“Nought May Endure but Mutability”: Spliceosome Dynamics and the Regulation of Splicing

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The spliceosome is both compositionally and conformationally dynamic. Each transition along the splicing pathway presents an opportunity for progression, pausing, or discard, allowing splice site choice to be regulated throughout both the assembly and catalytic phases of the reaction.

Introduction
A number of excellent reviews have been published over the past decade (including Staley and Guthrie, 1998; Burge et al., 1999; Cramer et al., 2001; Hastings and Krainer, 2001; Brow, 2002; Jurica and Moore, 2003; Nilsen, 2003; Stark and Lührmann, 2006; Valadkhan, 2007; Hertel, 2008; House and Lynch, 2008), addressing numerous aspects of pre-mRNA splicing and its coordination with other nuclear events. This review will focus on selected themes, highlighting the dynamic nature of both the assembly and catalytic phases. In the resulting view of the splicing reaction, unidirectional linear pathways describing progression are often insufficient, if not misleading. We aim to emphasize that multiple transitions in spliceosome assembly and catalysis can be modulated by alterations in the identity or activity of spliceosomal components or by tuning of the stability of interactions between the pre-mRNA substrate and the spliceosome. For a given intron and set of cellular conditions, one or more of these transitions will limit splicing and, thus, be available as a potential point of regulation. Changes in the efficiency of any transition in the splicing pathway can therefore result in regulated—and thus alternative—splicing.

The Dynamics of Spliceosome Assembly
The classical, sequential view of spliceosome assembly (reviewed in Burge et al., 1999) holds that the 5’ splice site (5’SS) is first bound by U1 snRNP, then the branch site (BS) and 3’SS by U2 snRNP and associated protein factors to form a prespliceosome, also known as complex A. The [U4/U6.U5] tri-snRNP joins the complex to form complex B, and a series of conformational and compositional changes result, including the loss of U1 and U4 snRNPs to leave U2/5/6. Recruitment of the CDC5L complex (known as the NTC in S. cerevisiae) follows (Makarov et al., 2002), giving rise to an activated spliceosome. Assembly can be stimulated or repressed by the binding of general or specific splicing factors to snRNPs and pre-mRNA. snRNPs can also interact both with pre-mRNA and with each other. Spliceosome assembly is, thus, highly cooperative, and the fact that many interactions can occur independently of one another results in an assembly cascade that does not follow a single obligatory trajectory but instead can occur via multiple pathways.

An extreme example of cooperative assembly is the penta-snRNP, which can be isolated from S. cerevisiae and contains all five snRNPs in the absence of pre-mRNA. Although the penta-snRNP alone is not competent to catalyze splicing, addition of nuclease-treated cell extract restores its activity without requiring disassembly and reassembly (Stevens et al., 2002). Together with the observation of splicing in the absence of U1 recruitment in multiple systems (reviewed in Burge et al., 1999), this finding suggests that there is unlikely to be any essential compositionally defined assembly intermediate prior to the activated spliceosome. Rather, a number of potential assembly pathways can lead to this same point. For a given substrate, the stabilities of intermediates along a given pathway might be such that this mode of assembly is observed to the exclusion of all others. In such cases, stable intermediates may be isolated (for example, the purification of complex A by Behzadnia et al., 2006), but this does not mean that the assembly pathway in question is an obligate one for any substrate.

Spliceosome assembly appears to be driven by the stochastic association of snRNPs with pre-mRNA. Data from photobleaching experiments using various GFP-tagged spliceosomal components are consistent with free diffusion of snRNPs within the nucleus and their transient random association with pre-mRNA (Rino et al., 2007 and references therein). Two major roles of factors stimulating or repressing assembly in such a system would be to increase or decrease the local concentration of snRNPs or near a transcript and to modulate the stability of snRNP pre-mRNA and inter-snRNP interactions. Given that the CTD of RNA polymerase II is one of the multiple regulators that can interact with snRNPs to modulate assembly (reviewed in Cramer et al., 2001), future studies must address how assembly is modulated in response to transcriptional events and chromatin structure. In addition, it is possible that future work will identify more entry points into the spliceosome assembly pathway: for example, “reinitiation” by postcatalytic U2/5/6 complexes.

Another convergence point during spliceosome assembly exists due to the two possible orientations of interaction between the U1 and U2 snRNPs. U1-U2 interaction can occur across the intron to be removed by the spliceosome that will ultimately contain these two snRNPs (intron definition). In multi-intron
genes, however, this interaction can also occur across an exon such that U1 bound at the 5’SS of the downstream intron interacts with U2 bound to the BS-3’SS of the upstream intron (exon definition). A reporter gene with long exons is spliced efficiently only if the flanking introns are short (Sterner et al., 1996), and increasing intron length in a model substrate favors exon definition (Fox-Walsh et al., 2005), consistent with the interpretation that simple binding kinetics determine the predominant assembly pathway for a given pre-mRNA substrate. It is not known whether multiple U1 snRNPs can interact with a single U2, or vice versa. If such polymeric interactions are not possible, inter-snRNP interactions promoting exon definition at a given splice site will preclude intron definition. An exon-defined assembly intermediate must therefore make the transition to an intron-defined state in order for functional splicing to proceed. Indeed, splicing can be inhibited by preventing the establishment of intron definition, for example, by hnRNP L binding to an exonic splicing silencer and stabilizing exon definition interactions between U1 and U2 (House and Lynch, 2006); hnRNP L has a stimulatory effect when bound within an intron, likely due to facilitation of a crossintron U1-U2 interaction (Hui et al., 2005). Similarly, the binding of SR proteins within an intron can inhibit splicing (Ibrahim et al., 2005), whereas SR binding within exons is generally stimulatory (reviewed in Hastings and Krainer, 2001). These observations are consistent with the existence of mutually exclusive interactions during exon- and intron-defined states.

Circular exons, the predicted products of splicing from an exon-defined state, have been detected in several systems (for example, see Bailleul, 1996). The formation of such products, even at a slow rate, suggests that the maintenance of U1-U2 interaction across an exon (i.e., with the wrong polarity) is not sufficient to prevent the formation of a catalytically competent spliceosome. It is therefore likely that a polarity-sensing mechanism normally exists to distinguish between exon- and intron-defined complexes. U1-U2 interactions, when intron-defined, may provide a binding surface for the [U4/U6.U5] tri-snRNP, with exon-defined complexes normally lacking such a surface. The recruitment of non-tri-snRNP proteins is also likely to play a role in this transition, with candidates for proteins involved in polarity sensing expected to bind only to intron-defined complexes. Although no mechanism or factors responsible for such polarity sensing are known, DEK, a chromatin-associated protein not required for early assembly but important for 3’SS definition and splicing catalysis (Soares et al., 2006), could represent one such factor. Proteomic analysis of human spliceosome assembly intermediates suggests that DEK joins the complex during or after the exon definition-intron definition transition (Sharma et al., 2008). In addition, there is no DEK homolog in S. cerevisiae, whose almost exclusively single-intron genes presumably lack an exon-defined stage.

The Role of RNA Structure

Consistent with many alternative splicing factors having a role in increasing or decreasing the local concentration of snRNPs on transcripts, a large number of sequence-specific RNA-binding proteins have been shown to modulate spliceosome assembly (for example Jensen et al., 2000). Several protein motifs that bind single stranded RNA have been characterized, and these are commonly found in splicing factors. Consistent with their action in a single-stranded state, a set of splicing enhancers and silencers has been confirmed bioinformatically to be more single stranded than bulk sequence, and to function more effectively when placed in the loop than the stem of a hairpin structure (Hiller et al., 2007). The formation and stabilization of secondary structure around such regulatory elements is therefore a potential mechanism to reduce their effects on splicing.

All evidence suggests that splice sites themselves must be single stranded in order to allow spliceosome assembly, with secondary structure inhibiting U1 and U2 snRNP binding. Inclusion of the 3’SS in a hairpin is inhibitory for splicing, although this can be overcome by the presence of a single stranded “helper” downstream of 3’SS, likely recognized during assembly (Liu et al., 1995). A particularly elegant example of alternative splicing regulation by direct modulation of secondary structure around splice sites is the control of alternative splicing by a thiamine pyrophosphate (TPP) riboswitch in Neurospora crassa (Cheah et al., 2007). When TPP concentration is low, the pre-mRNA adopts a structure such that an otherwise favored downstream 5’SS is occluded, and the branch region is flexible. Splicing proceeds using a suboptimal upstream 5’SS to produce mRNA encoding a functional NMT1 protein. When TPP concentration is high, however, conformational changes in the riboswitch cause structure around the favored downstream 5’SS to be disrupted, leading to the predominant production of a longer mRNA containing uORFs that prevent NMT1 translation. In addition, the branch region is partially occluded, yielding an overall decrease in splicing efficiency. It is likely that similar examples of splicing regulation, mediated by proteins or small molecules, will be discovered in other systems: how common such mechanisms of splicing regulation will prove to be remains an open question.

Secondary structure is not always inhibitory to splicing. For example, the S. cerevisiae RP51B intron contains complementary sequences close to the 5’SS and BS that bring the ends of the intron together and aid spliceosome assembly (Charpentier and Rosbash, 1996), and it is possible that this is a common way to increase the efficiency of U1-U2 binding and intron definition. The splicing of exon clusters 4 and 6 in the well-characterized Dscam gene in Drosophila provides two further examples of stimulatory secondary structures. Disruption of the iStem—a large hairpin loop downstream of exon 3—interferes with the splicing of all twelve exon 4 variants (Kreahling and Gravelay, 2005), although the mechanism by which this stem stimulates exon 4 splicing remains unclear. The basis of maintenance of mutually exclusive splicing in the exon 6 cluster, however, is better understood. Each exon 6 variant is preceded by a selector sequence complementary to a docking site downstream of exon 5. Interaction between a given selector sequence and the docking site leads to splicing of the following exon and, as the docking site is thus removed from the transcript, the inclusion of further exon 6 variants is suppressed under normal conditions (Gravelay, 2005). Knockdown of the hnRNP protein hpr36 leads to the inclusion of multiple exon 6 variants, suggesting that this protein mediates the repression of splicing across the cluster (Olson et al., 2007).

hnRNP proteins normally act as general inhibitors of splicing; they are antagonized by the generally activating SR proteins
There is increasing evidence that SR proteins exert at least some of their stimulatory effect via the stabilization of RNA-RNA interactions during both spliceosome assembly and splicing catalysis. A pre-mRNA with a 5′ exon as short as one nucleotide can undergo SR protein-dependent splicing in HeLa extract, suggesting a postassembly role for these proteins (Hertel and Maniatis, 1999). The arginine-serine rich (RS) domain of a natural SR protein or an artificial domain comprising multiple RS repeats, when tethered to pre-mRNA, directly contacts the branch site and facilitates pre-spliceosome formation (Shen et al., 2004). It is thought that the BS is already base paired to U2 snRNA in such assembly intermediates (Xu and Query, 2007). Defects due to SR protein deple

tion can be suppressed by increasing the strength of the interaction between the 5′SS and U6 snRNA that is required for the first catalytic step (Kim and Abelson, 1996; Shen and Green, 2006). This argues for the involvement of enhancer-recruited SR proteins not only in assembly, but also during the catalytic phase of splicing.

Data concerning the role of SR proteins in alternative splicing have generally been interpreted with assembly in mind; such data might now need to be reconsidered, as the stabilization of duplexes could produce diverse phenotypes during the dynamic and structurally complex catalytic phase. It is also possible that hnRNP proteins might exert some of their effect via the disruption of RNA-RNA interactions, most simply by sequence sequestration. The mechanistic basis for duplex stabilization by SR proteins remains unclear, as does the issue of whether this stabilization is general or protein-duplex specific, with defined SR proteins stabilizing only certain duplexes.

The Transition between the Two Chemical Steps

Splicing catalysis consists of two successive transesterification reactions: in the first step, the 2′ hydroxyl of the BS nucleotide nucleophilically attacks the 5′SS to yield a lariat intermediate and a free 5′ exon and, in the second step, this free exon nucleophilically attacks the 3′SS, producing mRNA and an excised lariat intron (Figure 1). The 3′SS remains sensitive to nuclease degradation until after the first step. This suggests that it enters the active site after first step catalysis (Schwer and Guthrie, 1992), a repositioning event that would require removal of the newly formed branch structure of the lariat intermediate from the catalytic center. Indeed, multiple lines of evidence suggest that the 3′SS replaces the branch structure, with the 5′ exon remaining in a fixed position relative to loop 1 of U5 snRNA. Crosslinks between U5 loop 1 and the terminal nucleotide of the 5′ exon can be chased through both steps of splicing. Those between loop 1 and position +2 of the intron, however, can be chased into a lariat near wild-type stability (Konarska et al., 2006). The splicing defect due to a hyperstabilized 5′SS-U6 helix can be suppressed not only directly by duplex destabilization, but also by the U6 U57A mutation, a variety of mutations in Prp8 (Konarska et al., 2006) or deletion of the NTC component Isy1 (Villa and Guthrie, 2005). Mutations that suppress the second step defect of an A3C intron also increase the efficiency of the second step for a variety of other intron mutations, including those at the branch site adenosine and 3′SS. Commensurate with their stimulation of the second step, these suppressors inhibit the first step of splicing. There also exists the opposite class of spliceosomal mutants: those that increase the efficiency of the first step at the expense of the second (Query and Konarska, 2004). The existence of two opposing classes of suppressor allele, each capable of suppressing a wide range of intron mutations, suggests that suppression is not necessarily via direct
reaction) is shown.

The combination of mutations in Prp8, a large and essential protein involved in splicing, has led to a refinement of the two-state model of splicing (Figure 1). First and second step suppressor point mutations in Prp8 have led to a two-state model of splicing where the two steps are in competition with one another. Mutations that stabilize the first step conformation relative to the second will stimulate the first step and inhibit the second, while the opposite will be true for those that cause relative stabilization of the second step conformation. A mutation that is more likely to disrupt an interaction than to form a new one, it is likely that relative stabilization takes the form of destabilization of the competing conformation, such that most first step suppressors would destabilize the second step conformation and vice versa. At present, however, the molecular basis of the action of these general suppressor mutations is unknown. This two-state model provides a mechanism by which modulation of the stabilities of conformational states of the catalytic spliceosome can impact the efficiency of splicing of suboptimal substrates. Such tolerance of suboptimal splice sites is manifested in metazoan alternative splicing, meaning that local or global modulation of conformational stability during catalysis could impact the splicing pattern of individual transcripts or of classes thereof, respectively.

Recent detailed analysis of the interplay between global suppressor mutations has led to a refinement of the two-state model (Figure 1). First and second step suppressor point mutations combined in the same molecule of Prp8, a large and exceptionally well-conserved U5 snRNP protein that makes contacts with the 5′SS, BS, and 3′SS (reviewed in Grainger and Beggs, 2005), cancel one another’s effects and produce a phenotype resembling wild-type Prp8. A different effect is observed, however, when Prp8 second step suppressor mutations are combined with first step suppressors in U6 snRNA or Prp16, the ATPase that modulates the first-to-second step transition (Burgess and Guthrie, 1993). In this instance, the suppressors act in concert such that both the first and second steps are improved. Cancellation by the opposing classes of prp8 allele indicates that these prp8 alleles act at the same kinetic step as one another, whereas the additive nature of the Prp16/U6-Prp8 suppressor pairs requires that they be affecting different kinetic steps in the transition. These observations have led to a model in which the transition between the two steps has multiple phases, requiring an “opening” step (affected by prp16 and U6 mutants) and a repositioning step (affected by prp8 mutants), followed by “closure” into the second step conformation (Liu et al., 2007).

The suggestion that transitions traditionally considered as one step actually comprise multiple phases has important implications for future proteomic, biochemical, and structural studies of the spliceosome. A complex containing a lariat intermediate and a free 5′ exon could plausibly be in one of at least four conformations that are currently compositionally and conformationally undefined: post-first step but preopening, open but unpositioned, open and repositioned, or closed pre-second step. In the first or fourth case the purified complex would be competent to carry out first or second step catalysis, respectively, whereas the open complexes are presumably not catalytically competent. Accurately ascertaining the state of purified complexes is therefore essential in order to allow coherent and reliable insights into the mechanism of splicing. As will be discussed below, there is an emerging view that multiple transitions along the splicing pathway resemble one another, so it is possible that the transition between the catalytic steps is not the only one composed of several smaller remodeling events.

**Conformational Toggling, Asymmetry, and the Reuse of Motifs**

Alleles of prp22, the ATPase involved in transitions during and after the second step of splicing, produce a cold-sensitive phenotype due to an mRNP release defect. A screen for suppressors of this phenotype identified a prp22 allele (Schneider et al., 2004) subsequently shown to also act as a general suppressor of first step splicing defects (Liu et al., 2007). Indeed, all known first step suppressor alleles of prp8 suppress prp22 defects. This observation is consistent with the hypothesis that these alleles destabilize the second step conformation of the spliceosome, thereby stimulating the surrounding steps on the splicing pathway (first step catalysis and mRNP release), but could also suggest some degree of similarity between these flanking states. Consistent with the existence of such similarity, a growing number of interactions appear to be disrupted and reform at defined stages of the splicing reaction—i.e., to toggle.

Many RNA-RNA interactions between snRNAs, as well as between snRNAs and pre-mRNA, have been identified and take the form of generally short intra- and intermolecular helices (reviewed in Brow, 2002). Some of these interactions are mutually exclusive with others, which suggests that they might exist only transiently or may toggle between competing conformations. As previously noted, during the first catalytic step the UGU trinucleotide at positions 4–6 of the 5′SS base pairs with a conserved ACA in U6 snRNA (positions 47–49) (Kim and Abelson, 1996). During the second step, when the 5′SS is in the branch of the lariat intermediate, it is in proximity to U6 positions...
42–44 (Sawa and Abelson, 1992). Interestingly, this region of U6 has also been shown to bind the 5’SS in early complexes (Johnson and Abelson, 2001; Chan et al., 2003). Thus, the same binding site may be used for the 5’SS before and after its involvement in first step catalysis. Indeed, spliceosomal states even further apart on the splicing pathway display surprising similarities: the ATPase Br2 disrupts the interaction between U4 and U6 snRNAs during spliceosome assembly, allowing U6 to interact with U2 and the 5’SS (Raghunathan and Guthrie, 1998). It has recently been demonstrated that Br2 is activated by the GTP-bound form of the U5 snRNP protein Snu114 and repressed by its GDP-bound form and that Br2 activation is required for spliceosome disassembly as well as U4/U6 unwinding (Small et al., 2006). GTP hydrolysis by Snu114 is not required for Br2 activation, suggesting a mechanism of action resembling that of classical G proteins. Although the GAP and GEF acting on Snu114 have not been identified, it is tempting to speculate that the relevant conformational states of the spliceosome might perform these roles, akin to the action of the signal recognition particle itself as the GEF for SR-β (Helmers et al., 2003).

Recent work provides another example of conformational toggling (Hilliker et al., 2007; Perriman and Ares, 2007). The dynamic stem II region of U2 snRNA can form two mutually exclusive interactions, known as stem IIa and stem IIc. Stabilization of stem II region of U2 snRNA can form two mutually exclusive interactions, known as stem IIa and stem IIc. Stabilization of stem IIa (and therefore relative IIc stabilization) suppresses second step splicing defects. These data are consistent with a model in which stems IIa and IIc toggle, coexisting with the previously discussed open and closed forms of the spliceosome, respectively, with stem IIc therefore present during catalysis and IIa during repositioning (Figure 1).

Although more examples of conformational toggling will likely be identified, many spliceosomal interactions are unlikely to reoccur once disrupted: for example, the Prp28-mediated replacement of U1 by U6 at the 5’SS (Staley and Guthrie, 1999). In fact, a general theme in intermolecular interactions involving U6 snRNA is that of asymmetry. U6 mutations that disrupt a structure often have a more severe phenotype than corresponding mutations in the interacting partner, and incomplete suppression by compensatory changes is common. For example, mutations on the U6 side of U2/U6 helix la are substantially more severe than those on the U2 side (Madhani and Guthrie, 1992). This suggests that helix la does not remain intact throughout the entire splicing reaction and that the interactions of the U6 component, when not engaged in this helix, are more critical for splicing than those of the U2 component. Similar asymmetry is observed for the conserved AGC triad of U6, which can interact with sequences in U4 snRNA, with U2 snRNA (to form helix Ib), and within U6 to extend the intramolecular stem loop (ISL) (reviewed in Brow, 2002). Most mutations in AGC are viable if accompanied by a compensatory mutation in U2 that restores helix Ib, but some substitutions, such as G60Y, cannot be suppressed in this manner (Hilliker and Staley, 2004). An identical AGC motif in domain 5 of group II self-splicing introns acts as a metal binding site crucial for catalysis (reviewed in Pyle, 2008) and, as noted by Hilliker and Staley, spliceosomal AGC mutations that cannot be suppressed by U2 are expected to interfere with metal binding. A second metal bound in domain 5 of group II introns is thought to mediate a docking interaction (reviewed in Pyle, 2008), and again, an analogous metal-binding motif exists in the ISL of U6. U80 (S. cerevisiae numbering) is bulged from the ISL and binds magnesium (Yean et al., 2000). The formation of this U80 bulge occurs after U4/U6 unpairing (reviewed in Brow, 2002), and suppression data indicate the delicate balance of relative stabilities required to allow both structures to form and, thus, permit splicing (McManus et al., 2007). However, the importance of specifying not only the nature but also the precise timing of interactions within the spliceosome is illustrated by the complex behavior of this nucleotide. When substituted by 4-thio-uridine, U80 forms a site-specific crosslink with a nucleotide well upstream of the branch site of actin pre-mRNA (Ryan et al., 2004). In addition, an Fe-BABE group tethered at the +10 position of the 5’SS stimulates cleavage at the human equivalent of U80 (U74) (Rhode et al., 2006). Although it is possible that these biochemical data correspond to off-pathway intermediates, it is also plausible that they indicate that at least some of the groups responsible for catalysis are sequestered by other interactions until the immediate precatalytic state of the spliceosome.

Spliceosome conformations can also be affected by transient protein modifications. For example, in addition to the known effects of the phosphorylation state of SR and other proteins on spliceosome assembly, Snu114 and the U2 snRNP protein SF3b155 appear to be dephosphorylated for the second step of splicing (Shi et al., 2006 and references therein). Similar effects are likely to be uncovered for many splicing factors, and indeed Prp8 ubiquitylation has recently been shown to affect spliceosome assembly (Bellare et al., 2008).

**ATPases and Fidelity**

Although the short duplexes involved in spliceosome assembly and catalysis may be able to unwind naturally to facilitate conformational changes, perhaps in concert with other remodeling events, DExD/H ATPases represent a major class of spliceosomal proteins (reviewed in Cordin et al., 2006). Many DExD/H ATPases have been shown to have RNA unwindase activity correlating with ATPase activity in vitro. Cyt19, a DExD/H protein that promotes group I intron splicing in vivo and in vitro, does so by acting as a nonspecific chaperone, resolving kinetic traps along the folding pathway (Mohr et al., 2006). Similarly, splicing of the ai5 group II intron in S. cerevisiae is stimulated by Ms116, and splicing activity in vitro correlates with Ms116 ATPase and unwinding activity (Del Campo et al., 2007). However, the same study reported a residual, ATPase-independent unwinding activity for Ms116, and recent data suggest that Dbp5, which is involved in mRNA export, functions to remodel RNPs only in its ADP-bound form, with ATP hydrolysis thus acting as a conformational switch rather than a power stroke (Tran et al., 2007). It therefore remains unclear whether ATPase/unwindase activity is the mechanistic basis of all, or indeed any, spliceosomal activity of DExD/H ATPases.

Most spliceosomal ATPases are currently thought to facilitate a single transition along the splicing pathway, although the
example of Brr2 demonstrates the possibility of a single ATPase acting multiple times (Figure 3). The question of how ATPase activity is limited to the correct stage(s) remains an open one. Binding of one ATPase to the spliceosome is not necessarily mutually exclusive with the presence of others, as illustrated by the persistent presence of Prp43 from early complexes until disassembly (reviewed in Jurica and Moore, 2003), during which time many other ATPases act. It is, however, possible that the binding sites for some ATPases share common elements, such that mutually exclusive subsets exist. A requirement for cofactors to stimulate ATPase activity is one mechanism by which activity could be temporally regulated: the helicase activity of Prp43 is stimulated by Ntr1, and this stimulation is required for Prp43’s role in spliceosome disassembly (Tanaka et al., 2007). The recruitment of a cofactor or, in the event of an ATPase interacting with multiple spliceosomal components, the conformation of the spliceosome itself could, therefore, activate an individual ATPase among several simultaneously bound and repress others such that inappropriate conformational changes are not induced.

By stimulating conformational transitions within the spliceosome, DExD/H ATPases play an integral role in the maintenance of splicing fidelity. Prp16, which as previously noted facilitates opening following the first step, was isolated as a suppressor of a branch site mutation in S. cerevisiae, and the isolation of the requirement for adenosine at the branch site was subsequently shown to be due to ATPase impairment, resulting in the kinetic proofreading model of splicing fidelity (Burgess and Guthrie, 1993). According to this model, functional progression into the second step occurs if catalysis precedes Prp16 ATP hydrolysis, whereas substrates discard results if catalysis has not occurred before ATP hydrolysis. ATPase-deficient prp16 alleles reduce fidelity and suppress splicing defects by allowing more time for catalysis to occur, resulting in the progression of suboptimal substrates, which would otherwise be discarded, through the first step.

An important prediction of the kinetic proofreading model is that each ATPase-mediated conformational change affords opportunity for such a progression/discard branch in the splicing pathway; recent work has indeed demonstrated analogous behavior for two more spliceosomal ATPases, consistent with the generality of this mechanism. S. cerevisiae spliceosomes assembled on 3’ SS mutant substrates and purified after the first step proceed through the second in the presence of mutant Prp22 ATPase deficient for ATPase and/or unwindase activity, or in the absence of ATP (Figure 3, Mayas et al., 2006). Genetic work in S. cerevisiae has also shown kinetic proofreading of BS-U2 snRNA interaction by Prp5 (Figure 3). ATPase-deficient prp5 alleles suppress mutations flanking the BS that destabilize its pairing to U2, but suppression can be superseded by restabilizing this interaction, either by compensatory mutations in U2 or intron mutations that generate extra upstream base pairs. The level of suppression by prp5 alleles correlates inversely with their ATPase activity. Prp5 proteins from organisms in which the branch site is less highly conserved than in S. cerevisiae have lower ATPase activity, thus providing a mechanism by which the fidelity of branch site selection is reduced in these organisms (Xu and Query, 2007). This work identified the first structure to be in direct competition

Figure 3. NTPase-Associated Steps during Splicing Offer Opportunities for Kinetic Discrimination of Suboptimal Pre-mRNA Substrates

Upper: schematic of transitions facilitated by DExD/H-box ATPases and the Snu114 GTPase during pre-mRNA splicing. SS, splice site; BS, branch site. Lower: characterized examples of kinetic proofreading mediated by spliceosomal ATPases. Left: altered competition between BS-U2 pairing and the conformational change mediated by the Prp5 ATPase changes the fidelity of BS selection. Center: altered competition between the first catalytic step and Prp16 ATPase activity affects the fidelity of splice site usage in this step. Right: altered competition between the second catalytic step and Prp22 ATPase affects second step splice site fidelity.
with the activity of a spliceosomal DExD/H ATPase, but the precise molecular consequences of ATPase activity remain to be elucidated for this and other ATPase-mediated transitions.

Important mechanistic questions regarding kinetic proof-reading remain to be addressed. For example, the direct targets of spliceosomal ATPases are unknown. In addition, although each ATPase-mediated step is an opportunity for discard, the nature of this discard is enigmatic. For example, it is possible that the conformational change resulting from ATPase activity is not compatible with binding of the substrate for the previous step, such that it would cause the spliceosome to fall apart. Alternatively, an active disassembly cascade could be triggered by such a conformational change. It is even possible that the spliceosome may need to undergo several conformational changes resembling functional progression along the splicing pathway. A role in discard for Prp43 and Ntr1, which cooperate in spliceosome disassembly (Pandit et al., 2006), might suggest that discard is mechanistically similar to progression. Some support for this model is provided by the observation that discarded intermediates are degraded in the cytoplasm (Hilleren and Parker, 2003). This finding implies that discard, like mRNA release, is coupled to nuclear export.

The Second Catalytic Step

The second step of splicing remains substantially less well characterized than the first. In addition to early recognition at the stage of complex A formation, the 3' SS has been proposed to be selected after the first step via a simple scanning mechanism as the first AG dinucleotide, or the second if sufficiently close to the first AG, downstream of the branch site (yeast) or polypyrimidine tract (metazoa) (Smith et al., 1993; Anderson and Moore, 1997; Chua and Reed, 2001). Distance from the branch site is an important determinant of 3' SS strength: Prp22, Slu7, and Prp18 are all dispensable in vitro for introns with short BS-3' SS distances (Schwer and Gross, 1998; references therein), and Prp22 mutants stimulate the use of non-AG splice sites closer to the branch (Mayas et al., 2006). In addition, the splicing of genes with short BS-3' SS spacing is unaffected by knockdown of the second step factor Prp17 in S. cerevisiae (Sapra et al., 2004). Aside from proximity to the branch site or polypyrimidine tract, what constitutes a favored 3' SS remains unclear. The YAG 3' SS consensus sequence has not been extended by bioinformatic investigation, although it seems possible that local RNA structure may play a role in determining the quality of a 3' SS for the second step. Such bioinformatics, along with genetic screens to search for possible determinants of YAG strength, may reveal higher complexity.

Experimental investigation of the second step is hindered by several factors: the apparent ability of the spliceosome to assemble on one 3' SS and use another on the same pre-mRNA is one such hindrance. Previously noted experiments in plants demonstrating that a 3' SS sequestered in a hairpin could act as a splice acceptor only in the presence of a downstream 3' SS suggested the possibility of specifying the 3' SS after early assembly (Liu et al., 1995). Work on autoregulatory splicing by sex-lethal in Drosophila has provided evidence that a 3' SS can play a critical role in exon definition, but not be preferred for catalysis (Penalva et al., 2001). It is therefore possible that 3' SS requirements for assembly and catalysis are, from an experimental point of view, at best not necessarily identical and at worst obligately distinct. An "ideal" 3' SS could thus represent a balance between assembly-competent and catalytically favored states. This and potential effects due to the almost unavoidable presence of nearby AG dinucleotides in natural genes must be taken into account in any systematic analysis of 3' SS quality.

A second obstacle to the investigation of the second step is that any active site component required for both steps will presumably exert its effects at the stage of first step catalysis, thus rendering the investigation of its role in the second step technically difficult. Although many spliceosomal components involved in the second step are dispensable for the first, such as the 3' SS itself (Chiara and Reed, 1995), Slu7 (Chua and Reed, 1999), and loop 1 of U5 snRNA (O'Keefe et al., 1996), several shared active site components might exist. Indirect information about the components of the second step active site could be derived from knowledge about the second step binding site of the lariat branch. The branch structure of the lariat intermediate must be repositioned and bound during the second step, consistent with the second step defect of 5' SS and BS mutants being suppressed by spliceosomal alleles that improve the second step (Query and Konarska, 2004; Konarska et al., 2006; Mayas et al., 2006). Although the 5' SS portion of the branch structure appears to reposition relative to U6 as previously discussed, the nature of this interaction as well as the interacting partner of the branch site region, which appears to unpair from U2 snRNA during or after first step catalysis (Smith et al., 2007), are unknown.

In fact, temporal epistasis, that is, the manifestation of mutations that cause multiple sequential defects in a pathway only at their earliest point of action, can also impede the study of first step catalysis due to the long preceding assembly phase. Systems in which assembly can be bypassed, at least in part (for example Konforti and Konarska, 1994; Anderson and Moore, 1997; Mayas et al., 2006; Valadkhan et al., 2007), represent possible ways to circumvent this problem.

Many proteins join the spliceosome between the two catalytic steps (reviewed in Jurica and Moore, 2003). In vitro depletion/reconstitution studies, together with genetic work in vivo, have provided clues as to the function of many second step factors. Loop 1 of U5 snRNA appears to act as a "platform" on which the 5' and 3' exons are juxtaposed for ligation (Crotti et al., 2007) and interacts functionally with Prp18, a protein that also binds Slu7 (Backova and Horowitz, 2002). Slu7 depletion leads to a loss of 3' SS fidelity and also appears to destabilize free 5' exon binding to the pre-second step spliceosome (Chua and Reed, 1999). Prp22 crosslinks directly to the 3' SS following the action of Prp16 (McPheeters et al., 2000) and is also involved in mRNA release (Schwer and Gross, 1998; Wagner et al., 1998). Interestingly, although all of these factors are essential in vivo, the requirement for each can be at least partially bypassed in vitro (Schwer and Gross, 1998; Chua and Reed, 1999; Segault et al., 1999; Crotti et al., 2007), suggesting both that the second step spliceosome is fairly robust and that these factors do not form interactions strictly necessary for catalytic events. Instead, a likely possibility is that they all contribute to stabilization of the second step conformation relative to the first. Elucidation of the interactions made by these factors will be
necessary to understand how they can impact 3′SS use and, therefore, alternative splicing.

Alternative Splicing

In the absence of repression, strong splice sites give rise to constitutive splicing. Alternative splicing, therefore, represents the suppression of optimal splice sites and/or the use of those that are suboptimal. Most alternatively spliced introns are thought to be controlled by multiple splicing enhancer and silencer elements whose activity depends on their location relative to splice sites. These regulatory elements are thought to affect splicing predominantly through corresponding RNA-binding proteins that exert their effects by altering a specific step in the spliceosome assembly pathway (reviewed in Black, 2003). However, modulation of splicing efficiency is theoretically possible at any stage of the splicing reaction. Each splicing event, depending on the introns and exons involved, will be limited by one or more transitions and will, as such, be sensitive to modulation of their efficiency while being insensitive to all but the most major changes in the efficiency of nonlimiting steps. Given the diversity of pre-mRNA substrates, the multistep nature of the spliceosome assembly and catalysis pathway, and the enormous number of factors involved in the splicing of every transcript, it is almost certain that examples exist in nature of splicing regulation at every possible stage. The existence of natural splicing events at least partially limited by postassembly transitions has already been demonstrated. The splicing of overlapping but nonidentical sets of endogenous introns is sensitive to the knockdown or mutation of core spliceosomal proteins important for the catalytic phase of the reaction (Park et al., 2004; Pleiss et al., 2007); this sensitivity affords opportunities for splicing regulation during catalysis. The ability to regulate the splicing of an individual intron by modulating the local activity or concentration of a given protein, or that of a class of introns via more global changes, together with the combinatorial effects of regulation of multiple transitions, can allow a robust and specific regulation of splicing events without a necessary requirement for large numbers of individual splicing factors to exert this control. Thus, in order to understand alternative splicing, specific factors acting on each transcript need not necessarily be sought, and the entire reaction through assembly, catalysis, and disassembly must be considered.

The stimulation or repression of spliceosome assembly naturally represents a common mechanism of splicing regulation and may indeed be the most prevalent in higher eukaryotes. Such assembly-based regulation might, however, be more subtle and complex than commonly thought. Virtually all metazoan transcripts contain multiple sequences capable of being recognized and used as splice sites, and much silencer-based repression may therefore require kinetic competition between these sites, serving predominantly to redirect rather than strictly to repress spliceosome assembly and/or catalysis.

Alternative splicing is not normally considered to occur in S. cerevisiae, but this organism’s ease of genetic manipulation and strong splice site consensus requirements allow many different events in splicing to artificially be made limiting for gene expression, facilitating detailed mechanistic analysis of suboptimal splice site use. Recent data showing that the splicing of meiosis-specific genes is repressed outside meiosis (Juneau et al., 2007) also potentially identify a system in which true alternative splicing can be studied in this organism.

Concluding Remarks

Enormous amounts of data exist regarding regulated and cell-type-specific splicing patterns (reviewed in Moore and Silver, 2007). Specific splicing factors that affect the splicing of small numbers of transcripts must exert their effects via spliceosomal transitions and through core components, both of which are finite in number. This imposes a limit on the number of possible unique mechanisms of splicing regulation. In addition to the discovery of more specific splicing factors, we anticipate that more widespread and varied regulatory roles will be discovered for core spliceosomal components themselves. A clear, mechanistic description of the splicing process is necessary to explain regulated splicing, and the detailed analysis of various alternative splicing systems is also likely to identify additional transitions in the splicing pathway.

Much work in the mechanistic splicing field is justifiably focused on the generation of spliceosome preparations suitable for X-ray crystallography. The high-resolution structures that will hopefully result from this work should resolve many of the questions posed in this review, and provide invaluable information about the mechanistic details of the splicing reaction. The preparation of large quantities of sufficiently pure, conformationally homogeneous spliceosomes, however, remains a serious challenge. Further and more detailed knowledge of the dynamic behavior of the spliceosome will aid such crystallographic efforts and will also be necessary for the rationalization and accurate interpretation of their results.

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