ATPγS stalls splicing after B complex formation but prior to spliceosome activation

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ABSTRACT

The ATP analog ATPγS inhibits pre-mRNA splicing in vitro, but there have been conflicting reports as to which step of splicing is inhibited by this small molecule and its inhibitory mechanism remains unclear. Here we have dissected the effect of ATPγS on pre-mRNA splicing in vitro. Addition of ATPγS to splicing extracts depleted of ATP inhibited both catalytic steps of splicing. At ATPγS concentrations ≥0.5 mM, precatalytic B complexes accumulate, demonstrating a block prior to or during the spliceosome activation stage. Affinity purification of the ATPγS-stalled B complexes (BγS) and subsequent characterization of their abundant protein components by 2D gel electrophoresis revealed that BγS complexes are compositionally more homogeneous than B complexes previously isolated in the presence of ATP. In particular, they contain little or no Prp19/CDC5L complex proteins, indicating that these proteins are recruited after assembly of the precatalytic spliceosome. Under the electron microscope, BγS complexes exhibit a morphology highly similar to B complexes, indicating that the ATPγS-induced block in the transformation of the B to B∗ complex is not due to a major structural defect. Likely mechanisms whereby ATPγS blocks spliceosome assembly at the activation stage, including inhibition of the RNA helicase Brr2, are discussed. Given their more homogeneous composition, B complexes stalled by ATPγS may prove highly useful for both functional and structural analyses of the precatalytic spliceosome and its conversion into an activated B∗ spliceosomal complex.

Keywords: pre-mRNA splicing; spliceosome; small molecule; ATPγS; Prp19/CDC5L complex

INTRODUCTION

Pre-mRNA splicing is catalyzed by the spliceosome, a dynamic highly complex RNP machine. Assembly of the spliceosome occurs by the stepwise interaction of the U1, U2, U4/U6, and U5 snRNPs, plus many additional non-snRNP proteins with the pre-mRNA (for review, see Wahl et al. 2009; Will and Lührmann 2011). The spliceosome carries out two successive transesterification reactions that lead to removal of an intron and the ligation of its flanking exons. Spliceosome assembly initiates by the ATP-independent association of U1 with the 5′ splice site (ss) and initial docking of the U2 snRNP, forming the spliceosomal E complex (Michaud and Reed 1991). Stable integration of the U2 snRNP via its interaction with the branch site (BS) of the pre-mRNA’s intron, generates the A complex. The recruitment and stable integration of the U4/U6,U5 tri-snRNP to the spliceosome follows, yielding the precatalytic B complex which does not yet have an active center. During the subsequent catalytic activation of the spliceosome, which generates the B∗ complex, the extensively base paired U4/U6 snRNAs are unwound, thereby releasing U4 and allowing the formation of new U2/U6 base-pairing interactions and a catalytically important U6 internal stem–loop (ISL) (Valadkhan 2013). Concomitant with or prior to this, the base-pairing interaction between the pre-mRNA’s 5′ss and the U1 snRNA must be disrupted, in order to allow base-pairing between U6’s highly conserved ACAGA sequence and the 5′ end of the intron (Staley and Guthrie 1998). This newly formed U2–U6-pre-mRNA RNA interaction network comprises the catalytic centre of the spliceosome (Valadkhan 2013). After additional RNP rearrangements, the catalytically active B∗ complex forms and catalyzes the first transesterification reaction of splicing. This involves cleavage at the 5′ss and ligation of

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the 5′ end of the intron to the branch site, and generates the splicing intermediates, the excised 5′ exon and the lariat-shaped intron-3′ exon. The subsequently formed C complex then catalyzes the second transsterification reaction of splicing, involving excision of the intron and ligation of the 5′ and 3′ exons to form the mRNA.

Over 170 proteins are associated at one time or another with the human splicing machinery and there is a dramatic exchange of proteins during the transitions from one spliceosomal complex to the next (Wahl et al. 2009; Wahl and Lührmann 2015). Our knowledge about the spliceosome’s highly complex compositional dynamics is, however, hampered by the limited number of distinct spliceosomal complexes that can be biochemically detected or isolated. Given the extensive exchange of proteins during the transitions from the B to B act , and B act to C complex, there are likely to be numerous additional, intermediate assembly stages in which only a subset of these proteins are released or recruited. Currently, many preparations of purified human spliceosomal complexes likely consist of a mixture of assembly stages, which prevents a clear picture of when some spliceosomal proteins are first recruited during assembly. To overcome this problem, small molecule inhibitors of splicing have more recently been used with the hope of obtaining more homogeneous populations of splicing complexes by stalling splicing at a highly specific stage (Kuhn et al. 2009; Berg et al. 2012; Effenberger et al. 2013, 2015; Pawallek et al. 2015).

ATP is required for each step of spliceosome assembly/disassembly and the catalytic steps of splicing, except for the association of U1 and U2 during the earliest stage of spliceosome assembly (i.e., E complex formation). However, the transsterification reactions per se do not consume ATP. At least eight conserved ATP-dependent DExD/H-box RNA/RNP remodeling enzymes drive the multiple rearrangements of the spliceosome during its working cycle and account for the ATP dependence of most structural rearrangements (Cordin and Beggs 2013; Liu and Cheng 2015). The DEAD-box helicases Sub2 and Prp5 are required for stable U2 snRNP integration during A complex assembly. During B complex formation, the DEAD-box protein Prp28 catalyzes the exchange of U1 with U6 at the 5′ss (Staley and Guthrie 1999), and during the transition from complex B to B act , the DEAD/H-box protein Brr2 of the Ski-2 helicase family catalyzes a key step in spliceosome activation, namely the unwinding of U4/U6 snRNA (Laggerbauer et al. 1998; Raghunathan and Guthrie 1998). The DEAD/H-box protein Prp2 promotes an RNP remodeling event that converts B act into the catalytically active B* complex (Kim and Lin 1996; Warkocki et al. 2009; Lardelli et al. 2010). The remaining DEAD/H-box proteins Prp16, Prp22 and Prp43 promote conformational rearrangements during the catalytic steps of splicing and/or are involved in spliceosome disassembly (for review, see Cordin and Beggs 2013; Liu and Cheng 2015).

ATP is required not only for the ATPases/RNA helicases that drive the conformational rearrangements in the spliceosome, but also for the many kinases and phosphatases involved in the phosphorylation/dephosphorylation of numerous spliceosomal proteins. For example, SR protein phosphorylation/dephosphorylation is essential for premRNA splicing (for review, see Misteli 1999; Soret and Tazi 2003), with SR protein phosphorylation shown to be required during spliceosome assembly (Mermoud et al. 1994; Xiao and Manley 1997). Dephosphorylation of several spliceosomal phosphoproteins is required for the catalytic steps of splicing, with PP1/PP2A phosphatases playing key roles at this stage (Mermoud et al. 1992; Shi et al. 2006).

The small molecule ATP analog ATPγS has been used to study the requirement for protein dephosphorylation during splicing, as thiophosphorylated proteins are poorly dephosphorylated by protein phosphatases (Li et al. 1988). Kinases can use ATPγS to thiophosphorylate proteins, making them resistant to rapid dephosphorylation. ATPγS can be hydrolyzed by some enzymes but usually at a much lower rate compared with ATP (Yount 1975). Addition of ATPγS to in vitro splicing reactions was initially reported to stall splicing during the second catalytic step (Tazi et al. 1992). In contrast, subsequent studies indicated that thiophosphorylation of the U1-70K protein (Tazi et al. 1993) or SR proteins (Cao et al. 1997), which were added back to nuclear or S100 extract lacking these proteins, blocked splicing prior to the first step of splicing, but after the B complex stage. However, the precise stage at which spliceosome assembly is blocked was not clear.

Here we have analyzed the effect of ATPγS on splicing in vitro, and characterized the composition and structure of splicing complexes that accumulate in the presence of ATPγS. Our studies indicate that ATPγS blocks splicing just prior to the action of Brr2, at a stage after B complex formation but prior to U4/U6 unwinding during spliceosome activation. In contrast to previous preparations of the human B complex, which consist of a mixture of assembly states due to the fact that they are kinetically stalled, B complexes that accumulate in the presence of ATPγS are essentially devoid of Prp19/CDC5L complex proteins and thus more homogeneous. Thus, our data indicate that highly pure B complexes lack the latter proteins and therefore that the hPrp19/CDC5 proteins first interact during B act formation, after/concomitantly with dissociation of U4. Taken together, our data also reveal that ATPγS can be used as an efficient small molecule inhibitor of splicing to generate spliceosomal complexes stalled at a very specific stage of the assembly process.

RESULTS AND DISCUSSION

To elucidate its effect on pre-mRNA splicing in vitro, we added increasing amounts of ATPγS to a splicing reaction containing HeLa nuclear extract and 32P-labeled MINX premRNA with MS2 aptamers for affinity purification (Fig. 1).
The extract was first depleted of ATP by preincubating with 2 mM glucose for 30 min at 30°C (Michaud and Reed 1991), which abolished both steps of splicing (Fig. 1A, compare lanes 16–18 with 19–21 where ATP/creatine phosphate, CP, were subsequently added), and inhibited splicing complex formation (Fig. 1B). Both catalytic steps of splicing were completely inhibited at all concentrations of ATPγS tested (0.25–4.0 mM), even after incubating for 3 h (Fig. 1A) or 6 h (data not shown). Addition of low concentrations (0.25–1.0 mM) of ATPγS led to an accumulation of A complexes, whereas at higher concentrations spliceosomal complexes migrating like a B complex accumulated (Fig. 1B). The accumulation of B complexes was specific for ATPγS, as the addition of the nonhydrolyzable AMP-PNP, which also inhibited splicing completely, abolished both A and B complex formation at 2 mM (Fig. 1C, lanes 10–12 and Fig. 1D, lanes 16–20).

These results suggest that the accumulation of precatalytic splicing complexes in the presence of ATPγS results either from the thiophosphorylation of one or more spliceosomal proteins or the inhibition of one or more ATP-dependent enzymes involved in the splicing process. Previous studies indicated that ATPγS blocks in vitro splicing first during the catalytic steps of splicing (Tazi et al. 1992). However, the splicing reactions performed by Tazi and colleagues contained creatine phosphate (CP) in addition to ATPγS; thus most of the hydrolyzed ATP analog could be regenerated to ATP, which could explain why their splicing reactions were not blocked at an earlier stage similar to ours. Indeed, thiophosphorylation of U1-70K, which would occur under our conditions, was shown to block splicing prior to step 1 but
affter assembly of the spliceosome (Tazi et al. 1993), consistent with ATPγS affecting splicing before the catalytic steps.

**Affinity purification of A complexes stalled with ATPγS**

To characterize the spliceosomal complexes stalled by ATPγS in more detail we affinity purified them by MS2 affinity selection. Complexes were assembled in splicing extract depleted of ATP and supplemented with 0.4 mM (for isolation of A complexes) or 2 mM ATPγS for the isolation of stalled B complexes. Splicing complexes were purified from the splicing reaction by MS2 affinity selection and then subjected to glycerol gradient centrifugation.

Two main peaks were observed in the presence of 0.4 mM ATPγS (Fig. 2A). The faster migrating peak contained complexes with equimolar amounts of the MINX pre-mRNA and U1 and U2 snRNPs as evidenced by their RNA composition (Fig. 2B), indicating that they are spliceosomal A complexes. Proteins of these affinity-purified complexes were

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**FIGURE 2.** Affinity purification of A complexes accumulating in the presence of ATPγS. (A) Splicing complexes were allowed to form on 32P-MINX pre-mRNA containing MS2 aptamers bound by MS2-MBP protein, in HeLa nuclear extract containing 0.4 mM ATPγS. Complexes were subsequently affinity purified on an amylose column and the eluate subjected to glycerol gradient centrifugation in buffer containing 150 mM NaCl. The number of cpm in each gradient fraction is plotted. (B) RNA composition of complexes in the faster migrating peak (fraction #13). (C) Proteins from affinity-purified A complexes formed in the presence of ATPγS (fractions #13–14) were separated by 2D gel electrophoresis, stained with RuBPS, and the identities of protein spots were determined by mass spectrometry. Proteins smaller than 25 kDa were not analyzed. Abundant proteins are indicated in black and less abundant ones in gray. (D) Summary of abundant/moderately abundant proteins identified in AATPγS complexes.
subsequently analyzed by 2D gel electrophoresis followed by mass spectrometry (MS) to identify abundant components (Fig. 2C.D). Abundant or moderately abundant proteins (with a molecular mass above 25 kDa) in the A complexes stalled by ATPγS (henceforth denoted AATPγS) included: (i) all 12S U1 and 17S U2 proteins, (ii) the 17S U2-related proteins U2AF65, U2AF35, and PUF60, which bind the polypyrimidine tract, (iii) the SR proteins SRSF1 and SRSF7, (iv) ASRB2 and finally S164/RBM25 (RNA-binding motif protein 25), which was shown to regulate the alternative splicing of the apoptotic factor Bcl-x pre-mRNA by promoting the recruitment of U1 snRNP to a weak 5′ss (Zhou et al. 2008). A highly similar set of abundant proteins was detected in affinity-purified A complexes formed in the presence of ATP (Agafonov et al. 2011). Thus, complexes with a protein composition generally characteristic for the A complex can be assembled with ATPγS as the sole energy source and these AATPγS complexes support efficient B complex formation if the concentration of ATPγS is above ~0.25 mM (Fig. 1B).

Phosphorylation of SR proteins is required for spliceosome assembly (Zhou and Fu 2013), and these results indicate that thio-phosphorylated SR (and other proteins) are still functional at this stage, consistent with previous observations (Tazi et al. 1993; Cao et al. 1997). Our results also indicate that low concentrations of ATPγS are sufficient for Prp5 and Sub2 activity at this stage of spliceosome assembly, consistent with a requirement for ATP binding, rather than ATP hydrolysis (or at least rapid hydrolysis) for their function in splicing. Indeed, DEAD-box proteins often can act after binding ATP (even without hydrolyzing it), which leads to a conformational change in the DEAD-box protein that can alter the local structure of bound RNA/RNPs (Cordin and Beggs 2013).

**Characterization of B complexes formed in the presence of ATPγS**

After MS2 affinity selection and subsequent glycerol gradient centrifugation, two peaks with S-values of ~40S and ~60S were consistently observed at an ATPγS concentration of 2 mM (Fig. 3A), where predominantly B complexes were observed on native agarose gels (Fig. 1B). The RNA composition of fractions comprising each peak was subsequently analyzed and revealed an identical snRNA composition. That is, complexes peaking in both regions of the gradient contained equimolar amounts of U2, U4, U5, U6, and MINX RNA, but almost no U1 snRNA, similar to affinity-purified B complexes generated by incubating the splicing reaction for 5–7 min (Agafonov et al. 2011). Analysis of the protein composition of the 40S and 60S complexes revealed a very similar protein composition (data not shown); thus, the faster migrating peak likely contains dimers of the B complexes that accumulate in the presence of ATPγS.

Abundant proteins present in the monomeric B complexes stalled in the presence of ATPγS (denoted BATPγS) were determined by 2D gel electrophoresis followed by mass spectrometry (MS) (Fig. 3C.D). Abundant or moderately abundant proteins (with a molecular mass above 25 kDa) in the BATPγS complexes included: (i) nearly all 17S U2, U5, and U4/U6 snRNP proteins, (ii) the tri-snRNP-specific proteins Snu66/110K and Sad1/65K, (iii) the so-called B-specific proteins hSmu1, MFAP1, RED, hSnu23, hPrp38, and FBP21, (iv) the Prp19 complex related proteins NPW38 and NPW38BP, (v) the 17S U2-related proteins U2AF65, U2AF35 and, as well as the DEXH-box protein Prp43, and (vi) the SR proteins SRSF1 and SRSF7. A very similar set of abundant proteins was previously reported for B complexes formed in the presence of ATP (Agafonov et al. 2011).

However, unlike kinetically stalled B complexes, BATPγS complexes contained little or no proteins of the Prp19/ CDC5L complex or Prp19-related proteins; only a low level of the Prp19 protein that is normally present in four copies in the Bact spliceosome was detected by 2D gel electrophoresis (Fig. 3C). Thus, BATPγS complexes appear to have a much more homogeneous protein composition than previously reported B complexes. Indeed, kinetically stalled B complexes (i.e., isolated after incubating for a defined period of time) most certainly contain a mixture of different assembly states of the spliceosome, leading to both compositional and structural heterogeneity. The near absence of Prp19/CDC5L proteins suggests that the vast majority of BATPγS complexes are stalled at a single assembly stage (i.e., prior to the docking/stable integration of the Prp19/CDC5L complex proteins) and/or that the presence of thio-phosphorylated spliceosomal proteins potentially inhibits the incorporation of Prp19/CDC5L complex proteins. In addition, our data indicate that the Prp19/CDC5L complex proteins are not required for the formation of B complexes that withstand gel electrophoresis in the presence of heparin (Fig. 1B) (i.e., highly stable B complexes), consistent with data showing the formation of heparin-resistant B complexes in extracts depleted of the Prp19/CDC5L complex (Makarova et al. 2004). Although proteins of the hPrp19/CDC5 complex are present in kinetically stalled B complexes, they are first enriched and stably associated in the spliceosomal Bact complex (Bessonov et al. 2010). Our data indicate that highly pure B complexes in fact do not contain these proteins and that the hPrp19/ CDC5 proteins first interact during Bact formation, after/concomitant with dissociation of U4.

**Structural characterization of the BATPγS complex by EM**

To determine whether ATPγS affects the structure of the B complexes that accumulate in its presence, we performed negative-stain electron microscopy (EM). Well-defined, single particles of similar shape and size were visible in the EM overview of the monomeric BATPγS complexes (Fig. 4A). Classification and class averaging of more than 10,000 single particle images revealed a triangular body comprised of a stump, a central mass and a foot, and an upper (head)
domain, which are connected via a slimmer neck domain (Fig. 4B,D). The overall structure of the BATPγ
\[\gamma\] complexes is indistinguishable from B complexes formed in the presence of ATP (Fig. 4C). Like wild-type, affinity-purified B complexes, almost identical views are seen in all class averages, indicating that the BATPγ complexes bind in a highly preferential orientation to the carbon film. Taken together these data indicate that the presence of ATPγS has little or no effect on the structural organization of the B complex. Thus, the ATPγS-induced block in the transformation of the B to Bact complex, leading to accumulation of B complexes, does not appear to be due to a major structural defect.

Our data indicate that B complex formation is not hindered if ATPγS is the sole source of energy in the in vitro splicing reaction, and is present at sufficient concentrations (i.e., >0.25 mM). The requirement for a higher concentration of ATPγS for B versus A complex formation may reflect the different ATPγ affinities of those enzymes required at each of these spliceosome assembly stages. The conversion of an A to B complex requires the action of the DEAD-box helicase Prp28, which exchanges U1 for U6 at the 5′ss (Staley and Guthrie 1999; Mathew et al. 2008). Phosphorylation of hPrp28 by the kinase SRPK2 also is required for B complex formation (Mathew et al. 2008). The tri-snRNP associated

FIGURE 3. Affinity purification of B complexes accumulating in the presence of ATPγS. (A) Splicing complexes were allowed to form on 32P-MINX pre-mRNA containing MS2 aptamers bound by MS2-MBP protein, in HeLa nuclear extract containing 2.0 mM ATPγS. Complexes were subsequently affinity purified on an amylose column and the eluate subjected to glycerol gradient centrifugation. (B) RNA composition of complexes migrating in the 60S and 40S peaks. (C) Proteins from ATPγS stalled complexes in the 40S gradient peak were separated by 2D gel electrophoresis, stained with Coomassie G-250, and the identities of protein spots were determined by mass spectrometry. (D) Summary of abundant/moderately abundant proteins present in monomeric BATPγ complexes.
factors hPrp31 and hPrp6 are phosphorylated by hPrp4 kinase at this stage and these post-translational modifications were also proposed to potentially contribute to stable tri-snRNP association during B complex formation (Schneider et al. 2010). Thus, one or more of these enzymes may require ATPγS concentrations above 0.25 mM to allow efficient B complex formation.

Like the DEAD-box proteins Sub2 and hPrp5, hPrp28 may not require rapid ATP hydrolysis for its function during splicing. However, in the presence of an ATPase deficient hPrp28 mutant, 37S human pre-B complexes accumulate (Boesler et al. 2016). The latter contain the U4/U6.U5 tri-snRNP, in addition to U1 and U2, but in contrast to B complexes, the tri-snRNP is not stably associated. Our data indicate, however, that the block in spliceosome assembly at low ATPγS concentrations occurs prior to the pre-B complex stage, as the vast majority of complexes that accumulate do not contain the tri-snRNP (Fig. 2). Thus, the block during the A to B complex transition must occur prior to the initial recruitment of the tri-snRNP.

As BγS complexes contain stoichiometric amounts of U4 snRNA (Fig. 3), ATPγS clearly blocks the transition from a B complex to a Bact complex during the activation stage, which is initiated by loss/distabilization of the U4 snRNA due to Brr2-mediated unwinding of the U4/U6 duplex present in the B complex. Our purified BγS complexes were also nearly devoid of the U1 snRNA (Fig. 3), suggesting that Prp28 has already facilitated the displacement of U1 from the 5′ss, leading to the base-pairing interaction between the U6 snRNA ACAGA box and intron nucleotides near the 5′ss. Thus, the action of these two spliceosomal helicases can apparently be uncoupled. The B complexes formed in the presence of ATPγS do not appear to be compositionally or structurally compromised. Thus, ATPγS most likely acts by inhibiting an enzyme required during the earliest stages of activation, most probably the ATP-dependent DEIH box RNA helicase Brr2. Indeed, previous studies have shown that Brr2 cannot use ATPγS for unwinding a U4/U6 duplex in vitro (Laggerbauer et al. 1998; Raghunathan and Guthrie 1998).

Small molecule inhibitors of splicing have proven to be valuable tools to study spliceosome assembly and function. To generate high-resolution structures of compositionally and conformationally dynamic, large (several MDa) ensembles such as the spliceosome, a major challenge is to obtain complexes that are biochemically and conformationally homogeneous. Our data show that ATPγS is an important tool to obtain more homogeneous B complexes, which are nearly devoid of Prp19 complex and Prp19-related proteins. This contrasts with those B complexes stalled by other small molecule inhibitors, such as SAHA and DHC (Kuhn et al. 2009). BγS complexes thus may prove highly useful for both functional and structural analyses of the precatalytic spliceosome, as well as its conversion into an activated Bact spliceosomal complex.

**MATERIALS AND METHODS**

**In vitro splicing and analysis of splicing complex formation**

Uniformly [32P]-labeled, m7G(5′)ppp(5′)G-capped MINX premRNA was synthesized in vitro by T7 runoff transcription and gel purified. HeLa nuclear extract was prepared essentially as previously described (Dignam et al. 1983). Nuclear extract was depleted of ATP by incubating in the presence of 2 mM glucose for 30 min at 30°C (Michaud and Reed 1991). A typical splicing reaction contained 10 nM of 32P-labeled pre-mRNA and 40% HeLa nuclear extract in buffer containing 3 mM MgCl2, 65 mM KCl, 20 mM HEPES-KOH pH 7.9, 2 mM ATP, and 2 mM creatine phosphate and was incubated at 30°C for the times indicated. Alternatively, instead of adding ATP and creatine phosphate, the reactions were supplemented with 0.25 to 4 mM ATPγS or 2 mM AMP-PNP. RNA was then isolated and analyzed by denaturing PAGE on a 6 M urea, 14% polyacrylamide gel, followed by autoradiography. Spliceosomal complexes were analyzed by agarose gel electrophoresis in the presence of TBE. As the bands corresponding to intermediates containing the tri-snRNP were broad and run well ahead of that containing the tri-snRNP, these complexes were not analyzed in detail. Spliceosomal complexes were not detected by this method, even when BATPγS complexes were prepared in the presence of 4 mM ATPγS, most likely due to the fact that BATPγS complexes are nearly devoid of Prp19 complex and Prp19-related proteins (Boesler et al. 2016).
of 0.65 μg/μL heparin (Das and Reed 1999) and bands were visualized with a Typhoon phosphoimager (GE Healthcare).

**MS2 affinity selection of splicing complexes**

Spliceosomal complexes were isolated by MS2 affinity selection. Briefly, MINX pre-mRNA containing three MS2 aptamers at its 3′ end was incubated with a 15-fold molar excess of MS2-MBP fusion protein and then added to a splicing reaction lacking ATP but supplemented with ATPγS. After incubating at 30°C for 180 min and centrifuging to remove aggregates, the reaction was loaded onto a MBP Trap HP column (GE Healthcare) after adding 100 mM NaCl. The column was washed with G-150 buffer (20 mM HEPES-KOH pH 7.9, 1.5 mM MgCl₂, 150 mM NaCl) and complexes were eluted with G-150 buffer containing 1 mM maltose. Eluted complexes were loaded onto a 14 mL linear 10%–50% (v/v) glycerol gradient containing G-150 buffer, centrifuged at 25,000 rpm for 15 h at 4°C in a Sorvall TST 41.14 rotor and fractions were harvested from the bottom. The distribution of 32P-labeled MINX RNA across the gradient was determined by Cherenkov counting. Peak fractions containing the spliceosomal complexes were pooled, and their RNA and protein compositions determined by denaturing PAGE and 2D gel electrophoresis, respectively.

**2D gel electrophoresis and mass spectrometry**

Two-dimensional gel electrophoresis of affinity-purified spliceosomal complexes was performed as described in Agafonov et al. (2011). For mass spectrometry, Coomassie-stained protein spots were cut out of the 2D gel and proteins were digested in-gel with trypsin and extracted. The extracted peptides were analyzed in a liquid chromatography coupled electrospray ionization quadrupole time of flight mass spectrometer (LTQ Orbitrap XL) under standard conditions. Proteins were identified by searching fragment spectra against the NCBI nonredundant (nr) database using Mascot as a search engine.

**Electron microscopy**

Purified spliceosomal complexes were loaded onto a 10%–30% (v/v) glycerol gradient in buffer G130 containing a linear gradient of 0%–0.1% glutaraldehyde (GraFix method) (Kastner et al. 2008). After centrifugation for 1:47 h at 60,000 rpm in a TH660 rotor, the gradient was harvested from the bottom in 120 μL fractions. Particles in peak gradient fractions were negatively stained with uranyl formate by the single-carbon film method adopted from Golas et al. (2003). Images were recorded using a CM 200-FeG electron microscope (FEI) at a magnification of 115,000× with a 4k × 4k digital camera (TVIPS). After performance of a reference-free alignment procedure (Dube et al. 1993), images were subjected to multivariate statistical analysis (MSA) and classification (van Heel and Frank 1981; van Heel 1984; Dube et al. 1993). The resulting class averages were used as reference images in subsequent rounds of alignment until the class averages were stable.

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