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Single amino-acid mutation in the *Drosophila melanogaster* ribosomal protein uL11: an insight in its transcriptional activity

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ABSTRACT

The ribosomal protein uL11 is located at the basis of the ribosome P-stalk and plays a paramount role in translation efficiency. In addition, no mutant for uL11 is available suggesting that this gene is haplo-insufficient as many other Ribosomal Protein Genes (RPGs). We have previously shown that overexpression of Drosophila melanogaster uL11 induces the transcription of many RPGs and Ribosomal Biogenesis genes (RiBis) suggesting that uL11 might globally regulate the level of translation through its transcriptional activity. Moreover, uL11 trimethylated on lysine 3 (uL11K3me3) interacts with the chromodomain of the Enhancer of Polycomb and Trithorax Corto. uL11, Corto and RNA Polymerase II co-localize at many sites on polytene chromosomes. These data have led to the hypothesis that the N-terminal end of uL11, and more particularly the trimethylation of lysine 3, supports the extra-ribosomal activity of uL11 in transcription. To address this question, we mutated the lysine 3 codon using a CRISPR/Cas9 strategy and obtained several lysine 3 mutants. We describe here the first mutants of *D. melanogaster uL11*. Unexpectedly, the *uL11^{K3A}* allele, in which the lysine 3 codon is replaced by an alanine, displays a genuine Minute phenotype known to be characteristic of RPG deletions (longer development, low fertility, high lethality, thin and short bristles) whereas the $uL11^{K3Y}$ allele, in which the lysine 3 codon is replaced by a tyrosine, is unaffected. In agreement, the translation rate slightly decreases in $uL11^{K3A}$ but not in $uL11^{K3Y}$. Deep-sequencing of RNA from wing imaginal discs shows enrichment in the GO categories "glutathione metabolism" for up-regulated genes in both mutants and "regulation of transcription" for down-regulated genes in *uL11^{K3A}*. Furthermore, analysis of deregulated genes' *cis*-regulatory sequences suggests that uL11 might regulate transcription of target genes in concert with the couple of transcription factors Mad/Med that mediate response to the Bone Morphogenetic Protein (BMP) signaling pathway.

INTRODUCTION

Loss of function alleles of *Ribosomal Protein Genes* (*RPGs*) have been studied for almost a hundred years in *Drosophila*, where they are known as *Minute* mutations (1). *Minute* mutants were first described for displaying thin and short bristles, *i.e. Minute* bristles, together with prolonged development (2). All *Minute* mutations are dominant and lethal when homozygous. The vast majority of them strongly impact viability and fertility, to the point that some could only be identified through transient aneuploidy experiments. *Minute* loci have been characterized over time and the *Minute* genes have now been identified to encode ribosomal proteins with very few exceptions (3,4).

While a substantial part of ribosomal proteins' contribution to cell metabolism has been attributed to their ability to alter ribosome behaviour with consequences on protein synthesis, some free ribosomal proteins are also known for a long time to carry regulatory activities, consequently termed "extra-ribosomal functions" [for a review see (5)] This is notably the case for RPL12, aka uL11 following the new nomenclature recently proposed to avoid confusion between species (6). Indeed uL11 was shown in *C. elegans* and mammals to bind its own messenger RNA and inhibit its splicing (7,8). Furthermore, in *S. cerevisiae*, uL11 was shown to be required for the transcription of a subset of PHO pathway genes that are inducible under low phosphate conditions (9).

Drosophila melanogaster uL11 is encoded by a unique gene (uL11/RPL12/CG3195/FBgn0034968) located on the right arm of chromosome 2 at cytogenetic position 60B7. Three annotated transcripts encode the same 165 amino acid protein. uL11 expression is ubiquitous and described as "very high" to "extremely high" in all tissues, developmental stages, and cell lines (10). Two deletions encompassing the whole uL11/RpL12 area have been described [i.e. Df(2R)bwVDe2LPxKR and Df(2R)Exel6081]. However, it was recently proposed that the cytological borders of the first one did not cover the uL11/RpL12 region (FBrf0230794) and the second one was lost (FBrf0206661). Thus, no evidence remains that flies can accommodate aneuploidy at this locus, which is indeed described as haplolethal (4,11). Another RPG (eL39/RpL39), and several essential genes (eIF5A, yki) are found in the vicinity of uL11/RpL12 (12,13). For this reason, the genes responsible for this haplolethality remain uncertain. However, it is likely that uL11/RpL12 contributes to this phenotype as no classical allele of uL11 has been described and we observed that ubiquitous RNAi mediated inactivation is lethal during the first larval instar.

The uL11 protein forms, together with the ribosomal protein uL10, the base of the P-stalk, a lateral protuberance of the 60S subunit which is a critical element of the ribosomal GTPase-associated center known to interact with translational elongation and termination factors (14). The uL11 protein consists of two globular domains connected with a hinge, a C-

terminal domain anchored to a conserved region of the 28S rRNA and a mobile N-terminal domain. The N-terminal domain interacts with several translation factors, notably eEF2 (14,15). As a consequence, uL11 has been shown to play an important role at many steps of the translation cycle. In yeast, deficiency of uL11 prevents the release of the ribosome associated protein Tif6, which is the last maturation step before the 60S subunit becomes functional (16). Yeast uL11 deficient ribosomes display a decrease in translational fidelity (16), and depletion of uL11 in cultured human Cystic Fibrosis bronchial epithelial cells reduces the rate of translation initiation and elongation (17).

We have recently established that *Drosophila melanogaster* uL11 is bound by a chromodomain-like structure of the epigenetic co-factor Corto. Chromodomains are known to specifically recognize trimethylated lysines on histones, and the interaction between uL11 and Corto was indeed shown to require uL11 lysine 3 to be trimethylated (uL11K3me3). Trimethylation of uL11 lysine 3 is very conserved and has been confirmed in *S. cerevisiae*, *S. pombe* and *A. thaliana* (18–20). In *D. melanogaster*, uL11 and Corto bind chromatin, colocalize on many sites on polytene chromosomes and are recruited on the *hsp70* gene upon transcriptional activation (21). Overexpression of the Corto chromodomain as well as uL11 induces the transcription of many *RPGs* and *Ribosomal Biogenesis* genes (*RiBis*) suggesting that uL11 might globally regulate the level of translation through its transcriptional activity. Thus, our previous study led us to hypothesize that uL11K3me3 is involved in transcriptional regulation.

To test this hypothesis, we designed a CRISPR/Cas9 strategy for mutating the uL11 lysine 3 codon. We describe here the $uL11^{K3A}$ and $uL11^{K3Y}$ mutants in which the lysine 3 codon of uL11 is replaced by an alanine and a tyrosine codon, respectively. $uL11^{K3A}$ displays Minute phenotypes i.e. longer development, low fertility, high lethality and thin bristles, but surprisingly $uL11^{K3Y}$ looks wild-type. RNA-seq analysis of wing imaginal discs from both mutants show enrichment of up-regulated genes in the GO category "glutathione metabolism" and down-regulated genes in the GO category "regulation of transcription". Analysis of deregulated genes' cis-regulatory sequences suggests that uL11 might act in concert with the couple of transcription factors Mad/Med that are primary effectors in the Bone Morphogenetic Protein (BMP) signaling pathway.

MATERIALS AND METHODS

Drosophila genetics

Drosophila melanogaster stocks and crosses were kept on standard yeast corn-meal medium (7.5 % yeast, 9 % cornmeal, 1 % agar, 0.27 % moldex) at 25 °C. For all experiments, crosses were set up with similar densities to prevent confounding effects of overcrowding.

 w^{1118} was used as the control strain. The $corto^{420}$ and $corto^{L1}$ alleles were described in (21).

Strategy of CRISPR/Cas9 mutagenesis

The *pU6-chiRNA:sgRNA* plasmid was obtained by incorporating the *sgRNA* sequence (obtained by annealing *phos-gRNA-F* and *phos-gRNA-R*, Supplementary Table 1) into *pU6-BbsI-chiRNA* (Addgene plasmid #45946) (22) following the protocol provided on http://flycrispr.molbio.wisc.edu/protocols/gRNA. Sequence was confirmed using the T3 universal primer.

A 123 nucleotides long single-stranded DNA (ssODN) carrying the lysine (AAA) to alanine (GCC) substitution flanked by two 60 nucleotide-long homology arms was used as a template for Homologous Directed Repair (HDR) (synthetized and purified by standard desalting by Integrated DNA Technologies, Inc) (Supplementary Table 1). The *uL11* region of the recipient line *vasa-Cas9* (BL-51324) was sequenced in order to respect possible polymorphisms. To prevent base pairing with the *sgRNA*, the *ssODN* was designed to be homologous to the PAM carrying strand.

Fly transgenesis

Two hundred vasa-Cas9 embryos were injected with a mixture containing 100 ng/µL pU6-chiRNA:sgRNA and 100 ng/µL ssODN (BestGene Inc.). Transformant G0 flies (48 females and 44 males) were individually crossed to w^{1118} ; In(2LR)Gla, wgGla1/SM5 flies (Gla/SM5). Among them, only 18 males and 11 females were fertile. Curly wing G1 siblings were individually crossed to Gla/SM5 flies. Once the G2 progeny born, G1 founding flies were harvested, genomic DNA extracted, and genotyping performed as described below. 294 G1 individuals were genotyped to detect the presence of uL11 mutant alleles. Curly wing G2 offspring of G1 flies carrying a mutant allele of uL11 were crossed with each other to establish mutant balanced strains. In order to eliminate potential unspecific mutations, balanced mutant females were crossed with w^{1118} males. Ten heterozygous mutant females from the offspring were then individually backcrossed with w^{1118} males and genotyped by HRMA after laying eggs. uL11 mutant females were kept and the whole procedure was repeated seven times.

Genomic DNA extraction

Genomic DNA was extracted by crushing single flies in 100 μ L SB buffer (10 mM Tris pH 8.0, 1 mM EDTA, 25 mM NaCl, 200 μ g/ μ L Proteinase K), followed by 30 min incubation at 37 °C. DNA was further purified by standard phenol-chloroform extraction followed by ethanol precipitation.

Locked Nucleic Acid allele-specific gPCR

Forward allele-specific primers with 3' end matching either wild-type (*LNAWT*) or mutated 3rd codon (*LNAK3A*) of *uL11* (AAA or GCC, respectively) and a Locked Nucleic Acid (LNA) nucleotide (23) at the second position of the mismatch codon were used in combination with a reverse primer (*CRISPR1_R*) to amplify a 219 nucleotide fragment (Supplementary Table 1). 25 μL reactions were set to contain 5 to 15 ng of genomic DNA, 0.5 μM forward and reverse primers, 0.4 nM dNTP, 0.75 μL SYBR green (Diagenode), and 2.5 units of DreamTaq polymerase (Thermo Fisher Scientific) in TMAC buffer (67 mM Tris pH 8.8, 6.7 mM MgCl2, 16.6 mM ammonium sulfate, 0.5 mM tetramethylammonium chloride, 0.17 mg/mL BSA) (24). qPCR plates were prepared and kept at 4°C until starting PCR cycles. 0.5 ng of plasmid containing the *uL11* coding region in which the AAA lysine 3 codon was replaced by GCC was used as positive control. qPCRs were carried out in a CFX96 system (BioRad) [95°C 3 min; 40 cycles (95 °C 20 s, 64 °C 20 s, 72 °C 30 s)]. To confirm the presence of the mutated allele, a 1.5 kb region centred on the lysine 3 codon was amplified from positive genomic DNA and sequenced.

High Resolution Melting Analysis (HRMA)

Genomic DNA was analysed by High Resolution Melting Analysis (HRMA) as described by (25). Briefly, oligonucleotides uL11-HRMA-F and uL11-HRMA-R (Supplementary Table 1) were used to amplify a 173 bp region centred on the uL11 lysine 3 codon. PCR were performed with Sso Fast Eva Green SupermixTM (BioRad) in 20 μ L reactions containing 2 to 15 ng genomic DNA and 0.5 μ M each oligonucleotide. Cycles were carried out in a CFX96 system (BioRad) [98 °C 3 min; 40 cycles (98 °C 2 s, 57.3 °C 15 s)]. Thermal melting profiles were obtained in the same device by increasing temperature from 75 to 95 °C using a temperature increment of 0.2 °C. They were normalized as described by (26).

uL11K3me3 antibodies

Polyclonal uL11K3me3 antibodies were generated in rabbit using a peptide corresponding to the first 16 amino acids of uL11 with methylated lysine 3 [PPK(me3)FDPTEVKLVYLRC] (Eurogentec). The serum was first loaded on a uL11K3me3 peptide affinity column which allowed to retain uL11K3me3 and uL11 antibodies. After elution, unmethylated uL11 and uL11K3me3 antibodies were separated by passage through an unmethylated uL11 peptide affinity column. Specificity of the antibodies was checked by dot blot (Supplementary Figure 1).

Proteins were extracted from third instar larvae in RIPA buffer (150 mM sodium chloride, 1 % NP40, 0.5 % sodium deoxycholate, 0.1 % SDS, 50 mM TrisHCl pH 8,0) supplemented with phosphatase and protease inhibitors (Roche). 30 µg of proteins were separated by SDS-PAGE electrophoresis on 15 % acrylamide gels. Western blots were

performed according to standard protocols using either goat anti-uL11 (SantaCruz sc82359, 1/1000), rabbit anti-uL11K3me3 (1/6000), or mouse anti-α-tubulin (DSHB E7c, 1/2500) as primary antibodies. Anti-goat (Jackson ImmunoResearch; 705035147; 1/10000), anti-rabbit (1/20000) or anti-mouse (Sigma NA931; 1/20000) were used as secondary antibodies.

Analysis of mutant life history traits

uL11 wild-type or mutant chromosomes were balanced with *CyO,Dfd-EYFP* (from strain BL-8578) or *SM5*. About 100 females and 60 males were placed in laying cages on agarose plates (2 % agarose, 5 % vinegar, neutral red) supplemented with yeast. To measure embryonic lethality, 100 embryos were collected from each laying cage, transferred on new agarose plates and emerging first instar larvae were counted. To measure larval and pupal lethality, 100 embryos were collected and transferred into yeast cornmeal medium tubes at 25 °C. Pupae and adults were then counted. Three independent experiments were performed and results were pooled. To measure developmental time, first instar larvae were collected and transferred into yeast cornmeal medium tubes at 25 °C (50 to 100 larvae per tube). Vials were checked from 9 days after egg laying until no more adults emerged. Chi² tests were performed to compare genotypes.

Measure of bristle length

Adult bodies free of wings, legs and heads were aligned on agar cups. Images were captured using a Leica Model MZ FLIII microscope equipped with a Leica Model DC480 camera. Scutellar bristles were measured using the ImageJ segmented line tool. Normality was checked by Shapiro-Wilk tests and homogeneity of variances by F tests. Student's t-tests were then set taking into account homo- or heteroscedasticity of variances.

Measure of wings

Adult flies were kept in 70 % ethanol for 48 h and transferred into PBS glycerol (1:1 v/v). Wings were dissected and mounted on glass slides, dorsal side up, in Hoyer's medium. Slides were scanned with a Hamamatsu Nanozoomer Digital Slide scanner, running the Nanozoomer software with a 20×10^{-5} x objective and an 8 bit camera. Wing pictures were separately exported into tif format using NDP.view and the 5×10^{-5} lens. Measurements of wing length were performed as described in (27).

Plasmids

uL11 was amplified from w^{1118} embryonic cDNAs and subcloned into pENTR/D-TOPO (Invitrogen) (21). $pENTR-uL11^{K3A}$ and $pENTR-uL11^{K3Y}$ were obtained by site-directed mutagenesis using the oligonucleotides described in (21) and in Supplementary Table 1,

respectively. The cDNAs were then transferred either into the *pAWM* or the *pAWH* Gateway® *Drosophila* vector allowing expression of fusion proteins with a C-terminal Myc or HA tag, respectively.

Cell transfection

S2 cells were cultured at 25 °C in Schneider's *Drosophila* medium supplemented with 10 % heat-inactivated foetal bovine serum and 100 units/mL of penicillin and streptomycin (Life technologies). To obtain cells permanently expressing $uL11^{K3A}$, a mix containing a 5:1 molar ratio of the $pA-uL11^{K3A}$ -HA expression vector and the selection plasmid pCoBlast was prepared. 10^6 cells were then transfected with 2 μ g of DNA using Effecten® transfection reagent (Qiagen) according to the manufacturer's instructions at a 1:10 DNA/Effecten® ratio. Selection was performed by addition of 10 μ g/mL of blasticidin after 48 h. After initial selection, stable cell lines were cultured in presence of 2 μ g/mL of blasticidin. For transient expression, 10^6 cells were transfected with 2 μ g of either $pA-uL11^{K3A}$ -Myc or $pA-uL11^{K3Y}$ -Myc vector using Effecten® at a 1:10 DNA/Effecten® ratio.

Polysome fractionation

Cells were harvested at 50 % confluence and washed in Schneider medium at room temperature to remove the foetal bovine serum. They were then resuspended in ice-cold lysis buffer (20 mM Hepes pH 7.5, 250 mM KCl, 10 mM MgCl2, 5 mM DTT, 1 mM EDTA, 0.5 % NP-40) supplemented with EDTA-free protease inhibitor cocktail (Roche Diagnostics) and 40 U/mL Ribolock RNAse Inhibitor (ThermoFisher). For EDTA treatment, the lysis buffer was replaced with (20 mM Hepes pH 7.5, 250 mM KCl, 5 mM DTT, 25 mM EDTA, 0.5 % NP-40). After centrifugation at 500 g for 5 min to pellet nuclei, supernatants were layered onto 10 to 50 % sucrose gradients in polyribosome buffer (20 mM Hepes pH 7.5, 250 mM KCl, 20 mM MgCl2, 2 mM DTT), supplemented with EDTA-free protease inhibitor cocktail (Roche Diagnostics) and 40 U/mL Ribolock RNAse Inhibitor (ThermoFisher). Gradients were centrifugated at 39,000 rpm for 165 min at 4 °C in a Beckman SW41-Ti rotor. Optical density at 254 nm was monitored using a density gradient fractionator (Teledyne Isco, Lincoln, NE).

Puromycin assays

Puromycin assays were adapted from (28) with the following modifications: 20 larvae were turned inside out and incubated for 1 h at 25 °C under gentle rotation in Schneider's medium supplemented or not with 10 mg/mL cycloheximide (Sigma). Puromycin (antpr1, InvivoGen) was then added at a final concentration of 0.28 mg/mL and incubation was continued for 2 h.

Total proteins were extracted from third instar larvae in buffer containing [30 mM Hepes pH 7.4, 0.1 % NP40, 150 mM NaCl, 2 mM Mg(OAc)₂] supplemented with phosphatase and protease inhibitors (Roche) (adapted from (29). 60 µg of protein extracts were deposited on a 12 % acrylamide gel.

Western blot analyses were performed according to standard protocols with mouse anti-puromycin (Kerafast, 3RH11; 1/500) or mouse anti-H3 (Diagenode; C15200011;1/1000) as primary antibodies and anti-mouse (Sigma; NA931; 1/20000) as secondary antibodies. Background, puromycin and H3 signals were measured using ImageJ. The puromycin signal was normalized towards the H3 signal. Results were submitted to Student's t-tests.

Co-Immunoprecipitation

Cells transiently transfected with either *pA-uL11^{K3A}-Myc* or *pA-uL11^{K3Y}-Myc* and *pA-FLAG-CortoCD* (21) were harvested after 48 h and washed in Schneider medium at room temperature. Co-immunoprecipitations were performed as described in (30) without fixation. 30 µl of Protein G coated Bio-Adembeads (Ademtech) were incubated with either 1 µg of mouse monoclonal anti-FLAG antibody (F3165, Sigma), or goat anti-HA antibody as mock (sc-805, Santa Cruz Biotechnology).

Western blot analyses were performed according to standard protocols with mouse anti-Myc (Abcam ab9132 1/10000) or anti-FLAG (Sigma F3165 1/5000) as primary antibodies and anti-mouse (Sigma NA931; 1/20000) as secondary antibodies.

RNA-seg and bioinformatic analyses

Wing imaginal discs of third instar female larvae (one disc per larva) were dissected by batches of 50 in ice-cold PBS and frozen in liquid nitrogen. 150 discs (three batches) were pooled and homogenized in lysis buffer by pipetting. Total RNAs were extracted using RNeasy kit (Qiagen).

Preparation of library and sequencing of cDNA from *corto*⁴²⁰/*corto*^{L1} wing imaginal discs were performed as described in (21). For *uL11* wing imaginal discs, library preparation was performed using the TruSeq® Stranded mRNA Library Prep kit (Illumina). Library preparation and Illumina sequencing were performed at the Ecole normale superieure genomics core facility (Paris, France) on a NextSeq 500 (Illumina). Three replicates were sequenced for each genotype. 75 bp single reads were trimmed using FastQC (Galaxy version 0.72). Reads were then aligned against the *D. melanogaster* genome (dm6 genome assembly, release 6.30) using STAR (Galaxy Version 2.6.0b-2). Reads were counted using featurecount (Galaxy Version 1.6.0.6). Differential analysis was performed using DeSeq2 version 1.32.0. Gene ontology was analysed with DAVID (https://david.abcc.ncifcrf.gov/home.jsp). Motif discovery was performed using MEME (31) and motif comparison using TomTom (32) [MEME Suite,

version 5.3.3, (33)]. The RNA-Seq gene expression data and raw fastq files are available at the GEO repository (https://www.ncbi.nlm.nih.gov/geo/info/seq.html) under accession number GSE181926.

RT-qPCR were performed on wing imaginal disc cDNAs as described in (27). Expression levels were quantified with the Pfaffl method (34) and normalized to the geometric mean of two reference genes, *GAPDH* and *Spt6*, the expression of which did not vary in the mutants. Sequences of primer couples are listed in Supplementary Table 1.

RESULTS

CRISPR/Cas9 editing of the uL11 lysine 3 codon

uL11 is located within a cluster of highly transcribed genes, many of which are also essential (*eIF5A*, *RpL39/eL39*, *yki...*) (12,13). Unsurprisingly, this cluster is part of the 1.6 % haplolethal regions of the euchromatic *Drosophila* genome (4). The immediate neighbourhood of *uL11* contains only two small intergenic sequences (465 and 620 bp, respectively) that might contain regulatory elements (Figure 1A). Thus, the insertion of a selection cassette at any position inside this locus would likely disturb gene expression and impede viability. We therefore opted to edit the *uL11* lysine 3 to alanine (K3A) by a single step CRISPR/Cas9 mediated HDR using a single-stranded DNA donor template (*ssODN*).

To recover the successful events, we set up a phenotype-independent screening protocol based on an allele specific amplification strategy. Discriminating power was increased by the substitution of the penultimate nucleotide of the screening primers with a locked nucleic acid (LNA) (Supplementary Figure 2A). The presence of a single LNA sufficiently improved specificity to allow us analyzing pools of flies for the presence of a single allele copy (Supplementary Figure 2B). 294 G1 individuals were tested for the presence of a mutated allele carrying the K3A substitution. Mixtures of genomic DNA from 4 to 5 individuals were prepared and the uL11 locus was amplified with either the lysine codon (LNA-WT) or the alanine codon (LNA-K3A) matching primer. While most genomic DNA mixtures displayed amplification kinetics close to the level of the negative control, six of them exhibited faster amplification (Δ Ct between 2 and 7). We thus repeated the experiment on individual genomic DNAs from 6 pools. Ten genomic DNA originating from three independent G0 founding flies, two males and one female, exhibited quicker amplification with the LNA-K3A forward primer than with the control primer (Δ Ct>5) (Supplementary Figure 2C). Sequencing the uL11 locus confirmed that they contained the recombinant allele $uL11^{K3A}$ at the heterozygous state.

To detect other mutations potentially resulting from non-homologous end joining (NHEJ) events, we also performed High Resolution Melting Analysis (HRMA) of a qPCR amplicon centred on the *uL11* lysine 3 codon. Denaturation kinetics of these PCR products

were analyzed individually for the 294 G1 flies. Among them, 36 denaturation profiles differed from the wild-type control. Sequencing of the amplicons confirmed the presence of a mutation at the uL11 locus in each of these 36 samples. Consistently, the 10 $uL11^{K3A}$ mutants identified with the allele-specific amplification strategy were also recovered by HRMA. Seven additional alleles were thus identified, that all carry a mutation impairing the lysine 3 codon: a single (K3Y) or double (P2LK3E, P2QK3R) amino acid substitutions, a single (Δ K3) or double (Δ K3F4) amino acid deletion, and an insertion or a deletion of 4 nucleotides (F-4 and F+2, respectively) (Supplementary Figure 3). Our first observations revealed that the mutants could be dispatched into two groups depending on the severity of their phenotypes. The first group contains the K3A, $\Delta K3$, $\Delta K3F4$, F+2 and F-4 alleles, and the second group the K3Y, P2QK3R and P2LK3E alleles. We choose to focus on two representative alleles, K3A and K3Y (Figure 1B and Supplementary Figures 2D and 3). They were introduced into the same controlled genetic background (w^{1118}) and expression of the uL11 proteins was confirmed by western blot (Figure 2). As expected, uL11K3me3 was not detected in the two homozygous mutants.

Lethality and developmental delay of *uL11* mutants

We first examined the lethality of uL11 mutants. The lethality rate depended on the mutant allele. Homozygous $uL11^{K3Y}$ flies were fully viable, whereas very few $uL11^{K3A}$ homozygotes emerged and almost all of them were males.

Next, we examined the lethality of the two mutants during development. During embryogenesis, they did not display more lethality than the w^{1118} control with the exception of $uL11^{K3A}/uL11^+$ whose lethality is slightly higher (Chi² test, p<0.05) (Figure 3A). By contrast, during larval life, lethality was very high for $uL11^{K3A}/uL11^+$, and $uL11^{K3A}/uL11^{K3A}$ (Chi² test, p<0.001), but did not increase neither for $uL11^{K3Y}/uL11^+$ nor for $uL11^{K3Y}/uL11^{K3Y}$ larvae. We did not observe any lethality during the pupal life for all genotypes. Similarly, developmental time from egg deposition to adult emergence was considerably extended for $uL11^{K3A}/uL11^+$ (up to 48 h) and $uL11^{K3A}/uL11^{K3A}$ (up to 96 h) but unaffected in $uL11^{K3Y}/uL11^+$ and $uL11^{K3Y}/uL11^{K3Y}$ flies (Figure 3B).

To summarize, the K3Y mutation had no effect on these life history traits whereas lethality and developmental time were increased both in heterozygous and homozygous K3A mutants, which characterized this allele as dominant.

Bristle and wing size of the *uL11* mutants

We noticed that the *uL11^{K3A}/uL11^{K3A}* homozygous mutants had thinner and shorter macrochaetes as compared to wild-type flies (Figure 4A and B). To determine whether the size of the macrochaetes was affected, we measured the anterior and posterior scutellar bristles in males (Figure 4C and D). Scutellar bristles were significantly shorter in

heterozygous and homozygous $uL11^{K3A}$ but unaffected in homozygous $uL11^{K3Y}$ males as compared to wild-type flies (Figure 4C and D). Similarly, we observed shorter bristles in heterozygous $uL11^{K3A}$ females while $uL11^{K3Y}$ homozygous females were only slightly affected (Supplementary Figure 4A and B). Wing length was also measured. Results show that wings were significantly shorter in heterozygous and homozygous $uL11^{K3A}$ while unaffected in $uL11^{K3Y}$ homozygous males (Figure 4E). Heterozygous $uL11^{K3A}$ females exhibited shorter wings whereas wings of homozygous $uL11^{K3Y}$ females were unaffected (Supplementary Figure 4C).

These results confirmed that K3A was a dominant allele and showed that the severity of the phenotypes depended on the mutation. The K3A mutation was highly detrimental while the K3Y mutation had almost no impact on the size of macrochaetes and wings. *Minute* mutants are known to be poorly fertile and viable, to exhibit developmental delay and have shorter and thinner bristles, all phenotypes that we observed in $uL11^{K3A}$ mutant flies. Moreover, *Minute* alleles are dominant which is also the case of the $uL11^{K3A}$ allele. All these data characterized the $uL11^{K3A}$ mutant as a *Minute* mutant.

Impact of the *uL11* mutations on translation

The strategic location of uL11 protein at the base of the P-stalk in the GTPase-associated center of the ribosome, suggests that its mutation might have a detrimental impact on translation. To test this hypothesis, we assessed the level of global translation in the *uL11* mutants. In order to label neo-synthesized proteins, third instar wild-type or mutant larvae were incubated with puromycin. Puromycin intake was normalized to histone H3 levels. A significant decrease in global translation level was observed in *uL11*^{K3A} heterozygous and homozygous larvae as compared to wild-type larvae, whereas it was unmodified in *uL11*^{K3Y} homozygous larvae (Figures 5A and B). These results were consistent with our previous phenotypic observations showing that *K3A* mutation was the most detrimental.

We then asked whether the uL11K3A variant protein would retain the ability to associate with translating ribosomes. We generated stable cell lines expressing either uL11K3A-HA or uL11-HA under the control of the *Actin* promoter. Cytoplasmic extracts were purified from both genotypes and lysates were loaded onto sucrose gradients for fractionation. As a control, an extract of each genotype was supplemented with 25 mM EDTA, a concentration that disrupts the interaction between ribosomal subunits and mRNA. The resulting fractions were analyzed by western blot to reveal the presence of uL11K3A-HA or uL11-HA (Figures 5C and 5D). These experiments revealed an enrichment of uL11-HA as well as uL11K3A-HA at the 60S and 80S ribosomal fractions and at polysomal fractions but not at the 40S fractions. In addition, the distribution of uL11K3A-HA was not biased towards inactive ribosomal fractions when compared to uL11-HA. Furthermore, EDTA treatment

triggered the relocation of uL11-HA and uL11K3A-HA towards lighter fractions, confirming that the sedimentation profiles truly resulted from their association with polysomes (Figures 5C and 5D). All these data showed that uL11K3A-HA was efficiently incorporated into translating ribosomes. Nevertheless, the decrease in puromycine incorporation observed in *uL11*^{K3A} mutants suggests that the yield of translation might be altered.

Interaction between the Corto chromodomain and uL11 mutant proteins

uL11 shares many transcriptional targets with Corto and interacts with its chromodomain through trimethylated lysine 3 (21). We thus hypothesized that the transcriptional activity of uL11 was impaired in the *uL11^{K3A}* and *uL11^{K3Y}* mutants. To address this, we first assayed the physical interactions between the chromodomain and the two uL11 mutant proteins by co-immunoprecipitation. *Drosophila* S2 cells were co-transfected with *pA-FLAG-CortoCD* and either *pA-uL11^{K3A}-Myc* or *pA-uL11^{K3Y}-Myc*. The FLAG-tagged chromodomain was immunoprecipitated using anti-FLAG antibodies. Contrarily to uL11-Myc that precipitated with FLAG-CortoCD, neither uL11K3A-Myc nor uL11K3Y-Myc co-immunoprecipitated with it (Figure 6). These results corroborated our previous results (21) and suggested that the transcriptional activity of uL11 might be at least partly impaired in the two mutants.

Transcriptomic analysis of *uL11* mutants

To address the role of uL11 in transcriptional regulation, we deep-sequenced transcripts from third instar larval wing imaginal discs of homozygous mutants for $uL11^{K3A}$, $uL11^{K3Y}$ or trans-heterozygous mutants for corto ($corto^{L1}/corto^{420}$). The w^{1118} line was used as reference (Table 1). Differential analyses were performed to obtain adjusted p-values associated to expression fold-changes for the three genotypes as compared to the reference. Taken a $\log_2(\text{fold-change}) < -0.5$ or > 0.50 and an adjusted p-value < 0.05, we found 458 down-regulated and 481 up-regulated genes in corto mutants (Supplementary Table 2). Using the same cutoffs, we found 152 down-regulated and 262 up-regulated genes in $uL11^{K3A}$, and 39 down-regulated and 45 up-regulated genes in $uL11^{K3Y}$ (Supplementary Table 2). Only 52 deregulated genes were shared by $uL11^{K3A}$ and $uL11^{K3Y}$, and 14 by the three genotypes, $uL11^{K3A}$, $uL11^{K3Y}$ and $corto^{L1}/corto^{420}$ (Figure 7A). qRT-PCR validations of RNA-seq data were performed for three genes: CG13516, one of the few genes to be deregulated in the three genotypes, Hsp67Bc, encoding a small heat-shock protein deregulated in $corto^{L1}/corto^{420}$ and $uL11^{K3Y}$, and GstE6, encoding a Glutathione S-transferase deregulated in $uL11^{K3A}$ and $uL11^{K3Y}$ (Figures 7B and C).

Strikingly, in *corto^{L1}/corto⁴²⁰* mutants, 52.7 % of the *RPGs* (89/169) were up-regulated (Supplementary Table 3). A similar enrichment of *RPGs* had been observed in flies

overexpressing the Corto chromodomain (21). Our results thus indicated that this overexpression behaved as a dominant negative allele.

Contrarily to what we observed following overexpression of uL11 (21), we did not find any deregulation of RPGs in imaginal discs of uL11 mutants. Gene ontology analyses revealed that genes up-regulated in $uL11^{K3A}$ imaginal discs are enriched in the category "glutathione metabolism" and down-regulated genes in the category "regulation of transcription" (Supplementary Table 3). The only enrichment found for $uL11^{K3Y}$ was for up-regulated genes in the category "glutathione metabolism" as for $uL11^{K3A}$ (KEGG pathway, Benjamini adjusted p-value 5.90e-03).

In an attempt to identify the transcriptional partners of uL11, we searched for conserved motifs in the *cis*-regulatory sequences of the deregulated genes. As *uL11^{K3A}* displayed a global decrease in translational rate that might interfere with the transcriptional response, we focused on the 84 genes deregulated in *uL11^{K3Y}*. Using MEME (32), we identified a 21 nucleotide-long consensus sequence present in the *cis*-regulatory sequences of 39 on the 84 genes (E-value: 2.1e-23). Using TomTom (31), we found that this motif significantly matched the 14 nucleotide-long binding site for Mad (p-value: 4.9e-07), as well as the 11 nucleotide-long binding site for its partner Med (p-value 1.11e-04 that together mediate the cellular response to the BMP pathway (Figure 7D and Supplementary Table 4).

DISCUSSION

We have previously shown that ribosomal protein uL11 interacts with the chromodomain of the *Drosophila* Enhancer of Trithorax and Polycomb Corto when trimethylated on lysine 3 (uL11K3me3) (21). uL11, Corto and RNA Polymerase II co-localize at many sites on polytene chromosomes and overexpression of *uL11* induces the transcription of many *RPGs* and *RiBi* genes. These data have confirmed that *Drosophila* uL11 is involved in transcription (35) and further suggest that the lysine 3 supports this extra-ribosomal activity. In the aim of testing this hypothesis, we generated mutant alleles of *uL11* using the CRISPR/Cas9 technology. By introducing a template to promote Homologous Directed Repair, we obtained a mutant in which the lysine was replaced by an alanine. However, we also obtained mutants harbouring indels probably obtained by Non-Homologous End Joining. Strikingly, the lysine 3 codon of uL11 was either deleted or substituted by another amino acid in all recovered mutants. Hence, it seems that a strong selection pressure occurs to maintain the *uL11* ORF, consistent with the haplo-insufficiency of this *RPG*.

A single amino acid substitution generates a *Minute* phenotype

The *uL11^{K3A}* allele produces very little homozygotes. Those are only males that hatch with a developmental delay larger than two days as compared to wild-type flies raised in the same conditions. Interestingly, *uL11*^{K3A} heterozygotes are also delayed but slightly less (about one day). In addition, *uL11*^{K3A} heterozygous females are frequently sterile making it necessary to carry out crosses of heterozygous *uL11^{K3A}* males with wild-type females and to genotype the offspring in order to maintain the stock. Hence, the $uL11^{K3A}$ allele clearly appears dominant, which is also visible for other phenotypes, such as the shorter and thinner macrochaetes and the global decrease in translational rate. Reduced viability, notably of females, delayed development, thin bristles and dominance are signatures of the Minute mutations that have been more recently shown to correspond to RPG deletions (3). The associated phenotypes are thought to reflect a defect in ribosomal proteins' stoichiometry resulting in a decreased capacity for protein synthesis. In accordance, tissues that are the most dependent on translation are likely to be the most affected. For instance, high ribosome biogenesis level is suggested to be necessary for the maintenance of germinal stem cells in the Drosophila ovarium, which could explain the reduced fertility of Minute females (36,37). Similarly, macrochaetes are described to require a very high amount of protein synthesis over a short developmental period (3). Unexpectedly, replacement of a single lysine by an alanine in the N-terminal tail of uL11 induces a *Minute* phenotype. Nevertheless, the global level of translation in the mutant is only slightly decreased and the mutated protein can be efficiently incorporated into translating ribosomes. However, translation speed or accuracy might be altered. Indeed, in yeast, uL11 loss-of-function has been shown to halve translation speed and to cause increased amino acid mis-incorporation and termination codon readthrough (16). Another possibility would be that uL11K3A ribosomes display altered affinity for specific mRNA, as has been described for ribosomes lacking RPL38/eL38 in mice (38).

Surprisingly, the $uL11^{K3Y}$ mutant does not display any *Minute* phenotype. The N-terminal extension of uL11 is supposed to be unstructured as neither the first 6 nor the first 9 amino acids were resolved in the structure of the *D. melanogaster* and *S. cerevisiae* 80S ribosome, respectively (15,39). This region is composed of hydrophilic amino acids and could thus be a hub for protein interactions. Addition of methyl groups on lysines, even if it has no effect on the overall charge of the residue, increases its hydrophobicity. Trimethylation of uL11 lysine 3 might thus modulate the activity of the N-terminal tail (40). However, in *S. cerevisiae* and *pombe*, deletion of the enzyme responsible for uL11 lysine 3 methylation has little impact on ribosome assembly and function or on cell viability (19,41) suggesting that the methylation of lysine 3 is not crucial for basic translational activity. In the $uL11^{K3Y}$ mutant, the tyrosine, a hydrophobic residue, might mimic the effect of methyl groups. If this is the case, in-depth comparison of translation between $uL11^{K3Y}$ and $uL11^{K3A}$ mutants, including sequencing of

polysomal mRNA, should permit to elucidate the role of the uL11 N-terminal tail methylation in translation.

Extra-ribosomal activities of uL11

Reduction in translation capacity might be indirect and due to extra-ribosomal activities of free ribosomal proteins (20,41). Moreover, in nematodes and mammals, uL11 autoregulates its own expression by interfering with the splicing of its mRNA. Indeed, an excess of uL11 induces aberrant retention of an intron and elimination of the corresponding mRNA by Non-sense Mediated Decay (8,7). We previously reported that overexpression of *uL11* increases the transcription of some *RPGs* and *RiBi* genes (21). The corresponding proteins might be produced, respecting the necessary stoichiometry for ribosome assembly, thus avoiding ribosomal stress. Our new data show that these genes are not down-regulated in the *uL11* lysine 3 mutants. Hence, the *Minute* phenotype of *uL11*^{K3A} does not result from transcriptional down-regulation of these genes, confirming that it should rather be due to an alteration in translation.

As the *uL11*^{K3Y} mutant displays neither *Minute* phenotypes nor a global decrease in translation, we assumed that solely its transcriptional activity would be affected. However, deep-sequencing of its transcriptome identified only 84 deregulated genes as compared to the reference line. Interestingly, the *cis*-regulatory sequences of a number of them share a nucleotide motif that significantly matches the binding site of the transcription factors Mad and Med, the effectors of the BMP pathway in *Drosophila*. Upon binding of the Decapentaplegic (Dpp) ligand to its heterodimeric serine/threonine kinase receptor, Mad is phosphorylated, translocates to the nucleus together with the Smad4 homolog Medea (Med) and regulates expression of target genes. This important pathway contributes notably to growth regulation and patterning [for a review see (44)]. Whether uL11 cooperates with Mad-Med in the regulation of transcription will remain to be demonstrated.

The absence of obvious phenotypes and the fact that only few genes are deregulated in $uL11^{K3Y}$ mutant could mean that the extra-ribosomal function of uL11 is not required under physiological conditions. Indeed, many extra-ribosomal functions are triggered under stress conditions. For example, in *S. cerevisiae uL11/RPL12* has been shown to regulate the PHO pathway in low phosphate conditions potentially at the transcriptional level. (9). It would thus be interesting to test whether the *Drosophila uL11* mutants, and especially $uL11^{K3Y}$, display altered resistance to stresses that affect ribosome biogenesis, for instance by raising them with specific food diets.

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Figure 1: Genomic organization of the *uL11* locus

A – Genomic locus of the 2R chromosome containing the *uL11/RpL12* gene (coordinates in grey).

B – Sequence of the $uL11^{K3A}$ and $uL11^{K3Y}$ mutants.

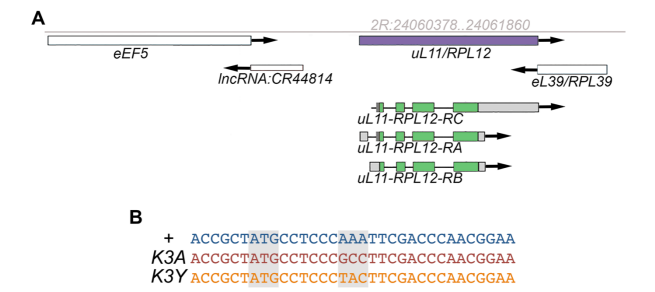


Figure 2: uL11K3me3 is undetectable in homozygous *uL11* mutants

Tubulin was used as a loading control. Whereas uL11 was present in hetero- and homozygous mutants, as revealed by the pan-uL11 antibody, uL11K3me3 was undetectable in the two homozygous mutants.

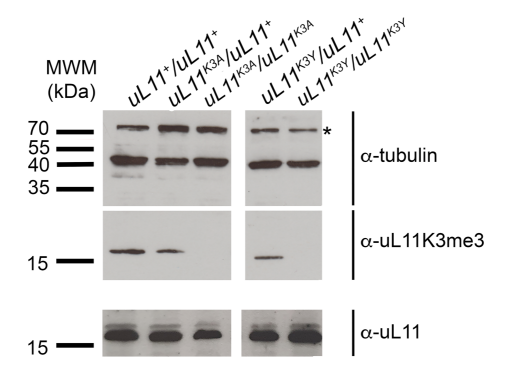


Figure 3: Life history traits of *uL11* mutants

A – Lethality of *uL11* mutants. $uL11^{x}/CyO$, Dfd-EYFP flies were crossed between them and 300 embryos per genotype were transferred on new medium ($uL11^{x}$: $uL11^{+}$, $uL11^{K3A}$ or $uL11^{K3Y}$, as indicated). **Left:** percentage of $uL11^{x}/uL11^{x}$ and $uL11^{x}/CyO$, Dfd-EYFP first instar larvae emerged from 300 embryos; **Right:** percentage of $uL11^{x}/uL11^{x}$ and $uL11^{x}/CyO$, Dfd-EYFP pupae coming from the emerged first instar larvae. Numbers of starting embryos and larvae are indicated.

 $uL11^{X}/uL11^{X}$ individuals: dark colour; $uL11^{X}/CyO,Dfd$ -EYFP individuals: light colour; blue: $uL11^{+}$, burgundy: $uL11^{K3A}$, orange: $uL11^{K3Y}$; grey: $uL11^{X}/uL11^{X}$ or $uL11^{X}/CyO,Dfd$ -EYFP.

B – Developmental time of uL11 mutants. The percentage of flies emerged from day 8 to 15 is shown. The total number of emerged flies is indicated. Solid lines: heterozygous flies; dotted line: homozygous flies; blue: $uL11^+$, burgundy: $uL11^{K3A}$, orange: $uL11^{K3Y}$.

Chi² test: *** p-value < 0.001; ** p-value < 0.01; * p-value < 0.05; only significant comparisons are shown.

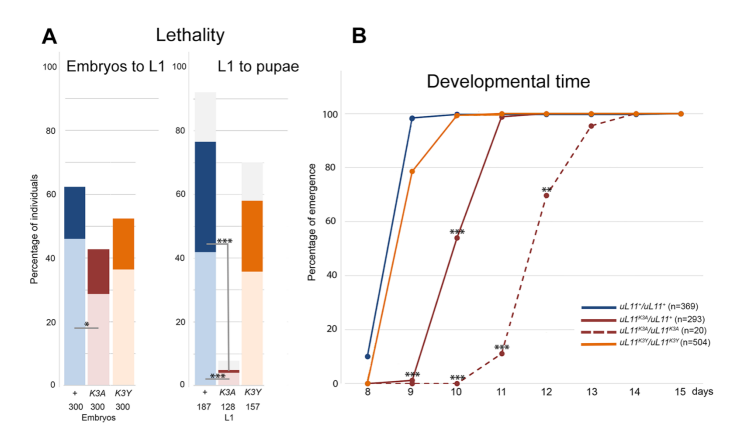


Figure 4: Analysis of bristles and wings in male *uL11* mutants

A – Thorax of a wild-type male. Anterior and posterior scutellar bristles are colorized.

B – Thorax of a $uL11^{K3A}/uL11^{K3A}$ male. Anterior and posterior scutellar bristles are colorized. Note that they look thinner and shorter that those of the wild-type male shown in A.

C – Length of anterior scutellar bristles of wild-type males (blue; n = 45), $uL11^{K3A}/uL11^+$ and $uL11^{K3A}/uL11^{K3A}$ (dark and light burgundy, n = 55 and n = 24, respectively) and $uL11^{K3Y}/uL11^{K3Y}$ (orange, n = 51).

D – Length of posterior scutellar bristles of wild-type males (blue; n = 44), $uL11^{K3A}/uL11^+$ and $uL11^{K3A}/uL11^{K3A}$ (dark and light burgundy, n = 53 and n = 13, respectively) and $uL11^{K3Y}/uL11^{K3Y}$ (orange, n = 50).

E – Wing length of *uL11* wild-type males (blue; n = 28), $uL11^{K3A}/uL11^+$ and $uL11^{K3A}/uL11^{K3A}$ (dark and light burgundy, n = 30 and n = 15, respectively) and $uL11^{K3Y}/uL11^{K3Y}$ (orange, n = 30).

t-tests: *** p-value < 0.001; ns: non significant.

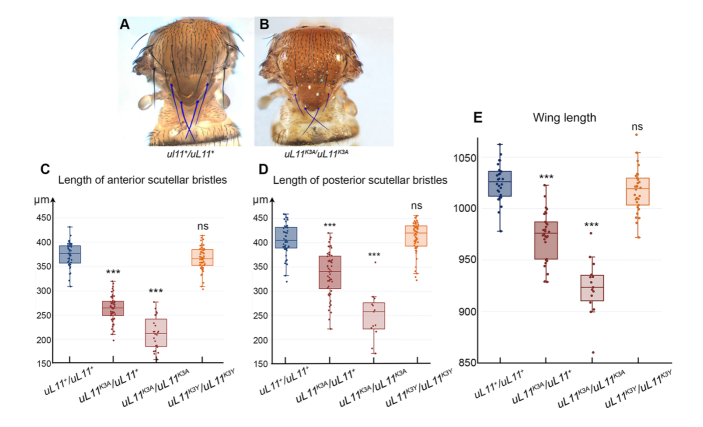


Figure 5: Translational rate of *uL11* mutants

A – Western blot showing puromycin incorporation in larvae of the two mutants as compared to wild-type larvae, in absence (-) or presence (+) of cycloheximide (CHX), an inhibitor of translation. Puromycin incorporation was revealed with an anti-puromycin antibody. Histone H3, revealed with an anti-panH3 antibody, was used as loading control. MWM: molecular weight marker.

B – Quantification of the puromycin signal in the 4 genotypes without (-) or with (+) CHX treatment. The puromycin signal was first normalized to the H3 signal, then to the wild-type signal. Student's t-tests were performed to compare puromycin incorporation in mutant and wild-type larvae. * p-value < 0.05; ns: non significant.

C – Polysome fractionation: cytoplasmic lysates (left) and EDTA-treated lysates (right) from S2 cells expressing uL11WT-HA (top) or uL11K3A-HA (bottom) were fractionated by centrifugation onto a sucrose gradient. Optical density at 254 nm was monitored during fractionation (top panels). The peaks observed in the gradient correspond to the different ribosomal complexes: 40S subunit, 60S subunit, 80S monosome, polysomes. Proteins extracted from fractions were analyzed by Western blotting with anti-HA antibody (lower panels). A vertical line indicates that different wells from the same gel were juxtaposed in the image for clarity. Images are representative for three obtained replicates.

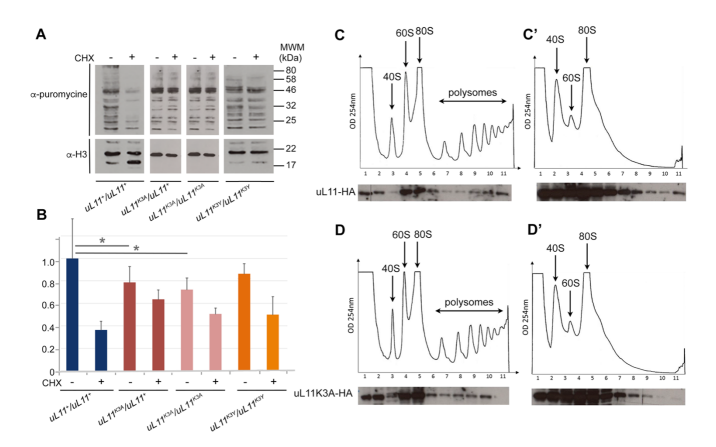


Figure 6: uL11 but neither uL11K3A nor uL11K3Y co-immunoprecipitates with the chromodomain of Corto.

S2 cells were co-transfected with plasmids expressing FLAG-CortoCD and uL11-Myc, uL11K3A-Myc or uL11K3A-Myc. Immunoprecipitations were performed with anti-FLAG antibodies (α -FLAG) or anti-HA antibodies (mock) and Western blot revealed using α -FLAG or anti-Myc antibodies (α -Myc). Spnt: supernatant, IP: immunoprecipitation. **A -** FLAG-CortoCD co-immunoprecipitated with uL11-Myc (arrow). **B -** FLAG-CortoCD did not co-immunoprecipitate with uL11K3A-Myc. **C -** FLAG-CortoDCD did not co-immunoprecipitate with uL11K3Y-Myc.

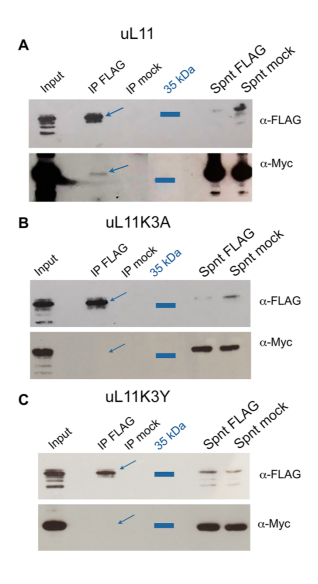


Figure 7: RNA-seq analyses of *corto* and *uL11* mutants.

A - Venn diagrams showing the intersection of genes deregulated in $corto^{L1}/corto^{420}$, $uL11^{K3A}$ and $uL11^{K3Y}$ up- and down-regulated genes together (cutoffs: adjusted p-value < 0.05; $log_2(fold-change) < -0.5$ ou > 0.5). See Supplementary Table 2 for detailed gene lists. The three genes validated by qRT-PCR are indicated.

B - RT-qPCR analysis of *CG13516*, *Hsp67Bc* and *GstE6* expression in $corto^{L1}/corto^{420}$, $uL11^{K3A}$ and $uL11^{K3Y}$ wing imaginal discs. Expressions were normalized on the geometric mean of *GAPDH* and *Spt6*. Mean of three replicates. Error bars correspond to standard deviations. * p-value < 0.05; ** p-value < 0.01; *** p-value < 0.001.

C – 21 nucleotide-long motif (top) found in the *cis*-regulatory sequences of 39 genes deregulated in the $uL11^{K3Y}$ mutant (see Supplementary Table 4 for details). This motif significantly overlaps the transcription factors Mad (idmmpmm2009 database, p-value 4.9e-07) and Med (dmmpmm2009 database, p-value 1.22e-04) motifs.

D – Snapshots showing the mapping of reads on CG13516, Hsp67Bc and GstE6 in the reference line w^{1118} , $corto^{420}/corto^{L1}$, $uL11^{K3A}$ and $uL11^{K3Y}$ wing imaginal discs.

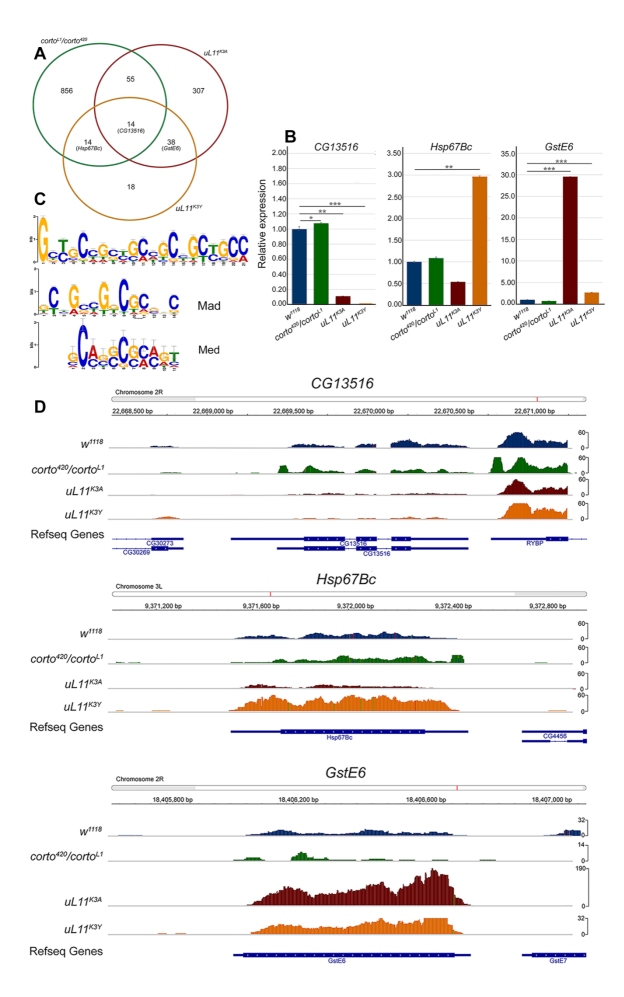


Table 1: RNA-seq of wing imaginal discs (GEO accession number GSE181926).

Sample name	Genotype	Total reads	Aligned reads	Unmap reads	Reads with multiple alignment	Reads used for analysis
wc_1	w 1118 ; corto +	1.50E+07	6.70E+06	9.00E+05	2.20E+06	6.71E+06
wc_2	w 1118 ; corto +	1.50E+07	6.40E+06	1.20E+06	2.20E+06	6.44E+06
cortoL1420_1	w 1118; corto L1/corto 420	2.71E+07	1.18E+07	1.20E+06	1.20E+07	1.18E+07
cortoL1420_2	w 1118 ; corto L1 /corto 420	2.71E+07	1.14E+07	1.90E+06	1.20E+07	1.14E+07
w_1	w ¹¹¹⁸ ; uL11 ⁺	2.58E+07	3.88E+07	5.39E+05	1.71E+07	2.08E+07
w_2	w ¹¹¹⁸ ; uL11 ⁺	3.22E+07	4.03E+07	3.50E+05	1.17E+07	2.79E+07
w_3	w ¹¹¹⁸ ; uL11 ⁺	3.49E+07	4.87E+07	6.90E+05	1.87E+07	2.86E+07
K3A_1	w ¹¹¹⁸ ; uL11 ^{K3A}	2.53E+07	3.34E+07	1.62E+06	1.14E+07	2.00E+07
K3A_2	w ¹¹¹⁸ ; uL11 ^{K3A}	4.01E+07	1.54E+07	4.96E+06	1.38E+08	1.06E+07
K3A_3	w ¹¹¹⁸ ; uL11 ^{K3A}	3.28E+07	8.30E+07	2.93E+06	6.18E+07	1.97E+07
K3Y_1	w ¹¹¹⁸ ; uL11 ^{K3Y}	3.07E+07	3.87E+07	4.65E+05	1.14E+07	2.64E+07