Transcriptomic comparison of *Drosophila* snRNP biogenesis mutants: implications for Spinal Muscular Atrophy

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Abstract

Spinal Muscular Atrophy (SMA) is caused by deletion or mutation of the Survival Motor Neuron 1 gene (SMN1), but the mechanism whereby reduced levels of SMN protein lead to disease is unknown. SMN functions in the assembly of spliceosomal small nuclear ribonucleoproteins (snRNPs) and potential splicing defects have been uncovered in various animal models of SMA. We used disruptions in Smn and two additional snRNP biogenesis genes, Phax and Ars2, to classify RNA processing differences as snRNP-dependent or Smn gene specific. Although more numerous, the processing changes in Ars2 mutants were mostly distinct from those identified in Phax and Smn animals. Phax and Smn null mutants exhibited comparable reductions in steady-state snRNA levels, and direct comparison of their transcriptomic profiles uncovered a shared set of alternative splicing changes. However, these RNA processing differences were largely absent in three different models of intermediate SMA. Instead, the three hypomorphic Smn point mutants displayed an activation of stress responsive transcripts that correlated with phenotypic severity. In summary, genome-wide analyses showed that mRNA expression and alternative splicing defects are not conserved between hypomorphic and complete loss-of-function Smn mutants. Because disease phenotypes are unlikely to result from global changes in RNA processing, additional hypotheses for the etiology of SMA should be considered.
Introduction

Spinal Muscular Atrophy (SMA) results from genetic loss of the human *Survival Motor Neuron 1* (*SMN1*) gene [1], but uncovering how SMN protein depletion leads to the characteristic neuromuscular pathology of SMA has proven difficult. SMN functions in the assembly of Sm-class small nuclear ribonucleoproteins (snRNPs), core components of the spliceosome [2,3]. Because of this well-established function, prior studies investigating SMA etiology have sought to connect SMN depletion to alternative splicing changes and downstream SMA symptoms [4-6]. However, these studies typically do not account for indirect effects, like developmental delays, that might be responsible for the observed differences [7,8]. Thus, the evidence linking a reduction in snRNP levels to spliceosomal dysfunction and specific splicing changes is confounded by false positives. Hence, whether or not SMA phenotypes are caused by disruptions to snRNP levels and pre-mRNA splicing remains unsettled.

*Drosophila* models of SMA exhibit numerous small changes in splicing patterns that are difficult to separate from biological noise and normal developmental fluctuations [7]. In the fruitfly, steady-state levels of small nuclear RNAs (snRNAs) appear to be uncoupled from SMA-like phenotypes [9]. Studies in SMA mouse models also failed to uncover clear connections between SMN depletion, snRNP-levels, and pre-mRNA splicing. Although late-stage symptomatic mice show widespread splicing changes [4], these differences are not apparent prior to the onset of symptoms [10]. In these studies, global alternative splicing patterns may have masked important cell-type specific differences. Recently, tissue-specific disruptions to the splicing patterns of genes in motor neurons and glial cells were found in pre-symptomatic SMA model mice [6]. However, it is not clear that decreased snRNP levels caused the observed changes to the splicing pattern. Thus, disruptions
to snRNPs and splicing in motor neurons and glial cells of the central nervous system (CNS) may not be the sole drivers of SMA pathology.

Evidence from fruitfly and mouse SMA models suggests that high SMN levels are required in cells and tissues outside of the CNS. A combination of pan-neuronal and mesodermal specific SMN expression is required to rescue lethality in *Drosophila* SMN mutants, as expression in either cell-type alone was insufficient for rescue [11]. Similarly, restoring full SMN expression solely within motor neurons of SMA mice does not rescue viability [12,13]. Furthermore, SMA-like phenotypes in severe mouse models that utilize a human SMN2 transgene can be rescued by peripheral administration of splice-correcting antisense oligonucleotides (ASOs) [14]. Intriguingly, restoration of SMN expression in the CNS was not essential for phenotypic rescue, as CNS-specific treatment with a blocking oligonucleotide complementary to the splice-correcting ASO did not hamper the rescue [15]. Additional work in fly SMA models has shown that the entire neuronal circuit, including proprioceptive interneurons, is important for motor function, indicating that SMN plays a non cell-autonomous role in the maintenance of neuromuscular connections [16]. Thus, SMA is not a cell-autonomous defect of motor neurons in mice and flies, and studies that focus solely on RNA changes in motor neurons would thereby neglect important changes in other cell types.

Although the evidence connecting SMN loss (and rescue) to particular splicing changes is unclear, observations in diverse eukaryotes suggest that perturbations of the core splicing machinery can affect specific alternative-splicing events. Indeed, mutations in yeast genes encoding core spliceosomal factors disrupted the splicing of only a subset of pre-mRNAs [17,18]. Inhibiting highly expressed ribosomal protein coding genes in yeast also perturbed numerous alternative splicing patterns, suggesting that spliceosomal accessibility controls pre-mRNA splicing patterns [19]. In *Drosophila*, core splicing factors were uncovered by an RNAi screen for regulators of a set of
alternative splicing reporters [20]. In cultured human cells, knockdown of Sm proteins led to numerous changes in alternative splicing [21]. Thus, evidence from multiple systems indicates that modulating snRNP levels can alter splice site choice, however, a clear connection between snRNP levels and human disease has yet to be made.

In this study, we carried out transcriptome profiling of Drosophila Smn mutants, along with two additional snRNP biogenesis mutants in order to identify snRNP-dependent vs. SMN-specific changes in gene expression, premature cleavage and polyadenylation, and alternative splicing. Specifically, mutations in genes encoding the snRNA-specific nuclear export adaptor Phax (Phosphorylated adaptor for RNA export) and the Cap Binding Complex (CBC) associated protein, Ars2 (Arsenite resistance 2), were used for comparison [22-28]. The Phax mutant was particularly useful in identifying an overlapping set of snRNP-dependent RNA changes in our SMA model flies. We found that alternative-splicing differences shared between the Phax and Smn mutants can be rescued at both the RNA and protein levels by expression of Phax or Smn transgenes, respectively. In combination with a modest restoration of steady-state snRNA levels, observations from these rescue lines provide a connection between snRNP levels and specific pre-mRNA splicing events. However, the link between these changes and SMA phenotypes remains unclear.
Results

Phax and Smn mutants display similar steady-state snRNA decreases

RNA-sequencing (RNA-seq) of poly(A)-selected RNA was used to identify mRNA changes in snRNP biogenesis mutants, and northern analysis coupled with RNA-seq of total ribosomal RNA-subtracted (rRNA-) RNA was used to quantify steady-state snRNA levels. Browser shots of Phax, Smn, and Ars2 gene loci confirmed the gene disruptions in the respective mutant lines (Fig. 1A; [7]). In our previous RNA-seq analysis of Smn null mutants, publicly available modENCODE data were used to determine the developmental stage at which the mutants arrested [7]. By the same method, the Phax and Ars2 mutants exhibited a similar timing of developmental delay as did the Smn animals (Fig. S1A&B). Using developmentally appropriate wild-type controls (Fig. S2A), northern blots demonstrated comparable, near two-fold decreases in steady-state U1, U2, and U4 snRNAs in the Phax and Smn mutants (Fig. 1B&C). When assayed by northern blotting, U5 snRNA levels were essentially unaffected in the Phax mutants (Fig 1B&C), as were levels of nearly all of the snRNAs in the Ars2 mutants (Fig. 1B&C). When RNA-seq data were used to quantify snRNA levels, each of the three mutants appeared to exhibit snRNA deficits, but the extent varied by the method of normalization (Figure S2B&C). Only small (<15%) changes were observed in specific snRNA paralogs relative to the total (Fig. S3), quantified by using reads covering isoform-specific nucleotides [29]. Overall, the Phax and Smn mutants displayed the most consistent decreases in steady-state snRNA levels.

snRNP biogenesis mutants share a small number of gene expression changes

To identify shared and distinct changes in gene expression between snRNP biogenesis mutants, we used the TopHat/Cufflinks pipeline to quantify mRNA differences from our poly(A)-selected RNA-seq data [30]. Raw reads from age-matched wild-type (WT) controls and modENCODE second (L2) and
early third instar (L3 12h) developmental controls were integrated with sequencing data from the mutants into a single analytical pipeline. For comparison, we focused on genes whose expression changed uniformly and significantly (false discovery rate-adjusted p-value < 0.05) relative to each of the controls (L2, L3 12h, and our WT). A relatively small number of genes (186 out of ~15,000) met these stringent criteria for differential expression, and most of these differences were mutant specific (Fig. 2A&C and Tables S1-S4). A number of stress-responsive transcripts were among those that were specifically elevated in Smn mutants, though some of these were also up slightly in the Ars2 mutants (Fig. 2B). Phax mutants shared only twenty-eight gene expression changes with Smn mutants (Fig. 2C and Table S2). This small list of genes was enriched for genes involved in oxidation-reduction, as measured by DAVID gene ontology term analysis (p-value = 0.003) [31-33]. A notable gene expression change shared by the Phax and Smn mutants was a 5-fold increase in the Activity-regulated cytoskeleton associated protein 1 (Arc1) transcript (Table S2). Arc1 functions in a well conserved hyperlocomotion response to starvation, and hence elevated levels in Phax and Smn mutants might be an indirect consequence of a disruption in nutrient acquisition in these animals [34].

In summary, gene expression analysis uncovered a small number of overlapping changes that exceeded normal developmental fluctuations. The relationship of these changes to SMA etiology and/or pathology remains to be determined.

**snRNP biogenesis mutants exhibit a trend toward shorter mRNAs**

In addition to its role in splicing, U1 snRNP has been reported to control mRNA length by preventing premature cleavage and polyadenylation [35,36]. To determine whether reduced U1 snRNA levels in our snRNP biogenesis mutants correlated with expression of shorter mRNAs, mapped RNA-seq reads were analyzed using the DaPars linear regression algorithm [37,38]. DaPars output is
displayed as a Percentage of Distal PolyA site Usage Index (PDUI) for each transcript, or as the
difference in the PDUI value between wild-type and mutant (ΔPDUI). Pairwise comparisons of PDUI
values for wild-type and mutants can be used to visualize changes in RNA length. Whereas the Phax
and Smn mutants expressed transcripts that were both longer and shorter than their wild-type
counterparts, there were greater numbers of shortened mRNAs (Fig. 3A-C and Tables S5-S7).
Focusing on differences that are greater than two-fold, Phax and Smn mutants had a relatively small
number of mRNA length changes in common with each other (34). However, the changes were
largely distinct from those observed in Ars2 animals, who had more than twice as many mRNA length
changes as did the Phax and Smn mutants combined (Fig. 3D and Table S8). The overall trend of
shorter transcripts in the mutants does not appear to be a consequence of developmental delay, as
parallel DaPars analyses of modENCODE datasets of second to third instar larvae showed a trend
toward shorter transcripts in the more mature animals (Fig. S4). Among the few overlapping mRNA
length changes between Phax and Smn, one of the largest and most apparent changes was a
shortened mRNA for CG9662, a putative oligosaccharyl transferase complex subunit (ΔPDUI = -0.96,
Table S5). These results are generally consistent with the notion that reductions in U1 snRNP levels
predispose mRNAs to premature cleavage and polyadenylation. Because the Ars2 mutants exhibited
only a small decrease in steady state U1 levels (<10%; Fig. 1C), the mRNA length changes in these
animals are more likely due to disruptions of the CBC, which are known to affect a subset of pre-
mRNA termination events [26,39,40].

**Phax and Smn mutants share a set of alternative splicing pattern changes**

Our previous RNA-seq analysis of Smn null animals revealed many small-amplitude alternative
splicing differences (compared to wild-type controls), but it was unclear whether any of these changes
might be due to decreases in snRNP levels [7]. To ascertain whether any of these splicing changes are snRNP-dependent, we used the Mixture of Isoforms (MISO) probabilistic framework to compare splicing differences in Smn mutants with those identified in Phax and Ars2 mutants [41]. Simply put, the MISO algorithm estimates the expression of alternatively spliced isoforms in RNA-seq data, and reports it as a fraction of alternatively spliced events relative to the total [41]. MISO denotes this fraction as the ‘percent spliced in’ or ‘Psi’ [41]. Differences are reported here as an absolute change relative to the wild-type control, |deltaPsi|. Similar to our previous results with the Smn null animals, we found mostly small changes (less than two-fold, |deltaPsi| < 0.5) in alternative-splicing events in the Phax and Ars2 (Fig. 4A-C and Tables S9-S11). More than half of the splicing changes identified in the Smn null animals overlapped with changes in the Phax mutants, but neither the Phax nor Smn mutants had considerable overlap with those observed in the Ars2 mutants (Fig. 4D and Table S11). Of the top 150 differences in the Ars2 background, close to ninety percent of these changes were located in the 5'-proximal intron (Table S11). These changes likely reflect the association of Ars2 with the CBC and the role of this complex in promoter proximal splicing events [24] rather than Ars2-related changes in snRNPs. In contrast, the changes that were common to Phax and Smn mutants are more likely to be due to decreases in snRNP levels.

Transgenic rescue of alternative splicing changes induced by loss of Phax or Smn

As described previously, Smn is essential for viability in the fly [11], and transgenic expression of Flag- or GFP-tagged SMN protein in an otherwise null mutant background fully rescues the larval lethality and other SMA-like phenotypes [42,43]. Here, we found that Phax is also an essential gene, and that ectopic expression of Phax-GFP using the GAL4-UAS system rescued the lethality of Phax^{SH/SH} mutants (Table S12). We next assayed (and confirmed by semi-quantitative RT-PCR) a
number of the most apparent alternative splicing changes identified by RNA-seq (Fig. S5). We then tested the ability of the Phax and Smn transgenes to rescue four of these splicing events by quantitative qRT-PCR (Fig. 5B&C). Ubiquitous expression of Phax using an Armadillo-GAL4 driver significantly rescued all of the splicing changes we tested (Fig. 5B). Transgenic expression of Smn, driven by its native promoter, also rescued these alternative-splicing events in Smn null animals (Fig. 5C). Together, the results confirm that the observed changes we identified by RNA-seq are indeed caused by loss of Phax or Smn expression.

One of the splicing changes we detected is expected to produce an alternative (larger) dUTPase protein isoform. Therefore, we assayed the impact of this alternative splicing change by western blotting, using an antibody that targets both dUTPase isoforms. As shown in Fig. 5D, the expected increase in the ratio of the long isoform relative to the short one was observed in the mutants, and this change was largely rescued in both transgenic lines. Thus, the transcriptomic changes we identified in SMA model animals can (and do) translate into detectable differences at the protein level.

An increase in steady-state snRNA levels correlates with rescue of alternative splicing

If the alternative splicing changes, observed in both Phax and Smn mutants, are truly caused by the decreased availability of snRNPs, then rescue of steady-state snRNA levels should mirror the rescue of these alternative splicing changes in the Phax and Smn transgenic lines. Northern blotting was used to determine whether the expression of Phax and Smn transgenes increased steady-state snRNA levels in the mutants (Fig. S6A&B). Expression of the Phax transgene by the Armadillo driver significantly rescued U1 and U4 snRNA levels in the Phax$^{SH/SH}$ mutant background (Fig S6A&B). All of the snRNAs tested here were significantly rescued by expression the wild-type Smn transgene in
the Smn null background (Fig. S6C&D). For the wild-type Smn rescue line, these observations differ from our previously published results [9], wherein we detected a modest increase in snRNA levels in the rescue line. This difference likely reflects minor improvements to our northern blotting protocol, specifically: using less total RNA per gel lane and more labeled probe in the hybridization reaction to ensure probe excess. The increases in steady-state snRNA levels that we observed in the Phax and Smn rescue lines correlated well with the phenotypic rescue at both the organismal (viability) and molecular (alternative splicing) levels.

**Splicing changes shared between Phax and Smn mutants are also caused by knockdown of splicing factors Prp6 and Prp8**

Although the correlation of snRNA levels with particular splicing events provides evidence that these events are acutely sensitive to disruption of the core spliceosomal machinery, it does not rule out other possibilities. For example, the covariance could be due to a direct interaction of splicing factors with Phax and SMN, or a result of indirect factors, such as altered developmental progression. To test whether the observed splicing changes are indeed linked to a reduction in snRNPs and a corresponding disruption in spliceosome function, we depleted the tri-snRNP and U5-associated proteins Prp6 and Prp8 using RNA interference. As shown in Fig. S7, SMN depletion caused a slight (<two-fold) perturbation in the splicing events tested, but the Prp6 and Prp8 knockdowns disrupted the splicing of these transcripts to a much greater extent (Fig. S7A-C). Knockdown of Prp8 disrupted the splicing of the events identified above (Abi, αCOP, dUTPase, and Rpn2 intron 1 (int. 1)), but it did not significantly disrupt the splicing of a downstream intron in Rpn2, which serves here as a control for broader spliceosomal dysfunction. The Prp6 and Prp8 knockdowns also altered dUTPase protein expression patterns to a greater extent than did knockdown of Smn (Fig. S7B). Steady-state snRNA
levels were not significantly affected by any of these knockdowns, with the possible exception of U5 snRNA, which was slightly reduced (~40%) in the Prp8 depletion (Fig. S7D&E). The lack of snRNA decreases in the SMN depleted samples is consistent with the well-known perdurance of snRNPs in cultured cells, and it likely mirrors the observed low-amplitude disruptions to splicing. Prp6 and Prp8 are core spliceosomal proteins, and their depletion is expected to severely compromise the spliceosome [44,45]. Thus, the splicing changes seen in these knockdowns are perhaps not unexpected, but they nevertheless support the conclusion that the splicing changes identified in the Phax and Smn null mutants are snRNP dependent.

Hypomorphic Smn missense mutants rescue snRNP levels and do not display defects in alternative splicing

As mentioned, the snRNP biogenesis mutants described above are severe loss-of-function alleles. SMA is, however, a hypomorphic condition, as complete loss of SMN activity is lethal in all organisms tested. To better model the disease in the fly, we previously engineered an allelic series of transgenic animals that express Smn missense mutations known to cause SMA in humans [9,46]. In addition to the wild-type rescue transgene ($Smn^{WT}$), we analyzed three hypomorphic lines that display intermediate (SMA type II) phenotypes: $Smn^{T205I}$, $Smn^{Y107C}$, and $Smn^{V72G}$ (44). These lines cover a range of phenotypic outcomes: $Smn^{T205I}$ is semi-viable (~30% eclose as adults), $Smn^{Y107C}$ is pharate-lethal (very few animals eclose), and $Smn^{V72G}$ mutants all die as pupae. Importantly, in this system the wild-type and mutant transgenes are integrated at the identical chromosomal locus and are expressed from the native promoter in an otherwise Smn null background [9,46]. Using qRT-PCR, we analyzed the alternative splicing patterns of the four transcripts described above. As shown in Fig. 6A, none of the Smn missense lines exhibited a greater than two-fold change (in either direction) relative
to Smn$^{WT}$. Consistent with this finding, the missense mutants also displayed little to no difference in steady state snRNA levels, compared to the wild-type rescue line (Figs. 6B&C). To ensure that differences in developmental timing did not confound the analysis, animals were harvested after they began to display the wandering behavior that is characteristic of the late third instar stage of Drosophila development. We also measured snRNA levels of animals at slightly earlier and later developmental time points, and found little difference between the mutant and the control lines (Fig. S8).

**Smn missense mutants display limited overlap in splicing pattern changes**

The large-amplitude changes identified in the severe loss-of-function mutants were not evident in the Smn hypomorphs, however these lines could exhibit other splicing pattern or gene expression changes that might account for differences in their phenotypic outcomes. To identify RNA changes in the hypomorphic lines, we sequenced total (rRNA-subtracted) RNA from whole animals isolated ~12hr after puparium formation, a developmental stage that Smn null animals never reach. The hypomorphs displayed numerous changes in their splicing patterns relative to the wild-type rescue line (1662 combined; Tables S13-16). However, the missense mutants displayed few large amplitude (greater than two-fold) changes (107/1662), and hardly any of these events were shared between them (42/1662; Fig. 7A and Table S13). Of the forty-two overlapping splicing changes, only six were previously identified as being snRNP-dependent (four) or Smn gene-specific (two) by comparison to data from the severe Phax and Smn loss-of-function lines (Fig. 7B). Browser shots of representative examples are shown in Fig 7C. We used qRT-PCR to validate one of the two Smn gene-specific changes. The scarface transcript (Fig. S9 and Table S13) exhibited an increase in intron retention between the third and fourth exons in the Smn$^{V72G}$ transgenic line (deltaPsi = -0.61 and a >100-fold
change by qRT-PCR). As measured RNA-seq, the $Smn^{Y107C}$ (deltaPsi = −0.23) and $Smn^{T205I}$ (deltaPsi = −0.07) mutants exhibited smaller changes; these events were below the limit of detection by qRT-PCR (Fig. S9A and Table S13). Consistent with the notion that this is not a snRNP-dependent change, an increase in scarface intron retention was not observed following knockdown of Prp8 (Fig. S9B). Thus, RNA-seq profiling of hypomorphic, SMA-causing missense mutations is in agreement with the qRT-PCR data described above, and together these findings demonstrate that the splicing changes identified in the $Smn$ null animals are not conserved among the intermediate SMA models.

**Differential expression analysis reveals an activation of stress signaling in $Smn$ missense mutant animals.**

In addition to changes in splicing patterns, our genome-wide analysis of RNA from the severe loss-of-function mutants revealed potential snRNP-dependent and $Smn$ gene-specific changes in steady-state mRNA levels. Differential expression analysis of the $Smn$ missense mutant lines revealed numerous changes in mRNA abundance in the most severe line, $Smn^{V72G}$ (599), but far fewer in the other less severe lines, $Smn^{Y107C}$ (88) and $Smn^{T205I}$ (59) (Fig. 8A&B and Table S17-S21). Despite the low number of changes in the least severe $Smn^{T205I}$ line, more than half of these overlapped with the other hypomorphic mutants (Fig. 8B). Surprisingly, many of the stress-responsive transcripts identified in the severe $Smn$ null mutants were also among the mRNAs that were upregulated in all three $Smn$ transgenic lines, relative to the wild-type rescue line (Fig. 8C and Table S18). As corresponding increases in mRNA levels were not found in the Phax mutants (Fig 2B), these events were classified as $Smn$ gene-specific. Thus, unlike the snRNP-dependent splicing changes (which are not conserved in the $Smn$ hypomorphs), the intermediate models of SMA exhibited clear RNA
signatures of stress that correlate with disease severity (Fig. 8C) and are independent of decreases in snRNP supply (Fig. 2B).

**Discussion**

Connecting SMA pathology to SMN’s function in snRNP assembly and downstream expression of specific mRNA isoforms remains a necessary benchmark for defining how splicing changes might contribute to disease etiology. Therefore, a first step toward uncovering the molecular mechanisms that elicit SMA outcomes would be to separate snRNP-dependent phenotypes from those that are SMN specific. Transcriptomic profiling of Smn and Phax mutants enabled us to identify a robust set of snRNP-dependent alternative splicing changes in SMA model flies. We show that reductions in steady-state snRNA levels correlate with the appearance of these alternative-splicing events, and that rescue of snRNA levels in transgenic animals correlates with the restoration of normal splicing. Furthermore, depletion of core splicing factors (Prp8 or Prp6) essentially bypasses the snRNP depletion step, causing these same alternative-splicing events, and providing a more direct link to spliceosomal dysfunction. Together, these observations support a close connection between snRNP levels and proper gene expression *in vivo*. Whether or not disruptions to this ubiquitously required process contribute to SMA phenotypes is less clear.

**Ars2, 5' exons, and U1 tele scripting**

Changes in splicing and mRNA length in Phax and Smn animals were largely distinct from the more abundant changes in the Ars2 mutants (Figs. 3D and 4D). Ars2 is a component of the 5' Cap Binding Complex (CBC), and this association may underlie the transcriptomic changes that we identified in
this mutant [47,48]. The Ars2 splicing changes were mostly in 5'-proximal introns, and this finding is consistent with CBC mutants in yeast, *Xenopus*, and plants [49-55]. The association of Ars2 with the CBC likely also underlies the numerous mRNA shortening events we observed. These findings are consistent with previous studies in yeast and human cells that suggest the CBC protects a subset of weak poly(A)-signals from premature cleavage and termination [26,39,40]. Importantly, both types of Ars2 dependent mRNA processing events were largely distinct from the snRNP-dependent changes observed in the *Phax* and *Smn* mutants.

The distinct mRNA processing changes in Ars2 mutants may also reflect species-specific differences in the role of the CBC and Ars2. Unlike *ARS2* knockdown experiments in mammalian cells [26,56], we did not observe defects in snRNA or histone mRNA processing. This finding is consistent with the observation that Ars2 protein is dispensable for proper histone mRNA maturation in *Drosophila* cells [57], and it suggests that aspects of Ars2 function are not conserved between vertebrates and invertebrates.

Our identification of a large number of mRNA shortening events that are shared between *Phax* and *Smn* mutants supports a recently discovered role for the U1 snRNP in the protection of pre-mRNAs from premature cleavage and polyadenylation (PCPA). PCPA was previously observed by subtractive hybridization PCR, coupled with high throughput sequencing of RNA from cells treated with antisense oligomers targeting the 5'-end of U1 snRNA [35,36]. This protective role for U1 snRNPs was termed ‘telescripting’ [35,36] and this study represents the first confirmation of the process outside of cell culture models. Although we observed an overall trend toward shorter transcripts in our datasets (Fig. 3), we found only a few large-magnitude (greater than two-fold) changes (Table S2). This is perhaps unsurprising, given that treatment of cultured cells with relatively high concentrations of anti-U1 oligomers provides an acute and nearly complete inhibition of U1
snRNA function, whereas the mutations in Phax and Smn do not perturb U1 to the same extent in the mutant animals.

**RNA signatures of disease**

Alternative splicing depends on the regulated accessibility of pre-mRNAs to the spliceosome. Studies in a variety of eukaryotes have shown that changing this accessibility, either directly or indirectly, alters the splicing patterns of specific pre-mRNAs [17-21]. The extent to which snRNP levels contribute to normal splicing regulation and/or disease pathology is still being worked out, however, changes in the pattern of alternative-splicing have been reported in various SMA model systems [4-6,10]. The splicing changes reported in these SMA models are typically assumed to stem directly from loss of SMN and snRNPs, but such changes are often difficult to distinguish from broad, indirect effects of disease pathology or developmental delay [7,10]. By identifying overlapping splicing changes in Phax and Smn mutants, we are now able to distinguish between snRNP-dependent changes and gene-specific effects.

Many of the alternative splicing changes shared by Phax and Smn occur in genes of known importance to neuromuscular biology. For example, the mammalian ortholog of the alphaCOP protein has a demonstrated interaction with SMN, and it co-localizes with SMN in neuronal projections [58,59]. This interaction was recently shown to be required for neurite outgrowth in mammalian cell and zebrafish models of SMA [60]. Orthologs of the Abi protein appear to have important roles in neurogenesis via the regulation of actin dynamics [61-63]. Dominant mutations in GARS cause the distal neuropathy Charcot–Marie–Tooth disease type 2D, and recent observations suggest that GARS may colocalize with SMN in human cells [64-66]. Mouse models of SMA exhibit
disruptions to ubiquitin homeostasis, and evidence of SMN interactions with proteasomal components, including Psmd1, the mouse ortholog of Drosophila Rpn2 [67,68].

Despite our finding of splicing disruptions in genes of known importance to SMA and neuromuscular biology in severe loss-of-function mutants, these splicing changes were absent from less severe models of SMA. Splicing patterns are not conserved from flies to humans, and we would not expect to find similar splicing disruptions in mammalian models of SMA. Finding these genes at the top of our dataset may merely reflect the sensitive nature of the splicing of genes that are important for neuromuscular biology, and the fact that transcriptome diversity is greatest in neuronal tissues, including the brain [69].

Although we found evidence for both snRNP-dependent and SMN-specific RNA changes in SMA models, it remains to be determined which of these, if any, contributes to disease pathology. Our analysis of severe and intermediate SMA models suggests that snRNP-independent, Smn gene-specific RNA changes may play a bigger role in SMA pathology than previously envisioned. The SMN-specific activation of stress signaling pathways is conserved in SMA models [70,71] and correlates well with disease severity in Smn hypomorphs (Fig. 8C). This activation of stress signaling appears to be a common signature of neuromuscular disease [72].

Based on our finding of activated stress signaling in fly models of SMA, an important question moving forward will be to understand how loss of SMN causes this stress response. By text mining with the stress signatures found in SMA models described here, we found that mutations in the Activating transcription factor 3 (Atf3) cause a similar dysregulation in immune and metabolic homeostasis [73]. Among other things, this signaling pathway appears to regulate the cytosolic formation of SMN-containing U-bodies in human cells in response to bacterial pathogens [74]. If SMN does indeed intersect with a normal host cell response to bacteria, this would provide a parsimonious
model for the activation of stress signaling in SMA models, and another avenue to explore potential connections to neuromuscular disease.

In conclusion, a direct comparison of snRNP biogenesis mutants revealed that defects in snRNP supply cause complex changes to the transcriptome. However, an analysis of three different Smn hypomorphic lines showed that RNA processing changes are unlikely to be the primary drivers of SMA pathophysiology. Our studies support a model of SMA etiology wherein general disruptions at the mRNA level and specific disruptions at the SMN protein-protein level are responsible for SMA phenotypes. In future experiments, the key will be to uncover the tissue specific molecular activities of SMN that are important for neuromuscular function. Ongoing work with these and other intermediate models of SMA should help clarify the relative contributions of the aforementioned factors to SMA pathology and etiology.
Materials and methods

Fly mutants and husbandry

Stocks were cultured on molasses and agar at room temperature (25±1°C) in half-pint bottles. Oregon-R was used as the wild-type allele. The SmnX7 microdeletion allele was a gift from S. Artavanis-Tsakonis (Harvard University, Cambridge, USA) [75]. Transposon insertion alleles SmnD (f01109), Ars2C (c00735), and Ars2E (e01128) were obtained from the Exelixis collection at Harvard Medical School [42,76]. The transposon insertion allele PhaxSH (P{lacW}PhaxSH064f) was a gift from S. Hou [77].

The wild-type and mutant Smn transgenes are expressed using the native Smn promoter. Transgenic constructs were integrated into the PhiC31 landing site located at 86F8 [9,46]. An intron-less Phax cDNA was sub-cloned into pBID-UASC-GV [78], and subsequently injected and integrated into the 51C1 PhiC31 landing site by BestGene Inc. (Chino Hills, CA). The UAS:Phax-mVenus transgene at 51C1 was then recombined into the PhaxSH background. The armadillo promoter-GAL4 driver P{GAL4-arm.S}11 [79] was also recombined into the PhaxSH mutant background.

RNA-seq & bioinformatics analyses

RNA was isolated from 74 ± 2-hour-old wild-type and snRNP biogenesis mutant larvae by homogenization in TRIzol® (Invitrogen™), according to the manufacturer’s protocol. Staged animals were isolated from either the wandering third instar stage of development or just before bubble migration in early tan pre-pupae (<12 hours after puparium formation). RNA was DNased with Amplification Grade DNase I (Invitrogen™) and TURBO™ DNase (Ambion™). A TruSeq RNA Sample Preparation Kit v2 (Illumina®) was used for: poly-(A)-enrichment or substituted here with Ribo-Zero™ (Epicentre®) rRNA subtraction, barcoding for multiplexing, and cDNA library preparation. Paired end (2x50) sequencing was performed on an Illumina® HiSeq 2000 platform. TopHat and Cufflinks were
used for gene expression analyses, according to the bioinformatic pipeline from Trapnell et al. (2012). DaPars was used to identify differences in RNA length and alternative polyadenylation [37,38]. MISO [41] was used to measure changes in annotated alternative-splicing events. For comparison, we used a ‘strong’ Bayes factor cutoff for our poly(A)-selected RNA-seq analysis [80]. We used a ‘positive’ Bayes factor as our cutoff for the total rRNA(-) RNA-seq analysis [81]. For developmental comparison, all publicly available RNA-seq data from L2 to L3 puff stage 7–9 were downloaded from the NCBI Sequence Read Archive (SRA) and converted to fastq format with the SRA toolkit [82] (http://www.modencode.org/). The heatmap was generated with the ‘pheatmap’ package from bioconductor (http://www.bioconductor.org).

**Northern blotting**

Northern analyses were performed using standard protocols [83]. Briefly, RNA was isolated as indicated above and approximately 300 ng of total RNA was separated on 10% Novex® TBE Urea gels (Invitrogen™), transferred to GeneScreen Plus® charged nylon membranes (PerkinElmer®), dried, and UV cross-linked. Northern blots were probed with 5’-end-labeled oligonucleotides for the respective snRNAs, labeled with T4 polynucleotide kinase (NEB®) and [γ-32P]ATP (Perkin-Elmer®). The snRNA-specific oligonucleotide probe sequences are listed in Table S22. Hybridizations were performed at 65°C, and sequential washes in 2x SSC and 0.33x SSC were performed at 60°C. Damp blots were exposed to storage phosphor screens (GE Healthcare) and analyzed with a Typhoon TRIO+ (GE Healthcare). Bands were quantified using ImageQuant TL (GE Healthcare), and p-values for three biological replicates were determined with a one-tailed Student’s t-test for unequal variance of two-samples.
RT-PCR analysis

Semi-quantitative RT-PCR was performed with intron flanking oligos, using the 2X Apex™ Taq Master Mix (Genesee Scientific). Quantitative RT-PCR experiments were carried out as in Praveen et al. (2012). Briefly, random hexamers were used to prime the reverse transcription of isolated RNA according to the manufacturers protocol SupersScript® III (Invitrogen™). Real-time PCR reactions of cDNA were carried out on a 7900HT Fast Real-time PCR machine (Applied Biosystems™), according to the Maxima® SYBR Green/Rox qPCR Master Mix (2X) (Thermo Scientific™) two-step protocol. Oligo pairs amplified cDNAs from intron inclusive versus spliced mature mRNAs. Three biological replicates were tested for each genotype. The ΔΔCt method was used to quantify differences, and p-values were determined with a one-tailed Student’s t-test for unequal variance of two-samples. Gene-specific primer sequences are listed in Table S22.

Western blotting

Western analyses were performed using standard protocols and modifications to the methods of Praveen et al. (2012). Briefly, larval protein lysates were prepared by crushing the animals in lysis buffer (50mM Tris-HCl [pH 7.5], 150 mM NaCl, 1mM EDTA, 1% NP-40, 0.1% SDS, 1% Deoxycholate) with 5X protease inhibitor cocktail (Invitrogen™) and clearing the lysate by centrifugation. The anti-dUTPase antibody was a generous gift from B.G. Vértessy, and it was used at a dilution of 1 to 50,000 [84]. Affinity purified anti-dSMN was used at a 1 to 2,500 dilution [9]. Anti-alpha-tubulin antibody (Sigma®) was used at a 1 to 50,000 dilution.

Cell culture and RNAi

The S2 cell line was obtained from the Drosophila Genome Resource Center (Bloomington, IN) and cultivated according to published protocols [85]. Gene-specific RNAi knockdown was also performed as described in Rogers and Rogers (2008). Briefly, S2 cells were maintained in Sf-900™ II SFM
media (Gibco™) supplemented with 1X Pen Strep (Gibco™). For RNAi, one milliliter of (1x10^6 cells/ml) was aliquoted into each well of a 6-well tissue culture plate. Double-stranded RNA (dsRNA) was in vitro transcribed overnight (~16 hours) from PCR templates with the T7 MEGAscript kit (Invitrogen™), and dsRNA products was subsequently treated with TURBO™ DNase (Ambion™), according to the respective manufacturers protocols. Oligonucleotide sequences for amplifying the PCR templates are listed in Table S22. PCR templates were amplified from the following plasmids: Smn, pAFW-SMN; Prp6/CG6841, cDNA clone OFa18909 (GenScript); or Prp8, cDNA clone OFa09689 (GenScript). Purified dsRNA was phenol-chloroform extracted, ethanol precipitated, resuspended in RNase-free water (Ambion™), and quantified by spectrophotometer. Approximately 15 µg of dsRNA was added to 1x10^6 cells per each well of a 6-well plate. The dsRNA was added once each day for three successive days. On the fourth day, cells were pelleted and resuspended in TRIzol® (Invitrogen™). RNA and proteins were extracted according to the manufacturer’s protocol.

Data deposition

The data from the Smn null animals were deposited in NCBI's Gene Expression Omnibus [86] and are accessible through GEO Series accession number GSE49587 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE49587). The deposition of the additional data discussed here in GEO is pending.
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References


44. Grainger RJ, Beggs JD. Prp8 protein: At the heart of the spliceosome. RNA. 2005 May 1;11(5):533–57.


52. Fresco LD, Buratowski S. Conditional mutants of the yeast mRNA capping enzyme show that the cap enhances, but is not required for, mRNA splicing. RNA. 1996 Jun;2(6):584–96.


Figure Legends

Figure 1. Analysis of three different snRNP biogenesis mutants. (A) Browser shots of larval RNA-seq reads at the three disrupted gene loci. Reads from Oregon-R (WT) versus Phax<sup>SH/SH</sup>, Smn<sup>X7/D</sup>, or Ars2<sup>C/E</sup> snRNP biogenesis mutants are shown. Mapped read tracks were normalized to the median of the middle two quartiles of the mapped WT sequence read counts. (B) Northern blot of RNA from snRNP biogenesis mutants and WT larvae. RNA was extracted from WT larvae at 60 ± 2 hours (60h) post egg-laying, and mutant RNA was extracted at 74 ± 2 hours to account for their delayed development (see text). (C) Quantification of northern blot in (B). Sm-class snRNAs were normalized to the Lsm-class U6 snRNA, and U6-normalized WT snRNA levels were set at 100. Asterisks are p-values from a Student’s t-test: * p-value ≤ 0.05, ** p-value ≤ 0.01, and *** p-value ≤ 0.001.

Figure 2. Gene expression differences in snRNP biogenesis mutants. (A) Heatmap comparison of Cuffdiff FPKM levels of differentially expressed transcripts. Heatmap colors were rescaled for each row and the rows were clustered based on pattern of gene expression between WT and mutants. (B) A heatmap of FPKMs from a set of stress-responsive transcripts in (A). (C) Venn diagram of the overlap in gene expression differences in the snRNP biogenesis mutants. Phax = Phax<sup>SH/SH</sup>, Smn = Smn<sup>X7/D</sup>, and Ars2 = Ars2<sup>C/E</sup>. Numbers in parentheses are totals.

Figure 3. mRNA length changes in snRNP biogenesis mutants. Pairwise comparison of significant (False discovery rate adjusted p-value < 0.05) differences in mRNA length between: (A) WT and Phax<sup>SH/SH</sup>, (B) WT and Smn<sup>X7/D</sup>, and (C) WT and Ars2<sup>C/E</sup>. RNA length differences were extracted from mapped RNA-seq read files by the DaPars linear regression algorithm (see text). Percentage of Distal PolyA site Usage Index = PDUI. (D) Venn diagram of RNA length differences in snRNP biogenesis mutants labeled as in Fig. 2.
Figure 4. Alternative-splicing differences among snRNP biogenesis mutants. Distribution of significant alternative-splicing changes from small |ΔPsi| near 0 to large |ΔPsi| near 1: (A) Phax<sup>SH/SH</sup> mutants relative to WT; (B) Smn<sup>X7/D</sup> mutants relative to WT; and (C) Ars2<sup>C/E</sup> mutants relative to WT. (D) Venn diagram of overlapping alternative-splicing changes in snRNP biogenesis mutants, labeled as in Fig. 2.

Figure 5. Transgenic rescue of alternative-splicing changes in Phax and Smn mutants. (A) UCSC browser shots of mapped read tracks for four of the top fifteen identified alternative splicing changes. (B) qRT-PCR analysis of intron retention in Phax<sup>SH/SH</sup> mutants versus Phax<sup>SH/SH</sup> mutants expressing a UAS-Phax transgene from the armadillo promoter driven GAL4 (Arm>Phax). (C) qRT-PCR of intron retention in Smn<sup>X7/D</sup> mutants versus Smn<sup>X7/X7</sup> mutants expressing a wild-type Smn transgene from its native promoter. For (B) and (C), levels of WT intron retention were set at 1. (D) Western blot for dUTPase. Alternate protein isoforms reflect alternative-splicing change in dUTPase mRNA. Wild-type Smn transgene in Smn<sup>X7/X7</sup> background = Tg:Smn<sup>WT</sup>. Long dUTPase protein isoform = PA and short = PB.

Figure 6. Analysis of steady-state snRNA levels and alternative splicing of target genes in SMN missense mutants (A) qRT-PCR analysis of intron retention of flies expressing SMA patient-derived missense mutations. Transgenic animals of the following generalized genotype were used: Smn<sup>X7/X7</sup>,Flag-Smn<sup>Tg/–</sup>, where Tg represents a WT, V72G, Y107C, or T205I transgene. Intron retention in the Smn<sup>WT</sup> transgenic rescue line was set at 1. (B) Northern blots of snRNA levels in the missense mutant lines. (C) Quantification of snRNA levels from panel B. RNA levels in the Smn<sup>WT</sup> (WT) transgenic line were set at 100.

Figure 7. Alternative splicing changes in Smn missense mutants. (A) Venn diagram of overlapping alternative-splicing changes in Smn missense mutants. (B) Venn diagram of the overlap in shared
splicing differences in the Smn missense mutants with either snRNP-dependent changes or Smn gene-specific changes categorized by comparison of the Phax and Smn total rRNA(-) RNA-seq. (C) Browser shot examples of an Smn gene-specific change in alternative-splicing in scarface (top) or a snRNP-dependent change in CG43394 within a PAPLA1 intron on the opposite strand (bottom). Genotypes are labeled by Smn

Figure 8. Differential gene expression in Smn hypomorphic animals. (A) Heatmap comparison of Cuffdiff FPKM levels of differentially expressed transcripts. Heatmap colors were rescaled for each row and the rows were clustered based on pattern of gene expression between Smn

and the Smn missense mutants. (B) Venn diagram of overlapping differences in mRNA levels. (C) A heatmap of FPKMs from a set of stress-responsive transcripts. Genotypes are labeled by Smn

which was expressed in the Smn
null background.
Supplemental Figure S1. Timing of the developmental delays exhibited by the snRNP biogenesis mutants. (A) Pairwise comparisons of Cuffdiff log_{10} FPKM levels from age-matched Oregon-R and snRNP biogenesis mutants relative to modENCODE developmentally-staged controls for second instar (L2), early 3rd instar (L3 12h), and a later “puff” stage of 3rd instar (L3 P1-2) wild-type larvae. (B) Dendrogram comparison of overall gene expression between age-matched Oregon-R, snRNP biogenesis mutants, and modENCODE developmentally staged animals. L3 P3-6 and L3 P7-9 are subsequent 3rd instar stages relative to L3 P1-2.

Supplemental Figure S2. snRNA levels in snRNP biogenesis mutants quantified by developmental northern (A) and ribo-minus total RNA-seq data (B-C). Note that the control flies in panels B and C are age-matched at 72 hrs, and thus RNA-seq reads are artificially high in the controls. (B) snRNA levels from aggregated mapped read counts, normalized to U6 snRNA mapped read counts. (C) snRNA levels from aggregated mapped read counts that were normalized to the median of the middle two quartiles of all mapped WT sequence read counts. Normalized age-matched Oregon-R (WT) snRNA levels were set at 100.

Supplemental Figure S3. Relative abundance of snRNA isoforms in snRNP biogenesis mutants: (A) U1 snRNA paralogs; (B) U2 snRNA paralogs; (C) U4 snRNA paralogs; and (D) U5 snRNA paralogs. Identical isoforms are listed together.

Supplemental Figure S4. Normal developmental changes in mRNA length. Pairwise comparison of significant differences in mRNA length between modENCODE developmentally staged animals: (A) L2 and L3 12h; (B) L3 12h and L3 P1-2; and (C) L2 and L3 P1-2.

Supplemental Figure S5. Semi-quantitative RT-PCR of alternative-splicing changes identified by RNA-seq. MISO alternative-splicing event annotations are listed.
Supplemental Figure S6. Analysis of snRNA levels in transgenic rescue lines. (A) Northern blot of snRNA in the Phax mutant and Phax rescue line (Arm>Phax) relative to WT 60h larvae; quantified in panel (B). (C) Northern blot of snRNA levels in the Smn mutant and Smn rescue line (Tg:Smn<sup>WT</sup>) relative to WT 60h; quantified in panel (D). U6-normalized snRNA levels of WT were set at 100.

Supplemental Figure S7. snRNP-specific protein knockdown in S2 cells. (A) qRT-PCR of untreated cells versus those treated with dsRNA for Smn, Prp6 or Prp8 mRNAs. (B) Western blot of dUTPase levels in RNAi-treated cells. Anti-SMN and anti-alpha-tubulin verify SMN knockdown and load, respectively. (C) qRT-PCR verification of Prp6 and Prp8 mRNA knockdowns. Levels in untreated cells were set at 100. (D) Northern blot of snRNA levels from S2 cell knockdowns, quantified in (E). U6-normalized snRNA levels of WT were set at 100.

Supplemental Figure S8. Lack of snRNA level decreases in Smn missense mutants over development. (A&C) Northern blots of snRNA levels in the SMN missense mutants over development, quantified in (B&D). snRNA levels from the Smn<sup>WT</sup> (WT) transgenic lines were set at 100. Pupal animals were staged by color and internal air bubble, and larval animals were from a timed collection (72-77 hours post egg laying). The small differences in the early larval stage likely reflect differences in developmental timing, as the Smn missense mutants exhibit slower developmental progression.

Supplemental Figure S9. Intron retention in scarface transcript. (A) qRT-PCR analysis of intron retention of flies expressing SMA patient-derived missense mutations. Transgenic animals of the following generalized genotype were used: Smn<sup>X7/X7</sup>,Flag-Smn<sup>Tg/-</sup>, where Tg represents a WT, V72G, Y107C, or T205I transgene. Intron retention in the Smn<sup>WT</sup> transgenic rescue line was set at 1. (B) qRT-PCR of untreated cells versus cells treated with dsRNA for Smn or Prp8 mRNAs.
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Relative intron retention (Tg:Smn<sup>WT</sup> = 1)

A

**Figure 6**

B

C

Relative snRNA levels

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Figure 7
Figure 8