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Pick one, but be quick: 5′ splice sites and the problems of too many choices

Xavier Roca, Adrian R. Krainer and Ian C. Eperon

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Splice site selection is fundamental to pre-mRNA splicing and the expansion of genomic coding potential. 5′ Splice sites (5′ss) are the critical elements at the 5′ end of introns and are extremely diverse, as thousands of different sequences act as bona fide 5′ss in the human transcriptome. Most 5′ss are recognized by base-pairing with the 5′ end of the U1 small nuclear RNA (snRNA). Here we review the history of research on 5′ss selection, highlighting the difficulties of establishing how base-pairing strength determines splicing outcomes. We also discuss recent work demonstrating that U1 snRNA:5′ss helices can accommodate noncanonical registers such as bulged duplexes. In addition, we describe the mechanisms by which other snRNAs, regulatory proteins, splicing enhancers, and the relative positions of alternative 5′ss contribute to selection. Moreover, we discuss mechanisms by which the recognition of numerous candidate 5′ss might lead to selection of a single 5′ss and propose that protein complexes propagate along the exon, thereby changing its physical behavior so as to affect 5′ss selection.

Questions about the mechanisms by which 5′ splice sites (5′ss) are selected are deeply rooted in the history of research on pre-mRNA splicing. Identification of the sequences associated with 5′ss triggered the first key insights into splicing mechanisms, efforts that are reflected now in the widespread use of genomic methods to quantify the contributions of other sequences and their cognate factors. The first factors shown to modulate alternative splicing affected 5′ss selection, and the difficulties of working out the molecular mechanisms involved provided a foretaste of the complexities awaiting investigations into other regulatory proteins. Despite the many insights resulting from such studies over the years, it is clear that our conceptual frameworks are not yet adequate. New ideas and models are needed for studies on splice site selection. One purpose of this review is to emphasize that developing new ideas may involve first the challenge of uprooting commonsense but unsubstantiated preconceptions hidden in established models.

The 5′ss is involved in both steps of splicing. In the first step, the 2′-hydroxyl group of the branchpoint adenosine attacks the phosphodiester bond at the 5′ss and displaces the 5′ exon; in the second step, the 3′-hydroxyl group of the 5′ exon attacks the phosphodiester bond at the 3′ splice site (3′ss) and displaces the lariat intron. Splicing was discovered just before new, gel-based DNA-sequencing methods transformed molecular biology. Thus, although the original discoveries were made without the benefit of sequence information (Berget et al. 1977; Chow et al. 1977), sequences emerged rapidly thereafter and revealed clear similarities among 5′ss. Moreover, the “consensus” sequence (comprising, at each position, the nucleotide most commonly found there) was complementary to the sequence at the 5′ end of U1 small nuclear RNA (snRNA), which immediately suggested a mechanism for recognition of the 5′ss (Fig. 1A; Lerner et al. 1980; Rogers and Wall 1980). An additional short stretch of complementarity between U1 snRNA and the 3′ss region does not mediate an interaction (Seraphin and Kandels-Lewis 1993). There were two interesting and important ways in which early preconceptions shaped the subsequent development of the field: The base-pairing between all 5′ss and U1 snRNA was assumed to be in a constant register, and the “consensus” sequence was assumed to be an optimal 5′ss (Rogers and Wall 1980).

The original sequence compilations suggested that a limited variety of sequences might be recognized as 5′ss, the sequence within the consensus region being sufficient to both define sites and ensure their usage. However, such a simple view became implausible with the ever-expanding lists of actual mammalian 5′ss sequences: A human 5′ss compilation now contains >9000 sequence variants in the −3 to +6 region of the 5′ss (Roca et al. 2012). Moreover, three observations showed the inadequacy of such a model. First, pre-mRNAs were found to contain sequences that matched the 5′ss consensus as well as or better than the actual 5′ss but were not used (now termed pseudo-5′ss), demonstrating that either sequence could not be the only determinant of use or the consensus might not be optimal. Second, some sequences in β-globin that

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**Keywords:** splicing, U1 snRNA; pre-mRNA; 5′ splice sites; exons

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"Pick one, but be quick: 5′ splice sites and the problems of too many choices"
recognise 5’ss were used when a natural 5’ss was inactivated [termed cryptic 5’ss] (Treisman et al. 1983; Wieringa et al. 1983), demonstrating that the use or avoidance of 5’ss could depend on other sites and that it was not an intrinsic property of any given sequence. Finally, an adenovirus gene [E1A] was shown to use both of two alternative 5’ss, and the ratio of use depended on the sequences of the sites—meaning that 5’ss were in competition [Montell et al. 1982]. These findings had important consequences.

The first consequence of the discovery of competitive alternative splicing for studies on 5’ss selection was that it allowed a genetic test in mammals of the role of U1 snRNA. By transfecting cells with a “suppressor” U1 gene containing appropriate mutations at positions hypothesized to be complementary to one of two alternative 5’ss, it was possible to shift the relative use of the two 5’ss in adenovirus E1A transcripts [Zhuang and Weiner 1986]. While it was not possible to complement bases at all positions in the 5’ss, the results confirmed the hypothesis that the 5’ end of U1 snRNA recognizes 5’ss by base-pairing and showed that the extent of base-pairing affects competition. Earlier tests of the role of the 5’ end of U1 snRNA by RNase H cleavage had shown that it was required for splicing [Kramer et al. 1984], but the absence of U1 small ribonucleoprotein particles (snRNPs) from some spliceosome preparations had raised doubts [Konarska and Sharp 1986]. The role of U1 snRNA in yeast was not clear at first because the 5’ end of U1 snRNA is perfectly conserved even though the consensus 5’ss has a mismatch to U1 snRNA [Siliciano et al. 1987]. The suppression of mutations in 5’ss by mutant U1 snRNA genes confirmed that U1 snRNA is essential in yeast as well (Seraphin et al. 1988; Siliciano and Guthrie 1988).

The second consequence of competitive alternative splicing was that it suggested that the relationships between 5’ss sequences, their strength, and their U1 base-pairing potential could be explored systematically. Previous work had shown that introducing mutations into a single 5’ss might inactivate it or lead to the use of cryptic 5’ss, but it was difficult to quantify the strength of each sequence [Aebi et al. 1986]. However, if there were an alternative 5’ss that was used to some extent, then it could provide a reference site against which test sites could be calibrated. The first system used a rabbit β-globin gene in which test sequences of 16 nucleotides [nt] were inserted to create potential alternative sites 25 nt upstream of the normal 5’ss. When the test sequence was a duplicate of the normal site’s sequence, both 5’ss were used after transient transfection of HeLa cells with the construct. The first experiments showed that the consensus sequence (CAG/GUAAGU) was the most potent and that it silenced the normal 5’ss [Eperon et al. 1986]. Much more extensive details on the 5’ss motif and its interaction with U1 as well as the influence of proteins in this recognition event are described below in “5’ss Preferences and U1 Base-Pairing Potential,” “5’ss Recognition Is Not Always Dependent on U1 snRNA Base-Pairing,” and “Extrinsic Factors Affecting 5’ss Choices.” However, even these initial results raised an interesting question: How can a perfectly good natural 5’ss, the reference site, be silenced? What, in fact, is the mechanism by which recognition by U1 snRNPs is turned into selection? These questions are addressed below in “How Does 5’ss Recognition Turn into Selection.”

**5’ss preferences and U1 base-pairing potential**

As outlined above, the role of base-pairing between the 5’ss and the 5’ end of U1 snRNA was firmly established in the mid-1980s [Zhuang and Weiner 1986]. Around the same time, 5’ss competition experiments provided the first functional tests to estimate 5’ss strength by comparing the splicing efficiency of test 5’ss sequences relative to a reference 5’ss. In these experiments, multiple 5’ss could be ranked based on their splicing efficiency, and their ranks correlated reasonably well with base-pairing potential to U1 snRNA, as estimated by thermodynamic
parameters. Such experiments provided a firm basis for models involving 5′ss recognition by U1 snRNA.

The dsRNA helix that forms upon base-pairing between a 5′ss and the 5′ end of U1 snRNA has a maximum length of 11 base pairs [bp] (Fig. 1A), as the 12th nucleotide of U1 is already engaged in an internal base pair in stem I. Not all base pairs at different 5′ss positions are equally important, and their contribution to splicing roughly correlates with their conservation [Fig. 1B]. The most conserved 5′ss positions lie at the first two intronic nucleotides (+1 and +2), which determine the 5′ss subtype. The GU subtype, with Watson-Crick complementarity with A7 and C8 in U1, accounts for ~99% of 5′ss. The minor subtypes have a mismatch to U1 at either +1 or +2 and include the GC [0.9%] and the very rare AU 5′ss recognized by the major spliceosome [only 15 cases in humans] (Sheth et al. 2006). The most next conserved 5′ss positions (>75% in humans) are −1G (the last exonic nucleotide) and +5G, which form strong G-C base pairs with U1, with three hydrogen bonds [Fig. 1B]. Consensus nucleotides −2A, +3A, +4A, and +6U are also conserved and have a lesser yet important contribution to 5′ss strength because their base pairs to U1 contribute only two hydrogen bonds. Gs at positions −2, +3, and +4 can also establish weaker wobble base pairs [G-U or G-Ψ] (see below), which are very frequent at position +3. The consensus −3C forms a C-G base pair with U1, but the conservation of this nucleotide and its contribution to splicing are less important, probably because this base pair is weakened by the adjacent U1 stem I. In invertebrates, the 5′ss motif is very similar yet with reduced conservation of exonic nucleotides (Sheth et al. 2006). In budding yeast, exonic nucleotides are not conserved at all, the intronic positions +1 to +6 are nearly invariant, and the +4A is replaced by +4U, which constitutes a mismatch to U1.

The 5′ss positions +7 and +8 do not exhibit substantial conservation in humans [Fig. 1B], yet several lines of evidence indicate that these positions can base-pair to U1 and contribute to splicing (Lund and Kjems 2002, Hartmann et al. 2008). In budding yeast, a hyperstable 5′ss/U1 helix with 10 or 12 bp [including those at positions +7 and +8] impedes splicing by reducing the off-rate of U1 (Staley and Guthrie 1999), whose displacement by U6 is a necessary step during spliceosome assembly. However, base pairs at +7 and +8 were shown to enhance splicing kinetics in human cells and extracts (Freund et al. 2005), indicating that the contribution of such 5′ss positions is species-specific.

The first 2 nt at the 5′ end of U1 have methylated riboses, but this modification is not expected to change the stability of the base pairs at these positions. A more important modification in this context is the replacement of the uridines at positions 5 and 6 with pseudouridines [Ψs] (Reddy et al. 1981). A Ψ is a regiosomer of uridine with analogous groups for hydrogen bond donors and acceptors and an extra imino group that contacts the sugar-phosphate backbone and stabilizes base stacking (Davis 1995). Consistently, thermodynamic experiments revealed a slightly higher stability of 5′ss/U1 helices with Ψs compared with those with unmodified Us (Hall and McLaughlin 1991; Roca et al. 2012).

Since the late 1980s, many algorithms have been developed to estimate 5′ss strength. These 5′ss scoring methods rely on either large-scale collections of genomic 5′ss or estimations of the 5′ss/U1 base-pairing stability. The methods in the first category assume that the most common 5′ss nucleotides and/or sequences are most efficient for splicing. The earliest and simplest algorithm used alignments of many 5′ss to derive position-weight matrices [PWMs], which account for the frequency of each nucleotide at each position (Shapiro and Senapathy 1987; Senapathy et al. 1990). Later PWMs were further processed using information content theory (Rogan and Schneider 1995). These methods assume independence between 5′ss positions, yet there is now ample evidence for complex associations between 5′ss positions (Burge 1998, Carmel et al. 2004, Roca et al. 2008). Other algorithms—like first-order Markov models, decision trees, and maximum entropy models—take into consideration these associations (Yeo and Burge 2004). Machine-learning approaches based on neural networks use overall sequence patterns to infer 5′ss strength (Brumak et al. 1991, Krawczak et al. 2007). Methods considering the frequency of the whole 5′ss sequence [excluding positions +7 and +8] in the collection of natural human 5′ss have also proved useful to estimate 5′ss strength (Sahashi et al. 2007). The second class of methods is based on the assumption that U1 binding is the only force governing 5′ss selection. The most common method estimates the minimum free energy of each 5′ss/U1 helix using experimentally derived thermodynamic parameters known as nearest-neighbor “Turner” rules (Mathews et al. 1999), although another algorithm using hydrogen-bonding patterns also exists (Freund et al. 2003; Hartmann et al. 2008). Overall, the scores correlate well among different algorithms and are useful in estimating 5′ss strength, yet they all have their limitations in matching the 5′ss strengths derived experimentally. Partly explaining their limitations, most [but not all] of these methods ignore the contribution of positions +7 and +8.

Another limitation of most methods is that they assume that all 5′ss are recognized by U1 using the same “canonical” base-pairing register, defined by U1 C8 nucleotide base-pairing to 5′ss +1G, and without bulged nucleotides. Recently, however, mutational analyses and suppressor U1 experiments demonstrated that subsets of 5′ss are recognized by U1 using alternative base-pairing registers. First, certain presumptively weak 5′ss were shown to be efficiently used because U1 base-pairs to them in a register that is shifted by 1 nt, so that 5′ss +1G base-pairs to U1 C9 (Fig. 1C; Roca and Krainer 2009). Secondly, many other 5′ss are more stably bound by U1 when a nucleotide is bulged at either the 5′ss (various positions) or the 5′ end of U1 [only the Ψs], and these base-pairing schemes were collectively termed bulge registers (Fig. 1D; Roca et al. 2012). While the shifted register was only estimated to apply to a few 5′ss [59 in humans], the bulge registers appear to be much more frequent, potentially accounting for the recognition of 5% of all human 5′ss. These registers highlight the flexibility of the in-
teraction between 5′ss and U1, allowing for many base-pairing arrangements to result in efficient splicing, and also provide a means for the efficient recognition of 5′ss that otherwise would be weakly bound by U1. Another implication of these registers is that the relevant 5′ss positions vary depending on the used register such that, for example, 5′ss +9 might be base-paired in shifted and some bulge registers. The redefinition of the length of the 5′ss motif as well as the consideration of these registers in new algorithms would certainly result in more accurate 5′ss scoring methods.

The 5′ss/U1 snRNA base-pairing appears to be the main determinant of 5′ss strength, yet this interaction can be affected by other factors binding at the same sequence. The U1C polypeptide, a specific protein of the U1 snRNP, has been shown to bind to 5′ss in the absence of U1 base-pairing [Du and Roshbash 2002]. A recent crystal structure of the U1 snRNP [Pomeranz Krummel et al. 2009] revealed certain U1C amino acids contacting the minor groove of the base pairs established by A7 and C8 at the 5′ end of U1, possibly explaining the nearly universal conservation of 5′ss positions +1G and +2U. Certain heterogeneous nuclear RNPs [hnRNPs] like hnRNP A1 and hnRNP H, which are abundant proteins extrinsic to the U1 snRNP, have been proposed to bind some 5′ss sequences, thereby competing with U1 base-pairing [Fig. 1F; Buratti et al. 2004; De Conti et al. 2012].

5′ss can also be involved in internal base-pairing interactions with other pre-mRNA sequences, and such secondary structures would compete with U1. The first proof for such steric hindrance models was provided by 5′ss competition studies, which showed that a 5′ss in an internal stem had a selective disadvantage over a free 5′ss [Eperon et al. 1986, 1988], and others in which an entire exon was skipped when located in an internal RNA loop [Solnick 1985]. After that, numerous studies provided evidence for the influence of secondary structures in 5′ss selection and splicing [Jin et al. 2011]. A remarkable example is the MAPT also known as tau alternative exon 10, whose inclusion levels are determined by the efficiency of 5′ss recognition, which is compromised because of an internal base-pairing structure involving downstream intronic nucleotides [Fig. 1G; Donahue et al. 2006]. Mutations weakening the structure increased exon 10 inclusion, thereby disrupting gene function and causing a neurodegenerative disease. The involvement of a pre-mRNA structure in a particular splicing event is typically modeled by RNA structure prediction tools [Jin et al. 2011]. However, such predictions can be inaccurate because of the high number of different structures with similar stability, the interference of RNA-binding proteins—mainly from the hnRNP family—in structure formation [Solnick and Lee 1987], and the likely dependence of the structures on the rates of transcription [Eperon et al. 1988]. Encouragingly, the traditional and labor-intensive methods for testing RNA structure, based on chemical or enzymatic probing, have been recently complemented by new technologies that allow high-throughput evaluation of structures [Underwood et al. 2010].

Overall, the pivotal role of 5′ss/U1 base-pairing in 5′ss recognition is well established. However, this short RNA duplex is far from a simple structure, as many variations and subtle determinants of 5′ss strength are being revealed. In “5′ss Recognition Is Not Always Dependent on U1 snRNA Base-Pairing,” “Extrinsic Factors Affecting 5′ss Choices,” and “How Does 5′ss Recognition Turn into Selection,” we discuss U1-independent mechanisms, the influence of proteins in 5′ss selection, and how all these processes are integrated to finally commit a 5′ss for splicing.

5′ss recognition is not always dependent on U1 snRNA base-pairing

Some studies have questioned the strict dependence of 5′ss use on U1 snRNA/snRNP in general or in particular cases. The most obvious examples of U1-independent 5′ss belong to the U12-type [or U12-dependent] introns [0.34% of all introns in humans] [Sheth et al. 2006]. Splicing of U12-type introns is catalyzed by the minor spliceosome, comprising U11, U12, U4atac, and U6atac in lieu of U1, U2, U4, and U6 snRNPs, respectively, and sharing the U5 snRNP with the major or U2-type spliceosome [Will and Luhrmann 2005]. The U12-type 5′ss conform to GU, AU, or noncanonical subtypes; have a distinct and highly conserved motif, and are recognized by base-pairing to U11 snRNA.

During the maturation of the spliceosome, U1 is replaced by U6 at the 5′ss [Wassarman and Steitz 1992; Kandels-Lewis and Seraphin 1993; Lesser and Guthrie 1993], with U6 establishing up to 5 hp with the consensus 5′ss (Fig. 2A). This change requires ATP and the DExD/H-box RNA helicase Prp28 [Staley and Guthrie 1999]. The U6—but not U1—snRNP is an integral component of the active spliceosomal complexes and has been proposed to catalyze both transesterification reactions [Valadkhan 2010]. U1 and U6 snRNAs can bind at nearby yet different sequences within the pre-mRNA, and the site of transesterification is ultimately determined by U6 and not U1. This phenomenon was shown first using artificial sub-
strates [Cohen et al. 1994; Hwang and Cohen 1996] and later in the natural human FGF1 pre-mRNA, in which U6 can support splicing via a noncanonical GA 5’ss in the presence of a nearby U1-binding site [Fig. 2B; Brackenridge et al. 2003]. Despite these and other cases in which suppressor U6 snRNAs enhance use of a 5’ss [Kubota et al. 2011], in general, U6 is thought to play only a minor role in initial 5’ss selection.

A few reports have questioned the necessity of either the U1 snRNP or the base-pairing of U1 snRNA to the 5’ss for certain U2-type introns. Strikingly, splicing that was abolished after either depletion of U1 snRNPs or addition of oligonucleotides complementary to the 5’ end of U1 snRNA could be restored by the addition of SR proteins [Fig. 2C; Crispino et al. 1994; Tarn and Steitz 1994]. The efficiency of splicing after depletion was affected by the level of complementarity to U6 snRNA [Crispino and Sharp 1995] and by non-5’ss sequences in the intron [Crispino et al. 1996]. The ability to circumvent a block on base-pairing might be attributed to U1C binding to 5’ss (Kubota et al. 2006). A recent study reports the detection of such splicing events and reveal the prevalence of these mechanisms. Finally, many U1 snRNA variant genes and pseudogenes can be found in the human genome [Kyriakopoulou et al. 2006]. A recent study reports the expression of a subset of these U1 variants [O’Reilly et al. 2012], but their involvement in splicing is not clear.

Most importantly, all of these studies suggest that there is functional redundancy in the recognition of U2-type 5’ss by U1 base-pairing and other mechanisms, be it other components of the U1 snRNP, U6, or other factors. Such alternative mechanisms could enhance fidelity and provide a platform for regulation of 5’ss recognition. The role of U1 and/or other factors in early 5’ss recognition could be seen as just a mark on the substrate that—along with other marks—triggers assembly of splicesomal complexes and then splicing catalysis. As part of active splicesomal complexes, U6 determines the final trans-esterification site in a narrow but flexible sequence window centered at the site of U1 binding. Below, we describe how 5’ss recognition by U1 is influenced by proteins bound at nearby sequences and how a U1-tagged 5’ss becomes committed to splicing.

### Extrinsic factors affecting 5’ss choices

**SR proteins bound to exon sequences favor the nearest 5’ss downstream**

SR proteins comprise one or two RNA recognition motif [RRM]-type RNA-binding domains and a C-terminal RS domain, a region rich in arginine and serine [mostly as RS dipeptides] [Fig. 3A]. They are involved in splicing, nuclear export of mRNA, the control of translation, and nonsense-mediated decay [for reviews, see Bourgeois et al. 2004; Long and Caceres 2009; Shepard and Hertel 2011]. In general, U6 is thought to play only a minor role in initial 5’ss selection.

![Figure 3](genesdev.cshlp.org/images/figure3.png)

**Figure 3.** Proteins that generally activate splicing on the basis of position or sequence of the sites. (A) Diagram of an SR protein. There are one or two RRM-type RNA-binding domains, which may also mediate interprotein contacts, and a C-terminal RS domain, rich in RS or SR dipeptides. SRSF1 has two RRM domains with an intervening linker comprising arginine-rich sequences flanking a central block of nine consecutive glycines. This linker plays an important role in binding to ESEs [Cho et al. 2011a]. (B) Some mechanisms by which RS domains bound to an ESE might stimulate exon inclusion. (Solid arrow) Interactions in early complexes between the RS domains of the SR protein [blue] and U2AF65 [binds the 3’ss AG; small green oval] become critical if the polypyrimidine tract and U2AF65 binding [larger green oval] are weak, [dotted arrows] interactions in later complexes with the duplexes formed by base-pairing of U2 or U6 snRNAs with pre-mRNA. (C) SR proteins generally promote a shift in splicing toward the intron-proximal 5’ss when either the concentrations of some SR proteins are raised (as with SRSF1) or an ESE is inserted between the alternative 5’ss [as shown]. The relative levels of use of the two 5’ss are shown by the thickness of the brown arrows. (D) Effect of an ESE to the 5’ side of alternative 5’ss. (E) Effects of phosphorylation of the RS domain of SRSF1 [Cho et al. 2011b]. A hypophosphorylated RS domain interacts with the RRMs and prevents them from binding to the RRM of U1-70K. (F) Binding of TIA-1 RM2 and RM3 to interrupted U tracts just 3’ of the 5’ss leaves RRM1 free to augment the interactions of the glutamine-rich domain [Q] with U1-C [Bauer et al. 2012].
A well-known activity of SR proteins is to stimulate the inclusion of exons with weak splice sites. This appears to involve binding to exonic splicing enhancer sequences (ESEs) in the exon, followed by direct (looping) or indirect interactions of the RS domain with the adjacent pre-mRNA. The relevance of enhanced U1 snRNP binding to splice site selection was called into question [Valcarcel and Green 1996] when it was shown that, unlike 5’ss switching, it involved the RS domain of SRSF1, which appeared to interact with a similar domain on the U1 snRNP 70-kDa subunit (U1-70K) [Kohtz et al. 1994; Jamison et al. 1995]. However, the interaction between the two RS domains could be explained by bridging through contaminating RNA [Xiao and Manley 1998]. Moreover, other assays showed that the RS domain was not required for the enhancement of U1 snRNP binding [Eperon et al. 2000]. Recent results showed that the interaction between SRSF1 and U1-70K can be mediated by the RRMs of the two proteins, and the RS domain interferes with this by binding intramolecularly if it is hypophosphorylated. This suggests that phosphorylation of the RS domain is a switch that exposes the RRMs for interaction with the U1 snRNP, resulting in enhanced U1 snRNP binding to 5’ss [Fig. 3E; Cho et al. 2011b].

It remains unclear whether either U1 snRNP or SRSF1 binds the pre-mRNA first and also whether specific binding sites for SRSF1 have to be present. Sequences other than the 5’ss are not required for formation of a U1-snRNP/SRSF1/5’ss complex [Jamison et al. 1995; Zahler and Roth 1995]. However, ultraviolet cross-linking and immunoprecipitation (CLIP) analysis in HEK293T cells confirmed that SRSF1 binds to sequences with a loose consensus of GAAGARR [Sanford et al. 2009], fitting previous results based on selection in vitro [Tacke and Manley 1995; Liu et al. 1998]. These sequence motifs are enriched in exons within ~200 nt of splice sites, peaking at 20–40 nt from the sites. Binding involves cooperation between the RRMs and the intervening linker [Cho et al. 2011a].

Other proteins modulating splicing act more simply, by modulating the affinity of binding by U1 snRNPs only at recognizable sites. In yeast, the selection of weak alternative 5’ss can be modulated by the presence of a U-rich tract just downstream from the 5’ss. This tract is bound by Nam8p, which is a component of the U1 snRNP in yeast [Puig et al. 1999]. Although there is no equivalent stable component of mammalian U1 snRNP, the existence of similar tracts downstream from human 5’ss led to the identification of TIA-1, a homolog of Nam8p, and a close relative named TIA-R. TIA-1 has a number of functions in the cytoplasm, where it is involved in translational repression, but in the nucleus, it has been shown to direct splicing to specific 5’ss in which a U-rich tract begins 5–9 nt downstream [Del Gatto-Konczak et al. 2000; Forch et al. 2000]. Binding is synergistic: TIA-1 facilitates binding by U1 snRNP to the adjacent 5’ss [Forch et al. 2000], and the 5’ss and U1 snRNP enhance the binding of TIA-1 [Del Gatto-Konczak et al. 2000]. TIA-1 binds directly to the N-terminal part of the U1-C polypeptide [Forch et al. 2002]. TIA-1 consists of three RRMs and a glutamine-rich C-terminal sequence. The primary contact with U1-C is mediated by the glutamine-rich sequence, but the interaction is strengthened by
RRM1 [Forch et al. 2002]. A recent structural study suggested that high-affinity binding to polyU involves all three RRMs but that interruptions to the continuity of U-tracts, as in natural pre-mRNA, prevent the binding of RRM1 to RNA, liberating it to reinforce the interaction with U1-C [Fig. 3F; Bauer et al. 2012].

Another important sequence element is the G-triplet. Although these triplets were first identified as elements characteristic of short mammalian introns [McCullough and Berget 1997], they occur widely and are found preferentially toward the 5′ end of an intron, being most frequent at only 20–30 nt from the 5′ss [Xiao et al. 2009]. When inserted between alternative 5′ss, G-triplets stimulate use of the upstream site [McCullough and Berget 1997]. These properties suggested that the triplets stimulate the use of an adjacent upstream 5′ss. It was initially suggested that the motifs were themselves bound by U1 snRNPs, base-paired via nucleotides 8–10 of U1 snRNA [McCullough and Berget 2000]. However, G-triplets can be bound by hnRNP H family proteins [Caputi and Zahler 2001; Dominguez et al. 2010], and there is evidence that these may play a role in recruiting U1 snRNPs [Wang and Cambi 2009]. A more detailed analysis suggests that SR proteins are involved also and that the outcome depends on the nature of the 5′ss, distance from the sites, and, perhaps, the ability of G-runs to form quadruplex structures [Xiao et al. 2009; Wang et al. 2011].

Proteins that inhibit 5′ss by stabilizing U1 snRNP binding

The ability to stabilize U1 snRNP binding is not confined to proteins that activate splicing: A number of examples have been discovered in which U1 snRNP is part of an unproductive complex either at pseudo or bona fide 5′ss. A common feature is that the binding site for the protein is juxtaposed to the 5′ss, although spliceosome assembly may be stalled at different stages. Proteins that inhibit splicing when their binding site is in the exon include HMGA1A, which interacts with the U1-70K polypeptide of U1 snRNP [Ohe and Mayeda 2010]; TIA-1 [Erkelenz et al. 2013]; and hnRNP A1 [Yu et al. 2008]. A systematic screen identified several other sequence classes that inhibit splicing when inserted at ~7 relative to the 5′ss, although cognate proteins could not be identified [Yu et al. 2008]. The nature of the inhibition has not yet been defined, although it appears to operate at an early stage after U1 snRNP binding, preventing either formation of complex E [Ohe and Mayeda 2010] or progression to complex A [Erkelenz et al. 2013]. Other proteins that inhibit when bound within an exon not only act on U1 snRNP but stabilize an exon-defining prespliceosomal complex containing U1 and U2 snRNPs, preventing the components from forming cross-intron interactions [House and Lynch 2006; Bonnal et al. 2008].

Proteins can also form unproductive complexes when juxtaposed to the 5′ss on the intron side. Polypyrimidine tract-binding protein [PTB] binds to pyrimidine-rich tracts flanking the N1 exon of Sce in noneuronal cells, preventing progression beyond the prespliceosomal A complex [Sharma et al. 2008]. The explanation for this appears to lie in the ability of RRM1 and RRM2 to bind to stem–loop 4 of U1 snRNA, which is exposed in the snRNP. This may lock the U1 snRNA in a conformation that blocks further interactions [Fig. 4A; Sharma et al. 2011]. SRSF7 is also inhibitory when bound in the intron, where it prevents progression beyond complex E [Erkelenz et al. 2013], and several other classes of sequence are inhibitory when inserted at +11 relative to the 5′ss, including a sequence bound by hnRNP A1 [Yu et al. 2008]. Interestingly, it has been suggested on the basis of tests with several proteins that it is a general property of SR and hnRNP proteins that they stimulate splicing when bound close to the 5′ss on the exonic or intronic sides, respectively, and inhibit splicing when bound on the opposite side [Erkelenz et al. 2013]. hnRNP A1 may be an exception, since it inhibits when bound to a high-affinity site on either side [Yu et al. 2008]. Whereas HMGA1A and PTB have specific mechanisms for inhibition, it may be that proteins generally considered to activate splicing act as repressors when juxtaposed to U1 snRNP on the “wrong” side because they interfere sterically with the interactions made by U1

![Image](genesdev.cshlp.org)
snRNP shortly after its binding. The U1 snRNP extends its interactions with the pre-mRNA such that ~20 nt on either side of the 5′ss are protected against nucleases (Chabot and Steitz 1987), although the molecular basis and roles of this extended footprint are unknown.

Steric interference with the development of interactions by U1 snRNP may account also for the drastic inhibition seen when two strong 5′ss are in close proximity. It has been shown by both ribonuclease protection [Nelson and Green 1988; Eperon et al. 1993] and single-molecule experiments [Hodson et al. 2012] that the 5′ss are both occupied by U1 snRNPs. It is likely that the two snRNPs mutually block further interactions [Fig. 4B].

Effects and mechanisms of competition

A protein purified as an antagonist to the effects of SRSF1 on 5′ss selection turned out to be hnRNP A1 (Mayeda and Krainer 1992). This protein shifts 5′ss preferences to the intron-distal site and favors exon skipping rather than inclusion [Mayeda and Krainer 1992; Mayeda et al. 1993; Yang et al. 1994]. The protein has two RRMAs and a C-terminal glycine-rich domain that promotes cooperative binding and interactions with other proteins [Casas-Finet et al. 1993; Cartegni et al. 1996]. Like SRSF1, hnRNP A1 appears to affect 5′ss selection with no absolute requirement for a high-affinity binding site and acts indiscriminately on U1 snRNP binding at candidate 5′ss—in this case, to reduce U1 snRNP binding [Eperon et al. 2000]. Rather than interacting directly with U1 snRNPs or the U1–pre-mRNA complex, hnRNP A1 appears to act by competition with SR proteins and U1 snRNPs for binding to pre-mRNA [Eperon et al. 2000; Zhu et al. 2001]. The presence of a high-affinity binding site in an exon promotes skipping [Caputi et al. 1999; Del Gatto-Konczak et al. 1999], and when it is placed between alternative 5′ss, the glycine-rich domain promotes a shift to the distal 5′ss [Fig. 4C; Eperon et al. 2000; Wang et al. 2006]. High-affinity sites nucleate cooperative binding of pure hnRNP A1 that can displace other proteins as it spreads along the RNA, a process that is more efficient between two high-affinity sites [Fig. 4D; Zhu et al. 2001; Okumula and Krainer 2009]. However, there is also evidence that the presence of multiple sites flanking a 5′ss can repress it by the formation of looping interactions mediated by the glycine-rich domain [Blanchette and Chabot 1999; Nasm et al. 2002], which may also involve interactions with hnRNP H [Fig. 4E; Fisette et al. 2010]. The formation of loops does not prevent binding of U1 snRNPs but prevents further spliceosome assembly for unknown reasons [Nasm et al. 2002]. It remains to be seen whether these two models for the effects of high-affinity sites indicate that there are different mechanisms or that there is an undiscovered unifying mechanism. One of the major difficulties in investigating the mechanisms of action of hnRNP A1, as with SRSF1, is that there are likely to be numerous low-affinity sites.

How does 5′ss recognition turn into selection?

The 9000 different bona fide human 5′ss sequences in the –3 to +6 region [Roca et al. 2012] represent over half of all possible sequences, assuming an absolute requirement for a GU at +1/+2. This means that the consensus region restricts the number of GU motifs in a sequence that could be potential 5′ss by only a factor of two. This range of possible sites is so broad that most genes will comprise many more pseudo-5′ss than actual 5′ss. We described above the evidence that most sites are recognized by U1 snRNP, even if this requires base-pairing to accommodate altered registers and bulges. We summarized the evidence that indicates that in most cases, those factors that modulate 5′ss usage act via a U1 snRNP by either stabilizing its interactions at a 5′ss, competing for binding, or binding adjacent to the U1 snRNP and stabilizing it in an inactive conformation. This leaves the focus of our attention on the question of how, if U1 snRNP recognizes so many sites, it can also be involved in selecting them. To put the same question differently: If U1 snRNPs mark the 5′ss to be used, then how does the mechanism ensure that only one U1 snRNP marks each intron?

U1 snRNPs may bind independently to multiple candidate 5′ss

If the main determinant of 5′ss use is the affinity of interactions with U1 snRNP, we need to ask how these affinities become apparent to the splicing apparatus. The answer will depend on how U1 snRNP binds the pre-mRNA. Multiple U1 snRNPs might interact independently with all candidate sequences [Fig. 5A], or a single U1 snRNP might be recruited by, for example, components at the 3′ss and then sample all local candidates [Fig. 5B]. The latter model fits well with what is known of complex E. This is an ATP-independent complex that is presumed to equate to the first complex that assembles on a pre-mRNA under normal conditions [Michaud and Reed 1991], although it has only been shown to be a prerequisite for in vitro spliceosome assembly on one substrate [Jamison et al. 1992]. It contains U1 snRNP, weakly associated U2 snRNP, and other proteins, including SRSF1 [Staknis and Reed 1994; Makarov et al. 2012]. Importantly, it is committed to splicing [Michaud and Reed 1991; Jamison et al. 1992], and there is a functional interaction in assembly between the 5′ss and 3′ss [Michaud and Reed 1993]. Hydroxy radical probes tethered near either the 3′ss or the 5′ end of U2 snRNA were used to show that the 5′ss and 3′ss are in close proximity [Kent and MacMillan 2002; Donmez et al. 2007]. Interestingly, the addition of Drosophila SR-related proteins [Tra/Tra2] promoted a switch between alternative 5′ss of fruitless pre-mRNA even after complex E had formed, whereas it did not do so after assembly of the first ATP-dependent complex A [Kotlajich et al. 2009]. Hence, candidate 5′ss may still be being explored by the bound U1 snRNP. Complex E is therefore consistent with the model in which a single U1 snRNP is recruited by 3′ss components. However, this model is contradicted by experiments showing that alternative strong 5′ss can be protected concurrently against ribonuclease digestion [Nelson and Green 1988; Eperon et al. 1993] and, more clearly, by single-molecule experiments showing that two U1 snRNPs are
associated with most molecules of pre-mRNA containing two strong 5' ss in the absence of ATP, whereas only one is bound in complex A [Hodson et al. 2012]. These results suggest that U1 snRNPs can interact independently with the candidate 5' ss and that selection is associated with the dissociation of the surplus U1 snRNPs [Fig. 5C].

**Selection by affinity may require low levels of occupancy by U1 snRNPs**

The affinity of binding determines the fraction of molecules in which a particular 5' ss is bound by U1 snRNP, which is equivalent to the probability that a particular site on a molecule is occupied at any given time. If the affinities are so low that only one 5' ss (or none) is occupied on any given molecule of pre-mRNA at the time when selection takes place, then affinity-based selection would become merely a matter of selecting whichever U1 snRNP is present: The probabilities that the sites are occupied will determine the relative use of the possible 5' ss. Clearly, such a model would not work if more than one site were bound per molecule of pre-mRNA.

There are several indications of the likely range of lifetimes of U1:pre-mRNA complexes. The lifetime of a complex between pure U1 snRNP and a consensus 5' ss is ~10 min [Eperon et al. 2000]. If base-pairing limits the dissociation rate, then the lifetimes at more typical 5' ss would be shorter by two or three orders of magnitude. In vivo measurements suggest that the lifetime of bound U1 snRNP averages <1 sec [Huranova et al. 2010]. In active yeast extracts with a substrate containing a highly conserved 5' ss, the lifetime of the complex formed between uncommitted U1 snRNP and pre-mRNA was estimated by single-molecule methods at ~0.1 min [Hoskins et al. 2011]. Given that transcription of a mammalian intron would generally take at least a few minutes, if not hours, a situation approaching equilibrium between the candidate 5' ss and U1 snRNPs might be established before the point is reached at which the 5' ss is selected.

The process of selection itself is unknown. If the 5' ss and 3' ss have been brought into close proximity in complex

Figure 5. Models for the effects of affinity for U1 snRNP and 5' ss position upon selection. (A,B) Possible mechanisms by which affinity determines outcome. The candidate 5' ss have low affinity, and the level of occupancy is low such that there is no multiple occupancy and the distribution of U1 snRNPs reflects the affinities for the 5' ss. (A) Independent binding by U1 snRNPs. (B) Recruitment of a single U1 snRNP by components assembled, for example, at the 3' ss and its rapid interactions with the 5' ss result in a distribution of occupancy that reflects the affinities. (C) With high-affinity sites, binding is saturating and there is multiple occupancy consistent with the model in (A) in complex E conditions; during formation of complex A, the intron-proximal U1 snRNP is selected, and the distal snRNP is displaced [Hodson et al. 2012]. (D) The effects of varying the distance between two alternative 5' ss suggest that the preference for an intron-proximal site in C is not accounted for by 3D diffusion of free RNA. [Green triangles] Experimental data from Cunningham et al. (1991); [purple squares] outcome expected from simulation with substrate as free RNA (from a distribution of possible conformations of a homogeneous random coil modeled as a freely jointed chain); [black circles] outcomes expected if the sequence between the 5' ss is rigid. From Hodson et al. (2012), by permission of Oxford University Press. (E) Insertion of a non-RNA linker between an ESE and the alternative 5' ss blocks its effects, arguing against a looping model [Lewis et al. 2012].
E, but the 5′ss to be used is still negotiable, then selection must involve more than simply forming contacts between the U1 snRNP and 3′ss components. It is associated with formation of complex A and ATP hydrolysis (Kotajic et al. 2009) and may include the process by which surplus U1 snRNPs are irreversibly displaced if there are multiple strong 5′ss (Hodson et al. 2012). A candidate component for such a role is DDX46, a DEAD-box helicase that interacts with U1 and U2 snRNPs (Xu et al. 2004). Its yeast homolog, Prp5, is a spliceosomal protein that interacts with U1-A (Shao et al. 2012) and has been proposed to recruit U2 snRNP and incorporate it stably into spliceosomes after ATP hydrolysis (Kosowski et al. 2009).

Higher levels of occupancy are associated with selection by position
In addition to selection by affinity, there is an additional method of selection based on the relative positions of candidate 5′ss. This is manifested as a strong preference for the intron-proximal 5′ss if the pre-mRNA contains two or more 5′ss with high affinity for U1 snRNP (Reed and Maniatis 1986; Cunningham et al. 1991, Yu et al. 2008; Hicks et al. 2010) or the concentration of some SR proteins, such as SRSF1, is increased [see “Extrinsic Factors Affecting 5′ss Choices” above]. With high-affinity candidate 5′ss, multiple sites on a molecule of pre-mRNA are occupied by U1 snRNPs at the same time (Eperon et al. 1993, 2000, Hodson et al. 2012), presumably because the higher affinities increase the independent chance that each site is occupied and therefore increase the proportion of molecules on which multiple sites are occupied. The same principle applies to the effects of SRSF1, which increases the level of binding by U1 snRNPs at intron-distal and intron-proximal 5′ss (Eperon et al. 1993). The result in either case is an increase in the proportion of pre-mRNA molecules bound by U1 snRNPs at more than one site simultaneously, in which case the basic condition for affinity-dependent selection is broken and there is a switch to position-dependent selection. hnRNP A1 has the opposite effect on splicing, and the likely explanation is that it reduces the proportion of molecules bound by multiple U1 snRNPs (Eperon et al. 2000).

It is not known why the intron-proximal 5′ss is favored when multiple sites are occupied by U1 snRNP. Simple explanations based on the relative probabilities that each 5′ss and its bound U1 snRNP will encounter the 3′ss by three-dimensional (3D) diffusion of an RNA chain (Yu et al. 2008) do not account for the extent of the preference or its dependence on the distance between the alternative 5′ss (Fig. 5D; Cunningham et al. 1991; Hodson et al. 2012). Interestingly, a closer approximation to the observed behavior is seen if the exon sequence is treated as a rigid body (Hodson et al. 2012), which is not unreasonable given the high density of associated proteins in exons (Beyer et al. 1981). The binding of U1 snRNP might recruit SR proteins, which in turn may nucleate the binding of further proteins to purine-rich sequences and other motifs characteristic of exons.

Enhancer sequences may not act by looping
The question of the adequacy of 3D diffusion or looping models was mentioned above when discussing the ability of enhancer sequences placed between alternative 5′ss to stimulate use of the intron-proximal 5′ss. A model in which SR proteins bound to ESEs make contacts by looping with splice site components has been invoked to explain the stimulation of U2AF binding at 3′ss as well as U1 snRNP at 5′ss [Fig. 3B; Lavigueur et al. 1993; Tian and Maniatis 1993, Wu and Maniatis 1993, Staknis and Reed 1994, Wang et al. 1995]. However, it has been supported directly only by two observations: cross-linking of an exon-tethered RS domain to pre-mRNA at splice site regions (Shen and Green 2004, 2006) and the attenuation of stimulation by increasing distance from a target 3′ss [Graveley et al. 1998]. Recently, looping was directly tested using an ESE at the 5′ end of pre-mRNA that stimulated the use of an upstream alternative 5′ss. Insertion of a flexible alkyl or PEG linker between the ESE and the rest of the pre-mRNA abolished its action, which is inconsistent with looping and implies that the elements have to be connected by RNA [Fig. 5E; Lewis et al. 2012]. Therefore, it is possible that the enhancer affects protein binding along the RNA.

Implications for genetic diseases
We reviewed a wide range of mechanisms that dictate 5′ss selection and their implication in multiple types of alternative splicing events, which highlights the importance of research on 5′ss recognition. In addition, elucidating such mechanisms is highly relevant for human genetics. Around 10% of all disease-causing mutations affect either of the two splice sites [Krawczak et al. 2007], and this percentage increases to nearly 50% in the particular cases of NF1 and ATM genes, whose inactivation cause neurofibromatosis type 1 and ataxia telangiectasia, respectively (Teraoka et al. 1999; Ars et al. 2000). About half of such mutations affect 5′ss.

The two most important parameters of a splice site mutation are (1) the severity, which refers to the extent of reduction of correct splicing, and (2) the molecular consequence, which in humans can be, by order of frequency, skipping of the exon [in the case of internal exons], activation of cryptic splice sites, and intron retention. These two parameters often correlate with disease severity. Ab initio predictions of mutation severity by 5′ss scoring methods are largely accurate; i.e., the higher the difference in scores between the wild-type and mutant 5′ss, the more severe the mutation. Predicting the precise consequence of the mutation is far more difficult, although recent analyses have shown some progress [Wimmer et al. 2007; Divina et al. 2009].

The most deleterious mutations at a 5′ss are those affecting the nearly invariant GU dinucleotide, and this is reflected by the high frequency of mutations at these positions causing genetic diseases. For the remaining nine positions, the diagnosis can be more difficult, often relying on analyses of the mRNA from patients and of
minigenes in vitro or in cultured cells. Most methods take into account the extent of conservation of the affected nucleotide, which correlates with the severity of the disruption. The effects of certain mutations are more complex, as is the case of +3A-to-G transitions causing genetic diseases, even though both A and G occur with almost equal frequency at this position in bona fide 5’s. Such transitions are deleterious when the affected 5’s has nonconsensus nucleotides at the adjacent positions +4 and +5, as proven by pairwise associations in genomic data sets and experimental analyses (Ohno et al. 1999; Madsen et al. 2006; Roca et al. 2008). In certain cases, the 5’s scores do not reflect the effect of the mutation on 5’s strength. For example, a 5’s +5 A-to-G transition in the RARS2 gene causes pontocerebellar hypoplasia, and the deleterious effects of this mutation can be explained by considering the shifted 5’s/U1 base-pairing register (Roca and Krainer 2009). Finally, a smaller group of disease-causing mutations create new 5’s, termed de novo 5’s, which are selected instead of the natural 5’s. Recent reports investigated the particular sequence patterns surrounding cryptic and de novo 5’s, which help to understand the selection of such aberrant 5’s (Kralovicova and Vorechovsky 2007). Furthermore, promising approaches are being developed to rescue splicing defects, including 5’s mutations not affecting positions +1 or +2. Such technologies are based on antisense oligonucleotides or larger RNA molecules that can affect splicing (Hammond and Wood 2011). These few examples illustrate that a better understanding of the mechanisms of 5’s selection will likely improve the molecular diagnosis of 5’s mutations and facilitate therapeutics development.

Finally, single-nucleotide polymorphisms (SNPs) can affect splicing signals, and >1000 SNPs in the human genome map to bona fide human 5’s (Roca et al. 2008). Whereas such variations do not substantially change the strength and use of the 5’s, a fraction of them do affect splicing (Lu et al. 2012). Thus, progress in the field of 5’s selection will also help in the identification of SNPs that affect splicing with potential phenotypic consequences.

Future perspectives

This review highlights a number of unsolved questions about 5’s selection, some of which we reiterate here as a spur to future work. Even the apparently simple issue of 5’s base-pairing potential to U1 snRNA [a short helix with a maximum of 11 bp] is not fully understood, as this interaction is very flexible, allowing for different registers, bulged nucleotides, and perhaps other subtle yet important modifiers of 5’s strength. Also the limitations of nearest-neighbor parameters for modified nucleotides (Ψ) or bulges limits the inference of 5’s strength based on the free energy of base-pairing to U1. Taking into account positions +7 and +8 as well as other base-pairing registers should improve existing 5’s scoring tools. The contribution of proteins, such as U1C and hnRNP, to 5’s strength is also poorly understood. Likewise, U6 replaces U1 at the 5’s in active splicingosomes, but its contribution to 5’s selection has been shown in very few cases.

The whole question of how the existence and perhaps recognition of many candidate 5’s is turned into selection of just one is now an urgent issue for all areas of research into mammalian splicing. Processes such as transcription can restrict the number of accessible sites, as suggested in the “first come, first served” model (Kuhne et al. 1983), but the clear dependence of 5’s selection on affinity, context, and the concentrations of factors shows that potential alternative 5’s are exposed to recognition. The simplest explanation of affinity-based selection is that free U1 snRNPs are at first able to interact reversibly and independently with potential 5’s, but it is difficult to reconcile this with what we know of the ATP-independent complex E. Is complex E a subsequent state in which one site has been selected, but events dependent on ATP hydrolysis have not yet removed surplus snRNPs or locked the U1 snRNP onto a specific site? Is the state of free but weak binding shown in Figure 5A followed by the state shown in Figure 5B?

The state of the pre-mRNA is hard to model at present. We referred to several proteins that affect 5’s selection, mostly via interactions with U1 snRNP. However, very comprehensive experiments on exon inclusion involving analyses of sequence conservation (Goren et al. 2006; Friedman et al. 2008; Barash et al. 2010), selection from large pools (Ke et al. 2011), or systematic mutagenesis (Singh et al. 2004) indicate that most nucleotides in an exon could and in fact probably do affect splicing, presumably by influencing the binding of the 100 or so pre-mRNA-binding proteins. Does this mean that the exon (and perhaps flanking intron sequences) is smothered in proteins either before or after splice sites are selected? Unfortunately, much of the evidence for protein interactions depends on cross-linking and immunoprecipitation. While these are invaluable methods, they provide no indication as to whether all, some, or just a few pre-mRNAs in the reaction mixture are bound by a particular protein at a particular site. Moreover, they provide no information about stoichiometry. This requires single-molecule methods (Cherny et al. 2010). Hence, we do not yet know whether, in vivo or in vitro, U1 snRNP encounters accessible RNA or an RNA–protein complex that might limit binding. We described experiments that suggest that free RNA is an inappropriate model to account for the selection of intron-proximal strong sites or ESE-proximal sites. Do stably bound U1 snRNPs and ESEs trigger the propagation of proteins that alter the physical behavior of exons? Nothing at all is known yet about the flexibility and other physical properties of exons and introns, but we predict that they will behave very differently. Gross changes in physical properties might be one way of integrating the influences of numerous proteins and sites.

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Comparative analysis detects 140 GENES

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