Pre-mRNA splicing alters mRNP composition: evidence for stable association of proteins at exon–exon junctions

Hervé Le Hir,1,2 Melissa J. Moore,2,3 and Lynne E. Maquat1,3

1Department of Cancer Genetics, Roswell Park Cancer Institute, Buffalo, New York 14263 USA; 2Department of Biochemistry, Howard Hughes Medical Institute/Brandeis University, Waltham, Massachusetts 02454 USA

We provide direct evidence that pre-mRNA splicing alters mRNP protein composition. Using a novel in vitro cross-linking approach, we detected several proteins that associate with mRNA exon–exon junctions only as a consequence of splicing. Immunoprecipitation experiments suggested that these proteins are part of a tight complex around the junction. Two were identified as SRm160, a nuclear matrix-associated splicing coactivator, and hPrp8p, a core component of U5 snRNP and spliceosomes. Glycerol gradient fractionation showed that a subset of these proteins remain associated with mRNA after its release from the spliceosome. These results demonstrate that the spliceosome can leave behind signature proteins at exon–exon junctions. Such proteins could influence downstream metabolic events in vivo such as mRNA transport, translation, and nonsense-mediated decay.

[Key Words: mRNP composition; pre-mRNA splicing; RNA–protein cross-linking; exon–exon junctions]

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Eukaryotic mRNAs are generated by a series of metabolic events including transcription, capping, splicing, and polyadenylation. Once made, mRNAs are transported to the cytoplasm where they undergo translation and, ultimately, decay. It is becoming increasingly clear that all of these metabolic steps are mechanistically linked in intact cells. For example, recent evidence suggests that the carboxy-terminal domain (CTD) of RNA polymerase II delivers proteins required for capping, splicing, and polyadenylation to the nascent transcript in vivo (Cho et al. 1997; McCracken et al. 1997a,b; Misteli and Spector 1999) and enhances splicing and polyadenylation in vitro (Hirose and Manley 1998; Hirose et al. 1999). Moreover, the nature of the promoter can affect alternative splicing patterns of transcripts (Cramer et al. 1999). Also, proteins acquired in the nucleus are essential for proper localization of certain mRNAs in the cytoplasm (Lall et al. 1999).

The action of spliceosomes also can influence downstream mRNA metabolism. The presence and position of pre-mRNA introns can affect the efficiency of mRNA transport (Chang and Sharp 1989; Legrain and Rosbash 1989; Pasquinelli et al. 1997; Saavedra et al. 1997; Luo and Reed 1999), the efficiency of mRNA translation [Matsumoto et al. 1998], and the rate of mRNA decay [Maquat 1995, 1996; Li et al. 1997; Hentze and Kulozik 1999]. Studies of nonsense-mediated mRNA decay indicate that mammalian cells can distinguish authentic from premature stop codons by their positions relative to the 3’-most exon–exon junction position in mRNA (Cheng et al. 1994; Carter et al. 1996; Nagy and Maquat 1998; Thermann et al. 1998; Zhang et al. 1998a,b).

The most probable means by which pre-mRNA splicing influences subsequent mRNA metabolism is by altering the structure of the messenger ribonucleoprotein particle (mRNP) [Nakielny and Dreyfuss 1997; Izaurralde and Adam 1998; Luo and Reed 1999]. Such alterations could consist of covalent nucleotide modifications [Rottman et al. 1994] or noncovalent associations of specific proteins, either of which could stay with an mRNP throughout all or a portion of its lifetime. In fact, Luo and Reed (1999) demonstrated recently that an mRNP generated by splicing in vitro has altered mobility in native gels and is exported more rapidly and efficiently from Xenopus laevis nuclei than an mRNP not generated by splicing. A number of proteins known to be involved in pre-mRNA splicing, such as hnRNP A1 [Dreyfuss et al. 1993; Nakielny and Dreyfuss 1997] and a subset of the SR splicing factors [Cáceres et al. 1998], shuttle between the nucleus and cytoplasm, making them potential mRNP components. HnRNP A1 clearly functions in mRNA export [Izaurralde et al. 1997], and
electron microscope tomography has shown that an hnRNP A1-related protein accompanies Balbiani ring granules, giant mRNPs of the dipteran Chironomus tentans, through nuclear pores and into cytoplasmic polyribosomes (Visa et al. 1996). However, despite its demonstrated importance, the extent to which mRNP composition is altered by splicing is currently unknown.

We report direct evidence that pre-mRNA splicing alters mRNP protein composition. We employed a new in vitro cross-linking strategy designed to identify proteins left behind by the splicing machinery specifically at mRNA exon–exon junctions. Dependent on the nature of the cross-linker, the position of the cross-linker relative to the exon–exon junction and the pre-mRNA sequence, we observed at least four different proteins that cross-link only to mRNAs generated by splicing in HeLa-cell nuclear extract. Immunoprecipitations suggested that these four proteins are part of a tight complex around the exon–exon junction. Two of the proteins were identified as SRm160, a nuclear matrix-associated splicing coactivator subunit, and hPrp8p, a highly conserved U5 snRNP protein. Glycerol gradient fractionation indicated that most of the proteins that cross-linked in a splicing-dependent manner remain associated with mRNA after its release from the spliceosome. These data provide a new view of the dynamic nature of protein-RNA interactions at exon–exon junctions after splicing and identify proteins that might influence subsequent mRNA fate.

Results

Cross-linking strategy

To identify proteins associated with mRNA specifically as a consequence of splicing, we constructed single intron-containing pre-mRNAs having two site-specific modifications: a photoreactive group near the intron–exon-containing pre-mRNAs having two site-specific modifications: a photoreactive group near the intron–exon boundary. In vitro splicing of such pre-mRNAs in HeLa-cell nuclear extract juxtaposes both groups at the exon–exon junction in the mRNA product (Fig. 1, right). After irradiation at a wavelength appropriate for the photoreactive group followed by ribonuclease treatment and then electrophoresis through a denaturing gel, only proteins attached to the cross-linkable moiety at the exon–exon junction, and therefore associated with the 32P, are detectable by autoradiography.

The first substrate we prepared was PIP:E1(B) pre-mRNA, a derivative of PIP.85B pre-mRNA (Query et al. 1994) that contained a benzophenone moiety (B) [MacMillan et al. 1994; Moore and Query 1998] at the penultimate nucleotide of exon 1 (E1, open box) and a single 32P [star] at the beginning of exon 2 (shaded box). Control mRNA (left) was structurally identical to the spliced product of experimental pre-mRNA. Control pre-mRNA (center) was identical to experimental pre-mRNA except that the 32P was positioned at the 5′ splice site.

Figure 1. Schematic structures of site-specifically modified PIP:E1(B) RNAs. The experimental pre-mRNA (right) contained a benzophenone (B) at the penultimate nucleotide of exon 1 (E1, open box) and a single 32P [star] at the beginning of exon 2 (shaded box). Control mRNA (left) was structurally identical to the spliced product of experimental pre-mRNA. Control pre-mRNA (center) was identical to experimental pre-mRNA except that the 32P was positioned at the 5′ splice site.

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The control pre-mRNA permitted identification of proteins that interact with the benzophenone only before lariat formation, because in this construct the benzophenone and 32P became separated during the first step of splicing and, therefore, were not juxtaposed in spliced mRNA.

Splicing and cross-linking of modified RNAs

To determine splicing kinetics and spliced product stability, each benzophenone-containing RNA was incubated under splicing conditions in HeLa-cell nuclear extract (Fig. 2A). Both pre-mRNAs spliced to similar levels, with splicing intermediates evident after 30 min and little additional product accumulation after 90 min (Fig. 2A, lanes 6–15). Control mRNA was degraded significantly over the course of the incubation (Fig. 2A, lanes 1–5). This facilitated differentiation of proteins that cross-link independent of splicing from those that cross-link dependent on splicing because the amount of mRNA that was or was not generated by splicing was comparable at both 90 and 120 min (Fig. 2A, cf. lanes 4 and 5 with 14 and 15). Neither splicing efficiency nor RNA stability was affected by the presence of the photoreactive group (data not shown, MacMillan et al. 1994).

To induce cross-links, each splicing reaction was incubated under splicing conditions (30°C) for 0, 45, or 90 min and then irradiated with a 302-nm lamp for 20 min on ice (4°C). No additional spliced product accumulated during the 4°C incubations, indicating that splicing was not ongoing during the period of UV irradiation (data not shown). Samples were subsequently treated with RNase A and analyzed by SDS-PAGE (Fig. 2B). Numerous labeled bands were observed with all three RNAs. Because no cross-linking was observed when the reactions were treated with proteinase K (Fig. 2B, lanes 4,9,14) or when RNAs lacked a benzophenone moiety (Fig. 2B, lanes 5,10,15; data not shown, except in lane 10, where the low level (<10%) of 254-nm light from the 302-nm lamp was...
sufficient to cross-link a small amount of a 220-kD protein to the 5’ splice site (see below)]. bands detected in other reactions represented proteins specifically cross-linked to the photoreactive group.

Multiple proteins cross-linked to the experimental PIP:E1(B) RNA after 45 and 90 min of splicing, times by which both modifications were juxtaposed at the mRNA exon–exon junction (Fig. 2B, lanes 12,13). The pattern of cross-linked proteins was the same regardless of whether RNase treatment was performed before protein denaturation or after boiling samples in the presence of 0.5% SDS (data not shown). However, the majority of these proteins were nonspecific because they also cross-linked to control mRNA, control pre-mRNA, or both. It should be noted that similar proportions of nonspecific protein cross-links have also been observed by others when highly reactive photo-cross-linking moieties, such as benzophenone, were employed with no affinity purification step after cross-linking (Wyatt et al. 1992; MacMillan et al. 1994; Moore and Query 1998). Nonetheless, careful comparison of the pattern obtained with or without splicing revealed four proteins, with apparent molecular masses of ~220, 160, 50, and 20 kD, that clearly cross-linked in a splicing-dependent manner [Fig. 2B, bands marked with an asterisk in lanes 11–13 and 16–18 (darker exposure)]. These proteins did not cross-link to control mRNA, control pre-mRNA, or both.

Figure 2. Splicing and cross-linking of benzophenone-containing PIP:E1(B) RNAs. (A) Splicing time courses. RNAs were incubated under splicing conditions for times indicated and separated by denaturing PAGE. Substrate and product structures are indicated above and to the right of each panel. Because all constructs contained only one 32P label (star), only a subset of splicing products are detectable. The small amount of RNA of similar mobility to mRNA in lanes 8–10 is debranched intron; this species contains label only when generated from control pre-mRNA. Species likely generated by exonuclease digestion of the lariat intron–exon 2 (Moore and Sharp 1992) (●) and four-way RNA ligation intermediates that copurified with full-length pre-mRNA (●●) are indicated. (B) Protein cross-linking. After incubation under splicing conditions for times indicated, samples were irradiated on ice at 302 nm and digested with RNase A. Reactions in lanes 4, 9, and 14 were treated with proteinase K after cross-linking; RNAs in lanes 5, 10, and 15 were subjected to splicing and cross-linking but contained no benzophenone. Lanes 16–18 are longer exposures of lanes 11–13. Molecular mass standards [kD] and apparent molecular masses of proteins that cross-linked in a splicing-dependent manner (●) are indicated to the left and right, respectively. (●) indicates position of the ~70-kD protein discussed in the text. (C) Side-by-side comparisons of protein cross-linking patterns. RNAs (ctl) control; (exp) experimental] were incubated under splicing conditions for 90 min, UV irradiated, and RNase A digested. Cross-linked proteins were separated in 7.5% [lanes 1–3], 10% [lanes 4–6], or 16% [lanes 7–9] denaturing polyacrylamide gels. Proteins that cross-link in a splicing dependent manner are indicated. (D,E) Dependence of cross-links on splicing. The indicated RNAs were incubated under splicing conditions for times shown and irradiated as in B. As specified, splicing was inhibited by ATP omission (−ATP) or blocking U2 snRNA with a complementary 2′-O-methyl oligonucleotide (+α-U2 oligonucleotide). (D) RNAs; (E) cross-linked proteins. ● and * are as in A and B, respectively.
control mRNA at any point [Fig. 2B, lanes 1–3] or to experimental RNA prior to spliceosome assembly [Fig. 2B, lanes 11,16]. Polyacrylamide gels of 7.5%, 10%, and 15% which allowed for greater separation in the >150-, 50-, and 20-kD ranges, respectively, corroborated these conclusions [Fig. 2C]. Moreover, 32P labeling of the specific proteins increased with longer splicing incubations, as would be expected for proteins that interact with the mRNA-spliced product (Fig. 2B, lanes 12,13, data not shown). When splicing of experimental pre-mRNA was inhibited by ATP omission [Fig. 2D, lane 3] or blocking U2 snRNA function with a 2′-O-methyl oligonucleotide complementary to the branch site binding region [Fig. 2D, lane 4], all four specific cross-links disappeared [Fig. 2E, lanes 5 and 6, respectively], additionally confirming their dependence on splicing. However, as expected, the cross-linking pattern observed with the mRNA control did not change significantly when U2 snRNA function was blocked [Fig. 2E, lanes 1,2].

A 220-kD protein also interacted with the 5′ splice site prior to lariat formation but dependent on spliceosome assembly [Fig. 2B, cf. lane 6 with lanes 7 and 8; data not shown], as did a protein of approximately 70 kD (band marked with a solid circle in Fig. 2B, lanes 7,8). As would be expected of proteins interacting with the photoreactive group before the first step of splicing, the extent of p220 and p70 cross-linking to control pre-mRNA decreased over time as the 32P label became separated from the benzophenone by 5′-methyl oligonucleotide (Fig. 2D, lanes 9,10). Moreover, the relative intensities of the four splicing-dependent cross-links were not affected by IP. No proteins that cross-linked to control mRNA were precipitated when RNase treatment preceded the IP (Fig. 2E, lane 3), and only the splicing-dependent 220- and 70-kD proteins that cross-linked to control pre-mRNA were precipitated under these conditions (Fig. 2A, lane 7).

Taken together, the above results demonstrate clearly that the protein cross-linking pattern around the exon–exon junction is different for an mRNA generated by splicing and one not undergoing this process. For PIP:E1(B) mRNA, four proteins with mobilities of ~220, 160, 50, and 20 kD cross-linked to the exon–exon junction specifically as a consequence of splicing.

**Immunoprecipitation of cross-linked proteins**

With the aims of separating the proteins that cross-linked dependent on splicing from those that cross-linked nonspecifically, as well as identifying the specific proteins, we performed a series of immunoprecipitations (IPs). Initially, we employed two monoclonal antibodies [mAbs], NM4 and B1C8, which had been shown previously to preferentially coimmunoprecipitate exon-containing RNA species, including the mRNA product, from in vitro splicing reactions [Blencowe et al. 1995]. NM4 recognizes the SR-related, nuclear matrix splicing coactivator subunits SRm300 and SRm160 [Blencowe et al. 1998; Eldridge et al. 1999] and most of the prototypic SR protein splicing factors [Zahler et al. 1992; Blencowe et al. 1995], whereas B1C8 specifically recognizes SRm160 [Blencowe et al. 1998].

Splicing reactions containing either experimental or control PIP:E1(B) RNAs were immunoprecipitated after UV irradiation. To examine the extent to which the IP pattern was dependent on intact RNA, RNase treatment was performed either before or after IP. RNase treatment prior to IP would be expected to eliminate any proteins complexes with the mAb epitope only by virtue of being bound to the same RNA. When mAb NM4 was employed and RNase treatment followed IP, numerous cross-linked proteins were precipitated from all three reactions [Fig. 3A, lanes 2,6,10]. However, when RNase treatment preceded the IP, all four proteins that cross-linked to experimental pre-mRNA in a splicing-dependent manner were efficiently precipitated, whereas other cross-linked proteins were not (Fig. 3A, cf. lane 11 with lanes 9 and 10). Moreover, the relative intensities of the four splicing-dependent cross-links were not affected by IP. No proteins that cross-linked to control mRNA were precipitated when RNase treatment preceded the IP [Fig. 3A, lane 3], and only the splicing-dependent 220- and 70-kD proteins that cross-linked to control pre-mRNA were precipitated under these conditions (Fig. 3A, lane 7).

The simplest interpretation of the above results is that the four proteins that cross-link to the PIP:E1(B) exon–exon junction dependent on splicing form a tight complex that is stable to RNase digestion, and the complex contains at least one protein bearing an NM4 epitope. However, the possibility that the nature of the complex varies between mRNA molecules so that individual molecules are bound by at least one protein bearing an NM4 epitope and any or all of the remaining three proteins cannot be discounted. Remarkably, IPs performed on experimental pre-mRNA using mAb B1C8 after RNase treatment, although less efficient overall, yielded a very similar pattern of precipitated proteins to that obtained with mAb NM4 [Fig. 3B, cf. lanes 4 and 7]. Because B1C8 is specific for SRm160, SRm160 must be a component of the protein complex at the exon–exon junction. To test whether the cross-linked 160-kD protein was SRm160, IPs were performed next with the two mAbs after both RNase digestion and protein denaturation. This reduced significantly the number of cross-links observed [Fig. 3B, lanes 5–6,8–9]. Under the most stringent denaturing conditions, the only splicing-dependent cross-linked protein immunoprecipitated with either mAb was the 160-kD protein [Fig. 3B, lanes 6,9], identifying it as SRm160.

Proteins with apparent molecular masses of ~55, 40, and 25 kD that cross-linked independent of splicing were also immunoprecipitated by mAb NM4 after stringent protein denaturation [Fig. 3B, lane 6]. This raised the possibility that these proteins were SRp55, SRp40, and either SRp30 or SRp20. We have confirmed that the 25-kD protein is SRp20, since SRp20 antiserum [Neugebauer and Roth 1997] immunoprecipitated the ca. 25-kD protein, and this protein comigrated with SRp20 when assayed by Western blot hybridization [data not shown].

Because the 220-kD protein was not immunoprecipitated by mAb NM4 after protein denaturation, it was unlikely SRm300 [Fig. 3B, lane 6]. Another possibility was hPrp8p, a 279-kD component of U5 snRNP [Luo et al. 1999], because a similar-sized protein also cross-linked to the 5′ splice site after spliceosome assembly [Fig. 2B, lanes 7,8], and other studies have shown that Prp8p cross-links in the vicinity of both the 5′ splice site...
prior to lariat formation (Wyatt et al. 1992; Reyes et al. 1996) and the 3’ splice site prior to exon ligation (Teigelkamp et al. 1995; Umen and Guthrie 1995; Chiara et al. 1996, 1997). Of all the proteins cross-linked to experimental PIP:E1(B) RNA, only the 220-kD band was immunoprecipitated with hPrp8p antiserum after stringent protein denaturation (Fig. 3C, lane 8). This protein was not immunoprecipitated by preimmune serum (Fig. 3C, lane 9). Therefore, the 220-kD protein is hPrp8p. As expected, the 220-kD protein that cross-linked to the 5’ splice site prior to lariat formation (Fig. 2B, lanes 7,8) is also hPrp8p (Fig. 3C, lane 5). No proteins were immunoprecipitated with hPrp8p antiserum from cross-linking reactions containing control mRNA (Fig. 3C, lane 2). In summary, the IP experiments indicate that proteins that cross-link to the PIP:E1(B) mRNA exon–exon junction dependent on splicing are part of a tight RNase-resistant complex. Additionally, two of these proteins are SRm160 and hPrp8p. The splicing-dependent ~50 and 20 kD proteins remain unidentified.

**Cross-linking to other exon–exon junctions**

To determine whether the pattern of splicing-dependent cross-linked proteins obtained with PIP:E1(B) pre-mRNA was unique to that construct or more general, we undertook additional cross-linking analyses with two other experimental pre-mRNAs, PIP:E1(S) and PIP:E2(S), and

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**Figure 3.** IP of proteins cross-linked to PIP:E1(B) RNAs. (A) IP of native complexes with mAb NM4. RNAs were incubated under splicing conditions for 90 min, irradiated, and treated with RNase without IP (+) [lanes 1, 5, and 9], treated with RNase after (+) IP [lanes 2,6,10], or treated with RNase before (+) IP [lanes 3,4,7,8,11,12]. IPs in lanes 4, 8, and 12 contained no antibody. (●) Indicates position of the ~70-kD protein discussed in text. (B) IP of native and denatured proteins with mAb NM4 or B1C8. Experimental pre-mRNA was spliced for 90 min, irradiated, and treated with RNase. Lanes 1 and 2 are shorter exposures of lanes 3 and 4. Samples in lanes 1 and 3 were not further purified. IPs with either mAb NM4 [lanes 2,4–6] or mAb B1C8 [lanes 7–9] were performed without prior protein denaturation (−) or after boiling in either 0.05% SDS (lanes 5,8) or 0.15% SDS (lanes 6,9). (C) IP of denatured proteins with hPrp8p antiserum or preimmune serum. After incubation of RNAs under splicing conditions for 90 min, irradiation, and RNase treatment, proteins were denatured by boiling in 0.25% SDS. Samples in lanes 1, 4, and 7 were not further purified. IPs were performed with either hPrp8p antiserum [lanes 2,5,8] or preimmune serum [lanes 3,6,9].
appropriate controls (Fig. 4). Relative to PIP:E1(B), both new constructs contained a different sequence at the 3’ end of exon 1, and a different photoreactive moiety, 4-thio-dU [S]. Compared to the benzophenone moiety, which was attached to the RNA via a long linker arm (MacMillan et al. 1994), 4-thio-dU has a much shorter reach and should therefore only react with proteins in close proximity to the RNA (Sontheimer 1994; Moore and Query 1998). PIP:E1(S) and PIP:E2(S) differed in the position of the 4-thio-dU, which was located at the 3’ end of exon 1 (E1) or the 5’ end of exon 2 (E2), respectively (Fig. 4A).

Both new pre-mRNAs spliced with similar kinetics to PIP:E1(B) pre-mRNA (data not shown). Like PIP:E1(B), the pattern of proteins that cross-linked to the exon–exon junctions of both PIP:E1(S) and PIP:E2(S) mRNAs was dependent on whether or not the mRNA was generated by splicing [Fig 4B]. Overall, the cross-linking patterns observed with experimental and control PIP:E1(S) RNAs were quite similar to those obtained for their respective PIP:E1(B) constructs (cf. Fig. 4B, lanes 1–3, with Figs. 2 and 3). Proteins migrating with apparent molecular weights of ~220, 160, and 20 kD whose cross-linking depended on splicing were clearly observable [Fig. 4B, cf. lane 3 with lanes 1 and 2]. Because the 220-kD protein also cross-linked to control PIP:E1(S) pre-mRNA [Fig. 4B, lane 2], this protein is most likely hPrp8p. However, there was no discernible protein of 50 kD that specifically cross-linked to experimental PIP:E1(S) RNA. It is possible that the 50-kD protein that cross-linked to experimental PIP:E1(B) RNA is relatively distant from the RNA or is sequence-specific. When the 4-thio-dU was moved to exon 2 [PIP:E2(S)], the overall cross-linking pattern was noticeably different. Although a splicing-specific p220 was still observed, another splicing-dependent protein migrated at ca. 55 kD.

These results indicate that splicing alters the complement of proteins associated with exon–exon junctions regardless of the nature of the photoreactive group, the position of this group relative to the exon–exon junction, or the exact sequence of the 5’ exon, even though changing the position or the nature of the photoreactive group can affect exactly which proteins are detected by this method.

Glycerol gradient fractionation

Although the experiments presented above identified proteins that cross-link to mRNA specifically as a consequence of splicing, they did not reveal which, if any, remain associated with mRNA after its release from the spliceosome. Previous studies have shown that mRNPs can be separated from spliceosomes by glycerol gradient fractionation (Cheng and Abelson 1987; Konarska and Sharp 1987). To determine the sedimentation profile of PIP RNAs and cross-linked proteins, experimental PIP:E1(B) pre-mRNA was incubated under splicing conditions for 2 hr, UV irradiated, and fractionated in a 10%–30% glycerol gradient. Fractions were then divided and the RNAs and cross-linked proteins analyzed in appropriate denaturing gels (Fig. 5A).

Splicing marks mRNP exon–exon junctions

![Figure 4. Protein cross-linking to 4-thio-dU-containing PIP RNAs. (A) Schematic structures of the three experimental PIP pre-mRNAs and sequences around the 5’ and 3’ splice sites. (Star) Positions of benzophenone [B], 4-thio-dU [S] and 32P. Underlined nucleotides correspond to those differing between constructs. (B) RNAs as indicated were incubated under splicing conditions for 90 min, UV irradiated, and treated with either RNase A or T1 [see Materials and Methods]. Proteins that cross-linked in a splicing-dependent manner are specified to the right of each panel.](https://genesdev.cshlp.org/1103/3/GENES-DEVELOPMENT-1103.png)

Spliceosomes sedimented toward the bottom of the gradient [fractions 8–15] and were defined by the presence of both pre-mRNA and lariat intermediate, a product of the first step of splicing [Fig. 5A, bottom]. The mRNA migrated as a broad peak toward the top of the gradient [fractions 3–8], indicating that nearly all of it had been released from the spliceosome. Notably, two of the mRNA-containing fractions were nearly free of both pre-mRNA and splicing intermediates [fractions 3 and 4]. All proteins that cross-linked independent of splicing cosedimented with mRNA. Cross-linked SRm160, p50, and p20 also cosedimented with mRNA. In contrast, cross-linked hPrp8p seemed to sediment in two peaks of
approximately equal intensity, one comigrating with released mRNA and another with intact spliceosomes. This suggests that only a portion of mRNA that was cross-linked to hPrp8p was released from the spliceosome. Furthermore, because only a small fraction of total mRNA cosedimented with spliceosomes, cross-linking of hPrp8p may be significantly more efficient prior to mRNA release. Interestingly, however, Northern blot analysis demonstrated that the peak of cross-linked hPrp8p that did cofractionate with released mRNA sedimented at a slightly lower velocity than the peak of U5 snRNA [Fig. 5A, cf. U5 snRNA, mRNA, and cross-linked hPrp8p]. This raised the intriguing possibility that hPrp8p cross-linked to released PIP mRNA was not as-sociated with fully intact U5 snRNP.

In theory, cross-linked proteins cosedimenting with released mRNA could be true components of PIP mRNP or could be present only because their association with the mRNA had been artificially stabilized by cross-linking. This issue was particularly pertinent to hPrp8p because U5 snRNA has been shown to remain associated with the spliced intron product in a spliceosome-sized complex after mRNA release [Cheng and Abelson 1987; Konarska and Sharp 1987]. To differentiate between these possibilities, we repeated the sedimentation analysis except that cross-linking was performed after sedimentation [Fig. 5B]. Under these conditions, cross-linked hPrp8p exhibited a bimodal distribution similar to that observed in Figure 5A. Similarly, cross-linked SRm160 and p50 cosedimented with mRNA as above. However, the distribution of cross-linked p20 was somewhat altered by the fractionation procedure, with its peak intensity moving to slightly higher velocity fractions. This suggests that p20 is not associated as stably with spliceosome-released mRNA as the other splicing-specific proteins. That protein–RNA associations could change within the gradient was confirmed by the presence of a strong, but previously unobserved, cross-link evident at ca. 75 kD [Fig. 5B, dot]. Thus, association of PIP mRNA with a previously undetected protein became apparent during sedimentation.

Taken together, results obtained by glycerol gradient fractionation indicate that hPrp8p, SRm160, and p50 are all genuine components of PIP mRNP produced in vitro.

Discussion

We provide evidence that some of the proteins that associate with mRNA exon–exon junctions in vitro do so only as a consequence of splicing. Several of these proteins form a stable complex that survives RNase treatment. Below we discuss how the identified proteins relate to the process of splicing and the current picture of mRNP structure and function.
Properties of proteins associated with exon–exon junctions specifically as a consequence of splicing

We observed four proteins that cross-linked to the exon–exon junction of PIP:E1(B) mRNA in a splicing-dependent manner. So far, we have identified two of these as SRm160 and hPrp8p. SRm160 is a nuclear matrix antigen [Blencowe et al. 1995, 1998] that contains multiple RS repeats but, unlike prototypic SR proteins [Zahler et al. 1992], lacks an RNA recognition motif [RRM]. However, SRm160 had not been shown previously to cross-link to RNA [Blencowe et al. 1998]. Instead, previous studies showed that SRm160 forms a complex with another nuclear matrix antigen, SRm300, and this complex serves to promote splicing through interactions with SR protein family members. Notably, association of SRm160/300 with specific pre-mRNAs is dependent on SR proteins and U1 snRNP and is stabilized by U2 sn-RNP [Blencowe et al. 1998; Eldridge et al. 1999]. This is consistent with our results showing that SRm160 only cross-links to an exon–exon junction that has been formed by the spliceosome. Moreover, our results suggest that SRm160 becomes closely associated with exonic RNA only after splicing, because SRm160 did not cross-link to the control pre-mRNA that allowed detection of protein–RNA interactions prior to lariat formation [Fig. 2B]. If the 160-kD cross-linked protein observed with experimental PIP:E1[S] RNA [Fig. 4B] is also SRm160, then it is closely associated with the mRNA and its ability to cross-link to the 5′ side of the exon–exon junction is not highly sequence-dependent.

hPrp8p cross-linked to the 5′ exon both before lariat formation and after exon ligation [Fig. 2B; data not shown]. It also cross-linked to the 3′ exon after exon ligation in the PIP:E2[S] construct [Fig. 4B]. This remarkably conserved U5 snRNP protein [Hodges et al. 1995; Luo et al. 1999] is a core component of the spliceosome that enters splicing complexes as part of the U4/U6:U5 tri-snRNP. Therefore, its association with the spliceosome substrates, intermediates, and products requires spliceosome assembly. The many documented interactions between Prp8p and sites in both pre-mRNA and snRNAs important for the transesterification reactions implicate Prp8p as a key active site component [Reyes et al. 1996; Collins and Guthrie 1999; Siatecka et al. 1999 and references therein]. Previous cross-linking studies have placed Prp8p at the 5′ splice site before lariat formation, consistent with our observations with the control pre-mRNA [Fig. 2B], as well as in the vicinity of the 3′ splice site before exon ligation [Wyatt et al. 1992; Teigelkamp et al. 1995; Umen and Guthrie 1995; Chiara et al. 1996, 1997; Reyes et al. 1996]. Our results now indicate that some of these interactions are maintained after exon ligation and can persist in the mRNP complex after spliceosome release (see below).

In our cross-linking reactions, we also observed numerous proteins that associated with the mRNA independent of splicing. So far, we have only identified one as SRp20 [Zahler et al. 1992; Neugebauer and Roth 1997]. This is consistent with previous studies showing that recombinant SRp20 can bind RNA independent of other splicing components [Cavaloc et al. 1999, Schaal and Maniatis 1999].

Some splicing-dependent proteins remain associated with PIP mRNA after spliceosome release

To determine whether any of the cross-linked proteins observed here were candidates for bona fide mRNP components, we also examined their association with PIP mRNA that had been released from the spliceosome. It was established previously that mRNA release is an active process requiring ATP hydrolysis by the Prp22/HRH1 RNA helicase [Company et al. 1991; Ohno and Shimura 1996; Schwer and Gross 1998; Wagner et al. 1998]. Release occurs efficiently in HeLa-cell splicing reactions, and the resultant mRNP can be separated from spliceosomes by glycerol gradient fractionation [Konarska and Sharp 1987]. Because most of the PIP:E1[B] mRNA present in splicing reactions after 2 hr had been released from spliceosomes [Fig. 5], it seemed likely that at least some of the proteins that cross-linked to the exon–exon junction at this time were mRNP components. This was clearly true for SRm160 and p50, which cosedimented exclusively with free mRNP regardless of whether cross-linking was performed before or after sedimentation [Fig. 5, cf. A and B].

In contrast, cross-linked hPrp8p was detected both in fractions containing spliceosomes and in fractions containing mRNP [Fig. 5A,B]. The presence of hPrp8p in mRNP was unexpected because it is known to interact directly with U5 snRNA [Dix et al. 1998], which remains bound to the spliced intron product after mRNA release (Cheng and Abelson 1987; Konarska and Sharp 1987). However, our observation that hPrp8p cross-linked to PIP mRNA even after glycerol gradient fractionation indicates that it is a component of the PIP mRNP. This mRNP is of somewhat lower density than free U5 snRNP, opening the possibility that hPrp8p in mRNP is no longer associated with fully intact U5 snRNP. Interestingly, upon U5 snRNP dissociation in vitro, hPrp8p can be found in RNA-free complexes with other U5 snRNP proteins including a 200-kD RNA helicase, a 116-kD EF-2 homolog, and a 40-kD WD-repeat-containing protein [Achsel et al. 1998]. Thus, it is possible that these other proteins are also mRNP components.

Evidence for a stable protein complex at the mRNA exon–exon junction

All four proteins that cross-linked to PIP:E1[B] mRNA in a splicing-dependent manner were immunoprecipitable with mAbs NM4 and B1C8, both before and after RNase treatment [Fig. 3A,B]. Among these proteins, hPrp8p does not contain an epitope recognized by these mAbs, and p50 and p20 also likely lack an appropriate epitope because they failed to be precipitated when both RNase digestion and protein denaturation preceded the IP. This suggests that all are part of a complex that remains
bound to the exon–exon junction as a signature of splicing. The glycerol gradient fractionation results indicate that at least hPrp8p, SRm160, and p50 remain stably associated with PIP:E1(B) mRNA after its release from the spliceosome (Fig. 5A,B).

While it remains to be determined whether or not the proteins we identified here typify all mRNPs, the type of complex observed could serve as a general mark of exon–exon junctions. It will now be of interest to determine how long these proteins remain associated with mRNA. Interestingly, SRm160 can be found in both nuclear and cytoplasmic fractions and can be visualized in nuclear tracks that branch and often terminate near nuclear pore complexes [J. Nickerson, pers. comm.]. This suggests that SRm160 may maintain its mRNA association even after mRNA export to the cytoplasm. Identification and functional characterization of other signature proteins will help to extend our understanding of how pre-mRNA splicing influences downstream mRNA metabolic events in vivo.

Materials and methods

**Synthesis of doubly modified RNAs**

The PIP:E1(B) experimental pre-mRNA was synthesized by splinted ligation of four separate RNA pieces [Moore and Query 1998]; [1] a GpppG-capped transcript corresponding to the 5' end of E1; [2] a synthetic oligomer containing a convertible adenosine near its 3' end (MacMillan et al. 1994); [3] a transcript comprising the entire intron, and [4] a 5'-32P-labeled transcript comprising E2. Each fragment is indicated within the pre-mRNA sequence below (where bold nucleotides represent exons, the asterisk specifies the convertible adenosine, underlining indicates intronic splice and branch site consensus sequences, and slashes separate the four fragments), which derived from PIP85.B (Query et al. 1994): 5'-GpppGGCGAUAUCUGCACUCUCUCUC CGACUCUGUCUCGAGGUAUCCACUGGCG/GGUUGU CGC(A*G)/GUGAUGAUAGUGUCCCUUAAAGGCGGC CGUGACUCUGAGCCAGGUGUUCGAGGCUU UCUCUGACGUGUACGUGCCUGGCGGCGUGCGU CUCUCUUCUCUCUCUCUCUCUCUCUCACAG/15PGUCCX CACAAACAACACGACAAAACUCUCGCGGCUCC UC GCAUGCAACGUU-3'. Following purification of full-length pre-mRNA, benzophenone derivatization of the convertible nucleotide was performed as described previously [MacMillan et al. 1994; Moore and Query 1998]. Control PIP:E1(B) pre-mRNA was generated by three-way ligation of fragments 1, 2, and a single 5'-32P-labeled transcript comprising fragments 3 and 4. Control PIP:E1(B) mRNA was synthesized by three-way ligation of fragments 1, 2, and a 5'-32P-labeled fragment 4. All three RNAs were also synthesized with an unmodified version of fragment 2.

Other experimental pre-mRNAs and corresponding controls were generated similarly. The relevant sequences around the 5' and 3' splice sites, as well as the position of the photoactive group (4-thio-dU; Glen Research) and 15P are indicated in Figure 4A for each experimental pre-mRNA.

**Nuclear extracts and splicing reactions**

HeLa-cell nuclear extracts [Dignam et al. 1983] were prepared with modifications described [Abmayr et al. 1988] from cells grown by Cellbi Biosciences. Splicing reactions [20 µl] containing ~10 fmole of labeled RNA were carried out in 40% nuclear extract, 2 mM MgOAc2, 20 mM potassium glutamate, 1 mM ATP, 5 mM creatine phosphate, and 0.05 mg/ml of *E. coli* tRNA.

Following incubation at 30°C for 5 min at 30°C without ATP and creatine phosphate prior to pre-mRNA addition, or by adding 5 µm of a 2'-O-methyl oligo-

nucleotide [5'-AGAUCACACUGUAAC-3'] complementary to nucleotides 27–41 of human U2 snRNA [Luo et al. 1999] and incubating for 10 min at 30°C prior to pre-mRNA addition. For RNA analysis, reactions were quenched with 10 volumes of splicing stop buffer [100 mM Tris at pH 7.5, 10 mM EDTA, 1% SDS, 150 mM NaCl, 300 mM NaO acetate], phenol/chloroform extracted and ethanol precipitated. RNAs were separated in 15% denaturing polyacrylamide gels and visualized by autoradiography or with a Molecular Dynamics PhosphorImager.

**Cross-linking**

Photoreactive substrates of high specific activity (8000 cpm/fmole) were incubated with splicing conditions. Cross-linking was performed on ice by irradiation for 20 min with a 302 or 365 nm hand-held lamp (UltraViolet Products) for benzophenone- and 4-thio-dU-derivatized RNAs, respectively. Reactions were digested with either 0.1 mg/ml RNase A [Sigma; for PIP:E1(B) and PIP:E2(S) RNAs] or 1 U/ml RNase T1 [Sigma; for PIP:E1(S) RNAs] for 30 min at 37°C. For RNase treatment after protein denaturation, samples were boiled for 2 min in the presence of 0.5% SDS prior to RNase addition. Proteinase K treatment was performed in 0.5% SDS after both cross-linking and RNase digestion by adding proteinase K to 1 mg/ml (Boehringer) and incubating for 5 min at 65°C and then 30 min at 30°C. Cross-linked proteins were separated in SDS–16% polyacrylamide (20:1 acrylamide:bis) gels or in SDS–7.5% and SDS–10% polyacrylamide (29:1 acrylamide:bis) and detected by autoradiography or PhosphorImaging. 14C-Labeled protein molecular weight standards [GIBCO BRL] were run as controls.

**Immunoprecipitations**

Antibodies were bound to protein A–Sepharose [PAS] beads [Pharmacia] via rabbit anti-mouse IgG + IgM [Pierce] for NM4, B1C8, and SRp20 antisera, and directly to the beads for hPrp8 antisera and preimmune serum. For IPs, 20 µl of cross-linked samples were diluted with 200 µl of IP100 [Luo et al. 1999] and combined with 40 µl of 50% slurry PAS-bound antibodies. Samples were denatured by boiling for 2 min in 0.05%, 0.15%, or 0.5% [wt/vol] SDS prior to dilution and IP, where indicated. Cross-linked proteins were bound over 3 hr at 4°C with gentle mixing, after which beads were washed three times with IP100, or twice with IP150-1 M urea [Luo et al. 1999] and once with IP100 for hPrp8 IPs.

**Glycerol gradient analysis**

Splicing reactions [30 µl] were incubated for 2 hr at 30°C, supplemented to 0.5 mg/ml heparin, and further incubated for 5 min at 30°C. Reactions were cross-linked either before or after sedimentation through a 10%–30% glycerol gradient [Beckman]. Gradients were manually fractionated into 16 × 25 µl aliquots. Typically, 5 µl of...
Splicing marks mRNP exon-exon junctions

each was extracted with phenol/chloroform, ethanol precipitated, and used for RNA analyses. For Northern analysis, RNAs were separated in an 8% denaturing polyacrylamide gel, transferred to an ICN nylon membrane, and hybridized with a probe complementary to U5 snRNA (Konarska and Sharp 1987). The remainder of each fraction was treated with RNase A, and cross-linked proteins were separated and detected as described above.

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Pre-mRNA splicing alters mRNP composition: evidence for stable association of proteins at exon–exon junctions

Hervé Le Hir, Melissa J. Moore and Lynne E. Maquat

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