The life of U6 small nuclear RNA, from cradle to grave

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ABSTRACT

Removal of introns from precursor messenger RNA (pre-mRNA) and some noncoding transcripts is an essential step in eukaryotic gene expression. In the nucleus, this process of RNA splicing is carried out by the spliceosome, a multi-megaDalton macromolecular machine whose core components are conserved from yeast to humans. In addition to many proteins, the spliceosome contains five uridine-rich small nuclear RNAs (snRNAs) that undergo an elaborate series of conformational changes to correctly recognize the splice sites and catalyze intron removal. Decades of biochemical and genetic data, along with recent cryo-EM structures, unequivocally demonstrate that U6 snRNA forms much of the catalytic core of the spliceosome and is highly dynamic, interacting with three snRNAs, the pre-mRNA substrate, and >25 protein partners throughout the splicing cycle. This review summarizes the current state of knowledge on how U6 snRNA is synthesized, modified, incorporated into snRNPs and spliceosomes, recycled, and degraded.

Keywords: U6 snRNA; U6 gene transcription; spliceosome; U6 snRNP

U6 snRNA IS A KEY COMPONENT OF THE SPLICEOSOME

U6 snRNA was discovered by virtue of its immunoprecipitation from mouse cell nuclear extract, along with U1, U2, U4, and U5 snRNAs, by anti-Sm patient serum (Lerner and Steitz 1979), and U6 from rat cells was subsequently sequenced (Epstein et al. 1980). It was later found that human U6 snRNA is stably base-paired with U4 snRNA (Bringmann et al. 1984; Hashimoto and Steitz 1984). Using human cell nuclear extract competent for pre-mRNA splicing, U4 and U6 snRNAs were found to be associated with affinity-purified spliceosomes (Grabowski and Sharp 1986), and oligo ablation studies revealed that both snRNAs are required for splicing (Black and Steitz 1986). Soon thereafter, U4 and U6 were found to be highly conserved and essential in yeast (Siliciano et al. 1987; Brow and Guthrie 1988), and both snRNAs were identified in yeast spliceosomes (Cheng and Abelson 1987), implying a central function in splicing.

The spliceosome is an intricate macromolecular machine that consists of five snRNAs, U1, U2, U4, U5, and U6, and their associated proteins (small nuclear ribonucleoprotein complexes or snRNPs), as well as a protein-only NineTeen Complex (NTC) and a number of accessory proteins (Wahl et al. 2009; Will and Lührmann 2011). A small percentage of human introns, as well as those of certain other metazoan and protozoan taxa, have noncanonical splice sites and are recognized and removed by the minor “U12-type” spliceosome. The minor spliceosome shares U5 snRNA with the major spliceosome, but contains the U11, U12, U4atac, and U6atac snRNAs, which replace U1, U2, U4, and U6 snRNAs, respectively (Tarn and Steitz 1996a,b). Both the major and minor spliceosomes assemble de novo on each intron and, after splicing, dissociate into their component snRNPs and accessory proteins. The dynamic nature of the spliceosome makes its study challenging.

U6 snRNA sits at the heart of the spliceosome, where it coordinates the magnesium ions required for splicing chemistry and, with help from U2 and U5 snRNAs, positions the substrate for the splicing reaction. The pre-mRNA splicing reaction consists of two transesterification steps and closely resembles the reaction carried out by group II self-splicing introns. First, the 2′-oxygen of a nucleotide near the 3′ end of the intron (the branch site, most often an adenosine) is activated for nucleophilic attack on the 5′-exon–intron junction (the 5′ splice site) to form a branched “lariat” intron–3′ exon intermediate. Subsequently, nucleophilic attack of the released 5′ exon on the intron–3′ exon junction (the 3′ splice site) produces a ligated 5′ exon–3′ exon product and free lariat intron (Wahl et al. 2009; Hoskins and Moore 2012; Shi 2017).
Early steps in recognition of the intron include base-pairing of U1 snRNA in the U1 snRNP to the 5’ splice site and U2 snRNA in the U2 snRNP to the branch site, forming the A complex (also known as the prespliceosome). The A complex is joined by a preassembled U4/U6.U5 triple (tri)-snRNP to form the B complex (Fig. 1). In tri-snRNP nomenclature, the slash indicates base-pairing between U4 and U6 snRNAs and the period conveys that U5 snRNP associates with the U4/U6 di-snRNP through protein–protein and RNA–protein interactions. During activation of the assembled spliceosome for the first catalytic step, large-scale conformational rearrangements cause U1 and U4 snRNPs to be ejected, leaving the U2/U6.U5 complex together with the NTC (B*act) and accessory factors to catalyze the chemical steps (Will and Lührmann 2011). U6 must undergo several RNA base-pairing exchanges in this transition: disruption of U4/U6 in favor of a mutually exclusive interaction with U2, and base-pairing to the 5’SS. The timing of these events and the interactions controlling activation are still being elucidated (Shcherbakova et al. 2013; Hoskins et al. 2016).

Recent cryo-EM structures of spliceosomal complexes, including the U4/U6.U5 tri-snRNP (Agafonov et al. 2016; Nguyen et al. 2016; Wan et al. 2016b), the precatalytic B complex (Bertram et al. 2017a; Plaschka et al. 2017), and the catalytic U2/U6.U5 spliceosome at different stages of splicing (Yan et al. 2015, 2016, 2017; Galej et al. 2016; Rauhut et al. 2016; Wan et al. 2016a, 2017; Bertram et al. 2017b; Fica et al. 2017; Zhang et al. 2017), have provided near-atomic level views of the extensive structural rearrangements that must take place during splicing. After each round of splicing, the snRNPs and NTC dissociate from the intron and are used again. U6, however, is released as a free RNA and so must bind its own snRNP proteins again before assembling with the U4 and U5 snRNPs to reconstitute the tri-snRNP (Fourmann et al. 2013). The splicing cycle is described in greater detail in recent reviews (Will and Lührmann 2011; Hoskins and Moore 2012).

A role for U6 in the catalytic center of the spliceosome had long been proposed based on a number of lines of evidence. The discovery that group II self-splicing introns use the same two-step chemical mechanism as the spliceosome (Peebles et al. 1986; van der Veen et al. 1986) suggested that the two enzymes share catalytic RNA elements (Sharp 1985; Cech 1986; Madhani and Guthrie 1992). The high degree of conservation in the U6 snRNA sequence (Brow and Guthrie 1988) and the presence of introns in the most conserved region of U6 in some organisms (Tani and Ohshima 1989, 1991), potentially due to reverse splicing, hinted at the proximity of U6 to the active site of the spliceosome (Brow and Guthrie 1989). Subsequent crosslinking and genetic studies indicated that the strictly conserved “ACAGA-box” sequence of U6 base pairs to the intron 5’ splice site in the catalytically active spliceosome (Sawa and Abelson 1992; Sawa and Shimura 1992; Wassarman and Steitz 1992; Kandels-Lewis and Séraphin 1993; Lesser and Guthrie 1993; Sontheimer and Steitz 1993), and mutational studies identified functional similarities between Domain V of group II introns and the U2/U6 base-paired region (Madhani and Guthrie 1992; Chanfreau and Jacquier 1994; Peebles et al. 1995; Shukla and Padgett 2002; Mefford and Staley 2009). Elegant biochemical experiments confirmed that U6 snRNA is responsible for coordinating catalytic metal ions essential for both transesterification steps (Yean et al. 2000; Fica et al. 2013).

At just over 100 nt in length, U6 snRNA is highly economical, as nearly every part of the RNA possesses one or more essential functions. In addition to its catalytic role at the heart of the spliceosome, U6 snRNA is notable for undergoing extensive structural rearrangements, including unwinding and reformation of stable internal secondary structure, and for directly interacting with >25 proteins.
during a single round of splicing. Herein, we summarize the entire life cycle of the U6 snRNA (Fig. 2), and highlight differences found among the eukaryotes. Commonalities in the U6 snRNA life cycle illustrate conservation across more than a billion years of eukaryotic evolution. We focus primarily on U6 snRNA from the best-studied organisms: yeast (S. cerevisiae, and to a lesser extent, S. pombe) and human.

U6 snRNA GENE TRANSCRIPTION

Number of U6 genes

S. cerevisiae has a single genomic locus for U6 snRNA, the SNR6 gene on chromosome XII (Brow and Guthrie 1988). However, a survey of 145 fungal genomes identified species with up to 20 U6 gene copies and an average of 2.3 per genome (Canzler et al. 2016). In contrast, there are >900 copies of U6 distributed throughout the human genome, although the majority of these are likely pseudogenes and not transcriptionally active (Doucet et al. 2015). At least four human U6 genes encoding identical RNAs are transcriptionally active to various degrees (Domitrovich and Kunkel 2003). Additionally, a variant of human U6 snRNA with nine substitutions and one nucleotide deletion is expressed under the control of an internal promoter, unlike other transcriptionally active human U6 genes (Tichelaar et al. 1994, 1998). The presence of multiple U6 genes of varying transcriptional activity has complicated their individual study, and whether paralogous but divergent U6 snRNAs exhibit differences in modification, localization or function is poorly understood.

The U6atac RNA is a paralog of U6 that functions in the minor spliceosome and is even further diverged in sequence from the other transcribed U6 snRNAs (Tarn and Steitz 1996a).

Transcription of U6 genes by RNA polymerase III

Unlike the other spliceosomal snRNAs, which are synthesized by RNA polymerase II (Pol II), U6 is synthesized by RNA polymerase III (Pol III) (Reddy et al. 1987; Moenne et al. 1990). While the sequence of U6 snRNA is highly conserved between yeast and humans, its Pol III promoter structure is divergent. In yeast, the U6 promoter region is similar to tRNA gene promoters (Eschenlauer et al. 1993) in that it contains A and B block elements (Brow and Guthrie 1990), as well as a TATA box that is bound by TATA-binding protein (TBP) (Fig. 3A; Margottin et al. 1991). In S. cerevisiae, the B block is located downstream from the U6 coding region. Consequently, the distance between the A and B blocks is atypical (~200 bp vs. the normal 30–90 bp in tRNA genes), and this extended spacing is important for transcription in vivo (Kaiser et al. 2004).

The B block is essential for transcription of the yeast U6 gene in vivo and is the binding site for transcription initiation factor TFIIIC, which also contacts the A block (Burnol et al. 1993; Eschenlauer et al. 1993; Kaiser and Brow 1995). TFIIIC recruits TFIIIB (composed of TBP, Bdp1, and Brf1), which binds over the TATA box and in turn recruits Pol III (Gerlach et al. 1995). Transcription is promoted by but not dependent on the presence of the TATA box and a
FIGURE 3. U6 snRNA gene promoter structure is divergent in eukaryotes. (A) U6 gene promoter structure in S. cerevisiae. The U6 gene is under the control of a Pol III Type II promoter, with an upstream TATA box, an internal A block, and downstream B block. The TFIIIC complex recognizes the A and B blocks and directs binding of the TFIIIB complex to the TATA box. Nhp6 also promotes transcription, but its binding site is uncertain. A possible nucleosome is indicated by a gray oval. (B) U6 promoter structure in humans. U6 synthesis is under the control of a Pol III Type III promoter, where promoter elements (in black) are exclusively upstream of the transcription start site. The TATA box is recognized by the TFIIIB2 complex, while the PSE is recognized by the SNAPc complex. Factors including OCT1, STAF, and CDH2 interact with the DSE. p38 inhibits Oct-1 binding to the DSE. A nucleosome between the DSE and PSE enhances Oct-1 and SNAPc binding.

stretch of thymidines just downstream from it (Burnol et al. 1993; Eschenlauer et al. 1993; Martin et al. 2001).

The nonhistone chromatin protein Nhp6 is also important for transcription of SNR6 (Kruppa et al. 2001; Martin et al. 2001). In a heterologous, in vitro chromatin assembly system, a nucleosome positioned between the A and B blocks brings the regions close together for optimal binding by TFIIIC (Shivashwamy et al. 2004), but the micrococcal nuclease footprint of native chromatin assembled between the A and B blocks in vivo is shorter than expected for an intact nucleosome (Gerlach et al. 1995). It is possible that Nhp6 modifies the structure of a nucleosome bound to SNR6 (Fig. 3A; Stillman 2010). Alternatively, Nhp6 may favor a bent conformation of DNA that promotes TFIIIB binding (Braglia et al. 2007).

U6 gene promoter structure in fungi is flexible and can include or exclude identifiable TATA boxes, intragenic A blocks, and downstream B blocks (Cantzler et al. 2016). S. pombe has a similar promoter structure to S. cerevisiae, with the exception that the B block element is located intragenically in an intron (Frendewey et al. 1990).

In humans, U6 RNA is transcribed from a Pol III Type III promoter, a promoter architecture that also drives 7SK RNA and RNase P RNA transcription. Most characterized mammalian U6 genes do not contain intragenic promoter elements, but rather exclusively contain upstream promoter elements (Das et al. 1988). These U6 genes have a TATA box 30 base pairs upstream of the transcription start site, a proximal sequence element (PSE) 50 base pairs upstream, and a distal sequence element (DSE; also called an OCT site) 250 base pairs upstream, which closely resembles the promoter structure for Pol II-transcribed genes (Fig. 3B). Interestingly, the PSE and DSE are similar to, and can be switched with, corresponding elements from the Pol II-synthesized snRNAs (Kunkel and Pederson 1988), and it is inclusion of the TATA box that specifies Pol III rather than Pol II transcription for snRNAs (Lobo and Hernandez 1989). Other organisms have divergent methods of specifying Pol II vs III transcription, such as the sequence of the PSE or the distance between the PSE and the TATA box (for review, see Hernandez 2001).

The PSE is required for basal transcription while the DSE is required for efficient transcription. The PSE is bound by the small nuclear RNA activating protein complex (SNAPc), which also binds the promoters of Pol II-transcribed snRNAs. The DSE is bound by the Oct-1 and Staf proteins (Fig. 3B). Cooperative binding of Oct-1 and SNAPc is enhanced by a nucleosome positioned between the DSE and PSE (Zhao et al. 2001). Oct-1 binding is negatively regulated by p38 kinase (Lin and Natarajan 2012), while Staf binding is influenced by chromatin-modifying enzymes (Yuan et al. 2007). The TATA box is bound by TFIIIB2, composed of TBP, Bdp1, and Brf2. TFIIIC is not required as there is no A or B block. Interestingly, yeast also possess an upstream PSE-like sequence that is not required for transcription (Eschenlauer et al. 1993) and an upstream positioned nucleosome (Arimbassi and Bhargava 2008), suggesting that while some similarities remain, the promoter structure in yeast and humans has diverged significantly.

Chromatin immunoprecipitation detects very low levels of Pol II over the S. cerevisiae U6 gene, similar to that detected over the Pol II-silenced rDNA and telomeres (Steinmetz et al. 2006). Furthermore, a hypomorphic mutation in the Sen1 helicase increases Pol II levels at all these loci (Steinmetz et al. 2006), and an anti-sense transcript of the U6 gene contains a high-affinity Nrd1 binding site that promotes Sen1-dependent Pol II termination (Steinmetz and Brow 1998). Thus, like the rDNA and telomeres (Vasiljeva et al. 2008), the S. cerevisiae U6 gene may be silenced for Pol II by an unknown mechanism coupled to Sen1-dependent termination of an anti-sense transcript.

Interestingly, in humans, transcription of U6 is dependent upon the interaction of Pol II at a site ∼300 bp upstream of the gene, a phenomenon shown to be generally true for Pol III-transcribed genes (Listerman et al. 2007; Barski et al. 2010; Oler et al. 2010). This may result from the influence of chromatin remodeling through recruitment of Pol II transcription factors that are also used in Pol III transcription (Raha et al. 2010). Transcription of U6atac is also dependent on both Pol II and Pol III (Younis et al. 2013). Thus, there is a complex interplay of Pol II and Pol III at U6 genes in a variety of organisms.

Transcription termination of U6 is caused by a stretch of dA’s in the template strand at the end of the gene, although
the number of dA’s for efficient termination vary in eukaryotes (Arimbasseri et al. 2013). The La protein (Lhp1 in yeast) binds the 3’ end of newly transcribed U6 RNAs (Rinke and Steitz 1985; Pannone et al. 1998) and has been implicated in transcription termination, RNA polymerase recycling, and transcription reinitiation (Gottlieb and Steitz 1989; Maraia et al. 1994; Maraia 1996; French et al. 2008). However, Pol III transcription in vitro is terminated efficiently without the help of additional termination factors. Weak base-pairing between the nascent oligo(rU) and the template strand oligo(dA) stretch, as well as interactions between the non-template strand oligo(dT) stretch and a subunit of Pol III, terminate transcription (Bogenhagen and Brown 1981; Hamada et al. 2000; Arimbasseri and Maraia 2015). Yeast U6 is terminated in a stretch of ten dAs, leaving an RNA product with a heterogeneous U-tail length consisting of 4–7 uridines (Brow and Guthrie 1990).

How U6 is transcriptionally regulated during the cell cycle, development, in different tissues, and in disease states has been poorly studied. It has been noted that tissue-specific differences of U6 expression levels exist (Spaniel et al. 2013). Whether this difference is due to tissue-specific stability or transcriptional activity is not clear (Spaniel et al. 2013). Tissue-specific differences in snRNP distribution have also been reported (Hamm and Mattaj 1989). Transcriptional activity represents the first step of regulation in the lifecycle of U6, yet little is known about the cellular regulation of U6 snRNA level and its effect on splicing efficiency or fidelity.

LOCALIZATION DURING BIOGENESIS

It is generally thought that U6 remains in the nucleus after transcription and for all biogenesis steps (Vankan et al. 1990; Pessa et al. 2008), while the Pol II-synthesized snRNAs have a cytoplasmic snRNP assembly step (for review, see Matera and Wang 2014). At some point after transcription, the La/Lhp1 protein is displaced from the 3’ end of U6 snRNA. Binding by La/Lhp1 is not sufficient for complete nuclear retention in Xenopus oocytes (Boelens et al. 1995). Instead, binding of the Lsm2-8 heteroheptameric protein ring acts as the primary nuclear retention signal (Spiller et al. 2007,a,b).

U6 is modified (pseudouridylated and 2’-O-methylated) in the nucleolus by snoRNPs (Tycowski et al. 1998; Ganot et al. 1999; Lange and Gerbi 2000), while the Pol II-synthesized snRNAs are modified in Cajal bodies by scaRNPs (Darzacq et al. 2002). U6 is not retained in the nucleolus, but rather transiently passes through to obtain modifications (Lange and Gerbi 2000). Nucleolar localization of Xenopus U6 does not depend on base-pairing interactions with U4 or U2, or formation of the U6 5’SL, but both nucleolar localization and Cajal body localization do depend on the 3’ oligo (U) tail (Gerbi and Lange 2002). Furthermore, localization does not depend on post-transcriptional modifications such as pseudouridylation or 2’-O-methylation. In S. cerevisiae, U6 is not constitutively pseudouridylated and is not 2’-O-methylated, and therefore is thought to be entirely nucleoplasmic (Bertrand et al. 1998).

The presence of snRNPs in subnuclear organelles was first identified in the early 1990s (Carmo-Fonseca et al. 1991a,b; Matera and Ward 1993). Work over the past decade has revealed that U6 can be assembled into the U4/U6 di-snRNP and U4/U6.U5 tri-snRNP in subnuclear compartments called Cajal bodies (CBs) (Stănêk et al. 2003; Stănêk and Neugebauer 2004; Klingauf et al. 2006; Novotny et al. 2011, 2015). CBs are also the compartment in which the Sm class snRNAs are post-transcriptionally modified by scaRNPs (Jâdy et al. 2003). In humans, U6 snRNA is localized to CBs through its interaction with the U6 snRNP protein SART3 (aka hPrp24, p110, or Tip110) (Stănêk et al. 2003). The N-terminal half-a-tetratricopeptide (HAT) domains of SART3 are necessary for U6 localization to CBs (Stănêk et al. 2003). Localization of SART3 depends upon its interaction with coilin, the major structural CB protein (Xu et al. 2005). Interestingly, U4/U6+SART3, rather than U6=SART3, accumulates in CBs, suggesting that CBs may be a site for snRNP assembly (Stănêk and Neugebauer 2004). While formation of U4/U6 is enhanced ∼10-fold in CBs, in part due to the increased local concentration of snRNAs (Klingauf et al. 2006; Novotny et al. 2011), snRNP formation can also occur elsewhere in the nucleus as CBs are nonessential (Lemm et al. 2006). SART3 localizes within CBs transiently (on the order of a few seconds) (Dundr et al. 2004) and re-enters new CBs after a period in the nucleolasm (Stănêk et al. 2008), suggesting that SART3 (and U6 along with it) localizes to CBs during both initial biogenesis and recycling after a round of splicing. Depletion of factors needed for tri-snRNP formation (hPrp6 and hPrp8) induces formation of CBs and results in accumulation of U4/U6 in CBs (Novotny et al. 2011, 2015). Because SART3 interacts with U6 and U4/U6 but is displaced upon formation of U4/U6.U5 (Bell et al. 2002), it can anchor immature U4/U6 di-snRNPs to CBs through its interaction with coilin (Novotny et al. 2015). Localization of free U6 snRNP in subnuclear CBs may prevent incorporation of immature splicing components into active spliceosomes. Retention of U6 snRNP in CBs is a convenient checkpoint for the cell to ensure that mature U6 (in the tri-snRNP) is used in splicing.

In yeast, the U6 gene is localized near the nucleolar periphery (Belagal et al. 2016). Although yeast also have nucleolar bodies akin to CBs (Verheggen et al. 2002), little work has been done to determine if and how subnuclear localization plays a role in U6 snRNP biogenesis in yeast. By analogy, work on the biogenesis of the U3 snoRNP has demonstrated that there are differences in localization during biogenesis in yeast and humans, but that overall the processes are similar (Verheggen et al. 2002). It remains to be determined if and why yeast U6 localizes to any subnuclear compartments, and if the U4/U6.U5 tri-snRNP is assembled in a specific region in yeast nuclei.
POST-TRANSCRIPTIONAL MODIFICATION

U6 snRNA is extensively modified in metazoans, and many of these post-transcriptional modifications are evolutionarily conserved even in S. pombe. In contrast, S. cerevisiae U6 snRNA contains very few modifications, and only one known obligate modification (3′ end processing). The known post-transcriptional modifications of U6 in S. cerevisiae and humans are summarized in Table 1 and discussed below. Post-transcriptional modifications of U6 are likely to be involved in modulation of U6 interactions with other RNAs and proteins throughout the splicing cycle. However, the precise effect that modifications have on U6 RNA structure or interactions with protein and other snRNAs is poorly understood, as is the timing and cellular location of where these modifications occur.

5′ Capping

Instead of the 2,2,7-trimethylguanosine (TMG) cap that the other snRNAs possess or the 7-methylguanosine (m7G/cap-0) cap of mRNAs, human U6 possesses a γ-monomethyl phosphate 5′ modification (Singh and Reddy 1989). The γ-monomethyl cap is shared by several other Pol III transcripts, including the noncoding 7SK RNA (Gupta et al. 1990a). However, capping of U6 small nuclear RNA in vitro is not dependent upon transcription, suggesting that capping of U6 is not obligatorily cotranscriptional in vivo (Gupta et al. 1990b). Capping is dependent upon the 5′ stem loop and is also sequence-dependent (Singh et al. 1990), unlike cotranscriptional capping of Pol II transcripts. It is not known if S. cerevisiae U6 has a γ-monomethyl cap, although deletion of its first 11 nt results in installation of a TMG cap despite still being transcribed by RNA Pol III (Kwan et al. 2000). This finding implies that yeast U6 normally receives a 5′ modification that prevents TMG capping, but that disruption of conserved sequences at the base of the 5′-stem, which are the determinant for γ-monomethyl capping in humans (Singh et al. 1990), blocks this modification.

The function of the U6 cap is not known, although it may be important for stability (Shumyatsky et al. 1993). TMG capping of other snRNAs is important for transport into the nucleus, but the γ-monomethyl cap of U6 likely does not impact transport (Shumyatsky et al. 1993) or retention in the nucleus (Spiller et al. 2007a). The γ-monomethyl cap could also play a role in displacing La/Lhp1 from U6 RNA, as La interacts with the 5′ triphosphate prior to capping (Bhattacharya et al. 2002).

A 130 kDa protein responsible for cap formation in U6 and 7SK was isolated in 1994 (Shimba and Reddy 1994), but the gene was not identified until 2007 (Jeronimo et al. 2007). Bin3 (renamed methyl phosphate capping enzyme, MePCE) is responsible for installation of the γ-monomethyl cap on 7SK and U6 RNA. While U6 RNA copurified with MePCE, knockdown of MePCE reduced levels of 7SK RNA but not U6 RNA (Jeronimo et al. 2007). It is currently not clear how or to what degree the 5′ cap of U6 contributes to stability or function. Interestingly, S. cerevisiae does not have a clear homolog of MePCE, and the methylation status of the 5′ triphosphate is unknown. For a comprehensive review on the enzyme Bin3/MePCE, see Cosgrove et al. (2012).

| Modification                      | Organism | Position       | Enzyme or guide RNA |
|----------------------------------|----------|----------------|---------------------|
| Terminal modifications           |          |                |                     |
| 5′ Capping                       | Hs       | 5′ γ-Monomethyl | Bin3/MePCE          |
| 3′ End trimming                  | Sc       | 3′ Terminal 3′ phosphate | Usb1/Mpn1 |
| 3′ End oligouridylation          | Hs       | 3′ End          | U6 TU Tase          |
| 3′ End oligoadenylation          | Hs       | 3′ End          | TRAMP?              |
| Internal modifications           |          |                |                     |
| Pseudouridylation (ψ)            | Sc       | U28            | Pus1                |
|                                  | Hs       | U31            | ?                   |
|                                  |          | U40            | ?                   |
|                                  |          | U86            | ?                   |
| Ribose 2′-O-methylation (2′-O-me)| Hs       | A47            | mgU6-47 (SNORD7)    |
|                                  |          | A53            | mgU6-53 (SNORD8), mgU6-53B (SNORD9) |
|                                  |          | G54            | ?                   |
|                                  |          | C60            | MBII-166            |
|                                  |          | C62            | ?                   |
|                                  |          | C63            | ?                   |
|                                  |          | A70            | ?                   |
|                                  |          | C77            | mgU6-77 (SNORD10)   |
| N-6-adenosine methylation (m^6A)| Hs       | A43            | METTL16             |
| N-2-guanosine methylation (m^2G)| Hs       | G72            | ?                   |
3’ End modifications

After synthesis by Pol III, U6 is left with a oligo(U) tail of variable length with a terminal 2’,3’ cis diol. The oligo(U) tail and terminal 2’ and 3’ hydroxyl groups are bound by the La protein (Lhp1 in yeast) (Stefano 1984; Rinke and Steitz 1985; Terns et al. 1992). While the length of the oligo(U) tail is initially heterogeneous in part due to Pol III termination, U6 is both post-transcriptionally shortened and extended (oligouridylated) (Reddy et al. 1987).

U6 snRNA is 3’ oligouridylated in many eukaryotes, including S. pombe and humans. A U6 terminal uridylyl transferase (TUTase) activity was identified in 1998 (Trippe et al. 1998) and the gene, TUT1, was identified in 2006 (Trippe et al. 2006). Characterization of this enzyme revealed that it requires the presence of uridine at the 3’ end of U6 and that it can add three additional uridines in vitro (Trippe et al. 2003). Interestingly, TUT1 may recognize additional secondary structure in U6 and therefore may preferentially oligouridylate U6 while the RNA is in a certain conformation (Yamashita et al. 2017). TUT1 localizes to the nucleolus, suggesting that oligouridylation may occur alongside other modifications such as pseudouridylation and 2’-O-methylation (discussed in the “Pseudouridylation” and “Ribose 2’-O-methylation” sections) (Trippe et al. 2006).

The enzyme responsible for 3’ tail shortening was characterized in the late 1990’s (Booth and Pugh 1997), but the gene was not identified for an additional 15 years (Mroczek et al. 2012; Shchepachev et al. 2012; Hilcenko et al. 2013). The enzyme was named Usix biogenesis protein 1 (Usb1), and is a 3’–5’ exonuclease (Mroczek et al. 2012; Shchepachev et al. 2012; Hilcenko et al. 2013). Mutations in Usb1 (also called Mpn1) are associated with the disease poikiloderma with neutropenia in humans (for review, see Mroczek and Dziembowski 2013). Usb1 trims back the oligo(U) tail and leaves a phosphate group on the terminal nucleotide. In humans, the terminal nucleotide has a 2’-O-cyclic phosphate, while in yeast it has a 3’ noncyclic phosphate (Lund and Dahlberg 1992; Didychuk et al. 2017). The majority of human U6 snRNAs end with five terminal uridines and a 2’,3’-cyclic phosphate (Shchepachev et al. 2015).

Some metazoa do not have a homolog of Usb1. These organisms, including C. elegans, do not possess the phosphoryl or cyclic phosphate modifications, but instead have either a 2’,3’-cis diol in their mature form or an intriguing “blocked” modification (Lund and Dahlberg 1992). Interestingly, C. elegans does not have a Usb1 homolog, but rather has a TUTase called USIP-1 (Rüegger et al. 2015). The USIP-1 enzyme interacts with the U6 snRNP protein SART3 (discussed in the “Prp24/SART3” section). The chemical identity of the 3′ end produced by USIP-1 is not known.

The 3’ tail of U6 can also be adenylated (Chen et al. 2000). Adenylation inhibits uridylation and may target U6 for degradation. In cells deficient for Usb1 activity, U6 with adenylated tails accumulate (Hilcenko et al. 2013; Shchepachev et al. 2015). Polyadenylation may result in targeting U6 to the nuclear exosome. Human Usb1 is active on polyadenylate tails, and therefore can counteract polyadenylation (Hilcenko et al. 2013). It is unclear what the polyadenylating enzyme is, or if it can function on the product of human Usb1, a 2’,3’-cyclic phosphate. An additional phosphatase enzyme may be necessary to remove the cyclic phosphate and allow the polyadenylating enzyme to function.

The timing and location of 3’ end modification by Usb1 is not known. An early study suggested that the length and modification of U6 changed during the splicing cycle (Tazi et al. 1993). Tazi et al. made the intriguing observation that 3’ end modification of U6 required pre-mRNA, raising the possibility that U6 may not be processed until after a round of splicing is complete. In this way, after a successful first round of splicing, the cell could mark functional U6 to be protected and kept for additional rounds of splicing. However, a later study showed that depletion of U1 and U2 snRNAs had no effect on U6 processing (Gu et al. 1997). It is possible that the 3′ end of U6 is altered throughout the splicing cycle, as the 3′ end binding Lsm2-8 complex (discussed in the “Lsm2-8” section) dissociates during spliceosome activation, which may render the U6 3′ end accessible for modification. Interestingly, Usb1 localizes to the nucleus but not the nucleolus (Mroczek et al. 2012). This finding implies that in humans, Usb1 recognizes a TUT1-extended U6 in the nucleoplasm after modification in the nucleolus but before it has traveled to Cajal bodies for di- and tri-snRNP assembly. It is unclear if TUT1 can function on the terminal 2’,3’-cyclic phosphate of mature, Usb1-processed U6, and it is also unclear if uridylation occurs only once after transcription termination, or if U6 is trimmed and extended repeatedly throughout its life cycle.

Modification of the U6 3′ end is important for recognition by U6 3′ end binding proteins. The La protein binds U6 immediately after transcription termination and specifically recognizes the 2’,3’-cis diol. Indeed, while a subset of U6 molecules found in the cell have cis diols and are bound by La (Rinke and Steitz 1985), maturation of U6 and incorporation into the U6 snRNP requires the 3′ modification by Usb1. Modification of the 3′ end reduces the ability of La to bind U6 (Terns et al. 1992) and is important for binding of the Lsm2-8 ring (Licht et al. 2008). Interestingly, human Lsm2-8 preferentially binds U6 containing a 2’,3’-cyclic phosphate modification (Licht et al. 2008), while S. cerevisiae Lsm2-8 prefers to bind a noncyclic phosphate, suggesting that Usb1 and Lsm2-8 coevolved to allow for tight binding by Lsm2-8 on the product of Usb1 activity (Didychuk et al. 2017).

Pseudouridylation

Pseudouridylation of RNA is thought to be structurally stabilizing because of increased base stacking (Davis 1995) as well as the added hydrogen bonding potential and conformational flexibility due to the C–C glycosidic bond (Charette and Gray...
The effects of pseudouridylation on structure and RNA–protein interactions in U2 snRNA have been well-studied (Yu et al. 1998; Newby and Greenbaum 2001; Wu et al. 2016; van der Feltz et al. 2017), but the effects of pseudouridylation on U6 RNA have not. In *S. cerevisiae*, U6 is not constitutively pseudouridylated (Massenet et al. 1999). Recently, position U28 was found to be inducibly pseudouridylated during filamentous growth by the pseudouridine synthase Pus1 (Basak and Query 2014). Interestingly, mutations within U6 far from U28 (U36C and G50U) increased pseudouridylation of U28. These mutations likely disrupt interactions of U6 with U6 snRNP protein Prp24 (Montemayor et al. 2014) and would also likely alter interactions within the activated spliceosome. It is not clear at which step in the U6 lifecycle U6–U28 is pseudouridylated, or how changing RNA–protein interactions elsewhere in the RNA increases pseudouridylation of U6–U28. Furthermore, it is not clear how pseudouridylation of U6–U28 would affect RNA–RNA or RNA–protein interactions in different splicing complexes. In the U4/U6.U5 tri-snRNP, U6–U28 interacts with U5–U99 (Nguyen et al. 2016), whereas in the C complex, U6–U28 is within 5 Å of Bud31 and Ecm2 (Galej et al. 2016). Understanding how pseudouridylation of U6–U28 affects RNP structure, and understanding how pseudouridylation of U6 induces filamentous growth in yeast, could reveal important new insights into spliceosome biology.

In contrast, human U6 and U6atac are constitutively pseudouridylated at several positions (Massenet and Branlant 1999; Massenet et al. 1999). These positions (U31, U40, and U86 in U6; U83 in U6atac) (Fig. 4; Massenet and Branlant 1999) are in functionally important parts of the RNA, although it is unclear if or how modifications in U6 can stabilize its structure. Recently solved cryo-EM structures have allowed for structural analysis of splicing complexes; however, they lack the resolution to unambiguously identify post-transcriptional modifications. Improved resolution may lead to a better understanding of how modifications in U6 function.

Localization in the nucleolus is sufficient for formation of both the three pseudouridines in human U6 and the eight 2′-O-methyl groups (discussed in the “Ribose 2′-O-
methylation” section), suggesting that all of the machinery required for these modifications is present in the nucleolus (Ganot et al. 1999). Localization of U6 during its biogenesis is discussed in the “Localization During Biogenesis” section.

**Ribose 2′-O-methylation**

Ribose 2′-O-methylation is a ubiquitous modification that can significantly stabilize RNA, either structurally (Sashital et al. 2007) or metabolically (by reducing hydrolytic or nucleolytic cleavage). Like pseudouridylation, ribose methylation occurs in the nucleolus (Ganot et al. 1999). Human U6 contains eight ribose methylations, at positions A47, A53, G54, C60, C62, C63, A70, and C77, which could stabilize secondary structure either in the U6 snRNP (the internal stem loop; ISL) or in U4/U6 (Stem I and Stem II) (Fig. 5). Ribose methylation requires a methyltransferase enzyme and “methylation guide” (mg) snoRNAs, including mgU6-47 and mgU6-77 (Tycowski et al. 1998), mgU6-53 (Ganot et al. 1999), and MBII-166 (which modifies U6-C60) (Hüttenhofer et al. 2001). Modification of many of the U6 ribose groups are conserved in plants (Kiss et al. 1987) and in *S. pombe* (Gu et al. 1996). The mgU6-47 snoRNA is conserved in *S. pombe* and is required for methylation. However, while disruption of the mgU6-47 gene resulted in complete loss of methylation, only a small cold-sensitive splicing defect was observed (Zhou et al. 2002). Interestingly, human U6atac has no ribose 2′-O-methyl groups, despite forming a homologous ISL in which ribose methylations in U6 are concentrated (Massenet and Branlant 1999).

Recent RiboMeth-seq results show that the level of ribose modification varies in different cell types (Krogh et al. 2017).
While ribose methylation may not be essential, its conservation across species suggests it (along with pseudouridylation and other modifications) may play a role in fine-tuning the stability of U6 and its complexes to modulate splicing, perhaps in response to variation in cellular temperatures that occur even in homeotherms.

**N-6-adenosine and N-2-guanosine base methylation**

Human U6 contains a single N-6-methyladenosine modification (m^6A) at position A43 (Shimba et al. 1995). Study of m^6A modification has recently undergone a renaissance due to transcriptome-wide mapping studies that show it is a ubiquitous and regulated modification in mRNA (for review, see Meyer and Jaffrey 2014). Most mRNA m^6A residues are found within a consensus sequence (GAG or AAG) and are modified by the METTL3 complex. U6–A43 does not lie within a consensus sequence and is not modified by METTL3, but rather by a recently identified methyltransferase, METTL16 (Pendleton et al. 2017; Warda et al. 2017). Interestingly, METTL16 activity is dependent on both sequence and secondary structure. This is consistent with earlier reports that modification of U6–A43 depends on formation of the adjacent ISL secondary structure within U6 (Shimba et al. 1995). It is unclear when A43 is modified to m^6A, but in vitro studies suggest that the conversion is rapid (Shimba et al. 1995). A recent report showed that METTL16 associates on U6 alongside the capping enzyme MePCE (see the “5′ Capping” section) and the 3′ end binding protein La (Warda et al. 2017). Determining the order of U6 modifications and the effect of multiple modifying enzymes competing for the same substrate may yield interesting insights into U6 biogenesis.

m^6A-modified U6 is incorporated into U4/U6 snRNPs, suggesting it is functional (Briggsman and Luhrmann 1987). The modification at position A43 lies in the center of the ACAGAGA motif, a perfectly conserved and essential motif that base pairs to the intron adjacent to the 5′ splice site and helps organize the spliceosome active site through formation of base triple interactions (Fica et al. 2014). Biophysical studies of m^6A modification within double-stranded RNA indicate that it destabilizes base-pairing, but at the ends of helices is stabilized when stacked and unpaired (Roos et al. 2015). In a recent structure of a human C+ complex, U6–A43 is unpaired across from an adenosine in the intron (Bertram et al. 2017b). The methylation state of the nucleotide could not be determined in the structure.

Human U6 also contains a single N-2-guanosine methylation (m^2G) at position G72 (Epstein et al. 1980). The role of this modification is unknown, as G72 is expected to fall within a G–C base pair in both the U6 ISL and within U4/U6 Stem II (Figs. 4, 5) and because m^2G is expected to be isoenergetic with guanosine in G–C base pairs (Rife et al. 1998). Like base methylation of A43, methylation of G72 may also depend on secondary structure (Ganot et al. 1999).

**Splicing of U6 snRNA**

The sole copy of the U6 gene in *S. pombe* has an intron that is removed by the spliceosome (Tani and Ohshima 1989). The intron is located immediately adjacent to the highly conserved catalytic AGC triad that is essential for splicing catalysis, leading to the hypothesis that the intron in *S. pombe* U6 RNA arose from reverse-spooling of an excised intron into a U6 molecule in the active spliceosome, followed by reverse transcription and incorporation into the genome (Brow and Guthrie 1989; Tani and Ohshima 1991). This observation suggested that U6 (and specifically this region of U6) is near the active site of the spliceosome and that splicing is reversible. Indeed, both of these hypotheses have been tested using biochemistry and structural biology. Splicing is reversible under the appropriate conditions (Tseng and Cheng 2008), and it is now well-established both biochemically and in near-atomic structural detail that this region of U6 forms the catalytic core of the spliceosome. Recent bioinformatics analysis shows that many yeast species have evolutionarily unrelated introns and that these introns are distributed throughout the U6 sequence (Canzler et al. 2016), although intron insertion sites are more concentrated near the catalytic core (U6 ISL and ACAGA box) and in a region upstream of the ACAGA box known to function as a branch acceptor for in vitro trans-splicing in nematodes (Yu et al. 1993). These introns (up to four introns in a single fungal U6 gene) possess canonical 5′ splice sites, branch sites, and 3′ splice sites, but are otherwise not conserved in sequence. Phylogenetic analysis suggests that while the presence of some introns in related species likely arose from a common ancestor, many of the U6 introns may be species-specific insertions (Canzler et al. 2016). Experimental validation that U6 genes containing introns are expressed and spliced is necessary. It is possible that species with multiple U6 paralogs of which some contain introns may use splicing as a method to regulate U6 levels.

Spliceosomal introns can also be found in U2 (Takahashi et al. 1993) and U5 snRNAs (Takahashi et al. 1996), supporting the hypothesis that RNAs close to the active site of the spliceosome can be targets of reverse splicing. However, spliceosomal introns have also been identified in U1 snRNA (Takahashi et al. 1996), which departs before the active site of the spliceosome is formed, and in rRNA genes (Bhattacharya et al. 2000). It is not clear what the determinants of novel spliceosomal intron insertion in ncRNAs are, but possibilities include RNA abundance, structure, or proximity to the spliceosomal active site.

**THE U6 SNRNP**

**U6 snRNA secondary structure**

U6 is the most conserved of the snRNAs in sequence and length (Brow and Guthrie 1988). The least conserved region is the 5′ stem–loop (5′SL), which varies in both sequence and
length across eukarya despite maintaining a stem–loop structure (Roiha et al. 1989). *C. merolae*, a moderate thermophile, contains a minimal spliceosome that lacks U1 snRNA, yet has an unusually stable 5′SL with 22 base pairs (versus 10 base pairs in *S. cerevisiae*) (Stark et al. 2015). The 5′SL persists throughout the various U6-containing splicing complexes. It does not have any known catalytic function, but does correspond to part of the A-block promoter element in *S. cerevisiae* and is necessary for efficient 5′ capping in humans (discussed in the “5′ Capping” section). The 5′ capping determinants are conserved in yeast phyla (Roiha et al. 1989), further evidence that 5′ capping is also conserved.

The remaining portion of U6 forms several mutually exclusive structures. While in the U6 snRNP, U6 consists of the 5′SL, the telestem, and the internal stem–loop (ISL, previously called the 3′ stem–loop or intramolecular stem–loop) (Figs. 4, 5). While the 5′SL likely remains stably base-paired throughout the splicing cycle, the less stable telestem and ISL helices make dynamic transitions into other secondary structures during spliceosome assembly and activation. Incorporation of U6 into the spliceosome requires unwinding of the ISL and subsequent base-pairing to U4, a mutually exclusive interaction. As observed in the U4/U6.U5 tri-snRNP structure (Agafonov et al. 2016; Nguyen et al. 2016; Wan et al. 2016b), the telestem is unwound when U6 is paired to U4 snRNA. However, in vitro data suggest the telestem can form transiently in free U4/U6 RNA (Brow and Vidaver 1995; Rodgers et al. 2016). During spliceosome activation, U4 is unwound from U4/U6 by the helicase Brr2 (Laggerbauer et al. 1998; Raghnunathan and Guthrie 1998a), U6 base pairs with U2 (Madhani and Guthrie 1992), and the U6 ISL reforms (Fig. 5). The telestem is mutually exclusive with formation of U2/U6 helix II (Fig. 5). The telestem enhances Prp24 affinity and Prp24-mediated annealing (Didychuk et al. 2016) and also destabilizes U4/U6 (Brow and Vidaver 1995; Rodgers et al. 2016). The dynamic telestem and U6 ISL structures may help U6 molecules avoid kinetic or thermodynamic folding traps during spliceosome assembly and activation.

Essential nucleotides in the U6 ACAGAGA box and the AGC triad were identified early on (Fabrizio and Abelson 1990; Madhani et al. 1990). During activation, the ACAGAGA box interacts with the 5′ splice site (Kandels-Lewis and Séraphin 1993; Lesser and Guthrie 1993; Johnson and Abelson 2001; Rauhut et al. 2016; Yan et al. 2016). The AGC triad is important for intramolecular base-pairing within U6 as well as pairing to U2 (in U2/U6 helix I) and U4 (in U4/U6 Stem I) (Fig. 5). The AGC triad makes intramolecular base-triple tertiary contacts with the last 2 nucleotides of the ACAGAGA sequence (Hilliker and Staley 2004; Fica et al. 2014, 2017; Galej et al. 2016; Rauhut et al. 2016; Wan et al. 2016a; Yan et al. 2016, 2017; Bertram et al. 2017b; Zhang et al. 2017). The AGC triad and base triple interactions are conserved in the self-splicing Group II introns, which share a common evolutionary ancestor with the spliceosome (Pyle 2016). A conserved bulged residue in the ISL (U80 in *S. cerevisiae*) is involved in catalysis by coordinating a catalytic metal ion and by forming a base triple with the C of the AGC triad (Fica et al. 2013, 2014). Mutation of U80 to C or A has no obvious effect on yeast growth, while mutation to a G is lethal unless U6–C67 is mutated to prevent pairing across the ISL and U4–G14 is mutated to maintain pairing with U6 residue 67 (McManus et al. 2007). Thus, the tertiary structure of this region of the ISL is more important than the primary structure.

U6atac lacks a 5′SL, but contains a similar central secondary structure (Fig. 4). Indeed, the yeast or human U6 ISL can functionally replace the U6atac ISL (Shukla and Padgett 2001). U6atac contains an additional stem loop 3′ of the ISL, which is important for targeting to the minor spliceosome (Dietrich et al. 2009).

**Prp24/SART3**

The yeast Prp24 protein was discovered through a genetic selection for suppressors of mutations in U4 RNA that disrupt U4/U6 pairing (Shannon and Guthrie 1991) as well as a screen for heat-sensitive mutations in splicing (Vijayraghavan et al. 1989). Mutations in Prp24 were also isolated in a selection for suppressors of a mutation that stabilizes the U6 ISL (Vidaver et al. 1999; Montemayor et al. 2014), underscoring its importance for the structural transitions of U6 snRNA. In *S. cerevisiae*, Prp24 contains three canonical RNA recognition motifs (RRMs) and a fourth “occluded” RRM in which the typical RNA binding surface, a four-stranded β-sheet, is masked by additional helices (Rader and Guthrie 2002; Kwan and Brow 2005; Bae et al. 2007; Martin-Tumasz et al. 2011; Montemayor et al. 2014). The human homolog of Prp24, SART3, has only two RRMs (which may correspond to yeast RRMs 1 and 2 or 2 and 3), and instead has multiple N-terminal half-a-tetracopeptide (HAT) repeats (Bell et al. 2002; Rader and Guthrie 2002; Kwan and Brow 2005). SART3 has recently been shown to dimerize through these HAT repeats (Park et al. 2016; Zhang et al. 2016), suggesting that Prp24 and SART3 may both require four RRMs to function efficiently. Prp24 from *S. pombe* may be an evolutionary intermediate in that it has four RRMs plus the HAT domains. All Prp24 homologs contain a highly conserved short peptide at the C terminus called the SNFFL box, which is important for interactions between Prp24 and Lsm2-8 (Fromont-Racine et al. 2000; Rader and Guthrie 2002). Prp24 is essential for unwinding the U6 ISL and annealing U4 to U6 (discussed in the “U4/U6 Annealing” section) (Ghetti et al. 1995; Rader and Guthrie 2002; Didychuk et al. 2016).

**Lsm2-8**

The Lsm proteins are paralogs of the Sm proteins that bind to U1, U2, U4, and U5 snRNAs, and are related to bacterial Hfq
and Sm-like archaeal proteins (Mura et al. 2013). Lsm proteins are small (9–21 kDa) and assemble into at least 2 different heterohtetrameric, ring-shaped complexes, Lsm1-7 and Lsm2-8. The Lsm1-7 ring localizes to the cytosol and is involved in the 5′–3′ mRNA decay pathway (Tharun et al. 2000). The Lsm2-8 ring localizes to the nucleus where it binds the 5′ U-tail of U6 (Achsel et al. 1999; Mayes et al. 1999). The \textit{S. cerevisiae} Lsm-2-8 complex preferentially binds U6 3′ ends modified with a terminal phosphate (Didychuk et al. 2017), while the metazoan Lsm2-8 complex preferentially binds a terminal 2′,3′-cyclic phosphate (Licht et al. 2008), as discussed above in the "3′ End Modifications" section. Structures of the Lsm1-7 and Lsm2-8 complexes revealed that the two rings are exceedingly similar, with <1 Å rmsd between subunits Lsm2-7 in each structure and between the core Sm folds of Lsm1 and Lsm8 (Sharif and Conti 2013; Zhou et al. 2014,a,b). The crystal structure of the \textit{S. cerevisiae} Lsm-2-8 bound to a short RNA oligo was determined and revealed that the 3′ end of U6 is recognized in the center of the ring in a manner distinct from Sm-RNA binding (Zhou et al. 2014,a). The Lsm2-8 complex can also be observed in cryo-EM structures of the U4/U6. U5 tri-snrnp, albeit at lower resolutions (Agafonov et al. 2016; Nguyen et al. 2016; Wan et al. 2016b).

Unlike the Sm ring, the Lsm2-8 ring is stable in the absence of RNA (Achsel et al. 1999) and assembles without assistance from other proteins. The Sm ring, in contrast, requires the SMN complex and Gemin proteins to assemble three Sm subcomplexes into a ring that encircles RNA in vivo (for reviews, see Battle et al. 2006; Matera and Wang 2014). The Lsm2-8 ring is likely imported into the nucleus via importin β/KAP95 (Spiller et al. 2007,a), where the N- and C-terminal domains of Lsm8 act as nuclear retention signals for U6 (Spiller et al. 2007,a,b; Reijns et al. 2009).

Structure of the U6 snRNP

The structure of the core of the U6 snRNP, with the majority of both U6 snRNA (nucleotides 30–101) and Prp24 (all four RRMs), has been determined (Montemayor et al. 2014). To date, this is the only structure of a protein containing four RRMs bound to RNA. It reveals how three of the RRMs cooperate to specifically and tightly bind U6 RNA. RRMs typically bind 3′–4′ single-stranded nucleotides across a β sheet surface using consensus RNP1 and RNP2 motifs (Afroz et al. 2015). In contrast, the Prp24 protein directly contacts 20 nt of U6 RNA via interactions with RRM domains 2–4. The first RRM does not contact U6 snRNA, but forms a large protein–protein interface with RRM2. RRMs 2 and 3 bind the asymmetric bulge in a relatively canonical RRM fashion, targeting the ACAGA box and sequences immediately upstream, respectively. The fourth RRM binds primarily to the ISL and telestem via the two amphipathic helices that occlude the β-sheet face of the RRM (Martin-Tumasz et al. 2011). Surprisingly, the RRM3/RRM4 linker, a single asparagine residue, passes through the asymmetric bulge and RRM4 makes tertiary interactions with RRM2, forming an entwined topology consisting of interlocked rings of protein and RNA (Montemayor et al. 2014). The entwined topology is further closed off by the ISL and telestem and is exceptionally stable, as the complex is resistant to 2 M monovalent salt (Shannon and Guthrie 1991). RRMs 1, 2, and 4 together form a large (~20 Å wide), positively charged groove. In the crystal lattice, the U6 ISL of a neighboring complex is packed into the groove, leading to the hypothesis that this groove might stabilize U4/U6 pairing during Prp24-mediated stimulation of U4/U6 di-snrnp formation (Montemayor et al. 2014).

The U6-Prp24 structure contained three substitutions to stabilize the RNA (Montemayor et al. 2014) including A62G, which stabilizes the base of the ISL and causes a cold-sensitive phenotype in vivo (Fortner et al. 1994). Interestingly, suppressors of U6–A62G cold sensitivity localize to the RNA–protein interface, suggesting that detrimental overstabilization of the U6 ISL structure can be neutralized by destabilization of the RNA–protein interface in vivo (Montemayor et al. 2014). A second structure with the wild-type ISL crystallized in a different space group, yet exhibits the same interlocked topology (Montemayor et al. 2017). Stabilization of the telestem with U100C/U101C mutations, which increase both Prp24 affinity for U6 as well as Prp24-mediated U4/U6 annealing activity (discussed in the "U4/ U6 Annealing" section), likely also stabilizes the interlocked topology (Didychuk et al. 2016). Together, these data strongly suggest that the interlocked topology is a biologically functional structure.

The structure of the entire snRNP, with full-length RNA and including both Prp24 and Lsm2-8, remains unsolved. It is known that Prp24 and Lsm2-8 have at least one protein–protein interaction (via the Prp24 SNFFL box), but where this interaction occurs on the Lsm2-8 ring and how the individual components are oriented with respect to one another has not been determined. Uncovering how Prp24 and Lsm2-8 interact will undoubtedly be vital to understanding the mechanism of Prp24-mediated U4/U6 annealing.

U4/U6 DI-SNRNP AND U4/U6.U5 TRI-SNRNP

U4/U6 annealing

In order to be incorporated into the assembling spliceosome, the ISL of U6 must be completely unwound and base-paired to U4 snRNA via the RNA chaperone activity of Prp24. This assembly step may proofread U6 structure by ensuring that only U6 that can bind Prp24 and base pair to U4 is incorporated into the spliceosome. In addition, U4 may act as an “antisense negative regulator” of the catalytic residues of U6 by sequestering them in the double-stranded helices of U4/U6 prior to spliceosome activation (Guthrie 1991).
It has long been known that the U6 snRNP protein Prp24 can anneal U4 and U6 RNAs in an ATP-independent manner (Ghetti et al. 1995; Raghunathan and Guthrie 1999b). The 20 Å wide electropositive groove of Prp24 is critical for U4/U6 annealing but not binding of U6, suggesting that it is the "active site" for U4/U6-annealing (Didychuk et al. 2016). Stabilization of the telomest in U6 also contributes to efficient annealing, possibly by promoting formation of the electropositive groove (Didychuk et al. 2016). The Lsm2-8 ring enhances Prp24-mediated annealing in vitro, but does not facilitate U4/U6 annealing by itself (Didychuk et al. 2016). Interestingly, Prp24 binds the U4/U6 product complex in vitro (Ghetti et al. 1995; Didychuk et al. 2016) yet does not stay associated with the U4/U6 di-snRNP in yeast in vivo (Shannon and Guthrie 1991). How the interlocked topology of U6•Prp24 is resolved during annealing, and how Prp24 is displaced from the U4/U6 di-snRNP, is unclear.

The upper ISL is not contacted by protein in either of the determined U6 snRNP core structures. In the ISL-stabilized (A62G) structure, the top of the ISL is disordered and not visible (Montemayor et al. 2014). In the wild-type U6 snRNP core structure, there are two complexes in the asymmetric unit (Montemayor et al. 2017). In one, the ISL is largely disordered and in the other, the ISL displays a 20° bend relative to the A62G ISL (Montemayor et al. 2017). These observations, along with NMR measurements and MD simulations (Blad et al. 2005; Venditti et al. 2009), suggest that the U6 ISL is inherently dynamic. Since the ISL must be completely unwound during U4/U6 annealing, dynamics likely play an important role in the annealing pathway. The stabilized A62G ISL in isolation has a melting temperature of ~63°C in 200 mM KCl (Sashital et al. 2003) while the wild-type ISL is less stable. Binding of Prp24 may facilitate U4/U6 formation by allowing the ISL to remain in a dynamic, meta-stable conformation while adjacent to the electropositive groove in Prp24. It is unlikely that the ISL needs to fully melt in order for annealing to be initiated, as local unwinding of a few base pairs could allow formation of a nascent U4/U6 duplex to invade the ISL and facilitate the subsequent exchange of intra- for inter-molecular base pairs via branch migration (Meselson 1972; Sigal and Alberts 1972).

In humans, SART3 (hPrp24) functions in a similar role in both U4/U6 annealing (Bell et al. 2002; Medenbach et al. 2004) and U4atac/U6atac annealing (Damianov et al. 2004). Although the HAT domains are dispensable for binding U6, both the HAT domains and the two RRMs are required for efficient U4/U6 recycling (Bell et al. 2002; Medenbach et al. 2004). Surprisingly, the C-terminal tail (the SNFFL box), which is known to interact with the Lsm2-8 ring (Rader and Guthrie 2002), is not necessary for efficient recycling in vitro, but an interaction with the U4 snRNP protein 90K (hPrp3) mediated by the HAT domains of SART3 is necessary (Medenbach et al. 2004). It has also been reported that human Lsm2-8 can recycle U4/U6 in the absence of other factors (Achsel et al. 1999). As monomeric SART3 has only two RRMs, it is unlikely to form an electropositive groove as observed in the yeast U6•Prp24 structure, although it may potentially do so via the aforementioned dimerization of the HAT domains (Park et al. 2016). Alternatively, the role of the electropositive groove of Prp24 may have been functionally replaced by protein–protein interactions between SART3 and hPrp3 (Medenbach et al. 2004).

The U4 snRNP proteins Snu13 and Prp31 and U4/U6 di-snRNP proteins Prp3/Prp4 (plus CypH in humans) may also contribute to the annealing process. Snu13 and Prp31 influence the conformation of the U4 5′SL, which may hold U4 RNA in a conformation amenable to annealing with U6 (Hardin et al. 2015). Indeed, the structure of protein-free U4/U6 (Cornilesu et al. 2016) is significantly different from its structure while in U4/U6.U5 tri-snRNP (Nguyen et al. 2016). Prp3 binds U4/U6 Stem II and a region on U6 3′ of the stem, suggesting that its binding site does not exist until U4/U6 is formed (Liu et al. 2015; Nguyen et al. 2016). However, it may transiently capture this single-stranded region of U6 from U6•Prp24 to accelerate U4/U6 annealing (Liu et al. 2015). The contribution of additional proteins to Prp24-mediated annealing has not been measured, but may reveal additional RNA–protein or protein–protein interactions that accelerate the assembly process. Understanding how the mechanism of Prp24-mediated annealing in both yeast and humans could provide new insight into conserved versus divergent mechanisms of RNA chaperones.

**Formation and structure of the U4/U6.U5 tri-snRNP**

The tri-snRNP is formed through protein–protein and protein–RNA interactions between the U4/U6 di-snRNP and the U5 snRNP. Prp31 is necessary for assembly of both major and minor U4/U6.U5 tri-snRNPs in humans (Schneider et al. 2002). Knockdown of both hPrp31 and the tri-snRNP-specific protein hPrp6 leads to an accumulation of U4/U6 di-snRNPs in Cajal bodies (Novotny et al. 2015) (discussed in the "Localization During Biogenesis" section). The structure of the tri-snRNP from yeast and humans has revealed myriad protein–protein contacts that hold the U4/U6 di-snRNP and U5 snRNP together (Agafonov et al. 2016; Nguyen et al. 2016; Wan et al. 2016b). Prp6 is the “glue” holding the tri-snRNP together, as its N-terminal region contacts U5 snRNP component Prp8 (and Brr2 in the human tri-snRNP) while its C-terminal region contacts U4 snRNP components Snu13, Prp31, Prp3, and Prp4 (Galisson and Legrain 1993; Agafonov et al. 2016; Nguyen et al. 2016; Wan et al. 2016b). In the presence of mutations that destabilize U4/U6 pairing and U6-Prp24 binding, tri-snRNP still assembles in the apparent absence of U4/U6 di-snRNP, suggesting there may be an alternate pathway for incorporating U4 and U6 into the tri-snRNP (Burke et al. 2015).
The structure of the U4/U6.U5 tri-snRNP revealed an unexpected short internal stem loop in U6 5′ of U4/U6 Stem I consisting of nucleotides 35–46 (Nguyen et al. 2016). Interestingly, this stem loop is immediately upstream of the ACAGAGA sequence, which must pair to the 5′ SS. This ACAGAGA stem loop (ASL) is stabilized by Dib1 and Prp8 and is also present in the B complex structure (Plaschka et al. 2017). Consistent with prior genetic and biochemical data indicating that the intron 5′ SS binds first to an upstream ACA sequence present in the loop of the ASL prior to binding the ACAGA box (Li and Brow 1996; Johnson and Abelson 2001), the 5′ exon–intron junction interacts with the ASL loop in the B complex (Plaschka et al. 2017). As Dib1 is not observed in B act or subsequent catalytic structures, it likely departs during activation and the ASL unwinds, promoting recognition of the intron by the ACAGA box and allowing catalysis to occur. The ASL was not resolved in the structure of the human tri-snRNP, despite Dim1 (the homolog of Dib1) binding near this region (Agafonov et al. 2016). The RNA–RNA and protein–RNA contacts in the tri-snRNP effectively sequester all of the catalytic elements of U6: The ACAGAGA sequence is partially occluded by the ASL and its associated proteins, the AGC triad is base-paired in U4/U6 Stem I, and U80 is base-paired in U4/U6 Stem II.

The most significant difference in the structures of U4/U6. U5 tri-snRNP from human and yeast is in the placement of the helicase Br2. In the yeast tri-snRNP, Br2 binds U4 in a single stranded region between the U4 3′ SL and U4/U6 Stem I, suggesting that it is poised for unwinding U4/U6 (Nguyen et al. 2016). Indeed, tri-snRNP isolated from yeast cell extract is disassembled in the presence of ATP (Raghunathan and Guthrie 1998a; Nguyen et al. 2015). In contrast, Br2 in the human tri-snRNP is on the opposite end of the complex ~20 nm away, far from its U4/U6 Stem I substrate, and the tri-snRNP is insensitive to ATP-dependent disassembly (Agafonov et al. 2016). This difference is likely due to the presence of Sad1 in the human tri-snRNP, where Sad1 both stabilizes the interaction between U4/U6 and U5 snRNPs and tethers Br2 in a “preactivation” position away from its substrate (Agafonov et al. 2016). Yeast Sad1, in contrast, does not copurify with the tri-snRNP (Stevens et al. 2001; Nguyen et al. 2016). Despite interacting weakly with the tri-snRNP in yeast, Sad1 still prevents Br2-mediated unwinding in whole cell extract (Huang et al. 2014).

The minor spliceosome tri-snRNP (U4atac/U6atac.U5) contains many of the same components as the major spliceosome U4/U6.U5 (Schneider et al. 2002). The additional 3′ stem loop in U6atac (Fig. 4) may differentiate the major and minor spliceosomes, as this stem loop is both important for targeting U6atac to the minor spliceosome (Dietrich et al. 2009) and for interacting with protein 65K, a U11/U12-specific protein (Singh et al. 2016). It remains to be seen if the major and minor spliceosomes have additional conformational or compositional differences, and how the structures of the major and minor tri-snRNP differ.

U6 in the Catalytic Spliceosome

Activation

U6 base pairs with the 5′SS after remodeling of the U1-5′SS interaction (Sawa and Abelson 1992) by the DEAD-box helicase Prp28 (Staley and Guthrie 1999). This is an important proof-reading step to ensure that stable and correct U6-5′SS duplexes are formed (Yang et al. 2013). U2 base pairs to U6 via formation of U2/U6 helix II (Fig. 5; Wassarman and Steitz 1992; Schneider et al. 2010). Then, U4/U6 Stems I and II are completely unwound in order for the U6 ISL to reform and for U6 to base pair with U2 (Fig. 5). Ejection of U4 depends on the U5-snRNP component helicase Br2, which unwinds U4/U6 (Raghunathan and Guthrie 1998a; for review, see Absmeier et al. 2016). Br2 activity is regulated by protein–protein contacts with itself and Prp8, which can both up- and down-regulate its unwinding activity (Maeder et al. 2009; Mozaffari-Jovin et al. 2012, 2013; Absmeier et al. 2015). The spliceosomal GTPase, Snu114, influences regulation of Br2 activity by Prp8 (Small et al. 2006). Intramolecular folding of U6 contributes to efficient unwinding (Rodgers et al. 2016; Theuser et al. 2016). Once unwound, U4 snRNA departs with Snu13 and Prp31 (and likely the U4 5′m ring), while Prp3/Prp4 are released separately (Theuser et al. 2016).

Br2-mediated unwinding and U4 snRNP departure typically occur before arrival of the NTC (Hoskins et al. 2016). Recent data on human spliceosomes chemically stalled between the transition from B to B act reveal that the Lsm2-8 ring is present on U6 after U4 departure (Sidorovich et al. 2017). The NTC is required for dissociation of the Lsm ring in yeast (Chan et al. 2003) and promotes interactions between U5, U6, and the 5′SS prior to first step chemistry (Chan and Cheng 2005). Recent structures of the spliceosome after activation show that the 3′ end of U6 (the binding site of the Lsm ring) is partially occluded by Syf1 of the NTC, suggesting that the NTC may directly displace the Lsm ring. It is not clear if 3′ end modifying enzymes such as Usb1 and U6-TUTase can access the 3′ end of U6 in these complexes, but high-throughput studies suggest that Usb1 may interact with components of the NTC (Hazbun et al. 2003; Krogan et al. 2006; Yu et al. 2008).

Conformational changes during catalysis

In the past two years, a multitude of spliceosomal structures have been determined at different catalytic steps. Structures of spliceosomes paused before activation (B complex) (Bertram et al. 2017a; Plaschka et al. 2017) the first catalytic step (B act complex) (Rauhut et al. 2016; Yan et al. 2016), after first step chemistry (C complex) (Galej et al. 2016; Wan et al. 2016a), remodeled before the second catalytic step (C* complex) (Bertram et al. 2017b; Fica et al. 2017; Yan et al. 2017; Zhang et al. 2017), and after ligated exon departure (ILS)
(Yan et al. 2015; Wan et al. 2017) reveal that, despite large-scale conformational changes elsewhere in the spliceosome (Table 2), the structure of U6 snRNA changes very little during these transitions. Indeed, the structure of U6 snRNA at each of the steps after activation of B complex is nearly indistinguishable (Fig. 6A), even when comparing structures isolated from S. cerevisiae and S. pombe. U2/U6 Helix Ia and Ib (composed of nucleotides 55–61 in U6) and Helix II (nucleotides 93–102 in U6) remain base-paired and unchanged throughout the structures (Fig. 6B). The ISL (nucleotides 63–84) is also structurally identical. Yeast U6 residue C66 is bulged from the ISL and sits in a pocket composed of the NTC and NTC-associated proteins Clf1, Cef1, Prp45, and Ecm2. The structure of the ISL within the spliceosomal complexes is significantly different from the free RNA or when within the U6 snRNP due to the many protein contacts surrounding it. The 3′ tail of U6 (nucleotides 105–112) is unstructured in all available cryo-EM structures and is not

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TABLE 2. RNA–RNA and protein–RNA contacts in U6-containing complexes

| U6 snRNP (PDB 4N0T) | U4/U6.U5 tri-snRNP (PDB 5GAN) | B complex (PDB 5NRL) | B* complex (PDB 5GM6) | C complex (PDB 5LJS) | C* complex (PDB 5MQ0) |
|----------------------|--------------------------------|---------------------|-----------------------|---------------------|-----------------------|
| U4 snRNA             | U2 snRNA                       | U2 snRNA            | U2 snRNA              | U2 snRNA            |
| U5 snRNA             | U4 snRNA                       | U5 snRNA            | U5 snRNA              | U5 snRNA            |
| U5 snRNA             | U6 snRNA                       | Substrate RNA       | Substrate RNA         | Substrate RNA       |

 RNAs and proteins in U6 snRNA-containing structures. RNAs are shown in italics. Components of individual snRNPs are highlighted in color, including U2 (blue), U4 (green), U5 (orange), and U6 (red). Protein-only complexes NTC (gray) and NTR (purple) are also included. Helicases that promote progression through the splicing cycle are starred. All structures are from S. cerevisiae; only proteins modeled in available structures are included in the table.

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modeled in any of the structures. Small local base-pairing changes do occur between the ACAGA box and the intron, especially U6 nucleotides G50 and A51. However, local changes in accessibility, flexibility, or structure may occur in solution, with important implications for activation and catalysis (Bao et al. 2017).

In contrast, U2 snRNA and the branch site helix, along with many protein partners, undergo >50 Å motions and large rotations. Analogy to group II introns, along with detailed genetic and biochemical studies, suggested that the active site for first and second step chemistry are the same (Chanfreau and Jacquier 1994; Marcia and Pyle 2012; Fica et al. 2013, 2014), and that large conformational changes are required in order to move the first step product (the 2′–5′ branch site) out of the single active site, followed by positioning of the 3′ exon in the active site for second step chemistry (exon ligation). The first hints of these large-scale motions came from single molecule experiments (Abelson et al. 2010; Crawford et al. 2013). Recent cryo-EM structures of splicing complexes before and after the first and second steps of splicing corroborate the hypothesis that U6 forms a rigid active site in the spliceosome and that its substrates must be positioned within it for catalysis to occur (Yan et al. 2015, 2016, 2017; Galej et al. 2016; Rauhut et al. 2016; Wan et al. 2016a, 2017; Bertram et al. 2017b; Fica et al. 2017; Zhang et al. 2017).

Metal ion binding in the spliceosome active site

A two-metal mechanism for RNA-mediated splicing catalysis was proposed in 1993 (Steitz and Steitz 1993). This model stated, by analogy to protein phosphoryl transfer mechanisms, that two divalent metal ions situated 3.9 Å apart stabilize the nucleophile and the leaving group during transesterification. Two groups used phosphorothioate substitutions to identify functionally important non-bridging oxygen atoms in the U6 phosphoribose backbone (Fabrizio and Abelson 1992; Yu et al. 1995), and subsequent thiophilic metal ion rescue experiments revealed that the U6 ISL coordinates the two Mg$^{2+}$ ions required for catalysis via five nonbridging phosphate oxygens (Yean et al. 2000; Fica et al. 2013). Electron density in the active site of recent spliceosome cryo-EM structures has been modeled as metal ions (Wan et al. 2016a; Yan et al. 2016, 2017; Fica et al. 2017; Zhang et al. 2017). However, the current resolution of these cryo-EM structures precludes the unambiguous placement of metal ions into the electron density, such that the exact location of magnesium ions in the active site, and positioning and identity of other structural metal ions, remains to be determined.

**RECYCLING AND DEGRADATION**

**Recycling**

After a round of splicing, the spliceosome undergoes an active disassembly process in which the snRNPs and NTC are recycled for a new round of splicing. The U2/U6.U5-intron lariat complex is disassembled by the ATP-dependent DEAH-box helicase Prp43 with cofactors Ntr1 and Ntr2 (Martin et al. 2002; Tsai et al. 2005). In in vitro assays, U6 is released as the free snRNA (Tsai et al. 2005; Fourmann et al. 2013, 2016). In the cell, U6 is presumably recognized during or after release from the ILS by Prp24 and/or Lsm2-8. Prp24 is required in yeast even when mutations in U4 and U6 allow for bypass of stable U4/U6 base-pairing,
indicating that Prp24 has an essential function other than U4/U6 annealing, perhaps in driving ILS dissociation by capturing U6 (Burke et al. 2015). It is not known if recycling of U6 after a round of splicing follows the same pathway as initial assembly of the U6 snRNP, i.e., U6 is bound by Prp24 and Lsm2-8, then assembled into the U4/U6 di-snRNP.

Degradation

The degradation pathway of U6 is poorly understood. The balance between uridylation, polyadenylation, and trimming by Usb1 (discussed in the “3′ End modification” section) affects the half-life of U6 in metazoa (Shchepachev et al. 2012, 2015; Hilcenko et al. 2013). End processing by Usb1 has a protective effect against degradation in S. cerevisiae, S. pombe, and human cells, suggesting that the modification may protect against other cellular nucleases (Mroczek et al. 2012; Shchepachev et al. 2012, 2015; Hilcenko et al. 2013). However, it is not known which nucleases are responsible for degrading U6 in Usb1-depleted cells or during the normal U6 life cycle. Additionally, other cellular deadenylases can interact with and influence the 3′ sequence of U6 snRNA (Shukla and Parker 2017). How the activities of different 3′ modifying enzymes affect the stability and functionality of U6 is poorly understood.

Recent data suggest that the exosome may play a role in normal turnover of splicedosomal RNAs, as knockdown of the exosome exonuclease Rrp44 resulted in higher levels of U6 (Zhang et al. 2015). The presence of a terminal phosphate (in yeast) or 2′,3′-cyclic phosphate (in humans) on U6 may inhibit degradation by the exosome, as the nuclear exosome-associated exonuclease Rrp6 is inactive on a 3′ phosphate-terminated RNA (Burkard and Butler 2000). Addtional enzymes may be necessary to dephosphorylate the 3′ end of U6 prior to adenylation and decay. It has been demonstrated that Mtr4 and Trf4 (components of the TRAMP complex) as well as Rrp6 interact with components of the U4/U6.U5 tri-snRNP (Nag and Steitz 2012). While the primary reason for recruitment of the exosome machinery may be to degrade improperly spliced mRNAs or excised introns, it may also have a role in degrading “expired” snRNAs. The TRAMP complex is recruited to the intron even before splicing is complete and may in fact help stimulate splicing (Kong et al. 2014). Thus, the components of the nuclear decay machinery are temporally and spatially near the splicedosomal RNAs. Supporting this idea, in S. cerevisiae, deletion of the poly(A) polymerase from the TRAMP complex, Trf4, results in a small increase in U6 levels (Copela et al. 2008). Moreover, Rrp44, Rrp6, and Trf4 can be crosslinked to U6 (Schneider et al. 2012). However, deletion of Trf4 in S. pombe does not affect polyadenylation of U6, suggesting that other poly(A) polymerases are involved or can complement loss of Trf4 (Shchepachev et al. 2015). Further study on the determinants of U6 decay, including the adenylation state of the 3′ tail, will help to define how this metabolically stable RNA is monitored, which may provide another mechanism for spliceosome fidelity.

CONCLUSIONS AND PERSPECTIVES

X-ray crystallography and cryo-EM approaches have led to all-atom models of multiple splicedosomal complexes that contain U6 snRNA. These structures allow us to directly visualize the vast conformational changes that U6 must undergo during spliceosome assembly and activation, and highlight the large number of dynamic steps that must occur to form such complex structures. Each assembly and processing step represents an opportunity for quality control, ensuring that functional splicosomes are prepared to carry out a step of gene expression that is essential in all eukaryotes. U6, U2, and U5 snRNA as well as many proteins must come together to form the catalytic core of the spliceosome, where U6 snRNA coordinates the magnesium ions required for splicing catalysis. Regulation and monitoring of U6 throughout its lifetime is crucial for retaining the spliceosome’s catalytic activity and to ensure that only full-length, properly modified U6 molecules are incorporated into the spliceosome. Research in the coming years will further illuminate how U6 is regulated (via transcriptional activity, post-transcriptional modification, RNA–protein interactions, and subnuclear localization) and assembled into the spliceosome.

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