Allele-specific alternative splicing of Drosophila Ribosomal protein S21 suppresses a lethal mutation in the Phosphorylated adaptor for RNA export (Phax) gene

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Abstract

Genetic disruptions to the biogenesis of spliceosomal small-nuclear ribonucleoproteins in Drosophila cause wide-spread alternative splicing changes, including changes to the splicing of pre-mRNA for Ribosomal protein S21 (RpS21). Using a transposon mutant for the Phosphorylated adaptor for RNA export (Phax) gene, we demonstrate that changes in the splicing of RpS21 transcripts have a strong influence on the developmental progression of PhaxSH/SH mutants. Different alleles of the Drosophila RpS21 gene are circulating in common laboratory strains and cell lines. These alleles exhibit differences in RpS21 intron retention and splicing efficiency. Differences in the splicing of RpS21 transcripts account for prior conflicting observations of the phenotypic severity of PhaxSH/SH mutant stocks. The alleles uncover a strong splicing enhancer in RpS21 transcripts that can fully suppress the larval lethality and partially suppress the pupal lethality exhibited by PhaxSH/SH mutant lines. In the absence of the splicing enhancer, the splicing of RpS21 transcripts can be modulated in trans by the SR-rich B52 splicing factor. As PhaxSH/SH mutants exhibit wide-spread splicing changes in transcripts for other genes, findings here establish the importance of a single alternative splicing event, RpS21 splicing or intron retention, to the developmental progression of Drosophila.

Keywords: phosphorylated adaptor for RNA export protein; Phax; ribosome; ribosomal protein; ribosomal protein S21; RpS21; alternative splicing; alternative polyadenylation; mRNA processing; snRNP biogenesis; Spinal Muscular Atrophy; oho23B; splicing factor; cis-element; exonic splicing enhancer; B52; survival motor neuron

Introduction

Depletion of core splicing machinery leads to specific, nonrandom changes in alternative splicing and gene expression, including changes to ribosomal protein (RP) coding genes (Pleiss et al. 2007; Saltzman et al. 2011; Garcia et al. 2016). Conversely, targeted inhibition of the transcription of highly expressed RP coding genes liberates the "hungry" spliceosome and results in distinct changes to alternative splicing (Munding et al. 2016; Talkish et al. 2019). Similarly, disease-associated mutations in RP coding genes and corresponding reductions in ribosomes cause specific changes in gene expression and the translation of select transcripts (Horos et al. 2012; Khajuria et al. 2018). The specificity of responses to different perturbations in the overall gene expression program supports an interconnected regulatory network for the biogenesis and homeostatic regulation of the spliceosome and the ribosome that intersects in the RNA processing of RP pre-mRNA.

Our prior transcriptomic profile of small-nuclear ribonucleo-protein (snRNP) biogenesis mutants in Drosophila identified wide-spread alternative splicing changes in Phosphorylated adaptor for RNA export (Phax) mutants and Survival motor neuron (Smn) mutants (Garcia et al. 2016). Phax functions as a small-nuclear RNA (snRNA)-specific adapter to export Sm-class snRNAs from the nucleus, and Smn functions to assemble exported snRNAs into snRNPs in the cytoplasm (Ohno et al. 2000; Ohno 2012; Matera and Wang 2014). Full loss of fly Smn function results in a recessive larval lethality in Drosophila (Chan et al. 2003; Rajendra et al. 2007; Chang et al. 2008). In contrast, lethality can be genetically separated from a disruptive P element insertion in the Drosophila Phax gene, an insertion previously classified as lethal (Oh et al. 2003; Kahsai et al. 2016). Our lab and others have observed adult flies that are homozygous for this P element insertion in the Phax gene (Kahsai et al. 2016). Nevertheless, both Phax and Smn mutants exhibit lower steady-state levels of Sm-class snRNAs and an overlapping set of alternative splicing changes that includes changes in RP pre-mRNA, including a large change in the alternative splicing of pre-mRNA for Ribosomal protein S21 (RpS21) (Garcia et al. 2016).

RpS21 is component of the small subunit of the ribosome with functions in translation initiation and ribosome biogenesis.
changes in pressing the lethality of different alleles uncover sequences in transcripts with dramatic differences in alternative splicing. The in phenotypes between native splicing in other snRNP biogenesis mutants, the similarity in the context of our identification of large changes in mutant overgrowth of hematopoietic tissues (Schenkel et al. 2002). In the context of our identification of large changes in RpS21 alternative splicing in other snRNP biogenesis mutants, the similarity

Here, we identify alternative alleles of RpS21 that encode transcripts with dramatic differences in alternative splicing. The different alleles uncover sequences in RpS21 exon 4 that function to enhance the splicing of exon 3 to exon 4. Remarkably, the RpS21 allele with the splicing enhancer sequences was capable of suppressing the lethality of a Phax P element insertion mutant. Hence, findings here reveal a discrete genetic background difference that can account for prior conflicting observations of the viability of different fly stocks from this Phaxmutant line. Importantly, in the background of wide-spread alternative splicing changes, restoration of a single splicing event in RpS21 was ultimately able to rescue the developmental progression of homozygous Phaxmutant flies.

Materials and methods
Fly strains and husbandry

Stools were maintained on comroal and agar at room temperature (25 ± 1°C) in half-pint bottles. For crosses, sorting and the isolation of larvae, stocks were cultured on molasses and agar at room temperature (23 ± 3°C) unless indicated otherwise. Stocks for wild-type Ore-R (Oregon-R-modENCODE) and the reference genome strain iso-1 (y[1]; Gr22b[1] Gr22d[1] cn[1] CG33964[R4.2] bw[1] speck[1] MatProx[1] GstD5[1] Rh6[1]) were obtained directly from the Bloomington Drosophila Stock Center (BDSC). Stocks for Canton-S (Can-S) and w1118 were obtained from D.A. Harrison. Recombination of the UAS:Phax-mVenusexpression line in the mutant (P[JacW] Phaxmutant) (Oh et al. 2003) was previously described (Garcia et al. 2016). Here, Phaxmutant refers to homozygous lines with the P element insertion in the Phax gene (P[JacW] Phaxmutant). The armadillo (arm) promoter-GAL4 driver (P[GAL4-arm. S11] (Sanson et al. 1996) was maintained in the Phaxmutant background.

To generate Phaxmutant mutant flies that were heterozygous and homozygous for the different RpS21 alleles, we crossed virgin female flies with a second chromosome balancer (Cy, Actin-GFP), the RpS21short allele, and the P element insertion in the Phax gene to Canton-S wild-type male flies, homozygous for the RpS21long allele, described below. Meiotic recombination was allowed to take place in noncurly virgin female progeny from this initial cross. These virgin females were crossed to curly male progeny with the RpS21allele and the P element in the Phax gene, using a proteinase K (New England Biolabs Inc) method for the nonlethal isolation of DNA from Drosophila wings (Carvalho et al. 2009). Separate crosses were set up with balanced flies, containing the P element insertion in the Phax gene and different combinations of the RpS21long and RpS21short allele. Through genotyping of unbalanced larvae, we isolated 3 separate stocks for each of the following genotypes: Phaxmutant, RpS21/−, Phaxmutant, RpS21/, and Phaxmutant, RpS21/−. For viability experiments, larvae were sorted on molasses-agar plates and transferred to cornmeal-agar vials. We sorted an average number (n) of 76 larvae for each replicate of the 23°C GAL4:UAS viability experiment, 100 larvae for the 27°C GAL4:UAS study, and 30 larvae for the viability study with homozygous or heterozygous RpS21 alleles in the Phaxmutant background. A Student’s t-test was used to calculate P-values.

Genotyping

DNA from whole animals or cells was isolated for genotyping with a Quick-DNA Tissue/Insect Miniprep Kit (Zymo Research) and a 4-place Mini Bead Mill Homogenizer (VWR) according to the manufacturer’s protocol. Oligonucleotides used for PCR with Apa Taq Red (Genesee Sci) of the long allele or the short allele are listed in Supplementary Table 1. Gene-flanking oligonucleotides were used to clone the RpS21 gene from Oregon-R flies, S2-DRSC cells, and Kc167 cells into the HindIII ampBamHI sites of pBlueScript II SK+ (Agilent) for sequencing with M13 forward and reverse primers. To genotype the Phaxmutant mutant, we used 2 sets of oligonucleotides. We used primers complementary to the Phax gene that flanked the P element insertion, and we used a P element out primer, an M13 sequencing primer, with a primer complementary to the 3’-end of the Phax gene. This oligonucleotide set confirmed the inverse directionality of the P element in relationship to the forward direction of the Phax gene. For primer design, gene and intergenic sequences were derived from Flybase and compatible with the current release FB2022_02 http://flybase.org/ (Larkin et al. 2021).

Splicing mini-gene reporter design

Different wild-type and mutant RpS21 gene sequences, surrounding intron 3, were subcloned into the pKm1 and BamHI sites in pAc5.1B-EGFP (Addgene plasmid # 21181; Elisa Izaaurralde depositor). Wild-type and mutant gene sequences were generated as double-stranded DNA gene fragments (IDT gblocks). Sequences for the minimal splicing reporter corresponded to the RpS21 gene from nucleotide 400 to nucleotide 611, based on the iso-1 reference sequence. These RpS21 gene sequences spanned the third intron from exon 3, 104 nucleotides upstream of the 5′-splice site, to immediately downstream of the AATAAA polyadenylation signal in exon 4, 50 nucleotides downstream of the intron 3 3′-splice site.

Cellfectin II Reagent (Gibco) was used for transfections of S2-DRSC according to the manufacturer’s protocol. RNA was isolated on day 3 post transfection. RT-PCR was performed as indicated below with mini-gene specific primers (Supplementary Table 1). PCR products were separated on an agarose gel, stained with GelRed (Biotium), and imaged on a Gel Doc Ez (BIO-RAD). Bands were quantified with Image Lab 6.0.1 software (BIO-RAD). Percent Spliced In (PSI) was quantified as the percentage of spliced over total transcripts, and P-values were determined with a Student’s t-test.

Cell culture and RNA interference

Drosophila S2-DRSC (DGRC Stock 181; https://dgrc.bio.indiana.edu/stock/181; RRID: CVCL_Z992) and Kc167 (DGRC Stock 1; https://dgrc.bio.indiana.edu/stock/1; RRID: CVCL_Z834) cells were acquired directly from the Drosophila Genomics Resource Center (DGRC) (Schneider 1972; Cherbas et al. 1988). Both cell
lines were cultured in Schneider’s Drosophila Medium supplemented with 10% fetal bovine serum and 1x penicillin–streptomycin–glutamine (Gibco). RNA from S2-DRSC cells was used for reverse transcription with Superscript III (Invitrogen) to make cDNA. This cDNA and Apex Taq Red (Genesee Sci) were used with T7 promoter-fusion primers (Supplementary Table 1) to amplify BS2. Sequences for T7-fusion primers were derived from cell-screening “R” and “S” primer sets from the Drosophila RNAi Screening Center (DRSC) (Hu et al. 2020). PCR products served as templates for the overnight in vitro transcription with MEGAscript (Invitrogen) to generate double-stranded RNA for RNA interference (RNAi). Approximately 15 µg of dsRNA was added to each sample in a separate well of a 6-well tissue culture dish. The dsRNA was added on the first, third, and fifth days of the treatment. Samples were taken for RNA isolation on day 6.

HeLa and HepG2 cell pellets were generously supplied by the labs of K.A. Fields and B.T. Spear, respectively. Human cell pellets were immediately put in TRIzol reagent (Invitrogen) and RNA isolated for RT-PCR with human specific primers, as indicated below.

Western blotting

Protein was isolated from wild-type and sorted larvae approximately 4 days post-egg laying. Protein was isolated by crushing larvae in 1x RIPA Buffer (Thermo Scientific) supplemented with Halt Protease (Thermo Scientific) and Halt Phosphatase (Thermo Scientific) inhibitor cocktails. Lysates were precipitated with trichloroacetic acid, washed with methanol, and dried (80°C for 10 min) before resuspension in 1x Sample Loading Buffer (LI-COR). Protein samples were separated on NuPAGE 4–12% Bis–Tris gels in LI-COR and blocked with Odyssey Blocking Buffer (TBS) (indicated above, as the percentage of spliced over total tran- 

tron of human RpS21 was used at a 1:30,000 concentration. A rabbit polyclonal antibody was generated to an antigen for the protein product of CG33057 by ABclonal. A CG33057 gene fragment (amino acids 1–212) was subcloned into the pGEX-4T-AB1 vector for protein expression and purification. Primary antibodies for CG33057, RpS21 (Abcam # ab90874), and RpS6 (C.896.4) (Invitrogen) were used here to the manufacturer’s protocol. Isolated RNA was subjected to an additional round of DNase with TURBO DNase (Invitrogen), followed by phenol chloroform extraction. Reverse transcription (RT) was performed with Superscript III (Invitrogen) and random hexamers. PCR was performed with Apex Taq Red (Genesee Sci), and limited cycles of amplification with intron flanking oligos. PCR products were separated on an agarose gel in 1x TBE, stained with GelRed (Biotium), and imaged on a Gel Doc Ez (BIO-RAD). Separated bands were quantified on unaltered images with Image Lab 6.0.1 software (BIO-RAD). PSI was quantified as indicated above, as the percentage of spliced over total transcripts. For statistical analysis, P-values were determined with a Student’s t-test. Real-time PCRs of cDNA were conducted on a StepOnePlus System (Applied Biosystems), using Maxima (Thermo Scientific) or PowerUp (Applied Biosystems) SYBR Green/ROX master mixes. Three biological replicates were tested for each genotype. The ΔΔCt method was used to quantify differences, and P-values were determined with a Student’s t-test. Gene-specific primer sequences are listed in Supplementary Table 1.

Results

RNA-seq analyses

RNA-seq analysis was performed on original fastq files that were previously deposited in the NCBI Gene Expression Omnibus (GEO) (Edgar et al. 2002). Files are indicated below. Transcript abundance was quantified with kallisto (Bray et al. 2016) and differential analysis was performed with sleuth (Pimentel et al. 2017). HISAT2 (Kim et al. 2019) and Samtools (Danecek et al. 2021) were used to align RNA-seq reads to Release 6 of the Drosophila melanogaster genome (Hoskins et al. 2015) for visualization and Sashimi plot comparison with the Integrative Genomics Viewer (Robinson et al. 2011, 2017).

The splicing pattern of the predominant terminal exon of RpS21 is highly conserved from flies to humans

Splicing of the third intron of RpS21 pre-mRNA modulates the carboxy-terminus of the encoded RpS21 protein (Fig. 2a). The last 3 amino acids at the carboxy-terminus of the RpS21 protein are lysine (K), asparagine (N), and phenylalanine (F). The splice junction between exons 3 and 4 spans the K codon. The 3 carboxy-terminal amino acids of fly RpS21 are conserved with the human RpS21e protein, and, most notably, the human RpS21e transcripts also include a splice junction that spans the terminal K codon (Fig. 2a). In unspliced RpS21 transcripts, the intron contains the third wobble base of the terminal K codon and a following stop codon. The same pattern is conserved in the corresponding intron of human RpS21e transcripts. Human cell lines also show evidence of retention of the corresponding intron of RpS21e (Fig. 2b). Hence, this is an exceptionally well-conserved splicing pattern from flies to humans.
Phax^{SH/SH} mutants exhibit an expected decline in viability but a surprising lack of any adult escapers

As this conserved splicing event in RpS21 transcripts was altered in Phax^{SH/SH} mutant flies, Phax^{SH/SH} mutants and a UAS-Phax rescue transgene were used to determine how changes in this splicing event might contribute to organismal viability. In agreement with our previous findings (Garcia et al. 2016), the Phax^{SH/SH} mutants exhibited a large decrease in viability, as compared to wild-type Oregon-R controls (Supplementary Fig. 1a). Also, as before (Garcia et al. 2016), this decrease in viability was partially rescued by the ubiquitous expression of a wild-type UAS-Phax transgene with an armadillo-GAL4 driver (Supplementary Fig. 1a). However, unlike the previous study, ubiquitous expression of the UAS-Phax transgene did not lead to the eclosure of any adult flies (Supplementary Fig. 1a). In our previous study, approximately 30% of Phax-transgene expressing larvae developed to adulthood (Garcia et al. 2016). As the yeast-derived GAL4-UAS system is optimal at approximately 27°C, we also assayed viability at 27°C. Temperature did not account for the discrepancies between our current and prior viability studies, as viability at 27°C was roughly equivalent to the viability at 23°C (Supplementary Fig. 1a and b). Prior to our rescue experiments and the site-specific integration of the UAS-Phax transgene into the Phax^{SH/SH} mutant...
background, adult escapers, homozygous for the P element insertion in the Phax gene, were intermittently detectable in different stocks of this Phax^{SH/SH} mutant line (Oh et al. 2003; Kahsai et al. 2016; Garcia et al. 2016). We hypothesized that subtle genetic background differences might contribute to the observed differences in the viability of different stocks of the Phax^{SH/SH} mutant and in our transgenic rescue experiments.

**Proper splicing across intron 3 of RpS21 is linked to Phax expression**

Changes in RpS21 intron 3 retention parallel the observed changes in viability of Phax^{SH/SH} mutant and rescue animals. The splicing across RpS21 intron 3 is normally inefficient, as wild-type Oregon-R larvae exhibit a high baseline level of intron 3 retention (Fig. 3a). In Oregon-R animals, the percentage of fully spliced RpS21 transcripts across intron 3 averaged a low 56 %SI (Fig. 3b). Splicing across intron 3 was worse in the Phax^{SH/SH} mutants, evident as a visible increase in the ratio of unspliced to spliced transcripts (Fig. 3a) and a decrease in the percentage of fully spliced transcripts to 34 %SI (Fig. 3b). These changes in RpS21 splicing were fully rescued by ubiquitous expression of the UAS-Phax transgene (Fig. 3, a and b). Nevertheless, the rescue of RpS21 alternative splicing did not fully restore the developmental progression in UAS-Phax expressing animals.

**Phax^{SH/SH} mutant changes in alternative splicing affect the steady-state levels of encoded proteins**

Anticipated changes in steady-state protein levels accompanied the changes in alternative splicing in the Phax^{SH/SH} mutants. Phax^{SH/SH} mutants exhibited a decrease in steady-state levels of RpS21 protein relative to a tubulin loading control (Fig. 3, c and d). Levels of RpS21 protein were rescued by expression of the UAS-Phax transgene (Fig. 3, c and d). Protein levels of another small subunit protein, ribosomal protein S6 (RpS6), were also down in the Phax^{SH/SH} mutants (Fig. 3, c and d). Although small changes in RpS6 transcripts were found in our transcript-specific analysis here, changes in RpS6 alternative splicing were not uncovered in our more stringent prior analysis (Garcia et al. 2016). Hence, changes in RpS6 protein levels could reflect the more dramatic changes in RpS21 alternative splicing or broader splicing disruptions exhibited by the Phax^{SH/SH} mutants. As an additional control for anticipated protein level changes, we blotted for the protein product of the computed gene CG33057, which encodes an ortholog of the yeast tRNA 2'-phosphotransferase protein (TPT1). Transcripts for CG33057 are contained entirely within an intron of transcripts for the monkey king protein (mkg-p) gene that is retained in snRNP biogenesis mutants (Garcia et al. 2016). We detected an increase in levels of a protein of the expected molecular weight for the fly TPT1 ortholog in lysates from Phax^{SH/SH} mutants (Fig. 3c). Protein levels of the putative fly TPT1 ortholog

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**Fig. 3.** Differential abundance of RpS21 transcripts and RpS21 protein levels in Phax^{SH/SH} mutant and rescue animals. a) Increase in RpS21 intron 3 retention in Phax^{SH/SH} mutants and corresponding decrease upon expression of the Phax transgene. Transgenic rescue animals (arm>Phax) utilized an armadillo-GAL4 driver (arm) and a UAS:Phax-mVenus transgene (Phax) in the Phax^{SH/SH} mutant background. Image of a GelRed-stained agarose gel of RT-PCR products, amplified with primers flanking the predominant terminal exon junction, encompassing intron 3 of Drosophila RpS21 transcripts. Gels include PCR products or negative controls from reactions with (RT +) or without reverse transcriptase (RT −), respectively. Gel image was adjusted as in Fig. 2b. Of note, RpS21 terminal intron 4 from Fig. 1a is rarely spliced out of RpS21 transcripts that utilize the distal polyadenylation signal. RpS21 distal polyadenylated transcripts are also relatively low compared to proximal polyadenylated transcripts in the whole animals and cells used here. b) Quantification of the percentage of splicing from exon 3 to exon 4 in (a). The graph is the percentage of spliced to total transcripts, shown as PSI. c) Western blots for the indicated proteins. d) Quantification of the RpS21 and RpS6 protein levels relative to α-tubulin from (c). Relative levels of these proteins in Oregon-R were set to 100%. *P < 0.05, **P < 0.001, and ***P < 0.0001.
returned to baseline levels upon expression of the UAS-Phax transgene (Fig. 3c). However, like the noted changes in RpS21 alternative splicing, rescue of protein levels was not sufficient to restore the developmental progression of UAS-Phax transgene expressing animals.

**Common Drosophila flies and cell lines have different alleles of the RpS21 gene**

DNA sequencing revealed a surprising allelic heterogeneity in exon 4 of RpS21, immediately downstream of intron 3, in common laboratory flies and cell lines. Using RpS21 gene-flanking primers, we subcloned and sequenced the RpS21 gene in Oregon-R flies, S2-DRSC cells, and Kc167 cells. Surprisingly, RpS21 exon 4 from S2-DRSC cells and Oregon-R flies contained a 7-nucleotide deletion and an additional adenine to thymine transversion relative to Kc167 cells and the Drosophila genome reference strain iso-1 (Fig. 4a). Available genome assemblies also support the identification of allelic heterogeneity in RpS21 exon 4 amongst different Drosophila cell lines (Supplementary Fig. 2). This subtle genetic difference was used to design allele-specific primers to genotype additional fly lines. The PhaxSH/SH mutants, Armadillo-GAL4 rescue line (arm), and the widely used w1118 mutant contained the short (S) RpS21S allele found in Oregon-R flies and S2-DRSC cells (Fig. 4b). The alternative wild-type Canton-S flies had the long (L) RpS21L allele like iso-1 flies and Kc167 cells (Fig. 4b). The genetic difference in exon 4 overlapped a predicted Exonic Splicing Enhancer (ESE) in the longer RpS21L allele (Cartegni et al. 2003). Therefore, we hypothesized that the discrete genetic differences in exon 4 could affect the splicing across intron 3 of RpS21 pre-mRNA.

**Allelic differences in the RpS21 gene correspond to differences in the splicing of RpS21 transcripts**

Indeed, the genetic differences in exon 4 correlated with the expected differences in intron 3 retention in RpS21 transcripts (Fig. 4, c and d). RpS21 transcripts from Oregon-R and w1118 flies exhibited a high level of intron 3 retention and a relatively low, approximately 60 PSI, level of spliced transcripts. In contrast, Canton-S and iso-1 flies with the longer RpS21 allele, containing the predicted ESE, had little to no intron 3 retention and a high, greater than 99 PSI, level of spliced transcripts (Fig. 4c). This pattern was also evident in Drosophila Kc167 cells with the long RpS21 allele vs. S2-DRSC cells, lacking theputative ESE (Fig. 4, d and e). Kc167 cells exhibited greater than 99 PSI with no RpS21 intron 3 retention, and S2-DRSC cells had a high level of intron 3 retention and a low, 55 PSI, level of spliced transcripts (Fig. 4, d and e). The increased splicing efficiency of RpS21 transcripts from the longer allele supports the prediction that the additional GC-rich sequences in exon 4 function as a strong ESE.

**Predicted ESE sequences in exon 4 of RpS21 transcripts enhance the splicing of exon 3 to exon 4**

We constructed an RpS21 mini-gene splicing reporter to further test the prediction that the additional sequences in exon 4 of the long RpS21 allele function as an ESE. The minimal RpS21 mini-gene spanned the 3′-end of exon 3 to the end of exon 4 with (miniS21S) and without (miniS21L) the putative ESE in exon 4 (Supplementary Fig. 3a). The splicing reporter utilized a heterologous Act5C promoter and an SV40 polyadenylation signal (Supplementary Fig. 3a). We created an additional mini-gene without the predicted ESE sequences that contains a consensus 5′-splice site (miniS21S A>G) (Supplementary Fig. 3a). The A>G mutation was designed to test the potential contribution of the unpaired A nucleotide in the 5′-spliced site to RpS21 intron 3 retention. As expected, the PSI of the miniS21L A>G reporter was significantly higher than the miniS21S reporter with the endogenous 5′-splice site, supporting a small contribution of this seemingly weak 5′-splice site to RpS21 intron 3 retention. However, the splicing efficiency of all 3 mini-gene reporters was greater than 80 PSI, which exceeded the inefficient splicing of endogenous transcripts from the RpS21 short allele (Fig. 4, c and d). This suggests that additional sequences that were not included in the mini-gene splicing reporter likely contribute to the high level of RpS21 intron 3 retention in transcripts encoded by the short allele. Nevertheless, the RpS21 mini-gene with the putative ESE in exon 4 (miniS21L) produced transcripts with significantly less intron retention and a greater PSI than transcripts from either mini-gene lacking the ESE sequences, which further supports the hypothesis that the additional nucleotides in the long allele function to enhance the splicing of RpS21 exon 3 to exon 4 (Supplementary Fig. 3, b and c).

**Knockdown of the B52 splicing factor decreases RpS21 intron 3 retention in RpS21 transcripts without the predicted ESE in exon 4**

A prior cross-comparison of published RNA-seq data for local splicing variations revealed a decrease in RpS21 intron 3 retention upon RNAi knockdown of mRNA for the splicing factor B52 in Drosophila S2 cells (Srivastava et al. 2021). As expected from these transcriptomic findings, B52 knockdown in S2-DRSC cells here led to a decrease in RpS21 intron 3 retention and a corresponding 30% increase in the percentage of spliced RpS21 transcripts (Fig. 5, a–c). However, B52 knockdown in Kc167 cells did not appear to affect the splicing across intron 3 of RpS21 in those cells, as the splicing efficiency of RpS21 in Kc167 cells without B52 knockdown is already very high (>99 PSI) (Fig. 5, a–c). In other words, the long allele of RpS21 with the predicted ESE in exon 4, appears to short circuit or counter the inhibitory regulation of pre-mRNA splicing of RpS21 exon 3 to exon 4 by B52.

TheRpS21 ESE-containing allele suppresses the pupal lethality of the PhaxSH/SH mutants

With mounting evidence for the high splicing efficiency of the long RpS21 allele, we asked whether this allele could restore the developmental progression of snRNP biogenesis mutants. Notably, the long allele of RpS21 was able to partially restore the developmental progression of PhaxSH/SH mutants (Fig. 6, a and b). At the pupal stage, the presence of the long RpS21 allele (RpS21L) significantly improved the viability of PhaxSH/SH mutants in an allele dose dependent manner (Fig. 6a). Furthermore, unlike PhaxSH/SH mutants with only the short RpS21 allele (RpS21S), PhaxSH/SH mutants that were either heterozygous for RpS21 (RpS21L/S) or homozygous for the long allele (RpS21L/L) produced adult escapers (Fig. 6b). To date, no progenies have been observed from any of the adult Phax escapers. Nevertheless, we conclude that different RpS21 alleles likely account for our prior observations of intermittent adult escapers in PhaxSH/SH mutant stocks.

The RpS21 ESE-containing allele decreases RpS21 intron 3 retention in the PhaxSH/SH mutants in an allele dose dependent manner

In the PhaxSH/SH mutant background, the presence of the long RpS21L allele rescues the splicing of exon 3 to exon 4 of RpS21 transcripts (Fig. 6c). PhaxSH/SH mutant larvae that are homozygous for the short RpS21S allele (RpS21S/S) exhibited the highest...
level of intron 3 retention and the lowest level of exon 3 and 4 splicing (21 PSI) relative to Oregon-R controls (Fig. 6, c and d) or larvae with even a single copy of the long RpS21L allele (Fig. 6, c and d). PhaxSH/SH mutants, which are heterozygous for the RpS21 short and long alleles (RpS21S/L), exhibited similar levels of intron 3 retention and splicing (58 PSI) as wild-type Oregon-R animals that are homozygous for the short allele (RpS21S/S) (Figs. 6, c and d and 3, a and b). PhaxSH/SH mutants that are homozygous for the long RpS21L allele (RpS21L/L) have levels of intron 3 retention and splicing (98 PSI) that are like wild-type Canton-S animals (99 PSI) with no genetic perturbations of snRNP biogenesis (Fig. 6, c and d). Hence, we conclude that the long RpS21L allele is sufficient to rescue the perturbation of RpS21 splicing caused by P element mutations in the Phax gene.

Discussion

The alternative splicing of pre-mRNA for RPs has the potential to regulate the overall abundance and composition of ribosomes. Changes in either ribosome abundance or composition can cause
specific changes to gene expression and affect the protein synthesis of select transcripts (Horos et al. 2012; Khajuria et al. 2018). Our prior transcriptome analysis of snRNP biogenesis mutants uncovered specific changes to the alternative splicing of RP pre-mRNA (Garcia et al. 2016). As shown here, the Phax^{R55H} mutant changes to RpS21 pre-mRNA correlated with a decrease in RpS21 and RpS6 protein abundance that is consistent with a putative decrease in ribosome numbers. In addition, RpS21 intron 3 retention encodes an RpS21 protein isoform that is 2 amino acids shorter at the carboxy-terminus. Thus, in addition to ribosome abundance, the modulation of the alternative splicing of RP pre-mRNA can subtly alter ribosome composition.

Disruptions to ribosome numbers have been previously linked to disease. Mutations in numerous RP coding genes, other than human RPS21e, are associated with the blood disorder Diamond-Blackfan anemia (DBA) (Narla and Ebert 2010; Horos et al. 2012; Mills and Green 2017; Khajuria et al. 2018; Aspesi and Ellis 2019; Costa et al. 2020). DBA-associated RP gene mutations have been linked to reductions in ribosome levels (Khajuria et al. 2018). A similar decrease in ribosome numbers has been observed in mouse models of the neuromuscular disease Spinal Muscular Atrophy (SMA) (Bernabò et al. 2017). SMA is caused by mutations in the human Survival Motor Neuron 1 (SMN1) gene (Lefebvre et al. 1995). In addition to a reduction in axonal ribosomes, SMA model mice exhibited lower levels of Rps6 protein and a wide-spread decrease in the translation efficiency of RP transcripts, including transcripts for mouse Rps21 and Rps6 (Bernabò et al. 2017). The reductions of fly RpS21 and RpS6 proteins in the Phax^{R55H} mutants are consistent with the finding of disrupted ribosome homeostasis in SMA model mice. How specific alternative splicing events contribute to RP levels and overall ribosome numbers remains to be determined, but findings here suggest that mis-splicing of RP coding transcripts may contribute to the observed reduction in RP levels and possibly ribosome numbers. Importantly, however, the long ESE-containing RpS21 allele has, to date, been insufficient to suppress the larval lethality of fly Smn null mutants.

In the absence of the strong ESE in exon 4, RpS21 alternative splicing was negatively regulated in trans by the SR-rich B52 splicing factor. The human ESEfinder predicts a Serine and Arginine Rich Splicing Factor 6 (SRSF6)-binding site, nucleotides –GGCGUA–, that overlaps the allelic difference in exon 4 of RpS21 transcripts from the longer RpS21e allele (Fig. 4a) (Cartegni et al. 2003). To date, our RNAi screens have yet to identify an orthologous Drosophila splicing factor and/or component of the 3’-cleavage and polyadenylation machinery that binds to this putative ESE in exon 4 to enhance RpS21 splicing. The negative regulation of RpS21 splicing by B52, occurring in RpS21 transcripts without this ESE, likely requires additional cis-regulatory sequences beyond intron 3 and proximal sequences in the adjacent exons. The minimal RpS21 splicing reporter without the ESE had a notably high PSI (>-80 PSI) relative to the low PSI (~58 PSI) of corresponding endogenous RpS21 transcripts. In addition to a heterologous promoter and polyadenylation signal, the minimal reporter included only intron 3, the short downstream exon 4, and a mere 100 out of 192 nucleotides of upstream exon 3. B52 has a demonstrated affinity for structured RNA with accessible single-stranded binding sites (Shi et al. 1997). Binding of B52 to RpS21 transcripts may therefore require additional upstream or downstream sequences necessary for proper RNA structure.

Overexpression of B52 has been linked to increased growth and depletion to decreased growth (Fernando et al. 2015; Wada et al. 2021). In addition to RpS21, third-party analysis of published RNA-seq data of B52 knockdowns in S2 cells uncovered alternative splicing of yorkie transcripts, a transcriptional co-activator in Hippo signaling (Srivastava et al. 2021). Hippo signaling is a highly conserved pathway that controls organ size (Halder and Johnson 2011; Harinaran 2015; Yu et al. 2015; Meng et al. 2016). Depletion of 1 copy of the yorkie gene enhanced the dysregulated cell-growth phenotype of heterozygous RP mutants (Wada et al. 2021). Through the concerted regulation of the alternative splicing of RpS21 and yorkie transcripts, splicing factors like B52 can synchronize the ribosome with cell growth signals.

Proteomic and phospho-proteomic studies of the human RPS21e protein have identified posttranslational modifications at the carboxy-terminus of the protein that are likely impacted by the alternative splicing of the terminal exon of RPS21e transcripts (Choudhary et al. 2009; Mertins et al. 2013, 2016; Akimov et al. 2018). Overall, our findings are consistent with a model for the regulation of RpS21 through the alternative splicing of the
terminal RpS21 coding exon and corresponding modulation of the carboxy terminus of the protein. The allele-specific RpS21 alternative splicing patterns uncovered here will aid the study of the alternative splicing control of RpS21 protein isoforms and their potential roles in ribosome biogenesis and homeostasis.

Data availability

All Drosophila stocks and plasmids are available upon request. The author affirms that all data necessary for confirming the conclusions of the article are present within the article, figures, and tables. RNA-seq analysis was performed on original fastq.
files from the following GEO series accession numbers: Oregon-R, GSE49587 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE49587 and PhaxSH/S, GSE81121 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE81121.

Supplemental material is available at G3 online.

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Conflicts of interest
None declared.

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