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Permalink
https://escholarship.org/uc/item/9d00d6sf

Journal
PloS one, 10(11)

ISSN
1932-6203

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Publication Date
2015

DOI
10.1371/journal.pone.0143026

Peer reviewed
Identification of Atg2 and ArfGAP1 as Candidate Genetic Modifiers of the Eye Pigmentation Phenotype of Adaptor Protein-3 (AP-3) Mutants in Drosophila melanogaster

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Abstract

The Adaptor Protein (AP)-3 complex is an evolutionary conserved, molecular sorting device that mediates the intracellular trafficking of proteins to lysosomes and related organelles. Genetic defects in AP-3 subunits lead to impaired biogenesis of lysosome-related organelles (LROs) such as mammalian melanosomes and insect eye pigment granules. In this work, we have performed a forward screening for genetic modifiers of AP-3 function in the fruit fly, Drosophila melanogaster. Specifically, we have tested collections of large multi-gene deletions—which together covered most of the autosomal chromosomes—to identify chromosomal regions that, when deleted in single copy, enhanced or ameliorated the eye pigmentation phenotype of two independent AP-3 subunit mutants. Fine-mapping led us to define two non-overlapping, relatively small critical regions within fly chromosome 3. The first critical region included the Atg2 gene, which encodes a conserved protein involved in autophagy. Loss of one functional copy of Atg2 ameliorated the pigmentation defects of mutants in AP-3 subunits as well as in two other genes previously implicated in LRO biogenesis, namely Blos1 and lightoid, and even increased the eye pigment content of wild-type flies. The second critical region included the ArfGAP1 gene, which encodes a conserved GTPase-activating protein with specificity towards GTPases of the Arf family. Loss of a single functional copy of the ArfGAP1 gene ameliorated the pigmentation phenotype of AP-3 mutants but did not to modify the eye pigmentation of wild-type flies or mutants in Blos1 or lightoid. Strikingly, loss of the second functional copy of the gene did not modify the phenotype of AP-3 mutants any further but elicited early lethality in males and abnormal eye morphology when combined with mutations in Blos1 and lightoid, respectively. These results provide genetic evidence for new functional links connecting the machinery for biogenesis of LROs with molecules implicated in autophagy and small GTPase regulation.
Introduction

The Adaptor Protein (AP)-3 complex is a component of the basic molecular machinery that mediates the trafficking of integral membrane proteins within the endosomal-lysosomal system [1]. It belongs to a family of heterotetrameric complexes, AP-1 through -5 and COPI-F, which originated through gene duplication events that took place before divergence of the last common ancestor of all modern eukaryotes [2]. The AP-3 complex, which is considered the most ancient of the five AP complexes [2], is composed of δ, β3, μ3 and σ3 subunits [1,3]. In mammals, each the β3, μ3 and σ3 subunits is encoded by a pair of paralog genes. Mutations in the gene encoding one of the two variants of the β3 subunit in humans cause Hermansky-Pudlak syndrome (HPS) type 2 [4–11]. Like all other forms of autosomal-recessive HPS, HPS type 2 is characterized by clinical manifestations that originate from defective formation of cell-type-specific, and functionally specialized, lysosome-related organelles (LROs). Thus, all forms of HPS are associated with oculocutaneous albinism and bleeding diathesis owing to abnormal biogenesis of melanosomes and platelet dense granules, respectively [12]. In addition, patients suffering from HPS type 2 are prone to recurrent infections owing to defective LROs in various cells of the innate immune systems [5–11].

Mechanistically, the AP-3 functions as a ‘sorting device’ that, upon recruitment to endosomal membranes from a soluble cytoplasmic pool, recognizes targeting determinants within the cytoplasmic aspect of integral membrane proteins [13]. This molecular interaction results in the incorporation of ‘cargo’ integral membrane proteins into membrane-bounded carriers, which are formed through membrane budding driven by assembly of AP-3 with other cytosolic factors into a protein coat [3,13–15]. The target organelles of this AP-3-dependent trafficking pathway, identified mainly through analyses of AP-3-deficient cells derived from HPS-2 patients and mouse models, turned out to depend on cell type [1,15]. For example, in fibroblasts and other mammalian cell types lacking LROs, AP-3 mediates delivery of integral membrane proteins to lysosomes [4,5,16–19], while in melanocytes it defines a route for the delivery of melanin-synthesizing enzymes to maturing melanosomes [20–22].

Several of the factors that cooperate with AP-3 in the trafficking of integral membrane proteins to lysosomes or LROs have been identified through protein-protein interaction analyses, either targeted towards candidates based on prior biological evidence or unbiased proteomic methods [23–28]. While successful, such approach is intrinsically limited to proteins that interact directly with AP-3 or indirectly through other components of the interaction network. Alternative approaches, such as those based on the analysis of genetic interactions, may uncover functional relationships–within the AP-3-dependent trafficking pathway or with other cellular activities–regardless of the existence or absence of direct physical interactions. However, few non-additive genetic interactions involving the AP-3 complex have been reported in metazoans. In mice, phenotypic enhancement effects were observed for homozygous double mutant mice deficient in AP-3 and each of three protein complexes named Biogenesis of Lysosome-related Organelles Complex (BLOC)-1 through -3 [26,29,30], which like AP-3 are involved in the biogenesis of LROs and associated with various forms of human HPS [12], as well as in double mutant mice deficient in AP-3 and OCA2 [31], with the latter being the product of the gene mutated in oculocutaneous albinism type 2 [32]. Understandably, these genetic interaction studies were limited to targeted approaches owing to the practical limitations of inter-breeding multiple mouse lines. Phenotypic enhancement (possibly additive) effects have also been observed in the worm, Caenorhabditis elegans [35–36], and the fruit fly, Drosophila melanogaster [37–39], through targeted analyses of double mutants deficient in the AP-3 complex and other factors previously implicated in LRO biogenesis. Additional targeted approaches have led to the observations of synthetic lethality in double-mutant worms lacking
AP-3 and the endocytic protein Disabled [40] as well as non-additive phenotypic enhancement effects in AP-3-deficient flies carrying a weak allele of white (w) [41] or misexpressing the endocytic protein Auxilin [39]. Finally, AP-3 mutant alleles have been identified as genetic modifiers of morphological phenotypes induced by misexpression in flies of Blue Cheese [42], which is a protein involved in selective disposal of ubiquitinated protein aggregates [43], the E3 ubiquitin ligase Deltex [44], and human β-amyloid peptide [45].

In this paper, we systematically searched for genetic modifiers of AP-3 function in the fly eye. Prior to the demonstration that mutations in genes encoding AP-3 subunits cause HPS-2 in humans [4] and related phenotypes in mouse models [17,46,47], the Drosophila gene garnet (g) had been shown to encode the fly counterpart of the δ subunit of AP-3 and to be required for normal biogenesis of eye pigment granules [48,49], which like mammalian melanosomes are LROs [50]. Using a hypomorphic allele of this gene for screening, we have identified distinct chromosomal regions on chromosome 3, containing the genes Atg2 and ArfGAP1, which upon gene dosage reduction ameliorated the eye pigmentation phenotype caused by AP-3 deficiency. To our knowledge, this is the first report of an unbiased, forward screening for genetic modifiers of AP-3 function.

Materials and Methods

Fly strains and collections

Flies were raised at 25°C using standard husbandry procedures [51]. The fly line g2, which was derived by multiple outcrosses of the triple mutant strain y1 wy2 g2 (from the Bloomington Drosophila Stock Center) into the genetic background of Canton-S, was kindly provided by Dr. David E. Krantz (University of California, Los Angeles, CA, USA). The fly strain ArfGAP1G3-85 (formerly Gap69CG3-85) was kindly provided by Dr. Vladimir E. Alatortsev (Russian Academy of Sciences, Moscow, Russia) [52]. The mutant fly line blos1ex2 had been generated in our laboratory [39]. All other fly strains, including Canton-S (used as ‘wild-type’ control), eyg1, ltd1, and those of the "classic" Bloomington Deficiency Kit for chromosomes 2 and 3 and the Bloomington Deficiency Kit of molecularly defined deletions for chromosome 4 [53], were obtained from the Bloomington Drosophila Stock Center.

Fly eye microscopy and imaging

Fly eyes were visualized on live adults anesthetized with CO2 using a Zeiss Stemi 2000-C Stereo microscope (Carl Zeiss, Thornwood, NY, USA). Images were acquired using a Nikon Coolpix P5000 digital camera adapted to the same microscope.

Quantification of eye pigments

Red (pteridines) and brown (ommochromes) pigments were extracted from pooled heads of four (for red pigments) or eight (for brown pigments) male flies at ~3 days after eclosion (range 2–5 days) and quantified as previously described [38]. In each experiment, pigment content values were normalized to those of White-null (w1118) flies and wild-type (Canton-S) flies, which were quantified in parallel (2–3 biological replicates per experiment) and served as 0% and 100% controls, respectively.

Statistical analyses

Statistical analyses were performed using GraphPad Prism 5.0b (GraphPad Software, San Diego, CA, USA). Comparisons between more than two groups of data were carried out using one-way ANOVA followed by either Dunnett’s (for comparison of multiple data groups to a
Results and Discussion

Primary and secondary screening for genomic regions carrying modifiers of AP-3 function

In order to identify regions within the autosomal chromosomes of *D. melanogaster* (chromosomes 2, 3 and 4) potentially bearing genes that, when in hemizygous form due to deletion of one copy, modify the severity of the phenotype of AP-3-mutant flies, a genetic screening was undertaken following the strategy depicted in Fig 1. In both primary (qualitative) and secondary (quantitative) steps of the screening, male flies carrying large multi-gene deletions (‘deficiencies’) from the so-called ‘Bloomington Deficiency Kit’ collections were crossed with female flies homozygous for the *g*2 mutation (on chromosome X) to obtain *F*1 males that were hemizygous for *g*2 and heterozygous for a given deficiency over a normal chromosome; for each deficiency, the eye color of these *F*1 males was compared to that of control *g*2 males first by light microscopy and, if selected for secondary screening, by pigment extraction and quantification (Fig 1). The *g*2 mutation in the gene encoding the δ subunit of AP-3 was chosen not only because of its genomic location on the X chromosome (thus minimizing the number of genetic crosses required to test each deficiency) but also because it represents a weak hypomorph of the *garnet* allelic series [54]. In our hands, the eyes of male adult flies carrying the *g*2 allele in the Canton-S genetic background contain 25–30% of the red pigment content of those of the control Canton-S line, whereas those of males carrying one of the strongest alleles of the same gene, *g*3<sub>d</sub>, contain less than 5% of wild-type levels of red pigments (see, for example, Ref. [39]). Thus, the *g*2 line was used as a sensitized strain with which both phenotypic enhancers and suppressors could potentially be identified.

Out of 213 deficiency lines screened, which together covered >92% of fly autosomal chromosomes 2–4, twenty were selected for secondary screening on the basis of apparent differences in eye color of the corresponding *F*1 males (*g*2 hemizygous carrying a copy of the deficiency) relative to *g*2 males upon visualization under a dissecting microscope. Of these twenty candidates, seven resulted in statistically significant changes in red pigment levels, upon correction for multiple testing, with all of these changes representing increased pigment content in the eyes of the corresponding *F*1 males relative to those of *g*2 controls (Fig 2). It is possible that some of the remaining candidate deficiencies might have modified the eye color phenotype of *g*2 mutants through quantitative changes in brown pigment levels only. Nevertheless, given that our goal was to identify modifier genes that might act at the level of pigment granule biogenesis and/or dynamics, thus affecting the accumulation of both pigment types in these granules, those candidate deficiencies that failed to elicit significant changes in red pigment content were not characterized any further. In addition, one of the seven candidates resulting in statistically significant differences, *Df(2R)PC4*, was excluded from further analysis because the relative increase in red pigment content was less than a threshold of 50% increase, which we had arbitrarily set to focus our efforts on the most promising candidates (Fig 2).

The six selected deficiencies were then tested for their ability to modify, when carried in single copy, the brown pigment content of *g*2 flies (to verify that both pigment types, not just red pigments, were affected) as well as the red pigment content of *rb*1 flies (which carry a hypomorphic mutation in the gene encoding the β3 subunit of AP-3 [55,56]) and wild-type (Canton-S) flies. The results of these experiments are listed in Table 1. Upon correction for multiple testing, the statistical analyses suggested that: (*i*) each of the six deficiencies modified the pigmentation phenotype of an independent AP-3 mutant (*rb*1); (*ii*) four of them also altered the
Fig 1. Strategy for the genetic modifier screening carried out in this study. The primary screening involved parental (P₀) crosses between male flies carrying a deficiency (Df) over a balancer in an autosomal chromosome (Chr. 2, 3 or 4) and female flies homozygous for the hypomorphic g² allele of the X-linked gene encoding the δ subunit of AP-3. The eye color of male progeny (F₁) carrying one copy of each deficiency (without the balancer) was compared with that of control g² males and, if deemed different, the corresponding deficiency was selected for a secondary screening involving the same P₀ cross followed by quantification of red and brown pigments in the F₁ males carrying the deficiency. In cases in which differences in both red and brown pigment content were statistically significant, further validation and fine mapping was attempted using independent deficiency lines in which the deleted genomic regions partially overlapped with that of the deficiency identified through screening. When successful, these steps allowed identification of a relatively small genomic region (rectangle) containing a modifier gene of interest (red arrow).

doi:10.1371/journal.pone.0143026.g001
Fig 2. Red pigment content in eyes of $g^{2}$ mutant flies carrying hemizygous deletions selected through primary screening. Red pigments were extracted from the heads of adult $g^{2}$ mutant males carrying no deletions (-) or a single copy of the indicated deficiencies, quantified as described under Materials and Methods, and expressed as percentages of the red pigment content of male flies of the wild-type line (Canton-S). Bars represent means + SD of 2–28 biological replicates. Grey bars denote values obtained for flies carrying the marker allele $w^{+mC}$ linked to the deficiency. Dashed lines indicate threshold values corresponding to 66.7% (2/3) and 150% (3/2) of the pigment content of control $g^{2}$ flies carrying no deletion (black bar). One-way ANOVA followed by Dunnett’s test of each group versus $g^{2}$ flies carrying no deletion: ***p<0.001.

doi:10.1371/journal.pone.0143026.g002
red pigment levels of wild-type flies; and (iii) all but one, Df(2R)CD21, modified the brown pigments levels of g2 flies (Table 1) in addition to modifying their red pigment content (Fig 2).

Together, the primary and secondary screening yielded one genomic region on chromosome 2, which is deleted in deficiency Df(2L)XE-3801, and four independent genomic regions on chromosome 3, which are deleted in deficiencies Df(3L)eygC1, Df(3R)Exel6195, Df(3L)ED4978, and Df(3L)BSC23, for validation and further characterization.

Attempts to validate candidate genomic regions deleted in deficiencies carrying the marker ‘mini-white’

Three of the five deficiencies selected for validation, namely Df(2L)XE-3801, Df(3R)Exel6195, and Df(3L)ED4978, carried the construct mini-white (w+mC) as a genetic marker. It is well known that expression of this construct, which represents a truncated version of the w gene lacking some regulatory sequences, varies significantly depending upon the site of chromosomal insertion [57]. Given that high levels of White protein overexpression had been reported to increase, albeit modestly, eye pigmentation of AP-3 δ mutants [41], the possibility of ‘false-positive’ hits in our screening owing to high expression of the w+mC construct deserved consideration. To begin to address this issue, genetic crosses were set up to generate male flies hemizygous for the White-null allele w1118 (on chromosome X) and heterozygous for each of the three deficiencies; in these flies, White function derived exclusively from expression of the w+mC construct. As shown in Fig 3A (arrow), the w+mC construct carried by one of the three deficiencies, Df(3L)ED4978, on a White-null background resulted in red pigment levels of almost 40% of wild-type levels. In further support of the notion that deficiency Df(3L)ED4978 might have represented a false-positive hit in our screening, none of two ‘overlapping’ deficiencies—with deletions that together covered the entire region deleted in Df(3L)ED4978—modified the eye pigmentation phenotype of g2 flies (Fig 3B). Although the w+mC construct carried by the other two deficiencies, Df(2L)XE-3801 and Df(3R)Exel6195, resulted in very little pigmentation on a White-null background (Fig 3A), attempts to validate their phenotypic modifier effect observed on the g2 background, using independent overlapping deficiencies, were nonetheless

Table 1. Pigment content in flies carrying deficiencies in different genetic backgrounds.

| Deficiency          | Brown pigment | Red pigment |
|---------------------|---------------|-------------|
|                     | g2 background | rb1 background | Canton-S background |
| —                   | 61.8 ± 2.8 (5) | 16.5 ± 1.7 (12) | 100 ± 8 (3) |
| Df(3L)eygC1        | 68.5 ± 3.7 (4)* | 33.7 ± 3.4 (6)** | 120 ± 8 (5)** |
| Df(2R)CB21         | 65.0 ± 4.8 (4)** | 26.1 ± 2.0 (8)** | 121 ± 13 (4)** |
| Df(3R)Exel6195     | 75.3 ± 1.2 (3)** | 31.0 ± 2.3 (7)** | 131 ± 12 (4)** |
| Df(3L)ED4978       | 74.3 ± 1.2 (3)** | 32.0 ± 3.7 (5)** | 128 ± 12 (4)** |
| Df(2L)XE-3801      | 68.8 ± 2.1 (3)* | 34.3 ± 3.4 (6)** | 123 ± 12 (5)* |
| Df(3L)BSC23        | 78.3 ± 4.5 (3)** | 39.2 ± 0.8 (9)** | 132 ± 12 (9)** |

Brown or red pigment content in the eyes of adult male flies carrying no deletions (—) or a single copy for each of the indicated deficiencies was measured as described under Materials and Methods and expressed as a percentage of the corresponding pigment content in the eyes of adult male flies of the control line Canton-S. Values represent means ± SD of the numbers of biological replicates indicated between parentheses. One-way ANOVA followed by Dunnett’s test of each group versus the corresponding control carrying no deficiency in the same background: NS, not significant

*p<0.05

**p<0.01

***p<0.001.

doi:10.1371/journal.pone.0143026.t001
Attempts to validate selected deficiencies carrying the \(w^{+mC}\) marker as genetic modifiers of the \(g^2\) mutation.

(A) Red pigments were extracted from the heads of AP-3-deficient \((g^2)\) or White-negative \((w^{1118})\) adult males carrying single copies of the indicated deficiencies with their associated \(w^{+mC}\) marker. The extracted pigments were quantified as described under Materials and Methods, and the resulting values expressed as percentages of the pigment content of wild-type (Canton-S) flies. Bars represent means + SD of 2–10 biological replicates. Notice that the activity of the \(w^{+mC}\) marker associated with deficiency \(Df(3L)ED4978\) resulted in a red pigment content (arrow) higher than that of \(g^2\) flies (black bar). (B-D) Analyses of red pigment content in the eyes of adult \(g^2\) mutant males carrying no deletions, single copies of the deficiencies \(Df(3L)ED4978\) (B), \(Df(3R)Exel6195\) (C) and \(Df(2L)XE-3801\) (D) that had been identified through screening, or single copies of deficiencies with partially overlapping deletions and devoid of the \(w^{+mC}\) marker. Schematic representations of the extent of overlap between the chromosomal regions deleted in the deficiencies identified through screening (blue) and the others (grey) are included in each figure panel. Notice in (C) that a small portion of the chromosomal region deleted in \(Df(3R)Exel6195\) (dashed box) did not overlap with any of those deleted in other available deficiencies. One-way ANOVA followed by Dunnett’s test of each group versus control \(g^2\) flies carrying no deletion (black bars): **\(p<0.01\); ***\(p<0.001\).

doi:10.1371/journal.pone.0143026.g003
unsuccessful (Fig 3C and 3D). It should be noted, however, that the deficiencies available for validation did not completely cover the genomic region deleted in Df(3R)Exel6195, thus leaving open the possibility that the partial phenotypic suppression effect elicited by this deficiency could have been caused by hemizygous deletion of a gene in the region that was not deleted in any of the other deficiencies tested (Fig 3C, dashed rectangle). Interestingly, this region turned out to contain a single gene, CG31145, which was reported to encode a Golgi-localized protein with casein-kinase activity [58]. Further investigation will be required to test the possibility of CG31145 being a genetic modifier of AP-3 function in the fly eye.

Validation and fine-mapping of the genomic regions responsible for the modifier effects elicited by Df(3L)eygC1 and Df(3L)BSC23

The partial suppressor effects elicited by deficiencies Df(3L)eygC1 and Df(3L)BSC23 on the eye pigmentation phenotype of AP-3-mutant flies could be successfully validated by independent deficiencies with partially overlapping deletions.

In the case of Df(3L)eygC1, four independent deficiencies were available to cover the entire genomic region deleted in the original deficiency, and two of them elicited partial suppressor effects on the phenotype of g2 flies that were akin to that observed for Df(3L)eygC1 (Fig 4). Although one of these two deficiencies, Df(3L)ED4483, carried the w+mC construct as a marker, expression of this construct was deemed low on the basis of the red pigment content of flies carrying such deficiency on a White-null background (Fig 3A). Through comparison of the genomic regions deleted in the deficiencies that elicited the modifier effect and those that failed to do so, a critical region comprising 17 annotated genes was defined (Fig 4, bottom box).

In the case of Df(3L)BSC23, seven independent deficiencies were available to cover the entire genomic region deleted in the original deficiency; three of them carried the w+mC construct (Fig 5), with expression levels that were deemed low to moderate on the basis of pigmentation of flies carrying them on a White-null background (Fig 3A). When assessed for their ability to modify the phenotype of g2 flies, only one overlapping deficiency carrying a ‘weak’ w+mC construct, Df(3L)Exel6091, and one deficiency lacking w+mC, Df(3L)BSC119, elicited a partial suppressor effect similar to that observed for the original deficiency (Fig 5). Through analysis of the genomic regions deleted in these deficiencies, a critical region encompassing six annotated genes was defined (Fig 5, left bottom box).

Practical limitations precluded a systematic investigation of each of the genes included in the critical regions derived from Df(3L)eygC1 and Df(3L)BSC23. These limitations were: the very limited availability of loss-of-function alleles of single genes, and the inconsistency of results that we obtained using available RNAi transgenes (expressed with the aid of a GMR-GAL4 driver [39]), with the latter likely arising from off-target effects [59]. For this reason, a candidate-gene approach was undertaken, as described in the following sections.

Atg2 as a genetic modifier of fly eye pigmentation

An obvious candidate out of the six genes located within the critical region derived from Df(3L)BSC23 was Atg2, which encodes a conserved protein involved in autophagy [60]. Genetic links between autophagy and pigmentation defects had been previously described in mammals [61] and flies [62], though none of them involved Atg2 in particular. A loss-of-function allele of this gene, Atg2EP3697, had been previously validated in assays of autophagy function in the larval fat body [63,64]. Homozygous Atg2EP3697 flies die before reaching adulthood [60], whereas heterozygotes are viable with no apparent eye morphological phenotypes. Although the Atg2EP3697 allele carries a copy of the w+mC construct, its expression was deemed to be quite
Fig 4. Validation and fine mapping of the critical region responsible for the modifier effect observed for Df(3L)eygC1. Red pigments were extracted from the heads of adult g^+^ mutant males carrying no deletions (---) or a single copy of the indicated deficiencies, quantified as described under Materials and...
low on the basis of the red pigment content of White-null flies carrying one copy of the allele (Fig 6).

Satisfyingly, a single copy of the Atg2EP3697 allele (in heterozygous form over a normal third chromosome) elicited statistically significant increases in the eye pigment levels of the AP-3 mutants g2 and rb1 (Fig 6). A single copy of the same allele also increased the red pigment content of Canton-S flies (Fig 6), an effect which was in line with that originally observed for the Df(3L)BSC23 deficiency (Table 1). In these three genetic backgrounds, the effect of the Atg2EP3697 allele on eye pigmentation was less pronounced than that observed for Df(3L)BSC23; we speculate that either Atg2EP3697 might not represent a true null allele of this gene or that other genes within the critical region might have contributed to the effect observed for the deficiency.

Next, we tested for effects of the Atg2EP3697 allele on the phenotypes of another two mutants of the pigment granule group, namely blos1ex2 and ltd1. The blos1 gene encodes a subunit of the fly counterpart of mammalian BLOC-1 [39,65], which like AP-3 is involved in intracellular protein trafficking and the biogenesis of LROs [22,25,26,66–68]. Although the two complexes have been shown to interact physically with each other [25–27,69–71], epistatic analyses in mice and flies have suggested that BLOC-1 and AP-3 can function, at least in part, independently of each other [26,30,39]. The lightoid (ltd) gene encodes the single fly counterpart of two closely related mammalian GTPases of the Rab family, named Rab32 and Rab38; both mammalian proteins as well as the Lightoid protein are involved in the biogenesis of LROs [37,72–74]. While mammalian Rab32 and Rab38 have been shown to interact physically and functionally with AP-3 [28], epistatic analyses have suggested that fly Lightoid and AP-3 can function, at least in part, independently of each other [37]. As shown in Fig 6, a single copy of the Atg2EP3697 allele elicited small but statistically significant increases in the eye pigment contents of homozygous blos1ex2 and ltd1 mutants.

Together, these results suggest that the gene dosage of Atg2 influences the eye pigmentation phenotype of AP-3 mutants as well as of other mutations affecting pigment granule biogenesis. Because the eye pigmentation of wild-type flies was likewise modified, it is tempting to speculate that the product of Atg2 may influence pigmentation by a mechanism distinct from those mediated by AP-3, BLOC-1 and Lightoid. Thus, one may envision that autophagy—at least a form of it that it is sensitive to partial reductions in Atg2 activity—could be involved in the turnover of pigment granules. In such a scenario, decreased pigment granule degradation in Atg2EP3697 heterozygotes would explain the increase in eye pigment content observed for all of the genetic backgrounds tested. Furthermore, the extent of pigment content increase could depend on how impaired autophagy of these granules might be in Atg2EP3697 heterozygotes as well as on the rate of pigment granule turnover in the different genetic backgrounds (e.g., abnormal pigment granules in mutant flies might be targeted for degradation at a rate higher than that of normal granules in wild-type flies). Although the notion of organelle turnover by autophagy is widely accepted, and extensively documented for compartments such as mitochondria or peroxisomes [75], evidence for a direct role of autophagy in the degradation pigment granules (or melanosomes, in mammals) remains quite limited. For example, autophagosomes containing melanosomal proteins were observed in human melanocytes.
Fig 5. Validation and fine mapping of the critical region responsible for the modifier effect observed for Df(3L)BSC23. Red pigments were extracted from the heads of adult g2 mutant males carrying no deletions (—) or a single copy of the indicated deficiencies, quantified as described under Materials and Methods, and expressed as percentages of the red pigment content of male flies of the wild-type (Canton-S) line. Bars represent means ± SD of 3–15 biological replicates. One-way ANOVA followed by Dunnett’s test of each group versus g2 flies carrying no deletion (black bar): **p<0.01, ***p<0.001.
deficient in the HPS1 subunit of BLOC-3 [76]–a protein complex that functions in the biogenesis of LROs through activation of Rab32 and Rab38 [77]–and compartments described as “autophagic melano-lysosomes” were found in retinal pigment epithelium from apparently normal rhesus monkeys [78]. In addition, it has recently been reported that melanosomes transferred from melanocytes to keratinocytes in human skin can be degraded by autophagy [79]. On the other hand, published work using mammalian melanoma cell lines has uncovered a surprising link between specific components of the machinery for autophagy and the regulation of melanosome biogenesis, with some of these components promoting (e.g., the human Atg18 homologue WIPI1 [80,81]) and others antagonizing (e.g., the human Atg1 homologue ULK1 [82]) melanin accumulation in cell culture. Given this reported link between autophagy and the biogenesis of mammalian melanosomes, a putative role for Atg2 in the regulation of fly pigment granule biogenesis should be considered.

From a genetic perspective, two aspects of the identification of Atg2 in our genetic modifier screening are worth noting. First, so far genetic defects in autophagy-related genes in humans or mice have been associated with reduced pigmentation [80,83,84], not with increased pigment content. Second, it is not clear why out of the large number of known autophagy-related genes [60] only Atg2 was isolated in a screening designed to cover >92% of fly autosomal chromosomes (a small subset of these genes, including Atg5 and Atg8a, were excluded from the screening because of their localization to fly chromosome X). Plausible explanations include genetic redundancy among groups of paralogs with partially overlapping functions, as it might be the case for the related Atg18a and Atg18b genes encoding alternative binding partners of Atg2 [64], and gene dosage sensitivity, i.e., loss of one of the two copies of a gene may not necessarily translate into significantly decreased physiological function of the gene product. These issues notwithstanding, the possibility of the Atg2 protein playing a specific function related to pigment granule dynamics (biogenesis or degradation), besides its role as a core component of the autophagy machinery [60], should deserve consideration for future studies.

ArfGAP1 as a dominant genetic modifier of AP-3 mutants

Two genes within the critical region inferred for the Df(3L)eygC1 deficiency (Fig 4) and with available loss-of-function alleles were considered as candidates of interest. The first candidate was eyegone (eyg), which encodes a member of the Pax6 family of transcription factors involved in retina development [85]. While the eyes of homozygous eyg1 mutants are drastically reduced in size [85], heterozygous mutants display apparently normal eye morphology. In contrast with the Df(3L)eygC1 deficiency, however, a single copy of the eyg1 allele failed to modify the eye pigmentation phenotype of g2 mutants. Thus, the red pigment contents (expressed as percentage of wild-type levels) were 29.5 ± 2.1 for g2 and 30.0 ± 3.4 for g2;eyg1/+ (means ± SD; n = 6 and 9 respectively). In light of this result, this candidate was not pursued any further.

The second candidate was ArfGAP1 (formerly known as Gap69C), which encodes a conserved GTPase-activating protein (GAP) with specificity to promote hydrolysis of GTP bound to members of the Arf protein family [52,86]. Flies carrying the null allele ArfGAP1G3-85, either in homozygous or heterozygous form, are viable and display no eye morphological abnormalities [52]. Here, a single copy of the ArfGAP1G3-85 allele elicited statistically significant increases in the pigment contents of the AP-3 mutants g2 (Fig 7A) and rb1 (Fig 7B). On the other hand,
Fig 6. Effects of the Atg2EP3697 allele on red pigment content in various genetic backgrounds. Red pigments were extracted from the heads of adult male flies of the indicated genetic backgrounds lacking (open bars) or carrying (filled bars) one copy of the Atg2EP3697 allele over a normal chromosome 3. The extracted pigments were quantified as described under Materials and Methods, and the resulting values expressed as percentages of the pigment content of wild-type (Canton-S) flies. Bars represent means ± SD of 7–15 biological replicates. One-way ANOVA followed by Bonferroni comparison of selected group pairs: *p<0.05, **p<0.01, ***p<0.001. Notice that a single copy of Atg2EP3697 over a normal Atg2 allele increased pigmentation of two AP-3-subunit mutants (g² and rb¹), a BLOC-1-subunit mutant (blos¹ex²), and a mutant in...
and consistently with the behavior observed for the $Df(3L)eyg^{C1}$ deficiency (Table 1), a single copy of the $ArfGAP1^{G3-85}$ allele failed to modify the red pigment content of wild-type flies (Fig 8A). Interestingly, the modifier effect elicited by the $ArfGAP1^{G3-85}$ allele in the $g^2$ background appeared to follow a dominant pattern. Thus, eye pigmentation increases similar to those observed for $g^2$ flies heterozygous for $ArfGAP1^{G3-85}$ over a normal third chromosome (Fig 7A) were observed not only for $g^2$ flies that were heterozygous for $ArfGAP1^{G3-85}$ over a control deletion (referred in the figure to as ‘Df3’) but also for those that were homozygous for the $ArfGAP1^{G3-85}$ mutation or compound heterozygous for $ArfGAP1^{G3-85}$ over either of two large deletions (‘Df1’ and ‘Df2’) encompassing the entire $ArfGAP1$ gene (Fig 7C).

Fig 7. $ArfGAP1$ as a modifier of the AP-3 eye pigmentation phenotype. (A-C) Red pigments were extracted from the heads of adult male flies of the indicated genetic backgrounds carrying wild-type (+) or mutant (G3-85) alleles of the $ArfGAP1$ gene on chromosome 3. The extracted pigments were quantified as described under Materials and Methods, and the resulting values expressed as percentages of the pigment content of wild-type (Canton-S) flies. Bars represent means ± SD of 3–17 biological replicates. Statistical analyses were performed by means of Student’s t-test (A and B) or one-way ANOVA followed by Dunnett’s test of each group versus $g^2$ flies carrying no deletion (C): *p<0.05, **p<0.01, ***p<0.001. Notice that a single copy of $ArfGAP1^{G3-85}$ mutant allele over wild-type $ArfGAP1$ was sufficient to ameliorate the pigmentation defects of both $g^2$ (A) and $rb^1$ (B) AP-3-subunit mutants. Notice in (C) that such partial suppressor effect was not exacerbated in flies homozygous for the $ArfGAP1^{G3-85}$ allele or heterozygous for this allele over any of two deficiencies in which the deleted genomic regions include the entire $ArfGAP1$ gene, namely $Df(3L)eyg^{C1}$ (Df1) and $Df(3L)BSC380$ (Df2). The deficiency $Df(3L)BSC413$, in which the deleting genomic region excludes the $ArfGAP1$ gene, was used as a control (Df3).

Suggested size: 2 columns
As described in the previous section for the \textit{Atg2EP697} mutant allele, we next tested for effects of the \textit{ArfGAP1G3-85} allele on the phenotypes of \textit{blos1ex2} and \textit{ltd1} mutants (deficient in BLOC-1 and the Rab GTPase Lightoid, respectively). The results were remarkable. On the one hand, males that were double homozygous for \textit{ArfGAP1G3-85} and \textit{blos1ex2} did not survive to adulthood (indicated with X). In those cases in which viable adult males were obtained, red pigments were extracted and quantified as described under Materials and Methods. Values were expressed as percentages of the pigment content of wild-type flies. Bars represent means + SD of 3–9 biological replicates. Statistical analyses were performed by means of Student’s t-test (B) or one-way ANOVA followed by Dunnett’s test of each group versus that of flies homozygous for the wild-type \textit{ArfGAP1} allele in the corresponding genetic background (A and C): ns, not significant; *p<0.05, ***p<0.001. (D and E) Eye morphology of adult flies homozygous for \textit{ltd1}’ (D) and for both \textit{ltd1}’ and \textit{ArfGAP1G3-85} (E), with the latter displaying a mild interommatidial facets phenotype.

As described in the previous section for the \textit{Atg2EP697} mutant allele, we next tested for effects of the \textit{ArfGAP1G3-85} allele on the phenotypes of \textit{blos1ex2} and \textit{ltd1} mutants (deficient in BLOC-1 and the Rab GTPase Lightoid, respectively). The results were remarkable. On the one
hand, a single copy of ArfGAP1\textsuperscript{G3-85} did not modify significantly the eye color phenotypes of homozygous \textit{blos1}<sup>x2</sup> and \textit{ltd}<sup>i</sup> flies (Fig 8B and 8C), suggesting that the dominant modifier effect observed for ArfGAP1\textsuperscript{G3-85} in the genetic backgrounds of \textit{g}<sup>1</sup> (Fig 7A) and \textit{rb}<sup>1</sup> (Fig 7B) reflected a specific functional interaction between ArfGAP1 and the AP-3 complex. On the other hand, in homozygous form the ArfGAP1\textsuperscript{G3-85} allele caused drastic effects, namely: synthetic lethality of males in the background of \textit{blos1}<sup>x2</sup> homozygous (i.e., no doubly homozygous males survived to adulthood), and abnormal eye morphology (i.e., disrupted organization of interommatidial bristles) with decreased pigmentation in the background of \textit{ltd}<sup>i</sup> homozygotes (Fig 8C–8E). These strong effects, which were also observed using compound heterozygous of ArfGAP1\textsuperscript{G3-85} over deletions of the entire \textit{ArfGAP1} gene [using deficiencies \textit{Df(3L)BSC380} and \textit{Df(3L)ED4483} for \textit{blos1}<sup>x2</sup> and the former for \textit{ltd}<sup>i</sup>], were at odds with those observed by ArfGAP1\textsuperscript{G3-85} in homozygous form in the genetic backgrounds of \textit{g}<sup>1</sup> (Fig 7C) or Canton-S (Fig 8A), where neither early lethality nor abnormal eye morphology were noted.

Taken together, the results suggest that ArfGAP1 may be a dominant genetic modifier of AP-3 function and interact genetically with \textit{blos1} or \textit{ltd} in a recessive fashion. Because of the multiple cellular functions ascribed to the conserved ArfGAP1 protein in various organisms [86, 87], several scenarios seem plausible. For example, given that the recruitment of the AP-3 complex to membranes is promoted by an Arf family member (possibly Arf1) in its GTP-bound form [88–91], one could envision that decreased Arf GAP activity (due to reduced dosage of the \textit{ArfGAP1} gene) might partially compensate for incomplete impairment of AP-3 function (as in the genetic backgrounds of \textit{g}<sup>1</sup> and \textit{rb}<sup>1</sup>) by slowing down inactivation of GTP-bound Arf molecules at the site(s) of AP-3 function. While this relatively simple scenario could explain the specificity and even the dominant pattern of the effects elicited by the ArfGAP1\textsuperscript{G3-85} allele on AP-3 mutants, a few potential shortcomings should be noted. Firstly, at least in mammalian cell lines the regulation of AP-3 function through inactivation of Arf-GTP has been ascribed to a different Arf GAP, named AGAP1 [24], for which a clearly recognizable homologue (encoded by \textit{ceng1A}) exists in flies [92]. Secondly, ArfGAP1 has well-documented activities besides inactivation of Arf GTPases, including serving as effectors of GTP-bound Arf proteins and having Arf-independent functions [86, 87]. Finally, such a simple model would not provide a straightforward explanation for the strong genetic interactions observed for homozygous double mutants in ArfGAP1 and either \textit{blos1} (synthetic lethality in males) or \textit{ltd} (abnormal eye morphology), as genetic interactions between AP-3 mutants and \textit{blos1} or \textit{ltd} alleles have been documented to affect eye pigment content but neither viability nor eye morphology [37, 39]. Along these lines, one should consider that a second Arf-dependent protein trafficking pathway, defined by the structurally related AP-1 complex, has been implicated in the biogenesis of mammalian melanosomes [21, 22, 28, 93] and might be likewise operational in the biogenesis of pigment granules in the fly eye. Accordingly, one could envision that reduced activity of ArfGAP1, which in mammals has been shown to interact physically with AP-1 [94], might exacerbate the activity of the AP-1-dependent pathway−at least in the context of impaired AP-3 function. Because AP-1, unlike AP-3, is essential for \textit{Drosophila} larval development [95], invoking a putative role for ArfGAP1 in the AP-1-dependent pathway could provide possible explanations for the abnormalities observed in homozygous double mutants for ArfGAP1 and \textit{blos1} as well as for ArfGAP1 and \textit{ltd}. These considerations notwithstanding, a scenario whereby ArfGAP1 would act on an AP-1-dependent pathway would not represent the only plausible alternative to that of a direct role in AP-3-dependent events. For example, recent studies have demonstrated that ArfGAP1 from mammals and flies can interact physically and functionally with a leucine-rich repeat kinase, named LRRK2 in mammals and Lrrk in flies [96, 97], which in turn has been shown to function in the endosomal-lysosomal system and autophagy of both flies and mammals [98–101] in part through interactions with Rab GTPases.
such as Rab7 [100] and—most notably—Lightoid and its mammalian counterparts [102,103]. Accordingly, decreased ArfGAP1 activity could have elicited the modifier effects reported in this paper through alteration of the function of proteins such as Lrrk. Future research will be required to test experimentally the validity of these and other plausible scenarios.

Conclusions

Our genetic modifier screening has led to the identification of distinct regions within fly chromosome 3, and in particular of two candidate genes, which upon deletion or mutation in heterozygous form modified the eye pigmentation phenotype of AP-3 mutants. One of these candidate genes encodes the single fly counterpart of Atg2, a conserved component of the molecular machinery for autophagy. The other gene encodes the single fly counterpart of ArfGAP1, a conserved protein implicated in Arf-dependent protein trafficking events as well as in Lrrk-dependent regulation of the endosomal-lysosomal system and autophagy.

Acknowledgments

We thank Verónica T. Cheli for her participation in early stages of the project, specifically, in a pilot screening using a subset of deficiency lines. We thank Drs. Vladimir E. Alatortsev and David E. Krantz for their generous gifts of fly strains. Stocks obtained from the Bloomington Drosophila Stock Center (NIH P40OD018537) were used in this study.

Author Contributions

Conceived and designed the experiments: ECD IAR. Performed the experiments: IAR. Analyzed the data: IAR ECD. Wrote the paper: ECD IAR.

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