Endogenous U2·U5·U6 snRNA complexes in S. pombe are intron lariat spliceosomes

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ABSTRACT
Excision of introns from pre-mRNAs is mediated by the spliceosome, a multi-megadalton complex consisting of U1, U2, U4/U6, and U5 snRNPs plus scores of associated proteins. Spliceosome assembly and disassembly are highly dynamic processes involving multiple stable intermediates. In this study, we utilized a split TAP-tag approach for large-scale purification of an abundant endogenous U2·U5·U6 complex from Schizosaccharomyces pombe. RNAseq revealed this complex to largely contain excised introns, indicating that it is primarily ILS (intron lariat spliceosome) complexes. These endogenous ILS complexes are remarkably resistant to both high-salt and nuclease digestion. Mass spectrometry analysis identified 68, 45, and 43 proteins in low-salt-, high-salt-, and micrococcal nuclease-treated preps, respectively. The protein content of a S. pombe ILS complex strongly resembles that previously reported for human spliced product (P) and Saccharomyces cerevisiae ILS complexes assembled on single pre-mRNAs in vitro. However, the ATP-dependent RNA helicase Brr2 was either substoichiometric in low-salt preps or completely absent from high-salt and MNase preps. Because Brr2 facilitates spliceosome disassembly, its relative absence may explain why the ILS complex accumulates logarithmically growing cultures and the inability of S. pombe extracts to support in vitro splicing.

Keywords: intron lariat spliceosome; proteomic; RNAseq; spliceosome footprint; S. pombe

INTRODUCTION
Removal of introns from precursors to messenger RNAs (pre-mRNAs) is an essential step in eukaryotic gene expression. Introns are excised by the spliceosome, a multi-megadalton machine composed of five uridine-rich small nuclear RNAs (U1, U2, U4, U5, and U6 snRNAs) and scores of polypeptides. Many proteins stably associate with the snRNAs to form small ribonucleoprotein particles (snRNPs), while many others function as transiently interacting “splicing factors.” The snRNPs and splicing factors are thought to assemble de novo on each new intron via a highly dynamic process involving multiple stable intermediates. Recruitment of a U1 snRNP to the 5′ end of the intron (5′ splice site; 5′SS) and U2 snRNP to the branch site (BS) near the 3′ end of the intron (3′ splice site; 3′SS) results in pre-spliceosome (complex A) formation. Addition of the U5·U4/U6 tri-snRNP forms the pre-catalytic spliceosome (complex B), which is then converted to the activated spliceosome (Bact) by a major structural rearrangement involving destabilization of U1 and U4 snRNPs and addition of the multiprotein NineTeen Complex (NTC). Further structural rearrangements result in the catalytically active complexes (B∗, C1, and C2) in which the first (5′SS cleavage and lariat formation) and second (3′SS cleavage and exon ligation) chemical steps of splicing occur (Jurica and Moore 2003; Wahl et al. 2009; Hoskins and Moore 2012; Li et al. 2013). Spliced exon release from the spliced product complex (P) (Ilagan et al. 2013) results in formation of intron-lariat spliceosome complex (ILS) (Yoshimoto et al. 2009; Fourmann et al. 2013), containing the lariat intron, U2, U5, and U6 snRNAs and the NTC. ILS complex disassembly and subsequent reassembly of the U5-U4/U6 tri-snRNP completes the spliceosome cycle and readies the components for another round. Because each step in the spliceosome cycle involves the comings and goings of multiple components, each stable and isolatable complex has a signature RNA and protein composition (Jurica and Moore 2003; Wahl et al. 2009; Hoskins and Moore 2012).
To date, most studies of spliceosome composition and structure have been performed on individual snRNPs or stable intermediate complexes assembled in vitro on a limited set of well-spliced pre-mRNAs (Will and Luhrmann 2011). The latter complexes are usually obtained by artificially stalling spliceosome assembly or disassembly at a particular step (Jurica and Moore 2002; Jurica 2008). When endogenous spliceosomes have been isolated, they often contain protein components representative of multiple assembly states (Stevens 2000; Ohi et al. 2002; Chen et al. 2007). One notable exception is a stable endogenous complex containing U2, U5, and U6 snRNAs observed in logarithmically growing *Saccharomyces pombe* (Huang et al. 2002). Previous purification of this complex (a.k.a., *S. pombe* CDC5 complex) via a TAP-tag on CDC5 revealed it to contain numerous orthologs of proteins associated with human C complex (McDonald et al. 1999; Ohi et al. 2002; Ren et al. 2011) and to structurally resemble human C complexes upon single particle reconstruction of electron microscope images (Ohi et al. 2007).

Here, we report large-scale isolation of these endogenous *S. pombe* U2·U5·U6 complexes using double-affinity purification via separate affinity tags affixed to two different proteins. Purifications were carried out under different stringencies to reveal both loosely and tightly bound factors. Comprehensive RNA and protein compositional analyses reveal this complex to consist largely of endogenous ILS complex. The absence of SF3A/B components coupled with the micrococcal nuclease cleavage pattern of U2 snRNA reveals new information about spliceosome conformational and compositional changes following spliced exon release. Finally, the relative absence of proteins required for spliceosome disassembly may explain why this complex accumulates in rapidly dividing *S. pombe* and why extracts from these cells fail to support in vitro splicing.

**RESULTS**

**Split TAP-tag purification of endogenous *S. pombe* U2-U5-U6 complexes**

*S. pombe* cell lysates contain significant amounts of a U2-U5-U6 snRNA complex indicative of post-activation spliceosomes (Huang et al. 2002; Ohi et al. 2002; Ren et al. 2011). To specifically purify this complex, we utilized a split TAP-tag approach, with protein A attached to U2 snRNP protein Lea1 (U2 A' in human) and calmodulin binding peptide (CBP) attached to U5 snRNP protein Snu114 (U5 116K in human). An additional GFP tag on Cdc5 (an NTC component), Spp42 (U5 snRNP protein Prp8), or Br2 (another U5 protein) allowed for real-time monitoring of later purification steps, as well as assessment of the extent to which the GFP-tagged species copurified with Lea1 and Snu114 (Fig. 1A). All three tags were introduced by homologous recombination into haploid cells, with tagged proteins replacing their untagged counterparts. Growth rates and cell morphologies of all triple-tagged strains were indistinguishable from the parental untagged strain (data not shown), indicating functionality of the fusion proteins.

To maximize yields, cells were grown in 5× YES media and harvested in late log phase (OD600 ≈ 23). Following cell lysis, immunoglobulin G (IgG)-sepharose and calmodulin affinity steps were carried out as described (Materials and Methods). Physiological (low salt; LS) complexes were obtained from 400 mM KCl cell lysate and purified in 150 mM NaCl. Increasing lysate KCl to 700 mM significantly increased yields, but 1 M total salt resulted in very low yields (Supplemental Fig. S1A). Salt-stable core (high salt; HS) complexes were thus obtained from 700 mM KCl lysates and exposed to 700 mM NaCl during early purification stages to remove loosely bound factors. Treatment of these HS complexes with MNase (MN) during the IgG-
sepharose step removed solvent-accessible RNA sequences. In all three preps (LS, HS, and MN), total protein and CDC5-GFP fluorescence exhibited identical calmodulin elution profiles (Fig. 1B; Supplemental Fig. S1B). Generally, 6 L of culture yielded ~250 g packed cell pellet from which we were able to purify ~2 mgs of endogenous U2·U5·U6 snRNA complexes.

To assess complex integrity and heterogeneity, HS complexes were analyzed by dynamic light scattering (DLS) and negative stain electron microscopy (EM). DLS revealed one major species with an estimated diameter of 34 nm and MW of ~2.5 MDa (Fig. 1C). Consistent with this, electron micrographs (Fig. 1D) revealed largely monodispersed (>95% monomer, <5% dimer) 30- to 35-nm particles of uniform size and reproducible shape similar to human C complex (Jurica et al. 2004) and S. pombe CDC5 complex (Ohi et al. 2007). Negative stain EM images of LS and MN particles yielded similar results (Supplemental Fig. S1C).

Protein composition of endogenous U2-U5-U6 complexes

SDS-PAGE revealed nearly identical protein banding patterns for the LS, HS, and MN preps (Fig. 2A, left). Quantitative analysis by liquid chromatography-coupled tandem mass spectrometry (LC-MS/MS) revealed 29 abundant proteins common to all three preps (Fig. 2A, right; Table 1). Consistent with other analyses of in vitro-assembled spliceosomes (Schmidt et al. 2010; Agafonov et al. 2011; Ilagan et al. 2013), the most abundant species was Prp19, having a three- to fourfold higher stoichiometry than any other polypeptide. In isolation, Prp19 is a known tetramer (Ohi et al. 2005). All other abundant species were present at an approximately 1:1 stoichiometry, with the exception of translation elongation factor EF1A, which was significantly reduced in the MN prep. Multiple substoichiometric proteins were also present, with a total of 68, 45, and 43 species being consistently quantifiable in the LS, HS, and MN preps, respectively (Table 1).
| Proteins | LS | HS | MN |
|----------|----|----|----|
| # | S. pombe | S. cerevisiae | Human | Accession number | MW (kDa) | # Peptides | RS<sup>b</sup> | # Peptides | RS<sup>b</sup> | # Peptides | RS<sup>b</sup> |
| 1 | Prp19 | Prp19 | hPRP19 | SPAC29A4.08c | 54 | 23 | 2.95 | 19 | 2.99 | 20 | 2.94 |
| 2 | Prp3 | Prp3 | hPRP3 | SPAC3A12.11c | 44 | 17 | 1.07 | 18 | 1.23 | 16 | 1.28 |
| 3 | Smd2 | Smd2 | Smd2 | SPAC2C4.03c | 13 | 7 | 1.03 | 6 | 1.10 | 6 | 1.03 |
| 4 | Svf3 | Svf3 | hSYF3 | SPBC31F10.11c | 81 | 40 | 1.00 | 39 | 1.03 | 39 | 0.98 |
| 5 | Prp5 | Prp46 | PRL1 | SPB22H7.07 | 52 | 25 | 1.00 | 22 | 1.00 | 23 | 1.00 |
| 6 | Cdc5 | Cef1 | Cdc5L | SPBC46.12 | 87 | 50 | 1.00 | 50 | 1.00 | 49 | 1.00 |
| 7 | Cwf15 | Cwc15 | CCAP2 | SPBC337.06c | 30 | 11 | 0.99 | 9 | 1.00 | 12 | 0.97 |
| 8 | Prp17 | Prp17 | hPRP17 | SPBC681.10 | 63 | 31 | 0.98 | 27 | 0.97 | 33 | 1.17 |
| 9 | Snu114 | Snu114 | U5-116K | SPBC215.12 | 111 | 49 | 0.98 | 48 | 0.96 | 51 | 1.25 |
| 10 | Spp42 | Spp8 | U5-220K | SPAC4F6.12c | 275 | 76 | 0.97 | 83 | 0.94 | 88 | 1.06 |
| 11 | Ef1a | Ef1 | Ef1-α | SPCC794.09c | 50 | 10 | 0.95 | 8 | 0.93 | 8 | 0.29 |
| 12 | Smd3 | Smd3 | Smd3 | SPBC19C2.14 | 11 | 8 | 0.94 | 6 | 0.92 | 6 | 0.76 |
| 13 | Cwf18 | CCDC12 | KIA0A560 | SPBC466.02 | 148 | 32 | 0.84 | 35 | 0.81 | 34 | 1.04 |
| 14 | Tdh1 | Tdh1 | U5-40K | SPBC1289.11 | 37 | 21 | 0.77 | 18 | 0.77 | 19 | 0.82 |
| 15 | Cwf18 | Cwc18 | CCDC12 | SPBC215.12 | 111 | 49 | 0.98 | 48 | 0.96 | 51 | 1.25 |
| 16 | Syf1 | Syf1 | Syf1 | SPAC611.02c | 93 | 39 | 0.85 | 36 | 0.84 | 40 | 0.78 |
| 17 | Cwf19 | CWF19L2 | CCAP2 | SPBC215.12 | 111 | 49 | 0.98 | 48 | 0.96 | 51 | 1.25 |
| 18 | Cwf16 | Cwf16 | CCAP2 | SPBC215.12 | 111 | 49 | 0.98 | 48 | 0.96 | 51 | 1.25 |
| 19 | Snu114 | Snu114 | U5-116K | SPBC215.12 | 111 | 49 | 0.98 | 48 | 0.96 | 51 | 1.25 |

(continued)
The additional polypeptides in the LS prep were predominantly ribosomal proteins.

With the exception of EF1A and Tdh1, all abundant polypeptides were proteins previously known to associate with spliceosomes. Most highly represented were the Sm snRNP core proteins, the NTC, the NTC-related complex, and U5 snRNP. Together, the latter three species comprise the 35S U5 complex (Makarov 2002; Makarova et al. 2004). The only abundant U2-specific protein was Lea1 (U2 A′), to which the protein A tag was attached. Notably, no substituents of SF3A or SF3B (U2 snRNP-associated complexes required for early spliceosome assembly steps) were detectable in either the HS or MN prep; this was consistent with the predominant purified species being a post-activation complex, as SF3A and B are destabilized coincident with spliceosome activation (Bessonov et al. 2008; Lardelli et al. 2010). Also notably absent from the HS prep were the second step factors Slu7, Prp22, and Prp18, all of which are abundant components of Saccharomyces cerevisiae C complex purified under LS-like conditions (Fabrizio et al. 2009). Slu7 was found in human P complex (Ilagan et al. 2013), and S. cerevisiae ILS complex (Fourmann et al. 2013). Thus, with regard to their protein content, endogenous S. pombe U2-U5-U6 complexes are similar, but not identical, to human C and P complexes and S. cerevisiae C and ILS complexes.

RNA composition of purified complexes

RNAs in the LS, HS, and MN preps were analyzed both by denaturing PAGE and high-throughput sequencing (RNAseq) (see Materials and Methods). SYBR Gold staining of gels revealed three major 100- to 200-nt bands in the LS and HS preps (Fig. 2B, left). Northern blotting confirmed that these were U2, U5, and U6 snRNAs (Fig. 2B, center). SYBR Gold staining and RNAseq revealed three snRNAs to be present at comparable ratios (Fig. 2B, right). The next most abundant small RNA species were U4 and U1, with relative stoichiometries of 0.05 and 0.01, respectively. Other small RNAs (e.g., snoRNAs and tRNAs) collectively constituted <0.5% of the total. Thus, the bulk of our purified complexes constituted post-activation spliceosones (i.e., subsequent to U1 and U4 departure).

In the MN sample, both U5 and U6 snRNAs were quantitatively converted to faster migrating species, but their relative stoichiometries remained largely unchanged. RNAseq revealed that MNase removed 4 nt from the 5′ end of U5 and 5 nt from the 3′ end of U6 (Fig. 3A,B). Although migration of U2 was apparently unaffected by MNase treatment, its relative stoichiometry was reduced by nearly half. Analysis of the 5′ and 3′ termini of U2-mapping RNAseq tags in the MN library indicated two major cleavage sites: (1) within the GUAGUA branch recognition sequence; and (2) at the base of the penultimate stem–loop (Fig. 3B). The region between these two cleavage sites (within which our Northern probe

| # | S. pombe | S. cerevisiae | Human | Accession numbera | MW (kDa) | LS Peptides | HS RSb | MN RSb | LS Peptides | HS RSb | MN RSb |
|---|----------|--------------|-------|------------------|----------|-------------|--------|--------|-------------|--------|--------|
| 60 | Sme1     | Sme1         | Sme   | SPBC11G1.166c   | 100      | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 61 | Cip1     | Pin4         | Pin4  | SPBC16A3.18     | 53       | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 62 | Rps8-2   | Rps8b        | Rps8  | SPAC521.05      | 23       | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 63 | Sap155   | Hsh155       | S3b155| SPAC27F1.09c    | 135      | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 64 | Sum3     | Ded1         | Ddx3  | SPCC1795.11     | 70       | 1           | 0.08   | 1      | 2           | 0      | 0      |
| 65 | Cip2     |              |       | SPAC12G12.03    | 62       | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 66 | Rps15-2  | Rps15a       | Rps15 | SPAC1071.07c    | 18       | 2           | 0.08   | 1      | 2           | 0      | 0      |
| 67 | Rps3     | Rps3         | Rps3  | SPBC16G5.14c    | 28       | 1           | 0.08   | 1      | 2           | 0      | 0      |
| 68 | Sap130   | Prp12        | S3b130| SPAP698.03c     | 135      | 2           | 0.08   | 1      | 2           | 0      | 0      |

aThe accession numbers were taken from www.pombase.org.
b[RS] Relative stoichiometry to CDC5 protein.
was located) was twofold underrepresented in the RNAseq data compared to the 5′ and 3′ ends. Additional Northern blotting with probes complementary to other regions in U2 confirmed the existence of numerous fragments in the MN sample (Supplemental Fig. S2). Thus, our purified complexes either contain two distinct U2 conformers (one fully protected from nuclease digestion and another in which the region between the branch recognition sequence and the penultimate stem–loop is nuclease-accessible), or this region of U2 snRNA is kinetically less accessible to MNase digestion than are the 5′ and 3′ ends of U5 and U6 snRNA, respectively.

### Genome-wide analysis of RNAseq reads

In addition to U2, U5, and U6 snRNAs, heterogeneous higher molecular weight species were clearly present in all three samples (Fig. 2B, left). To identify these, we performed genome-wide analysis of our RNAseq tags using a stringent mapping algorithm (see Materials and Methods; see Supplemental Table S1A for data statistics). Of reads mapping to the S. pombe genome, 98%, 98%, and 99% (LS, HS, and MN libraries, respectively) mapped both within and sense to annotated genes. Further, of the nongenome mapping reads, 94%–99% mapped sense to exon-exon junctions in the predicted transcriptome. Thus, almost all of our mappable reads corresponded to the annotated S. pombe transcriptome (Rhind et al. 2011).

Known S. pombe genes can be divided into seven broad classes: CDS, rRNAs, snRNAs, snoRNAs, tRNAs, long terminal repeat (LTR)-containing transcripts, and other miscellaneous RNAs. The vast majority (≥99.5%) of tags mapping to known genes (including introns) mapped to the snRNA, rRNA, or CDS class, but the relative abundances among these classes varied among the three preps (Fig. 4A). Consistent with higher ribosomal protein abundance in the LS prep (Table 1), 40% of LS tags were from rRNA, compared to only 10% and 20% of HS and MN tags. Fewer tags in the MN library mapped to the CDS class (6%) than in the LS (17%) and HS (14%) libraries. This was also expected, as MNase should eliminate any sequences not directly protected by the spliceosome (see below).

There was excellent correlation between tag abundances mapping to individual CDSs across all three preps (r > 0.8),
indicating the high quality of our data (Fig. 4B). Out of the 2345 annotated intron-containing CDS genes in *S. pombe*, only 73 had no uniquely mapping reads in the MN data set and <5 reads in the HS data set. Close reexamination of the *S. pombe* transcriptome RNAseq data (Rhind et al. 2011) revealed that 70 of these are misannotated introns, while the remaining three are poorly expressed in late log phase (W Chen, H Shulha, N Rhind, Z Weng, M Moore, in prep.). Thus, our data cover 99.9% of intron-containing CDS genes.

Among CDS mapping tags, the fraction mapping to introns and exons of single-intron genes varied among the three preps in expected ways (Fig. 4C; Supplemental Table S1C). The vast majority of MN tags were intronic, consistent with the idea that only regions protected by spliceosomes survive MNase digestion. In the LS and HS preps, both of which initially contained intact RNAs, the percentage of total tags mapping to exons was higher in the LS prep than in the HS prep. This suggests that exonic regions were preferentially dissociated upon high-salt treatment. All of these data features were readily apparent on individual genes (Fig. 4D).

To more rigorously examine the possibility that exonic regions were less stably bound to our purified complexes than intronic regions, we calculated intron and exon tag densities for the most abundant single-intron CDS genes represented in the LS and HS preps (Table 2). In both preps, 5′ and 3′ exon tag densities were highly similar, indicating no preferential dissociation of the 5′ exon as might be expected for splicing intermediate-containing complexes. Further, the higher intron:exon ratio in the HS prep (12.7–14.4) than the LS prep (6.5–7.4) indicates that exonic sequences were prone to dissociation under high-salt conditions. Thus, the predominant CDS RNA species (93% for the HS prep) consisted of introns no longer associated with exons. This indicates that endogenous *S. pombe* U2·U5·U6 snRNA complexes are predominantly ILS complexes.

**DISCUSSION**

In this paper, we describe large-scale purification and comprehensive proteomics and ribonomics characterization of an abundant endogenous splicing complex containing U2, U5, and U6 snRNAs from *S. pombe* (Fig. 1A). RNAseq revealed that these endogenous complexes contain predominantly spliced intron products representing 99.9% of intron-containing CDS genes. Consistent with this, the set of proteins stably associated with this endogenous U2·U5·U6 complex strongly resembles the proteins associated with intron lariat spliceosomes assembled in vitro on a single intron from *S. cerevisiae* (Fourmann et al. 2013). Thus, spliceosome disassembly from intron lariats is likely rate-limiting for pre-mRNA splicing during logarithmic

**TABLE 2. Intron/exon tag densities and 5′SS, 3′SS numbers**

| Items                        | LS  | HS  |
|------------------------------|-----|-----|
| Average intron tag density/gene | 0.390 | 0.663 |
| Average 5′ exon tag density/gene | 0.053 | 0.052 |
| Average 3′ exon tag density/gene | 0.060 | 0.046 |
| Intron tag density:5′ exon tag density | 7.4  | 12.7 |
| Intron tag density:3′ exon tag density | 6.5  | 14.4 |
| ILS complex/all purified complexes (%) | 88  | 93  |

Data from the top 10% of single intron genes with signal. *Per million mappable reads.*
growth in *S. pombe* cells. A potential explanation for slow ILS complex disassembly is the relative absence of the ATPase Brr2, which is responsible for facilitating structural transitions involving U2 and U6 snRNAs.

**Protein compositional similarities and differences between C, P, and ILS complexes**

In comparison to the stable complexes observable during spliceosome assembly and activation, the complexes involved in spliceosome disassembly have been much less studied. One paper described isolation of two post-splicing complexes, Intron Large (IL) and Intron Small (IS), from in vitro splicing reactions in HeLa nuclear extract containing a chicken c-crystallin splicing substrate with three MS2 binding sites in the intron (Yoshimoto et al. 2009). Purified IL complex contained U2, U5, and U6 snRNAs and a number of bound proteins, whereas IS complex consisted largely of the naked intron. More recently, two other studies also employing the MS2 purification approach from in vitro splicing reactions reported much more comprehensive proteomics analyses of two late-stage splicing complexes: (1) HeLa P complex stalled on a splicing substrate derived from AdML pre-mRNA containing a very short (13-nt) 3′ exon that does not allow for efficient spliced exon release (Ilagan et al. 2013); and (2) *S. cerevisiae* ILS complex assembled on a splicing substrate derived from the Actin7 (Act7) pre-mRNA (Fourmann et al. 2013).

As expected, the protein composition of endogenous *S. pombe* ILS complexes closely resembles that of in vitro-assembled *S. cerevisiae* ILS complex (Fig. 5). Proteins shared between human P, *S. cerevisiae* ILS, and endogenous *S. pombe* ILS complexes include (with notable exceptions as discussed below) most U5, NTC, and NTC-related proteins. Taking into account the expected twofold and three- to fourfold stoichiometries of the Sm proteins and Prp19, respectively, and the snRNAs and protected intronic regions, the calculated MW of *S. pombe* ILS complexes is 2.1 MDa (MN) to 3.5 MDa (LS). This is very similar to human C complex (estimated MW 2.6 MDa), consistent with previous EM reconstructions showing strong structural similarities between these two complexes (Jurica et al. 2004; Ohl et al. 2007).

Nonetheless, numerous proteins stably associated with human C complex are missing from all three post-splicing complexes characterized to date. Several are displaced upon spliced exon dissociation by Prp22 (Fourmann et al. 2013). These include the second step factors S1u7 and Prp18 (Zhang and Schwer 1997), the RES complex (Lst3, Bud13, Pml1) that functions in nuclear retention of spliceosomes (Dziembowski et al. 2004), and NTC components Cwc21 and Cwc22. Cwc21 interacts directly with Prp8 and influences the fidelity of splicing (Grainger et al. 2009). Cwc22 was recently shown to serve as the binding platform for exon junction complex (EJC) assembly on the spliced exons (Barbosa et al. 2012; Steckelberg et al. 2012). As expected for complexes predominated by lariat introns, we detected no EJC proteins in any of our endogenous ILS complex preparations.

**U2 snRNP proteins and U2 snRNA conformation**

In comparison to human P complex and *S. cerevisiae* ILS complex, a number of proteins are significantly underrepresented in or completely absent from *S. pombe* ILS complexes. Of particular note are all specific components of 17S U2 snRNP (Brosi et al. 1993a; Krämer et al. 1999) except for U2A′ (Lea1), to which one of our affinity tags was attached. One missing protein is U2B′′ (Msl1), an RNA recognition motif (RRM)-containing protein that binds the 3′-most structural element in U2 snRNA (stem–loop IV) as a heterodimer with U2A′ (Scherly et al. 1990; Williams and Hall 2011). The complex of U2 snRNA, the Sm proteins, and the U2A′:U2B′ heterodimer comprise 12S U2 snRNP (Boelens et al. 1991). Although Msl1 is essential for vegetative growth in *S. pombe* (Kim et al. 2010), it is inessential in *S. cerevisiae* (Tang et al. 1996). Together with Lea1, Msl1 has been implicated in pre-spliceosome formation and exhibits synthetic lethality with Mud2, a splicing factor involved in initial recruitment of U2 snRNP to the branch site (Abovich et al. 1994; Tang et al. 1996; Caspary 1998). Taken together, these data suggest that Msl1 functions primarily in early spliceosome assembly stages but is not required for overall spliceosome integrity. Although we observed no growth defects in our tagged *S. pombe* strains compared to the parental strain, it is possible that the presence of the protein A tag on Lea1 destabilized Msl1 association with U2 snRNA.

The remainder of 17S U2 snRNP is comprised of the nine SF3A and B proteins (SF3A: SF3a60, SF3a66, SF3a120; SF3B: SF3b14b, SF3b14a, SF3b49, SF3b130, SF3b145, and SF3b155) (Behrens et al. 1993; Hodges and Beggs 1994; Bessonov et al. 2008). During U2 snRNP assembly in vitro, SF3B proteins bind first and then recruit the SF3A proteins (Behrens et al. 1993; Brosi et al. 1993a; Krämer et al. 1999). SF3B components bind U2 snRNA stem–loop IIa, whereas SF3A complex binds stem–loops IIa and IIb (Yan and Ares 1996; Lin and Xu 2012). SF3A and B also protect the region around the GUAGUA branch recognition sequence from chemical modification (Behrens et al. 1993; Yan and Ares 1996).

Both SF3A and SF3B are required for early spliceosome assembly (Brosi et al. 1993a,b). At that stage, SF3B components make extensive interactions with the pre-mRNA to either side of the branch site and are important for U2 snRNP recruitment and retention (Brosi et al. 1993a; Gozani et al. 1996). Three studies had suggested that SF3B’s interaction with the rest of the spliceosome is destabilized upon catalytic activation of the human and *S. cerevisiae* spliceosomes (Bessonov et al. 2008, 2010; Lardelli et al. 2010). More recently, however, Coltri et al. (2011) identified SF3B proteins in human C complex; they proposed that SF3A and SF3B release their hold on U2 snRNA during spliceosome activation but maintain their grip on the intron, where they help to hold the
splicing intermediates in place. Consistent with this, SF3b145 and SF3b130 were recently detected in human P complex (Ilagan et al. 2013), and most of the SF3A/B proteins were identified in *S. cerevisiae* ILS complex (Fourmann et al. 2013).

Our data indicate that SF3A components are completely absent from endogenous *S. pombe* ILS complexes and that the SF3B proteins are only loosely bound—they were substochiometric in LS complexes and completely absent from HS and MN complexes (Fig. 2B; Table 1). Consistent with this, our RNAseq results revealed two major MNase cleavage sites (↓) in U2 snRNA: one within the GUAGUA branch recognition sequence and another at the 5′ end of stem–loop IV (Fig. 3B). Protection and retention of stem–loop IV is expected because this is the binding site of Lea1 (see above).

Unexpected from previous data, however, is nuclease accessibility of the GUAGUA branch recognition sequence. A recent detailed study of snRNA nucleotide accessibility to chemical modification revealed only minimal modification of the GUAGUA sequence in pre- and post-activation spliceosomes (human B, B*hs, and C complex) (Anokhina et al. 2013).
compared to downstream nucleotides (i.e., 3′ from the GUAGUA region). Our data thus suggest that after spliced exon release, the GUAGUA region becomes more solvent accessible, perhaps due to SF3 protein release.

**Spliceosome disassembly: a slow step in the S. pombe splicing cycle?**

Despite sustained efforts by several laboratories, no conditions have yet been found that allow complete spliceosome assembly and catalysis in *S. pombe* extracts. Whereas *S. cerevisiae* and human extracts contain significant amounts of U4/5/6 tri-snRNP ready to engage in new spliceosome assembly, crude *S. pombe* lysates are predominated by ILS complex (Huang et al. 2002). Therefore, spliceosome disassembly seems to be a slow step in the *S. pombe* splicing cycle. ILS complex disassembly requires the abundant DEAH ATPase, the U5 snRNP protein and DEIH box ATPase Brr2 (Chen et al. 2010). Ntr1 bridges Prp43 to Ntr2, which, in turn, interacts with Ntr2 (Boon et al. 2006; Tanaka et al. 2007; Tsai et al. 2007). Step spliceosomes by the NTC-related proteins Ntr1 and Ntr2 (Boon et al. 2006; Tanaka et al. 2007; Tsai et al. 2007). Ntr1 bridges Prp43 to Ntr2, which, in turn, interacts with the U5 snRNP protein and DEIH box ATPase Brr2 (Chen et al. 2013). Although *S. pombe* contains no clear Ntr2 homolog, it is of note that Ntr1, Prp43, and Brr2 were all significantly substoichiometric in our purified complexes (Table 1).

During spliceosome activation, Br2 functions to unwind the U4/U6 helices to enable release of U4 snRNA and formation of the catalytic core in which U6 is extensively base-paired with U2 (Laggerbauer et al. 1998). Br2’s ATPase activity is also required for spliceosome disassembly (Small et al. 2006; Hahn and Beggs 2010), and its ATPase activity in vitro is stimulated by hybridized U2 and U6 snRNAs (Xu et al. 1996). Thus, Br2 likely functions to disrupt base-pairing interactions between U2 and U6 at this stage (Small et al. 2006). Because of its stable association with U5 snRNP as well as all major spliceosomal complexes in the human and *S. cerevisiae* systems, Brr2 was not previously thought to be a dynamically interacting splicing factor (Hahn and Beggs 2010). It was, therefore, surprising to us that Brr2 was significantly substoichiometric in (LS preps) or completely absent from (HS and MN preps) our purified endogenous *S. pombe* U2-U5-U6 complexes. Perhaps this unstable interaction of Brr2 with U2-U5-U6 complexes explains both ILS accumulation in logarithmically growing yeast and the inability of *S. pombe* extracts to support splicing in vitro (Huang et al. 2002).

**MATERIALS AND METHODS**

**S. pombe strain construction**

All strains derived from yFS 104 (ura4-D18 leu1-32 h+) and yFS 105 (ura4-D18 leu1-32 h–). The *lea1* and snu114 chromosomal loci fused at their C termini either to a TEV cleavage site followed by protein A or to the calmodulin binding peptide were constructed by PCR-based gene targeting (Bahler et al. 1998) using plasmid pFA6a-CTAP-MX6 (Tasto et al. 2001) as a PCR template. *cdc5, spp42 (prp8),* and *brr2* chromosomal loci were individually 3′ end GFP-tagged by PCR-based gene targeting using pFA6a-GFP (F64L, S65T)-kanMX6 (Bahler et al. 1998) as a PCR template. Strain MJM01 (Cdc5-GFP, Lea1-protein A, Snu114-CBP, ura4-D18 leu1-32 h–) was generated by mating of appropriate parent strains, followed by sporulation and PCR validation of haploid derivatives. Strains MJM02 and MJM03 encoded Spp42-GFP and Brr2-GFP, respectively.

**Complex purification**

Purification of endogenous U2-U5-U6 complexes was carried out as previously described for TAP-tagged complexes (Puig et al. 2001) with the following modifications. For LS preps, cells from high-density cultures (5×YES; OD600 = 23–25) were washed in Buffer A (10 mM K-HEPES, pH 7.9, 40 mM KCl, 1.5 mM MgCl2, 0.5 mM DTT, 0.5 mM PMSF, 2.0 mM benzamidine), repelleted, frozen as nucleotides in liquid nitrogen, and pulverized to a powder in a ball mill (PM 100, Retsch). Frozen cell powder was resuspended in Buffer B (Buffer A with 400 mM KCl) and the supernatant used for TAP purification in 150 mM NaCl as previously reported (Puig et al. 2001). Complexes were recovered from calmodulin resin by applying elution buffer containing 6 mM EGTA and collecting eight 400-μl fractions. HS preps were identical except that frozen cell powder was resuspended in Buffer C (Buffer A with 400 mM KCl and 300 mM NaCl), and the protein A binding and washing buffers contained 700 mM NaCl. MN preps were the same as HS preps except complexes bound to IgG beads were treated with MNaSe prior to TEV cleavage. For all preps, elution fractions were assessed by GFP fluorescence and Bradford assays. Detailed purification protocols, dynamic light scattering, and negative stain electron microscopy are described in Supplemental Materials and Methods.

**Electron microscopy**

For EM, purified complex (200 μg/mL) was applied to glow-discharged, carbon-coated grids and negatively stained with 2% uranyl acetate. Micrographs (66,000× magnification) were taken on a Philips CM12 microscope operating at 120 kV.

**Mass spectrometry**

Fifty micrograms total protein from each prep were digested with trypsin (Promega), and *S. cerevisiae* alcohol dehydrogenase (ADH) tryptic digest (P00330; Waters Inc.) was added to the digest as an internal standard (Silva et al. 2006). Each sample had 500 ng of tryptic peptides injected in triplicate, and nanoflow separation was performed on a nanoACQUITY Ultra Performance Liquid Chromatography (UPLC) system (Waters) equipped with a 75-μm × 150-mm analytical column packed with 1.7-μm bridged-ethyl hybrid (BEH) C18, by application of a linear 60-min gradient from 3% to 90% acetonitrile (ACN) in 0.1% formic acid. A lock mass solution of 200 fmol/μl Glu-fibrinopeptide was delivered via the auxiliary solvent manager at 500 nL/min into the reference sprayer of the NanoLockSpray source. Analysis of nano-UPLC elutes were performed on a Q-Tof Premier mass spectrometer (Waters) operated in the V-mode, and alternating scans were used to detect precursor ions.
ions and then fragment ions. The masses of the precursor ions were detected with a low collision energy (CE) of 5 eV, which was followed by a scan where the CE was ramped from 15 to 40 eV (MS² scan). MS data were analyzed using ProteinLynxGlobalServer (PLGS) Identity³ Version 2.4 software (Waters) against the protein database Sanger S. pombe (www.sanger.ac.uk) which had been combined with an equal-sized database consisting of the randomized sequences for the estimation of false-positive (Li et al. 2009). For database search and protein acceptance, the following criteria were imposed: three fragment ions per peptide and seven fragment ions per protein with at least one unique peptide per protein; within 10 and 20 parts per million (ppm) of the theoretical masses for precursor and fragment ions, respectively; full tryptic specificity with one missed cleavage; variable modifications of oxidation (methionine), deamidation (asparagine, glutamine), as well as static modification of carbamidomethylation (cysteine). The database search returned results for each injection until 4% of the identified proteins were false (random sequence). After all of the data were processed, a further requirement was made that, for acceptance, a protein must be identified in at least two of three replicate injections of the same sample. Absolute quantitation of identified peptides was achieved with the ion chromatographic peak areas of the three most intense tryptic peptides for each protein relative to that of the signal intensity of the three most intense ions for the internal standard (ADH). For each accepted protein, the relative standard deviation of the femtomole amount from the three replicate runs was calculated (Silva et al. 2006).

**Western blots**

Cell extracts from strain MJM01 (Cdc5-GFP, Lea1–protein A, Snu114-CBP), MJM02 (Brr2-GFP, Lea1–protein A, Snu114-CBP), and MJM03 (Spp42-GFP, Lea1–protein A, Snu114-CBP) were used for Western blots. Total proteins were separated by 4%–12% SDS-PAGE and transferred to PVDF membrane. Western blots were carried out by using anti-GFP (SC-9996, Santa Cruz) with 800RD anti-mouse IgG, and detected with a low collision energy (CE) of 5 eV, which was followed by a scan where the CE was ramped from 15 to 40 eV (MS² scan). MS data were analyzed using ProteinLynxGlobalServer (PLGS) Identity³ Version 2.4 software (Waters) against the protein database Sanger S. pombe (www.sanger.ac.uk) which had been combined with an equal-sized database consisting of the randomized sequences for the estimation of false-positive (Li et al. 2009). For database search and protein acceptance, the following criteria were imposed: three fragment ions per peptide and seven fragment ions per protein with at least one unique peptide per protein; within 10 and 20 parts per million (ppm) of the theoretical masses for precursor and fragment ions, respectively; full tryptic specificity with one missed cleavage; variable modifications of oxidation (methionine), deamidation (asparagine, glutamine), as well as static modification of carbamidomethylation (cysteine). The database search returned results for each injection until 4% of the identified proteins were false (random sequence). After all of the data were processed, a further requirement was made that, for acceptance, a protein must be identified in at least two of three replicate injections of the same sample. Absolute quantitation of identified peptides was achieved with the ion chromatographic peak areas of the three most intense tryptic peptides for each protein relative to that of the signal intensity of the three most intense ions for the internal standard (ADH). For each accepted protein, the relative standard deviation of the femtomole amount from the three replicate runs was calculated (Silva et al. 2006).

**RNA and Northern blots**

Total RNA from purified complexes was resolved on an 8% polyacrylamide–8 M urea gel and the gel stained with SYBR Gold (Invitrogen). For Northern blotting, RNAs were transferred to Biotrans (+)™ NYLON membranes by semidry electroblotting. snRNAs were detected with complementary, ³²P-labeled oligonucleotides; the U6 probe spanned its exon–exon junction.

**Deep sequencing**

RNaseq libraries were generated as previously described (Ingolia et al. 2009) with the following modifications. Total RNA from each prep was subjected to alkaline hydrolysis followed by incubation with T4 polynucleotide kinase (NEB) in the absence of ATP to remove 2′, 3′-cyclic and 3′ phosphate groups. Following denaturing gel electrophoresis, the 30- to 80-nt region was excised, and recovered RNAs were poly[A]-tailed and converted into libraries for deep sequencing on the Illumina GAIL platform as previously described (Ingolia et al. 2009).

**RNAseq data analysis**

Raw 36-bp reads were searched for the poly(A) adapter sequence. If a run of six or more adenosines was detected, the sequence prior to the first A was excised and mapped to the S. pombe genome and transcriptome (Sanger Sept. 2011 version) with Bowtie (version 0.11.3). To avoid mapping to spurious locations in the genome, we only retained sequences that met the following criteria: (1) sequences 12- to 15-nt-long had to map perfectly to a single location in the genome or transcriptome; (2) sequences 16- to 19-nt-long had to map perfectly to the genome or transcriptome; (3) sequences ≥20 nt had to map to the genome or transcriptome with no more than one mismatch. If a sequence ≥16 nt mapped to multiple locations, each location was allocated an equal share. The UCSC genome browser (http://genome.ucsc.edu/) was used for making numerous figures. All other analyses were performed with custom-built PERL scripts.

Quantitative analyses employed five classes of CDS mapping reads: (1) exonic = reads mapping entirely inside an exon; (2) intronic = reads mapping entirely inside an intron; (3) 5′SS = reads straddling a 5′ splice site; (4) 3′SS = reads straddling a 3′ splice site; and (5) EEJ = reads straddling an exon–exon junction. For correlation with Pol II occupancy and mRNA abundance, we summed the total reads of all five types for each gene. Peak calling in intergenic regions was performed by extending each intergenic position with ≥11 tags in both directions until a position having no tag was encountered. Overlaps between peaks and annotated genes were performed in a strand-specific manner.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

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