Post-transcriptional modifications modulate conformational dynamics in human U2–U6 snRNA complex

KRISHANTHI S. KARUNATILAKA¹ and DAVID RUEDA¹,²,³,⁴
¹Department of Chemistry, Wayne State University, Detroit, Michigan 48202, USA
²Department of Medicine, Section of Virology, Imperial College London, London W12 0NN, United Kingdom
³Single Molecule Imaging Group, MRC Clinical Sciences Center, Imperial College London, London W12 0NN, United Kingdom

ABSTRACT

The spliceosome catalyzes precursor-mRNA splicing in all eukaryotes. It consists of over 100 proteins and five small nuclear RNAs (snRNAs), including U2 and U6 snRNAs, which are essential for catalysis. Human and yeast snRNAs share structural similarities despite the fact that human snRNAs contain numerous post-transcriptional modifications. Although functions for these modifications have been proposed, their exact roles are still not well understood. To help elucidate these roles in pre-mRNA splicing, we have used single-molecule fluorescence to characterize the effect of several post-transcriptional modifications in U2 snRNA on the conformation and dynamics of the U2–U6 complex in vitro. Consistent with yeast, the human U2–U6 complex reveals the presence of a magnesium-dependent dynamic equilibrium among three conformations. Interestingly, our data show that modifications in human U2 stem I modulate the dynamic equilibrium of the U2–U6 complex by stabilizing the four-helix structure. However, the small magnitude of this effect suggests that post-transcriptional modifications in human snRNAs may have a primary role in the mediation of specific RNA–protein interactions in vivo.

Keywords: splicing; human U2–U6 snRNAs; single-molecule FRET; structural dynamics; post-transcriptional modifications

INTRODUCTION

Precursor mRNA (pre-mRNA) splicing is an essential process of eukaryotic gene expression, during which noncoding intron sequences are removed from primary transcripts to form mature mRNA. It consists of two transesterification steps that resemble analogous steps in group II intron splicing (Newman 1998; Sontheimer et al. 1999; Gordon et al. 2000). Splicing regulation is critical to maintain proper cellular function, and defects in splicing have been linked to neurodegenerative disorders and various cancers (Kalnina et al. 2005; Licatalosi and Darnell 2006; Pettigrew and Brown 2008).

Splicing is highly conserved in all eukaryotes (from yeast to humans) and is catalyzed by the spliceosome, a dynamic ribonucleoprotein (RNP) complex assembled from five small nuclear RNAs (U1, U2, U4, U5, and U6 snRNAs) and a large number of proteins (Stark and Luhrmann 2006; Ritchie et al. 2009; Wahl et al. 2009). In addition to catalyzing pre-mRNA splicing, the spliceosome plays an important central role in alternative splicing regulation (Saltzman et al. 2011). Following a highly orchestrated assembly, the activated spliceosome maintains only the U2, U5, and U6 snRNPs and releases U1 and U4 (Wahl et al. 2009). However, several lines of evidence have suggested that only U2 and U6 are critical for catalysis (Parker et al. 1987; Lesser and Guthrie 1993; O’Keefe et al. 1996; Segault et al. 1999; Valadkhan and Manley 2001, 2003). Furthermore, a recent crystal structure of a large Prp8 fragment (a U5 snRNP protein) further supports RNA-based catalysis (Galej et al. 2013). Although the U2–U6 complex is essential for pre-mRNA splicing, its structure and role in catalysis are still not well understood (Madhani and Guthrie 1992; Wolff and BINDEREIF 1993; Sun and Manley 1995; Sashital et al. 2004; Mefford and Staley 2009).

The U6 snRNA contains two highly conserved sequences that are involved in catalysis (Fig. 1): the 5′ splice site recognition sequence (ACAGAGA box) and the AGC triad (Lesser and Guthrie 1993; Hilliker and Staley 2004). In addition, it also contains the highly conserved U74 (analogous to U80 in yeast) in the U6 intramolecular stem–loop (U6 ISL), which is involved in metal ion binding and coordination for splicing (Huppler et al. 2002; Valadkhan et al. 2009). Based on in vivo genetic and in vitro NMR studies (Madhani and Guthrie 1992; Sun and Manley 1995; Sashital et al. 2004) two different secondary structures for the yeast U2–U6 (yU2–U6) snRNA

© 2013 Karunatilaka and Rueda This article is distributed exclusively by the RNA Society for the first 12 months after the full-issue publication date (see http://rnajournal.cshlp.org/site/misc/terms.xhtml). After 12 months, it is available under a Creative Commons License (Attribution-NonCommercial 3.0 Unported), as described at http://creativecommons.org/licenses/by-nc/3.0/.

16 RNA 20:16–23; Published by Cold Spring Harbor Laboratory Press for the RNA Society
Conserved AGC triad in U6 and three residues in U2 snRNA that extend the so-called Helix Ib that forms between the highly conserved residues in intramolecular base pairs within U6 snRNA that extend the U6 ISL (Sashital et al. 2004). Interestingly, compensatory mutational genetic studies have shown that Helix Ib residues from both U2 and U6 participate in intramolecular rather than intermolecular base-pairing in humans, suggesting that, in humans, the U2–U6 complex forms the four-helix junction (Wolff and Binder ef 1993; Sun and Manley 1995). Since then, several ensemble-averaged studies have provided evidence that support a hypothesis in which the U2–U6 complex can adopt multiple conformations corresponding to different activation states of the spliceosome (Rhode et al. 2006; Sashital et al. 2007). Recently, we have shown by single-molecule fluorescence that a protein-free U2–U6 snRNA complex from yeast can adopt at least three conformations, consistent with the previously proposed structures (Guo et al. 2009). Consistent with this notion, a recent NMR and enzymatic probing study has shown that a protein-free U2–U6 complex from humans can adopt primarily the four-helix junction structure but also the three-helix junction (Zhao et al. 2013).

Although the primary sequences of U2 and U6 are highly conserved from yeast to humans (Brow and Guthrie 1988), the human snRNAs contain numerous post-transcriptional modifications such as 2′-O-methyl groups and pseudouridines (Massenet et al. 1998; Yu et al. 1998; Sashital et al. 2007). The majority of these conserved modifications are distributed in functionally important regions of the snRNAs, where they can affect pre-mRNA splicing by regulating RNA–RNA and RNA–protein interactions (Newby et al. 2002b; Karijolich and Yu 2010). Out of the five snRNAs in humans, U2 is the most highly modified, containing 10 2′-O-methylated residues and 13 pseudouridines (Fig. 1A; Donmez et al. 2004). Recent NMR and UV melting studies have shown that although the human and yeast U2 stem I structures are nearly isosteric in vitro, post-transcriptional modifications in human U2 significantly increase its stability (Sashital et al. 2007). Since the formation of helix Ib requires the melting of base pairs in stem I, the increased stability of the modified U2 stem I may hinder formation of Helix Ib in human U2–U6 (hU2–U6).

Here, we have used single-molecule Förster resonance energy transfer (smFRET) to characterize the structural dynamics of a protein-free U2–U6 from humans (Fig. 1A,C). smFRET experiments help dissect RNA folding pathways by revealing the presence of transient folding intermediates that can be hidden in ensemble-averaged experiments and by providing real-time dynamic information (Rueda and Walter 2005; Aleman et al. 2008; Karunatilaka and Rueda 2009; Zhao and Rueda 2009). Our data show that, similarly to the yeast complex, the hU2–U6 also adopts at least three conformations in dynamic equilibrium, and that Mg2+ ions modulate their relative stability. Furthermore, post-transcriptional modifications in the U2 stem I stabilize the four-helix structure. However, the relatively small energetic magnitude of this effect raises the interesting possibility that post-transcriptional modifications in U2 play a more important role in protein-specific recognition in vivo.

RESULTS

hU2–U6 snRNA adopts three dynamic conformations

To characterize the structural dynamics of the unmodified hU2–U6 complex, we have used a previously characterized labeling strategy (Fig. 1A,C; Guo et al. 2009). The U6 strand was labeled with a FRET pair (Cy3, Cy5) at the 5′ end and at uracil 64, respectively, and a 3′-biotin for surface immobilization. This labeling scheme enables monitoring of the dynamics of helix III relative to the U6 ISL, a motion that is thought to bring in close proximity the highly conserved residues involved in the first step of splicing (AGC triad, ACAGAGA loop, and U74) (Fig. 1A). Differing from our previous
work, here we have used a single-stranded U6 snRNA, which
simplifies the experiments and their interpretation.

We first monitored the folding dynamics of the surface-
immobilized fluorophore-labeled construct in standard buff-
er (50 mM TRIS-HCl at pH 7.5, 100 mM NaCl, and 10 mM
MgCl2) using smFRET (Fig. 2A). Similar to our previous
work with yeast (Guo et al. 2009), the resulting time trajecto-
ries (Fig. 2A) clearly reveal the presence of three distinct
FRET states (∼0.6, ∼0.4, and ∼0.2), corresponding to at least
three different conformational states in dynamic equilibri-
um. A time-binned FRET histogram from 154 trajectories
(Fig. 2B) confirms the presence of the three FRET states.
Similar to the yeast construct, the histogram in 10 mM
Mg2+ shows that the high FRET conformation is the least
populated (13%), while the intermediate and low FRET con-
formations are approximately equally populated (43% and
44%, respectively).

Based on the similarities between the human and yeast re-
results (Guo et al. 2009), we propose a two-step folding path-
way for the hU2–U6 snRNAs (Figs. 1C, 2D), where the
high and low FRET conformations (N and G) may corre-
spond to the four- and three-helix structures, respectively.
As for the yeast construct, out of >350 observed trajectories,<2% exhibited direct transitions from high (G) to low (N)
FRET states. Therefore, we assign the intermediate confor-
mation (I) as an obligatory folding intermediate.

A dwell time analysis of the time trajectories yields the
folding rate constants (k1, k−1, k2, and k−2) (Fig. 2C), which
are in close agreement with those previously obtained with
the yU2–U6 complex, except k2, which is ∼70-fold faster
for the human construct. This difference may arise from
the larger number of mismatched nucleotides in the human
ISL, which may be less stable than the yeast ISL, and the pres-
ence of fewer base pairs in the human stem I. Overall, in the
absence of post-transcriptional modifications, these data sug-
gest that the hU2–U6 snRNA complex behaves similarly to
the yeast complex, consistent with the idea that splicing is a
highly conserved process amongst all eukaryotes (Will and
Luhrmann 2011).

Characterization of the low FRET conformation
To support that the low FRET state corresponds to the three-
helix structure with helix Ib formed, as observed for the yU2–
U6 complex (Guo et al. 2009), we used a fourfold mutant that
prevents helix Ib formation, while maintaining the junction
structure of the four-helix structure (Fig. 3A). We flipped
four bases in U2 stem I to prevent base-pairing with the cat-
alytic AGC triad of U6. The resulting FRET trajectories and
histogram (Fig. 3B,C) clearly show that the fourfold muta-
tion significantly increases the mid FRET population while
the low FRET population decreases, as expected. The FRET
time trajectories confirm that excursions to the low FRET
become less frequent and transient than for the wild-type
complex (cf. Figs. 2A and 3B). These results are consistent
with the formation of helix Ib in the low FRET conformation
(Fig. 1B), similar to the yeast results. However, we cannot for-

Mg2+ ions modulate the U2–U6 conformational
dynamics
Magnesium ions play an essential role during pre-mRNA
splicing by facilitating RNA folding and catalysis (Yean et al.
2000; Butcher 2011). We have previously shown that Mg2+
ions stabilize the intermediate and low FRET conformation

FIGURE 2. Magnesium stabilizes the low FRET conformation. (A) Typical FRET trajectories of the unmodified U2–U6 complex in 10 mM (top) and
40 mM Mg2+ (bottom). (B) FRET histograms of the unmodified U2–U6 complex obtained in 10 mM (top) and 40 mM Mg2+ (bottom). Both FRET
trajectories and histograms exhibit three FRET states: low FRET (∼0.2, G), mid FRET (∼0.4, I), and high FRET (∼0.6, N). (C) Observed folding rate
constants (k1, k−1, k2, and k−2) of the hU2–U6 complex without post-transcriptionally modified bases in 10 mM (red) and 40 mM Mg2+ (purple). (D)
Effect of Mg2+ on the folding energy landscape of the hU2–U6 complex.
in the yU2–U6 construct. To test the role of Mg²⁺ ions on the unmodified hU2–U6 complex, we repeated the smFRET measurements in high [Mg²⁺] (40 mM). At this concentration, the trajectories (Fig. 2A) show that the molecules remain predominantly in the low FRET state and excursions to the intermediate and high FRET states are less frequent and more transient. A time-binned FRET histogram from 179 trajectories (Fig. 2B) confirms that, in high [Mg²⁺], the low FRET conformation becomes the most populated (69%), indicating that Mg²⁺ ions stabilize this conformation relative to the intermediate (26%) and the high (5%) FRET states.

A dwell time analysis of 99 and 108 trajectories at 10 and 40 mM, respectively, (Fig. 2C; Supplemental Fig. 1) shows that the rate constants $k_1, k_{-1},$ and $k_2$ do not change significantly ($P > 0.02$) with increasing [Mg²⁺]. However, the rate constant $k_2$ increases by approximately twofold when Mg²⁺ concentration increases from 10 mM to 40 mM (Fig. 2C). This result confirms that in the 10–40 mM range Mg²⁺ ions preferentially stabilize the low FRET conformation (Fig. 2D).

Taken together, these data indicate that in the absence of post-transcriptional modifications, a high concentration of divalent ions stabilizes the low FRET state, which we attribute to the three-helix conformation of the hU2–U6 complex similar to that of yeast snRNAs (Guo et al. 2009).

**Post-transcriptional modifications modulate the U2–U6 conformational dynamics**

Human U2–U6 snRNA complex contains 23 post-transcriptionally modified nucleotides in both strands (Fig. 1A), in contrast to the single modified nucleotide (pseudouridine at position 35 in U2) for the equivalent complex in yeast (Yu et al. 1998; Donmez et al. 2004; Sashital et al. 2007; Karijolich and Yu 2010). To understand the effect of these modifications on the structural dynamics of the hU2–U6 complex, we have focused on four modifications in U2 stem I (Fig. 4A) that have been shown to be important for splicing (Donmez et al. 2004; Karijolich and Yu 2010): 2′-O-methylguanosines (Gm) at positions 11, 12, 19, and a
pseudouridine at position 15. Several studies have suggested that modifications in U2 snRNA are involved in spliceosomal assembly (Karjolich and Yu 2010). Specifically, 2′-O-methylations at positions 12 and 19 have been individually shown to be required for early spliceosomal E complex formation (Donmez et al. 2004).

The observed time-trajectories in 40 mM [Mg2+] with the fully modified U2 stem I reveal that the snRNA complex resides for less time in the low FRET state and more time in the intermediate FRET conformation (cf. Figs. 2A and 4B, bottom). A time-binned FRET histogram from 144 trajectories (Fig. 4C), confirms a significant population decrease in the low FRET state (from 69% to 47%) accompanied by a population increase in the intermediate FRET state (from 26% to 44%). Dwell time analysis of the time trajectories shows that while the presence of the modifications does not affect appreciably the rate constants $k_1$ and $k_{-1}$, they decrease $k_2$ by approximately twofold and increase $k_{-2}$ by approximately twofold (Fig. 4D). These effects in rate constants clearly explain the population decrease in the low FRET state and population increase in the intermediate FRET state. However, consistent with comparable rate constants of $k_1$ and $k_{-1}$, we did not observe a substantial change in the high FRET population with modifications (8% with modifications and 5% without modifications).

Collectively, these results indicate that the presence of post-transcriptional modifications destabilize the low FRET state that has been proposed as the three-helix conformation (helix Ib) and stabilize the intermediate conformation (stem I). Based on the FRET histograms, we estimate that these modifications stabilize the U2 stem I by $\sim{0.5}$ kcal/mol. These results are in qualitative agreement with previous UV melting studies that have shown a modification-mediated stabilization of the U2 stem I helix (Sashital et al. 2007). Lowering the [Mg2+] to 10 mM results in similar but less prominent effects (Fig. 4B,C).

To understand the effect of pseudouridine 15 and 11Gm on the structural dynamics of the U2–U6 complex, we performed single-molecule experiments with a U2–U6 snRNA construct containing only Gm at positions 12 and 19 ($12Gm + 19Gm$) (Fig. 5A; Supplemental Fig. 2). We used 40 mM Mg2+ because the stabilizing effect of the modifications is most pronounced at this concentration (Fig. 4D). A time-binned histogram from 102 FRET trajectories reveals almost identical distribution to the fully modified U2 construct (cf. Figs. 4C and 5A), indicating that the absence of pseudouridine 15 and 11Gm does not significantly affect the relative stability of the U2–U6 conformations. An estimate of the relative free-energy difference between states (Table 1, $\Delta\Delta G$) confirms that most of the stabilization of the U2 stem I arises from $12Gm$ and $19Gm$ alone.

To assess the contribution from each of the remaining modifications ($12Gm + 19Gm$), we repeated the measurement with only one of them at a time (Fig. 5B,C). A time-binned histogram from 118 single-molecule trajectories of only $19Gm$ (Fig. 5B) reveals a population distribution that resembles the one obtained with the unmodified construct (cf. Figs. 2B and 5B), indicating that $19Gm$ plays a minor role in stabilizing the U2 stem I. An estimate of the relative free-energy difference between states (Table 1) shows that 0.19 of the 0.53 kcal/mol stabilization of the modified U2 stem I arises from $19Gm$ alone. In the presence of only $12Gm$ (Fig. 5C), a time-binned histogram from 151 single-molecule trajectories reveals a population distribution that is intermediate between the fully modified construct and the unmodified construct.

| TABLE 1. Post-transcriptional modifications destabilize the low FRET conformation ($\psi$ = pseudouridine and $Gm$ = 2′-O-methylguanosine) |
|-----------------|
| U2–U6 snRNA complex | $\Delta\Delta G$ (kcal/mol) |
| Unmodified | 0 |
| $11Gm + 12Gm + 19Gm + 15\psi$ | $+0.53 \pm 0.17$ |
| $12Gm + 19Gm$ | $+0.51 \pm 0.18$ |
| $12Gm$ | $+0.30 \pm 0.16$ |
| $19Gm$ | $+0.19 \pm 0.13$ |

Relative free-energy difference ($\Delta\Delta G$) between the unmodified and modified hU2–U6 complexes. $\Delta\Delta G > 0$ indicates that the modification destabilizes the low FRET state. Error associated with $\Delta\Delta G$ was calculated using the bootstrapping method.
Additional information about the U2–U6 snRNA complex and its role in the spliceosome is provided in this section. The U2–U6 complex is a key component of the spliceosome, involved in the formation of the splicing reaction. The complex is highly conserved across species, and its structure and dynamics have been extensively studied. The U2–U6 complex interacts with other components of the spliceosome, including U5, U6, and U7 snRNAs, to facilitate the splicing reaction.

The U2–U6 complex is composed of two RNA molecules, U2 and U6, which are joined by a series of base pairs to form a stable structure. The U2 and U6 RNAs are known to undergo conformational changes, which are crucial for the proper function of the spliceosome. These conformational changes are mediated by various factors, including Mg2+ concentration, post-transcriptional modifications, and protein interactions.

In high magnesium concentrations, the U2–U6 complex prefers the low FRET state, which has two fluorophores far away from each other. As we previously observed in yeast, the hU2–U6 complex may also adopt the three-helix structure with helix Ib by rearranging the junction structure. The existence of the three-helix structure with helix Ib was further supported using a fourfold mutant containing flipped bases in the U2 stem I that prevent the formation of helix Ib. Although it can maintain the junction structure, the mutant U2–U6 construct greatly decreases the population of the low FRET state. Thus, smFRET data of the fourfold mutant provide evidence for the observed low FRET state as the previously proposed three-helix structure.

In addition to the conformations discussed above, smFRET data of U2–U6 snRNAs exhibited the existence of a previously unidentified folding intermediate with FRET ~0.4. Since previous studies have shown that the stability of this conformation is independent of the mutations at U80 in yeast (comparable to U74 in humans) U6 snRNA (Guo et al. 2009), we propose this mid FRET state has the extended U6 ISL structure, but lacks the tertiary interactions between the ACAGAGA sequence and the U6 ISL. However, the role of this conformation in pre-mRNA splicing is not well understood in both humans and yeast. Although we have assigned the observed FRET states as the three-helix and the four-helix conformations, it is important to note that the hU2–U6 complex may fold into alternative three-dimensional structures based on different coaxial and side-by-side stacking of helices in solution.

Consistent with yeast smFRET data, the population of the low FRET state (FRET ~0.2) increases with Mg2+ concentration, indicating a magnesium-mediated stabilization of the three-helix structure (Guo et al. 2009). As a result, under high Mg2+ concentrations, the hU2–U6 complex without modified bases showed a higher population of molecules in the three-helix structure than the four-helix structure. In contrast, the hU2–U6 snRNAs with post-transcriptionally modified bases in U2 stem I (11Gm, 12Gm, 19Gm, and pseudouridine at position 15) show a clear stabilization effect of the mid FRET state while lowering the population of the low FRET state corresponds to the three-helix structure. Interestingly, smFRET data obtained from the U2–U6 construct with 2′-O-methylguanosines at positions 12 and 19 (12Gm+ 19Gm) exhibited folding dynamics similar to the fully modified construct. Therefore, these results suggest that 11Gm and pseudouridine at position 15 of U2 stem I do not exhibit an overall effect on the folding dynamics of the U2–U6 complex. This unique behavior can be due to either cancellation of opposite effects or due to lack of direct stabilization/destabilization effect from the two individual modifications. Furthermore, compared with the modification at position 19, the 2′-O-methylguanosines at position 12 showed a higher destabilization effect on the low FRET state of the hU2–U6 complex. This result is in agreement with previous measurements of Holliday Junction dynamics which show that the first base pair near the junction plays an important role in its stability and dynamics (McKinney et al. 2005).

Taken together, the smFRET data suggest that modified bases collectively affect the structural dynamics of the U2–U6 complex and may prevent the formation of the proposed
three-helix structure consisting of helix I by stabilizing U2 stem I. 2′-O-methylations can alter the structural stabilization of U2 stem I by blocking sugar edge interactions and by reducing the reactivity of the RNA backbone (Auffinger and Westhof 1997; Helm 2006). In the case of pseudouridine, this may result from improved base stacking (Davis 1995) and from formation of a water-mediated hydrogen bond between the additional protonated imino group and adjacent phosphate residues, as reported in functionally important RNAs including tRNAs (Arnez and Steitz 1994; Newby et al. 2002a).

Although the modified bases destabilize the low FRET state, the effect of modifications on the stability of U2 stem I is less significant in the U2–U6 complex than in the U2 stem I alone, as shown in UV melting studies (Sashital et al. 2004). Therefore, we propose that post-transcriptionally modified residues may be involved in protein-dependent functions rather than direct stabilization of U2–U6 RNA structures in vivo. Hence, we suggest that modified bases in U2 snRNA may play a major role in protein recognition and interactions in cells. Thereby these modifications can directly participate in the spliceosome assembly process as suggested previously (Donmez et al. 2004). This hypothesis is further supported by the presence of a large number of spliceosomal proteins associating with human snRNAs, which are highly post-transcriptionally modified compared with the yeast system.

In summary, the hU2–U6 complex follows the minimal two-step folding pathway consisting of the previously proposed two structures and at least one obligatory intermediate conformation. Furthermore, RNA constructs without post-transcriptionally modified bases exhibited similar dynamics as yeast spliceosomal RNAs (Guo et al. 2009). Although modified bases can affect the structural dynamics of hU2–U6 snRNAs, the structural effects of the U2–U6 complex were less than that of U2 stem I alone, suggesting a protein-dependent role rather than direct stabilization of RNA structures in cells.

**MATERIALS AND METHODS**

**RNA purification and labeling**

5′-Cy3 labeled U6 RNA (5′-AUACAGAGAGAUUGCAU GGCCTCCTTGCGCAAGGACAGACAGCAAAUUAGUUGAAGC-3′) Biotin, where dT contains an internal, free primary amine for Cy5 labeling) and U2 RNA (5′-GCUUCUGGCGCUCUUGCUGAAG AUCAAGUGAUAGAU-3′) with or without modified bases (pseudouridine and 2′-O-methyl guanosines) were purchased from the Howard Hughes Medical Institution Biopolymer/Keck Foundation Biotechnology Resource Laboratory (Yale University). Synthetic RNAs were 2′-OH deprotected, labeled, and purified by denaturing 18% polyacrylamide gel electrophoresis and C8 reversed-phase HPLC as described (Steiner et al. 2008; Zhao and Rueda 2009). Internal dT1 of U6 was Cy5 labeled following the manufacturer’s protocol and purified by reversed-phase HPLC.

**Single-molecule FRET (smFRET)**

Single-molecule FRET experiments were performed on a home-built, prism-based, total internal reflection microscope as described (Guo et al. 2009; Zhao and Rueda 2009). Briefly: Biotinylated and fluorophore-labeled U6 (1 μM) and U2 (10 μM) were heated (90° C for 45 sec) in standard buffer (50 mM TRIS-HCl at pH 7.5, 100 mM NaCl, and variable [MgCl2]) and annealed (15–20 min) at room temperature. The resulting U2–U6 RNA complex was diluted to ~25 pM and surface immobilized onto a streptavidin-coated quartz slide via a biotin-streptavidin linkage. Unbound RNA was washed away with standard buffer. To reduce fluorophore photobleaching, an oxygen-scavenging system (OSS) consisting of 10% (w/v) glucose, 2% (v/v) 2-mercaptoethanol, 50 µg/mL glucose oxidase, and 10 µg/mL catalase was used in all experiments. All TIRF-based single-molecule experiments were performed using a 532-nm laser with 33-msec time resolution at room temperature.

**Data analysis**

FRET and dwell times histograms were built by time binning >100 dynamic single-molecule time trajectories. Kinetic rate constants were obtained by fitting the resulting dwell times histograms to exponential decays as described (Guo et al. 2009). Differences between free-energy changes associated with the formation of the low FRET state for unmodified (U) and modified (M) U2–U6 snRNA constructs were calculated as ∆∆G = ∆Gm − ∆G13, where ∆G = −RT ln(K), K = Ar/Am and A represents the area of a given FRET state in a FRET histogram. Error associated with ∆∆G was analyzed using the bootstrap method as described (Efron and Gong 1983).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

**ACKNOWLEDGMENTS**

We thank N. Greenbaum and S. Butcher for critically reading the manuscript. The Rueda lab is funded by grants from the NIH (GM085116) and a CAREER award from the NSF (MCB-0747285). 

Author contributions: K.S.K. and D.R. designed research; K.S.K. performed research and analyzed data; K.S.K. and D.R. wrote the paper.

Received August 8, 2013; accepted October 2, 2013.

**REFERENCES**

Alien EA, Lamichhane R, Rueda D. 2008. Exploring RNA folding one molecule at a time. Curr Opin Chem Biol 12: 647–654.

Arnez JG, Steitz TA. 1994. Crystal structure of unmodified tRNA<sub>Glu</sub> complexed with glutaminyl-tRNA synthetase and ATP suggests a possible role for pseudo-uridines in stabilization of RNA structure. Biochemistry 33: 7560–7567.

Auffinger P, Westhof E. 1997. Rules governing the orientation of the 2′-hydroxyl group in RNA. J Mol Biol 274: 54–63.

Brow DA, Guthrie C. 1988. Spliceosomal RNA U6 is remarkably conserved from yeast to mammals. Nature 334: 213–218.

Burke JE, Sashital DG, Zuo X, Wang YX, Butcher SE. 2012. Structure of the yeast U2/U6 snRNA complex. RNA 18: 673–683.
Butcher SE. 2011. The spliceosome and its metal ions. *Met Ions Life Sci* **9**: 233–251.

Davis DR. 1995. Stabilization of RNA stacking by pseudouridine. *Nucleic Acids Res* **23**: 5020–5026.

Donmez G, Hartmuth K, Luhrmann R. 2004. Modified nucleotides at the 5′ end of human U2 snRNA are required for splicosomal E-complex formation. *RNA* **10**: 1925–1933.

Efron B, Gong G. 1983. A leisurely look at the bootstrap, the jackknife, and cross-validation. *Am Stat* **37**: 36–48.

Galej WP, Oubridge C, Newman AJ, Nagai K. 2013. Crystal structure of Prp8 reveals active site cavity of the spliceosome. *Nature* **493**: 638–643.

Gordon PM, Sontheimer EJ, Piccirilli JA. 2000. Metal ion catalysis during the exon-ligation step of nuclear pre-mRNA splicing: Extending the parallels between the spliceosome and group II introns. *RNA* **6**: 199–205.

Guo Z, Karunatilaka KS, Rueda D. 2009. Single-molecule analysis of protein-free U2–U6 snRNAs. *Nat Struct Mol Biol* **16**: 1154–1159.

Helm M. 2006. Post-transcriptional nucleotide modification and alternative folding of RNA. *Nucleic Acids Res* **34**: 721–733.

Hilliker AK, Staley JP. 2004. Multiple functions for the invariant AGC triad of U6 snRNA. *RNA* **10**: 921–928.

Huppert A, Nikstad LJ, Allmann AM, Brow DA, Butcher SE. 2002. Metal binding and base ionization in the U6 RNA intramolecular stem-loop structure. *Nat Struct Biol* **9**: 431–435.

Kalinina Z, Zayakin P, Silina K, Line A. 2005. Alterations of pre-mRNA splicing in cancer. *Genes Chromosomes Cancer* **42**: 342–357.

Karijolich J, Yu YT. 2010. Splicosomal snRNA modifications and their function. *RNA Biol* **7**: 192–204.

Karunatilaka KS, Rueda D. 2009. Single-molecule fluorescence studies of RNA: A decade’s progress. *Chem Phys Lett* **476**: 1–10.

Lesser CF, Guthrie C. 1993. Mutations in U6 snRNA that alter splice site specificity: Implications for the active site. *Science* **262**: 1982–1988.

Licatalosi DD, Darnell RB. 2006. Splicing regulation in neurologic disease. *Neuron* **52**: 93–101.

Madhani HD, Guthrie C. 1992. A novel base-pairing interaction between U2 and U6 snRNAs suggests a mechanism for the catalytic activation of the spliceosome. *Cell* **71**: 803–817.

Massenet S, Mougin A, Branlant C. 1998. Posttranscriptional modifications in the U small nuclear RNAs. In *The modification and editing of RNA*. (ed. Grosjean H, Benne R), pp. 201–228. ASM Press, Washington, DC.

McKinney SA, Freeman AD, Lilley DM, Ha T. 2005. Observing spontaneous branch migration of Holliday junctions one step at a time. *Proc Natl Acad Sci* **102**: 5715–5720.

Mefford MA, Staley JP. 2009. Evidence that U2/U6 helix I promotes both catalytic steps of pre-mRNA splicing and rearranges in between these steps. *RNA* **15**: 1386–1397.

Newby MI, Greenbaum NL. 2002a. Investigation of Overhauser effects between pseudouridine and water protons in RNA helices. *Proc Natl Acad Sci* **99**: 12697–12702.

Newby MI, Greenbaum NL. 2002b. Sculpting of the splicososomal branch site recognition motif by a conserved pseudouridine. *Nat Struct Biol* **9**: 958–965.

Newman A. 1998. RNA splicing. *Curr Biol* **8**: R903–R905.

O’Keefe BT, Norman C, Newman AJ. 1996. The invariant U5 snRNA loop 1 sequence is dispensable for the first catalytic step of pre-mRNA splicing in yeast. *Cell* **86**: 679–689.

Parker R, Siliciano PG, Guthrie C. 1987. Recognition of the TACTAAC box during mRNA splicing in yeast involves base pairing to the U2-like snRNA. *Cell* **49**: 229–239.

Petitgrew CA, Brown MA. 2008. Pre-mRNA splicing aberrations and cancer. *Front Biosci* **13**: 1090–1105.

Rhode BM, Hartmuth K, Westhof E, Luhrmann R. 2006. Proximity of conserved U6 and U2 snRNA elements to the 5′ splice site region in activated splicosomes. *EMBO J* **25**: 2475–2486.

Ritchie DB, Schellenberg MJ, MacMillan AM. 2009. Splicosomal structure: Piece by piece. *Buchim Biophys Acta* **1789**: 624–633.

Rueda D, Walter NG. 2005. Single molecule fluorescence control for nanotechnology. *J Nanosci Nanotechnol* **5**: 1990–2000.

Saltzman AL, Pan Q, Blencowe BL. 2011. Regulation of alternative splicing by the core splicosomal machinery. *Genes Dev* **25**: 373–384.

Sashital DG, Cornilescu G, McManus CJ, Brow DA, Butcher SE. 2004. U2–U6 RNA folding reveals a group II intron-like domain and a four-helix junction. *Nat Struct Mol Biol* **11**: 1237–1242.

Sashital DG, Venditti V, Angers CG, Cornilescu G, Guthrie SE. 2007. Structure and thermodynamics of a conserved U2 snRNA domain from yeast and human. *RNA* **13**: 328–338.

Seghalt V, Will CI, Polycarpou-Schwarz M, Mattaj IW, Branlant C, Luhrmann R. 1999. Conserved loop 1 of U5 small nuclear RNA is dispensable for both catalytic steps of pre-mRNA splicing in HeLa nuclear extracts. *Mol Cell Biol* **19**: 2782–2790.

Sontheimer EJ, Gordon PM, Piccirilli JA. 1999. Metal ion catalysis during group II intron self-splicing: Parallels with the spliceosome. *Genes Dev* **13**: 1729–1741.

Stark H, Luhrmann R. 2006. Cryo-electron microscopy of splicosomal components. *Annu Rev Biophys Biomol Struct* **35**: 435–457.

Steiner M, Karunatilaka KS, Sigel RK, Rueda D. 2008. Single-molecule studies of group II intron ribozymes. *Proc Natl Acad Sci* **105**: 13853–13858.

Sun JS, Manley JL. 1995. A novel U2–U6 snRNA structure is necessary for mammalian mRNA splicing. *Genes Dev* **9**: 843–854.

Valadkhani S, Manley JL. 2001. Splicing-related catalysis by protein-free snRNAs. *Nature* **413**: 701–704.

Valadkhani S, Manley JL. 2003. Characterization of the catalytic activity of U2 and U6 snRNAs. *RNA* **9**: 892–904.

Valadkhani S, Mohammadi A, Jaladat Y, Geisler S. 2009. Protein-free small nuclear RNAs catalyze a two-step splicing reaction. *Proc Natl Acad Sci* **106**: 11901–11906.

Wahl MC, Will CI, Luhrmann R. 2009. The spliceosome: Design principles of a dynamic RNP machine. *Cell* **136**: 701–718.

Will CI, Luhrmann R. 2011. Splicosomal structure and function. *Cold Spring Harb Perspect Biol* **3**: a037070.

Wolff T, Bindereif A. 1993. Conformational changes of U6 RNA during the splicosomal cycle: An intramolecular helix is essential both for initiating the U4–U6 interaction and for the first step of slicing. *Genes Dev* **7**: 1377–1389.

Yean SL, Wuenschell G, Termini J, Lin RJ. 2000. Metal-ion coordination required for snRNP assembly and pre-mRNA splicing. *EMBO J* **17**: 5783–5795.

Zhao C, Bachu R, Popovic M, Devany M, Brenowitz M, Schletterer JC, Greenbaum NL. 2013. Conformational heterogeneity of the protein-free human splicososomal U2–U6 snRNA complex. *RNA* **19**: 561–573.