A splicing enhancer exhibits both constitutive and regulated activities

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The Drosophila proteins Transformer (Tra) and Transformer2 (Tra2) regulate the sex-specific alternative splicing of Drosophila doublesex (dsx) female-specific pre-mRNA by specifically binding to a splicing enhancer (dsxRE) located 300 nucleotides (nt) downstream from a female-specific 3′ splice site. In this paper we show that the dsxRE can function as a Tra and Tra2-independent splicing enhancer in vitro when located within 150 nucleotides of the 3′ splice site. Based on the relative levels of SR proteins that bind stably to the dsxRE in the presence or absence of Tra and Tra2, we propose that the constitutive splicing activity of the dsxRE is mediated by its weak interactions with SR proteins and possibly other general splicing factors. In contrast, Tra and Tra2 allow the dsxRE to function at a distance from the intron by stabilizing the interactions between these proteins and the dsxRE.

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The Drosophila proteins Transformer (Tra) and Transformer2 (Tra2) mediate the positive regulation of doublesex (dsx) female-specific pre-mRNA splicing by binding to a regulatory element located in the exon down-stream of a female-specific 3′ splice site [Nagoshi and Baker 1990; Hedley and Maniatis 1991; Hoshijima et al. 1991; Ryner and Baker 1991, Tian and Maniatis 1992, 1993]. This regulatory sequence, which is characterized by six copies of a 13-nucleotide repeat sequence, is called the dsx repeat element (dsxRE). The dsxRE has been shown to be both necessary and sufficient for Tra and Tra2-dependent, female-specific splicing of dsx pre-mRNA in vivo [Hedley and Maniatis 1991; Hoshijima et al. 1991; Ryner and Baker 1991] and in vitro [Tian and Maniatis 1992]. In addition, the dsxRE is capable of suppressing both branchpoint sequence and pyrimidine tract mutations in heterologous introns [Tian and Maniatis 1992]. Thus, the dsxRE functions as a Tra and Tra2-dependent splicing enhancer.

Both Tra [Boggs et al. 1987] and Tra2 [Amrein et al. 1988; Goralski et al. 1989] contain arginine/serine-rich (RS) domains, which are also found in the SR family of general splicing factors (Ge et al. 1991; Krainer et al. 1991; Fu and Maniatis 1992; Zahler et al. 1992), the splicing factor U2AF [Zamore et al. 1992], the U1 small nuclear ribonucleoprotein [snRNP] 70K protein [Theissen et al. 1986; Mancebo et al. 1990], and the negative regulator of alternative splicing suppressor of white apicot [su(w)] [Chou et al. 1987]. Recent studies have shown that RS domains mediate associations between proteins containing this motif [Wu and Maniatis 1993; Amrein et al. 1994; Kohtz et al. 1994] and are responsible for directing splicing factors to subnuclear regions [speckles] where many splicing factors are concentrated [Fu and Maniatis 1990; Li and Bingham 1991; Spector et al. 1991; M.L. Hedley, H. Amrein and T. Maniatis, in prep.]. Tra2 also has an RNA recognition motif (RRM) that mediates protein–RNA interactions [for review, see Bandziulis et al. 1988, Kenan et al. 1991]. In contrast, Tra does not contain an RRM. Both the RS domain and the RRM are essential for the in vivo functions of Tra2 [Amrein et al. 1994].

Although the dsxRE can function as a regulated splicing enhancer, another type of splicing enhancer (the purine-rich element) can function constitutively [Lavigne et al. 1993; Sun et al. 1993a,b; Watakabe et al. 1993; Xu et al. 1993; Dirksen et al. 1994; Tanaka et al. 1994]. Although distinct from each other in sequence, the dsxRE and the purine-rich elements share several properties. Both types of elements are located in the exon and activate the splicing of upstream introns. In addition, both the dsxRE and the purine-rich elements can mediate the activation of heterologous splice sites. Finally, both elements serve as binding sites for splicing factors. Tra and Tra2 bind cooperatively with members of the SR family of general splicing factors to the dsxRE and promote the formation of a splicing enhancer complex [Tian and Maniatis 1993]. SR proteins and U1 snRNA have been shown to interact with purine-rich elements [Lavigne et al. 1993; Sun et al. 1993b; Watakabe et al. 1993].

The primary difference between these two types of
splicing enhancers is that the dsxRE requires Tra and Tra2, whereas the activity of the purine-rich element is constitutive. An additional difference is the relative positions of these elements with respect to the regulated introns. The dsxRE is separated from the female-specific 3' splice site by 300 nucleotides, whereas the purine-rich elements are generally located closer to the regulated intron. A recent study showed that a purine-rich element in the human fibronectin alternate ED1 exon loses its activity when placed beyond 293 nucleotides downstream of the 3' splice site (Lavigneur et al. 1993).

In this study we have investigated the characteristics of the dsxRE and its relationship to the purine-rich splicing enhancers. We find that the activity of the dsxRE is a function of its distance from the 3' splice site. When the element is located at a distance from the 3' splice site, its activity depends on Tra and Tra2. When placed close to the 3' splice site the dsxRE can function as a constitutive splicing enhancer, analogous to the purine-rich elements. The constitutive activity of the dsxRE may be mediated by SR proteins that can bind weakly to the dsxRE in the absence of Tra and Tra2.

Results

Tra and Tra2-independent activity of the dsxRE

To determine whether the activity of the dsxRE is affected by its distance from the female-specific 3' splice site, we analyzed the effects of altering this distance on dsx female-specific splicing in HeLa cell nuclear extracts. As shown previously, activation of the female-specific 3' splice site requires Tra and Tra2 when the dsxRE is located in its original position, 300 nucleotides downstream of the 3' splice site (Tian and Maniatis 1992; Fig. 1A, D1, lanes 1 and 2). Reducing the length of the spacer sequence to 190 nucleotides appeared to destabilize the pre-mRNA in nuclear extracts, but female-specific splicing remained Tra and Tra2-dependent (Fig. 1A, D2, lanes 3 and 4). Remarkably, when the length of the spacer sequence was reduced to 150 nucleotides, a significant level of splicing was observed in the absence of Tra and Tra2 (Fig. 1A, D3, lane 5). Addition of Tra and Tra2 appeared to stabilize the RNA but did not significantly change the ratio of splicing product to unspliced precursor (Fig. 1A, D3, lane 6). Further truncation of the spacer sequence to 100 nucleotides resulted in efficient female-specific splicing in the absence of Tra and Tra2, and the addition of Tra and Tra2 had no effect (Fig. 1A, D4, lanes 7 and 8). The constitutive female-specific splicing observed with D3 and D4 was dependent on the repeat sequence because female-specific splicing was not observed in the absence of the dsxRE [Tian and Maniatis 1992, see also Fig. 3A, D16, lane 1 [below]]. These results show that the dsxRE can function as a constitutive splicing enhancer when located near the female-specific 3' splice site. An alternative possibility is that the deleted spacer sequence normally blocks the constitutive activity of the dsxRE, and Tra and Tra2 are required to overcome this block. This possibility was ruled out by showing that dsxRE remained Tra and Tra2-dependent when the normal spacer sequence was substituted by a number of different sequences of the same size [data not shown].

Compared with the Tra and Tra2-dependent regulation, the constitutive activity of the dsxRE is somewhat more efficient in activating the female-specific 3' splice site (Fig. 1B, cf. lanes 1–5 with 6–10). To compare the constitutive activities of the dsxRE with that of a purine-rich element, we replaced the dsxRE with three copies of a purine-rich sequence from the IgM M2 exon [Watakabe et al. 1993]. As measured by the ratio of unspliced precursor to splicing product, the purine-rich element appeared to activate splicing to a similar extent as the dsxRE, although the RNA containing the purine-rich element was less stable during the incubation in nuclear extracts (Fig. 1B, cf. lanes 11–15 with 6–10).

In spite of similar constitutive activities, the dsxRE is distinguishable from the purine-rich element by Tra and Tra2 regulation, which allows the dsxRE to function at a distance from the regulated intron. Consistent with published reports, the purine-rich element lost its activity when placed 300 nucleotides downstream of the 3' splice site, and Tra and Tra2 had no effect on its function (Fig. 1C, lanes 1,2). In contrast, Tra and Tra2 could efficiently activate the female-specific 3' splice site when the dsxRE was placed at the same distance from the 3' splice site (Fig. 1C, lanes 3,4). To test whether Tra and Tra2 can function at a distance >300 nucleotides, we moved the dsxRE farther away from the female-specific 3' splice site by inserting a 200-nucleotide fragment. Significantly, Tra and Tra2 could efficiently activate the female-specific 3' splice site when the dsxRE was 500 nucleotides downstream of the 3' splice site (Fig. 1C, lanes 5,6). Furthermore, the splicing efficiency remained essentially unchanged with the insertion, implying that the Tra and Tra2-dependent regulation may function at even greater distances from the 3' splice site (Fig. 1C, cf. lanes 5–6 with 3–4).

In addition to analyzing the effect of distance on positive regulation, we also tested whether the dsxRE can function at locations other than downstream of the 3' splice site. Specifically, we found that the dsxRE failed to function when located either in the 5' exon upstream of the 5' splice site or within the intron [data not shown].

Suppression of 3' splice site mutations by the dsxRE

The weak constitutive activity of the dsx female-specific 3' splice site is attributable to a nonconsensus pyrimidine tract that is interrupted by purines (Burris and Baker 1989; Hoshijima et al. 1991). Tra and Tra2 activate the female-specific 3' splice site by suppressing the adverse effect of purine substitutions in the pyrimidine tract. In this study we show that the constitutive activity of the dsxRE can also activate the female-specific 3' splice site in the absence of Tra and Tra2, and exhibits somewhat higher activity compared with the Tra and Tra2-dependent regulation. To further compare the constitutive and Tra and Tra2-dependent activities of dsxRE, we tested
Positive regulation of pre-mRNA splicing

Figure 1. Tra and Tra2-independent activity of the dsxRE. (A) Autoradiograph of denaturing polyacrylamide gel analysis of in vitro splicing reactions using pre-mRNAs in which the distance between the 3' splice site and the dsxRE is progressively decreased. The positions of the unspliced pre-mRNAs and ligated exons are diagramed to the right of the autoradiograph. Because of the deletions, the precursors and splicing products decrease in size from the left to right and are labeled at an angle. Because the splicing reactions were incubated for 2.5 hr, the amount of splicing intermediates is low and the intermediates are not labeled. (cr) The splicing products involving cryptic splice sites. (Top) - and + indicate the absence or presence of Tra and Tra2 in the splicing reactions, respectively; the pre-mRNAs (D1-D4) used in the reactions are also indicated. The relevant region of clsx pre-mRNA from which the in vitro splicing substrates are derived is diagramed below the autoradiograph and includes the last common exon (E3), the intron, and the female-specific exon (E4). The slashed region in E4 represents the dsxRE. The number in E4 indicates the number of nucleotides between the female-specific 3' splice site and the first copy of the repeat sequence. The in vitro splicing substrates (D1-D4) are diagramed below dsx. (B) A time course of the in vitro splicing reactions. The autoradiographs are labeled as in A. The time of incubation in hours is indicated above each lane. The shaded area in D15 represents the purine-rich element. (C) Analysis of the effects of increasing the distance between the dsxRE (D27) or the purine-rich element (D26) and the 3' splice site. Because of the instability of D26 RNA, lanes 1 and 2 were exposed five times longer than lanes 3-6.
their abilities to suppress more extensive purine substitutions in the pyrimidine tract.

The longest stretch of pyrimidines in the nonconsensus pyrimidine tract of dsx female-specific 3' splice site contains 6 continuous pyrimidines. We disrupted this 6-pyrimidine stretch with 3 purine-substitutions [UC-UCUGAUCAUUAAACCGG (wild type) was changed to UCgaUGAUCAUUAAACCGG; the mutations are in lowercase letters]. Significantly, these mutations completely abolished the activation of the female-specific 3' splice site by Tra and Tra2 when the dsxRE was located beyond 190 nucleotides downstream of the 3' splice site (Fig. 2A, D7, D8, lanes 1–4). The small amount of remaining splicing products and intermediates results from splicing events involving cryptic 3' splice sites in the female-specific exon. In contrast to the complete inhibition of Tra and Tra2-dependent regulation, these pyrimidine tract mutations had only a mild effect on the constitutive activity of the dsxRE. A significant level of splicing was observed when the dsxRE was placed near the mutant 3' splice site (Fig. 2A, cf. lanes 5 and 8 with 7). The addition of Tra and Tra2 has a small inhibitory effect on splicing, presumably because of steric hindrance caused by the assembly of a large Tra and Tra2-dependent complex near the splice site (Fig. 2A, lane 6). A similar negative effect of Tra and Tra2 on splicing was occasionally observed for D4 as well [data not shown].

Previous studies demonstrated that the first step of the splicing reaction is independent of the 3' AG in the presence of a strong pyrimidine tract [Reed 1989]. Similarly, when the 3' AG of the female-specific 3' splice site was deleted, efficient accumulation of splicing intermediates, the released 5' exon and the lariat intermediate, was observed (Fig. 2B, lane 1). We noticed that the dsxRE contains a stretch of pyrimidines [UUCUCUUCUC-UUC]. It was therefore possible that this sequence could function as a pyrimidine tract in the first step of the splicing reaction. Furthermore, the dsxRE used in these experiments is 220 nucleotides in length and may contain other unidentified sequence elements responsible for the constitutive activity of dsxRE. To rule out these possibilities, we replaced the repeat-containing fragment with four copies of an oligonucleotide corresponding to the repeat sequence but lacking the surrounding sequences. We found that the synthetic dsxRE functions indistinguishably from the original dsxRE (Fig. 2B, lanes 2,3). Therefore, the repeat sequence is sufficient to mediate the constitutive activity of the dsxRE. Similarly, the purine-rich element can also efficiently suppress the same pyrimidine tract mutations and mediate the first step of splicing when the 3' AG is deleted (Fig. 2C, lanes 1–3). These results show that the constitutive activities of the dsxRE and the purine-rich elements can suppress more extensive purine substitutions in the pyrimidine tract than the Tra and Tra2-dependent activity. As negative controls, we replaced the dsxRE with a variety of sequences [data not shown]. We found that the antisense of dsxRE and several other unrelated sequences did not activate splicing. However, we did find that a few sequences can weakly stimulate splicing. In these cases, the RNA was unstable, and the splicing product was detected only after long exposure of the gel. Given the rather promiscuous RNA-binding property of SR proteins, we believe that these weak stimulatory activities could be ascribed to fortuitous sequence homology to SR protein-binding sites.

Introduction of the purine substitutions in the female-specific 3' splice site resulted in a sequence containing equal numbers of purines and pyrimidines and no continuous pyrimidine stretch. However, the mutant 3' splice site was still activated by the constitutive splicing enhancers. This observation led us to test the possibility that the dsxRE may completely substitute for the function of pyrimidine tract. Thus, we introduced additional purines into the pyrimidine tract. [UCUCUGAUCAUUAAACCGG (wild type) was changed to UCgaUGAUCAUUAAACCGG; these substitutions rendered the sequence preceding the 3' AG purine-rich, and they completely inhibited the constitutive activation of the female-specific 3' splice site by the dsxRE (Fig. 2A, cf. lanes 9 with 7 and 8). Thus, although the constitutive activity of the dsxRE can suppress severe mutations in the pyrimidine tract, it cannot substitute completely for the pyrimidine tract.

Suppression of 5' splice site mutations by the dsxRE

To determine whether the dsxRE can also suppress 5' splice site mutations, we mutated the 5' splice site of the dsx minigene transcript. To eliminate or minimize the effect of the dsxRE on the weak female-specific 3' splice site, we substituted the original nonconsensus pyrimidine tract [UCUCUGAUCAUUAAACCGG] with the sequence UCUCUGAUCAUUAAACCGG/G. This substitution increases the pyrimidine content of the 3' splice site and results in constitutive splicing of the intron in the absence of the dsxRE (Fig. 3A, D17, lane 2; cf. with D16, lane 1). The original 5' splice site matches the consensus at eight of nine positions [5' splice site consensus, (C/A)AG/GU(A/G)AGU; dsx 5' splice site, AG/GUAAGU]. In the absence of the dsxRE, one point mutation in the 5' splice site [AGc/GUAGU; the mutation is the lowercase letter] significantly decreased the level of in vitro splicing activity (Fig. 3A, D18, lane 3). Moreover, no splicing was observed when two point mutations were introduced into the 5' splice site [AGc/GUAaAGU; Fig. 3A, D19, lane 4].

When the dsxRE was placed near the 3' splice site of the intron containing 5' splice site mutations, it not only stimulated the splicing of the intron containing a single point mutation (Fig. 3B, D20, lane 1) but also led to detectable splicing of the intron with two point mutations in the 5' splice site (Fig. 3B, D21, lane 2). One complication in comparing the splicing of 5' splice site mutants with or without the dsxRE is the different stability of the RNAs. In the absence of the dsxRE, the RNAs were very unstable. However, even after long exposures of the gel, we could not observe splicing products with D19. Thus, we believe that the splicing product observed with D21
Figure 2. Functional interaction between the pyrimidine tract and the dsxRE. [A] Analysis of the effects of purine substitutions in the pyrimidine tract of the female-specific 3' splice site on in vitro splicing. Py represents the pyrimidine tract in the female-specific 3' splice site. The shaded Py in D7–D9 represents a mutant pyrimidine tract in which three pyrimidines are replaced by purines (see text). The solid box in D25 represents the pyrimidine tract substituted further with purines (see text). [B] Analysis of the effect of pyrimidine tract mutations in the absence of 3' AG. The accumulated lariat intermediate and the excised 5' exon are diagramed together with unspliced pre-mRNA and ligated exons. The shaded AG indicates that the 3' AG of the female-specific 3' splice site has been deleted. The stippled region in D11 and D12 represents four copies of synthetic repeat sequence. [C] Analysis of the suppression of 3' splice site mutations by the purine-rich sequence. The shaded area in D13–D15 represents the purine-rich sequence.
was attributable to splicing activation, rather than an increase in RNA stability. We also tested whether Tra and Tra2 can activate the mutated 5′ splice site when the dsxRE is in its original position, 300 nucleotides downstream of the female-specific 3′ splice site (Fig. 3C). Because of the increase in pyrimidine content in the female-specific 3′ splice site, constitutive splicing was observed for the pre-mRNAs with either a wild-type 5′ splice site or a 5′ splice site containing one point mutation (Fig. 3C, D22 and D23, lanes 3 and 5; cf. D1, lane 1). Tra and Tra2 could further stimulate the splicing of both pre-mRNAs (Fig. 3C, D22 and D23, lanes 4 and 6; cf. with lanes 3 and 5, respectively). No splicing was observed when the 5′ splice site contained two point mutations, and the addition of Tra and Tra2 resulted in barely detectable splicing (Fig. 3C, D24, lanes 7 and 8). These results show that both the regulated and constitutive activities of the dsxRE can suppress mutations in the 5′ splice site. This suppression could be a direct effect on the 5′ splice site or an indirect consequence of enhancing the strength of the 3′ splice site. The failure of Tra and Tra2 to suppress the more severe 5′ splice site mutation could simply be attributable to the fact that the Tra and Tra-2-dependent activity at a distance is weaker than the constitutive activity of the dsxRE.

To determine whether the dsxRE must be positioned near the 5′ splice site for efficient suppression of 5′ splice site mutations, we placed the dsxRE in the 5′ exon upstream of the mutated 5′ splice site. We found that the dsxRE failed to activate the 5′ splice site at this position (data not shown). The suppression of 5′ splice site mutations by the dsxRE was significantly less than that observed with 3′ splice site mutations, suggesting that the dsxRE may act primarily on the 3′ splice site.

**Association of SR proteins with the dsxRE in the presence and absence of Tra and Tra2**

We showed previously that Tra and Tra2 promote the assembly of a multicomponent complex (the dsx repeat...
compared on the dsxRE, and this complex contains a subset of SR proteins in addition to Tra and Tra2 (Tian and Maniatis 1993). In this study, we found that the dsxRE can mediate positive regulation in the absence of Tra and Tra2 when placed close to the 3' splice site. This observation suggests that general splicing factors may associate with the dsxRE in the absence of Tra and Tra2. The association of general splicing factors with the dsxRE in the absence of Tra and Tra2 may be weak and may therefore have eluded detection in the previous study (Tian and Maniatis 1993). To test this possibility, we scaled up the affinity purification of the repeat complex (Tian and Maniatis 1993) and used the monoclonal antibody mAb 104 [Roth et al. 1991] to probe Western blots of the associated proteins. mAb 104 recognizes all of the known SR proteins through a common phosphoepitope [Zahler et al. 1992]. As shown in Figure 4, low levels of SRp30 could indeed be detected in the repeat complex formed in the absence of Tra and Tra2 (lane 2). The association of SRp30 with the repeat complex was specific, because it was not detected in the control complex formed on an RNA without the repeat sequence (Fig. 4, cf. lanes 1 with 2). Consistent with our previous report, Tra and Tra2 significantly increased the amount of SR proteins that stably associate with the complex (Fig. 4, cf. lanes 2 with 3). The prominent band comigrating with SRp40 is most likely attributable to the nonspecific staining of glutathione S-transferase (GST)–R17 protein used in the affinity purification of the complex. The specific association of SR proteins with the dsxRE in the absence of Tra and Tra2 is also consistent with observations made in previous UV cross-linking experiments [Tian and Maniatis 1992]. In the absence of Tra and Tra2, two proteins from nuclear extracts (33 and 25 kD, respectively) could be specifically cross-linked to the dsxRE. These two proteins most likely correspond to SRp30 and SRp20, because the two cross-linking activities copurify with the SR proteins during the fractionation of nuclear extracts [data not shown]. The failure to detect SRp20 in the Western analysis may be attributable to the small amount of this protein in the complex in the absence of Tra and Tra2. SRp30 is a mixture of two SR proteins, SF2/ASF and SC35 [Ge et al. 1991; Krainer et al. 1991; Fu and Maniatis 1992; Zahler et al. 1992]. At present, we do not know whether one or both of these proteins are required for the constitutive activity of the dsxRE.

**Discussion**

In previous studies, the dsxRE was shown to function as a splicing enhancer in the presence of Tra and Tra2, and to exhibit no activity in the absence of these splicing regulatory proteins. In contrast, the purine-rich elements were known to activate splicing constitutively. For this reason, the dsxRE was considered distinct from the purine-rich elements. However, in this study we show that the dsxRE can also act as a constitutive splicing enhancer indistinguishable from the purine-rich elements. When placed within 150 nucleotides of the 3' splice site, the dsxRE can activate splicing in a Tra and Tra2-independent manner. When separated from the 3' splice site by 190–500 nucleotides, the dsxRE requires Tra and Tra2 for its activity. Thus, the dsxRE exhibits both constitutive and regulated activities, depending on its distance from the 3' splice site.

We compared the effect of the constitutive and Tra and Tra2-dependent activities of the dsxRE on mutations in both 3' and 5' splice sites. We found that the constitutive activity of the dsxRE could suppress more extensive purine substitutions in the pyrimidine tract than the Tra and Tra2-dependent regulation. Whether this difference is attributable to the relative strengths of the two enhancer activities or reflects a difference in mechanism remains to be determined.

We also showed that both the constitutive and the Tra and Tra2-dependent activities of the dsxRE could suppress mutations in the 5' splice site. However, the efficiency of suppression of 5' splice sites was significantly less than that observed with 3' splice site mutations. Recently, the purine-rich element of bovine growth hormone pre-mRNA was shown to activate the splicing of an upstream intron containing a weak 5' splice site [Dirksen et al. 1994]. In both of these cases, it remains to be determined whether the enhancers act directly on the weak 5' splice site or indirectly by increasing the strength of the 3' splice site. Similarly, we demonstrated previously that the Tra and Tra2-dependent activity of the dsxRE could suppress the effects of mutations in the branchpoint sequence [Tian and Maniatis 1992]. Again, this effect could be an indirect consequence of strengthening interactions at the pyrimidine tract rather than
directly promoting the interactions of splicing components such as U2 snRNP with the branchpoint sequence.

The striking similarity in the ability of the constitutive activity of the $dsxRE$ and the purine-rich enhancer to suppress purine substitutions in the pyrimidine tract suggests that the two elements may function through a common mechanism. We showed previously that Tra and Tra2 promote the formation of a multiprotein complex on the $dsxRE$, and this complex contains SR proteins and other unidentified proteins (Tian and Maniatis 1993). In this study we show that SR proteins can specifically associate with the $dsxRE$ in the absence of Tra and Tra2, but the level of SR proteins bound to the $dsxRE$ is substantially increased by the addition of Tra and Tra2. SR proteins have also been shown to bind to purine-rich elements (Lavigueur et al. 1993; Sun et al. 1993b).

Tra, Tra2, and SR proteins may interact with splicing factors that recognize the pyrimidine tract and thereby stabilize their association with the pyrimidine tract. Two splicing factors have been shown to bind specifically to the pyrimidine tract, U2AF (Zamore and Green 1989, 1991; Zamore et al. 1992) and pyrimidine tract-binding protein [PTB]-associated splicing factor [PSF] (Patton et al. 1993). The splicing factor U2AF is present in the earliest prespliceosome complex E (Bennett et al. 1992) and is required for the formation of the first ATP-dependent prespliceosome complex A (Zamore and Green 1989, 1991; Zamore et al. 1992). In contrast, the splicing factor PSF specifically associates with C complex and is essential for the second catalytic step in the splicing reaction (Gozoni et al. 1994). The intron-binding protein [IBP] also binds to the pyrimidine tract, but if it is a functional splicing component it will not act at the earliest steps of spliceosome assembly, because it is specifically associated with U5 snRNA (Gerke and Steitz 1986; Tazi et al. 1986). Finally, a number of heterogeneous nuclear RNP [hnRNP] proteins have been shown to interact with pyrimidine-rich sequences (Swanson and Dreyfuss 1988; Gil et al. 1991; Patton et al. 1991), but these proteins dissociate from pre-mRNA during spliceosome assembly (Staknis and Reed 1994).

Thus, the most likely target for splicing enhancers is U2AF, which consists of two subunits, U2AF$^{65}$ and U2AF$^{35}$ (Zamore and Green 1989, 1991; Zamore et al. 1992). This possibility is consistent with a number of observations. First, the purine-rich enhancer has been shown to promote A complex formation (Sun et al. 1993a) and could also be required for E complex assembly. Second, U2AF is the only known pyrimidine tract-binding protein that has been detected in E complex (Bennett et al. 1992). Third, Tra, Tra2, and SR proteins can interact with one another through the RS domain (Wu and Maniatis 1993; Amrein et al. 1994), and each of these proteins can interact, in turn, with U2AF$^{35}$ (Wu and Maniatis 1993). Thus, the SR proteins bound to the splicing enhancers may interact with U2AF$^{35}$ to stabilize the binding of the U2AF heterodimer to the pyrimidine tract.

The interaction between the SR proteins and U2AF may also provide an explanation for the observation that when it is close to the 3' splice site, the activity of $dsxRE$ is independent of Tra and Tra2 and can suppress more extensive purine substitutions in the pyrimidine tract. According to the model of Figure 5, the intrinsic affinity of U2AF for the weak 3' splice site and that of the SR proteins for the $dsxRE$ is low (Fig. 5A). The close proximity of the $dsxRE$ and the 3' splice site may facilitate cooperative interactions between U2AF and the SR proteins (Fig. 5B). This cooperative interaction would mutually stabilize their individually weak binding to their respective binding sites. In contrast, when the $dsxRE$ is at a distance from the 3' splice site, this cooperative interaction would not occur in the absence of Tra and Tra2 (Fig. 5C). Thus, the SR proteins and U2AF would dissociate from the RNA before they could interact. However, according to the model, Tra and Tra2 would

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**Figure 5.** Model for the mechanism of the constitutive and the Tra and Tra2-dependent activities of the $dsxRE$. (A) Individually, U2AF and SR proteins bind weakly to a nonconsensus pyrimidine tract [Py] and the $dsxRE$ (RE), respectively. (B) Cooperative interactions between SR proteins and U2AF stabilize their association with the $dsxRE$ [RE] and the pyrimidine tract [Py], respectively, when the two binding sites are in close proximity. (C) Tra and Tra2 stabilize the binding of SR proteins to the $dsxRE$ [RE] and facilitate the cooperative interactions between the splicing enhancer complex with U2AF at the distant 3' splice site.
stabilize the association of SR proteins with the dsxRE, thus facilitating cooperative interactions between the splicing enhancer complex and U2AF.

Materials and methods

In vitro splicing reactions

In vitro splicing reactions were carried out as described previously (Tian and Maniatis 1992).

In vitro splicing substrates

D1 contains 84 nucleotides of dsx exon 3, the complete intron between exon 3 and 4 (114 nucleotides), and 536 nucleotides of exon 4. D2, D3, and D4 contain progressively larger deletions starting 30 nucleotides from the first copy of the repeat sequence. The number of nucleotides between the repeat sequence and the female-specific 3′ splice site after the deletion is indicated in the figures. D7, D8, and D9 are derived from D1, D2, and D4, respectively, with mutations in the pyrimidine tract of the female-specific 3′ splice site [TCTCTCGATCT-GATCTAAACCAG/G is mutated to TCgATGCATCTGATC- TCTAAACCAG/G]. D10 is derived from D9 by deleting a 30-nucleotide fragment containing the 3′ AG. D11 and D12 are derived from D10 and D9, respectively, by replacing the repeat region with underlined). D13, D14, and D15 are derived from D9, D10, and D4, respectively, by replacing the repeat region with three copies of an oligonucleotide corresponding to the S3 element in the IgM M2 exon (CATGGGAGGACAGGACAG; the 3′ element is underlined). D16 contains 84 nucleotides of exon 3, the intron between exon 3 and 4 (114 nucleotides), and 70 nucleotides of exon 4. D17 is derived from D16 with 5′ splice site mutations as in D18 and D19, respectively. D20 is derived from D17 with pyrimidine substitutions in the female-specific 3′ splice site and the repeat region.

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