMs of fed *Acn*<sup>wt</sup> or *Acn*<sup>D527A</sup> fat bodies. Examples of dense lysosomes (arrowheads) and membrane-enriched autolysosomes (arrows) are shown at higher magnification. (N) Quantification of percentage of autolysosomal area averaged from 25 images per genotype. (O) Quantification of diameters of  $\geq 100$  lysosomes and autolysosomes per genotype. Detailed genotypes are given in Table S2. wt, wild type. Error bars show means  $\pm$  SD. \*\*,  $P < 0.01$ ; \*\*\*,  $P < 0.001$ .



to *Acn* levels changed dynamically in photoreceptor nuclei. In emerging R3/R4 cells, identified by their shape and position in ommatidia close to the furrow (Fig. 2 D'; Wolff and Ready, 1993), the ratio of Myc to *Acn* was high. That ratio dropped in older R3/R4 cells more posterior to the furrow. Loss of the Myc-tagged N-terminal part of *Acn* relative to the C-terminal region is consistent with cleavage of *Acn* and preferential degradation of the N-terminal fragment. In *Acn*<sup>D527A</sup> eye discs, in contrast, Myc immunoreactivity was increased in R3/R4 as well as in R2/R5/R8 cells (Fig. 2, E–G). Stabilization of *Acn*<sup>D527</sup> was also revealed by the increased correlation of Myc and *Acn* immunoreactivity in early photoreceptor cells when compared with *Acn*<sup>wt</sup> (Fig. 2 H). These data indicate that in developing photoreceptors *Acn* is cleaved at aspartate<sup>527</sup>.

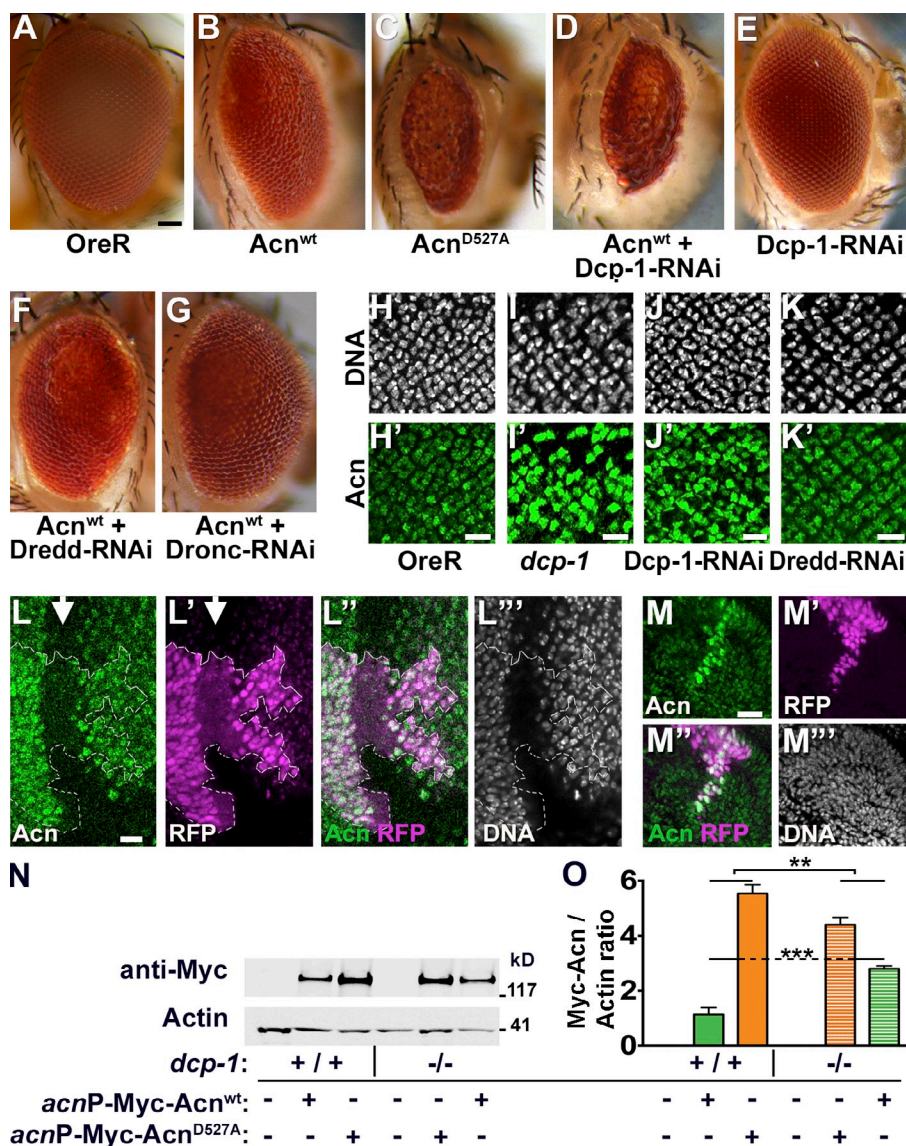
In apoptotic mammalian cells, a chromatin-modifying function of *Acn* is activated by caspase-mediated cleavage (Sahara et al., 1999; Hu et al., 2005; Joselin et al., 2006). To test whether *Drosophila* *Acn* was activated by its cleavage, we compared the effects of UAS-controlled transgenes expressing full-length *Acn* or truncated versions that would be generated by cleavage at aspartate<sup>527</sup> (Fig. 2, I–N). These UAS transgenes

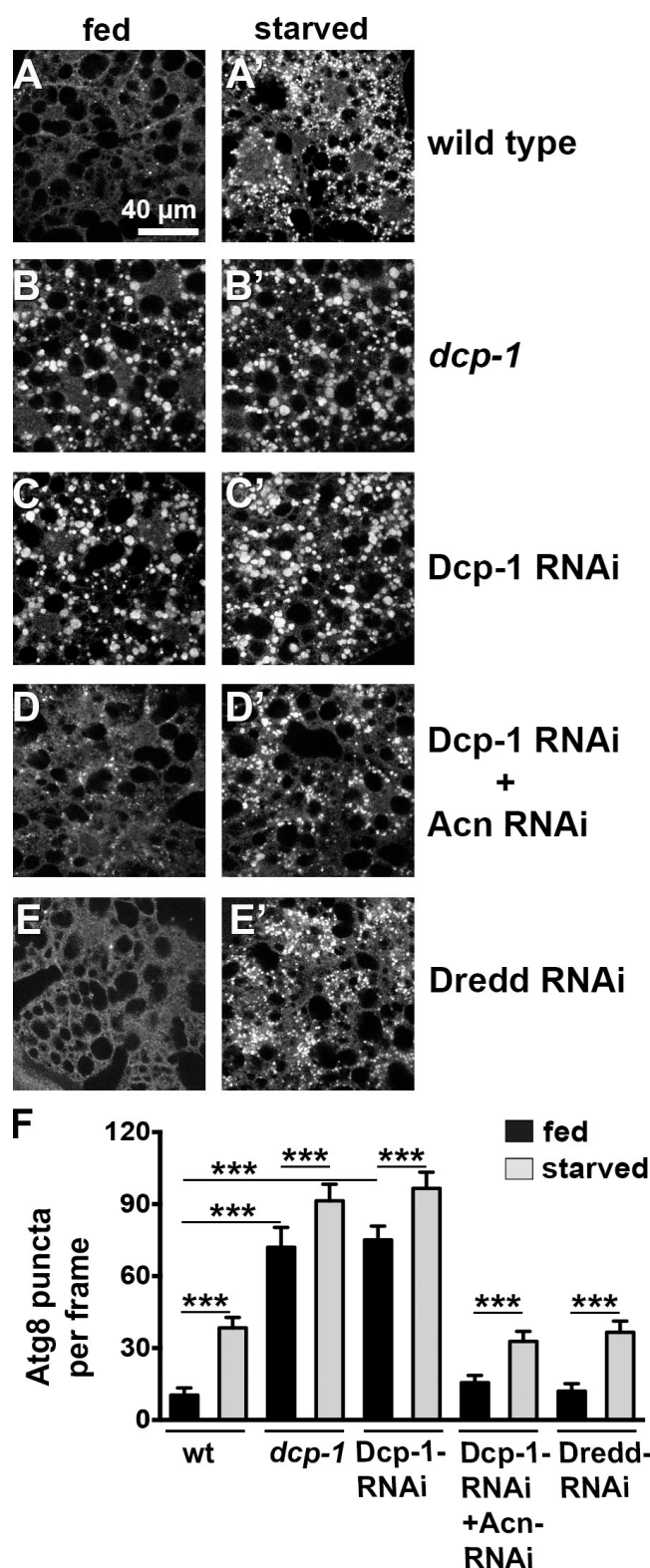
were expressed under control of the eye-specific GMR-Gal4 driver. Compared with wild-type (Fig. 2 I) or GMR-Gal4 control eyes (Fig. 2 J), eyes expressing wild-type UAS-*Acn* were mildly rough (Fig. 2 K). Expression of stabilized *Acn*<sup>D527A</sup> caused enhanced roughness (Fig. 2 L). In contrast, eyes expressing the truncated UAS-*Acn*<sup>1–527</sup> (Fig. 2 M) or UAS-*Acn*<sup>528–739</sup> (Fig. 2 N) appeared unchanged from controls. Furthermore, Daughterless (Da)-Gal4-driven expression of truncated *Acn* proteins, unlike the corresponding UAS-*Acn*<sup>wt</sup>, could not rescue lethality of *acn*-null alleles. In summary, our data indicate that in developing photoreceptors, *Acn* is inactivated by cleavage at aspartate<sup>527</sup>.

### **Acn cleavage regulates autophagy**

We have previously shown that *Acn* function is required downstream of TOR for autophagy, whereas elevated levels of *Acn* induce autophagy independent of TOR activity (Haberman et al., 2010). We therefore tested whether larvae expressing only stabilized *Acn*<sup>D527A</sup> from the endogenous promoter exhibited altered levels of autophagy. In these *Acn*<sup>D527A</sup> eye discs, Atg8-positive autophagosomes formed at rates significantly higher than in *Acn*<sup>wt</sup>, indicating increased autophagy (Fig. 3, A–C).







**Figure 5. Dcp-1 regulates basal autophagy.** (A–E') Micrographs show fat bodies from 96-h fed and starved size-matched larvae stained for Atg8a. Genotypes are wild type (A), *dcp-1* (B), Da-Gal4, UAS-*Dcp-1*-RNAi (C), Da-Gal4, UAS-*Dcp-1*-RNAi + UAS-*Acn*-RNAi (D), and Da-Gal4, UAS-*Dredd*-RNAi (E). Each image encompasses six to eight cells. (F) Quantification of Atg8a puncta averaged from five larvae from one representative experiment out of three repeats. wt, wild type. See also Fig. S1 H for efficiency of *Acn* knockdown. Detailed genotypes are given in Table S2. Error bars show means  $\pm$  SD. \*\*\*,  $P < 0.001$ .

GMR-Gal4–driven expression of UAS-*Acn*<sup>wt</sup> at elevated temperature (28°C; Fig. 4, A and B). This phenotype was further enhanced when stabilized UAS-*Acn*<sup>D527A</sup> was expressed instead (Fig. 4 C). We reasoned that reduced activity of the protease responsible for *Acn* cleavage should enhance the roughness induced by UAS-*Acn*<sup>wt</sup> to a similar level to that of cleavage-resistant UAS-*Acn*<sup>D527A</sup>.

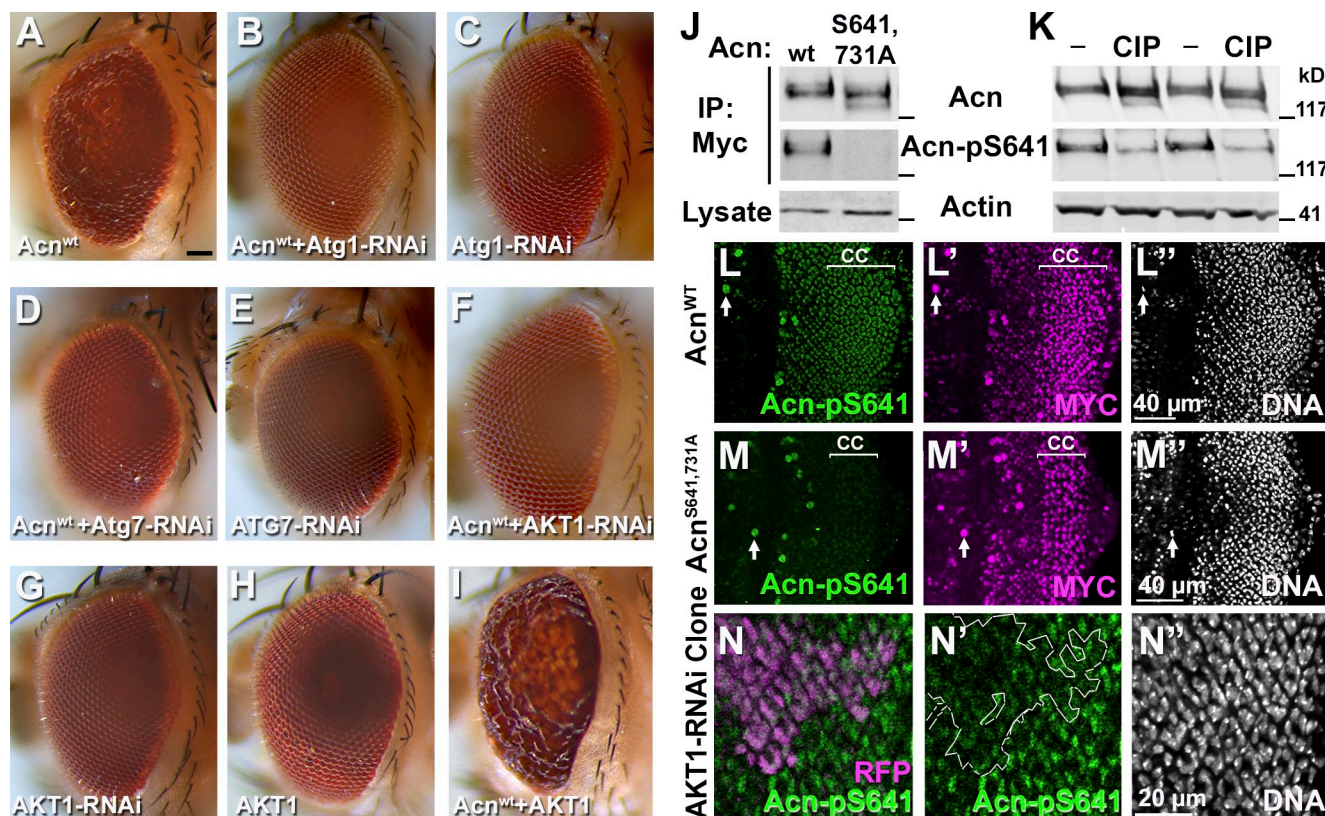
Among the eight caspases tested (Table S1), only *dcp-1* (*death caspase-1*) genetically interacted with *Acn*. Two different UAS-*Dcp-1*-RNAi transgenes, when coexpressed with UAS-*Acn*<sup>wt</sup> using the GMR-Gal4 driver enhanced roughness and reduced eye size similar to UAS-*Acn*<sup>D527A</sup> (Fig. 4 D and not depicted). Importantly, expression of *Dcp-1* RNAi by itself did not cause eye roughness (Fig. 4 E). For all other caspases tested in the eye, RNAi coexpression with UAS-*Acn*<sup>wt</sup> did not obviously modify the eye roughness (Fig. 4, F and G, shows *Dredd*-RNAi and *Dronc*-RNAi as examples).

To investigate the role of *Dcp-1* further, we analyzed its effect on *Acn* levels. Eyes mutant for *dcp-1* (Fig. 4 I'; Laundrie et al., 2003) or expressing *Dcp-1*-RNAi (Fig. 4 J'), but not *Dredd*-RNAi (Fig. 4 K'), exhibited increased *Acn* levels in photoreceptor cells in larval eye discs compared with wild type (Fig. 4 H'). Moreover, flippase-induced *Dcp-1* knockdown clones (marked by RFP) in eye-antennal discs demonstrated enhanced *Acn* levels compared with surrounding wild-type tissue (Fig. 4, L and M). Furthermore, we compared the level of Myc-tagged *Acn*<sup>wt</sup> and *Acn*<sup>D527A</sup> in both wild-type and *dcp-1* larvae by Western blotting. In a wild-type background, the level of Myc-tagged *Acn*<sup>D527A</sup> is higher than *Acn*<sup>wt</sup> (Fig. 4, N and O). This difference is significantly reduced, and the level of *Acn*<sup>wt</sup> is significantly higher in the *dcp-1* background (Fig. 4, N and O). These data identify the Caspase-3 homologue *Dcp-1* as the main protease necessary for cleavage of *Acn* at aspartate<sup>527</sup> in nonapoptotic cells, such as developing photoreceptors.

#### Dcp-1-dependent cleavage of *Acn* regulates autophagy

Next, we tested whether loss of *Dcp-1* function and the resulting *Acn* stabilization induces autophagy. We compared autophagosome levels in 96-h fat bodies using Atg8a staining. Compared with wild type, *dcp-1* mutant or *Dcp-1* knockdown fat bodies displayed robustly induced autophagy in fed larvae (Fig. 5, A–C and F). A similar enhancement was observed for the formation of autolysosomes detected by LysoTracker staining (Fig. S2, F–K). Enhanced autophagy was specific for *Dcp-1* caspase, as knockdown of *Dredd* did not induce autophagy in fed larvae (Fig. 5, E and F; and Fig. S2, J and K). For mutant and RNAi-induced *dcp-1* loss-of-function larvae, the number of Atg8a-positive puncta was further increased upon starvation (Fig. 5, B' and C'). To test whether the increased basal autophagy in response to loss of *Dcp-1* function is mediated through elevated levels of *Acn*, we generated animals in which *Dcp-1* and *Acn* were knocked down together. In those larvae, fed or starved, autophagy was reduced back to wild-type levels (Fig. 5, D and F; and Fig. S2, I and K), indicating that the remaining protein after *Acn* knockdown (Fig. S1 H) is sufficient to support starvation-induced autophagy (Fig. 5, D' and F; and Fig. S2, I' and K).





**Figure 6. AKT1 phosphorylates Acn.** (A–I) Micrographs of eyes expressing the indicated UAS transgenes under control of GMR-Gal4. Bar, 50  $\mu$ m. (J) Western blots of anti-Myc immunoprecipitates of Myc-Acn<sup>wt</sup> and Myc-Acn<sup>S641,731A</sup> expressed in S2 cells and detected with antibodies against Acn or Acn-pS641. S2 cell lysates were probed for Actin. IP, immunoprecipitation; wt, wild type. (K) Western blots of lysates from S2 cells expressing Myc-Acn<sup>wt</sup> that were treated without (–) or with calf intestinal phosphatase (CIP) and probed for Acn, Acn-pS641, or Actin. (L–N'') Apical most regions of eye discs that include cone cell nuclei (marked CC) are depicted in z projections of confocal sections. Eye discs were from larvae expressing Myc-Acn<sup>wt</sup> (L–L'') or Myc-Acn<sup>S641,731A</sup> (M–M'') and were stained for Acn-pS641, Myc, and DNA. Arrows point to unspecific staining in mitotic cells. (N–N'') Projection of confocal sections of eye disc expressing AKT1-RNAi within RFP-marked clones (magenta) and stained for Acn-pS641 (N') and DNA (N''). The broken line indicates the clone boundary. Detailed genotypes and roughness quantification are given in Tables S1 and S2.

but not sufficient to drive elevated levels of basal autophagy in a fed *dcp-1* background (Fig. 5, D and F; and Fig. S2, I and K). Together, these data are consistent with a requirement for Dcp-1 in nonapoptotic cells to control the level of Acn protein and thereby the basal starvation-independent level of autophagy.

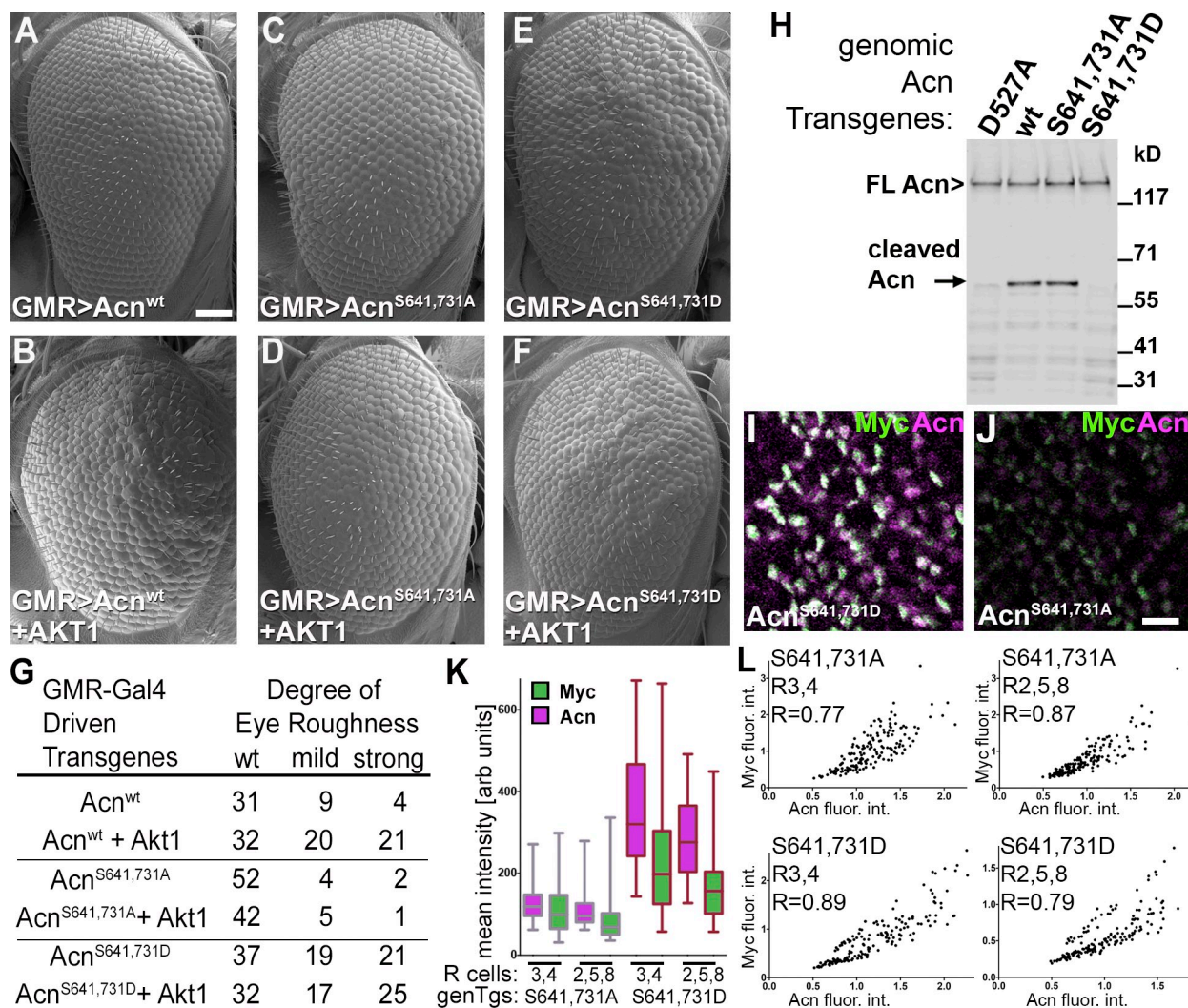
### AKT1 regulates Acn stability

High-level overexpression of Acn causes autophagy-dependent lethality (Haberman et al., 2010). Therefore, we tested whether the Acn-induced rough eye (Figs. 2 K and 6 A; compare with wild-type eyes in Figs. 2 I and 4 A) is also a reflection of overactive autophagy. We found that knockdown of Atg1 or Atg7 suppressed the UAS-Acn<sup>wt</sup>-induced rough eye (Fig. 6, B and D; and Table S1), whereas expression of RNAi for Atg1 or Atg7 by itself resulted in an eye indistinguishable from wild type (Fig. 6, C and E). This indicates that the Acn-induced rough eye, at least in part, is caused by overactive autophagy.

Next, we used this sensitized system to test whether AKT1 functions to regulate Acn-mediated autophagy. Two findings suggested such a role of AKT1: (1) Acn contains two conserved AKT1 consensus targets sites (RxRxxS/T) at S641 and S731 (Fig. 1 A), and (2) AKT1 can phosphorylate mammalian Acn in vitro and reduce its proteolytic cleavage (Hu et al., 2005).

Importantly, the UAS-Acn<sup>wt</sup>-induced rough eye (Fig. 6 A) was suppressed when UAS-AKT1-RNAi was coexpressed (Fig. 6 F). In contrast, by themselves, AKT1 knockdown (Fig. 6 G) or UAS-AKT1 overexpression in the eye (Fig. 6 H) did not result in visible phenotypes. GMR-Gal4-driven coexpression of UAS-Acn<sup>wt</sup> and UAS-AKT1, however, enhanced the rough eye (Fig. 6 I). These strong genetic interactions between Acn and AKT1 are consistent with AKT1 phosphorylating Acn and thereby enhancing its activity.

To further analyze this possibility, we generated a phospho-specific antibody against Acn-pS641, one of the two conserved AKT1 target sites. To confirm its specificity, we used S2 cells to express AKT1 together with either Myc-tagged Acn<sup>wt</sup> or Acn<sup>S641,731A</sup>, which eliminates both potential AKT1 target sites. After immunoprecipitation using anti-Myc antibodies, both Acn proteins were detected by anti-Acn antibodies on Western blots. The phosphospecific Acn-pS641 antibody, however, detected only wild-type Acn but not dephosphomimetic Acn<sup>S641,731A</sup> (Fig. 6 J). Furthermore, in lysates of Myc-Acn<sup>wt</sup>-expressing S2 cells, treatment with calf intestinal phosphatase substantially reduced the signals obtained with Acn-pS641 on Western blots but not those of anti-Acn or Actin control antibodies (Fig. 6 K). Together, these data indicate that Acn is phosphorylated at serine<sup>641</sup>



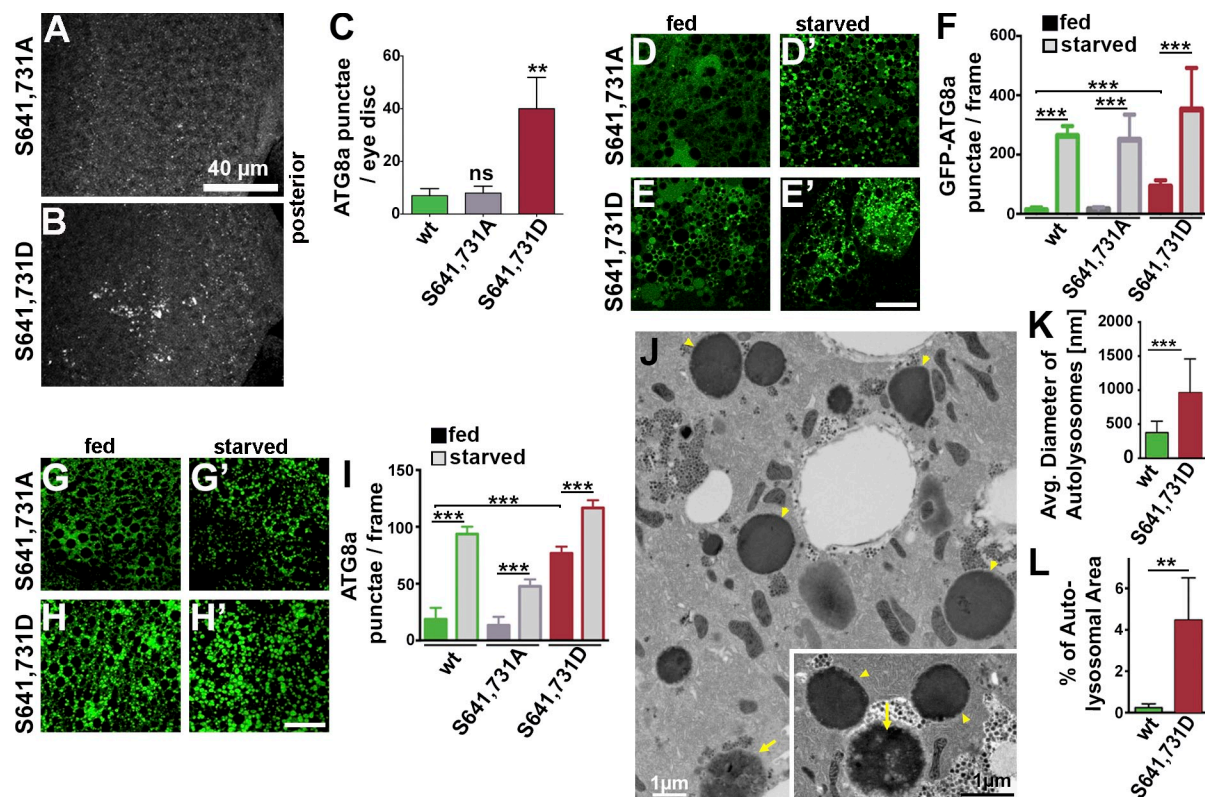
**Figure 7. AKT1 phosphorylation regulates Acn levels.** (A–F) SEMs of eyes expressing the indicated UAS-Acn transgenes with or without UAS-AKT1 under control of GMR-Gal4. Bar, 50  $\mu$ m. (G) Number of flies with the indicated degree of roughness for genotypes as in A–F. (H) Western blot of lysates from fed 75-h larvae with the indicated Acn transgenes probed for Acn. Bands of full-length and cleaved Acn are indicated. FL, full length; wt, wild type. (I and J) Acn expression in developing photoreceptors visualized in projections of confocal sections of eye discs from *acn*P-Myc-Acn<sup>S641,731D</sup> (I) or *acn*P-Myc-Acn<sup>S641,731A</sup> (J) larvae stained for Myc and Acn. Bar, 10  $\mu$ m. (K and L) Quantification of Acn and Myc levels in early R3/4 and R2/5/8 cells from  $\geq 120$  ommatidia from three eye discs. (K) Box and whisker graphs of integrated intensities of Acn and Myc immunoreactivity for individual groups of R3/4 and R2/5/8 cells in discs with the genomic transgenes (genTgs) Myc-Acn<sup>S641,731A</sup> and Myc-Acn<sup>S641,731D</sup>. Box and whisker graphs show boxes with medians, 25 and 75 percentiles, and whiskers extending to minimal and maximal values. arb units, arbitrary units. (L) Intensity blots of normalized integrated intensities of Acn and Myc immunoreactivity for individual groups of R3/4 and R2/5/8 cells in Myc-Acn<sup>S641,731A</sup> and Acn<sup>S641,731D</sup> discs and calculated Pearson correlation coefficients *r*. flour. int., fluorescence intensities. Detailed genotypes are given in Table S2.

and that our Acn-pS641 antibody specially recognizes phosphorylated Acn.

To explore the role of Acn phosphorylation, we generated flies in which endogenous Acn was replaced by Myc-Acn<sup>S641,731A</sup> or Myc-Acn<sup>S641,731D</sup> substituting either an inert alanine or a phosphomimetic aspartate residue for the two potential AKT1 phosphorylation sites. Both mutant Acn proteins were expressed from genomic transgenes inserted at the same 96F3 landing site as described previously for Acn<sup>wt</sup> and Acn<sup>D527A</sup> (Fig. 1 A), and both rescued lethality and endocytic trafficking defects of *acn*-null mutations (Fig. S1, A–F). Acn-null flies rescued by the phosphomimetic Acn<sup>S641,731D</sup> or the dephosphomimetic Acn<sup>S641,731A</sup> did not display externally visible phenotypes.

We used these flies to explore phosphorylation of Acn at serine<sup>641</sup> in eye discs. In cone cells, Acn<sup>wt</sup> is more consistently expressed at high levels than in the dynamically changing expression of photoreceptors (Haberman et al., 2010). Cone cells (Fig. 6 L, marked CC) also show strong staining with Acn-pS641 antibodies. This staining is dramatically reduced in larvae expressing only dephosphomimetic Acn<sup>S641,731A</sup> (Fig. 6 M), although anti-Myc antibodies detect Acn<sup>S641,731A</sup> expression in cone cells (Fig. 6 M'). To investigate whether this phosphorylation is caused by AKT1 activity, we generated AKT1 knockdown clones in eye discs marked by RFP. In such clones, Acn-pS641 antibody staining was reduced compared with surrounding wild-type tissue (Fig. 6 N), consistent with AKT1 phosphorylating Acn.





**Figure 8. AKT1-mediated phosphorylation of Acn modulates basal autophagy.** (A and B) Micrographs of fed  $Acn^{S641,731A}$  or  $Acn^{S641,731D}$  eye discs stained for Atg8a. (C) Quantification of Atg8a punctae in eye discs from five larvae. (D–E' and G–H') Micrographs of  $Acn^{S641,731A}$  or  $Acn^{S641,731D}$  fat bodies encompassing six to eight cells from 96-h fed or starved size-matched larvae. Bars, 40  $\mu$ m. (D–E') GFP-Atg8a fluorescence detected in live fat bodies. (F) Quantification of GFP-Atg8a punctae averaged from five larvae from one representative experiment out of three repeats. (G–H') Fat bodies stained for Atg8a. (I) Quantification of Atg8a punctae averaged from five larvae from a single representative experiment out of three repeats. (J) TEM images of fed 96-h fat bodies expressing  $Acn^{S641,731D}$ . Examples of dense lysosomes (arrowheads) and membrane-enriched autolysosomes (arrows) are shown at higher magnification (inset). (K) Quantification of diameters of  $\geq 100$  lysosomes and autolysosomes diameters per genotype. (L) Quantification of percentage of autolysosomal area averaged from 25 images per genotype. Wild-types images are shown in Fig. 3 L. Detailed genotypes are given in Table S2. wt, wild type. Error bars show means  $\pm$  SD. \*\*,  $P < 0.01$ ; \*\*\*,  $P < 0.001$ .

The aforementioned results suggest that phosphorylation of the AKT1 target sites in Acn is the main basis for the genetic interactions between AKT1 and Acn. To further test this possibility, we expressed  $Acn^{wt}$  or the AKT1 target site mutants  $Acn^{S641,731D}$  and  $Acn^{S641,731A}$  from UAS-controlled transgenes. GMR-Gal4-driven coexpression of AKT1 enhanced the rough eye phenotype of the  $Acn^{wt}$  transgene (Fig. 7, A, B, and G) but not roughness of either of the transgenes with mutated AKT1 target sites (Fig. 7, C–G).

Next, we tested whether this phosphorylation affects Acn levels by comparing  $Acn^{S641,731A}$  and  $Acn^{S641,731D}$  flies. Western blotting of whole 96-h larvae or fat bodies revealed that, compared with  $Acn^{wt}$ , the level of dephosphomimetic  $Acn^{S641,731A}$  was reduced, and phosphomimetic  $Acn^{S641,731D}$  was increased similar to the cleavage-resistant  $Acn^{D527A}$  (Fig. 2, B and C; and Fig. 3, D and E). This accumulation was not yet detectable in 75-h larval lysates (Fig. 7 H). However, in these early larvae, an intermediate cleavage product detectable for  $Acn^{wt}$  and  $Acn^{S641,731A}$  was drastically reduced for phosphomimetic  $Acn^{S641,731D}$ , similar to caspase cleavage-resistant  $Acn^{D527A}$  (Fig. 7 H, arrow). These findings suggest that the AKT-mediated phosphorylation of Acn stabilizes the protein against proteolytic degradation. Stabilization of phosphomimetic  $Acn^{S641,731D}$  was also evident in developing

photoreceptors in third instar discs (Fig. 7, I–L). Similar to the effect of caspase cleavage-resistant  $Acn^{D527A}$  (Fig. 2), phosphomimetic  $Acn^{S641,731D}$  exhibited higher levels of staining with Acn and Myc antibodies (Fig. 7, I and K), and the correlation of staining for N-terminal Myc and C-terminal Acn epitopes was increased (Fig. 7 L). Dephosphomimetic  $Acn^{S641,731A}$  (Fig. 7 J) displayed low levels of Acn and Myc staining, but the remaining protein displayed elevated levels of correlation between Acn and Myc staining (Fig. 7 L) when compared with wild type (Fig. 2 H). Together, these data support the notion that in fat bodies and developing photoreceptor cells, Acn stability is modified by AKT1-dependent phosphorylation.

As increased levels of Acn can enhance autophagy, we examined Acn proteins with mutated AKT1 target sites for their effect on autophagy. Compared with  $Acn^{wt}$  (Fig. 3 A) or dephosphomimetic  $Acn^{S641,731A}$  (Fig. 8, A and C), eye discs expressing  $Acn^{S641,731D}$  at endogenous levels exhibited increased numbers of autophagosomes (Fig. 8, B and C). Furthermore, fat bodies of fed  $Acn^{S641,731D}$  larvae exhibited significantly enhanced basal autophagy at early and late stages of the autophagy pathway as detected by increased GFP-Atg8a punctae, punctae of antibody-stained endogenous Atg8a (Fig. 8, D–I), and LysoTracker staining (Fig. S2, D and E). In addition, TEM revealed larger and

more abundant autolysosomes (Fig. 8, J–L). Interestingly, these effects closely resembled those observed in larvae expressing stabilized  $\text{Acn}^{\text{D527A}}$  resistant to Dcp-1 cleavage (compare with Fig. 3). In contrast, basal levels of autophagy were unaltered for destabilized  $\text{Acn}^{\text{S641,731A}}$  (Fig. 8, A–I; and Fig. S2 C). Nevertheless,  $\text{Acn}^{\text{S641,731A}}$  and  $\text{Acn}^{\text{S641,731D}}$  both supported significant increases in autophagy upon starvation (Fig. 8, D–I; and Fig. S2, C' and D'), arguing against a significant contribution of AKT1-mediated phosphorylation of Acn to the regulation of starvation-induced autophagy. Together, these data indicate that AKT1-mediated phosphorylation regulates Acn stability and, thereby, the basal levels of autophagy.

### Acn gain of function enhances quality control autophagy and improves life span

Next, we explored the physiological consequences of altered Acn regulation and the resulting changes in autophagy. The most striking effect of stabilized Acn proteins was the significant increase in basal autophagy (Figs. 3 and 8), which is important for quality control and maintenance of muscles (Demontis and Perrimon, 2010; Bai et al., 2013) and neurons (Hara et al., 2006; Komatsu et al., 2006; Mizushima et al., 2008). Neurodegeneration in flies can be efficiently followed by electroretinograms (ERGs) as a sensitive measure of the health of photoreceptor neurons (Williamson et al., 2010). At 1 or 2 wk of age, wild-type flies exhibit ERGs with a mean sustained depolarization of  $\sim 10$  mV. Acn knockdown significantly reduced depolarization after 1 wk and even further after 2 wk (Fig. 9, A and B). Expression of the Huntingtin-derived HttQ93 protein in photoreceptor cells using GMR-Gal4 triggers strong degeneration in flies with only a small detectable depolarization remaining after 1 wk (Fig. 9, A and B). This was further reduced by Acn knockdown (Fig. 9, A and B), prompting us to test whether increasing autophagy through stabilized versions of Acn could improve retinal degeneration induced by HttQ93. We found that in 2-wk-old flies, one copy of the genomic  $\text{Acn}^{\text{D527A}}$  transgene significantly improved depolarization compared with  $\text{Acn}^{\text{wt}}$  (Fig. 9, C and D) and also reduced the load of aggregated HttQ93 protein as measured by dot blots (Fig. 9 E).  $\text{Acn}^{\text{S641,731D}}$  also lowered aggregated HttQ93 protein levels (Fig. 9 E) but not sufficiently to reduce neurodegeneration (Fig. 9 D).

To test whether the beneficial long-term effects of stabilized Acn extend beyond the eye, we measured survival of Acn transgenic flies. First, we tested a role of Acn in survival under starvation conditions (Fig. 9 F). When resistance to starvation was compared with  $\text{Acn}^{\text{wt}}$ , flies expressing only stabilized  $\text{Acn}^{\text{D527A}}$  ( $P < 0.0001$  log rank) or  $\text{Acn}^{\text{S641,731D}}$  ( $P < 0.0001$  log rank) survived significantly longer in the absence of food. In contrast, survival of  $\text{Acn}^{\text{S641,731A}}$  flies was not significantly different.

Next, we tested whether Acn stability can modify longevity. Compared with  $\text{Acn}^{\text{wt}}$  or  $\text{Acn}^{\text{S641,731A}}$ , we found significantly enhanced life spans for stabilized  $\text{Acn}^{\text{D527A}}$  and  $\text{Acn}^{\text{S641,731D}}$  (both  $P < 0.0001$  log rank), with the median life span extended by 50% from 38 to 58 d for  $\text{Acn}^{\text{D527A}}$  (Fig. 9 G). Together, these results are consistent with a model in which AKT1 phosphorylates Acn to inhibit its caspase-mediated cleavage, and resulting elevated levels of Acn enhance basal autophagy with beneficial effects for neural maintenance and prolonged life span.

## Discussion

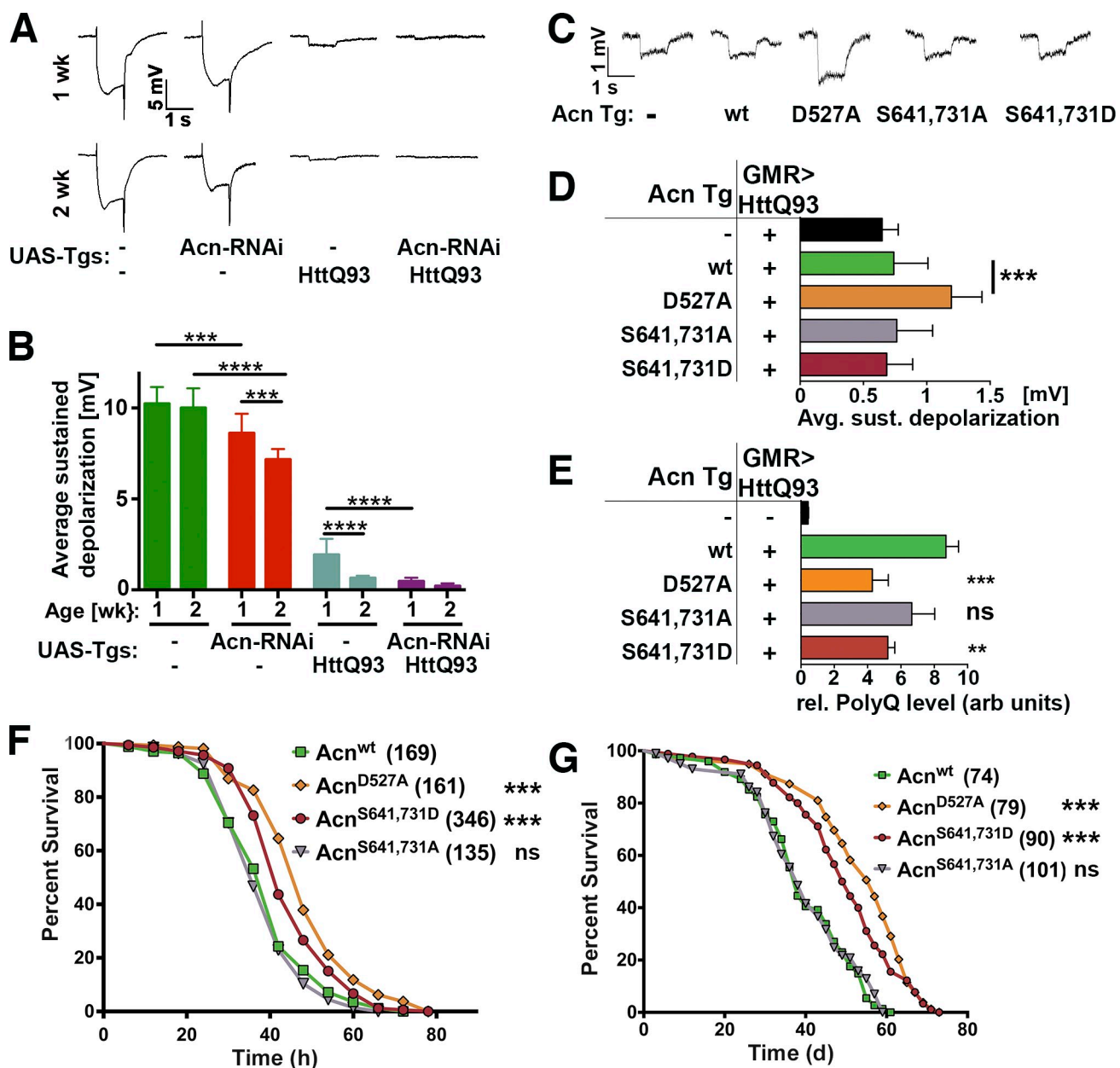
We have previously shown that Acn is a regulator of autophagy in *Drosophila* as loss or gain of Acn function results in reduced or exaggerated autophagy (Haberman et al., 2010). In this study, we examine the mechanisms by which Acn levels are fine-tuned in developing tissues to determine the level of basal autophagy. We find that Acn levels are negatively regulated by Dcp-1-mediated cleavage. Elevated levels of Acn, whether caused by loss of Dcp-1 function, mutations modifying its target site D<sup>527</sup>, or mimicking AKT1-mediated phosphorylation, enhanced basal autophagy. The requirement of basal autophagy for neuronal maintenance and normal longevity is well established in flies (e.g., Simonsen et al., 2007; Juhász and Neufeld, 2008). Corresponding benefits resulted when stabilized Acn proteins enhanced basal autophagy: such flies lived longer, were more resistant to starvation stress, and exhibited reduced neurodegeneration in a *Drosophila* Huntington's disease model (Scherzinger et al., 1997). We cannot exclude that roles in other pathways, such as endocytic trafficking (Haberman et al., 2010) or RNA metabolism (Murachelli et al., 2012), may contribute to the benefits of enhanced Acn function. The importance of elevated basal autophagy is consistent, however, with the beneficial outcomes of its up-regulation in neurons or muscles by increased levels of Atg8a in flies (Simonsen et al., 2008; Bai et al., 2013) or Beclin 1 in mice (Spencer et al., 2009).

The window for beneficial effects is narrow, however. Eye-specific overexpression of Acn disrupts retinal development in an autophagy-dependent manner (this study), and systemic high-level expression can cause lethality as a result of excessive autophagy (Haberman et al., 2010). Detrimental consequences of autophagy also result from overexpression of Atg1 (Scott et al., 2007). In a more limited way, autophagy contributes to the elimination of cells of salivary glands or the midgut during metamorphosis (Berry and Baehrecke, 2007; Denton et al., 2009). A similarly ambivalent picture arises from genetic activation of autophagy in *Caenorhabditis elegans* (Kang et al., 2007) and from pharmacological studies in murine cells (Sarkar et al., 2009; Liu et al., 2013; Shoji-Kawata et al., 2013), with low levels of autophagy being protective for cells but high levels causing cell death. Intriguingly, changes in Acn levels, its phosphorylation, or epigenetic status have been observed in different cancer cells (Shu et al., 2006; Jang et al., 2008; Patwa et al., 2008) and linked to differential cell survival (Park et al., 2009).

Our data suggest that these links between cancer and Acn may reflect its conserved function in autophagy. In the past, however, much of these data have been attributed to a role of Acn in apoptosis caused in part, by the initial discovery of Acn as a factor promoting apoptotic DNA condensation or fragmentation after being activated by caspase-mediated cleavage (Sahara et al., 1999). We find that caspase-mediated cleavage of Acn as well as its stabilization by AKT-mediated phosphorylation (Hu et al., 2005) is conserved in flies, but our results differ in two aspects.

First, *acn*-null cells still display DNA fragmentation and condensation (Haberman et al., 2010), and here, we show that eye-specific overexpression of either of the fragments predicted





**Figure 9. Acn regulates quality control autophagy and longevity.** (A and B) ERGs of 1- or 2-wk-old flies expressing the indicated UAS transgenes (Tgs) under control of GMR-Gal4. (B) Quantification of sustained negative photoreceptor response averaged from three replicate experiments including at least 64 traces from eight flies. (C and D) ERGs of 2-wk-old flies all expressing UAS-HttQ93 under control of GMR-Gal4 (GMR>HttQ93) and the indicated genomic Acn transgenes. (D) Quantification of sustained negative photoreceptor response from three replicate experiments including at least 64 traces from eight flies. Avg. sust., average sustained. (E) Quantification of dot blots measuring aggregated HttQ93 protein in fly heads expressing UAS-HttQ93 under control of GMR-Gal4 (GMR>HttQ93) and the indicated genomic Acn transgenes. Flies expressing neither indicate background level of dot blot measurements ( $n = 3$ ). rel., relative; arb units, arbitrary units. (F) Starvation-induced mortality of flies expressing the indicated Acn proteins. In the parenthesis shown are the numbers of initial flies in a single representative experiment. (G) Survival curves of flies expressing the indicated Acn proteins. In the parenthesis shown are the numbers of initial flies in a single representative experiment. Detailed genotypes are given in Table S2. wt, wild type. Error bars show means  $\pm$  SD. \*\*,  $P < 0.01$ ; \*\*\*,  $P < 0.001$ ; \*\*\*\*,  $P < 0.0001$ .

from caspase-mediated cleavage at Acn<sup>D527</sup> does not disrupt normal development. Furthermore, elevated levels of Acn in the eye, which are sufficient to induce increased autophagy, are not sufficient to increase levels of apoptosis (Fig. 3 B and not depicted). Moreover, developmental defects induced by GMR-Gal4-driven overexpression of Acn are suppressed by reduced function of core autophagy genes but not by reduced function of caspases, including the apical caspase Dronc (Figs. 4 and 6). Together, these data

suggest that the role of Acn in DNA condensation or fragmentation in apoptotic mammalian cells (Sahara et al., 1999; Joselin et al., 2006) may not be conserved in *Drosophila*. In contrast, Acn function in autophagy is conserved between *Drosophila* and human cells (Haberman et al., 2010; Orvedahl et al., 2011).

Second, cleavage of Acn by Dcp-1 is not restricted to apoptotic cells but is found in developing photoreceptors and fat body cells. In apoptotic cells, caspases gain access to nuclear

proteins upon breakdown of the nuclear envelope (Faleiro and Lazebnik, 2000). The cleavage of nuclear-enriched Acn by cytoplasmic Dcp-1 in nonapoptotic photoreceptor precursor cells thus suggests that Acn shuttles in and out of the nucleus. This is consistent with Acn being detected in the cytoplasm in a subset of eye discs cells (Haberman et al., 2010) and with a predicted nuclear export signal at amino acids 691–698 of Acn (la Cour et al., 2004).

Caspase-mediated cleavage in nonapoptotic cells is not unique to Acn, other examples include dSREBP (*Drosophila* sterol regulatory element-binding protein) activation by Drice (Amarneh et al., 2009) and local caspase activation during axon branching (Campbell and Okamoto, 2013). Subapoptotic activation of caspases contributes to tumor invasion and metastasis (Rudrapatna et al., 2013). Atg16l1 levels, which are critical for the regulation of autophagy in Paneth cells (Cadwell et al., 2008), are modulated by stress-induced, but subapoptotic, levels of Caspase-3 (Murthy et al., 2014). Mutations in *Atg16l1* that enhance its cleavage by Caspase-3 thus impair autophagy and pathogen clearance and thereby contribute to Crohn's disease (Hampe et al., 2007).

Other proautophagic targets of caspases include Atg4 (Betin and Lane, 2009) and Beclin 1 (Luo and Rubinsztein, 2010; Wirawan et al., 2010). Their cleavage by caspases also reduces autophagy and suppresses its cytoprotective function (Kroemer et al., 2010).

A surprising finding of our study is the proautophagy function that AKT1 can exert through its effect on Acn levels. This observation contrasts with previous results in flies and mammals, which firmly established an inhibitory role of AKT1 signaling in autophagy (Kroemer et al., 2010; Mariño et al., 2014). In the context of the canonical PtdIns3K–Akt–TOR pathway, AKT1 activates the mTOR complex directly by phosphorylation of Raptor and indirectly via the inhibitory phosphorylation of the TSC1/2 inhibitory complex (Neufeld, 2010). Furthermore, AKT1-mediated phosphorylation of Beclin 1 inhibits its proautophagy function (Wang et al., 2012). In contrast, our results imply a proautophagy role for AKT1 signaling as a result of its conserved phosphorylation of Acn and the resulting inhibition of its proteolytic cleavage (Hu et al., 2005; Park et al., 2009; this study).

How can we resolve the conundrum of these seemingly opposing AKT1 functions in the regulation of autophagy? A possible explanation is provided by the presence of distinct pools of AKT1 kinase that may be engaged in different contexts (Martelli et al., 2012). For example, although Beclin 1, mTOR, and TSC1/2 are primarily cytoplasmic proteins, Acn is mostly nuclear in flies and mammals (Schwerk et al., 2003; Haberman et al., 2010). Therefore, a distinct nuclear pool of AKT1 may be responsible for Acn regulation (Martelli et al., 2012). This is not without precedent. Nuclear AKT1 has been suggested to phosphorylate Foxo proteins (Martelli et al., 2012) and the coactivator p300 (Huang and Chen, 2005), which have been implicated in regulating autophagy (Demontis and Perrimon, 2010; Ferdous et al., 2010; Lo Ré et al., 2012). Consistent with the notion of two distinct AKT1 pools regulating separate modes of autophagy, we find that AKT1-dependent activation of Acn does not occlude additional enhancement of autophagy via the conventional

starvation-induced mTOR pathway. Physiologically, the availability of independent pathways for the induction of autophagy may be advantageous for cells as they respond to distinct challenges. Acn activation may provide an alternative pathway to engage basal, starvation-independent autophagy without invoking the complex metabolic changes triggered by altered mTOR signaling (Laplanche and Sabatini, 2012; Costa-Mattioli and Monteggia, 2013).

How elevated levels of Acn induce autophagy is not clear. It may increase the activity of the ASAP complex that Acn forms with SAPI8 and RNPS1 (Schwerk et al., 2003) or reduce an inhibitory function of the alternative Pinin-containing ASAP complex, in which Acn is replaced by Pinin (Murachelli et al., 2012). These complexes have been implicated in the regulation of splicing, nuclear export, translation and decay of mRNAs through their interactions with spliceosomes, the exon junction complex, and messenger ribonucleoprotein particles (Tange et al., 2005; Joselin et al., 2006; Singh et al., 2010, 2012; Hayashi et al., 2014; Malone et al., 2014), but the primary function of ASAP or Pinin-containing ASAP complexes is still poorly understood. Alternatively, the regulation of autophagy by Acn may be independent of these complexes and instead relay on its interaction with other binding partners, such as AAC-11. Inhibition of their interaction reduced resistance to chemotherapeutic drugs possibly as a result of altered apoptosis or autophagy (Rigou et al., 2009). Chemotherapy is one of many stress situations in which it is important to understand how the proper balance between apoptosis and autophagy is regulated (White, 2012; Mariño et al., 2014). The mechanisms by which Acn contributes to that regulation will be an important goal of future research.

## Materials and methods

### Fly work

Flies were maintained using standard conditions. RNAi lines were obtained from Vienna *Drosophila* RNAi Center or the Bloomington Stock Center, which also provided Da-Gal4, Lsp2-Gal4, and GMR-Gal4 driver lines, *w<sup>1118</sup>*, UAS-GFP-KDEL, UAS-AKT, P[*hs-FLP*]<sup>1</sup>, *y<sup>1</sup> w<sup>1118</sup>*; Dr<sup>1</sup>/TM3, *Sb<sup>1</sup>*, P[*hs-FLP*]<sup>1</sup>, *w<sup>1118</sup>*; *Adv<sup>1</sup>/CyO*, and *w<sup>1118</sup>*; P[GAL4-Act5C(FRT.CD2).P]<sup>5</sup>, P[UAS-RFP.WJ<sup>3</sup>/TM3, *Sb<sup>1</sup>*. Other fly strains used were *dcp-1* (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; Landrie et al., 2003; gift from K. McCall, Boston University, Boston, MA), UAS-*htt-exon1-Q93* (expresses Huntington's disease exon1 peptide with 93 glutamines under UAS control; Steffan et al., 2001; gift from R. Hiesinger, University of Texas Southwestern Medical Center, Dallas, TX), and UAS-GFP-Atg8a (expresses GFP-tagged Atg8a protein under Gal4/UAS control; Juhász et al., 2008; gift from T. Neufeld, University of Minnesota, Saint Paul, MN). RNAi clones were generated by heat shocking larvae carrying *hs-Flp*; Actin>CD2>Gal4, UAS-RNAi, and UAS-RFP at 38°C for 1 h for a consecutive 2 d. Transgenic flies were generated by BestGene, Inc. DNA constructs in the context of genomic *acn* were generated by standard mutagenesis of the 4-kb Acn DNA fragment sufficient for genomic rescue (Haberman et al., 2010), confirmed by sequencing, cloned into an *AttB* vector, and inserted into the 96F3 *attP* landing site (Venken et al., 2006). Similarly, UAS-controlled wild-type and mutant Acn transgenes were generated by standard mutagenesis from full-length Acn cDNA, confirmed by sequencing, and inserted into pUAS vectors. An initial set of UAS-Acn<sup>wt</sup> and UAS-Acn<sup>D527A</sup> transgenes was generated by *P* element-mediated transformation. Subsequently, transgenes—including new lines of UAS-Acn<sup>wt</sup> and UAS-Acn<sup>D527A</sup>—were generated in a pUAS(l) variant with an added *AttB* site and inserted into the 53B2 or 28E7 landing sites (Venken et al., 2006) as indicated (Table S2).

Starvation resistance was measured in vials with 4–5-d-old virgins at 25°C at 6-h intervals. To measure life spans, males that emerged within a



2-d period were pooled and aged for an additional 3 d, and their survival at 25°C was recorded as described in Stenlesen et al. (2013). In brief, 70–100 flies were placed in demography cages (~1 liter). Every other day, food vials were changed, and dead flies were counted and removed. Initial life span measurements were performed with the *acn* transgenes in an *acn*<sup>1</sup>/*acn*<sup>27</sup> background with both chromosomes containing background lethals (Haberman et al., 2010) and, in the absence of the TM6, Hu balancer. Results in multiple experiments in that background were qualitatively the same as for the experiments shown (Fig. 9 G) for which the chromosome carrying the *acn*<sup>27</sup>-null allele was cleaned from other mutations before crossing the different *acn* transgenes into the isogenized *acn*<sup>27</sup> stock for the longevity measurements.

### Biochemistry

Acn-pS641 antibody was raised in rabbits by Genemed Synthesis against the RSRSGS(p)PASKTKKC peptide and double affinity purified. For Western blots, five 96-h or ten 75-h larvae were crushed in 300  $\mu$ l lysis buffer (10% SDS, 6 M urea, and 50 mM Tris-HCl, pH 6.8) at 95°C, boiled for 2 min, and spun for 10 min at 20,000 g to remove larval cuticle. 20  $\mu$ l lysate was separated by SDS-PAGE. Proteins were transferred to nitrocellulose membranes or, for detection of Atg8 proteins, onto 0.2- $\mu$ m polyvinylidene fluoride membranes and probed with mouse anti-Actin (1:2,000; JLA 20; Developmental Studies Hybridoma Bank), guinea pig anti-Acn (aa 423–599; 1:3,000; Haberman et al., 2010), mouse anti-Myc (1:2,000; 9E10; Covance), rabbit anti-Acn-pS641 (1:1,000), or rabbit anti-Atg8a (1:5,000; Nagy et al., 2014; gift from G. Juhász, Eötvös Loránd University, Budapest, Hungary). Bound antibodies were detected and quantified by comparison to Actin using fluorescence-labeled antibodies and the Odyssey scanner (LI-COR Biosciences). Prestained molecular weight markers were obtained from Invitrogen (HiMark) or New England Biolabs, Inc. (ColorPlus).

For expression in S2 cells, plasmids encoding Myc-tagged versions of Acn<sup>wt</sup> and Acn<sup>S641,731A</sup> under control of the metallothionein promoter were transfected using the TransIT-2020 reagent (Mirus Bio LLC) followed by induction of expression, cell lysis, and immunoprecipitation with anti-Myc beads (Pulipparacharuvil et al., 2005). For phosphatase treatment, Acn<sup>wt</sup>-transfected S2 cell lysates were treated with 400 U calf intestinal phosphatase (New England Biolabs, Inc.).

Larval fat bodies were dissected from 10 larvae in ice-cold 50  $\mu$ l larval lysis buffer (25 mM Tris-HCl, 1 mM EDTA, 0.1 mM EGTA, 5 mM MgCl<sub>2</sub>, 150 mM NaCl, 2 mM Na<sub>2</sub>VO<sub>4</sub>, 10% glycerol, 1% NP-40, 1 mM DTT, and protease and phosphatase inhibitor mixes [Roche]). Dissected fat bodies were sonicated for 5 min on ice and mixed with 50  $\mu$ l Freon. The mixture was centrifuged at 10,000 g for 10 min, and supernatants were collected for Western blots.

Transcript levels of Myc-tagged Acn transgenes were measured using quantitative RT-PCR as previously described (Akbar et al., 2011). In short, RNA was isolated using TRIzol (Ambion) according to the manufacturer's instructions. 2  $\mu$ g RNA was reverse transcribed using High-Capacity cDNA Reverse Transcription kit (Applied Biosystems) using random hexamer primers. Quantitative PCR was performed using the Fast SYBR Green Master Mix in a real-time PCR system (Fast 7500; Applied Biosystems). Each data point was repeated three times and normalized with rp49 as an internal control. Primers were left, 5'-CTGGAGGAGCAGAAGCTGAT-3', within the Myc and right, 5'-GGAGTCTCGACCTCGGTCTT-3', within the Acn coding regions, and rp49 primers were left, 5'-ATCGGTTACGGATC-GAACAA-3', and right, 5'-GACAATCTCCTTGCGCTTCT-3'.

### Histology

Micrographs of eyes were obtained on a microscope (Stereo Discovery V12; Carl Zeiss) with a camera (AxioCam MRC 5; Carl Zeiss) using AxioVision image acquisition software (Carl Zeiss) at 72 $\times$  magnification. Images of eyes are a composite of pictures taken at multiple z positions and compressed using CZFocus software.

For TEM, size-matched 96-h fed larvae were dissected, inverted to expose fat bodies, and fixed in 2% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.2. Samples were postfixated with 2% OsO<sub>4</sub> and 1.5% KFeCN in the same buffer. Samples were embedded in epoxy resin. A series of ultrathin sections (60–80 nm) containing fat body were cut, and sequential sections were collected on mesh and formvar-coated slot grids. Sections were stained with uranyl acetate and lead citrate to enhance contrast, examined with a transmission electron microscope (120 kV; Tecnai G2 Spirit BioTWIN; FEI), and images were captured with an 11-megapixel camera (Morada; Olympus). From TEMs, measurements of autolysosomal diameters and areas were obtained using Macnification software (Orbicle).

SEM of fly eyes were obtained as previously described (Wolff, 2011). In short, eyes were fixed in 2% paraformaldehyde, 2% glutaraldehyde, 0.2% Tween 20, and 0.1 M cacodylate buffer, pH 7.4, for 2 h. Samples were processed in a series of four washes with increasing ethanol (25–100%) for 12 h, each followed by a series hexamethyldisilazane washes (25–100% in ethanol) for 1 h each. Flies air dried overnight were mounted on SEM stubs and coated in fast-drying silver paint on their bodies only. Flies were sputter coated with a gold/palladium mixture for 90 s and imaged at 1,000 $\times$  magnification, with extra high tension set at 3.0 kV on a scanning electron microscope (SIGMA; Carl Zeiss). The microscope was equipped with the InLens detector (Carl Zeiss).

Whole-mount tissues were prepared for immunofluorescence staining as previously described (Akbar et al., 2011). In short, dissected samples were fixed in periodate-lysine-paraformaldehyde, washed in PBS, permeabilized with 0.3% saponin in PBS (PBSS), blocked with 5% goat serum in PBSS, and stained with the indicated primary antibodies: guinea pig anti-Acn (1:1,000; Haberman et al., 2010), mouse anti-Myc (9E10; 1:1,000), rabbit anti-dAtg8a (1:500; Barth et al., 2011; Abcam; gift from K. Koehler, Eidgenössische Technische Hochschule Zürich, Zürich, Switzerland), rabbit anti-Boss (1:1,000), rabbit anti-activated Caspase (1:300; Cell Signaling Technology), or rabbit-GFP (1:500; Invitrogen), and secondary antibodies were labeled with Alexa Fluor 488, 568, or 647 (1:500; Molecular Probes) and mounted in Vectashield (Vector Laboratories). Fluorescence images were captured with 63 $\times$ , NA 1.4 or 40 $\times$ , NA 1.3 Plan Apochromat lenses on an inverted confocal microscope (LSM 510 Meta; Carl Zeiss). Confocal z stacks of eye discs were collected at 1- $\mu$ m step size.

To quantify Acn/Myc immunoreactivity in eye discs, confocal stacks were imported to ImageJ (National Institutes of Health), and apical slices containing nuclei of early photoreceptors were projected into a single slice by summation of pixel intensities. Masks for individual groups of R3/4 and R2/5/8 cells were generated in ImageJ based on nuclear GFP expression (Fig. 2 D'). Integrated pixel intensities for Acn and Myc immunoreactivity were determined for  $\geq 120$  ommatidia from at least three eye discs per genotype. To compare different eye discs and measure the correlation coefficients for individual R-cell groups, integrated densities for each R-cell group were normalized to the mean value of Acn intensities in R2/5/8 cells for a given eye disc. For statistical evaluation and computation of Pearson correlation coefficients, data were imported into Prism software (GraphPad Software).

LysoTracker (GFP-Certified Lyso-ID red lysosomal detection kit; Enzo Life Sciences) staining of size-matched 90–96-h fat bodies from fed and starved larvae was performed as previously described (Rusten et al., 2004; Scott et al., 2004). In brief, larvae were dissected in Schneider's *Drosophila* media (Gibco), inverted to expose fat bodies, and incubated in 100  $\mu$ M LysoTracker Red DND-99 for 1 min. Inverted carcasses were then washed in PBS and placed onto a droplet of Vectashield (Vector Laboratories) for fat body separation and mounting. Samples were imaged immediately using 63 $\times$ , NA 1.4 Plan Apochromat lens on an inverted confocal microscope (LSM 510 Meta). LysoTracker and Atg8a punctae in fat bodies were quantified using Imaris software (Bitplane) from z projections of three optical sections of fat body tissue, each 1  $\mu$ m apart. The number of punctate was quantified either per fat body cell or per frame, in which one frame represents a volume of (471  $\mu$ m)<sup>2</sup>  $\times$  2  $\mu$ m = 44  $\times$  10<sup>3</sup>  $\mu$ m<sup>3</sup>. Digital images for display were imported into Photoshop (Adobe) and adjusted for gain, contrast, and  $\gamma$  settings.

### ERGs

ERGs were recorded as previously described (Williamson et al., 2010). In brief, female flies were immobilized with nontoxic Glue-All (Elmer's). Recording and reference electrodes containing 2 M NaCl were placed on the fly's corneal surface and inserted into the thorax, respectively. Voltage measurements were filtered through an electrometer (IE-210; Warner Instruments), digitized with a Digidata 1440A and MiniDigi 1B system (Molecular Devices), and recorded using Clampex (version 10.2; Axon Instruments). 1-s light pulses were provided by a computer-controlled white light-emitting diode system (MC1500; Schott). Eight ERG traces from at least eight individual flies of each genotype were used for quantification with Clampfit software (Axon Instruments).

### Poly-Q dot blot filter retardation assay

Polyglutamine aggregates were detected with a modified filter assay (Scherzinger et al., 1997). In brief, 25 heads were homogenized in 200  $\mu$ l Cytoplasmic Extraction Reagent I buffer and fractionated using NE-PER Nuclear and Cytoplasmic Extraction Reagents following the manufacturer's protocol (Thermo Fisher Scientific). Cytosolic fractions were adjusted to 1%

SDS, incubated for 15 min at room temperature, denatured at 95°C for 5 min, and filtered through a 0.2- $\mu$ m cellulose acetate membrane (Sterlitech Corporation) preequilibrated with 1% SDS. Filters were washed twice with 0.2% SDS and blocked in TBS (100 mM Tris-HCl, pH 7.4, and 150 mM NaCl) containing 3% nonfat dried milk before development with mouse 1C2 antibody (1:1,000; MAB1574; EMD Millipore) and detection and quantification with Odyssey scanner and software.

### Statistical methods

Statistical significance was determined in Prism using log-rank for survival assays and one-way analysis of variance for multiple comparisons, followed by Tukey's test. Pearson correlation coefficients for Acn and Myc double-stained eye discs were calculated in Prism from normalized integrated intensities measured in ImageJ. All bar graphs resulting from these comparisons show means  $\pm$  SD. For quantifications of fluorescence images and Western blots, at least three independent experiments were used. Box and whisker graphs show boxes with median, 25 and 75 percentiles, and whiskers extending to minimal and maximal values. P-values smaller than 0.05 are considered significant, and values are indicated with one (<0.05), two (<0.01), three (<0.001), or four (<0.0001) asterisks.

### Online supplemental material

Fig. S1 shows the characterization of genomic Acn transgenes and effectiveness of Acn-RNAi. Fig. S2 shows autophagy levels in different Acn transgenes as measured by lysoTracker staining of fat bodies or ATG8 Western blots. Table S1 shows the quantification of the roughness of eyes with different genotypes for which representative examples are shown in Figs. 2, 4, and 6. Table S2 shows the complete genotypes of flies used in different figures. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201404028/DC1>. Additional data are available in the JCB DataViewer at <http://dx.doi.org/10.1083/jcb.201404028.dv>.

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