Changes in endolysosomal organization define a pre-degenerative state in the *crumbs* mutant *Drosophila* retina

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Running title: Endolysosomes in pre-degenerative retina

Keywords: Arl8, Rab7, Atg8, lysosome, photoreceptor cells, degeneration, trafficking

Abstract

Mutations in the epithelial polarity gene *crumbs* (*crb*) lead to retinal degeneration in *Drosophila* and in humans. The overall morphology of the retina and its deterioration in *Drosophila crb* mutants has been well-characterized, but the cell biological origin of the degeneration is not well understood. Degenerative conditions in the retina and elsewhere in the nervous system often involve defects in degradative intracellular trafficking pathways. So far, however, effects of *crb* on the endolysosomal system, or on the spatial organization of these compartments in photoreceptor cells have not been described. We therefore asked whether photoreceptors in *crb* mutants exhibit alterations in endolysosomal compartments under pre-degenerative conditions, where the retina is still morphologically intact. Data presented here show that, already well before the onset of degeneration, Arl8, Rab7, and Atg8-carrying endolysosomal and autophagosomal compartments undergo changes in morphology and positioning with respect to each other in *crb* mutant retinas. We propose that these changes may be early signs of the degeneration-prone condition in *crb* retinas.

Introduction

Intracellular protein trafficking is essential for the maintenance of cell and tissue homeostasis. A multitude of functions are dependent on intracellular transport, including signal transduction, secretion or membrane remodelling, to mention just a few. Therefore, it is not surprising that impaired trafficking, induced by aging, environmental factors or mutations in genes encoding important components involved in trafficking, is associated with numerous detrimental human diseases. These include neurodegenerative diseases in particular, characterized by the progressive loss of neuronal function, such as Alzheimer's and Parkinson's disease, or retinal degeneration, leading to blindness [1, 2]. Important insight into the regulation of intracellular trafficking has been obtained by studies in model organisms, notably the fruit fly *Drosophila melanogaster*. The enormous number of mutants collected over the years, the sophisticated genetic and molecular toolkit as well as the ease of performing unbiased genetic screens [3, 4] have uncovered novel molecular pathways and identified a plethora of genes safeguarding neuronal homeostasis, including those regulating intracellular trafficking [5, 6]. Importantly, roughly 75% of genes associated with human diseases have functional orthologues in the fly genome, many of which are associated with neurodegeneration [7, 8]. The fly eye is an ideal organ to study the basis of neurodegeneration and to identify the players involved: it is a non-essential organ, in which genetic mosaics can be induced, and is easily accessible to cell biological and electrophysiological analyses [3, 4, 9]. In addition, retinal photoreceptor cells are heavily reliant on the membrane trafficking system due to the need for constant renewal and repair of light-induced damage to membranes, and hence are very sensitive to any impairment in trafficking.

Phototransduction in the *Drosophila* retina takes place in a specialized organelle called the rhabdomere. It is formed at the apical surface of each photoreceptor by $\sim 30,000$ densely packed microvilli [10, 11]. Maintenance of this organelle depends on extensive membrane turnover to constantly replenish molecules of the microvillar membrane, including the light sensitive molecule rhodopsin. This process is particularly challenged during light exposure due to continual endocytosis of light-activated rhodopsin, followed by recycling to the rhabdomere or by degradation via the endolysosomal trafficking pathway [12, 13]. These

observations underscore the importance of all trafficking compartments, including those involved in autophagy, degradation and recycling. Failure in any step of these pathways often results in aberrant rhodopsin accumulation in a late endosomal, Rab7-positive compartment, and represents one cause of retinal degeneration [14–16].

One of the central players in the breakdown of biomolecules is the lysosome. Lysosomes are dynamic, membrane-bound organelles, which were initially identified as sites for degradation of intracellular components. There is increasing evidence that lysosomes are similarly important for a plethora of other functions, including metabolic signalling and plasma membrane repair [17]. Lysosomal functions are closely associated with autophagy, a conserved cellular process required for degradation and recycling of nutrients and aged or damaged organelles, whereby cellular components are enclosed in double-membrane vesicles, the autophagosome [18]. Fusion of the autophagosome with the lysosome results in formation of the autophagolysosome, in which biomolecules are degraded and/or recycled to support energy production [19].

Lysosomes are not static entities, but highly dynamic structures. They can change their intracellular position by moving bidirectionally along microtubules, and their position within a cell can have a major effect on their function [20–22]. Depending on the effector bound, late endolysosomes associated with the small GTPase Rab7 can be transported to either the microtubule minus or plus end by interactions with dynein or kinesin, respectively. Plus-end directed transport of endolysosomes is facilitated by Arl8, a small GTPase of the Arf (ADP ribosylation factor) family, which seems to engage in a "tug of war" with Rab7, and together with Rab7 can lead to tubulation of lysosomes [23–27]. In addition to Arl8 and Rab7, the retromer and its interactions with the lysosome play important roles in the degradative process important in maintaining photoreceptor homeostasis and integrity [16, 28].

Besides genes encoding components of the trafficking machinery such as the retromer, many other fly genes are required to ensure photoreceptor survival [29]. One of these is *crumbs* (*crb*), mutations in which result in light-dependent retinal degeneration [30–32]. Similarly, mutations in one of the three human orthologues, *CRB1*, result in Retinitis pigmentosa (RP12) and Leber congenital amaurosis, two of the most severe retinal dystrophies associated with blindness [33–35]. Retinal degeneration in *crb* mutant photoreceptors is strongly attenuated when flies are kept in the dark, or when raised and kept on food lacking vitamin A,

a cofactor necessary for functional rhodopsin [31]. In addition, lack of *crb* impairs MyosinV-dependent rhodopsin trafficking to the rhabdomere [36]. We recently showed that the retina of a hypomorphic *crb* allele, *crb*^{P13A9}, degenerates upon constant exposure to light [32]. The retina of this allele lacks the morphological defects in cell shape observed in other *crb* alleles [31, 37, 38], suggesting that retinal degeneration associated with lack of *crb* is independent of its developmental phenotype. These observations, together with the fact that *crb*^{P13A9} mutant flies are homozygous viable, makes this allele ideally suited to address the question to what extent a reduction of *crb* function affects the endolysosomal system and thereby could make photoreceptors more prone to damage by light.

Here, we show that photoreceptors in hypomorphic crb^{P13A9} mutant flies exhibit alterations in endolysosomal compartments under normal, non-degenerative light conditions, in which the overall morphology of the retina is still intact. Moreover, the spatial relationships between different components associated with endolysosomal and autophagosomal compartments is altered in crb mutant photoreceptors. From these results we hypothesize that spatial changes in endolysosomal organization could define a "predegenerative condition", which makes mutant photoreceptors prone to degeneration upon light stress.

Materials and Methods

Fly stocks and culture

Fly stocks used in this study were: w^* [32] (used as control), w; crb^{P13A9} [32], w; Rh1-Gal4 [39], w; UAS-LAMP-GFP (a gift of H. Krämer;[40]), GeneTrap CG6707 [41], w^* ; UAS-Rab7-GFP (a gift of M. Gonzalez-Gaitan;[42]), yw; $otd^{1.6}$ -Gal4 pWIZ8 (UAS- w^{RNAi}) (a gift of T. Cook;[43], yw; UAS-spinster-RFP (a gift of S. Sweeney; [44]), UAS-Atg8a-mCherry (a gift of T. Neufeld; [45]). For expression of transgenes encoding fluorescent protein (FP) in the crb background, the stock w; Rh1-Gal4; crb^{P13A9} was crossed to UAS-FP; crb^{P13A9} , and Rh1-Gal4/UAS-FP; crb^{P13A9} adults were selected. Unless otherwise stated, flies were raised on standard cornmeal medium, and kept at 25 °C either on a 12h light, 12h dark cycle, or in constant light at ~2100-2500 lux, or in complete darkness during pupariation and molting, until the time of dissection. For experiments with flies raised on vitamin A-depleted food, larvae were raised and flies were maintained on carotenoid-depleted food [31, 46].

Dissection and preparation of retinas

Adult females were decapitated under CO₂ and heads were bisected between the eyes with a razor blade before fixing them in cold, freshly prepared 4% formaldehyde solution for 2h on ice. Eyes of dark-reared flies were prepared in dim room-light without microscope illumination, and fixation was carried out in the dark. Preparation and fixation of retinas from flies raised in constant light was done under bright light. For 12 h light/12 h dark experiments, flies were dissected at roughly the same time of day, i.e. after ~6h of light exposure. Fixed eyes were washed in PBS/0.3% TritonX-100 (PBTx) and dissected in phosphate-buffered saline (PBS), using an insect pin with the tip bent, so that intact retinas could be scooped out of the lens. 8-10 retinas per condition were transferred individually using a P20 tip into glass wells with PBTx, and blocked in 5% normal goat serum 30 min at room temperature. For VitaminA-depletion experiments, flies were hatched and reared completely on carotenoid-free medium [46].

Retinas were incubated for at least 15h at 4°C in primary antibodies diluted in PBTx with 1% normal goat serum. Primary antibodies used in this study were: rat anti-Crb (batch 2.8; [47], 1:1000); rabbit polyclonal anti-Arl8 (1:200)[48]; Developmental Studies Hybridoma Bank (DSHB), Univ. Iowa); mouse monoclonal (MAb) anti-alpha Sodium-potassium ATPase (1:200)(AbCam #7671); mouse MAb anti-Rab7 (1:5)([48]; DSHB); rabbit anti-GFP (1:1000, Invitrogen/ThermoFisher, Cat. no. A11122); mouse MAb anti-mCherry (1:1000, MPI monoclonal antibody facility, Dresden); mouse anti-Rhodopsin1 (1:200, Millipore, Cat. no. MAB5356). Alexa 488-, 568-, and 647-coupled secondary antibodies (ThermoFisher) were used at 1:1000 dilution, and DyLight 405 secondary antibody (Jackson ImmunoResearch) at 1:500. Alexa680-coupled Phalloidin (ThermoFisher, Cat. no. A22286) was used at 1:200 to detect F-actin, together with secondary antibodies.

Microscopy and image analysis

Retinas were mounted in Vectashield (VectaLabs), under sealed coverslips supported with two streaks of nail polish to avoid flattening of the tissue. High-resolution imaging was done on a Zeiss LSM 880 Airy microscope with a 63x/1.4 NA lens. Imaging for quantification of spot sizes and intensities and for co-localization of Arl8-Rab7 was done on a Yokogawa Spinning Disk confocal microscope with a 100x/1.4 NA Silicon immersion lens, at 117 nm pixel dimensions. Laser and exposure settings were kept constant and below-saturation for all experiments.

The procedure for quantification of size and intensity of Arl8 regions was as follows: for each retina or retina fragment, between 1-6 confocal image stacks of >200 images each, spanning the depth of the retina, were acquired with an Andor iXon Ultra 888 cooled EMCCD camera at 230 nm z-distance, and processed in Fiji before deconvolution with Huygens software. For each image slice at a minimum of 1 µm z-spacing, where longitudinally oriented ommatidia were present, judged by anti-Crb staining, Arl8-positive regions within the Region Of Interest (ROI) to be analyzed were determined by automatic thresholding to detect signal above background, yielding "spots". These spots, or Arl8 regions (outlined in beige, Fig. 2C"), were defined as having an intensity over background (Signal – Noise, S-N) at least 5 standard deviations from the average intensity over the whole ROI. Average sizes and

intensities of individual Arl8 spots (or other antigen of interest) within the region were recorded using a script based on the Find Maxima plug-in in Fiji. The average for all spots in an entire stack was taken as one data point, and all data points for one experiment were normalized to the w^* control for each experiment, yielding values for w^* close to 1. Normalized data from up to three experiments were pooled for each graph. Control and crb^{P13A9} samples were always processed in parallel and data were plotted in RStudio.

Pearson's correlation coefficient (R coloc) was calculated for Arl8/Rab7 from deconvolved spinning disk image pairs using the Coloc2 plugin in Fiji, with thresholding, on ROIs drawn around longitudinal regions of ommatidial organization, judging by anti-Crb staining. Optical image slices at ~1 μm intervals were analyzed in order to avoid redundant sampling. Atg8-mCherry co-localization with Arl8 was calculated from Airy processed confocal images, using the thresholded Manders' coefficient (tM2) for the Atg8-mCherry channel alone, based on the rationale that Arl8 would be found in all cells, whereas Atg8-mCherry was only driven in photoreceptors. Isolated Atg8-mCherry compartments were chosen as ROIs in the mCherry channel, and these were then evaluated for tM2.

Results

Positioning of the Arl8-positive compartment in adult photoreceptor cells depends on light

We hypothesized that reduction of *Drosophila crb* in photoreceptor cells could render the cells susceptible to increased light stress, which ultimately results in retinal degeneration. Since impaired trafficking in photoreceptors can result in retinal degeneration [16], which is often dependent on increased light intensity, we first aimed to determine the role of light on the organization/positioning of the endolysosomal system in wild-type photoreceptors. The Arf-like small GTPase Arl8, defined as a bona-fide lysosomal marker [24, 48], has been linked to various aspects of the endolysosomal system, such as intracellular transport and fusion with the autophagosome [23, 26]. We noted that the intracellular positioning of Arl8 within the retina was altered upon changes in light conditions. When flies were exposed to a 12 hour light/12 hour dark cycle (defined as "normal light conditions" throughout the text), a proximo-distal alignment of the Arl8-positive compartments was evident in photoreceptors of eyes from the control strain (w^*) (Fig. 1A, A'). Optical cross-sections (Fig. 1B, B') revealed a predominant positioning of the Arl8-positive compartments (red) nearer to the base of the rhabdomere, the actin-rich, light sensitive organelle of the photoreceptor (white), than to the basolateral surface of the photoreceptors (green; Fig. 1B, B'). This alignment was less pronounced in photoreceptors of flies that had been kept in constant intense light for 3 days (Fig. 1C, C'), or kept in complete darkness during pupation and until 10 days after hatching (not shown). In addition, constant light induced the formation of large Arl8-positive clusters, which were less frequently located close to the base of the rhabdomere, compared to normal light conditions (Fig. 1C-D'). These results show that light conditions influence the spatial organization and positioning of the Arl8-positive lysosomal compartment in w^* control retinas.

To determine the identity of the Arl8-positive compartment in the retina, we co-labelled retinas with anti-Arl8 antibodies together with other markers of the endolysosomal trafficking pathway. The markers used included antibodies against endogenous Rab7 to label late endosomes [16, 42], and antibodies to detect overexpressed GFP-Rab7, mCherry-tagged Atg8 to label the autophago-lysosome [45], RFP-tagged Spinster (a gift of Sean Sweeney;[44]), a presumptive transporter involved in autophagosomal-lyososomal regeneration, and GFP-tagged Lysosome Associated Membrane Protein (LAMP), which is

targeted to the lysosome [40] (Suppl. Fig. S1A-E). Retinal overexpression of fusion proteins was achieved using either *Rh1*-Gal4 or *Otd*-Gal4. We also used a protein trap line, carrying an insertion in CG6707, which encodes a lysosomal phosphatidyl-inositol bisphosphatase (PIP2ase) (http://flybase.org/reports/FBal0185170.html) and labels the lysosomes [41] (Suppl. Fig. S1F)). The endogenous late endosomal protein Rab7, as well as Rab7-GFP expressed via Rh1-Gal4 in photoreceptors only (Suppl. Fig. S1A, B), are often found in Arl8-positive compartments. This contrasts with the autophagolysosomal markers LAMP-GFP, Atg8-mCherry, Spinster-RFP and LysoPIP2-ase, which occupy only partially overlapping or adjacent, but not identical compartments (Suppl. Fig. 1C-F). From these results we conclude that under normal light conditions, Arl8 is most closely associated with a late endosomal or endolysosomal compartment that also carries Rab7.

Arl8-positive compartments exhibit aberrant clustering in crb mutant retina under normal light conditions

The strong loss of function allele crb^{11A22} shows severe photoreceptor degeneration in response to light stress [30, 31]. However, the retina of these flies show additional developmental defects due to impaired elongation of the rhabdomere during pupal stages [31, 38]. In contrast, flies homozygous mutant for the hypomorphic crb^{P13A9} allele are viable and fertile, and their retina does not exhibit any of the developmental or polarity defects caused by loss-of-function crb alleles. Yet, the retina of homozygous mutant crb^{P13A9} flies undergoes degeneration under prolonged intense light conditions in a white-eyed (white, w^*) background [32]. Therefore, this allele is ideally suited to further study the effects linked to degeneration, but independent from any developmental effect.

When kept under normal light conditions (\sim 2100-2500 lux, 12 h light/12 h dark cycle), crb^{P13A9} mutant retinas reveal a striking difference in the distribution of Arl8-positive compartments compared to that of control retinas kept under the same conditions. Arl8 compartments were no longer regularly aligned along the proximo-distal axis of the cells and showed aberrant clusters. Many of these clusters looked similar to those observed in w^* retinas when exposed to constant light (compare Fig. 2B' with Fig. 1C'), but were not present in w^* retinas under normal light conditions (Fig. 2A'). In addition to large Arl8-positive

clusters, crb^{P13A9} mutant retinas frequently displayed even larger patches of Arl8. These appeared to consist of assemblies of vesicles, rather than enlarged individual vesicles (Fig. 2C' (arrows) and E). It is noteworthy that these patches are present in the crb^{P13A9} retina under non-degenerative conditions, and in the absence of any obvious loss of rhabdomeres indicative of the onset of degeneration [32]. The patches were not restricted to photoreceptors (arrow, Fig. 2G), but could also be detected in surrounding support cells (arrowheads, Fig. 2G).

To characterize the observed differences in the Arl8-positive compartment, we concentrated on the size and intensity of Arl8 spots, since these two features differed the most between mutants and controls, whereas other features, such as aspect ratios and spot numbers, were found not to correlate with any particular genotype. To quantitatively assess the differences, we devised a protocol for detecting and quantifying the Arl8-positive regions in terms of spot size and intensity, using a spot detection script in Fiji (explained in detail in Methods, and see Fig. 2C-C''). In brief, retinal regions from plane longitudinal sections (judged by Crb staining, green, in Fig. 2C) were demarcated manually in the green channel as Regions of Interest (ROIs; Fig. 2C, C'). Arl8-positive spots were then detected within the ROI for each image ("Arl8 Regions"; beige outlines in fig. 2C''). Sizes and intensities of the Arl8 regions thus defined were then calculated and averaged for each image stack. crb^{P13A9} mutant retinas of flies kept under normal 12h/light-dark conditions for 5 days showed a consistent and significant increase in the size and intensity of Arl8-positive regions compared to those of w^* mutant retinas (Fig. 2H).

Increase in size and intensity of Arl8-positive compartments in crb^{P13A9} mutant retinas depends on light and dietary vitamin A

To test the effect of light on the Arl8-positive compartments in crb^{P13A9} mutant retinas, we next modified light conditions by keeping crb^{P13A9} flies either in normal light conditions (12h light/12h dark) or in complete darkness (Fig. 3A-B'). The differences in size and intensity of Arl8-positive regions observed between crb and w^* after 10 days of normal light-dark conditions (Fig. 3E) disappeared when flies were kept in complete darkness for 10 days (Fig. 3F; similar results were found after 5 days of complete darkness; not shown).

Retinal degeneration in crb mutants is a consequence of light exposure and is rescued in the absence of vitamin A [31, 32], a precursor to the chromophore retinal, which is in turn a component of the light-activatable photopigment rhodopsin. In the absence of vitamin A, Rh1 levels are strongly reduced [46]. Therefore, we asked whether the endolysosomal abnormality reflected by an Arl8 clustering defect in crb^{P13A9} mutants could also be influenced by reducing the amount of Rh1 in the retina. The increase in size and intensity of Arl8-positive clusters in crb vs. control retinas observed under normal light conditions was abolished or even slightly reversed when flies were raised and kept in vitamin A depleted food (Fig. 3C, C', E vs. G). To summarize, the observed increase in size and intensity of Arl8-positive compartments in crb^{P13A9} vs. w^* controls depends on light and rhodopsin.

Because previous studies identified a correlation between the aberrant accumulation of rhodopsin 1 (Rh1) in Rab7-positive compartments and retinal degeneration in the fly [14, 15, 16], we examined whether the aberrant Arl8 clusters in crb^{P13A9} retinas contained Rh1. In retinas of crb^{P13A9} mutant flies raised under normal light conditions, we found that Rh1 accumulated in enlarged intracellular punctae, but these only rarely co-localized with Arl8 (arrowhead, Fig. 4B). Large Arl8 clusters and patches in crb^{P13A9} never contained rhodopsin, nor was an extensive overlap seen between Arl8 vesicles and rhodopsin in control retinas (w^*) under normal light conditions (Fig. 4A). Similarly, no localization of Rh1 was found in either genotype under constant intense light conditions (Suppl. Fig. S2).

Rab7 is abnormally distributed in crb^{P13A9} and moves to the Arl8 compartment in light

We showed that Arl8 often occupies the same compartment as the late endosomal marker Rab7 in control retinas (Suppl. Fig. S1) and that the Arl8-positive compartment undergoes changes in morphology and distribution in crb^{P13A9} mutant retinas. Therefore, we next asked whether Rab7 distribution is similarly affected in crb^{P13A9} mutant retinas. No difference in colocalization of Arl8 and Rab7 was observed in control (w^*) and crb^{P13A9} mutant retinas under normal light conditions (Fig. 5A, C; Fig. 5E, brown box plots). Rab7 was observed in some large Arl8-positive clusters of mutant retinas (arrowhead, Fig. 5C). When exposed to constant light for two days, the degree of co-localization between Arl8 and Rab7 increased significantly in both genotypes compared to the values determined under normal light conditions (Fig. 5B, D; Fig. 5E, yellow box plots). However, in constant light, total

Arl8/Rab7 co-localization was significantly less in crb^{Pl3A9} mutant retinas compared to control retinas (Fig. 5E, yellow box plots). In addition, large crescent-shaped or hollow vesicles containing both Rab7 and Arl8 were often observed in crb^{Pl3A9} mutant retinas upon two days of constant light exposure, but not in control retinas (Fig. 5F, G). Surprisingly, however, two or six days of constant light exposure did not result in increased size and intensity of Arl8-regions in crb^{Pl3A9} mutant retinas relative to w^* controls (Suppl. Fig. S3), as we had observed after five or ten days of normal light exposure (compare to Fig. 2H and Fig. 3E). To summarize, constant exposure to light promotes co-localization of Arl8 with endogenous Rab7 in control, and to a lesser extent, in crb^{Pl3A9} mutant retinas.

crb affects the distribution of the Atg8-positive compartment and its co-localization with Arl8

Arl8 function is closely linked with the autophagosome [49, 50], a compartment that is central to the prevention of retinal degeneration [15]. Since reduction of *crb* induces changes in the Arl8-positive compartment in retinal cells, we monitored possible changes in autophagosomes in the retina of control and crbP13A9 mutant flies. To do this, we examined Atg8 (Autophagy-related protein 8), which plays a pivotal role during autophagy in the formation of the autophagosomal membrane [51]. We used Atg8-mCherry (a gift from Thomas Neufeld [45]), a marker for the autophagosome and autolysosome. Atg8-mCherry expressed in control photoreceptors using Rh1-Gal4 revealed a proximo-distal distribution in photoreceptors (Fig. 6A, A') under normal light conditions, reminiscent of that of Arl8. Notably, Atg8-mCherry vesicles were adjacent to or overlapping with Arl8-positive vesicles in controls (Fig. 6C, E). In contrast, Atg8-mCherry-positive compartments in crb^{P13A9} mutant photoreceptors clustered (Fig. 6B, B') and were separated from Arl8 (Fig. 6D, E). This was reflected in reduced co-localization of individual Atg8-mCherry compartments with Arl8, assessed by measuring Manders' coefficient for the mCherry channel (tM2), using the Fiji plugin Coloc2 (Fig. 6E). Co-localization between Atg8-mCherry and Arl8, quantified as Pearson's R coloc for individual compartments, was also reduced in *crb*^{P13A9} (not shown). These results indicate that reducing *crb* function in photoreceptors lowers the spatial proximity of Arl8 to an Atg8-carrying autolysosomal compartment.

Discussion

Retinal degeneration is in most cases a gradual process, which raises two questions: i) what are the factors that drive the process of degeneration? And ii) what makes photoreceptor cells carrying mutations in certain genes more prone to degeneration, and how is this predisposition manifested at the cellular level prior to the onset of the degenerative process? In other words, how do extrinsic and genetic, intrinsic factors impair retinal homeostasis? To date, about 100 genes have been linked to retinal degeneration in human. Strikingly, the majority of these have orthologs in *Drosophila* [53]. Many of these genes encode proteins involved in trafficking, which is not surprising, given the high membrane turnover observed in photoreceptors required to maintain homeostasis [54–57]. Although the correlation between a given mutation and the human disease is often well established, we know very little about whether mutations in these genes affect cellular physiology/homeostasis before the onset of degeneration, and if so, how. We hypothesize that such alterations could render photoreceptors more susceptible to external stress factors and hence prone to degeneration.

Using the fly eye as a system to study the genetic control preventing retinal degeneration, we explored the effect of a mutation in $Drosophila\ crumbs\ (crb)$ on the organization of the endolysosomal system in the retina prior to any sign of degeneration. $Drosophila\ crb$ is a well-established model for Leber's congenital amaurosis and RP12-linked retinitis pigmentosa, two severe forms of retinal dystrophy leading to blindness. The recently identified hypomorphic crb^{P13A9} allele of Drosophila used in this study is particularly suited for these analyses, since unlike other crb alleles, it is homozygous viable, shows no developmental retinal phenotype and undergoes retinal degeneration only under constant light stress [32].

The major observations of our study were the following: i) Abnormal clustering of Arl8 was observed in retinas of crb^{P13A9} mutant flies under normal light conditions (12 hours dark/12 hours light). Unexpectedly, similar looking clusters also appeared in photoreceptors of w^* controls, but only under constant light stress. ii) Coincidence of Arl8-positive compartments with late endosomal Rab7 increased under constant light stress in both genotypes (but significantly less so in crb). In addition, crb^{P13A9} mutant photoreceptors kept in constant light

formed large, balloon-like vesicular structures that contained Rab7 on the membrane, but only small foci of Arl8. iii) Finally, coincidence of Arl8-positive compartments with autophagosomal Atg8mCherry decreased in crb^{Pl3A9} under normal light conditions. Taken together, data presented here allow us to conclude that retinas with reduced crb function exhibit a "pre-degenerative" endolysosomal phenotype, entailing changes in the association of Arl8 with two of its known interactors, Rab7 and autophagosomes. It should be noted that these defects occur prior to any obvious signs of degeneration, such as rhabdomere loss.

These results raise several questions: How does a reduced function of crb influence localization of endolysosomal small GTPases and their associated compartments? To what extent do the morphological alterations of the endolysosomal system affect its function? How do the changes observed make crb^{P13A9} mutant photoreceptors prone to light-induced degeneration?

Degeneration in the vertebrate retina has been subdivided into three phases, with the first phase characterized by increased cell stress due to protein mis-localization (reviewed in [1]). Our data suggest that Drosophila retinas mutant for crb reveal a pre-degenerative (or degeneration-prone) phenotype when kept under normal light conditions. At least two mechanisms, which have previously been linked to the *crb* mutant phenotype, could promote increased cell stress and hence could be responsible for the induction of such a predegenerative state. First, complete loss of crb function causes accumulation of intracellular rhodopsin-1 (Rh1)-containing vesicles due to impaired Rh1-transport to the rhabdomere [36]. Defective anterograde trafficking of rhodopsin is a well-established feature associated with retinal degeneration also in the vertebrate retina [57, 58]. Second, loss of crb induces an upregulation of NADPH-oxidase activity, resulting in higher levels of reactive oxygen species (ROS) [30]. Elevated ROS levels play a critical role in the pathogenesis of many neurodegenerative diseases, including retinal degeneration [59-62]. There is increasing evidence suggesting a crucial role of ROS in activating autophagy, one of the main degradative pathways critical not only for neuronal health, but also for the survival of other cells, e. g. muscles [60, 63]. During autophagy, engulfed organelles, nutrients, and protein aggregates are delivered to the lysosome to be broken down and degraded or eventually recycled [64-67]. Autophagy is necessary to maintain retinal integrity, and upon its impairment, retinal degeneration occurs in the fly eye, with rhodopsin accumulating in Rab7positive compartments [15]. However, we did not observe marked co-localization of Rh1-positive vesicles with Arl8 under either normal or light stress conditions in crb^{P13A9} mutant retinas, suggesting that the defects observed, i. e. the modification of the intracellular distribution of the Arl8-, Rab7-, and Atg8-positive compartments, represent another aspect of the mutant phenotype.

Both Arl8 and Rab7 are involved in autophagosome-lysosome fusion, a process that is influenced, among other factors, by the positioning of these compartments [19, 49, 50, 68] (reviewed in [23, 69]). In HeLa cells, for example, lysosomes concentrated in either a perinuclear pool or in the periphery differ in their pH and hence in their functionality [25]. Similarly, the change in positioning of the Arl8- and Rab7-positive compartments under stress (constant light in control flies and 12 h light/ 12 h dark in crb mutant flies) and the transformation of the small Rab7-positive compartments into large, balloon-like vesicles with only small foci of Arl8 could indicate a change in their functionality. Likewise, the formation of large Atg8-mCherry clusters and their reduced co-localization with Arl8 in crb^{P13A9} mutant retinas suggest a possible change in function of the autophagosome. Since autophagy is required for degradation of light-activated Rh1 in order to prevent retinal degeneration [15, 70], we tentatively propose that the pre-degenerative state of *crb* mutant retinas is defined by reduced autophagosomal activity, which makes cells prone to degeneration upon increased light stress. Future work will reveal whether the displacement of the Arl8-positive compartments and morphological change in the Rab7-positive vesicles are associated with defects in the degradative machinery, which in turn promotes degeneration [71].

Intracellular positioning and fusion of lysosome depends on both microtubules and the actomyosin network [72]. This raises the question whether Crb might indirectly be involved in regulating lysosomal positioning and function by controlling cytoskeletal organization. In wild-type photoreceptors, Crb protein is restricted to the stalk membrane, a portion of the apical plasma membrane between the rhabdomere and the adherens junctions [31, 38]. The stalk membrane borders a specialized cortical cytoplasm abutting the base of the rhabdomere, called ectoplasm, which is defined by an actin-rich terminal web also loaded with MyosinV. The ectoplasm is involved in shaping the subrhabdomeral cisternae (SRC), a distinct ER compartment next to the rhabdomere base [73]. Interestingly, proximo-distal alignment of

Arl8 in control retinas is just next to the base of the rhabdomere, suggesting a localization at or close to the SRC.

In photoreceptors completely lacking crb function or mutant for crb^{P13A9} the stalk membrane is reduced by about 40% and 20%, respectively [32, 38]. In addition, the amount of β_{H} -spectrin at the stalk membrane is strongly reduced in loss of function alleles [38]. The spectrin cytoskeleton recruits microtubule-binding proteins to the apical pole, such as Spectraplakin (called Shortstop in flies) [74, 75]. In this way, the ectoplasm facilitates the movement of cargo, which is required for building the rhabdomere during development and its maintenance in the adult. These cargo include rhodopsin and other molecules destined for the rhabdomeric membrane, but also, notably, pigment granules. Pigment granules are lysosome-related organelles, whose biogenesis and transport pathways share that of lysosomes [40, 76, 77]. In strong light, they align along the rhabdomere base, where they act as a shield from light stress [73]. Since our experiments were performed in a w background, the effect of crb reduction on pigment granule migration could not be analyzed.

Taken together, we hypothesize that reduced *crb* function results in impaired organization and/or function of the ectoplasm and the SRC. This influences the spatial organization of the endolysosomal system, which, in turn, affects Rab7-Arl8 coordination. These alterations may predispose the hypomorphic *crb* retina toward a degeneration-prone state, which leads to degeneration when subjected to light stress.

Acknowldgements

The authors would like to thank Sarita Hebbar, Ruchi Jhonsa, and Kassiani Skouloudaki for helpful discussions, Robert Haase and Benoit Lombardot from the MPI Scientific Computing Facility for the spot analysis script and help with image analysis, Britta Schroth-Diez and the MPI imaging facility for help with spinning disk and confocal imaging, and Jona Drushku for initial screening of compartment markers in retina. This work was supported by the Max-Planck Society. The authors declare that all data and material reported in this manuscript can be made available upon request, and that they have no competing interests.

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Figure legends

Fig. 1. The distribution of Arl8-positive lysosomal compartments depends on light conditions in w^* control photoreceptor cells.

- (**A-B'**) Lysosomal compartment marked with Arl8 in longitudinal (**A, A'**) and cross-sectional (**B, B'**) optical sections of w^* retinas after 5 days exposure to 12h light/ 12h dark (half sun).
- (C-D') Lysosomal compartment marked with Arl8 in longitudinal (C, C') and cross-sectional (D, D') optical sections of w^* retinas after 3 days of constant bright light (full sun). Stalk membranes of photoreceptor cells in longitudinal sections are marked with anti-Crb (green; A, C), basolateral domains are labelled with anti-Na⁺-K⁺-ATPase in cross-sections (green; B, D), and actin-rich rhabdomeres are labelled with phalloidin (white; B, B', D, D'). Note that the alignment of the Arl8-positive vesicles along the base of the rhabdomeres under dark/light conditions is abolished under constant light. Scale bars: 5 μ m.

Fig. 2. Arl8-positive lysosomal compartments cluster abnormally in crb^{P13A9} mutant photoreceptor cells under normal light conditions.

- (A-B') Optical longitudinal sections of w^* retinas after 5 days of 12h light/12h dark conditions, stained with anti-Arl8 (red) and anti-Crb (green).
- (C-C'') Depiction of the procedure for quantification of size and intensity of Arl8 regions (see Methods section for details). Areas of longitudinally oriented ommatidia were selected as ROIs (thin teal lines), judged by anti-Crb (green). Arl8-positive regions (red; arrows, C') within the ROIs were determined by an automatically thresholded Find Maxima plugin to detect signal above background, yielding the "Arl8 regions" outlined in beige (C'').
- (**D**, **E**) Optical cross-sections of w^* (**D**) and crb^{PI3A9} (**E**) mutant retinas in the same conditions as above show positioning of Arl8 clusters (red) in crb^{PI3A9} with respect to rhabdomeres, which are labelled for actin (white).
- (**F**, **G**) Optical cross-sections of $w^*(\mathbf{F})$ and $crb^{P13A9}(\mathbf{G})$ mutant retinas in the same conditions as above, labelled for basolateral domains (Na⁺-K⁺-ATPase; white) and rhabdomeres (actin; blue), where Arl8-positive clusters (red) can be detected in photoreceptors (arrow) and support cells (arrowheads). (**G**) is a montage from two different images of crb^{P13A9} rhabdomeres. Scale bar: 5 μ m.
- (H) Comparison of the size and intensity (brightness) of Arl8 regions in w^* (gray box plots) and crb^{Pl3A9} (orange box plots) retinas, with averaged data from each image stack shown by a dot (for detailed procedure, see Methods). Under 5 days of normal light/dark conditions, Arl8-positive clusters are detected as larger and brighter regions in crb^{Pl3A9} retinas (visible in C^*), compared to those in w^* retinas. Size and brightness values were normalized to w^* controls for both groups, and w^* and crb^{Pl3A9} retinas were always processed in parallel in

each experiment. Normalized data from up to three experiments were pooled for each box plot. P values for each pairwise comparison were calculated in RStudio using Student's t-test, and are encoded by stars as follows for this and subsequent plots: * P<0.01; **P<0.001; ***P<0.0001; ****P<0.00001; ****P<0.00001.

Fig. 3. Arl8 abnormalities in crb^{P13A9} retinas depend on light and dietary carotenoids.

- (A-C') Longitudinal optical sections of crb^{P13A9} retinas of flies kept for 5 days in 12h light/12h dark (A, A'), for 10 days in constant darkness (B, B'), or 10 days in 12h light/12h dark without carotenoids (-VitA) (C, C'), compared with w^* retinas (D, D') kept in the same conditions as (C), stained for Arl8 (red; gray in A'-D') and Crb (green). Scale bar: 5 μ m.
- (**E-G**) Quantification of Arl8-positive regions (defined as in Fig. 2) shows significant differences in size and intensity between w^* (gray box plots) and crb^{Pl3A9} (orange box plots) in retinas of flies kept for 10 days in the light conditions indicated below. The difference between control and mutant retinas was abolished when flies were raised in constant darkness for 10 days (**F**) or raised for 10 days in light/dark conditions, but without carotenoids in the food (**G**; -VitA). "ns" indicates not significant (P>0.01).

Fig. 4. Arl8- and Rh1-positive compartments are largely non-overlapping in crb^{Pl3A9} and w^* under normal light conditions.

Longitudinal optical sections of $w^*(\mathbf{A})$ and $crb^{P13A9}(\mathbf{B})$ retinas of flies kept for 5 days in 12h light/12h dark, stained for Arl8 (red), and Rh1 (green). Note that the mutant retina contains more Rh1-positive vesicles, which are rarely positive for Arl8 (arrowheads). Scale bar: 5 μ m.

Fig. 5. Exposure to constant light causes increased co-localization of Arl8 with late endolysosomal Rab7 and abnormal Rab7 compartment shapes in crb^{P13A9} .

- (A, C) w^* and crb^{P13A9} retinas, respectively, under 5 days normal light conditions (12h light, 12h dark) show some overlap of large endogenous Rab7-carrying vesicles (green) with Arl8 (red) (arrowhead). Additionally, many normal-sized compartments in w^* (A) and abnormal patches in crb^{P13A9} (C) are non-colocalizing (arrows), resulting in a relatively low, positive Pearson correlation coefficient (R coloc) of ~0.25 (on a scale of -1 to +1) for both genotypes over the whole retina (E, brown boxes).
- **(B, D)** Constant light stress causes an increase in co-localization between Rab7 and Arl8 (arrowheads) in w^* and crb^{P13A9} retinas, respectively (R coloc = \sim 0.5 for w^* ; \sim 0.33 for crb^{P13A9} ; **E**). Scale bars for **A-D**: 5 µm.

- **(E)** Quantification of Rab7 and Arl8 co-localization. 5 days of normal light conditions does not reveal any significant difference in Arl8/Rab7 co-localization (brown box plots). Two days of constant light stress causes a significant increase in the Pearson correlation coefficient in w^* , and a smaller but significant increase in crb^{P13A9} . Quantification was done on ROIs from image stacks, as described in Fig. 2.
- (**F**, **G**) Enlarged images showing numerous large round lysosomal compartments in w^* positive for both Arl8 and Rab7. In contrast, large, distended Rab7-positive compartments in crb^{P13A9} can be seen in contact with small foci of Arl8 (**G**). Scale bars: 1 μ m.

Fig. 6. Autophago-lysosomal marker Atg8-mCherry loses its association with Arl8 in crb^{PI3A9} mutant retinas.

- (**A-B'**) Longitudinal optical sections of $w^*(\mathbf{A}, \mathbf{A'})$ and $crb^{P13A9}(\mathbf{B}, \mathbf{B'})$ retinas of flies expressing Rh1-Gal4-mediated Atg8-mCherry (red). Retinas of flies kept for 6 days at 12h light/12h dark show differences in morphology of autophago-lysosomes in $crb^{P13A9}(\mathbf{B}, \mathbf{B'})$. Rhabdomeres are labelled for F-actin (phalloidin; white). Scale bar: 5 µm.
- (C, D) Examples of single autophagosomal Atg8-mCherry compartments (red) in w^* (C) and crb^{P13A9} (D). Atg8-mCherry is often directly adjacent to or surrounded by Arl8 (green) in the control (C), whereas it is non-overlapping with Arl8 in crb^{P13A9} (D). Scale bars: 1 μ m.
- **(E)** Quantification showing loss of Atg8-mCherry co-localization with Arl8 in crb^{P13A9} in individual compartments (n >25 compartments similar to those shown in **C** and **D**) measured by Manders' thresholded co-localization coefficient tM2 for the Atg8-mCherry channel.

Fig. S1. The Arl8-positive compartment partially overlaps with late endosomal and autophagosomal markers.

Longitudinal optical sections of w^* retinas after 5 days of 12h light/12h dark conditions, stained with anti-Arl8 (red in all panels) and other lysosomal markers (green in all panels).

- (A, B) Arl8 occupies the same compartment as endogenous Rab7 (A) and Rh1-Gal4-mediated Rab7-GFP (B) in a sub-population of vesicles.
- **(C)** Arl8-positive compartments are often nearby, but non-identical to those labelled by Atg8-mCherry (green), an autophagosomal marker expressed in all photoreceptors via *Otd*-Gal4.
- **(D)** Arl8-positive compartments are often nearby, but non-identical with those labelled by the putative lysosomal transporter spinster (spin)-GFP (green), expressed by *Rh1*-Gal4 in photoreceptors R1-6.

- **(E)** Arl8-positive compartments are similarly adjacent, but largely non-identical to those labelled by the *bona fide* lysosomal marker, LAMP-GFP, expressed by *Rh1*-Gal4 in photoreceptors R1-6.
- **(F)** Arl8-positive compartments are often nearby, but distinct from those carrying an intrinsic lysosomal marker, a GFP gene-trap insertion in the PIP₂ phosphatase gene (Lyso PIP2-ase), CG6707.

Scale bars: 5 µm.

Fig. S2. Arl8 compartments overlap little with Rh1 even under constant light stress.

Longitudinal optical sections of w^* and crb^{P13A9} retinas of flies kept for 6 days in constant light, stained for Arl8 (red) and Rh1 (green). Very few small Arl8-positive compartments are also Rh1-positive (**A-A''**; arrowhead), as is the case in normal light conditions (see Fig. 4). In contrast, large Arl8-positive patches in crb^{P13A9} are always negative for Rh1 (**B**).

Fig. S3. Constant light stress reduces differences in Arl8 compartment size and intensity between w^* and crb^{P13A9} .

Arl8 compartments, quantified as in Figs. 2 and 3, show no significant differences in size and fluorescence intensity between w^* (gray box plots) and crb^{P13A9} (orange box plots) after two (left graph) or six days (right graph) of constant intense light stress. "ns" indicates not significant (P>0.01).

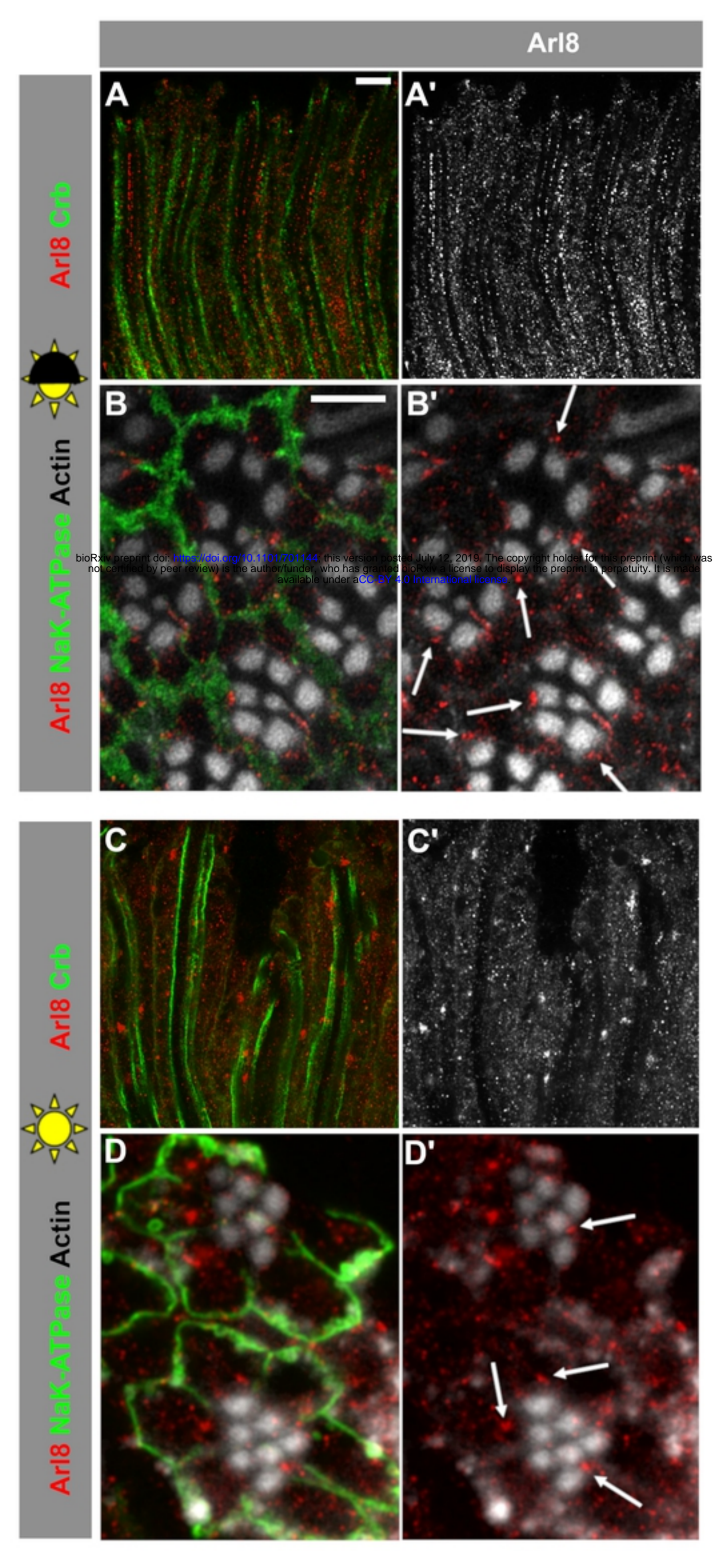


Fig. 1

Figure 1

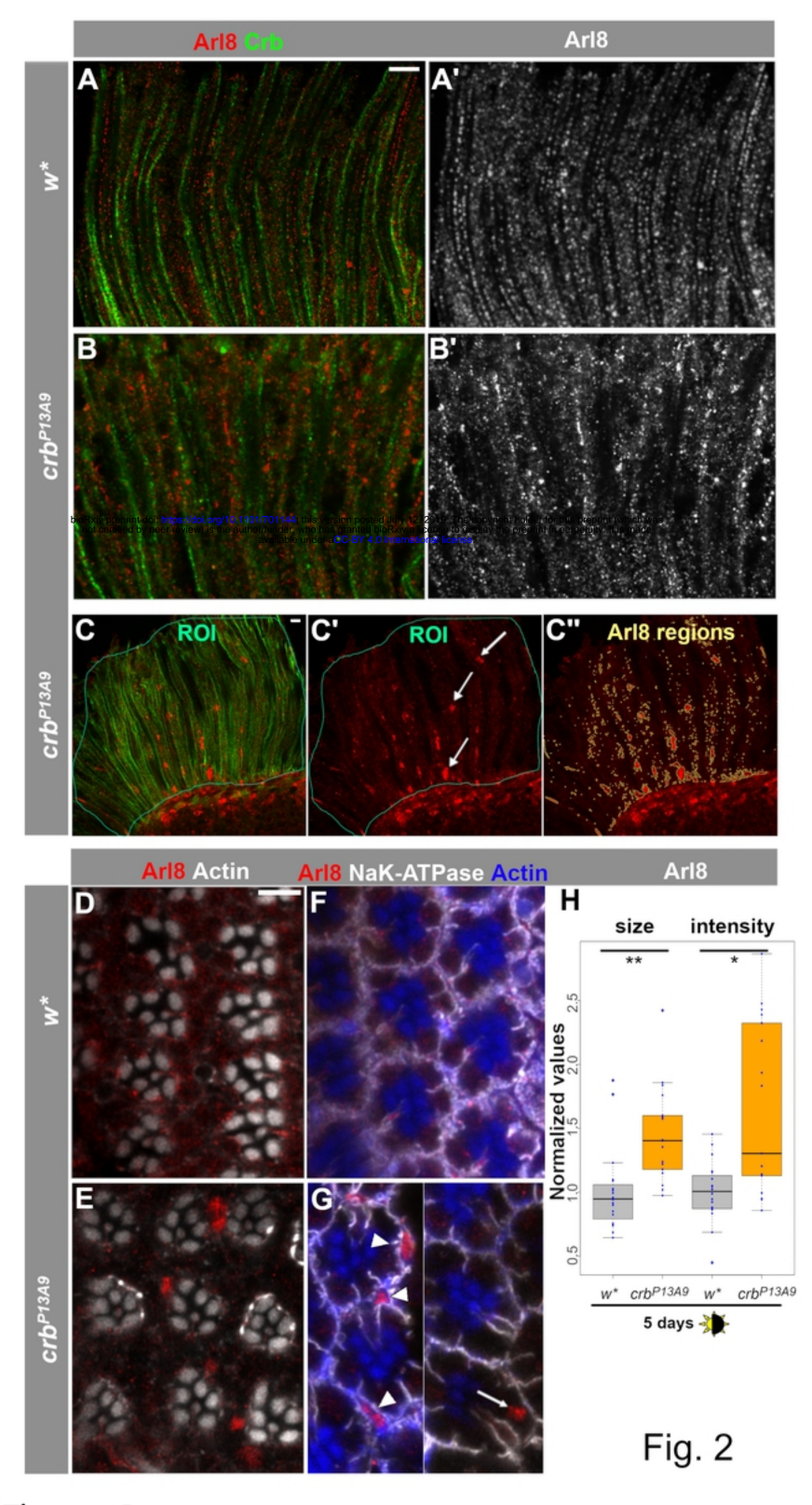


Figure 2

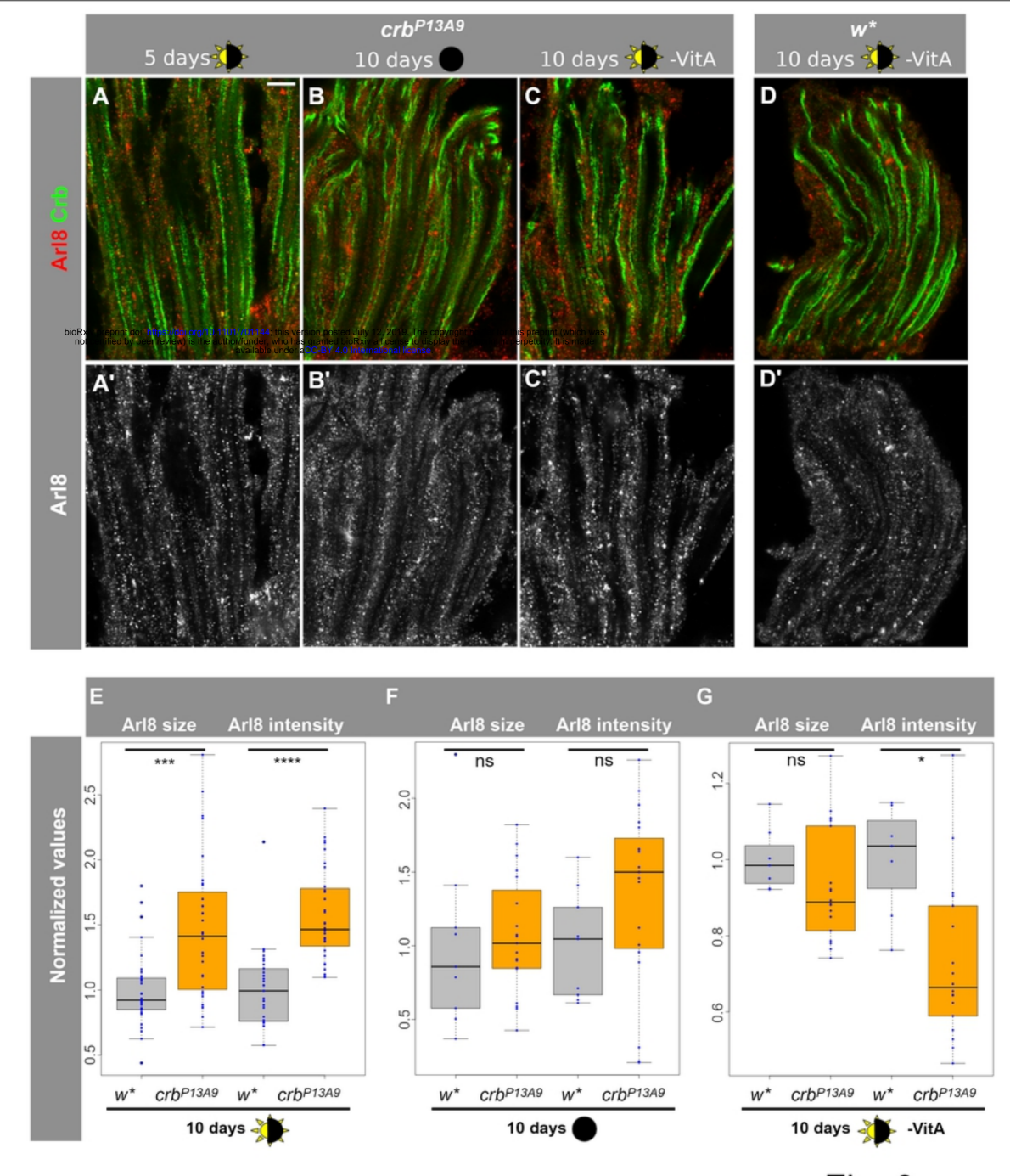


Fig. 3

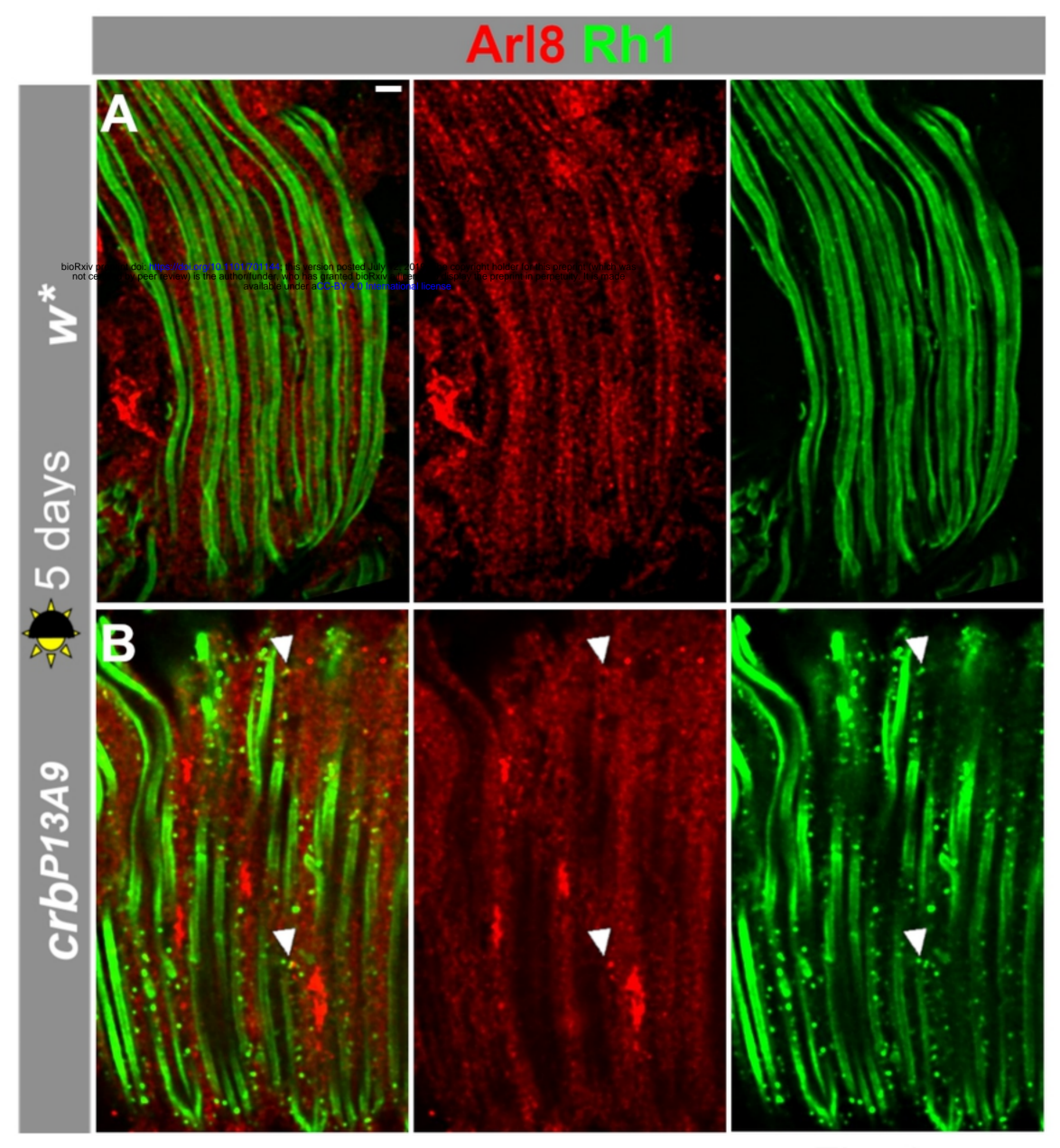


Fig. 4

Fig. 5

Figure 5

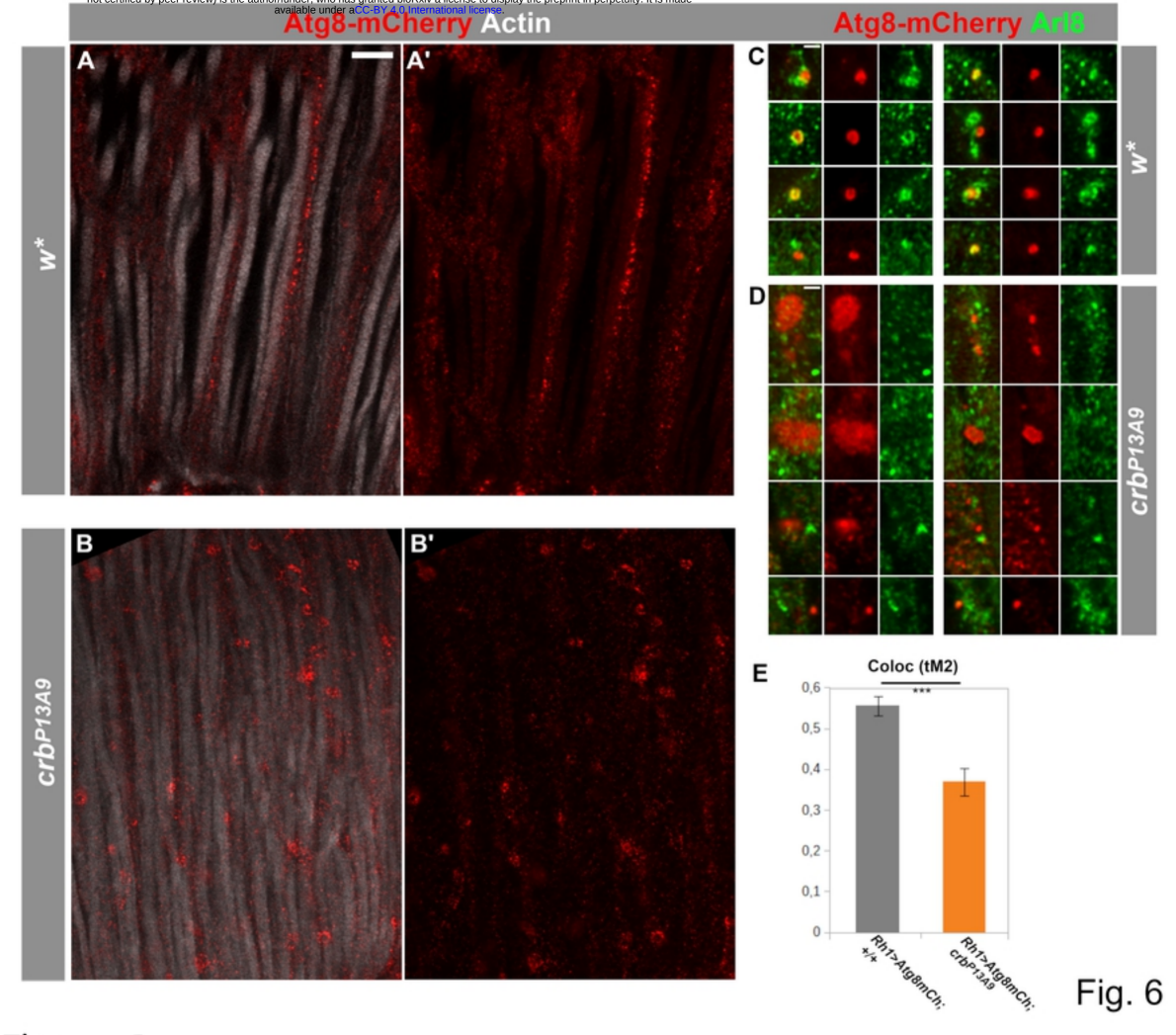


Figure 6

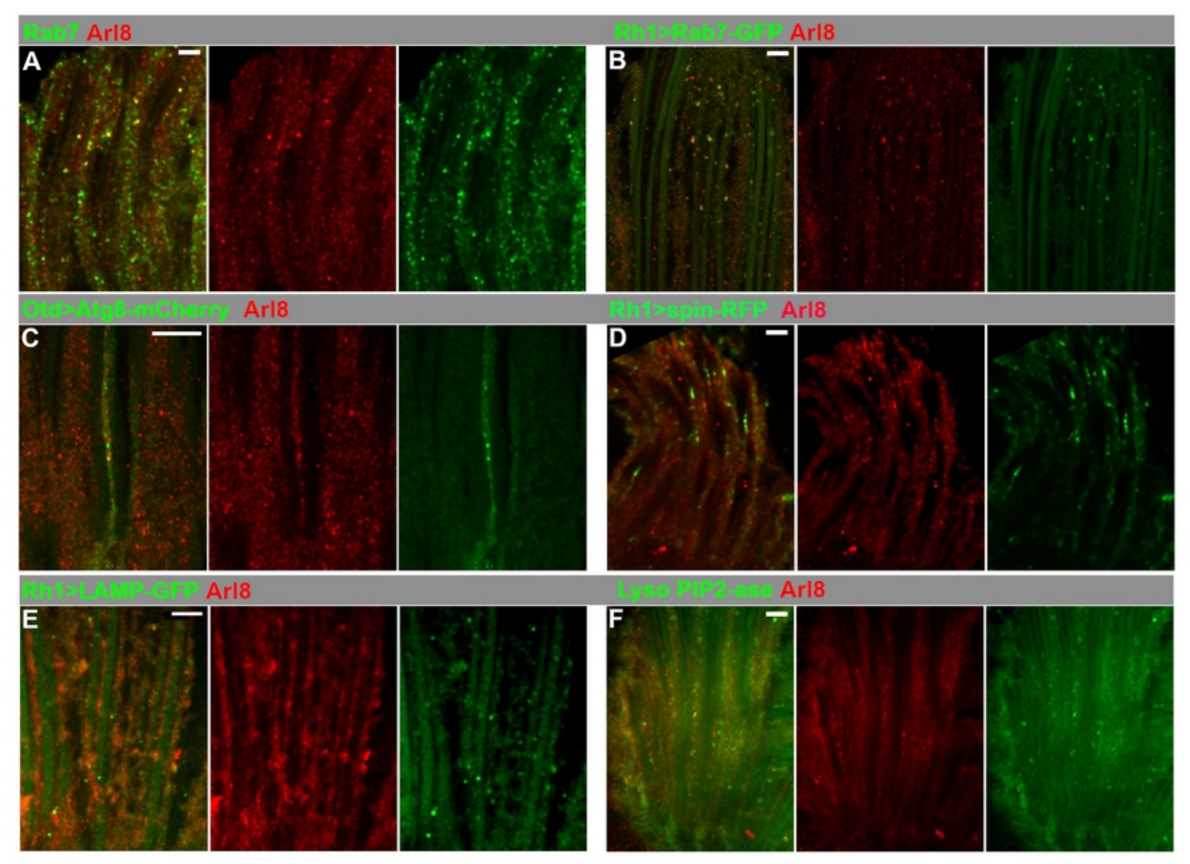


Fig. S1

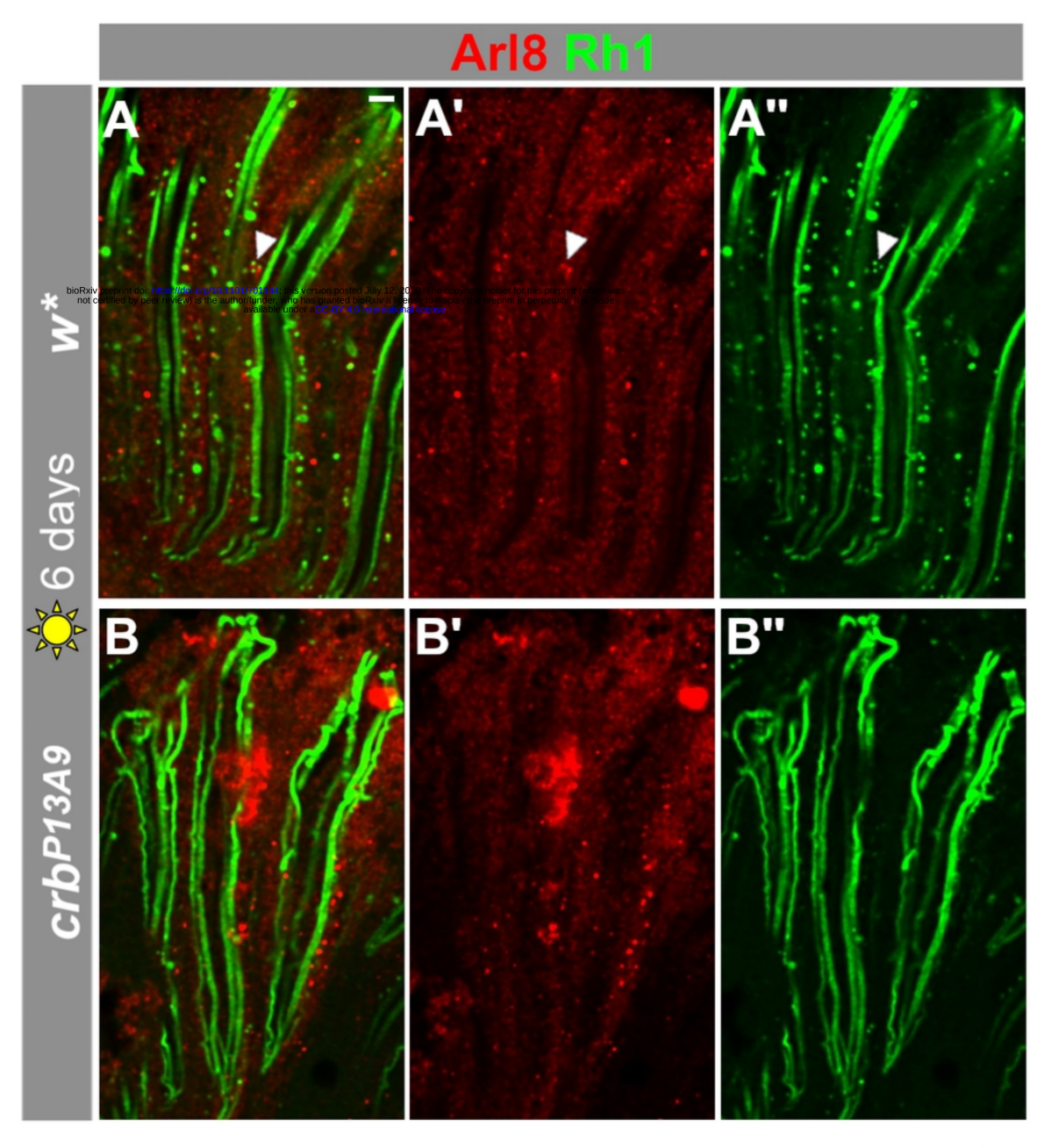


Fig. S2

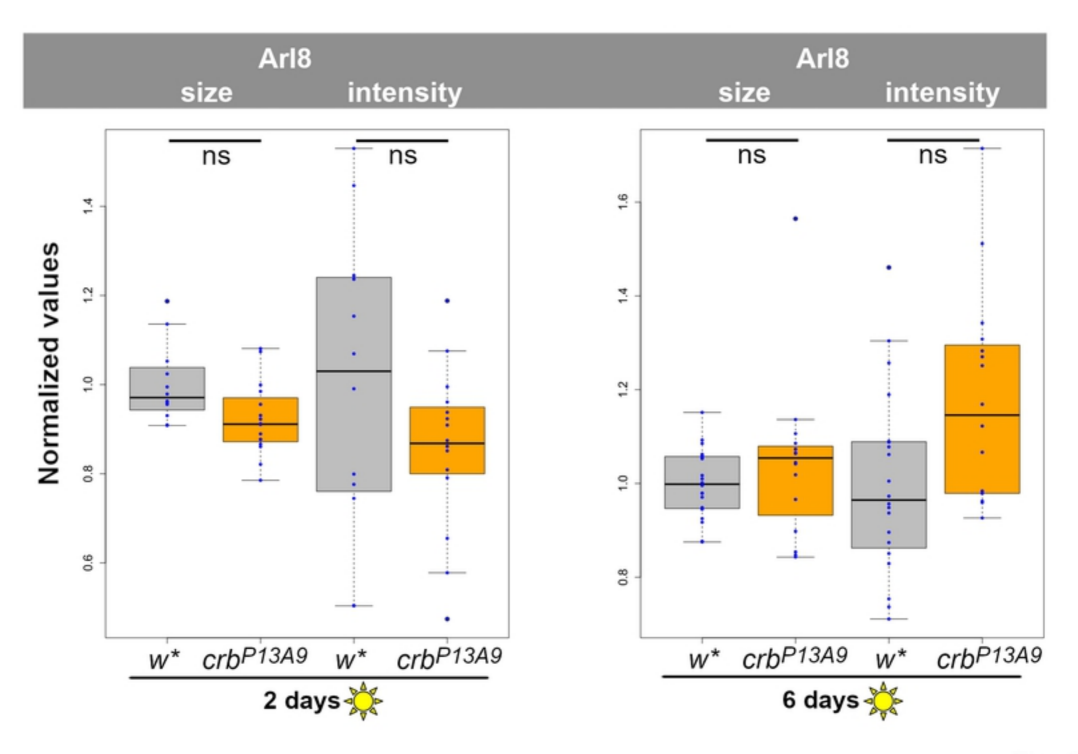


Fig. S3