

Srrm234, but not canonical SR and hnRNP proteins drive inclusion of *Dscam* exon 9 variable exons

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ABSTRACT

Q2 Alternative splicing of pre-mRNA is a major mechanism to diversify protein functionality in metazoans from a limited number of genes. In the *Drosophila melanogaster* Down syndrome cell adhesion molecule (*Dscam*), which is important for neuronal wiring, up to 38,016 isoforms can be generated by mutually exclusive alternative splicing in four clusters of variable **Q3** exons. However, it is not understood how a specific exon is chosen from the many variables and how variable exons are prevented from being spliced together. A main role in the regulation of *Dscam* alternative splicing has been attributed to RNA binding proteins (RBPs), but how they impact on exon selection is not well understood. Serine–arginine-rich (SR) proteins and hnRNP proteins are the two main types of RBPs with major roles in exon definition and splice site selection. Here, we analyzed the role of SR and hnRNP proteins in *Dscam* exon 9 alternative splicing in mutant *Drosophila melanogaster* embryos because of their essential function for development. Strikingly, loss or overexpression of canonical SR and hnRNP proteins even when multiple proteins are depleted together, does not affect *Dscam* alternative exon selection very dramatically. Conversely, noncanonical SR protein Serine–arginine repetitive matrix 2/3/4 (*Srrm234*) is a main determinant of exon inclusion in the *Dscam* exon 9 cluster. Since long-range base-pairings are absent in the exon 9 cluster, our data argue for a small complement of regulatory factors as main determinants of exon inclusion in the *Dscam* exon 9 cluster.

Keywords: alternative splicing; *Dscam*; RNA binding proteins; SR proteins; SRm300; hnRNP proteins

INTRODUCTION

During alternative splicing, the combination of exons can be varied to generate multiple different transcripts and proteins from one gene (Soller 2006; Nilsen and Graveley 2010; Fiszbein and Kornblihtt 2017). In humans, 95% of genes, and in *Drosophila melanogaster* 63% of genes are alternatively spliced, respectively (Wang et al. 2008; Fu and Ares 2014). Among the genes where alternative

splicing generates the greatest diversity of isoforms is the *Drosophila melanogaster* homolog of human Down syndrome cell adhesion molecule (*Dscam*), which encodes a cell surface protein of the immunoglobulin superfamily. The *Dscam* gene comprises 95 alternatively spliced exons that are organized into four clusters, namely 4, 6, 9, and 17, which contain 12, 48, 33, and two variables, respectively. Hence, the *Dscam* gene can generate up to 38,016 different proteins (Schmucker et al. 2000; Neves et al. 2004; Hemani and Soller 2012; Sun et al. 2013). *Dscam* is

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functionally required for neuronal wiring in the nervous system, but also for phagocytosis of invading pathogens in the immune system (Schmucker et al. 2000; Watson et al. 2005). Interestingly, *Dscam* in mosquitos changes its splicing pattern upon pathogen exposure to produce isoforms with higher binding affinity for binding pathogen (Dong et al. 2006). However, despite intense research, relatively little is known about how *Dscam* alternative splicing is regulated in flies.

Pre-mRNA splicing is a multistep process catalyzed by the spliceosome sequentially assembled from five U snRNPs together with numerous proteins. Spliceosome assembly initiates by the recognition of the 5' splice site by U1 snRNP and of the 3' splice site by U2 snRNP, together with U2AFs recognizing the branchpoint and the polypyrimidine tract and the AG of the 3' splice site, respectively. Then the U4/5/6 tri-snRNP is recruited, and upon several structural rearrangements where U4 snRNP leaves the spliceosome, catalysis takes place by two transesterification reactions (Lührmann and Stark 2009).

Alternative splicing is to a large degree regulated at the level of splice site recognition involving base-pairing of U1 snRNP to the 5' splice site YAG/GURAGU and U2 to the branchpoint WNCUAAU (W: A or U, *Drosophila melanogaster* consensus [Lim and Burge 2001]) whereby splice sites closer to the consensus are preferably used (Soller 2006). Splice site selection is critically assisted by RNA binding proteins (RBPs) that support or inhibit recognition of splice sites.

Serine-arginine rich (SR) and heterogenous nuclear ribonucleoproteins (hnRNPs) are two prominent classes of RBPs involved in alternative splicing regulation (Busch and Hertel 2012; Fu and Ares 2014; Bradley et al. 2015). Humans have twelve and flies have eight SR proteins, each having one or two RNA recognition motifs (RRM) and RS domain rich in serines and arginines (Busch and Hertel 2012). In addition, RS domains are present in some other splicing factors lacking RNA binding domains such as Tra2, SRRM1 (SRm160), and SRRM2 (SRm300), SRRM3, and SRRM4 (nSR100), which are termed noncanonical SR proteins (Blencowe et al. 1999; Long and Caceres 2009; Busch and Hertel 2012; Best et al. 2014). In contrast, hnRNP proteins are more diverse in their modular assembly containing RNA binding domains (e.g., RRM, KH, or RGG domains) and various auxiliary domains (Geuens et al. 2016). In humans, the most prominent hnRNPs are the abundantly expressed hnRNP A and C family (Busch and Hertel 2012; Geuens et al. 2016).

SR proteins mostly bind to exonic splicing enhancers (ESEs) and recruit spliceosomal components to splice sites through their RS domains. SR protein binding sites are present in both alternatively spliced and constitutive exons to promote exon inclusion, but they can also repress inclusion of alternative exons (Black 2003; Shen et al. 2004; Wang et al. 2006; Chen and Manley 2009; Pandit et al. 2013).

Although SR proteins recognize similar sequences, distinct functions have been shown either by binding distinct sites or when bound to the same site through differential regulation mediated by combinatorial interactions with other splicing regulators (Gabut et al. 2007; Änkö et al. 2012; Pandit et al. 2013; Bradley et al. 2015).

In contrast to SR proteins, hnRNP proteins mostly bind to intronic splicing silencers (ISSs) and repress inclusion of alternative exons (House and Lynch 2006; Wang et al. 2008). In addition, they can act antagonistically to SR proteins by binding to exonic splicing silencers (ESSs) and compete with SR proteins for binding (Soller 2006; Long and Caceres 2009). However, a more comprehensive analysis revealed that SR and hnRNP proteins can also act coordinately in many instances in exon inclusion or repression (Brooks et al. 2015).

With regard to the alternative splicing in the *Dscam* gene, a model has been proposed for the exon 6 cluster involving long-range base-pairing. Here, a conserved docking sequence in the first intron of the exon 6 clusters can base-pair with complementary selector sequences in front of each variable exon to bring a chosen variable exon into the proximity of the proximal constant flanking exon for splicing (Graveley 2005). This model also requires that the entire cluster is maintained in a repressed state and variable exons are selected under the control of RBPs. However, the architecture in the exon 4 and 9 clusters is different and conserved "docking site" sequences are found at the end of the exon 4 and 9 clusters, but the support for this model based on evolutionary sequence conservation is weak (Yang et al. 2011; Haussmann et al. 2019).

RNAi knockdown of RBPs in cell culture in *Drosophila melanogaster* S2 cells revealed little changes for most variable exons in the *Dscam* exon 4 cluster, but this result could be due to residual protein that is left (Park et al. 2004). Hence, we wanted to investigate the role of SR and hnRNP proteins in *Drosophila melanogaster Dscam* alternative splicing more comprehensively at an organismal level using knockout mutants and overexpression. Most SR and hnRNP proteins are essential for development to adult flies, but embryonic development proceeds such that *Dscam* alternative splicing could be analyzed in late-stage embryos in SR and hnRNP mutants or when overexpressed. Unexpectedly, we find that inclusion of *Dscam* exon 9 variables is affected little in loss or gain of function (GOF) conditions. Likewise, even upon removal even of multiple factors, *Dscam* exon 9 alternative splicing changes little. However, the noncanonical SR protein Serine-arginine repetitive matrix 2/3/4 (Srrm234) is required for inclusion of most exon 9 variables. We further find that long-range base-pairing is not supported as a general model. Hence, our results argue that a small complement of RBPs are the main regulators of *Dscam* exon 9 alternative splicing.

RESULTS

Analysis of *Dscam* exon 9 alternative splicing by restriction digests

All 33 exons in the *Drosophila melanogaster Dscam* exon 9 variable exon cluster have about the same length and run as a single band on an agarose gel (Fig. 1A,B). The sequences in variable exons, however, differ enough such that a complement of restriction enzymes can digest the complex mix of PCR products to identify a majority of isoforms on sequencing type gels using one ³²P labeled primer after reverse transcription of the mRNA (Fig. 1C,D; Haussmann et al. 2019). Using a combination of *Sac*II, *Cl*al, *Psh*AI, *Hae*III, *Mse*I, *Bsr*I, and *Bst*NI yields 26 fragments of unique size identifying 20 variable exons (Fig 1D).

Alteration of SR proteins has little impact on *Dscam* exon 9 alternative splicing

SR proteins are organized into five families and representative orthologues are present in *Drosophila melanogaster* (Fig. 2; Busch and Hertel 2012). To determine the role of SR proteins in *Drosophila melanogaster Dscam* exon 9 alternative splicing we obtained mutants and overexpression lines for most of the canonical SR proteins as well as for general splicing factor SF1 and noncanonical SR protein *Srrm1* (SRm160) and *Srrm234* (SRm300, CG7971) (Fig. 2A; Supplemental Fig S1). For loss of function (LOF) alleles of SR genes, five from the ten analyzed, which are *X16^{GS1678}*, *SF2^{GS22325}*, *B52²⁹*, *Srrm1^{B103}*, and *Srrm234^{AN}*, are required to reach adulthood (Fig. 2A,B). Gain of function conditions by pan-neural *elavGAL4* mediated over-expression via *UAS* was lethal in larval instars for *UAS GFP-X16*, *UAS RSF1*, *UAS GFP-SC35*, *UAS GFP-SF2*, and *UAS GFP-B52*, while over-expression of *UAS SF1* and *UAS Srrm1* from *EP* lines did not result in a phenotype (Fig. 2A,C).

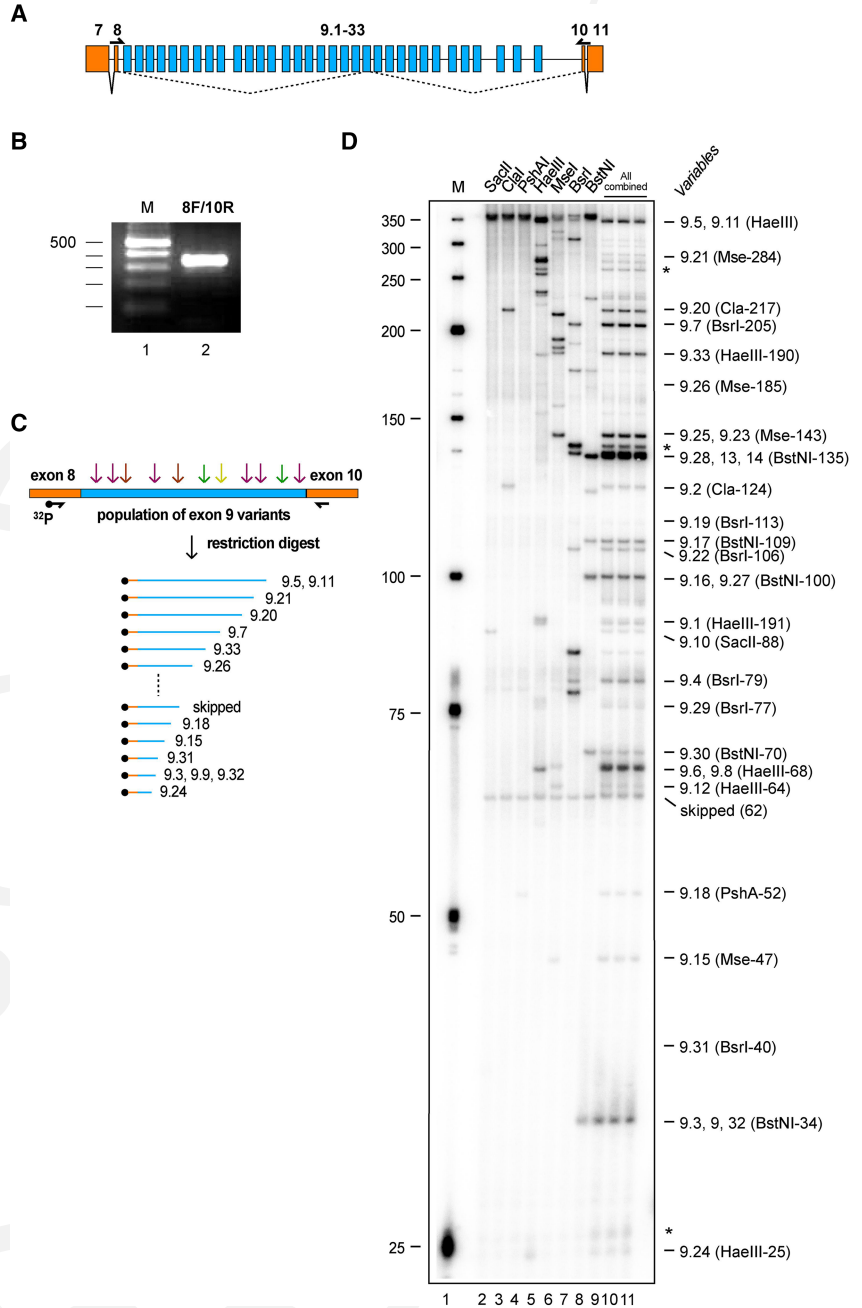


FIGURE 1. Analysis of *Dscam* exon 9 alternative splicing by restriction digestion of PCR products. (A) Schematic of the *Dscam* exon 9 variable cluster gene region. Constitutive exons are shown in orange and variable exons in blue. Primers to amplify the variable part are shown on top of the exons. (B) RT-PCR product for variable exon cluster 9 shown on a 3% agarose gel. (C) Schematic of the method used to resolve inclusion levels of variable exons using a ³²P labeled forward primer in combination with a set of restriction enzymes followed by separation of a denaturing polyacrylamide gel. (D) Denaturing acrylamide gel (6%) showing a restriction digest (*Sac*II, *Cl*al, *Psh*AI, *Hae*III, *Mse*I, *Bsr*I, *Bst*NI) of *Dscam* exon 9 variables amplified with a ³²P labeled forward primer from 14–18 h *Drosophila* embryos. Single enzyme reference digests are shown on the left (lanes 2–8) and the combination of all enzymes on the right (lanes 9–11).

Next, we analyzed inclusion levels of exon 9 variables in embryos for LOF and GOF conditions of canonical SR proteins and SF1. For LOF alleles *X16^{GS1678}*, *RBP1^{HP37044}*,

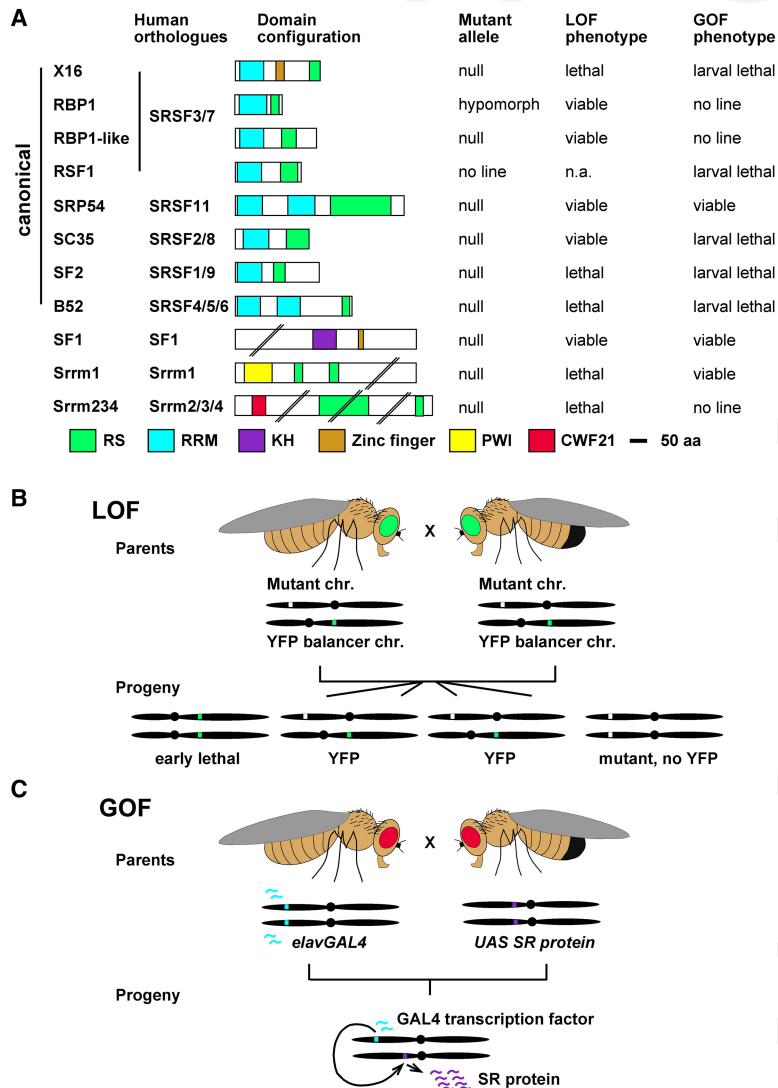


FIGURE 2. Protein domain structure of *Drosophila* SR proteins and phenotype of LOF and GOF mutants. (A) Evolutionary relationship of *Drosophila* SR proteins with human homologs is shown on the left and the domain structure is indicated by colored boxes. Arginine-serine-rich domain (RS, green), RNA recognition domain (RRM, light blue), hnRNP K homology domain (KH, purple), zinc finger domain (ochre), Proline-Tryptophan-Isoleucine domain (PWI, yellow) and CWF21 domain (red). The type of allele obtained, and the loss (LOF) and GOF phenotypes are indicated on the right. (B) Viability was determined from stocks that harbor a zygotically expressing GFP marked balancer chromosome, which contains a set of inversions to suppress recombination, and a recessive lethal mutation. If a gene is essential, only heterozygous flies will survive. Homozygous mutant embryos were identified in the progeny of these stocks by the lack of GFP and advanced development as homozygous balancer carrying embryos die early before GFP expression. (C) To obtain embryos overexpressing SR proteins, flies carrying the yeast GAL4 transcription factor under the control of the pan-neuronal *elav* promoter were crossed with lines harboring SR proteins under the control of the yeast *UAS* promoter.

RBP1-like^{NP0295}, *Srp54*^{GS15334}, *SC35*^{KG02986}, *SF2*^{GS22325}, *B52*²⁸, and *SF1*^{G14313} the *Dscam* exon 9 splicing pattern unexpectedly remained largely unchanged and significant changes are prominent in exon 9.4, 9.21, and 9.5/9.11 for mutants compared to wild-type (marked in red for decrease and blue for increase, Fig. 3A,B). Similar results

were obtained for over-expression of *UAS GFP-X16*, *UAS RSF1*, *UAS GFP-SC35*, *UAS GFP-SF2*, *UAS GFP-B52*, and *UAS-Srrm1* with significant changes only prominent in exon 9.19 and 9.21 for GOF conditions compared to wild-type (marked in red for decrease and blue for increase, Fig. 3C,D).

Noncanonical SR protein *Srrm234* is required for the selection of *Dscam* exon 9 variables

The canonical SR proteins contain an RRM and bind to RNA. In contrast, the large *Srrm1* and *Srrm234* proteins contain RS domains, but seem not to bind RNA and exert splicing enhancing functions through association with proteins bound to ESEs (Fig. 2; Blencowe et al. 1998, 2000; Eldridge et al. 1999; Szymczyna et al. 2003).

We obtained null mutants for both genes, *Srrm1*^{B103} and *Srrm234*^{ΔN}, which are late embryonic lethal (Fan et al. 2014). While loss of *Srrm1* had little effect on *Dscam* exon splicing, loss of *Srrm234* resulted in significant reduction in inclusion for many variable exons (marked in red, 9.4, 9.7, 9.10, 9.16/9.27, 9.17, 9.18, 9.20, 9.23/9.25, 9.29, and 9.33) that is compensated by increased inclusion of a few variable exons (marked in blue, 9.3/9.32, 9.6/9.8, 9.12, and 9.30) (Fig. 4A,B). Consistent with a role in *Dscam* exon 9 alternative splicing regulation, *Srrm234* is also expressed in the nervous system of *Drosophila melanogaster* embryos in the same pattern as *Dscam* (Supplemental Fig. S2).

Alteration of hnRNP proteins has little impact on *Dscam* exon 9 alternative splicing

Since alterations of individual canonical SR proteins had little impact on selection of *Dscam* exon 9 variables, we focused on hnRNP proteins as candidates for repressing inclusion of exon 9 variables (Olson et al. 2007; Chen and Manley 2009; Fu and Ares 2014). *Drosophila melanogaster* has four members of the highly expressed hnRNP A family (*Hrp36*, *Hrp38*, *Rb97D*, and *Hrp48*) and one member of the

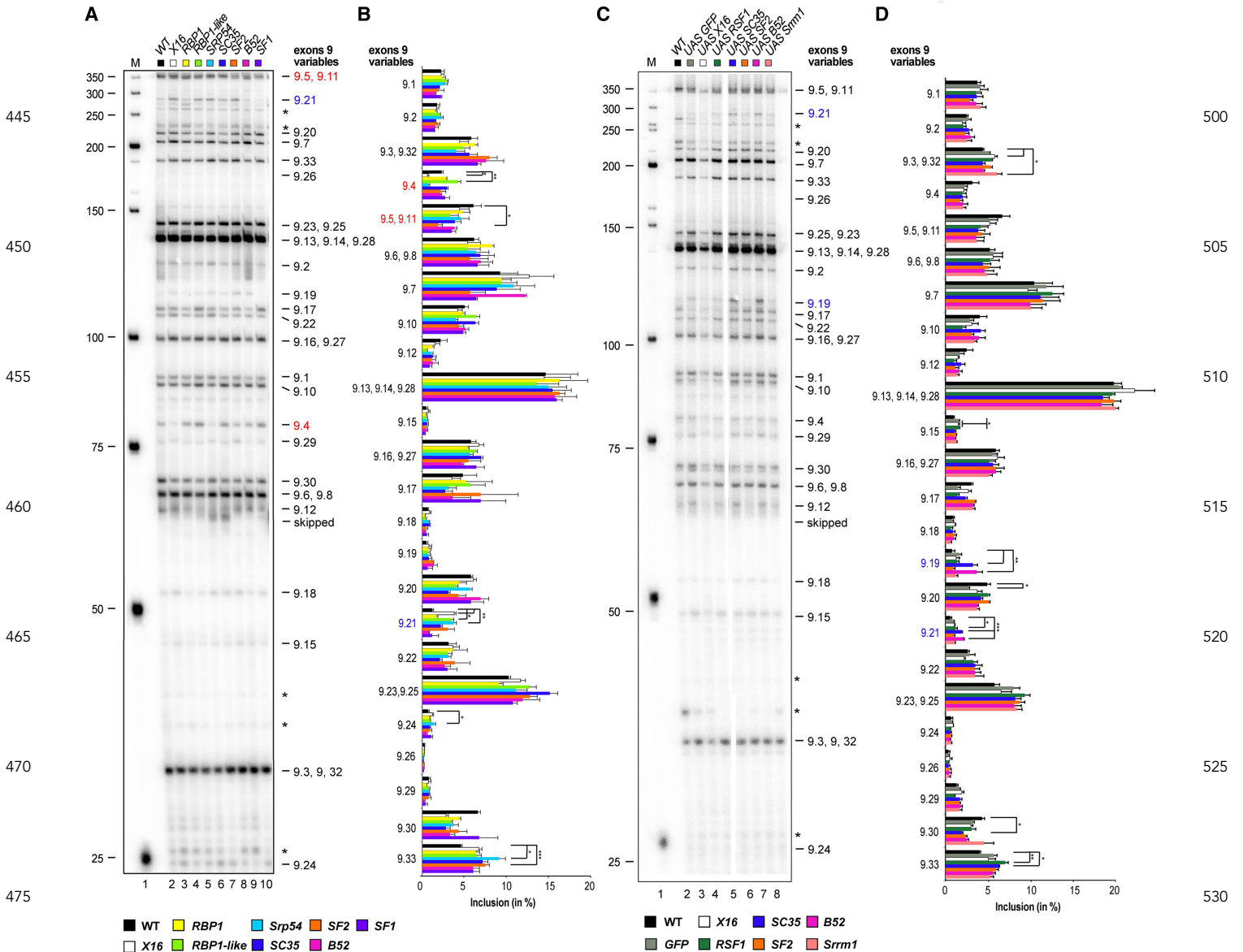


FIGURE 3. Analysis of *Dscam* exon 9 alternative splicing in canonical SR protein LOF and GOF mutants. (A,C) Denaturing acrylamide gel showing restriction digests of *Dscam* exon 9 variables amplified with a ³²P labeled forward primer from 14–18 h *Drosophila* embryos for canonical SR protein LOF (A) and *elavGAL4* UAS GOF mutants (C). Quantification of inclusion levels are shown as means with standard error from three experiments for canonical SR protein LOF (B) and GOF mutants (D). Prominent changes in inclusion levels in mutants compared to wild-type are marked with red letters for a decrease and in blue for an increase, and statistically significant differences are indicated by asterisks. (***) $P \leq 0.001$, (**) $P \leq 0.01$, (*) $P \leq 0.05$.

485 hnRNP C, although the *Drosophila melanogaster* ortho-
F5 logue is considerably longer (Fig. 5; Appocher et al. 2017). Other highly expressed hnRNP proteins are Hrp40 (hnRNP D), Glorund (hnRNP F/H), Hrb57A (hnRNP K), and Hrb59 (hnRNP M).

490 To determine the role of hnRNP proteins in *Drosophila*
melanogaster *Dscam* exon 9 alternative splicing we
could obtain null-mutants for all major hnRNP proteins
(Hrp36^{BG02743}, Hrp38^{MI1059}, RB97D¹, Hrp48^{GS14498},
Hrp40^{GS18188}, glo^{t02674}, Hrb57^{G13574}, and Hrp59^{GS6029})
495 and gene-switch or EP over-expression lines for

most (Hrp36^{GS15926}, Hrp38^{GS12795}, Hrp48^{EY12571}, and
Hrp59^{GS6029}, Figs. 5, 7A; Supplemental Fig. S3). Half of
the tested major hnRNPs are required for viability
(Rb97D, Hrp40, glo, and Hrp59), while only over-expres-
sion of Hrp36 was lethal (Fig. 5).

Then, we analyzed *Dscam* exon 9 inclusion levels for
LOF and GOF of hnRNPs. For LOF alleles, the *Dscam*
exon 9 splicing pattern also unexpectedly remained large-
ly unchanged and significant changes are prominent in
exon 9.17, 9.19, and 9.21 for mutants compared to wild-
type (marked in red for decrease and blue for increase,

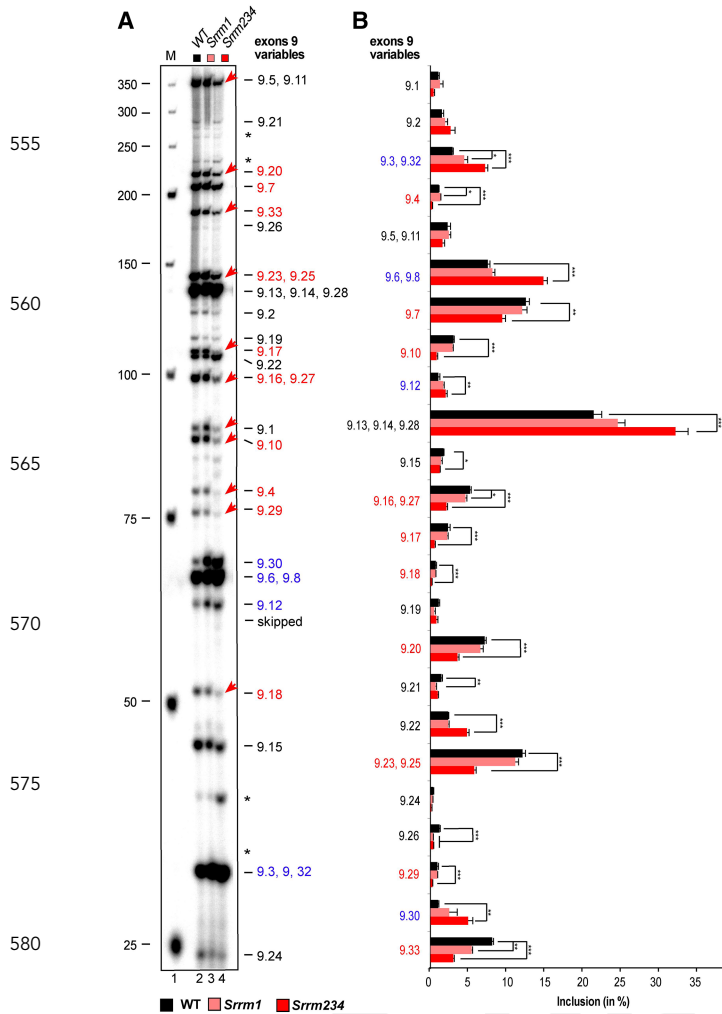


FIGURE 4. Analysis of *Dscam* exon 9 alternative splicing in noncanonical SR proteins *Srm1* and *Srm234*. (A) Denaturing acrylamide gel showing restriction digests of *Dscam* exon 9 variables amplified with a ³²P labeled forward primer from 14–18 h *Drosophila* embryos for *Srm1* and *Srm234* protein LOF mutants. Quantification of inclusion levels is shown as means with standard error from three experiments (B). Red arrows point toward exons with reduced inclusion levels in the *Srm234*^{ΔN} mutant compared to wild-type. Prominent changes in inclusion levels are marked with red letters for a decrease and in blue for an increase in the *Srm234*^{ΔN} mutant compared to wild-type. Statistically significant differences are indicated by asterisks. (***) $P \leq 0.001$, (**) $P \leq 0.01$, (*) $P \leq 0.05$.

Fig. 6A,B). Similar results were obtained for GOF conditions with significant changes only prominent in exon 9.2, 9.4, 9.17, 9.19, 9.21, and 9.30 (marked in red for decrease and blue for increase, Fig. 6C,D).

Binding of Hrp36, Hrp38, Hrp48, and Hrp40 to RNA has been analyzed globally in *Drosophila melanogaster* and binding motifs have been established by SELEX (Blanchette et al. 2009). When we reanalyzed these data we did find significantly increased binding for all four Hrp proteins ($P < 0.05$) to the *Dscam* exon 9 variable cluster compared to averaged binding, but the binding curves did

not overlap with changes in exon inclusion in LOF or GOF conditions (Supplemental Fig. S4). Likewise, we did not find an overlap of SELEX motif enrichments with changes in exon inclusion in LOF or GOF conditions (Supplemental Fig. S4).

Removal of multiple SR and hnRNP proteins has little impact on selection of *Dscam* exon 9 variables

hnRNP36 and SR protein B52 genes lie next to each other in the *Drosophila melanogaster* a genome and potentially could cross-regulate to compensate for each other (Fig. 7A). Therefore, we generated a double knockout of *hnRNP36* and *B52* genes (*hnRNP36/B52*^{Δ1}). In addition, we also combined this double knockout with the *hnRNP38* mutant as *hnRNP36* and *hnRNP38* are closely related and since they are highly expressed, they could act redundantly.

Surprisingly, even in *hnRNP36/B52*^{Δ1} *hnRNP38*^{d05172} triple mutants, *Dscam* exon 9 was robustly spliced with only differences in inclusion levels of variables 9.4, 9.16/9.27, 9.17, 9.18, 9.19, and 9.21 compared to controls (marked in red for decrease and blue for increase, Figure 7B,C).

Variable exon selection is not explained by long-range base-pairing in *Dscam* exon 9

In the exon 4 cluster, we could not detect conserved sequences adjacent to every variable exon that could mediate long-range base-pairing arguing that such a mechanism is not involved in variable exon 4 selection (Haussmann et al. 2019). A sequence alignment between *Drosophila melanogaster* and *Drosophila virilis* showed strong conservation in the coding sequences and the architecture of the exon 9 cluster with only few insertions and deletions of exons (Supplemental Fig. S5). There are conserved sequence elements in the intron before constant exon 10 that potentially could serve as a docking site (Yang et al. 2011), but we did not find conserved sequence elements between every variable exon by manual inspection (Supplemental Fig. S6). A systematic bioinformatics comparison of this potential docking site with arbitrary sampled sequences of the same length and sequence complexity further did not reveal a special propensity of the docking site to form more or stronger complementary alignments within *Dscam* intronic sequences in the variable exon 9 cluster. The same picture arises when comparing predicted energies of the best secondary duplex structures formed by the reverse intronic sequence, the docking site, and sampled sequences (Supplemental Fig. S7). In addition, we did not find other instances of the docking site in any other gene of *D. melanogaster* (except in the anti-sense RNA *CR45129* of *Dscam*), indicating that the sequence is not recurrent in other genes.

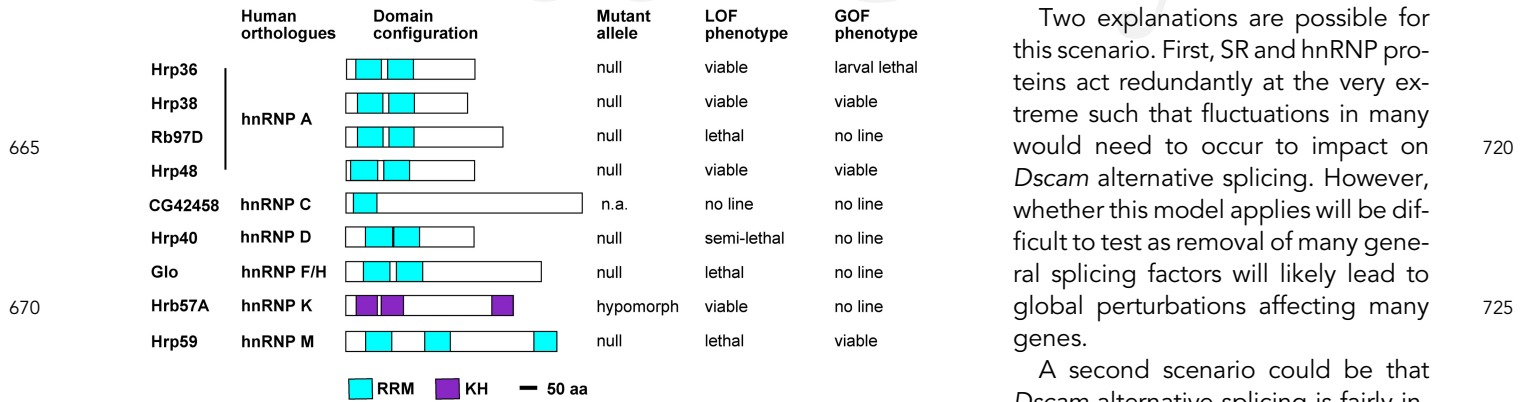


FIGURE 5. Protein domain structure of *Drosophila* hnRNP proteins and phenotype of LOF and GOF mutants. Evolutionary relationship of *Drosophila* hnRNP proteins with human homologs is shown on the left and the domain structure is indicated by colored boxes. RNA recognition domain (RRM, light blue) and hnRNP K homology domain (KH, purple). The type of allele obtained, and the loss (LOF) and gain of function (GOF) phenotypes are indicated on the right.

DISCUSSION

Although the sequence determinants that direct the spliceosome to its correct position are very degenerate, splicing needs to occur precisely and with high accuracy to prevent disease (Cooper et al. 2009; Zaharieva et al. 2012). It is therefore thought that RBPs play key roles in localizing functional splice sites. In particular, the abundant SR and hnRNP proteins have been attributed key roles in this process by forming RNA–protein complexes cotranscriptionally to recruit early splicosomal components for defining splice sites. Although SR proteins were initially viewed as binding ESEs to activate splicing, and hnRNP proteins to bind ISSs for antagonizing SR proteins, a number of genome-wide studies draw a more complex picture for both SR and hnRNP binding and function (Blanchette et al. 2009; Änkö et al. 2010, 2012; Huelga et al. 2012; Pandit et al. 2013; Brooks et al. 2015). In fact, both SR and hnRNP proteins can have very specific functions in one context, but also redundant functions in another context.

In this regard, we hypothesized that an array of similar exons as found in the *Drosophila melanogaster* *Dscam* gene would provide a platform for SR and hnRNP proteins to evolve exon-specific functions to regulate their inclusion. Surprisingly, however, *Dscam* exon 9 alternative splicing is exactly the opposite and the splicing pattern is very robustly maintained when SR and hnRNP proteins were either removed or overexpressed. In particular, the advances of *Drosophila melanogaster* genetics allowed us to use complete knockouts of most canonical SR and general hnRNP proteins and thus avoid the ambiguity of RNAi that would leave residual protein. Hence, despite complete loss of individual SR and hnRNP proteins, or combinations thereof, the *Dscam* splicing pattern is robustly maintained.

Two explanations are possible for this scenario. First, SR and hnRNP proteins act redundantly at the very extreme such that fluctuations in many would need to occur to impact on *Dscam* alternative splicing. However, whether this model applies will be difficult to test as removal of many general splicing factors will likely lead to global perturbations affecting many genes.

A second scenario could be that *Dscam* alternative splicing is fairly independent of general splicing factors. This would imply a more specific mechanism. Initially, it has been thought that long-range base-pairing would provide such a mechanism as conserved sequences have been

found in the *Dscam* exon 6 cluster (Graveley 2005). Our previous analysis of the exon 4 cluster, and the in-depth analysis of the exon 9 cluster in this paper, however, rule out such mechanism in these two clusters (Hausmann et al. 2019). Hence, the question remains whether two independent mechanisms arose to regulate mutually exclusive alternative splicing in *Dscam* variable clusters. Potentially, the conserved sequences present in the variable clusters could provide binding sites for RBPs that have adopted cluster specific roles. In this context, it is interesting to note that deletion of the docking site in exon 6 leads to the inclusion of mostly the first exon in the cluster (May et al. 2011). This is unexpected as removal of the splicing activating mechanism should result in skipping of the entire variable cluster, because the proposed repressor hnRNP36 would still be present. Accordingly, the docking site in the exon 6 cluster also exerts a repressive role in maintaining the entire cluster in a repressed state.

Likewise, our finding that noncanonical *Srrm234* regulates the inclusion of many variables in the *Dscam* exon 9 cluster suggests a mechanism in *Dscam* mutually exclusive alternative splicing that differs from more general splicing rules directed by canonical SR and general hnRNP proteins. One of the human homologs of *Srrm234*, *SRRM4* has key roles in the regulation of microexons (Irimia et al. 2014). Due to their small size, microexons cannot be defined through the standard mechanism of exon definition. Hence, a distinct mechanism must apply, that can accurately direct splicing of microexons. Because microexons are often found in large introns, a robust process must underlie their selection and involves a newly described enhancer of microexons domain (eMIC), present in human *SRRM3* and *SRRM4*. Intriguingly, in vertebrates, the ancestral prohomolog *Srrm234* has duplicated into three genes to adopt distinct functions through dedicated protein domains, but these features are maintained by alternative

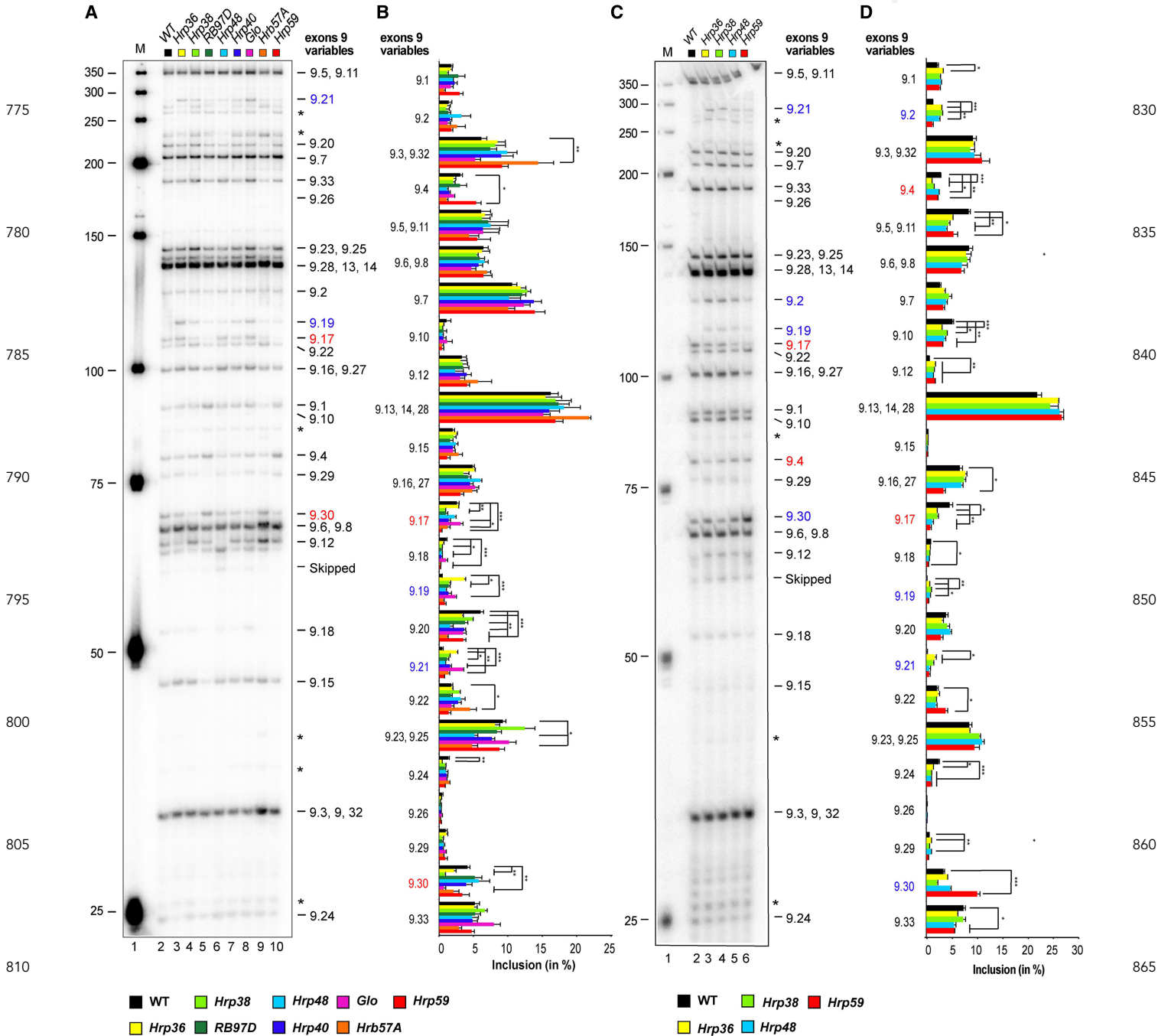
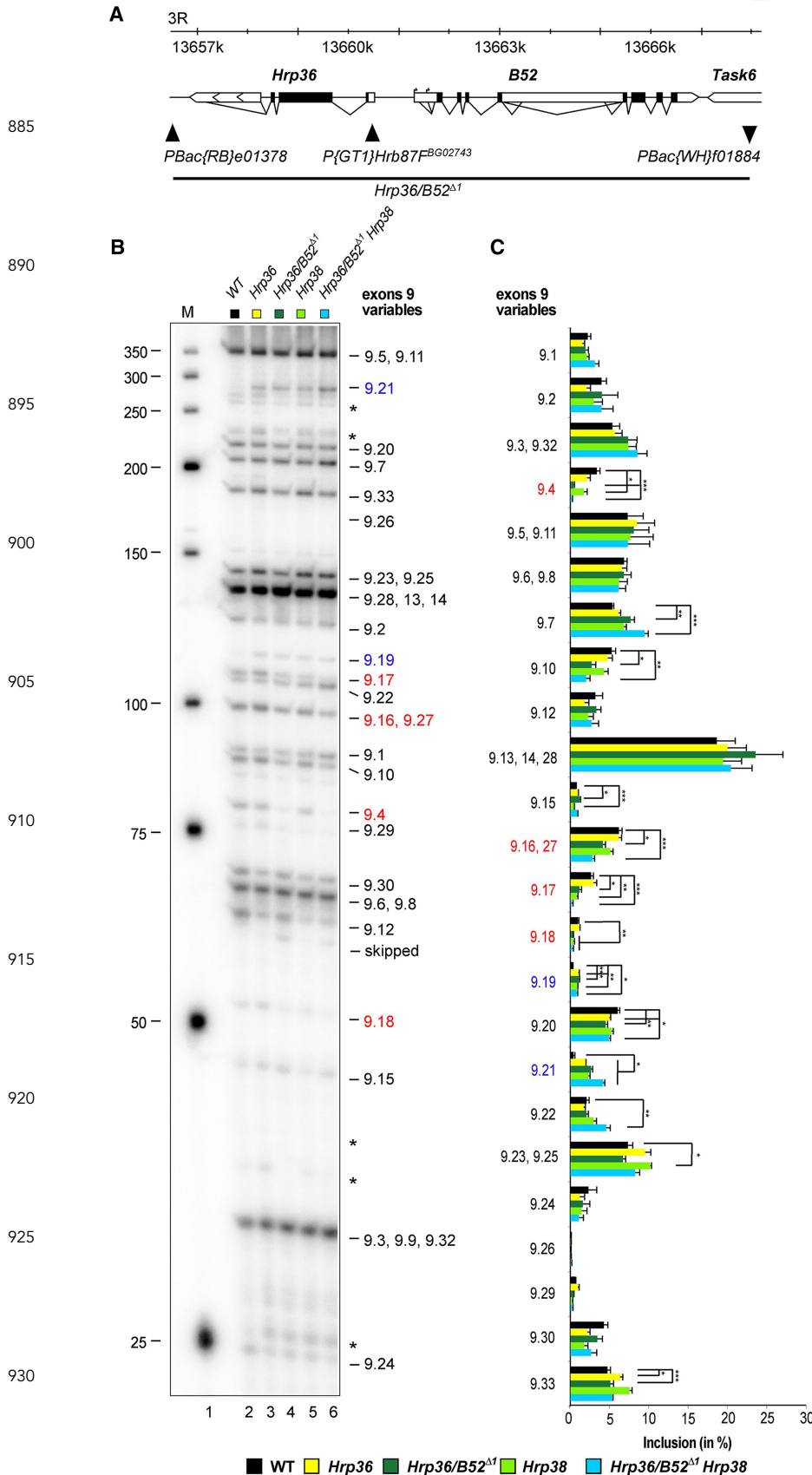


FIGURE 6. Analysis of *Dscam* exon 9 alternative splicing in general hnRNP protein LOF and GOF mutants. (A,C) Denaturing acrylamide gel showing restriction digests of *Dscam* exon 9 variables amplified with a ³²P labeled forward primer from 14–18 h *Drosophila* embryos for general hnRNP protein LOF (A) and *elavGAL4* UAS GOF mutants (C). Quantification of inclusion levels are shown as means with standard error from three experiments for canonical SR protein LOF (B) and GOF mutants (D). Prominent changes in inclusion levels are marked with red letters for a decrease and in blue for an increase, and statistically significant differences are indicated by asterisks. (***) $P \leq 0.001$, (**) $P \leq 0.01$, (*) $P \leq 0.05$.

mRNA processing in *Drosophila melanogaster* *Srrm234* to include the eMIC at the carboxyl terminus of the protein in neuronal tissue (Torres-Méndez et al. 2019).

Dscam alternative exons comply with the general average length of exons and thus the mechanism of their regulation is likely distinct from microexons. *Dscam* exon 9

cluster regulation by *Srrm234* seems to involve its Cwf21 domain, which is not required for microexon inclusion (Torres-Méndez et al. 2019). Transposon inserts in the middle of the *Srrm234* gene resulting in a truncated protein, that contains the Cwf21 domain do not affect exon 9 diversity (*Mi{ET1}CG7971^{MB07314}* and *Mi{Liang et al. 2003}*)



CG7971^{M104068}, data not shown). The Cwf21 domain, which is homologous to the yeast Cwc21 domain, has been attributed key roles in splicing due to copurification of the human ortholog SRRM2 with active spliceosomes and its localization in the catalytic center of the spliceosome (Bessonov et al. 2008). Furthermore, the Cwc21 domain interacts with the U5 snRNP core components Snu114 and Prp8 involved in key structural rearrangements in the spliceosome during catalysis (Grainger et al. 2009; Gautam et al. 2015). Interestingly, Cwc21 has been attributed roles in splicing of meiotic genes which are regulated differently of general intron-containing genes (Gautam et al. 2015). *Srrm2* forms a complex with *Srrm1* to promote alternative splicing of *Drosophila melanogaster* *doublesex* required for sex determination, but the mechanism in *Dscam* is different as loss of *Srrm1* does not impact on alternative splicing in the exon 9 cluster (Blencowe et al. 1998; Eldridge et al. 1999). In addition, *Srrm234* also interacts with U1 70K, so potentially could act to activate 5' splice site and initiate the splicing process from a repressed state of the variable cluster (Guruharsha et al. 2011).

A *Dscam* cluster-specific role has been suggested for *Hrp36* acting as a repressor preventing splicing together of exon 6 variables in S2 cells, but whether the results are the same in flies remains to be tested (Olson et al. 2007). Such a factor has not been identified for exon 4 and 9 clusters, and for the factors tested, we did not observe splicing together of variable exons. Intriguingly, a newly annotated exon 4.0 in the beginning of the bee *Dscam* exon 4 cluster is spliced to exon 4.6 (Decio et al. 2019). This unexpected finding, however, is not compatible with the model described for *Hrp36* in the exon 6 cluster, but might involve a repressive sequence element around exon 4.0 similar the element discovered in the beginning of the exon 6 cluster, which

FIGURE 7. (Legend on next page)

when deleted results only in inclusion of exon 6.1 (May et al. 2011). Likewise, factors like *Srrm234* would then act as activators to drive inclusion of variables.

Taken together, *Dscam* exon 9 mutually exclusive alternative splicing is robust against fluctuations of in canonical SR and general hnRNP proteins arguing for a specific mechanism regulating inclusion levels of variable exons. Indeed, noncanonical SR protein *Srrm234* plays a key role in increasing inclusion of many exon 9 variables. However, since *Srrm234* does not have one of the classic RNA binding domains, additional RBPs likely connect *Srrm234* to *Dscam* exon 9. Hence, our data obtained from knockouts of general splicing factors indicate that a small complement of RBPs are likely key regulators of *Dscam* mutually exclusive alternative splicing.

The gene structure of invertebrate *Dscam* harboring arrays of variable exons for mutually exclusive splicing is at the extreme, but arrays of several alternative exons are common to many genes in metazoans. Likely, a yet to be discovered feature of the spliceosome has been exploited in mutually exclusive alternative splicing of *Dscam* such that only one exon is chosen. Likewise, since sequences that look like splice site are common in large introns, such a mechanism could be broadly relevant for robust select of isolated exons.

MATERIALS AND METHODS

Fly genetics

Flies were maintained on standard cornmeal agar food as previously described (Hausmann et al. 2013). *CantonS* was used as a wild-type control. The following LOF mutants were used as depicted in Supplemental Figures S1, S2, and Figure 7A: *X16* (*P{GSV6}Hrb27C^{GS16784}*, DGRC Kyoto #206763), *RBP1* (*P{EPg}mRpL37^{HP37044}*, BDSC #22011), *RBP1-like* (*P{GawB}Rbp1-like^{NP0295}*, DGRC Kyoto #103580), *Srp54* (*P{GSV6}Srp54^{GS15334}*, DGRC Kyoto #206174), *SC35* (*P{SUPor-P}SC35^{KG02986}*, BDSC #12904), *SF2* (*P{GSV7}SF2^{GS22325}*, DGRC Kyoto #203903), *B52²⁸* (Gabut et al. 2007), *SF1* (*P{EP}SF1^{G14313}*, BDSC #30203), *Hrp36* (*P{GT1}Hrb87F^{BG02743}*, BDSC #12869), *Hrp38* (Mi(Liang et al. 2003)*Hrb98DE^{MI10594}*, BDSC #55509), *Rb97D* (*P{PZ}Rb97D¹*, BDSC #11782), *Hrp48* (*P{GSV6}Hrb27C^{GS14498}*, DGRC Kyoto #205836), *Hrp40* (*P{GSV6}sqd^{GS18188}*, DGRC Kyoto #201020), *Glo* (*PBac{WH}glo^{f02674}*, BDSC #18576), *Hrb57A*

(*P{EP}HnRNP-K^{G13574}*, BDSC #29672), *Hrp59* (*P{GSV3}rump^{GS6029}*, DGRC Kyoto #200852), *Srrm1^{B103}* (*SRm160^{B103}*, (Fan et al. 2014), and *Srrm2340^{AN}*. Since the transposon used for mutagenesis are large (~10 kb), inserts in the transcribed part were considered to be null alleles, while inserts in promoter regions were considered hypomorphic alleles. If inserted in an intron transposons disrupt splicing, if inserted in the 5'UTR they will prevent translation of the ORF, or if inserted in the ORF lead to a truncated nonfunctional protein. If inserted in the promoter region, transposon inserts reduce transcription. Whether lethality of mutants mapped to the locus was tested by crossing to chromosomal deficiencies.

The null-allele *Srrm234^{AN}* (CG7971) was generated by GenetiVision CRISPR gene targeting services. The 3.2 kb deletion at the amino terminus of the gene was generated using sgRNAs AGTCTGCTGGGGACTGCT and CGCCGCAGGACATATAA CAG together with donor template harboring two homology arms flanking a *loxP* 3xP3-GFP *loxP* cassette. Left and right homology arms of the donor were amplified using primers CG7971-LAF1 (GTTCCGGTCTCTTAGCCCTGCAGCAGCTTCT GCTTG) and CG7971-LAR1 (TCCAAGGTCTCACAGTTTATAT GTCCTGCGGCGTGC), and CG7971-RAF2 (GTTCCGGTCTCT GTCAGCTGGGAGCCGGCAGTGC) and CG7971-RAR2 (TCC AAGGTCTCAATCGAGTGGAGAACCCATACGTACTTAGATCC), respectively. Successful deletion and integration of the cassette was validated by PCR and Sanger sequencing using primers CG7971-outF1 (CATCGATTGTGTGATGAAGTTTAC) and CG7971-outR2 (GGGGAGTATCTGTGAGCAGTTGTATC), and LA-cassette-R (AAGTCGCCATGTTGGATCGACT), and Cassette-RA-F (CCTGGGCATGGATGAGCTGT), respectively. For the analysis of *Dscam* alternative splicing the 3xP3-GFP marker was removed by Cre mediated recombination using an insert on a third chromosome balancer (*TM6B, P{w⁺mC}=Crew}D^{H2}, Tb*, BDSC #1501) and the resulting chromosome was rebalanced with a zygotically YFP-expressing balancer (*TM6B, P{Dfd-EYFP}3 Sb, Tb*, BDSC #8704) to collect the embryonic lethal homozygous mutants.

For GOF experiments the following UAS lines, gene switch vector inserts and EP lines were used: UAS GFP control line (Gabut et al. 2007), UAS GFP-X16 (Gabut et al. 2007), UAS RSF1 (Labourier et al. 1999), UAS GFP-SC35 (Gabut et al. 2007), UAS GFP-SF2 (Gabut et al. 2007), *Srrm1* (*P{EP}Srrm1^{G18603}*, BDSC #26938), UAS GFP-B52 (Gabut et al. 2007), *Hrp36* (*P{GSV6}GS15926*, DGRC Kyoto #206416), *Hrp38* (*P{GSV6}GS12795*, DGRC Kyoto #204283), *Hrp48* (*{EPgy2}Hrb27C^{EY12571}*, BDSC #20758), *Hrp59* (*P{GSV3}GS6029*, DGRC Kyoto #200852).

Hrp36 and *B52* genes lie next to each other. The *Hrp36/B52^{Δ1}* double mutant was generated by FRT/FLP mediated recombination using *PBac{RB}e01378* and *PBac{WH}f01884* transposon insertions as previously described (Zaharieva et al. 2015). The lethal *Hrp36/B52^{Δ1}* allele was balanced and validated using primers e01378 Rev (GCCACATTTAGATGATTCAGCATTAT), f01884 Rev (GATTTCCAATAGATCCCAAC CGTTTCG), and RB 3' MINUS (TCCAAG CGGCGACTGAGATG).

Lethal lines were rebalanced with balancers expressing YFP zygotically, but not maternally under a *Dfd* promoter

FIGURE 7. Analysis of *Dscam* exon 9 alternative splicing in double and triple mutants for general splicing regulators. (A) Schematic of the genomic region of *Hrp36* and *B52* genes. Transposon inserts are shown with triangles and the deletion *Hrp36/B52^{Δ1}* is indicated at the bottom. (B) Denaturing acrylamide gel showing restriction digests of *Dscam* exon 9 variables amplified with a ³²P labeled forward primer from 14–18 h *Drosophila* embryos for *Hrp36^{BG02743}*, *Hrp36/B52^{Δ1}*, *Hrp38^{d05172}*, and *Hrp36/B52^{Δ1} Hrp38^{d05172}* mutants. (C) Quantification of inclusion levels is shown as means with standard error from three experiments. Prominent changes in inclusion levels are marked with red letters for a decrease and in blue for an increase; statistically significant differences are indicated by asterisks. (***) *P* ≤ 0.001, (**) *P* ≤ 0.01, (*) *P* ≤ 0.05.

(CyO, P{Dfd-EYFP}2, BDSC #8578, TM6B, P{Dfd-EYFP}3 Sb, Tb, BDSC #8704) to allow for the selection of homozygous lethal mutants. Non-GFP expressing 14–18 h embryos were further selected according to the morphology of the auto-fluorescing gut to distinguish them from homozygous balancer carrying animals, which die before they express GFP (Haussmann et al. 2008). For overexpression, a third chromosomal *elavGAL4* insert was used (P{w[+mmC]=GAL4-elav.L}3, BDSC #8760).

RNA extraction, RT-PCR, restriction digestion, and denaturing acrylamide gels

Total RNA was extracted using Tri-reagent (SIGMA) and reverse transcription was done with Superscript II (Invitrogen) as previously described (Koushika et al. 1999) using primer *Dscam* 11RT1 (CGGAGCCTATTCCATTGATAGCCTCGCACAG, 1 pmol/20 μ L reaction). PCR to amplify *Dscam* exon 9 cluster was done using primers 8F1 (GATCTCTGGAAGTGCAAGTCATGG) and 10R1 Δ ST (GGCCTATCGGTGGCAGGTTCCATCTGGGAGGTA) for 37 cycles with 1 μ L of cDNA. Primers were labeled with 32 P γ -ATP (6000 Ci/mmol, 25 mM, PerkinElmer) with PNK (NEB) to saturation and diluted as appropriate. From a standard PCR reaction with a 32 P labeled forward primer, 10%–20% were sequentially digested with a mix of restriction enzymes (NEB) according to their buffer requirements and temperatures. PCR reaction and restriction digests were phenol/CHCl₃ extracted, ethanol precipitated in the presence of glycogen (Roche) and analyzed on standard 6% sequencing type denaturing polyacrylamide gels. After exposure to a phosphorimager (BioRad), individual bands were quantified using ImageQuant (BioRad) and inclusion levels for individual variable exons were calculated from the summed up total of all variables. Statistical analysis was done by one-way ANOVA followed by Tukey–Kramer post-hoc analysis using Graphpad prism. Percent inclusion levels of exon 9 variables of embryos were calculated from the total sum of variables. RNA in situ were obtained from flybase as previously described (Haussmann et al. 2008).

Sequence analysis

The in silico analysis for SELEX motif occurrence was done by plotting the scores from sliding window using a position weight matrix of the *Drosophila melanogaster* consensus sequence for a given RBP to the *Dscam* exon 9 variable cluster ($P < 0.05$, Blanchette et al. 2009; Korhonen et al. 2009). The CLIP data was downloaded from GEO as deposited in (Blanchette et al. 2009), and the TiMAT window-scores have been used to generate the plot based on the version 4 of the genome assembly. Tiling arrays scores within the *Dscam* variable exon 9 cluster were plotted against all scores to determine whether the four Hrp proteins show significantly increased binding.

Vista alignments were generated as previously described (Haussmann et al. 2011). The exon 9 docking site was scanned against all gene sequences (as downloaded from FlyBase on the 26th of February 2019) using Blat (Version 35, parameters: -stepSize = 1 tileSize = 6 -minScore = 0 -minIdentity = 0, filtering for hits of at least bit-score of 30 and a coverage over the docking sequence of at least 20 residues) (Kent 2002).

In order to compare the docking site's propensity to form potential long-range base-pairing with other intronic *Dscam* sequences of the same length, we sampled 100 strictly intronic sequenced of the same or higher sequence complexity as determined by Shannon entropy. We then used the BioPython Bio.pairwise2 local alignment implementation to get pairwise alignments against all (excluding the sampled sequenced and the docking sequence) reverse and reverse-complement subsequences of the same length in any *Dscam* intron, using a custom substitution matrix to allow for G-U pairings (scoring scheme: G \rightarrow U 0.8, A \rightarrow U: 1, G \rightarrow C: 1.2, gap opening penalty 0.1, gap extension penalty 0.1) (Cock et al. 2009). We then retained all hits with less than three gaps and normalized the alignment scores by each queries' potential highest score. To assess the best potential secondary structure formed by the docking sequence with any intronic sequence, we run the same sampled sequenced against the concatenated reverse and reverse complement intronic sequences of *Dscam* (masking each query) using RNA duplex from the Vienna package and obtained for each sequence the predicted lowest energy secondary structure prediction (Lorenz et al. 2011).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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Author contributions: M.S. and P.U. designed and directed the project. P.U., M.S., I.H., and H.L. performed experiments. A.T.-M. and M.I. generated and validated the *Srrn234*^{4N} allele. R.A. performed bioinformatic analysis. P.U. and M.S. wrote the manuscript. All authors read and approved the final manuscript.

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