Stages in the second reaction of pre-mRNA splicing: the final step is ATP independent

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We have analyzed pre-mRNA splicing in yeast extracts immunodepleted of the PRP18 protein. We find that while the first step of splicing (cleavage at the 5' splice site, and generation of the exon 1 and lariat intermediates) is unaffected by the absence of PRP18, the second step of splicing (excision of the lariat intron and formation of mRNA) is substantially slower in the absence of PRP18. The splicing intermediates that are formed in the absence of PRP18 can be rapidly chased into products by the addition of purified PRP18 protein. This chasing is not dependent on ATP, implying that ATP is not required during the second cleavage-and-ligation reaction. This result suggests that there are ordered stages within the second step of splicing and that PRP18 acts late in the second step, perhaps during the catalytic step. The ATP independence also supports the idea that this reaction proceeds by a transesterification mechanism.

[Key Words: Pre-mRNA splicing; U5 snRNP; PRP18; Saccharomyces cerevisiae]

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The splicing of pre-mRNA takes place on a large ribonucleoprotein particle called the spliceosome, which consists of five small nuclear RNAs (snRNAs), U1, U2, U4, U5, and U6, and a large number of proteins [for review, see Green 1991; Guthrie 1991; Ruby and Abelson 1991]. Splicing of pre-mRNA is a two-step process. In the first step, the pre-mRNA is cleaved at the 5' splice site, releasing the first exon; concomitantly, the 5' phosphoryl group at the end of the intron is ligated to the 2' hydroxyl of an internal adenosine, yielding a branched RNA termed a lariat. In the second step, cleavage at the 3' splice site is accompanied by ligation of the two exons, yielding the product mRNA and releasing the intron as a lariat. These two steps are formally the same as those that occur in the self-splicing of group II introns (Peebles et al. 1986; van der Veen et al. 1986). From the analogy with group II self-splicing, each of the two steps of pre-mRNA splicing has been hypothesized to be a transesterification reaction, in which the number of phosphodiester bonds is conserved (Cech 1986; Jacquier 1990). The analogy also suggests that RNA may play an important role in the catalysis of the splicing reaction. In contrast to self-splicing, ATP is required for the splicing of pre-mRNA. It has been hypothesized that ATP hydrolysis is involved in proofreading or in conformational changes within the spliceosome. In the splicing of pre-tRNA, ATP hydrolysis is coupled to the ligation of the two exons, but it has not been determined whether ATP might have a similar role in pre-mRNA splicing.

In this paper we focus on the second step of splicing in Saccharomyces cerevisiae. At least four proteins, called PRP16, PRP17, PRP18 (for pre-mRNA processing), and SLU7 (for synergistically lethal with U5) [for synergistically lethal with U5], appear to be required for the second step. PRP16 was found as a second site suppressor of a mutation in the invariant TAC-TAAC sequence located at the branchpoint in all S. cerevisiae introns (Couto et al. 1987). PRP16 is an RNA-dependent ATPase (Burgess et al. 1990; Schwer and Guthrie 1991), which appears to mediate a conformational change in the spliceosome preceding the second reaction of splicing (Schwer and Guthrie 1991), which appears to mediate a conformational change in the spliceosome preceding the second reaction of splicing (Schwer and Guthrie 1992). PRP17 was found in a screen for yeast strains that were temperature sensitive for splicing (Schwer and Guthrie 1992), and was found again as a synthetic lethal with U5 [SLU4] [Frank et al. 1992], but has not yet been further characterized. PRP18, the focus of this study, was found in the same screen that yielded PRP17 [Vijayraghavan et al. 1989]. Initial work showed that PRP18 was involved in the second step of splicing [Vijayraghavan and Abelson 1990], and we have recently found that PRP18 is associated with the U5 snRNP [D. Horowitz and J. Abelson, in prep.]. SLU7 is a synthetic lethal with some mutations in the U5 snRNA [Frank et al. 1992] and appears to be involved in the recognition of the 3' splice site [Frank and Guthrie 1992]. Two snRNAs, U2 and U6, have been shown to be directly involved in the second step of splicing. Fabrizio and Abelson (1990) analyzed the effects of a large number

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of mutations in the U6 snRNA in vitro; McPheeters and Abelson (1992) carried out a similar analysis of U2. In both cases, a small number of the mutations analyzed resulted in the accumulation of intermediates in the splicing reaction, showing that both snRNAs are important for the second step. A role for the U1 and U5 snRNAs in 3' splice site recognition has been inferred from genetic experiments. Reich et al. (1992) have suggested that U1 base-pairs to the conserved AG at the 3' splice site, whereas Newman and Norman (1992) have found evidence that U5 base-pairs to the first two nucleotides in exon 2. Whether these base-pairing interactions are important during the actual splicing reaction is unknown.

In the present work, we show that PRP18 is involved only in the second step of splicing. In splicing reactions carried out in vitro, intermediates accumulate in the absence of PRP18. These intermediates can be chased to products by the addition of PRP18. We find that this chase does not require ATP, providing strong support for a transesterification mechanism for the second step of pre-mRNA splicing, and suggesting that PRP18 acts during the second transesterification reaction.

Results

Immunodepletion of PRP18

Antibodies against recombinant PRP18 were raised in rabbits (D. Horowitz and J. Abelson, in prep.). Immunodepletion of a yeast extract was carried out by incubating extracts with protein A-bound αPRP18. Some antibody was released from the resin during this incubation, and most of the released antibody was removed by a subsequent incubation with protein A-Sepharose. A Western blot of immunodepleted extract, probed with αPRP18, is shown in Figure 1. PRP proteins are typically difficult to detect (Jackson et al. 1988; Banroques and Abelson 1989; Petersen-Björn et al. 1989); so a high background in the blot is not unexpected. Quantitation of the blot pictured showed that >95% of the PRP18 had been removed. A faint band seen at the position of PRP18 in the αPRP18-depleted extract is also seen with similar intensity in extract from the strain DH120R, in which the PRP18 gene has been disrupted. None of the other proteins seen by blotting could be immunoprecipitated by αPRP18 (data not shown).

Kinetics of splicing

Initial work with extracts from prp18 temperature-sensitive yeast showed that PRP18 was involved in the second step of splicing (Vijayraghavan et al. 1989; Vijayraghavan and Abelson 1990). We also found this result by αPRP18 inhibition of splicing (D. Horowitz and J. Abelson, in prep.), but neither of these studies showed whether PRP18 might also be involved in the first step of splicing or whether PRP18 was absolutely required for the second step. To answer these questions, we have analyzed the kinetics of splicing in wild-type extracts depleted of PRP18. We used standard splicing conditions at 23°C (Lin et al. 1985). A gel of a time course of splicing in preimmune antibody-depleted and αPRP18-depleted extracts is shown in Figure 2A. Quantitative analyses of the gel are plotted in Figure 2B–D. In Figure 2B the kinetics of disappearance of pre-mRNA in the two extracts are compared; in Figure 2, C and D, the kinetics of formation of 2/3 lariat, exon 1, and mRNA are shown for each of the two extracts. The kinetics of splicing in the preimmune antibody-depleted extract and an untreated extract were essentially identical (data not shown). It is apparent from the gel (Fig. 2A) that the second step of splicing is inhibited in the PRP18-depleted extract. The 2/3 lariat and exon 1 intermediates accumulate; however, the second step of splicing is not completely inhibited, and mRNA is produced slowly. There are two interpretations of this result: Either PRP18 has not been completely depleted or PRP18 is not absolutely required for splicing. We favor the latter interpretation (see Discussion; D. Horowitz and J. Abelson, in prep.) Two as-
Figure 2. Kinetics of splicing and chasing of intermediates to products by PRP18. (A) A gel showing the time courses of splicing in preimmune antibody-depleted (lanes B–G) and αPRP18-depleted (lanes H–M) extracts. The time points are indicated in minutes at the top of each lane. Lane A shows pre-mRNA only. Two other results are also presented. The first is a splicing reaction with an αPRP18-depleted extract to which PRP18 had been added before the start of the splicing reaction (lanes N,O). The second is a chase of intermediates to products by the addition of PRP18; for this experiment, an aliquot of the splicing reaction using αPRP18-depleted extract (shown in lanes H–M) was removed at 11 min, PRP18 was added to this aliquot, and splicing was assayed at the indicated times (lanes P–R). All times have been rounded to the nearest minute. The positions of the 2/3 lariat, intron, pre-mRNA, mRNA, and exon 1 are indicated symbolically. A single exposure of one gel is shown; the lanes have been rearranged for clarity. (B) A comparison of the rates of disappearance of pre-mRNA in the splicing reactions displayed in A. (C–D) Graphs of the kinetics of splicing in preimmune antibody-depleted (C) and αPRP18-depleted (D) extracts. Pre-mRNA [□], 2/3 lariat [○], exon 1 [△], and mRNA [■] are shown. Values for 11 min were interpolated from D. (The interpolated values are <15% different from those at 10 min; the conclusions discussed in the text do not depend on the precision of the interpolations.) All values graphed in B–E were obtained from PhosphorImager quantitation of the gel shown in A; all RNAs are in relative molar amounts (arbitrary units), obtained by dividing the measured quantity from the PhosphorImager analysis (in arbitrary units) by the molecular weight of the RNA species. The vertical scales in B–E are in the same units.

Aspects of the analysis show that the first step of splicing is unaffected by the depletion of PRP18. The rates of disappearance of pre-mRNA in the two extracts are identical; because this process may be dominated by degradation of the pre-mRNA, it is not conclusive evidence that the first step is unaffected [Fig. 2B]. More convincingly, the amounts of pre-mRNA that have undergone the first step of splicing (given by the sum of the molar amounts of 2/3 lariat and mRNA in Fig. 2C,D) are essentially the same in both extracts.

The apparent yield of products in our splicing reactions was 15–20%, although 98% of the pre-mRNA is conserved during the reaction. Much of the RNA added to the extract has been degraded, and the stabilities of the various RNA species are different. Lariat intron is clearly less stable than mRNA [in Fig. 2A, six times more mRNA than lariat intron RNA (by molar amount) was present in the preimmune antibody-depleted extract after 20 min], and pre-mRNA also appears to be relatively unstable [see Fig. 2C,D]. Exon 1 is more stable than 2/3 lariat, perhaps owing to the difference in sizes. The variable stabilities of the RNAs and the complexity of the splicing reaction precluded any attempt to derive kinetic rate constants for the full splicing reaction. However, with some approximations, we can estimate a rate constant for the second step of splicing, as detailed in Materials and methods. We find that the second step is 80-fold slower in the PRP18-depleted extract than in the wild-type extract.

Activity of PRP18 protein
To determine whether purified PRP18 protein was active, we assayed splicing in a PRP18-depleted extract to
which purified recombinant PRP18 had been added. PRP18 was added to the PRP18-depleted extract immediately before the addition of the splicing cocktail. Essentially wild-type splicing was seen [Fig. 2A, lanes N,O]. Comparison of the splicing in the preimmune-depleted extract [Fig. 2A, lanes E,G], the αPRP18-depleted extract [Fig. 2A, lanes K,M], and the PRP18-return extract [Fig. 2A, lanes N,O] shows that normal splicing activity had been restored. This result demonstrates both that the purified PRP18 protein is active and that the PRP18-depleted extract has been specifically depleted of PRP18. About 40 times as much PRP18 as is present in a wild-type extract was added in this experiment [and in subsequent chasing experiments] to ensure rapid and complete reconstitution of activity. We have not determined the specific activity of our PRP18.

Chasing of intermediates to products by PRP18

Figure 2A shows the accumulation of intermediates in a PRP18-depleted extract; we wished to determine whether these intermediates could be chased to products by the addition of PRP18 protein. Splicing was carried out in a PRP18-depleted extract for 11 min for optimal formation of intermediates (Fig. 2). A portion of the reaction shown in Figure 2A (lanes H–K) was removed after 11 min for the chasing experiment. PRP18 was added, and aliquots were periodically removed for assay, as shown in Figure 2A (lanes P–R). Rapid chasing of the 2/3 lariat and exon 1 intermediates to mRNA and lariat intron was seen (Fig. 2A, cf. lanes K and P); the results are shown graphically in Figure 2E. Although lane K shows the splicing reaction 1 min before the chase was initiated, it is a suitable reference lane, as the amounts of the RNA species were changing slowly (Fig. 2D). About half of each intermediate was converted to product within 1 min, although the apparent yield of mRNA from 2/3 lariat is only ~60%, presumably owing to the instability of the RNA. This chase confirms the idea that the lariat 2/3 and exon 1 are intermediates in the splicing pathway. Comparison of the amounts of RNA [Fig. 2E] at the beginning of the chase at 11 min [Fig. 2A, lane K] and the first time point at 12 min [Fig. 2A, lane P] shows that the changes in the amounts of RNA are Δpre-mRNA = −0.3, Δ2/3 lariat = −1.7, Δexon 1 = −1.0, and ΔmRNA = +1.1 (relative molar amounts). The amount of mRNA produced is fourfold larger than the amount of pre-mRNA consumed during the chase; hence, the mRNA must have been produced from the 2/3 lariat and exon 1.

To determine whether the chasing of intermediates to products required ATP, we first developed a method for depleting ATP in yeast splicing reaction mixtures. The initial concentration of ATP in splicing reactions was 2 mM; after 10 min, at which time we wanted to remove the ATP, its concentration had fallen to 1 mM. We initially used hexokinase and glucose to deplete ATP (Black and Pinto 1989) but found that the hexokinase was superfluous; addition of 2 mM glucose resulted in destruc-

Figure 3. Thin-layer chromatograph showing the fates of the four NTPs and dATP in a splicing reaction and following glucose-caused depletion. For each of the five reactions, the α32P-labeled nucleoside triphosphate was added to a standard splicing reaction (with unlabeled transcript in place of α2P-labeled transcript) at the outset. Time points were taken at 1, 10, and 20 min. At 10 min, 2 mM glucose was added to an aliquot of each splicing reaction, and the NTP in each was analyzed by thin-layer chromatography (TLC) in 1.2 M LiCl (Randerath and Randerath 1964). A typical result is shown in Figure 3. ATP was destroyed rapidly (<3 min); ATP was not detected either by long exposure or with the PhosphorImager. The TLC results showed that the concentration of ATP remaining after the depletion was <5 μM. We have also measured the concentration of ATP using the luciferase–luciferin bioluminescence assay, which gave an ATP concentration of 2–3 μM, consistent with the TLC results. We are uncertain what the processes leading to the depletion of ATP are; some AMP, but very little ADP, was seen after the ATP was depleted. Endogenous hexokinase and nucleoside monophosphate kinase may be responsible for these results. Similar experiments to track the other NTPs, as well as dATP, were done (Fig. 3). In these experiments, [α-32P]NTP [or [α-32P]dATP] was substituted for [α-32P]ATP. Following incubation under splicing conditions, the NTP was analyzed by TLC. In all cases, substantial depletion of the NTP was seen. The concentration of CTP was reduced by >400-fold, GTP by 40-fold, UTP by >50-fold, and dATP by 75-fold. The total NTP concentration was mea-
sured using bioluminescence; for this measurement, NTPs in a phenol-extracted extract were converted to ATP using nucleoside diphosphate kinase. The concentration of NTPs other than ATP was \( \sim 1 \mu \text{M} \). The depletion of the NTPs may result from their rapid conversion to ATP. We found that 2 mM glucose was sufficient for the depletion of ATP; 2 mM ATP added after the depletion was relatively stable. When 5 mM glucose was used for the depletion, additional ATP was not stable.

Using this method for removing ATP, we carried out chase experiments analogous to those described above. An outline of the experiment is shown above the gel in Figure 4A. Splicing was carried out under normal conditions in a PRP18-depleted extract. At \( t = 10 \text{ min} \) or \( t = 9 \text{ min} \) a portion of this splicing reaction was transferred to a tube containing sufficient glucose to bring the final concentration of glucose to 2 mM (or 5 mM). After a 4-min incubation to allow depletion of the ATP, either PRP18 or PRP18 and ATP were added. A gel showing the time course of each of the different reactions is pictured in Figure 4A; a single splicing reaction was the starting point for all of the samples shown in Figure 4. The main conclusion from this experiment is that ATP is not required for chasing of the intermediates to products by PRP18. This conclusion can be seen from a comparison of the amounts of the spliced RNAs in the ATP-depleted splicing reaction before the addition of PRP18 (lane F) and 3 min after the addition of PRP18 (lane I). In the absence of ATP, the addition of PRP18 caused a reduction in the amounts of 2/3 lariat and exon 1 intermediates, and an increase in the amounts of mRNA and lariat intron. There was a twofold increase in the amount of mRNA, and a threefold increase in the amount of intron. About 35% of the 2/3 lariat consumed in the reaction is found in mRNA. When ATP was added with PRP18, the chase was apparently more complete, and the apparent yield of mRNA was higher (55%).

Figure 4, B and C, shows graphically the amounts of
2/3 lariat and mRNA present in lanes A–M in the gel shown in Figure 4A. The time course of splicing in the PRP18-depleted extract (lanes A–D) is essentially the same as that shown in Figure 2C. Following depletion of ATP, the pre-mRNA was stable (lanes E–G and N–P). In these ATP-depleted extracts, 2/3 lariat and exon 1 were not entirely stable (Fig. 4B), and a small amount of mRNA was produced (Fig. 4C). This observation is consistent with the idea that neither PRP18 nor ATP is required (although both are beneficial) for the second reaction. Addition of PRP18 (lanes H–J) or PRP18 and ATP resulted in a rapid decline in the amounts of 2/3 lariat (Fig. 4B) and exon 1. [The decrease in the amount of exon 1 occurs at ~65% of the rate of that of 2/3 lariat in all of the chase experiments in Figs. 2 and 4.] mRNA (Fig. 4C) and intron are produced during the chase. The twofold increase in the amount of mRNA produced in the absence of ATP is reproducibly seen. Essentially identical chasing was seen when 5 mM glucose was used to deplete the ATP (Fig. 4A, lanes N–S), in this case, ATP added after the depletion is not stable because of the excess of glucose.

In the control experiments for the ATP-independent chasing experiment, we particularly wished to be able to determine whether ATP might be bound to the spliceosome before the second step of splicing occurs and then used during the reaction, and whether the concentration of ATP might still be high enough after the depletion of ATP to support splicing. We carried out an experiment identical to that in Figure 4 except that a preimmune antibody-depleted extract was used in place of the αPRP18-depleted extract (data not shown). After depletion of the ATP, the pre-mRNA, the splicing intermediates, and both products of the splicing reaction were stable, declining by <15% during the 8 min incubation. Figure 2D shows that the levels of all of the RNAs except the pre-mRNA change only slowly in the interval from 10 to 20 min in the presence of ATP, so little can be attributed to their increased stability. However, there was no indication that 2/3 lariat and exon 1 could be further spliced following the depletion of ATP, as might be expected if sufficient ATP remained to carry out the second step after depletion of ATP. We do not know why the pre-mRNA is stable in the absence of ATP; it may be protected against attack, because the debranched intron is rapidly degraded in the absence of ATP.

A second control experiment is shown in Figure 5. Extract was incubated under splicing conditions in the absence of pre-mRNA. Glucose was added to deplete the ATP, and then either pre-mRNA or pre-mRNA and ATP were added, as outlined in Figure 5. No splicing of the added pre-mRNA was seen unless ATP was also added, showing that the remaining ATP was insufficient to support the complete splicing reaction.

Discussion

We have analyzed pre-mRNA splicing in yeast extracts depleted of the PRP18 protein. We showed that PRP18 is involved only in the second step of splicing. Intermediates accumulated in splicing reactions in the absence of PRP18, and these intermediates could be rapidly chased to products by added PRP18 even in the absence of ATP.

We carried out an analysis of the kinetics of splicing in a PRP18-depleted extract. Depletion of PRP18 had no effect on the first step of splicing but did slow the second step substantially, showing that PRP18 is involved only in the second step of splicing. The second step was not, however, completely inhibited. While this remaining splicing activity may have been the result of a trace of PRP18 in the extract, we believe that this activity indicates that PRP18 is not absolutely required for the second step of splicing. As reported (D. Horowitz and J. Abelson in prep.), yeast strains bearing a null allele of PRP18 are viable at 23°C, but grow extremely slowly, and do not grow at 37°C; even at 23°C these yeast strains accumulate unspliced RNA. Corresponding results were
obtained in vitro; in extracts depleted of PRP18, the rate of the second step of splicing declined as the temperature was increased. We concluded that PRP18 was not absolutely essential for splicing, and, hence that the residual splicing activity at 23°C in the PRP18-depleted extracts represents PRP18-independent splicing. As discussed in D. Horowitz and J. Abelson (in prep.), where we showed that PRP18 is associated with the U5 snRNP, the results of the kinetic analysis imply that the U5 snRNP is involved in the second step.

Wild-type splicing activity can be restored to PRP18-depleted extracts by the addition of purified PRP18 protein. Analogous experiments with other PRP proteins have generally, but not always, been successful [Chang et al. 1988]. The 2/3 lariat and exon 1 intermediates that accumulate during splicing in a PRP18-depleted extract can be chased to products by the addition of purified PRP18 protein. This chase shows that the PRP18-binding site remains accessible in the spliceosome; similar experiments have been reported with PRP2 and PRP16 proteins [Lin et al. 1987; Schwer and Guthrie 1991]. As described in Results, the chasing of the accumulated 2/3 lariat and exon 1 to products is unambiguous and proves that they are intermediates in the splicing pathway.

The chasing of intermediates made in a PRP18-depleted extract to products was not dependent on ATP. Following the formation of intermediates in a PRP18-depleted extract, the ATP was depleted by the addition of glucose. PRP18 was then added, leading to chasing of the 2/3 lariat and exon 1 to mRNA and lariat intron [Fig. 4]. This experiment strongly suggests that ATP hydrolysis is not coupled to the second cleavage-and-ligation reaction in pre-mRNA splicing. Depletion of ATP was followed by TLC and bioluminescence; we found that the ATP concentration had been reduced ~700-fold, from 2 mM to ~3 µM. PRP2 and PRP16, which are both ATPases required for splicing, have been shown to hydrolyze NTPs other than ATP [Kim et al. 1992, Schwer and Guthrie 1992]; we therefore measured the concentrations of other NTPs and found that all were substantially depleted when the ATP was depleted. Control experiments showed that the amount of ATP remaining was insufficient to support splicing. Experiments on PRP2, which is required for the first step of splicing, have shown that 20 µM ATP is needed for good PRP2 activity (S.-H. Kim and R.-J. Lin, personal communication). For PRP16, which is required for the second step of splicing, 100 µM ATP is sufficient for good splicing activity in chasing experiments [Schwer and Guthrie 1992]. Our measured ATP concentration is significantly lower than that needed by PRP2 or PRP16, but comparisons of these numbers, derived from different procedures using different splicing extracts, are problematic. There is a possibility that ATP is stably bound to the spliceosome in our reaction before the depletion takes place and that this bound ATP is then hydrolyzed during the second reaction. Although this possibility is not addressed by our experiments, it would require that ATP have a high affinity for the spliceosome. A dissociation constant of 3 µM corresponds to a 7 kcal-binding energy (using $\Delta G = -RT \log K$), about half of the energy available from ATP hydrolysis. A more difficult question to answer is whether there is energy stored in the spliceosome. Energy could be stored as chemical energy, as in an adenylylated protein, or as mechanical energy. Our experiments do not address this question.

Following the depletion of ATP, the chasing of intermediates to products was apparently more efficient if ATP was included in the chase. This result is puzzling if the chase reaction is independent of ATP. One explanation for this observation is that in the absence of ATP there is no driving force for the reaction. We expect that in the second reaction the reactants and products are similar in free energy. Therefore, in the absence of ATP in the chase, the intermediates and products may be approaching equilibrium. The disassembly of the spliceosome is very likely an active process involving ATP hydrolysis by the PRP22 protein [Company et al. 1991]. Thus, in the presence of ATP, the products are pushed forward in the splicing pathway by ATP hydrolysis, and the equilibrium in the second reaction of splicing is shifted far toward the formation of products. Similar energetic schemes apply to self-splicing RNAs, in which splicing is driven by cyclization of the intron after the splicing reaction is completed [Zaug et al. 1983]. ATP might also lead to improvement in the chasing efficiency if some of the 2/3 lariat and exon 1 seen is present in complexes that are stopped at steps before, for example, the PRP16-requiring step, as might occur if PRP18 has some effect on the early stages of the second step. Two other factors also may complicate the comparison of chasing in the absence and presence of ATP. First, the stabilities of the various RNAs are different and are affected by ATP; and, second, in the presence of ATP, pre-mRNA may be spliced to mRNA during the chase. Therefore, some of the difference in the efficiencies of chasing may be misleading.

Vijayraghavan and Abelson (1990) carried out a different type of chasing experiment using extracts from prp18 temperature-sensitive strains. In their experiments, spliceosomes formed in heat-inactivated extracts from a prp18 temperature-sensitive strain were purified in glycerol gradients. Chasing of the intermediates to products, done with heat-inactivated extracts from other prp temperature-sensitive strains, or with fractionated extracts, required ATP. Our results show that the second step of splicing continues in PRP18-depleted extracts lacking ATP; we suggest that during the sedimentation of spliceosomes, all of the PRP18-chatable intermediates were spliced, leaving only intermediates trapped at earlier stages of the second step. Chasing from these stages requires ATP [Schwer and Guthrie 1991].

The finding that the second cleavage-and-igation reaction of splicing proceeds in the absence of ATP strongly supports the idea that this reaction is a transesterification [Cech 1986; Jacquier 1990]. In the absence of ATP hydrolysis or other energy source, the number of phosphodiester bonds must be conserved during the reaction, as is the case in a transesterification. Both group I and group II self-splicing occurs by transesterification.
mechanisms, and the similarity between the intermediates and products in pre-mRNA splicing and group II splicing has led to suggestions that these two types of splicing are evolutionarily or mechanistically related (Cech 1986; Jacquier 1990). U6 snRNA has often been proposed as the RNA catalyst of the transesterification reaction, owing to its high degree of evolutionary conservation (Brow and Guthrie 1988), its involvement in both steps of splicing (Fabrizio and Abelson 1990), and its similarity to other self-splicing RNAs (Tani and Oshihama 1991). A recent proposal suggests that the second reaction of splicing is analogous to group I splicing, with ATP hydrolysis carried out by PRP16 (Schwer and Guthrie 1991). Our findings imply that hydrolysis of ATP precedes the actual transesterification reaction, at least for the second step. In yeast, ATP hydrolysis by two proteins, PRP2 and PRP16, which both RNA-dependent ATPases, has been studied extensively, and the results of these studies are consistent with our conclusions. PRP16 is required for the second step of splicing, and we would expect it to act before the second reaction. Spliceosomes assembled without PRP16 carry out the first step of splicing, and have recently shown that addition of PRP16 and ATP to these spliceosomes causes a conformational change in the spliceosome, which makes the 3′ splice site inaccessible for oligonucleotide-directed cleavage by RNase H. This conformational change, which requires hydrolysis of ATP, occurs in mutant substrates that do not undergo the second step of splicing, suggesting that the ATP hydrolysis carried out by PRP16 precedes the transesterification reaction. The PRP2 protein is required for the first step of splicing (Lin et al. 1987). Spliceosomes assembled without PRP2 require both PRP2 and ATP, but no other extrinsic factors, to carry out the first step of splicing (Kim and Lin 1993). However, no direct connection between ATP hydrolysis and the splicing reaction has been demonstrated.

Our results have a number of implications for the role of PRP18 in pre-mRNA splicing. As discussed above, PRP18 is involved exclusively in the second step of splicing; it is not absolutely required for splicing at low temperature, although the second step is greatly slowed in the absence of PRP18. Several other factors are also involved in the second step of splicing, including three proteins, PRP16 (Schwer and Guthrie 1991), PRP17 (Vijayraghavan et al. 1989), and SLU7 (Frank et al. 1992) and three U snRNAs, U2 (McPheeters and Abelson 1992), U5 (Newman and Norman 1992), D. Horowitz and J. Abelson (in prep.), and U6 (Fabrizio and Abelson 1990). The ATP independence of the chasing of intermediates into products by PRP18 allows us to order the actions of some of these factors. It is clear that PRP18 acts after PRP16. The ATP dependencies for the chasing of intermediates into products for other splicing factors have not yet been determined, so they cannot presently be ordered.

Our findings lead us to suggest that PRP18 functions during the second transesterification reaction, in contrast to PRP16 and SLU7, which appear to act before the actual cleavage-and-ligation reaction. This conclusion is suggested primarily by the absence of an ATP requirement for the chasing of intermediates to products by PRP18. This finding restricts the number and kinds of conformational changes that can occur between the addition of PRP18 and the second transesterification, precluding, for example, proofreading steps. We think that it is plausible that PRP18 acts during the transesterification reaction. The precise role of PRP18 in the reaction remains to be defined; it may play a largely structural role, stabilizing a conformation of the U5 snRNP, or it may be a more active participant.

Materials and methods

Materials and strains

Yeast splicing extracts [Lin et al. 1985] were made from CBO18 [a CR1 pep4A::HIS3 prbl::hisG prclA::hisG] (from J. Robinson, California Institute of Technology, Pasadena). Labeled nucleotides were from New England Nuclear. The luciferin-luciferase ATP Assay Kit, nucleoside diphosphate kinase, and protein A-Sepharose were from Sigma. The Enhanced Chemiluminescence (ECL) system was from Amersham. cPRP18 antiserum is described by D. Horowitz and J. Abelson (in prep.), in this study we used the antibody referred to as αA.

Depletion of PRP18 from yeast extracts

Fifty microliters of protein A-Sepharose were incubated with 240 μl of αPRP18 or preimmune serum for 1 hr and then washed extensively. Yeast splicing extract (300 μl) containing 20 mM K2HPO4/KH2PO4 at pH 7.4, 100 mM KC1, and 0.05% NP-40 was added to the resin, incubated for 70 min at 4°C, recovered by filtration, and added to 15 μl of protein A-Sepharose, incubated for 30 min at 4°C, and recovered by filtration.

Western blotting

Western blotting was carried out as described by D. Horowitz and J. Abelson (in prep.), using the ECL system.

Splicing reactions

Splicing was carried out as described in Lin et al. (1985). The total concentration of K2HPO4/KH2PO4 was 40 mM (to account for the additional KC1 present in the depleted extracts) when antibody-depleted extracts were assayed. To deplete ATP, glucose in a minimum volume was added to a final concentration of 2 mM (or 5 mM). All incubations were carried out at 23°C. For reconstitution of activity with PRP18, 15–30 ng of PRP18 per microliter of yeast extract was added, similar amounts of PRP18 were used in the chasing experiments. PRP18 was expressed and purified to near homogeneity from Escherichia coli, as described by D. Horowitz and J. Abelson (in prep.). All gels were quantitated on a Molecular Dynamics Phosphoimager.

The rate constant for the second step of splicing was esti-
imated as follows. We assumed that the second step could be modeled as a single-step, unimolecular reaction, and that degradation of mRNA could be neglected. The rate equation for the production of mRNA is \( \frac{\Delta[mRNA]}{\Delta t} = k_2[\text{exon} 1 \cdot 2/3 \text{lariat}] \), where exon 1 \( \cdot 2/3 \text{lariat} \) represents the spliceosome containing the two intermediates. This equation was approximated as \( \Delta[mRNA]/\Delta t \approx k_2[\text{exon} 1 \cdot 2/3 \text{lariat}]_{avg} \) for the time intervals shown in Figure 2, allowing us to calculate values for \( k_2 \).

**NTP assays**

Thin-layer chromatographs were run on polyethyleneimine-cellulose with 1.2 M LiCl, as described in Randerath and Rendarath [1964]. Luciferin-luciferase assays for ATP were carried out according to the manufacturer’s recommendations. ATP was assayed directly in splicing reaction mixtures depleted of erath (1964). Luciferin-luciferase assays for ATP were carried out according to the manufacturer’s recommendations. ATP with 2 mM or 5 mM glucose, yielding an ATP concentration of 3 mM. For measurement of the total NTP concentration, 20 \( \mu \text{M} \) ADP and nucleoside diphosphate kinase were added to phenol- and ether-extracted reaction mixtures to convert NTPs to ATP, whose concentration was measured. Where possible, internal standards were used to calibrate the measurements.

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