Biased Brownian ratcheting leads to pre-mRNA remodeling and capture prior to first-step splicing

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The spliceosome is a dynamic ribonucleoprotein (RNP) machine that catalyzes the removal of introns during the two transesterification steps of eukaryotic pre-mRNA splicing. Here we used single-molecule fluorescence resonance energy transfer to monitor the distance of the 5′ splice site (5′ SS) and branch point (BP) of pre-mRNA in affinity-purified spliceosomes stalled by a mutation in the DExD/H-box helicase Prp2 immediately before the first splicing step. Addition of recombinant Prp2 together with NTP and protein cofactor Spp2 rearranges the spliceosome-substrate complex to reversibly explore conformations with proximal 5′ SS and BP that accommodate chemistry. Addition of Cwc25, a small heat-stable splicing factor, then strongly biases this equilibrium toward the proximal conformation, promoting efficient first-step splicing. The spliceosome thus functions as a biased Brownian ratchet machine where a helicase unlocks thermal fluctuations subsequently rectified by a cofactor ‘pawl’, a principle possibly widespread among the many helicase-driven RNPs.

Introns are removed by the spliceosome, a large RNP complex, in a two-step transesterification process. In the first step, the 2′ OH of the BP adenosine attacks the phosphodiester bond at the 5′ SS, releasing the 5′ exon and creating the branched, lariat structure. In the second step, the 3′ hydroxyl of this exon attacks the phosphodiester bond at the 3′ splice site (3′ SS), releasing the lariat intron and creating the spliced mRNA product. The most conspicuous feature of this enzyme is that it lacks a preformed catalytic core, which is created in a stepwise fashion, beginning with the assembly of the U1 and U2 small nuclear RNPs (snRNPs), at the 5′ SS and BP, respectively, to form the pre-spliceosome (A complex). The U4–U6•U5 tri-snRNP then binds to create the mature spliceosome (B complex). Notably, however, U1 and U4 snRNPs must be removed before catalysis, creating the activated B (B act) complex and then, after additional rearrangements, the catalytically active B (B*) complex. The resulting post–first-step (C) complex then undergoes further remodeling required for the second step of splicing and the formation of mature mRNA.

The highly dynamic process of spliceosome assembly and catalysis is guided by a set of RNA-dependent ATPases in the DExD/H-box helicase family that collectively function to insure the fidelity of splicing. A major experimental challenge has been to understand the precise conformational rearrangements of RNA and protein that accompany each ATP-dependent step. The DExD/H-box helicase Prp2 is required for the first chemical step of splicing, and recent proteomic analyses of the B act, B* and C complexes revealed that its action results in the destabilization of the U2 snRNP–associated proteins SF3a and SF3b. An attractive hypothesis is that SF3b sequesters the BP adenosine to prevent a premature attack on the 5′ SS, in which case the ATP-dependent action of Prp2, together with its cofactor Spp2, would be required to initiate catalysis. In a biochemical tour de force, successful reconstitution of both steps of splicing with the addition of recombinantly expressed proteins to immunopurified splicing complexes has been demonstrated. In particular, first-step chemistry has been achieved with the addition of ATP, Prp2, Spp2 and Cwc25 (ref. 6).

We set out to investigate the roles of Prp2, Spp2 and Cwc25 in activating the spliceosome for the first step of splicing, developing an approach that couples the purification of specific splicing complexes with single-molecule fluorescence resonance energy transfer (FRET). We used the resulting single-molecule pulldown FRET (SiMPull-FRET) technique to analyze a functional B act complex assembled on pre-mRNA with fluorophores near the scissile bonds. When this complex is assembled in an extract with a temperature-sensitive allele of Prp2 (prp2-1) (ref. 12), the B act spliceosome is stalled and can catalyze the first step of splicing only through formation of the B* complex, upon addition of ATP, Prp2 and Spp2, and then the C complex, upon further addition of Cwc25. Using SiMPull-FRET, we show that ATP-dependent action of Prp2 and its cofactor Spp2 unlocks reversible switching of the intron between splicing-active and splicing-inactive conformations. Cwc25 then rectifies this thermal Brownian ratcheting by stabilizing the conformation in which the 5′ SS and BP are in close proximity, driving the equilibrium toward catalysis.

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RESULTS
Purifying Bact in complex with FRET-labeled Ubc4 pre-mRNA
It has been known for almost 25 years that a Prp2-1 yeast splicing extract can be heat inactivated. In this extract, the spliceosome is fully assembled but cannot carry out the first step of splicing. The immature Prp2-1 spliceosome purified by gradient centrifugation (B\(^{\text{act}}\)) was shown to proceed through the first step of splicing only upon the addition of Prp2 protein and heat-stable factor(s). More recently, this experiment was repeated with purified Prp2, Spp2 and the (since identified) heat stable factor Cwc25 (ref. 6). Such a purified system is ideal for exploring substrate dynamics using single-molecule FRET. To this end, we constructed a yeast strain containing the prp2-1 mutation and a tandem affinity purification (TAP) tag derivative of one of the NTC components, Cef1, known to be present in the spliceosome at this stage. This approach allowed us to purify the stalled B\(^{\text{act}}\) complex via the Cef1-TAP tag using biotin-IgG bound to streptavidin.

The Ubc4 pre-mRNA used in this study was synthesized chemically and labeled with the FRET donor Cy3, which is 6 nucleotides downstream from the BP adenosine, and with the FRET acceptor Cy5, which is 6 nucleotides down from the Cef1-TAP tag using biotin-IgG bound to streptavidin.

Figure 1 The SiMPull-FRET approach used to interrogate active splicing complexes. (a) Schematic showing the affinity-purified B\(^{\text{act}}\) complex immobilized to a streptavidin-coated quartz slide through biotinylated IgG. The green and red circles represent Cy3 and Cy5, respectively. (b) A 15% urea-polyacrylamide gel scanned by a variable-mode Typhoon imager shows pre-mRNA and first-step products, rendered using an overlay of the Cy3 and Cy5 scans (top, intron-lariat; middle, pre-mRNA; bottom, the 5′ exon). Bottom plot, quantification of normalized first-step product from each corresponding lane. Error bars indicate s.d. from triplicate experimental sets. Sample sizes for the different conditions are comparable and effects measured are a reflection of the changes in the distribution of the population.

Bact complex–bound pre-mRNA shows a static low-FRET state
To obtain mechanistic insight into pre-mRNA splice-site juxtaaction during the Prp2-driven restructuring of the B\(^{\text{act}}\) complex into the catalytically activated B* complex, we carried out SiMPull-FRET on the slide-bound B\(^{\text{act}}\) complex (Fig. 1a) in standard splicing buffer. After verifying that each selected pre-mRNA molecule contained one Cy3 and one Cy5 fluorophore, we collected FRET values over the first 100 video frames (at 100-ms time resolution) from 297 molecules. Histograms of the FRET values indicated a single Gaussian distribution with an average FRET value of 0.3 ± 0.15 (s.d.) (Fig. 2a). Given the background noise inherent to any single-molecule experiment, we used hidden Markov modeling (HMM) to find the underlying FRET states. Using a K-means approach, we then clustered the HMM-assigned states using all the experimental conditions from this study into four macro states with FRET values of 0.0–0.23 (state L1), 0.23–0.42 (L2), 0.42–0.60 (M) and 0.6–1.0 (H) (Supplementary Fig. 3 and Supplementary Table 2). The dominant behavior in the B\(^{\text{act}}\) complex is a static low-FRET state (L2) (Fig. 2b). Transition occupancy density plots (TODPs), which are scaled to emphasize the transitions found to be most common among a molecule population, indicate that the static L2 state represents the only behavior in ~52% of all B\(^{\text{act}}\) molecules (Fig. 2c and Supplementary Table 3). In addition, molecules in this state have few transitions (Fig. 2b, HMM fit, cyan line). To test for dynamics that would be too fast for detection by HMM, we performed cross-correlation analysis between the donor and acceptor trajectories of each molecule, and in the resulting scatter around 0, found no evidence for rapid transitions (Fig. 2b and Supplementary Fig. 3). Although splice-site recognition begins in the splicing cycle as early as the commitment complex, our results suggest that the 5′ SS and BP in the B\(^{\text{act}}\) complex are kept stably apart, probably not close enough for splicing chemistry to occur.

Prp2 mediates NTP-dependent remodeling of the pre-mRNA
The ATPase action of Prp2 has been shown to catalyze a large conformational change that activates the spliceosome for the first step of splicing.9 Spliceosomal binding of Prp2 is dependent on its interaction with the G patch domain of its cofactor protein Spp2 (refs. 16,17). The addition of Prp2, Spp2 and ATP transforms the precatalytic at 23 °C for 30–40 min, within two-fold of that of other labeled and unlabeled Ubc4 constructs. Taken together, these experiments establish that FRET labeling of the pre-mRNA substrate is compatible with the expected assembly and splicing activity of the immunopurified B\(^{\text{act}}\) complex, paving the way for SiMPull-FRET interrogation.
Figure 2 In the Prp2-stalled B\textsuperscript{act} complex, the pre-mRNA is restricted predominantly to a static low-FRET state. (a) FRET probability distribution of the raw single-molecule FRET trajectories from the purified B\textsuperscript{act} complex. (b) Representative time traces of the B\textsuperscript{act} complex with raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories and their idealized HMM (cyan). The corresponding cross-correlation of donor and acceptor intensities is shown (right). (c) TODP showing the fraction of B\textsuperscript{act} complex molecules that either do not transition and thus lie on the diagonal (dotted white line) or transition from one indicated FRET state to another. Sample sizes for the different conditions were comparable and effects measured are a reflection of the changes in the distribution of the population. a.u., arbitrary units.

B\textsuperscript{act} complex into the catalytically active, distinctly sedimenting B\textsuperscript{*} complex and results in low levels of first-step splicing. To investigate the role of Prp2 in pre-mRNA remodeling during this step, we incubated the B\textsuperscript{act} complex assembled on the slide surface with Prp2, Spp2 and 2 mM ATP (henceforth referred to as B\textsuperscript{*} complex conditions). B\textsuperscript{*} complex conditions resulted in a substantial shift in the FRET histogram toward a new ~45% population with a mean FRET value of 0.71 ± 0.01 (s.d.), diminishing the lone 0.33 ± 0.01 FRET distribution observed for the B\textsuperscript{act} complex (Fig. 3a). In contrast to the predominantly static L2 state of the B\textsuperscript{act} complex, molecules under B\textsuperscript{*} conditions show dynamic (reversible) excursions to high-FRET states, indicating that the 5′ SS and BP can now reach the close proximity required for first-step chemistry. More specifically, the B\textsuperscript{*} condition comprises the L2, M and H FRET states, where the H state is accessed from either the L2 or M states (Fig. 3b). TODPs show that only ~11% of molecules retained the static L2 state characteristic of the B\textsuperscript{act} complex, whereas ~39% of molecules exhibited at least one L2-to-H transition (Fig. 3c and Supplementary Table 3). Notably, transitions into the H state were short lived in the majority of molecules (Fig. 3b). However, 12% of molecules showed a static high-FRET state (Fig. 3c), indicating that they made a transition through the low levels of first-step splicing observed under B\textsuperscript{*} conditions, after which the labeled 5′ SS and BP become covalently linked (Fig. 1b). To verify that this static high-FRET state corresponds to the

Figure 3 Upon the addition of ATP, Prp2 and Spp2, the pre-mRNA is able to explore splice-site proximity. (a) FRET probability distribution of the raw single-molecule FRET trajectories upon the addition of Prp2, Spp2 and ATP to the B\textsuperscript{act} complex, leading to formation of the B\textsuperscript{*} complex. (b) Representative single-molecule FRET trajectories from the B\textsuperscript{*} condition with the raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories and idealized HMM models (cyan). (c) TODP generated from the idealized HMM for molecules in the B\textsuperscript{*} condition. L1, L2, M or H refers to the four states resulting from clustering analysis. (d) FRET probability densities generated from molecules in B\textsuperscript{act} incubated with various combinations of components required for formation of B\textsuperscript{*} (Prp2, Spp2 and ATP). In addition, B\textsuperscript{act} was incubated with one of two NTP analogs, non-hydrolysable AMP-PNP or UTP. Sample sizes for the different conditions were comparable and effects measured are a reflection of the changes in the distribution of the population. a.u., arbitrary units.
pre-mRNA substrate configuration after the first step of splicing, we trapped this configuration using a dominant-negative Prp16 mutant (Prp16DN; K379A)\(^5\) added to the (non–heat-inactivated) Cef1-TAP–tagged Prp2-1 yeast extract in the presence of 2 mM ATP. This protocol is expected to enrich for the post–first-step C complex, which was then immobilized on the slide surface, washed and imaged. The resulting histogram showed a dramatic enrichment to ~76% of a high-FRET population with a mean FRET value of 0.7 ± 0.01 (s.d.) (Supplementary Fig. 4 and Supplementary Table 3). TODP analysis revealed that ~70% of molecules adopt the same static high-FRET state first observed under B* conditions, strongly supporting the notion that these molecules indeed have undergone the first chemical step of splicing.

Prp2 can directly bind a region in the pre-mRNA downstream of the BP adenosine, even in the absence of ATP\(^20,21\). To investigate whether Prp2 alone can induce the observed pre-mRNA remodeling, we omitted Prp2 or Spp2 from our B* conditions and found the resulting FRET histograms to be indistinguishable from those of the starting B\(^{act}\) complex (Fig. 3d), average FRET value of 0.31 ± 0.16 and 0.33 ± 0.13 (s.d.), respectively. Next, we studied the role of ATP in the remodeling of the B\(^{act}\) complex. For most splicingosomal DExD/H-box helicases, both ATP-dependent and ATP-independent roles have been proposed\(^22–26\). Prp2 in particular has been shown to cause extensive conformational remodeling of the spliceosome in the absence of ATP, whereas the displacement of SF3b is ATP dependent\(^5\). To determine whether the pre-mRNA remodeling observed here requires ATP, we incubated the B\(^{act}\) complex with Prp2 and Spp2 in the absence of ATP and observed no appreciable change in the FRET histogram (Fig. 3d, average FRET value of 0.33 ± 0.12 (s.d.)). When the non-hydrolyzable ATP analog AMPPNP was used instead of ATP, the FRET histogram was again similar (average FRET value of 0.37 ± 0.11) to that of the B\(^{act}\) complex, with no notable excursion to higher FRET states, showing that ATP hydrolysis is required for these excursions to occur (Fig. 3d). (We note that the minor upward shift observed in the histogram may be due to binding of AMPPNP to Prp2, resulting in a slight conformational change.) Finally, the DExD/H-box helicases involved in splicingosomal reorganization are either integral components of snRNPs or extrinsic components, as is Prp2. MS studies have shown that the stalled B\(^{act}\) complex contains stoichiometric amounts of the DExD/H-box helicase Br2, the integral component of the U5 snRNP responsible for U4–U6 unwinding\(^27\). Direct interactions between Prp2 and the C terminus of Br2 have recently been discovered\(^21\), suggesting a possible role for Brr2 in first-step catalytic activation. To test this possibility, we exploited the fact that Brr2 is a strict ATPase\(^28\), whereas Prp2 is a broad NTPase, and supplemented the B\(^{act}\) complex under B* conditions with UTP instead of ATP. The resulting FRET efficiency histogram is clearly distinct from that of B\(^{act}\) and overlays well with that of the ATP-mediated B* condition, with a slightly less-efficient shift toward the higher FRET population (Fig. 3d). This lower efficiency is consistent with the ~2-fold reduction in activity of Prp2 in the presence of NTPs other than ATP\(^29\).

Collectively, these results indicate that the NTP-driven helicase activity of Prp2 in complex with its activator Spp2 causes a large structural reorganization of the pre-mRNA that allows the distal 5’ SS and BP of the B\(^{act}\) complex to reversibly access proximal conformations, which in turn enable first-step splicing.

**Cwc25 enhances first-step splicing by H state stabilization**

Although the pre-mRNA is remodeled by the NTPase action of Prp2–Spp2, it does not undergo efficient first-step catalysis. Further enhancement of first-step splicing efficiency requires addition of Cwc25 to the B\(^{act}\) complex incubated with Prp2, Spp2 and ATP (Fig. 1b). Cwc25 was identified as one of a group of proteins in complex with Cef1–Ntc85 of the NTC complex\(^30\). To determine the role of Cwc25 in remodeling of the pre-mRNA, we performed SiMPull-FRET on the purified B\(^{act}\) complex supplemented with Prp2, Spp2, ATP and Cwc25 (henceforth referred to as C complex conditions). This resulted in a FRET histogram with an enhanced ~73% population with a mean FRET value of 0.75 ± 0.01 (s.d.) (Fig. 4a). We found the FRET states under C complex conditions to be the same as those under B* conditions, with the L2-to-H and M-to-H transitions prevalent; however, the occupancy in the H state was considerably enhanced under C conditions (Fig. 4b). TODP analysis revealed the fraction of molecules displaying at least one L2-to-H and one M-to-H transition to be similar under B* and
Cwc25 enhances the first step of splicing by stabilizing the H state. (a) A comparison of the aggregate molecular behavior before (B* condition) and after (C condition) Cwc25 addition through PSHs with all trajectories synchronized to start from either the M (top) or H state (bottom). (b) A comparison of the rate constants of the observable transitions under B* and C conditions. The thickness of the arrows corresponds to the relative rate constants. (c) Representative single-molecule FRET trajectories showing transition dynamics from the same molecules imaged before (B* conditions) and after (C conditions) Cwc25 addition with the raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories. The axis breaks represent 10 min of incubation after Cwc25 addition, a.u., arbitrary units.

Figure 5 Cwc25 enhances the first step of splicing by stabilizing the H state. (a) A comparison of the aggregate molecular behavior before (B* condition) and after (C condition) Cwc25 addition through PSHs with all trajectories synchronized to start from either the M (top) or H state (bottom). (b) A comparison of the rate constants of the observable transitions under B* and C conditions. The thickness of the arrows corresponds to the relative rate constants. (c) Representative single-molecule FRET trajectories showing transition dynamics from the same molecules imaged before (B* conditions) and after (C conditions) Cwc25 addition with the raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories. The axis breaks represent 10 min of incubation after Cwc25 addition, a.u., arbitrary units.

Figure 5 Cwc25 dynamically interacts near the BP upon B* formation

Previous studies have shown that Cwc25 binds stably to the spliceosome after Prp2-mediated SF3a–SF3b destabilization. It seems likely that Cwc25 enhances first-step chemistry by binding to the pre-mRNA, as mutation at the BP abolishes this interaction. Cwc25 was also recently shown to cross-link near the BP of the pre-mRNA. To directly observe the binding of Cwc25 to the pre-mRNA, we labeled the protein’s C terminus with Cy5. The Cy5 near the 5′ S5 of the pre-mRNA was pre-bleached so that the pre-mRNA had a single fluorescence signal from Cy3 label near the BP. We tested the activity of the Cy5-tagged Cwc25 (Cwc25-Cy5) using our bead pull-down assay and found it to be fully functional. SimPull-FRET experiments were then carried out with Cwc25-Cy5 added to the B* complex with and without Prp2, Sp3 and ATP (Fig. 6a). We observed repeated binding and dissociation of Cwc25-Cy5 and resulting FRET with the BP under B* conditions (Fig. 6b,c). On the basis of the FRET distribution of these binding events, centered around 0.37 ± 0.03 (s.d.), we estimate that Cwc25 binds within FRET distance of the Cy3-Cy5 pair (<100 Å), roughly 5 Å from the BP adenosine (Fig. 6b). The lower FRET peak, centered around 0.15, represents background signal. By contrast, there was little observable FRET between Cwc25 and the pre-mRNA BP in the absence of Prp2–Sp3 and ATP (Fig. 6d,e, B* condition). We conclude that Cwc25 activates the spliceosome for the first step by dynamically binding to the pre-mRNA near the BP.
**Figure 6** Prp2-mediated spliceosome remodeling creates a binding site for Cwc25 near the BP. (a) Schematic of our SiMPull-FRET experiment to test for binding of Cy5-tagged Cwc25 near the BP of spliceosome-associated pre-mRNA containing a single active Cy3 fluorophore. (b) FRET probability distribution of the raw single-molecule FRET trajectories under B* conditions (in the absence of Prp2-mediated remodeling). (c) FRET probability distribution of the raw single-molecule FRET trajectories under Bact conditions (in the presence of Prp2-mediated remodeling). (d) Representative single-molecule FRET trajectories showing the binding and associated FRET between the Cy5 on Cwc25 and the Cy3 near the pre-mRNA BP under B* conditions with the raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories. Arrows indicate binding events with close proximity to the BP. (e) Representative single-molecule FRET trajectory showing the absence of FRET between the Cy5 on Cwc25 and the Cy3 near the pre-mRNA BP under Bact conditions with the raw donor (Cy3, green), acceptor (Cy5, red) and FRET (black) trajectories. Sample sizes for the different conditions were comparable and effects measured are a reflection of the changes in the distribution of the population. a.u., arbitrary units.

**DISCUSSION**

Here we have combined single-molecule FRET between fluorophores attached near the 5′ SS and BP of the pre-mRNA substrate with affinity purification, in a technique we term SiMPull-FRET, to study the spliceosomal Bact complex stalled by heat inactivation of Prp2. Stepwise addition of ATP and the recombinant proteins Prp2, Spp2 and Cwc25 revealed the role of each factor in pre-mRNA remodeling (Fig. 7). We find that the pre-mRNA remains in the static low-FRET L2 state of the Bact complex (which, for clarity, we term L2act) with distal 5′ SS and BP until activation by Prp2–Spp2 in the presence of ATP produces the B* complex. Prp2-mediated hydrolysis of ATP (or UTP) in this step weakens the binding of some seven proteins, including SF3a and SF3b7,7, that bind the pre-mRNA upstream and downstream of the BP adenosine, presumably preventing its premature nucleophlic attack on the 5′ SS. Accordingly, the B*-associated low-FRET state (L2*) allows the pre-mRNA to transiently and reversibly visit two new states of either mid- (M*) or high-FRET (H*) and more proximal 5′ SS and BP. First-step splicing now proceeds with low efficiency, leading to post-catalytic C complex formation signified by a static high-FRET state (H5). This finding indicates that the increased proximity of the reactive sites is sufficient for catalysis. However, reaction chemistry is greatly enhanced by the addition of Cwc25, which binds the pre-mRNA substrate near the BP and slows particularly the rate constant of the high- to mid-FRET transition, leading to a longer dwell time in the precatalytic, stabilized FRET state H5-pre. In turn, this event leads to enhanced progression to the static high-FRET state associated with the post-catalytic C complex (H5) (Fig. 7).

Our data show that before the action of Prp2, Spp2 and ATP, the spliceosome keeps the reactive sites of the pre-mRNA strictly apart. This observation is consistent with and refines a recent report suggesting that stable splice-site juxtaposition occurs at some point after the NTC assembles on the pre-mRNA 33. Furthermore, it has previously been speculated that the catalytically activated B* complex may shift back and forth between inactive and active conformations 8. We have presented direct evidence for this hypothesis by showing that only in the B* state are dynamic excursions between low- and high-FRET

**Figure 7** Model for the conformational mechanism of first-step splicing. The 5′ SS and BP of the pre-mRNA in the Bact complex reside predominantly in the static distal L2act conformation with low FRET (i.e., high-Cy3 and low-Cy5 fluorescence, indicated by the green and red circles, respectively). Upon ATP hydrolysis and conversion into the B* complex, Prp2, along with its cofactor Spp2, unlocks the B*-associated low-FRET state L2* to reversibly sample the mid- and high-FRET (spatially proximal) conformations M* and H* (and H5-pre under C complex conditions). Cwc25 binds near the BP of the pre-mRNA, thus reducing the rate constant of the high- to mid-FRET transition and enhancing first-step chemistry, whereinupon the pre-mRNA adopts the static high-FRET state H5.
states observable, and only the high-FRET state places the reactive 5’ SS and BP in close enough proximity for subsequent catalysis, correlated with the appearance of the static high-FRET state H'C. The same authors also proposed that Cwc25 binding may shift the equilibrium between inactive and active conformations toward the latter, which we directly observe and assign to a marked increase of the dwell time in the active conformation H C-pre with proximal 5’ SS and BP.

The behavior of the spliceosome resembles that of a classical biased Brownian ratchet machine that draws path directionality from the random thermal fluctuations, which it constantly experiences, through a form of directional ‘rectification’ or ‘biasing’. In fact, the ribosome has been described as a biased Brownian ratchet machine and our previous single-molecule FRET probing of pre-mRNA dynamics in whole yeast cell extract suggested that the spliceosome, like the ribosome, works close to thermal equilibrium. We therefore propose that the ATP-driven helicase activity of Prp2-Spp2 acts to remove SF3a–SF3b as an impediment to the intrinsic thermal fluctuations of the spliceosome-substrate complex and Cwc25 provides directionality to the reaction pathway by then acting as a ‘pawl’ to stabilize the catalytically competent conformation. Perhaps the closest known analogy is found in translocation of the ribosome, where the random conformational ratcheting between the two ribosomal subunits at thermal equilibrium is rectified by GTP-bound elongation factor-G in conjunction with the interchange of a conserved two-nucleotide 16S ribosomal RNA pawl into the mRNA, which appears to prevent it from ratcheting back. We note that alteration between reversible thermal motion and irreversible NTP hydrolysis steps is thought to form the basis for repeated proofreading by the ribosome and may do so for the spliceosome. Future studies will probably further illuminate the molecular mechanisms of these events.

Finally, DEXD/H-box helicases such as Prp2 are widespread enzymes that participate in many aspects of RNA processing. In general, they are thought to use ATP hydrolysis to remodel RNA and RNP complexes by binding, unwinding and releasing the RNA. Unlike previous single-molecule approaches, our SiMPull-FRET approach has not only allowed us to unveil the dependence of pre-mRNA ratcheting on the NTPase activity of Prp2, which is kinetically biased by Cwc25 binding, but to do so in a well-controlled purified system. Biased Brownian ratcheting may be widespread among helicase-driven RNPs, and SiMPull-FRET will allow us to test this hypothesis further.

METHODS

Methods and any associated references are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

R.K., M.R.B. and M.L.K. performed the biochemical and single-molecule experiments and performed data analysis. M.L.K. cloned, expressed and labeled the proteins. R.K., M.R.B., M.L.K., J.A., C.G. and N.G.W. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Affinity purification of the B43 complex. Extracts were prepared from a prp-2-1 cef1-1-TAP yeast strain (ATCC 201388: MATa his3Δ1 leu2Δ0 met15Δ ura3Δ0)66 and heated at 37 °C for 40 min to inactivate Prp2 and stall the spliceosome at the B43 complex. In a final volume of 135 µl, 40% (v/v) of this heat-treated extract was incubated with ~50 pmol FRET-labeled Ubc4 pre-mRNA13 in the presence of 2 mM ATP in splicing buffer (8 mM HEPES-KOH, pH 7.0, 2 mM MgCl2, 0.08 mM EDTA, 60 mM K3[PO4], 20 mM KCl, 8% (v/v) glycerol, 3% (v/v) PEG, 0.5 mM DTT) and incubated at 23 °C for 35 min. For biochemical experiments, streptavidin-coated magnetic beads (Dynabeads MyOne Streptavidin C1, Invitrogen) were handled as per the manufacturer’s recommendation. For each splicing reaction, 200 µl of the suspended beads were equilibrated in 200 µl of T50 buffer (50 mM Tris-HCl, pH 7.5, 50 mM NaCl). An equal volume of 0.5 mg/ml biotin-IgG (ZyMAX rabbit anti-mouse IgG (H+L) − BT (ZyMAX Grade)) in T50 was added and incubated in a tube rotator at 23 °C for 30 min. The beads were then pulled down using a magnet, and the supernatant was discarded. To block any streptavidin not bound by biotin-IgG, the beads were incubated with excess free biotin at 1.5 mg/ml in T50 buffer in a tube rotator at 23 °C for 20 min. After equilibration in splicing buffer, the independently assembled splicing reactions were added and incubated in a tube rotator for 30 min at 23 °C to allow the protein A of the Cef1-TAP tag in the spliceosome complex to bind the biotin-IgG. Upon removal of the supernatant, the beads were further washed three times with buffer A (20 mM HEPES-KOH, pH 7.9, 120 mM KCl, 0.01% NP40, 1.5 mM MgCl2, 5% (v/v) glycerol) and once with splicing buffer to further purify the B43 complex. The reactions were scaled up for reconstitution reactions pursued in parallel and split at this step. Prp2, Spp2 and Cwc25 were added at 90–120 nM final concentration in splicing buffer in the presence or absence of 2 mM ATP or AMPNP or UTP and incubated in the tube rotator for 30–40 min for various degrees of reconstitution. RNA was isolated and products of splicing were analyzed on a denaturing gel (7 M urea, 15% polyacrylamide) and scanned on a Typhoon variable mode imager (GE Healthcare). Normalized product for Ubc4 was calculated by taking the amount of free 5’ exon and dividing by the total amount of pre-mRNA and free 5’ exon. Normalized product for actin was calculated by taking the amount of lariat intermediate and dividing by the total amount of pre-mRNA and lariat intermediate.

Cloning, expression and purification of splicing factor proteins. The full-length prp2 gene was PCR-amplified and ligated into plasmid pRSETA (Invitrogen) with a C-terminal His6 tag. The N-terminally truncated form of SPP2 (encoding amino acids 37–185) containing a C-terminal His6 tag, and the full-length Cwc25 gene were obtained from R. Lührmann (Max Planck Institute for Biophysical Chemistry, Germany). Cwc25 was subcloned into a pRSETA plasmid containing a single cysteine residue and His6 tag at the C terminus. The constructs were then transformed into Escherichia coli strain Rosetta II (Novagen). Cultures were grown in 2–4 L of TB medium and induced with 125 µM IPTG. Cultures were then incubated at 20 °C for 18 h. Cells were harvested, washed and the pellets stored at ~80 °C.

Purification of Cwc25-His and Spp2-His was performed as described67. Protein purity was confirmed by 16% SDS-PAGE (Supplementary Fig. 1) and proteins were either first fluorescence labeled or directly divided into aliquots, flash frozen in liquid nitrogen and stored at ~80 °C. Protein concentrations were determined by Bradford assay and measurement at A280. His-tagged Prp2 obtained from E. coli cell lysate was purified as described67. Protein purity was confirmed by 10% SDS-PAGE (Supplementary Fig. 1), and the final product was divided into aliquots, flash frozen in liquid nitrogen and stored at ~80 °C. Protein concentrations were determined by Bradford assay and measurement at A280. RNA sequences used in the study are reported in Supplementary Table 1.

Single-molecule FRET of purified splicesomal complexes. For the single-molecule FRET experiments on affinity-purified complexes, we prepared slides using previously published procedures14. In short, the surface of a quartz slide was amino functionalized, PEGylated and reacted with 0.2 mg/ml streptavidin in T50 buffer for 15 min at 23 °C. We flowed 100 µl of 0.5 mg/ml biotin-IgG in T50 onto the slide and incubated it for 20 min, followed by free biotin at 1.5 mg/ml in T50 buffer for 15 min. B43 splicesomal complexes were assembled and stalled as described above by incubation of FRET-labeled Ubc4 pre-mRNA with heat treated Prp2-1 Cef1-TAP yeast splicing extract in splicing buffer supplemented with 2 mM ATP and an oxygen scavenger system (OSS) composed of protocatechuate dioxygenase, protocatechuate and Trolox. These complexes were then flowed onto the slide surface and incubated for 15–20 min to allow the Cef1-TAP on the spliceosome to bind biotin-IgG. The slide surface was washed rigorously and reconstituted (in the presence of OSS) as described for the biochemical purification and incubated for 10–40 min before acquiring data. A home-built prism-based TIRF microscope was used to collect data as described13,48,49. To obtain FRET data, we directly excited the Cy3 donor near the BP adenosine with a 532-nm laser, and we recorded emission by Cy3 and Cy5 fluorophores at 100-ms time resolution using an intensified CCD camera (I-Pentamax, Princeton Instruments).

Fluorescent labeling of Cwc25 and distance estimation from FRET. The single-cysteine mutant of Cwc25 was labeled with Cy5-maleimide (GE Healthcare). Labeling was performed using 0.150 µmol of Cwc25 in storage buffer and 0.5 mg of dye containing 10 µM reducing agent Tris(2-carboxyethyl)phosphine (TCEP) (Sigma). Reactions were incubated at 23 °C for 1 h followed by overnight at 4 °C. Free dye was removed by re-purification of protein on an Ni2+ column and dialysis back into storage buffer. The degree of labeling was determined using GE Healthcare’s protocol and was found to be 70%. Protein functionality was confirmed using an ensemble pulldown assay as described above. The fluorescent distance, R, and the apparent FRET efficiency, Eapp, were calculated as described48,53,54 from the equations

\[ E_{app} = \frac{1}{1 + \left(\frac{R}{R_0}\right)^6} \]

where \( c = 0.69 \) and \( R_0 = 54 \) Å, and

\[ E_{app} = \frac{\frac{I_{C5}}{I_{C5} + I_{C3} \times \left(\frac{\eta_{C5}}{\eta_{C3}}\times \frac{\phi_{C5}}{\phi_{C3}}\right)}}{I_{C5} + I_{C3} \times \left(\frac{\eta_{C5}}{\eta_{C3}}\right)} \]

\( \varphi \) and \( \eta \) signify fluorophore quantum yields and detector channel efficiencies, respectively. The donor and acceptor intensities \( I_{C3} \) and \( I_{C5} \), respectively, were corrected for leakage of donor photons into the acceptor channel.

Single-molecule data analysis. Cross-correlation analysis was carried out using customized MATLAB scripts with built-in xcorr function. Time lags for the cross-correlation ranged from 0 to 5 s (0–50 frames of each 0.1-s integration time). Quality control for the raw smFRET trajectories obtained from the experimental conditions was performed as described13. Histograms for data sets measuring pre-mRNA dynamics were constructed by sampling 100 frames of data from each molecule. Histograms for Cwc25-Cy5 and BP-Cy3 FRET experiments were constructed by sampling the entire length of data. Hidden Markov modeling (HMM) was performed on trajectories using the vbFRET software suite14. Each trajectory was individually fit with models ranging from one to five states with the optimal number of states determined by the vbFRET algorithm. The inherent experimental variations of the FRET signal between single molecules leads to a slightly different state assignment for similar states across different molecules. A K-means clustering approach was therefore performed in MATLAB to group similar states into larger macro states (L1, L2, M and H). A matrix cataloging the HMM assigned FRET state, raw FRET level and difference in donor and acceptor intensities for each HMM derived event was used as input for the K-means algorithm (Supplementary Fig. 3 and Supplementary Table 2). Four macro states were identified whose boundaries were used to assign the original HMM idealized FRET states. The number of macro states was determined using the Bayesian information criterion (BIC) as a model selection tool. K-means and BIC have been used previously to group and accurately determine the number of clusters in single-molecule data13-15. The BIC score was calculated in MATLAB as follows:

\[ BIC = -(2 \times LLF) + (\text{NumParams} \times \log(\text{NumObs})) \]

where LLF is the log-likelihood function, NumParams the number of parameters and NumObs the number of observations. The number of clusters (K) varied from two to eight. The lowest BIC score was achieved for K = 4 (Supplementary Fig. 3c). Once identified, the clusters were tested for similarity using the multi-comparison Tukey test in GraphPad PRISM. The results of this analysis indicate...
that there is a significant difference ($P < 0.001$) for the pairwise comparison of all four clusters (Supplementary Fig. 3d).

Transition occupancy density plots (TODPs) were used to plot the fraction of molecules that contain any given HMM transition at least once\textsuperscript{14}. Molecules that did not exhibit any transitions were plotted along the diagonal at their respective positions. For kinetic rate calculations, transition density plots (TDPs) that are scaled by the number of times a transition occurs irrespective of how many molecules exhibit that transition were used as described\textsuperscript{14}. A cumulative histogram scatter plot was then fit with a double-exponential association equation in MicroCal Origin (Supplementary Fig. 4). A weighted average ($k_{\text{observed}}$) of the two rate constants from the double-exponential fits was calculated based on the amplitude values of the exponential equation. To correct for bias introduced by the limited observation window used to measure dwell times, the measured $k_{\text{observed}}$ values were corrected by subtracting the photobleaching rate constant and the reciprocal of the observation window to yield $k_{\text{actual}}$ as described\textsuperscript{55}. Equilibrium constants ($K_{\text{eq}}$) were calculated by taking a ratio of the forward and backward rate constants for a set of state-to-state transitions. Post-synchronized histograms (PSHs) were constructed by synchronizing individual FRET events to the time where one of the macro-states (M or H) was achieved. The color bar in Figure 5a represents the fraction of FRET events that exhibits a certain FRET state at a given time.

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