

UCSF

UC San Francisco Previously Published Works

Title

An unanticipated early function of DEAD-box ATPase Prp28 during commitment to splicing is modulated by U5 snRNP protein Prp8

Permalink

<https://escholarship.org/uc/item/96k5k2t6>

Journal

RNA, 20(1)

ISSN

1355-8382

Authors

Price, Argenta M
Görnemann, Janina
Guthrie, Christine
[et al.](#)

Publication Date

2014

DOI

10.1261/rna.041970.113

Copyright Information

This work is made available under the terms of a Creative Commons Attribution-NonCommercial License, available at <https://creativecommons.org/licenses/by-nc/4.0/>

Peer reviewed

An unanticipated early function of DEAD-box ATPase Prp28 during commitment to splicing is modulated by U5 snRNP protein Prp8

ARGENTA M. PRICE,¹ JANINA GÖRNEMANN,^{2,3,4} CHRISTINE GUTHRIE,^{1,5} and DAVID A. BROW^{2,5}

¹Department of Biochemistry and Biophysics, University of California, San Francisco, California 94143, USA

²Department of Biomolecular Chemistry, School of Medicine and Public Health, University of Wisconsin, Madison, Wisconsin 53706, USA

³Max Planck Institute of Molecular Cell Biology and Genetics, 01307 Dresden, Germany

ABSTRACT

The stepwise assembly of the highly dynamic spliceosome is guided by RNA-dependent ATPases of the DEAD-box family, whose regulation is poorly understood. In the canonical assembly model, the U4/U6.U5 triple snRNP binds only after joining of the U1 and, subsequently, U2 snRNPs to the intron-containing pre-mRNA. Catalytic activation requires the exchange of U6 for U1 snRNA at the 5' splice site, which is promoted by the DEAD-box protein Prp28. Because Prp8, an integral U5 snRNP protein, is thought to be a central regulator of DEAD-box proteins, we conducted a targeted search in Prp8 for cold-insensitive suppressors of a cold-sensitive Prp28 mutant, *prp28-1*. We identified a cluster of suppressor mutations in an N-terminal bromodomain-like sequence of Prp8. To identify the precise defect in *prp28-1* strains that is suppressed by the Prp8 alleles, we analyzed spliceosome assembly in vivo and in vitro. Surprisingly, in the *prp28-1* strain, we observed a block not only to spliceosome activation but also to one of the earliest steps of assembly, formation of the ATP-independent commitment complex 2 (CC2). The Prp8 suppressor partially corrected both the early assembly and later activation defects of *prp28-1*, supporting a role for this U5 snRNP protein in both the ATP-independent and ATP-dependent functions of Prp28. We conclude that the U5 snRNP has a role in the earliest events of assembly, prior to its stable incorporation into the spliceosome.

Keywords: pre-mRNA splicing; Prp28; Prp8; commitment complex; pre-spliceosome

INTRODUCTION

The spliceosome is a macromolecular machine that locates and excises introns with single-nucleotide precision. To accurately identify and remove introns, the spliceosome undergoes a stepwise series of binding events and conformational rearrangements, guided by proteins of the DExD/H-box family of RNA-dependent ATPases. In the canonical model of spliceosome assembly (for review, see Will and Lührmann 2011) the U1 snRNP first recognizes the 5' splice site (5'ss), forming commitment complex 1 (CC1). The branch site and 3' end of the intron are then recognized by two proteins, BBP and Mud2 in yeast, forming commitment complex 2 (CC2). Next, the first ATP-dependent step of splicing occurs: DEAD/DECD-box proteins Prp5 and Sub2 promote rearrangements that replace BBP and Mud2 with the U2 snRNP at the branch site, forming the pre-spliceosome or A com-

plex. The U4/U6.U5 triple-snRNP (tri-snRNP) then joins to form the assembled spliceosome or B complex.

Once assembled, the spliceosome undergoes activation for catalysis. DEAD-box protein Prp28 promotes release of the U1 snRNP, allowing U6 to base-pair with the 5'ss. In coordination with Prp28, DEIH-box helicase Brr2 (a stable component of the U5 snRNP) unwinds U4 from U6, allowing U6 to base-pair with U2. Together, these ATP-dependent activities of Prp28 and Brr2 form the U2/U6.U5 spliceosome (B^{act} complex) competent for the final steps of activation, which include binding of the NineTeen Complex (NTC) and additional rearrangements catalyzed by the DEAH-box protein Prp2. While this general sequence of events is well supported, several studies have also implicated the U5 snRNP in very early steps of spliceosome assembly, before stable incorporation of the tri-snRNP (Wassarman and Steitz 1992; Wyatt et al. 1992; Newman et al. 1995; Ast and Weiner 1997; Maroney et al. 2000). In addition, the mechanisms underlying the

⁴Present address: Department of Cellular and Molecular Medicine, KU Leuven, BE-3000 Leuven, Belgium

⁵Corresponding authors

E-mail christineguthrie@gmail.com

E-mail dabrow@wisc.edu

Article published online ahead of print. Article and publication date are at <http://www.rnajournal.org/cgi/doi/10.1261/rna.041970.113>.

© 2013 Price et al. 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/>.

timing and coordination of each assembly step remain poorly understood. In this study, we focus on Prp28 and its interactions with Prp8, an integral component of the U5 and tri-snRNPs, thought to be a central regulator of spliceosome activity (Kuhn et al. 2002; Grainger and Beggs 2005; Galej et al. 2013).

Several studies have provided evidence that Prp28 promotes U1 snRNP release and is involved in stable tri-snRNP incorporation into the spliceosome. The cold-sensitive *prp28-1* mutation, G279E, is in motif 1b of the DEAD-box domain (Strauss and Guthrie 1991, 1994), which likely contacts an as yet unknown RNA substrate (Fairman-Williams et al. 2010). It causes a broad splicing defect (Pleiss et al. 2007) and exacerbates mutations that hyperstabilize the base-pairing between U1 and the 5' splice site (Staley and Guthrie 1999). Other mutations in Prp28 reduce fidelity of U6/5' splice site base-pairing, the interaction that replaces U1/5' splice site pairing (Yang et al. 2013). Further supporting a role in U1 snRNP release, Prp28 is nonessential in strains with mutations that are predicted to destabilize the U1/5' splice site interaction (Chen et al. 2001; Hage et al. 2009; Schwer et al. 2013). Finally, U1/5' splice site hyperstabilization and *prp28-1* cause similar in vitro spliceosome assembly defects, which include inhibition of U4/U6 unwinding and reduced stability of tri-snRNP association (Staley and Guthrie 1999). In mammalian cells, Prp28 is a stable component of the tri-snRNP (Teigelkamp et al. 1997) and is directly involved in its recruitment to the spliceosome (Ismaili et al. 2001; Mathew et al. 2008). However, yeast Prp28 appears to be less stably associated with tri-snRNP (Strauss and Guthrie 1994; Gottschalk et al. 1999; Stevens et al. 2001; Small et al. 2006), so it is unclear whether the defect in tri-snRNP incorporation in the *prp28-1* strain is direct or is an indirect consequence of the block to U1 snRNP release.

The Prp28- and Brr2-catalyzed releases of the U1 and U4 snRNPs appear to be coordinated: mutations that inhibit either step also block the other (Kuhn et al. 1999; Staley and Guthrie 1999). Prp8 is a good candidate for regulating Prp28 and coordinating its activity with that of Brr2. The Jab1/MPN domain at the C terminus of Prp8 is known to regulate Brr2 activity (Maeder et al. 2008; Mozaffari-Jovin et al. 2012, 2013). Furthermore, the N-terminal domain (NTD) of Prp8 harbors a mutation that suppresses the cold-sensitivity of the *prp28-1* mutation (Kuhn et al. 2002) and also physically interacts with two U1 snRNP proteins (Abovich and Rosbash 1997; van Nues and Beggs 2001). A recent study found that Prp8 cross-links to the U1 snRNA, and mutations in U1 that reduce this cross-link also reduce triple-snRNP association with the pre-mRNA (Li et al. 2013). Therefore, Prp8 is perfectly positioned to promote tri-snRNP joining through interactions with U1 and then regulate Prp28-directed-U1 release.

To define Prp28's interaction with Prp8, we conducted a targeted selection for cold-insensitive suppressors of *prp28-1* in the Prp8-NTD. We identified 15 suppressor substitutions that cluster in a proposed bromodomain (Dlakić and

Mushegian 2011). To determine the precise order of spliceosome assembly defects caused by *prp28-1* and to identify which defects are suppressed by the Prp8 alleles, we monitored the kinetics of cotranscriptional spliceosome assembly in vivo and post-transcriptional spliceosome assembly in vitro. As expected, *prp28-1* blocks U1 and U4 release and reduces stable tri-snRNP association. Surprisingly, however, *prp28-1* also inhibits formation of the pre-spliceosome and ATP-independent CC2. Therefore, we propose that Prp28 has at least two distinct roles in spliceosome assembly and activation: an ATP-independent role in CC2 formation and a subsequent ATP-dependent role in U1 release. A *prp28-1*-suppressor mutation in the *PRP8-NTD* partially alleviates all of the spliceosome assembly and activation defects caused by *prp28-1*, including the early CC2 formation defect. Our results provide evidence that Prp8 and the U5 snRNP have a role in the earliest events of spliceosome assembly, prior to the stable incorporation of the tri-snRNP into the spliceosome.

RESULTS

prp8-tes alleles: substitutions in the N-terminal quarter of Prp8 that suppress *prp28-1*

In order to characterize the interactions between Prp28 and its potential regulator, Prp8, we screened the NTD of Prp8 for suppressors of *prp28-1*. Prp8 is a 280-kDa protein that is an integral component of the U5 and tri-snRNPs. Substitutions in distinct regions of Prp8 suppress mutations in a variety of spliceosomal proteins as well as mutations in snRNAs and conserved splice signals in the pre-mRNA itself (Grainger and Beggs 2005). Recent studies have provided deep insight into the structure of the C-terminal two-thirds of Prp8 (Galej et al. 2013). The C-terminal domains of Prp8 contact the 5' splice site with an RNase-H-like fold (Pena et al. 2008; Ritchie et al. 2008; Yang et al. 2008; Schellenberg et al. 2013) and regulate Brr2 through a Jab/MPN domain (Pena et al. 2007; Maeder et al. 2008; Mozaffari-Jovin et al. 2012, 2013). The N-terminal portion of Prp8 remains structurally elusive, but is particularly interesting in connection with Prp28. In a prior selection for suppressors of a cold-sensitive mutation in U4 RNA (U4-cs1) that blocks U4 and U1 release at 16°C, we isolated an N-terminal Prp8 substitution (L280P) that suppressed both U4-cs1 and *prp28-1* (Kuhn et al. 2002). This substitution was the only U4-cs1-suppressor of 13 tested that also suppressed *prp28-1*, implying that suppression of *prp28-1* is due to alteration of a specific interaction rather than a general effect of decreased Prp8 activity.

Therefore, we sought to map this interaction surface in the N-terminal quarter of Prp8 by mutagenizing codons 1–660 of *PRP8* by error-prone PCR and selecting alleles that allow a *prp28-1* strain to grow at 16°C (Fig. 1A,B; Table 1). Thirty-seven cold-resistant clones were isolated, 16 of which contained the substitution D273G, either as the sole substitution (eight clones) or as one of two or more substitutions.

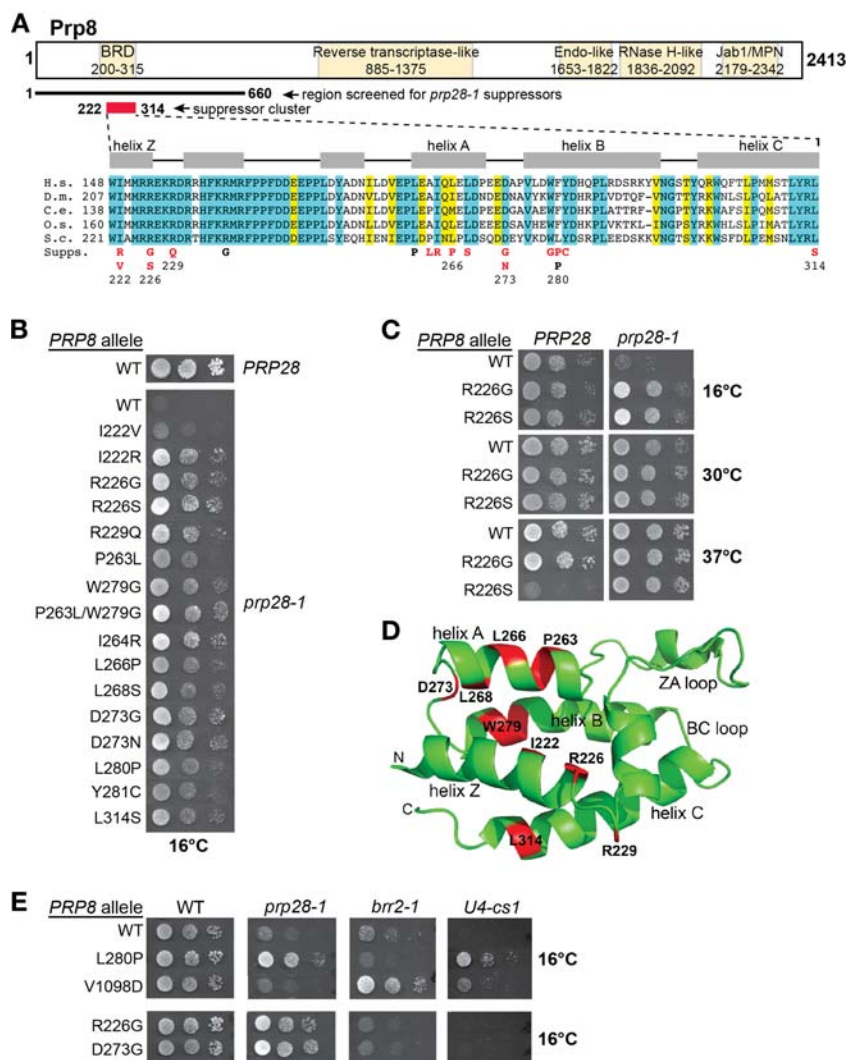


FIGURE 1. Suppressors of *prp28-1* selected in the N-terminal quarter of Prp8. (A) Map of Prp8, showing domains identified by sequence or structural homology (Grainger and Beggs 2005; Dlakić and Mushegian 2011; Galej et al. 2013). Numbers indicate amino acid residues. (BRD) Bromodomain-like (Dlakić and Mushegian 2011). The region that was screened for *prp28-1*-suppressor mutations and the subregion in which such mutations were obtained are indicated. Sequences of human (H.s.), fly (D.m.), worm (C.e.), rice (O.s.), and budding yeast (S.c.) Prp8 are shown below, aligned in the region containing suppressor mutations, with the position number of the first residue and selected altered residues indicated. Residues highlighted in blue are identical in all five sequences, while those highlighted in yellow are highly similar. Substitutions that suppress *prp28-1* are shown in red below the alignment, and substitutions that suppress U4-cs1 (Kuhn and Brow 2000) are shown in black. Locations of putative α -helices (see D) are indicated above the sequences. (B) *prp8-tes* alleles suppress the cold-sensitive growth defect of *prp28-1*. Tenfold serial dilutions of wild-type, *prp28-1*/PRP8, and the indicated *prp28-1*/*prp8-tes* strains were spotted on YEPD medium and incubated at 16°C for 10 d. (C) The *prp8-R226S* mutation confers a heat-sensitive growth defect that is suppressed by *prp28-1*. Another substitution at the same position, *prp8-R226G*, is not heat sensitive. Tenfold serial dilutions on YEPD are shown. (D) Location of *prp8-tes* substitutions mapped onto a structural model of the putative Prp8 bromodomain (model from Dlakić and Mushegian 2011). Residues altered by *prp8-tes* mutations are shown in red and selected residues and secondary structure elements are labeled. (E) *prp8-tes* mutations do not suppress the cold-sensitive phenotype caused by mutations in *BRR2* or the U4 RNA gene. Tenfold serial dilutions of strains were spotted on YEPD medium and incubated at 16°C for 10 d, except for U4-cs1 strains (18°C for 12 d). Controls *prp8-L280P* and *prp8-V1098D* behaved as in Kuhn et al. (2002).

Three additional suppressors contained a D273N substitution, once as the sole mutation. Seven clones had an R226G substitution, in two cases as the sole mutation, and another clone had the R226S substitution. Of the remaining 10 clones, three had at least one substitution between residues 222 and 229, six had at least one change between residues 263 and 281, and one had the substitution L314S. Silent (synonymous) and secondary mutations spanned codons 20–595 in the mutagenized region, indicating that the tight clustering of the suppressor mutations is not an artifact of the PCR mutagenesis or in vivo recombination. Furthermore, the most common suppressor mutations were isolated in three independent selections; therefore, it is unlikely that they were over-represented in the mutant pool by chance.

Working from the collection of suppressor alleles, we generated PRP8 alleles containing 15 different single amino acid substitutions at 12 positions between residues 222 and 314. We refer to these collectively as “*prp8-tes*” alleles, for *twenty-eight-1 suppression*. The *prp8-tes* alleles were scored for their strength of suppression of *prp28-1* (Fig. 1B; Table 1). P263L and W279G gave stronger suppression when present in combination, as isolated in the selection, than when present individually. The previously isolated L280P substitution conferred a level of cold-resistant growth similar to that of the newly selected suppressors of *prp28-1*.

To assess the general effects of the suppressor substitutions on Prp8 function, we also tested the *prp8-tes* alleles for growth phenotypes in the presence of wild-type PRP28. The R226S substitution conferred a strong heat-sensitive growth defect at 37°C, but supported normal growth at 16°C and 30°C (Fig. 1C). All other substitutions exhibited no qualitative growth defect at 16°C, 30°C, or 37°C. Importantly, the *prp8-R226S*, *prp28-1* double-mutant strain was not heat sensitive (Fig. 1C), indicating that these two mutations mutually suppress one another.

Of the 12 residues altered by *prp8-tes* mutations, nine are invariant in yeast, rice, fruit fly, worm, and human Prp8

TABLE 1. Suppressors of *prp28-1* in the N-terminal quarter of Prp8

Substitution	No. of independent clones	Isolated as single substitution	Strength of suppression
I222R	1	1	+++
I222V	1	1	+
R226G	7	2	+++
R226S	1	0	++++
R229Q	1	0	+++
P263L	1	0 ^a	++
I264R	1	0	++++
L266P	1	0	++
L268S	2	2	+++
D273G	16	8	++++
D273N	3	1	++++
W279G	1	0 ^a	+++
L280P ^b	NA	NA	++
Y281C	1	0	++
L314S	1	0	+++
P263L/W279G	1	NA	++++

(NA) Not applicable.

^aP263L/W279G were isolated as double substitution.^bIdentified previously (Kuhn et al. 2002).

(Fig. 1A), consistent with an important function. Intriguingly, the region encompassing the *prp8-tes* substitutions corresponds to a recently proposed bromodomain (BRD) in Prp8 (Dlakić and Mushegian 2011). Bromodomains bind acetylated lysine residues in histones and other proteins (Filippakopoulos and Knapp 2012). The putative Prp8 bromodomain lacks several residues important for acetylated-lysine binding (Dlakić and Mushegian 2011) and so may serve as a general protein–protein interaction domain instead. Strikingly, when the suppressor residues are mapped on the structural model of the bromodomain from Dlakić and Mushegian (2011), the residues cluster tightly in the four α -helices of a helical bundle (Fig. 1D), in contrast to their dispersal in the primary structure. All but three of the 15 substitutions substantially alter the side-chain structure, and thus are likely to destabilize the putative four-helix bundle itself and/or its interaction with another protein or RNA.

Substitutions in Prp8 that suppress *prp28-1* do not generally suppress U4-cs1

Since the original *prp28-1* suppressor, Prp8-L280P, was isolated in a selection for suppressors of the cold sensitivity of U4-cs1, we hypothesized that mutations in the Prp8-NTD that decrease reliance of the spliceosome on Prp28 may also counteract the effects of U4-cs1. To test this hypothesis, we assessed the ability of *prp8-tes* alleles to suppress U4-cs1. None other than L280P suppressed U4-cs1, nor did any suppress the cold-sensitive growth phenotype conferred by *brr2-1*, which blocks U4 release (Fig. 1E; Raghunathan and Guthrie 1998; data not shown). This result indicates that suppression of the growth defect of *prp28-1* is not sufficient for

suppression of U4-cs1, and suggests that L280P is unusual in its ability to suppress both mutations.

prp8-tes substitutions do not promote U1/5'ss destabilization or alter the interaction of Prp8-NTD with Prp39, Prp40, or Prp28 in vitro

We next tested the hypothesis that disruption of Prp8–U1 snRNP interactions by the *prp8-tes* alleles could bypass the growth requirement for Prp28 function. The *prp8-tes* mutations cluster in a potential protein–protein interaction interface, and the Prp8-NTD is known to interact with U1 snRNP proteins Prp39 and Prp40 (Abovich and Rosbash 1997; van Nues and Beggs 2001). Further, Prp28 is nonessential in the presence of mutations in U1 snRNP proteins or the U1 snRNA that are expected to destabilize the U1/5'ss interaction (Chen et al. 2001; Hage et al. 2009; Schwer et al. 2013). Therefore, if loss of Prp8–U1 interactions similarly leads to destabilization of U1 snRNP from the 5'ss, they might bypass the requirement for Prp28. However, we found that a *prp8-tes* allele did not rescue the lethality caused by a complete absence of Prp28 protein or by a substitution predicted to destroy the catalytic activity of Prp28 (*prp28-D341N*) (Fig. 2A). Nor did a *prp8-tes* allele suppress the low-temperature growth defect caused by hyperstabilization of U1/5'ss base-pairing (Fig. 2B). These results indicate that *prp8-tes* alleles do not significantly destabilize the U1/5'ss interaction.

To more directly test the effect of *prp8-tes* alleles on binding of the Prp8-NTD to Prp39 or Prp40, we used in vitro binding assays. An N-terminal fragment of Prp8 (Prp8 residues 1–330) with an N-terminal glutathione S-transferase (GST) tag and either wild-type sequence or one of three *prp8-tes* substitutions was expressed in *E. coli*. Full-length Prp39 and Prp40 with C-terminal His₆ tags were also expressed in *E. coli*. Using a standard GST pull-down method, we reproduced the known interactions of the Prp8-NTD with Prp39 and Prp40. However, wild-type Prp8-NTD and the three *prp8-tes* alleles all pulled down similar amounts of both Prp39 and Prp40 (Fig. 2C). Taken together, our results indicate that *prp8-tes* alleles suppress *prp28-1* by a means other than altering interactions between Prp8 and Prp39 or Prp40, or by directly promoting destabilization of the U1/5'ss interaction.

If the *prp8-tes* mutations are not suppressing *prp28-1* indirectly by destabilizing the U1 snRNP–5' splice-site interaction, they may be acting directly by altering a Prp8–Prp28 interaction. Although a direct interaction between Prp28 and the Prp8-NTD has not been reported, we were able to detect binding by GST pull-down using moderately stringent wash conditions (200 mM KOAc at room temperature). However, the *prp8-tes* substitutions did not alter that interaction under the conditions of our assay (Fig. 2D). These results do not exclude the possibility that the *prp8-tes* substitutions alter Prp8–Prp28 interactions in a way that alters regulation of Prp28 activity by Prp8.

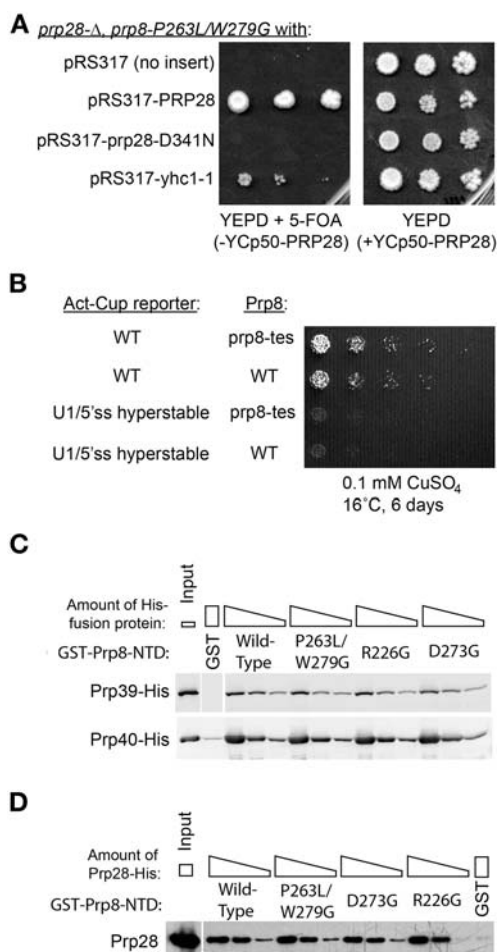


FIGURE 2. *prp8-tes* does not bypass Prp28, suppress U1/5'ss hyperstabilization, or alter interactions with U1 proteins Prp39 or Prp40 in vitro. (A) *prp8*-P263L/W279G does not bypass the requirement for Prp28. A *prp28* Δ /*prp8* Δ strain (JG140) containing wild-type PRP28 on a URA3-marked plasmid and pRS313-*prp8*-P263L/W279G was transformed with pRS317 (empty vector), pRS317-PRP28, pRS317-*prp28*-D341N (should be catalytically inactive), or pRS317-*yhc1*-1, which is known to bypass Prp28 (Chen et al. 2001). Tenfold serial dilutions were plated on YEFD or YEFD + 5-FOA and grown at 30°C for 3 d. Only strains with pRS317-PRP28 or pRS317-*yhc1*-1 were able to grow in the absence of the URA3-marked PRP28 plasmid. (B) *prp8-tes* does not suppress U1/5'ss hyperstabilization. Fivefold serial dilutions of strains with PRP8 or *prp8*-P263L/W279G and wild-type or U1/5'ss hyperstabilized Act-Cup splicing reporters (Staley and Guthrie 1999) were plated onto SD-Ura medium with 0.1 mM copper sulfate, and grown at 16°C for 6 d. In the presence of the U1/5'ss hyperstabilized Act1-Cup1 reporter, wild-type PRP8 and *prp8-tes* strains grow equally poorly at 16°C on 0.1 mM copper sulfate. (C) Prp39 and Prp40 bind similarly to wild-type Prp8-NTD, Prp8-NTD(P263L/W279G), Prp8-NTD(R226G), and Prp8-NTD(D273G). His₆ fusion Prp39 or Prp40 proteins were added to bead-bound GST-fusion Prp8-NTD proteins in a threefold dilution series. GST-only beads were incubated with the highest amount of His-fusion protein, and Input lanes contain 25% of the lowest amount of His-fusion protein used in the binding reactions. (D) Prp28 binds similarly to Prp8-NTD and Prp8-NTD-*tes* alleles. His₆ fusion Prp28 was added to bead-bound GST-fusion Prp8-NTD proteins in a fourfold dilution series. GST-only beads were incubated with the highest amount of Prp28 protein, and Input lane contains 65% of the lowest amount of Prp28 used in the binding reactions.

A *prp8-tes* allele partially suppresses the *prp28-1* global splicing defect

To measure the effect of a *prp8-tes* mutation on the global splicing defect caused by *prp28-1* in vivo, we used splicing-specific DNA microarrays (Pleiss et al. 2007). The *prp8-tes* strain P263L/W279G was chosen for this and subsequent assays, as it displays strong suppression of the *prp28-1* growth defect. Cy3- or Cy5-labeled cDNA was made from total cellular RNA isolated from wild-type, *prp28-1*, *prp8-tes*, or *prp28-1/prp8-tes* double-mutant strains, then was competitively hybridized to DNA microarrays as indicated in Figure 3.

Genome-wide splicing improved in the *prp28-1/prp8-tes* strain relative to *prp28-1* alone. The *prp28-1* strain exhibited a broad splicing defect at 16°C, characterized by increases in intron levels and decreases in spliced RNA levels relative to wild-type, consistent with our previous report (Fig. 3, lanes 6,10; Pleiss et al. 2007). The *prp8-tes* strain by itself behaved similarly to wild type (lanes 1,5,9), as expected given its lack of growth defect. Suppression of the *prp28-1* splicing defect can be seen by comparing the *prp28-1* versus wild-type pattern to the *prp28-1/prp8-tes* versus wild-type pattern (cf. lanes 6 and 7), or by directly testing suppression on the array by competitively hybridizing cDNA from *prp28-1* against cDNA from *prp28-1/prp8-tes* (lanes 4,8,12). Intron-containing RNA accumulated to some extent in the suppressed double mutant relative to wild type (lane 7). A mild level of suppression is consistent with the incomplete suppression of the *prp28-1* growth defect (Fig. 1B).

U2 snRNP joining and subsequent steps of spliceosome assembly are inhibited by *prp28-1* in vivo

In order to determine the precise defect in the *prp28-1* strain that is suppressed by *prp8*-P263L/W279G, we monitored the kinetics of cotranscriptional spliceosome assembly in *prp28-1* and *prp28-1/prp8-tes* strains using chromatin immunoprecipitation (ChIP). We measured the occupancy of several HA-tagged splicing factors (representatives of the U1, U2, and U5 snRNPs and of the NTC) at five positions along an intron-containing gene, *ECM33*. As pre-mRNA is synthesized, splicing factors bind and can be cross-linked to nearby DNA. Thus, the position of cross-linking along a gene can be used as a readout for splicing factor association with the nascent pre-mRNA (Fig. 4A; Kotovic et al. 2003; Görnemann et al. 2005; Lacadie and Rosbash 2005). To compare spliceosome assembly at permissive and nonpermissive temperatures, cultures were grown at 30°C, then divided and incubated at either 30°C or 16°C for 45 min.

Our results for a wild-type strain at 30°C are consistent with previous reports (Görnemann et al. 2005; Lacadie and Rosbash 2005). U1 snRNP association peaked near the region of *ECM33* DNA that corresponds to the 3'ss, then decreased by 500 bp downstream to a level similar to that at the 5'ss,

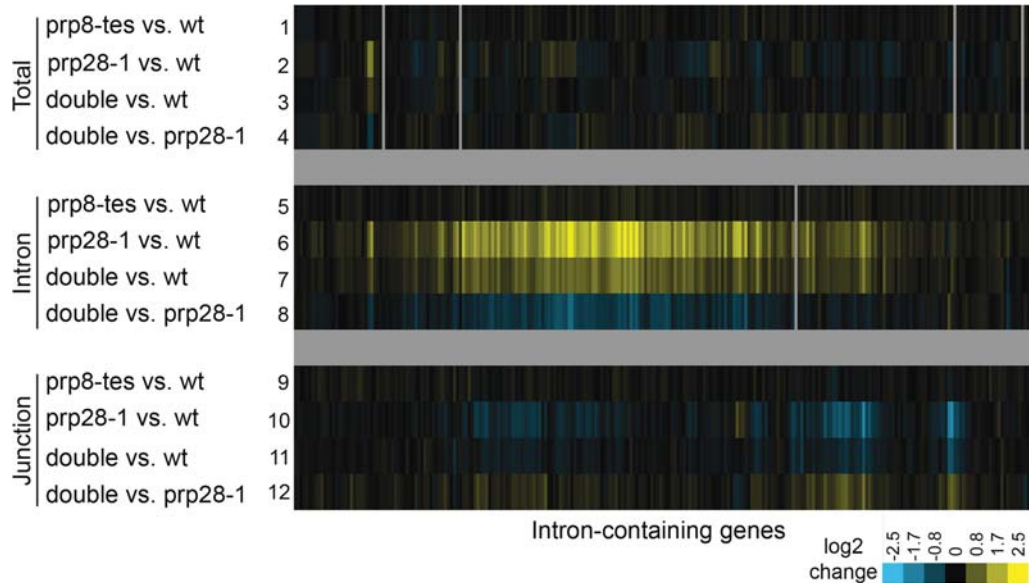


FIGURE 3. The global splicing defect of the *prp28-1* mutant is partially suppressed by *prp8-P263L/W279G*. Each of 253 introns is represented by three probes on a DNA microarray (Pleiss et al. 2007): one in an exon to detect total RNA (lanes 1–4), one in the intron to detect unspliced pre-mRNA (lanes 5–8), and one across the exon–exon junction to detect spliced mRNA (lanes 9–12). Differentially labeled cDNA samples from pairs of strains were competitively hybridized to each array and log₂-transformed normalized fluorescence ratios are presented. Yellow indicates an increase (ratio >0 [log₂]) and blue a decrease (ratio <0 [log₂]) in signal. (Lanes 1,5,9) *prp8-P263L/W279G* hybridized against wild-type; (lanes 2,6,10) *prp28-1* against wild type; (lanes 3,7,11) *prp28-1 + prp8-P263L/W279G* (“double”) against wild type; (lanes 4,8,12) *prp28-1 + prp8-P263L/W279G* against *prp28-1* to directly measure suppression. Results presented are the average of two biological replicates, each with a dye-flip technical replicate.

consistent with release of the U1 snRNP upon activation of the spliceosome (Fig. 4B, black). U2 snRNP occupancy was highest near the 3′ss, then slowly decreased over the next 1000 bp (Fig. 4C). U5 and NTC associations were similar: they peaked over the second exon, consistent with tri-snRNP and NTC addition following U2 snRNP binding (Fig. 4D,E). When wild-type cultures were shifted to 16°C prior to cross-linking for ChIP, the patterns of U1 snRNP and NTC binding were similar to those at 30°C (Supplemental Fig. 1). U2 and U5 snRNP associations were slightly delayed compared with 30°C, but reached similar maximum levels (Supplemental Fig. 1).

The *prp28-1* strain shifted to restrictive temperature (16°C) yielded a dramatically different ChIP pattern (Fig. 4B–E, orange). In agreement with the function of Prp28 in displacing the U1 snRNP from the 5′ss, we observed persistence of the U1 snRNP signal in the *prp28-1* background up to 1500 bp downstream from the 5′ss (Fig. 4B). This delayed release was most pronounced at 16°C, but was evident even at permissive temperature (30°C). U2 snRNP recruitment was predicted to be unaltered by the *prp28-1* mutation because the U2 snRNP is thought to bind the branch point well before Prp28 removes the U1 snRNP from the 5′ss (for review, see Will and Lührmann 2011). Surprisingly, shifting the *prp28-1* strain to 16°C caused a pronounced delay in U2 snRNP recruitment and decreased the maximum U2 snRNP signal (Fig. 4C). We also observed a severe delay in U5 snRNP binding at 16°C: the highest signal was at the

end of the gene and was well below the peak signal observed at 30°C (Fig. 4D). This finding was confirmed by analyzing the U5 snRNP pattern along *SEC27*, a gene with a longer second exon than *ECM33* (Supplemental Fig. 2). We conclude that *prp28-1* causes a delay in recruitment of the U2 and tri-snRNPs to introns in vivo.

The *prp28-1* strain exhibited even more dramatically delayed/reduced association of NTC, even at permissive temperature (Fig. 4E). The NTC joins the spliceosome soon after the tri-snRNP (Hoskins et al. 2011) and appears to be more stably associated after the U4 snRNP is released from the tri-snRNP (Fabrizio et al. 2009). Thus, the result that the NTC is even more sensitive to *prp28-1* is consistent with the allosteric cascade model, which posits that later binding events are dependent on conformational changes induced by earlier binding events (Brow 2002). Similarly, the inhibition of U2 snRNP joining by *prp28-1* may be a consequence of decreased commitment complex 2 formation (see below).

The *prp8-tes* allele partially suppressed all of the cotranscriptional assembly defects caused by *prp28-1* at the nonpermissive temperature. U1 snRNP release was still somewhat delayed in the *prp28-1/prp8-tes* strain, but substantially less so than in the absence of the suppressor mutation (Fig. 4B). A corresponding increase in the rate and magnitude of recruitment of U2 snRNP, U5 snRNP, and NTC was observed at 16°C (Fig. 4C–E). The ability of *prp8-tes* to suppress the U2 association defect suggests that Prp8, a component of

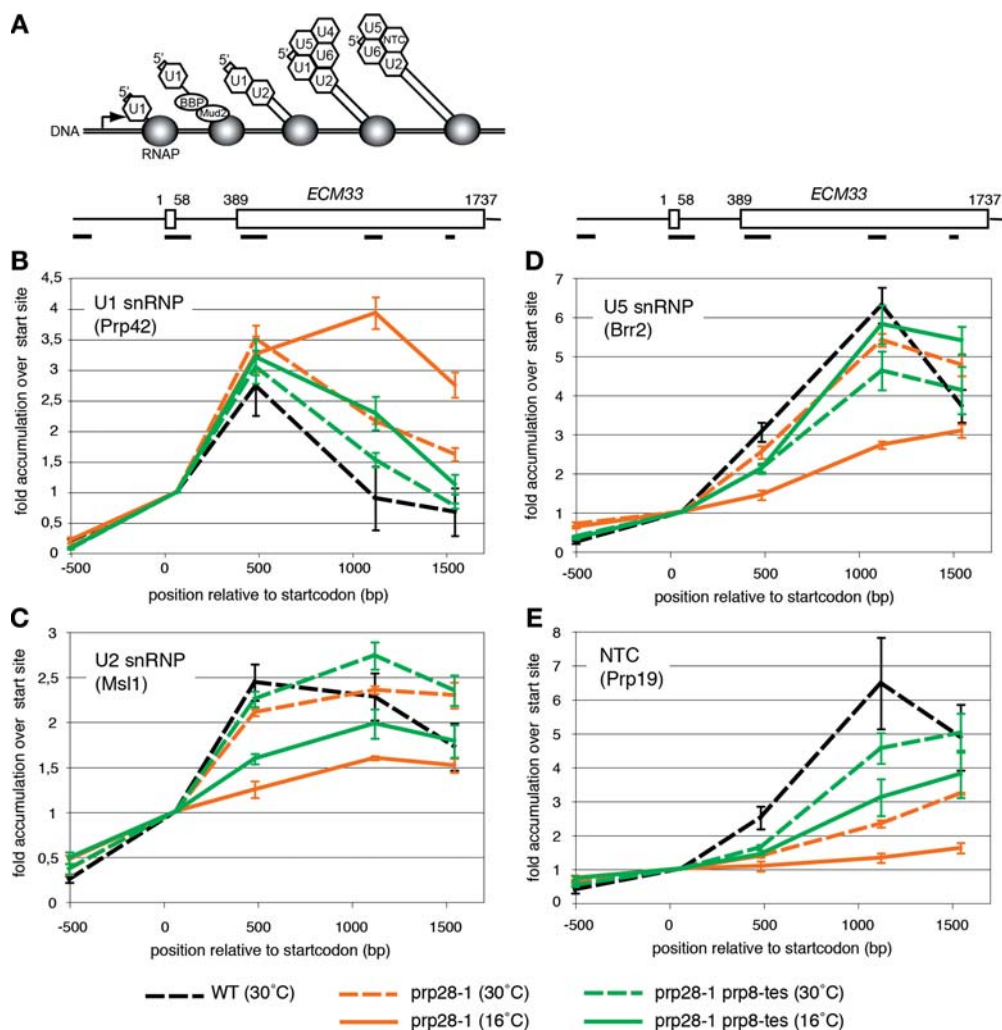


FIGURE 4. The *prp28-1* mutation delays cotranscriptional spliceosome assembly at the *ECM33* gene. (A) Diagram of the expected order of snRNP and NTC occupancy of nascent transcripts along the *ECM33* gene. The short and long open rectangles represent the 5' and 3' exons, respectively. (RNAP) RNA polymerase II. In order, from the transcription start site (bent arrow), the splicing complexes are as follows: CC1, CC2, pre-spliceosome, complete spliceosome, and active spliceosome. (B–E) Data for four HA-tagged splicing factors, which are components of three snRNPs and the NineTeen Complex (NTC), as indicated on each panel. Diagrams at *top* show the positions of qPCR amplicons used for ChIP analysis of the gene *ECM33*. In the panels aligned *below*, data points are placed according to the positions of the PCR products along the gene. Cross-linking of each factor was assessed in strains with wild-type *PRP28* and *PRP8* (black line), *prp28-1* and wild-type *PRP8* (orange lines), and *prp28-1* and *prp8-P263L/W279G* (green lines). Strains were grown at 30°C (dashed lines) or 16°C (solid lines). The data represent the average of at least three independent experiments. Error bars, SEM.

the tri-snRNP, can influence a step of splicing that occurs prior to tri-snRNP incorporation into the spliceosome.

U2 and tri-snRNP association are inhibited by *prp28-1* in vitro, and inhibition is suppressed by *prp8-tes*

To determine whether the effects of *prp28-1* and *prp8-tes* on U2 snRNP and U5 snRNP association also occur in the absence of chromatin and transcription, we monitored the kinetics of spliceosome assembly in vitro at 16°C (Fig. 5). snRNP association with biotinylated pre-mRNA as a function of time was determined by streptavidin affinity purification, followed by RT-qPCR for each snRNA. All five snRNPs asso-

ciated rapidly with pre-mRNA in wild-type splicing reactions, reaching maximum accumulation at 5–15 min. Using a pull-down procedure with low stringency washes to maintain U1 snRNP association, we were able to see that *prp28-1* blocks release of U1 snRNP in vitro (Fig. 5A) as proposed, but not directly observed previously (Staley and Guthrie 1999). At most time points, about twofold more U1 was pulled down from reactions containing *prp28-1* extract than from reactions containing wild-type extract (Fig. 5A). We also confirmed the previous report that U4 is released in wild-type reactions, but retained in *prp28-1* reactions at late time points (Fig. 5B; Staley and Guthrie 1999). Consistent with our in vivo ChIP results, we found that association

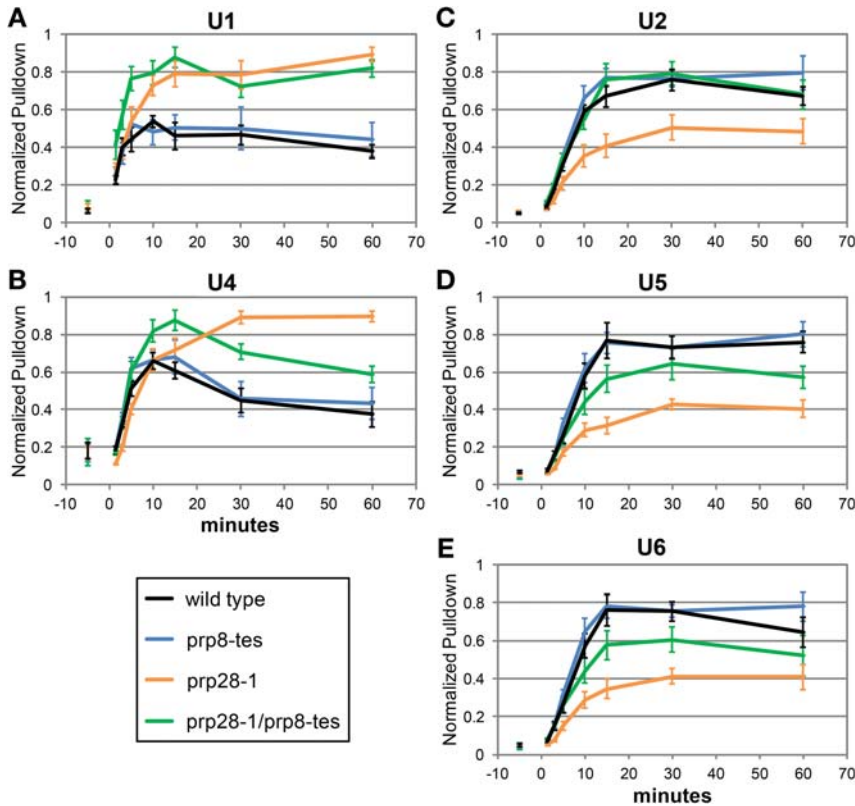


FIGURE 5. In vitro, *prp28-1* blocks U1 and U4 release and reduces stable U2, U5, and U6 association. Splicing extracts made from four strains (wild type: black, *prp28-1*: orange, *prp8-P263L/W279G*: blue, *prp28-1 + prp8-P263L/W279G*: green lines) were incubated with biotinylated actin pre-mRNA at 16°C. Aliquots were taken at the indicated times and snRNAs bound to the pre-mRNA were detected by RT-qPCR. (A–E) Pull-down results for U1, U4, U2, U5, and U6 snRNAs, respectively. Reactions with the four extracts were performed in parallel. The “–5 min” time point represents background (mock pull-down without pre-mRNA). The data were normalized as described in Materials and Methods. Error, SEM after normalization.

of U2, as well as U5 and U6, was inhibited in reactions containing *prp28-1* relative to wild type (Fig. 5C–E). The accumulation of excess U4 RNA in spliceosomes assembled in *prp28-1* extract, despite the decreased level of U5 and U6 RNAs, implies that decreased tri-snRNP recruitment is accompanied by decreased Brr2 activity, and thus increased U4 snRNP retention as observed previously (Staley and Guthrie 1999). Our in vivo and in vitro results suggest that Prp28 has at least two functions: one important for early spliceosome assembly, affecting U2 and subsequent tri-snRNP association with pre-mRNA, and one required for U1 and U4 release after spliceosome assembly.

The *prp8-tes* allele was able to suppress most of the in vitro spliceosome assembly defects (Fig. 5A–E, green). Importantly, the reduction in U2 snRNA pull-down was completely suppressed, which indicates that the functional role of Prp8 early in spliceosome assembly is not specific to cotranscriptional spliceosome assembly. The defects in U5 and U6 binding and U4 release were also partially suppressed. However, U1 release was not improved in the *prp28-1/prp8-tes* double-mutant extract (see Discussion).

prp28-1 inhibits ATP-independent CC2 formation

The surprising in vivo and in vitro defect in U2 snRNP recruitment caused by *prp28-1* could result from inhibition of an even earlier assembly step, so we investigated commitment complex and pre-spliceosome formation directly. Splicing reactions with radiolabeled RP51A pre-mRNA were incubated at nonpermissive temperature (16°C) and run on native gels that resolve CC1 from CC2 (Seraphin and Rosbash 1989, 1991). Wild-type splicing reactions depleted of ATP formed roughly equal amounts of CC1 and CC2, and very little pre-spliceosome or spliceosome (P/SP; these complexes comigrate) (Fig. 6, lane 1). Addition of ATP and incubation for an additional 20 min converted most of the CC2 into P/SP (Fig. 6, lane 5). In contrast, *prp28-1* splicing reactions without ATP accumulated the majority of pre-mRNA in CC1 (Fig. 6, lane 3), and the CC1 band appeared more diffuse. Upon addition of ATP to the *prp28-1* splicing reaction, only about half as much pre-mRNA was converted to P/SP than was with wild-type extract, and most of the pre-mRNA that remained in commitment complexes remained in CC1 (Fig. 6, lane 7). Thus, *prp28-1* not only inhibits U2 snRNP joining but also reduces BBP

and Mud2 association with CC1 (see Fig. 4A). The decrease in U2 joining, seen in this assay and in Figures 4 and 5, is thus likely a downstream result of inefficient CC2 formation.

Splicing is not completely blocked at 16°C by the *prp28-1* mutation (Strauss and Guthrie 1991; data not shown), so we repeated the commitment complex gels with extracts from a strain in which Prp28 expression can be turned off by growth in glucose-containing media (P Raghunathan and C Guthrie, unpubl.). After 3.5–5 h of growth in the presence of glucose, splicing extract made from these cells no longer contained HA-tagged Prp28 detectable by Western blot for HA and failed to splice at 16°C (data not shown). This extract formed even less CC2 (–ATP) and subsequent splicing complexes (+ATP) than *prp28-1* extract (Fig. 6, lanes 9–10). This result confirms that Prp28 contributes to the ATP-independent formation of CC2, and shows that the defect in CC2 formation is not specific to the *prp28-1* allele.

We also assayed the ability of *prp8-tes* to suppress the *prp28-1* defects in CC2 and pre-spliceosome formation (Fig. 6, lanes 4,8). Only partial suppression was observed, but *prp8-tes/prp28-1* extract reproducibly formed CC2 and

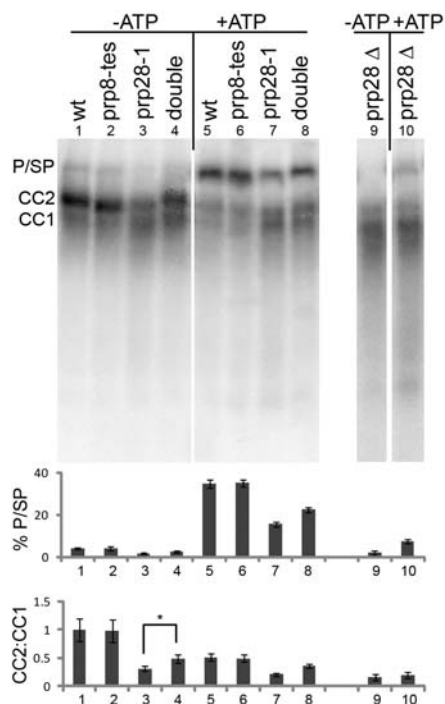


FIGURE 6. *prp28-1* inhibits formation of CC2 and the transition from commitment complex to pre-spliceosome. Complexes assembled on 32 P-labeled RP51A pre-mRNA substrate in the presence or absence of ATP were resolved on a non-denaturing acrylamide/agarose composite gel. (Lanes 1–4,9) no ATP added; (lanes 5–8,10) 2 mM ATP added. (Lanes 1,5) wild-type extract; (lanes 2,6) *prp8-P263L/W279G* extract; (lanes 3,7) *prp28-1* extract; (lanes 4,8) *prp28-1 + prp8-P263L/W279G* extract; (lanes 9,10) extract from cells genetically depleted of Prp28. (CC1) Commitment complex 1; (CC2) commitment complex 2; (P/SP) pre-spliceosome/spliceosome, which comigrate on these gels. Percentage of substrate in P/SP band, relative to the sum of P/SP, CC1, and CC2 bands, and ratio of CC2:CC1 are presented below. Error, SEM. (*) Paired *t*-test conducted, pairing samples run on the same gel; significantly different, $P = 0.00023$.

pre-spliceosome better than *PRP8/prp28-1* extract. Therefore, Prp8, in addition to Prp28, influences this ATP-independent step of spliceosome assembly.

DISCUSSION

A novel, ATP-independent function of Prp28 in CC2 formation

Here we have shown that the DEAD-box ATPase Prp28 has an unanticipated function early in spliceosome assembly. Specifically, we found that Prp28 promotes formation of CC2 prior to the first ATP-dependent step of splicing, since ATP-depleted splicing reactions containing *prp28-1* or depleted of Prp28 accumulated CC1 instead of CC2 (Fig. 6). We also observed *prp28-1*-dependent decreases in U2 and tri-snRNP association in vitro and in vivo, which are likely downstream effects of the failure to efficiently or stably form CC2 (Figs. 4, 5). Previous studies on Prp28 indicated

that it acts during spliceosome activation to promote U1 snRNP release (Staley and Guthrie 1999; Chen et al. 2001; Ismaili et al. 2001), and our current results confirm this conclusion. Therefore, we propose a model in which Prp28 has at least two roles during spliceosome assembly and activation: one ATP-independent role in which Prp28 helps stabilize U1, BBP, and Mud2 on the pre-mRNA during CC2 formation, and a subsequent ATP-dependent role in which Prp28 facilitates the release of U1 following tri-snRNP association (Fig. 7).

Prp28 is an essential gene, but becomes nonessential in the presence of mutations predicted to destabilize the U1/5' ss interaction and facilitate U1 release independent of Prp28 (Chen et al. 2001; Hage et al. 2009; Schwer et al. 2013). Therefore, while Prp28 strongly stimulates CC2 formation on the gene we tested, it is likely not absolutely required for this assembly step. Indeed, cells that have lost *PRP28* in the presence of the *yhc1-1* bypass mutant grow very slowly (Fig. 2B), consistent with the low level of CC2 formation that we observed in extract depleted of Prp28 (Fig. 6).

It is not surprising that a substitution in a conserved helix motif of Prp28 inhibits its ATP-dependent function in U1 snRNP displacement, but why would this substitution affect an ATP-independent role in CC2 formation? Motif Ib, in which the *prp28-1* substitution lies, is implicated in RNA binding (Fairman-Williams et al. 2010), and so may

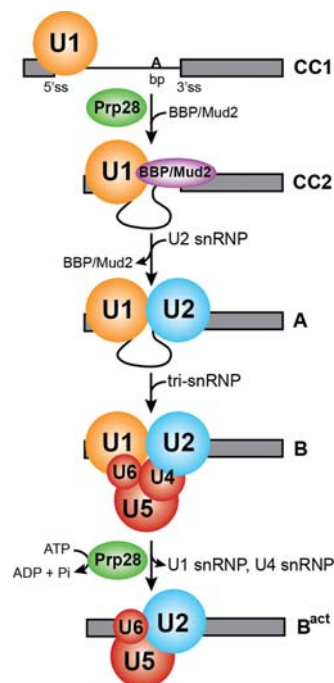


FIGURE 7. Model for the dual functions of Prp28 in spliceosome assembly and activation. Prp28 has an early, ATP-independent function in assembly of commitment complex 2 (CC2) and the pre-spliceosome (A), as well as a later, ATP-dependent function in conversion of the complete spliceosome (B) into the active spliceosome (B^{act}). Only selected splicing factors are shown.

influence the ATP-independent interaction with an RNA substrate.

The DExD/H-box splicing factors Prp5 and Sub2 also have both ATP-dependent and ATP-independent roles in spliceosome assembly. Prp5 uses ATP hydrolysis to promote rearrangements in the U2 snRNP that allow U2 RNA to base-pair with the branchsite, but it first helps recruit U2 to the pre-mRNA in an ATP-independent manner (Perriman et al. 2003; Xu et al. 2004; Kosowski et al. 2009). In addition, extracts depleted of Prp5 are less able to form commitment complexes (Kosowski et al. 2009). Sub2 uses ATP hydrolysis to release BBP and Mud2 during pre-spliceosome formation (Kistler and Guthrie 2001; Libri et al. 2001), but is also required for CC2 formation: Depletion of Sub2 results in spliceosomes that are stalled at CC1 (Zhang and Green 2001). Our finding that Prp28 promotes CC2 formation suggests that Prp28, Sub2, and Prp5 all collaborate to promote formation of CC2, whose components they later destabilize.

Substitutions in the U5 snRNP protein Prp8 promote formation of early spliceosomal complexes

Our finding that substitutions in the N terminus of Prp8, an integral U5 snRNP and tri-snRNP protein, suppress the early assembly defects caused by *prp28-1*, supports the hypothesis that the U5 snRNP functions in the earliest steps of spliceosome assembly, prior to its stable incorporation into the spliceosome. We identified 15 “*prp8-tes*” substitutions spanning a recently proposed bromodomain in the N terminus of Prp8, each of which partially suppressed the cold-sensitive growth defect of a *prp28-1* strain. We tested one of the strongest *prp8-tes* alleles in subsequent assays, and it was able to partially suppress every *prp28-1* splicing defect discussed above, including the CC2 and pre-spliceosome formation defects (Figs. 4–6), except the in vitro U1 snRNP release defect (Fig. 5A). Suppression of the U1 snRNP release defect could be substrate dependent, or could require that additional factors specific to cotranscriptional splicing collaborate with Prp8 and Prp28 to promote U1 release in vivo. The strongest *prp8-tes* allele did not bypass the growth requirement for Prp28, nor did it physically interact differently with two U1 snRNP proteins or suppress the growth defect caused by hyperstabilizing U1/5′ss base-pairing (Fig. 2). Together, our results indicate that *prp8-tes* suppresses *prp28-1* by promoting formation of early splicing complexes, not by destabilizing U1 from the spliceosome.

The finding that substitutions in Prp8 suppress the earliest spliceosome assembly defects caused by *prp28-1* raises the intriguing possibility that both Prp28 and Prp8 act on CC1 in the context of the U5 snRNP. Yeast Prp28 is associated only weakly with U5 snRNP (Gottschalk et al. 1999; Stevens et al. 2001; Small et al. 2006) and our results do not determine whether it promotes CC2 formation as a component of the U5 snRNP or tri-snRNP, or as a free protein. Prp8, however, is an integral component of both the U5 snRNP

and tri-snRNP and is unlikely to exist as free protein, so it presumably acts as part of the U5 snRNP or tri-snRNP when influencing CC2 formation. Whether the U5 snRNP or the tri-snRNP interact with the pre-mRNA early in spliceosome assembly has been a controversial topic in the splicing field. Several prior observations are consistent with an early, transient role for Prp8 and the U5 snRNP. In vitro cross-linking data place the U5 snRNA and Prp8 at the 5′ss at early timepoints in the splicing cycle, both in human (Wasarman and Steitz 1992; Wyatt et al. 1992; Ast and Weiner 1997) and yeast (Newman et al. 1995) cell extracts. Also, affinity-purified human pre-spliceosomes contain U5 snRNP proteins, including Prp28 (Hartmuth et al. 2002). Further characterization of an early interaction between Prp8 and the 5′ss showed that it required intact U4 and U6 RNAs and ATP, but was independent of U2 snRNP binding to the branch point (Maroney et al. 2000). Therefore, tri-snRNP/pre-mRNA interactions apparently occur prior to U2 binding. However, more recent CHIP and single-molecule studies found that in wild-type yeast cells or yeast cell extract, spliceosome assembly usually takes place in a stepwise sequence according to the canonical assembly pathway (Kotovic et al. 2003; Görnemann et al. 2005; Lacadie and Rosbash 2005; Lacadie et al. 2006; Hoskins et al. 2011). These latter studies showed that the spliceosome is unlikely to bind an intron as a pre-formed penta-snRNP, but they do not rule out the transient U5–U1-pre-mRNA interactions that are suggested by our results and others.

While our results implicate the U5 snRNP in early spliceosome assembly events, we cannot rule out the possibility that Prp28 promotes CC2 formation as a free protein and the *prp8-tes* substitutions destabilize Prp28–U5 snRNP interactions, thereby increasing the pool of free Prp28 and compensating for reduced Prp28 function. We found that the N-terminal domain of Prp8 binds Prp28 in vitro, and the *prp8-tes* alleles do not alter the pull-down of Prp28 with Prp8-NTD (Fig. 2D). However, *prp8-tes* alleles could interact differently with Prp28 in the context of full-length Prp8 and the U5 snRNP. Alternatively, the *prp8-tes* mutations may alter interactions in the Aar2–U5 snRNP, which is a precursor of the U5 snRNP that lacks Brr2 and copurifies with the U1 snRNP (Gottschalk et al. 2001; Weber et al. 2011). If the U1 snRNP’s interaction with the Aar2–U5 snRNP competes with stable CC2 formation, weakening this interaction via substitutions in Prp8 could promote CC2 formation.

Prp8 as a master regulator of spliceosome assembly

Prp8 sits at the heart of the spliceosome starting from the time the tri-snRNP binds until after the second step of catalysis, has an extensive set of physical and genetic interactions with other spliceosome components, and has been shown to regulate Brr2 for U4/U6 unwinding (Maeder et al. 2008; Mozaffari-Jovin et al. 2012, 2013). Because of these many interactions, Prp8 is perfectly situated to be a master regulator of

spliceosome assembly, activation, and disassembly. Here, we have shown that substitutions in the N terminus of Prp8 promote CC2 and pre-spliceosome formation in a *prp28-1* background. Thus, Prp8 may also regulate the earliest steps of spliceosome assembly, even before stable tri-snRNP joining. This regulation could be through physical contacts that Prp8 makes with U1 snRNP proteins or the U1 snRNA (Li et al. 2013), although the Prp8/U1 protein interactions tested here were not affected by *prp8-tes* alleles (Fig. 2). Prp8 may also exert its influence on this step by regulating the activity of Prp28; for example, by switching Prp28 between two states: one that promotes a U1 conformation that is favorable for CC2 formation, and one in which Prp28's ATPase activity is activated to promote U1 release after tri-snRNP joining. Additional structural and biochemical studies will be required to discriminate between such possibilities.

MATERIALS AND METHODS

Yeast strains

Yeast strains used in this study are listed in Supplemental Table 1. For strains JG111–JG116 and JG121–JG126, a C-terminal HA₆ tag was added to splicing factors by one-step gene replacement using *K. lactis TRP1* as a selection marker (Knop et al. 1999). Wild-type strains for ChIP are as described (Görnemann et al. 2011). Strains JG110/AMP27 and JG120/AMP28 were generated by replacing YCp50-PRP8 in ANK828 with pRS313-PRP8 (Kuhn and Brow 2000) or pRS313-prp8-P263L/W279G by plasmid shuffle. AMP25 and AMP26 were generated by crossing ANK828 to EJS54 (a gift from E. Strauss: wild-type sister spore of the parent strain of ANK828, EJS51) (Strauss and Guthrie 1991) and selecting for *prp8Δ::ADE2 PRP28* spores, then replacing YCp50-PRP8 with pRS313-PRP8 or pRS313-prp8-P263L/W279G by plasmid shuffle.

Selection for suppressors of *prp28-1* in *PRP8*

We used error-prone PCR and gap repair as described previously (Umen and Guthrie 1996) to introduce mutations into the first quarter of the *PRP8* gene. A region of pRS313-*PRP8* including 233 bp of vector sequence as well as 719 bp upstream of the *PRP8* open reading frame and the first 1980 bp of the open reading frame were amplified with Taq DNA polymerase and cotransformed into ANK828 with pRS313-*PRP8* that was gapped by restriction digest with XhoI and SalI between base pairs (–717) and base pairs 1715 relative to the start codon. Functional *PRP8* alleles were selected by their ability to lose the wild-type copy of *PRP8* on medium containing 5-FOA, then suppressors of *prp28-1* were selected by replica plating to YEPD and incubating at 16°C. Colonies present after 14 d were transferred to fresh medium. Plasmids were isolated from clones that had grown at 16°C within 10 d, their suppression capacity confirmed by retransforming ANK828, and then sequenced from 50 bp upstream of the Prp8 open reading frame to codon 660 to identify mutated residues. Three independent selections were performed. The ApaI/SalI fragments (base pairs –726 to +1715) were subcloned into pRS313-*PRP8* to ensure that suppression was due to mutations within this region. Selected mutations from clones with more than one amino acid substitution were introduced into

pRS313-*PRP8* as single substitutions using the QuikChange technique (Stratagene) and analyzed for their ability to suppress the cold-sensitivity of *prp28-1*. All suppressor alleles were also introduced into ANK800 (wild type), ZRL102 (U4-cs1), and ANK821 (*brr2-1*), and the wild-type allele of *PRP8* was shuffled out to test for genetic interactions. Growth at different temperatures was assayed by spotting 10-fold serial dilutions of OD₆₀₀ = 0.25 cultures on solid YEPD medium, followed by incubation as indicated.

Test for bypass suppression of *prp28-1*

Strain JG140 was constructed by one-step gene disruption of the *PRP28* locus in strain JG120, using the PCR-amplified kanMX6 cassette of pYM1 (Knop et al. 1999) adding 45-bp homologous overhangs to delete base pairs –2 to +1650 of the *PRP28* locus. The PCR product was cotransformed with YCp50-PRP28. Resulting strains were transformed with pRS317-PRP28, pRS317-prp28-D341N, or pRS317-yhc1-1 to create strains JG141, JG142, and JG144. Tenfold serial dilutions of OD₆₀₀ = 0.25 cultures were then plated onto YEPD or YEPD + 5-FOA to select for loss of YCp50-PRP28, and grown at 30°C.

Test for suppression of U1/5's hyperstabilization

A *prp8Δ::LYS2* strain, TB72 (a gift from T. Brenner), was crossed to a *cup1Δ::ura3-52* strain, YS72 (Burgess and Guthrie 1993), and a *prp8Δ, cup1Δ* spore was selected. This strain, AMP24, was transformed with pRS313-PRP8 or pRS313-prp8-P263L/W279G, and with Act1–Cup1 splicing reporters (Lesser and Guthrie 1993) pCG90 (wild type) or pCG91 (5'ss modified to extend U1/5'ss base-pairing to 10 bp) (Staley and Guthrie 1999), generating AMP39–42. Cultures were grown to OD₆₀₀ 0.6, diluted back to OD₆₀₀ 0.1, and fivefold serial dilutions were plated onto SD -Ura medium with varying concentrations of copper and were grown for 6 d at 16°C.

GST pull-down assay to test in vitro protein–protein interactions

Prp28 protein was expressed in *E. coli* BL21(DE3) RIPL (Agilent) from plasmid pRSetA-Prp28 (a gift from Nils Walter), in which Prp28 has a C-terminal His₆ tag. Three liters of bacteria containing the Prp28 plasmid were grown to OD₆₀₀ 0.6 and induced with 1 mM IPTG at 18°C for 15 h. Cells were harvested and frozen at –80°C. Cells were then resuspended in 50 mM NaH₂PO₄ (pH 8.0), 300 mM NaCl, 5 mM β-mercaptoethanol, 10 mM imidazole, and sonicated in five 30-sec bursts at 10% intensity with 30-sec rests. Lysate was clarified for 30 min at 20,000 RPM in a JA25.5 rotor, then the supernatant was passed through a 0.22-μm filter and flowed over a column of Ni-NTA Agarose (Qiagen). Beads were washed with 50 mM NaH₂PO₄, 300 mM NaCl, 5 mM β-mercaptoethanol, and 20 mM imidazole (pH 8), then a second wash with imidazole increased to 36 mM. Prp28 protein was eluted with 50 mM NaH₂PO₄, 300 mM NaCl, 5 mM β-mercaptoethanol, and 250 mM imidazole (pH 8.0), then dialyzed into 20% glycerol, 100 mM KCl, 20 mM HEPES (pH 7.9), 0.2 mM EDTA, 0.5 mM DTT, and stored at –80°C.

Full-length Prp39 and Prp40 were tagged with His₆ by cloning into pET21b (Novagen) at the BamHI and XhoI restriction sites. Prp39 and Prp40 proteins were expressed in *E. coli* Rosetta strain

(Novagen) by growing 100-mL LB cultures to OD₆₀₀ 0.8 and then inducing with 400 μM IPTG for 3 h at 30°C. Cells were harvested, washed once with 1x PBS (pH 7.2), resuspended in 1.5 mL lysis buffer (according to Qiagen protocol for native conditions), and frozen at -20°C. After thawing, lysozyme was added to 1 mg/mL and cells were sonicated in two 15-sec bursts at 25% intensity, with a 30-sec break. His₆-fusion Prp39 and Prp40 were batch purified with Ni-NTA Agarose (Qiagen), following the manufacturer's protocol for native conditions, and stored at 4°C.

Prp8-NTD (1–330) was GST tagged by cloning into pGEX-KG (Guan and Dixon 1991) at the XmaI and XhoI restriction sites. GST-Prp8-NTD was expressed as described for Prp39 and Prp40, except 100 μM IPTG was used for induction and the cell pellet was resuspended in pull-down buffer (20 mM Hepes at pH 7.2, 200 mM potassium acetate, 2 mM magnesium acetate, 0.5% Tween 20, 1 mM DTT). After two clearing spins at 4°C (10 min each at 10,000 rcf), GST-Prp8-NTD lysate was frozen in liquid nitrogen and stored at -80°C for up to 3 mo. For pull-downs, 100 μL of GST fusion protein lysate per sample was added to 25 μL of a 50% slurry of glutathione-Sepharose 4 fast flow (GE Healthcare). For Prp28 pull-downs, 25 μL of GST fusion protein plus 75 μL pull-down buffer was used. GST fusion protein was shown to be in saturating amounts by checking the supernatant by Western blot for residual GST-tagged protein after incubation (data not shown). Following 1 h incubation at 4°C, supernatants were removed, the beads were washed twice with pull-down buffer and used for the pull-down assay. Purified His₆-fusion proteins were added in 350 μL pull-down buffer and incubated for 3 h at 4°C with rotation. Beads were then washed 4× with pull-down buffer (4°C for Prp39 and Prp40, room temperature for Prp28), spun for 1 min at 750 rcf, and boiled with 30 μL 2x SDS sample buffer. Samples were analyzed by Western blot, detecting bound His₆-fusion Prp39 or Prp40 with mouse-anti-His₅ serum (Qiagen), then goat-anti-mouse IgG IR-Dye 680 (LI-COR Biosciences). Bound Prp28 was detected with anti-Prp28 antibody (Strauss and Guthrie 1994), then goat-anti-mouse-HRP secondary antibody (Biorad), and detected by Amersham ECL (GE Healthcare). Equal loading of GST-fusion proteins was checked either by Ponceau stain prior to blocking or detection with goat-anti-GST antiserum (Pharmacia), then donkey anti-goat-HRP (Santa Cruz) and ECL.

Microarray analysis of in vivo splicing defects

Microarray samples were prepared essentially as in Pleiss et al. (2007). Briefly, strains AMP25–28 were grown in YEPD at 30°C to OD₆₀₀ 0.4–0.5, then shifted to 16°C for 50 min. Cells from 15 mL of culture were collected by centrifugation. cDNA was prepared from 50 μg of total cell RNA using random 9-mer primers and labeled with Cy3 or Cy5. Labeled cDNA from each mutant was competitively hybridized to the array with cDNA from wild-type or another mutant as indicated in Figure 3. Arrays were analyzed as in Plocik and Guthrie (2012). The results presented are the average of two biological replicates, each with a dye-flipped technical replicate.

Chromatin immunoprecipitation and quantitative PCR (ChIP-qPCR)

ChIP-qPCR analysis of the genes *ECM33* and *SEC27* was done essentially as described (Görnemann et al. 2005). To cross-link splic-

ing factors to DNA, 4.0 mL 37% formaldehyde solution was added to 150 mL yeast culture, incubated for 15 min at room temperature, then quenched by addition of 8.1 mL 2.5 M glycine. For parallel analyses at permissive and nonpermissive temperatures, cells were grown in 200 mL YPD at 30°C to an OD₆₀₀ of 0.6–0.8. The cultures were divided, 100 mL added to 50 mL 30°C YPD or 4°C YPD, and incubated for 45 min at 30°C or 16°C, respectively, before cross-linking. Cells were harvested by centrifugation at 3500g for 5 min, washed twice with PBS, and then frozen. Cell pellets were thawed on ice, resuspended in 1 mL buffer FA-1 (50 mM HEPES-KOH at pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton-X 100, 0.1% sodium deoxycholate) and lysed on a Vortex Disruptor Genie with glass beads for 40 min at 4°C at maximum speed. Lysates were recovered, including two washes of the glass beads with 1 mL FA-1 each and subjected to sonication on ice to fragment the chromatin to an average of 500 bp. With the 250D Sonifier Ultrasonic Processor Cell Disruptor (Branson) used, this required 6 min of 15-sec on/off cycles at 30% intensity. Lysates were subsequently cleared by centrifugation at 3500g for 5 min. Supernatants were cleared prior to immunoprecipitation by addition of 200 μL of a 50% slurry of Sepharose CL-4B beads (Sigma-Aldrich), previously washed with FA-1 buffer, and rotation at 4°C for 1 h. Eight micrograms of 12CA5 antibody was then added to 700 μL of lysate, followed by rotation at 4°C for 2 h, then 50 μL of washed GammaBind G beads (GE Healthcare) were added for 1 h. Mouse IgG (Sigma) served as a negative control. Beads were washed three times with buffer FA-1, once each with buffer FA-2 (FA-1 with 500 mM NaCl) and FA-3 (20 mM Tris-Cl at pH 8.0, 250 mM LiCl, 0.5% NP-40, 0.5% sodium deoxycholate, 1 mM EDTA), and twice with TE buffer (pH 8.0), transferring the beads to a fresh tube. For elution and uncross-linking, 250 μL TE/1% SDS was added, followed by incubation at 65°C overnight. In parallel, 20 μL of lysate in 200 μL TE/1% SDS was uncross-linked to serve as input control. Eluates were transferred to a fresh tube, incubated with 10 μL Proteinase K (20 mg/mL) at 55°C for 2 h, and purified with a PCR purification kit according to instructions (Qiagen). The data in Figure 4 represent the average of three independent experiments, except: Prp42 (all strains): *n* = 4; *prp28-1*—Brr2 at 16°C and 30°C: *n* = 6; *prp28-1* + *prp8- P263L/W279G*—Brr2 at 16°C and 30°C: *n* = 4; Msl1, Brr2, Prp19 in WT: *n* = 4.

Primers used for ChIP-qPCR

ECM-UP-F: GCAGTATCATCCTTCACGACCC; ECM-UP-R: GCG TCTTCCCGTTTTGTC; ECM9-31: CAAGAACGCTTTGACTGC TACTG; ECM145-123: GAAGAGGACCACGAATCTACTCG; ECM 430-451: ACTTCTGCCACTGCTACTGCTC; ECM562-539: AGG AACCATCAATCTCTTGGATAC; ECM1073-1097: TTGGTCAAT CTTTGTCTATCGTCTC; ECM1173-1150: TGTGTTGTTAGCAA TGATGAAACC; ECM1531-1555: TCTAAGAAGTCTAAGGGTGC TGCTC; ECM1582-1561: TGAATGAAGTGGCTGGAACAAG; SE C280-304: GGGTTTTGACTACCTTGACTCTGG; SEC370-349: GGAGTTTCCGTAACCTTGGATGG; SEC931-950: TTTCTGGTT CCGAAGATGGC; SEC1043-1022: TGTGGATGGGTAGCGATA CAC; SEC1309-1330: TTGTTACAGTTGTTGGGGATGG; SEC 1395-1372: CAAAGTCTTGACATTTACCGAAGG; SEC1969-1992: GAGAGGTCCATGTTTATGGTTACG; SEC2057-2037: AATGGC TTCTTCAATTTCCCC; SEC2599-2622: CTGTATCAGAAAGAG TTTGTGGGG; SEC2692-2669: GCTGGAGTGGAAATCTAAGTC AATG.

In vitro splicing extract preparation

A total of 2–3-L cultures of strains AMP25–28 were harvested at OD₆₀₀ 1.7–1.9, and splicing extract was prepared as described (Umen and Guthrie 1995), except that frozen cell pellets were ground with a ball mill (Retch MM 301 mixer-mill; 3 × 3 min at 11 Hz and 2 × 3 min at 12 Hz) and dialyzed twice in 2 L buffer D. Prp28-depleted extract was made from strain PR88 (a gift from P. Raghunathan) in which genomic *PRP28* is deleted and complemented by a plasmid with the *GAL10* promoter controlling a copy of *PRP28* with a 3x HA tag inserted in the *ClaI* site near the start codon of *PRP28*. The strain was grown to an OD₆₀₀ of 0.5–0.7 in YEP-galactose, then filtered and shifted to YEPD for 3.5–5 h. Cells were collected and splicing extract prepared as above. Extracts were checked by Western blot with anti-HA antibody to confirm that Prp28 was no longer detectable.

Biotinylated pre-mRNA pull-down experiments

Biotinylated pre-mRNA pull-down experiments were adapted from previous protocols (Ruby et al. 1990; Kuhn et al. 1999; Staley and Guthrie 1999; Brenner and Guthrie 2006). ActinΔ6 pre-mRNA (Vijayraghavan et al. 1986) was synthesized with Megascript T7 RNA polymerase (Ambion) using biotin-11-UTP (Ambion) as 5% of the UTP. Seven pull-down time course experiments were conducted using four different extract preparations from each of the four strains (AMP25–28); 120–140 μL splicing reactions were prepared; 1.9 μL was taken for a “0 min input” sample, and in three experiments, a 19-μL “no pre-mRNA” sample was removed to 50 μL stop buffer for background pull-down measurement (shown as “–5 min” time point). Reactions were started by adding 2.5 fmol pre-mRNA/μL and ATP to 2 mM final concentration, and were incubated at 16°C. Twenty microliter aliquots were removed at indicated times, added to 50 μL ice cold stop buffer (20 mM EDTA, 40% Buffer D, 60 mM KPO₄ at pH 7, 3% PEG 8000), and incubated on ice until the end of the time course. A 2-μL “input” sample was taken at 30 min. A total of 50 μL of streptavidin-agarose beads (Thermo Scientific) (50% slurry in stop buffer, after being blocked in stop buffer plus 10 μg/mL each glycogen, tRNA, and bovine serum albumin) were added to each reaction except inputs, and incubated at 4°C, rocking for 1 h. For most experiments, 100 μL wash buffer (150 mM NaCl, 60 mM HEPES at pH 7.6, 3 mM MgCl₂, 15% glycerol, 0.05% NP40, 0.5 mM DTT) was added before supernatant was removed. Beads were then washed 3× with 500 μL wash buffer. snRNAs and pre-mRNA were eluted by adding 50 μL formamide +40 mM EDTA and incubating at 90°C for ~10 min followed by washing beads with 50 or 100 μL TE +1% SDS. In two experiments, RNA was instead eluted from beads with Proteinase K as in Kuhn et al. (1999). Input samples were added to 5 μL stop buffer, and were incubated on ice until the elution step, at which point they were treated the same as pull-down samples. RNA was then phenol-chloroform extracted, ethanol precipitated, and resuspended in 20 μL H₂O. Eight microliters of RNA were reverse transcribed with primers specific to the snRNAs as described in Brenner and Guthrie (2006) and detected with qPCR as described, except the standard curve was generated from serial dilutions of a pool of input samples. qPCR values used are the average of 2–3 qPCR technical replicates.

qPCR values from each individual time course were normalized to the average of the respective 0 and 30-min input samples. All of

the values in a single experiment (time courses with all four extracts done in parallel) were further normalized to the maximum value for that experiment, which allowed all experiments to be averaged together. In three of the experiments, 1- and 2-min time points were taken instead of 1.5 and 3 min. In order to include data from all repeats, 1.5- and 3-min values were interpolated, assuming a linear trend between 1 and 5 min. Similarly, in one experiment the 30-min value was interpolated from 25- and 50-min time points, and the 50-min measurement was used as an approximation of the 60-min point. These interpolated values were included in data for Figure 2, but leaving them out did not significantly change the results. Including interpolated values, $n = 7$ for 1.5, 3, 5, 10, 15, and 30 min, and $n = 6$ for 60 min. Exceptions: wt 30 min: $n = 6$; *prp8-tes* 30 min: $n = 6$, *prp28-1* 5, 10, and 30 min: $n = 6$, *prp28-1/prp8-tes* 1.5 min: $n = 6$, background (no pre-mRNA added “–5 min”): $n = 3$. Error, SEM of final values after all described normalization.

Native gel analysis

Commitment complex gels were based on a published protocol (Seraphin and Rosbash 1989). Twenty-two-microliter standard splicing reactions (Lin et al. 1985) containing 8.8 μL of extract were depleted of ATP by adding glucose to 2 mM and incubating at 25°C for 20 min. A total of 1–2 μL (~15 fmol or ~30,000 cpm) radiolabeled RP51AΔ2 pre-mRNA (Seraphin and Rosbash 1991) was added to each reaction, and reactions were incubated for 20 min at 16°C. Ten microliters was removed to 10 μL ice cold buffer R (2 mM MgOAc, 50 mM Hepes (pH 7.5), 1 mg/mL tRNA, 50-fold molar excess cold Ubc4 or Actin pre-mRNA). Then, ATP (to 2 mM) was added to the remaining reaction volume, and reactions were incubated for another 20 min at 16°C before the final 10 μL was removed to 10 μL ice cold buffer R. Five microliters of loading dye (2.5x TBE, 50% glycerol, bromophenol blue, and xylene cyanol dyes) was added, and the reaction was loaded on a 0.5x TBE, 3% acrylamide, 0.5% agarose, 5% glycerol gel, with a plug of 10% acrylamide gel at the bottom ~3 cm (1x TBE = 89 mM Tris-borate, 2 mM EDTA). The 26-cm gel was run in 0.5x TBE at 120 V for 20–24 h at 4°C. Bands were detected by PhosphorImager (Molecular Dynamics) and quantified with ImageQuant v.5.2 (Molecular Dynamics). Error bars in Figure 6 show SEM from $n = 5$ (–ATP: *prp28Δ*), $n = 6$ (+ATP: *prp28Δ*), $n = 8$ (–ATP: wt, *prp28-1*, *prp28-1/prp8-tes*), $n = 9$ (–ATP: *prp8-tes*), $n = 11$ (+ATP: *prp28-1/prp8-tes*), or $n = 12$ (+ATP: wt, *prp8-tes*, *prp28-1*) replicates. To determine whether the CC2:CC1 ratio was different between *prp28-1* and *prp28-1/prp8-tes*, we conducted a paired *t*-test, pairing reactions that were prepared together and run on the same gel.

DATA DEPOSITION

Microarray data generated in this study are available for download from the Gene Expression Omnibus (GEO), under accession no. GSE42754.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

ACKNOWLEDGMENTS

We thank Karla Neugebauer for supporting the ChIP experiments, Thomas Hoffmann for data analysis advice, Matthew Kahlscheuer and Nils Walter for the Prp28 expression plasmid, members of the Brow and Guthrie labs for helpful suggestions, and Sam Butcher, Tien-Hsien Chang, Charles Query, and Joan Steitz for comments on the manuscript. This study was supported by an NSF GRFP fellowship to A.M.P., a postdoctoral fellowship from the Deutsche Forschungsgemeinschaft to J.G., and NIH grants to D.A.B. and C.G. C.G. is an American Cancer Society Research Professor of Molecular Genetics. This work is dedicated to the memory of Stephanie W. Ruby, who did pioneering studies on the role of U1 snRNP in spliceosome assembly.

Received August 14, 2013; accepted October 17, 2013.

REFERENCES

- Abovich N, Rosbash M. 1997. Cross-intron bridging interactions in the yeast commitment complex are conserved in mammals. *Cell* **89**: 403–412.
- Ast G, Weiner AM. 1997. A novel U1/U5 interaction indicates proximity between U1 and U5 snRNAs during an early step of mRNA splicing. *RNA* **3**: 371–381.
- Brenner TJ, Guthrie C. 2006. Assembly of Snu114 into U5 snRNP requires Prp8 and a functional GTPase domain. *RNA* **12**: 862–871.
- Brow DA. 2002. Allosteric cascade of spliceosome activation. *Annu Rev Genet* **36**: 333–360.
- Burgess SM, Guthrie C. 1993. A mechanism to enhance mRNA splicing fidelity: The RNA-dependent ATPase Prp16 governs usage of a discard pathway for aberrant lariat intermediates. *Cell* **73**: 1377–1391.
- Chen JY-F, Stands L, Staley JP, Jackups RR Jr, Latus LJ, Chang T-H. 2001. Specific alterations of U1-C protein or U1 small nuclear RNA can eliminate the requirement of Prp28p, an essential DEAD box splicing factor. *Mol Cell* **7**: 227–232.
- Đlakić M, Mushegian A. 2011. Prp8, the pivotal protein of the spliceosomal catalytic center, evolved from a retroelement-encoded reverse transcriptase. *RNA* **17**: 799–808.
- Fabrizio P, Dannenberg J, Dube P, Kastner B, Stark H, Urlaub H, Lührmann R. 2009. The evolutionarily conserved core design of the catalytic activation step of the yeast spliceosome. *Mol Cell* **36**: 593–608.
- Fairman-Williams ME, Guenther U-P, Jankowsky E. 2010. SF1 and SF2 helicases: Family matters. *Curr Opin Struct Biol* **20**: 313–324.
- Filippakopoulos P, Knapp S. 2012. The bromodomain interaction module. *FEBS Lett* **586**: 2692–2704.
- Galej WP, Oubridge C, Newman AJ, Nagai K. 2013. Crystal structure of Prp8 reveals active site cavity of the spliceosome. *Nature* **493**: 638–643.
- Görnemann J, Kotovic KM, Hujer K, Neugebauer KM. 2005. Cotranscriptional spliceosome assembly occurs in a stepwise fashion and requires the cap binding complex. *Mol Cell* **19**: 53–63.
- Görnemann J, Barrandon C, Hujer K, Rutz B, Rigaut G, Kotovic KM, Faux C, Neugebauer KM, Séraphin B. 2011. Cotranscriptional spliceosome assembly and splicing are independent of the Prp40p WW domain. *RNA* **17**: 2119–2129.
- Gottschalk A, Neubauer G, Banroques J, Mann M, Lührmann R, Fabrizio P. 1999. Identification by mass spectrometry and functional analysis of novel proteins of the yeast [U4/U6·U5] tri-snRNP. *EMBO J* **18**: 4535–4548.
- Gottschalk A, Kastner B, Lührmann R, Fabrizio P. 2001. The yeast U5 snRNP coisolated with the U1 snRNP has an unexpected protein composition and includes the splicing factor Aar2p. *RNA* **7**: 1554–1565.
- Grainger RJ, Beggs JD. 2005. Prp8 protein: At the heart of the spliceosome. *RNA* **11**: 533–557.
- Guan KL, Dixon JE. 1991. Eukaryotic proteins expressed in *Escherichia coli*: An improved thrombin cleavage and purification procedure of fusion proteins with glutathione S-transferase. *Anal Biochem* **192**: 262–267.
- Hage R, Tung L, Du H, Stands L, Rosbash M, Chang T-H. 2009. A targeted bypass screen identifies Ynl187p, Prp42p, Snu71p, and Cbp80p for stable U1 snRNP/Pre-mRNA interaction. *Mol Cell Biol* **29**: 3941–3952.
- Hartmuth K, Urlaub H, Vornlocher H-P, Will CL, Gentzel M, Wilm M, Lührmann R. 2002. Protein composition of human prespliceosomes isolated by a tobramycin affinity-selection method. *Proc Natl Acad Sci* **99**: 16719–16724.
- Hoskins AA, Friedman LJ, Gallagher SS, Crawford DJ, Anderson EG, Wombacher R, Ramirez N, Cornish VW, Gelles J, Moore MJ. 2011. Ordered and dynamic assembly of single spliceosomes. *Science* **331**: 1289–1295.
- Ismaili N, Sha M, Gustafson EH, Konarska MM. 2001. The 100-kDa U5 snRNP protein (hPrp28p) contacts the 5' splice site through its ATPase site. *RNA* **7**: 182–193.
- Kistler AL, Guthrie C. 2001. Deletion of *MUD2*, the yeast homolog of U2AF65, can bypass the requirement for Sub2, an essential spliceosomal ATPase. *Genes Dev* **15**: 42–49.
- Knop M, Siegers K, Pereira G, Zachariae W, Winsor B, Nasmyth K, Schiebel E. 1999. Epitope tagging of yeast genes using a PCR-based strategy: More tags and improved practical routines. *Yeast* **15**: 963–972.
- Kosowski TR, Keys HR, Quan TK, Ruby SW. 2009. DEXD/H-box Prp5 protein is in the spliceosome during most of the splicing cycle. *RNA* **15**: 1345–1362.
- Kotovic KM, Lockshon D, Boric L, Neugebauer KM. 2003. Cotranscriptional recruitment of the U1 snRNP to intron-containing genes in yeast. *Mol Cell Biol* **23**: 5768–5779.
- Kuhn AN, Brow DA. 2000. Suppressors of a cold-sensitive mutation in yeast U4 RNA define five domains in the splicing factor Prp8 that influence spliceosome activation. *Genetics* **155**: 1667–1682.
- Kuhn AN, Li Z, Brow DA. 1999. Splicing factor Prp8 governs U4/U6 RNA unwinding during activation of the spliceosome. *Mol Cell* **3**: 65–75.
- Kuhn AN, Reichl EM, Brow DA. 2002. Distinct domains of splicing factor Prp8 mediate different aspects of spliceosome activation. *Proc Natl Acad Sci* **99**: 9145–9149.
- Lacadie SA, Rosbash M. 2005. Cotranscriptional spliceosome assembly dynamics and the role of U1 snRNA:5' ss base pairing in yeast. *Mol Cell* **19**: 65–75.
- Lacadie SA, Tardiff DF, Kadener S, Rosbash M. 2006. In vivo commitment to yeast cotranscriptional splicing is sensitive to transcription elongation mutants. *Genes Dev* **20**: 2055–2066.
- Lesser CF, Guthrie C. 1993. Mutational analysis of pre-mRNA splicing in *Saccharomyces cerevisiae* using a sensitive new reporter gene, *CUP1*. *Genetics* **133**: 851–863.
- Li X, Zhang W, Xu T, Ramsey J, Zhang L, Hill R, Hansen KC, Hesselberth JR, Zhao R. 2013. Comprehensive *in vivo* RNA-binding site analyses reveal a role of Prp8 in spliceosomal assembly. *Nucleic Acids Res* **41**: 3805–3818.
- Libri D, Graziani N, Saguez C, Boulay J. 2001. Multiple roles for the yeast *SUB2/yUAP56* gene in splicing. *Genes Dev* **15**: 36–41.
- Lin RJ, Newman AJ, Cheng SC, Abelson J. 1985. Yeast mRNA splicing *in vitro*. *J Biol Chem* **260**: 14780–14792.
- Maeder C, Kutach AK, Guthrie C. 2008. ATP-dependent unwinding of U4/U6 snRNAs by the Brr2 helicase requires the C terminus of Prp8. *Nat Struct Mol Biol* **16**: 42–48.
- Maroney PA, Romfo CM, Nilsen TW. 2000. Functional recognition of the 5' splice site by U4/U6.U5 tri-snRNP defines a novel ATP-dependent step in early spliceosome assembly. *Mol Cell* **6**: 317–328.
- Mathew R, Hartmuth K, Möhlmann S, Urlaub H, Ficner R, Lührmann R. 2008. Phosphorylation of human PRP28 by SRPK2 is required

- for integration of the U4/U6-U5 tri-snRNP into the spliceosome. *Nat Struct Mol Biol* **15**: 435–443.
- Mozaffari-Jovin S, Santos KF, Hsiao H-H, Will CL, Urlaub H, Wahl MC, Lührmann R. 2012. The Prp8 RNase H-like domain inhibits Brr2-mediated U4/U6 snRNA unwinding by blocking Brr2 loading onto the U4 snRNA. *Genes Dev* **26**: 2422–2434.
- Mozaffari-Jovin S, Wandersleben T, Santos KF, Will CL, Lührmann R, Wahl MC. 2013. Inhibition of RNA helicase Brr2 by the C-terminal tail of the spliceosomal protein Prp8. *Science* **341**: 80–84.
- Newman AJ, Teigelkamp S, Beggs JD. 1995. snRNA interactions at 5' and 3' splice sites monitored by photoactivated crosslinking in yeast spliceosomes. *RNA* **1**: 968–980.
- Pena V, Liu S, Bujnicki JM, Lührmann R, Wahl MC. 2007. Structure of a multipartite protein-protein interaction domain in splicing factor Prp8 and its link to retinitis pigmentosa. *Mol Cell* **25**: 615–624.
- Pena V, Rozov A, Fabrizio P, Lührmann R, Wahl MC. 2008. Structure and function of an RNase H domain at the heart of the spliceosome. *EMBO J* **27**: 2929–2940.
- Perriman R, Barta I, Voeltz GK, Abelson J, Ares M. 2003. ATP requirement for Prp5p function is determined by Cus2p and the structure of U2 small nuclear RNA. *Proc Natl Acad Sci* **100**: 13857–13862.
- Pleiss JA, Whitworth GB, Bergkessel M, Guthrie C. 2007. Transcript specificity in yeast pre-mRNA splicing revealed by mutations in core spliceosomal components. *PLoS Biol* **5**: e90.
- Plocik AM, Guthrie C. 2012. Diverse forms of RPS9 splicing are part of an evolving autoregulatory circuit. *PLoS Genet* **8**: e1002620.
- Ragunathan PL, Guthrie C. 1998. RNA unwinding in U4/U6 snRNPs requires ATP hydrolysis and the DEIH-box splicing factor Brr2. *Curr Biol* **8**: 847–855.
- Ritchie DB, Schellenberg MJ, Gesner EM, Raithatha SA, Stuart DT, MacMillan AM. 2008. Structural elucidation of a PRP8 core domain from the heart of the spliceosome. *Nat Struct Mol Biol* **15**: 1199–1205.
- Ruby SW, Goelz SE, Hostomsky Z, Abelson JN. 1990. Affinity chromatography with biotinylated RNAs. *Methods Enzymol* **181**: 97–121.
- Schellenberg MJ, Wu T, Ritchie DB, Fica S, Staley JP, Atta KA, LaPointe P, MacMillan AM. 2013. A conformational switch in PRP8 mediates metal ion coordination that promotes pre-mRNA exon ligation. *Nat Struct Mol Biol* **20**: 728–734.
- Schwer B, Chang J, Shuman S. 2013. Structure-function analysis of the 5' end of yeast U1 snRNA highlights genetic interactions with the Msl5•Mud2 branchpoint-binding complex and other spliceosome assembly factors. *Nucleic Acids Res* **41**: 7485–7500.
- Seraphin B, Rosbash M. 1989. Identification of functional U1 snRNA-pre-mRNA complexes committed to spliceosome assembly and splicing. *Cell* **59**: 349–358.
- Seraphin B, Rosbash M. 1991. The yeast branchpoint sequence is not required for the formation of a stable U1 snRNA-pre-mRNA complex and is recognized in the absence of U2 snRNA. *EMBO J* **10**: 1209–1216.
- Small EC, Leggett SR, Winans AA, Staley JP. 2006. The EF-G-like GTPase Snu114p regulates spliceosome dynamics mediated by Brr2p, a DExD/H box ATPase. *Mol Cell* **23**: 389–399.
- Staley JP, Guthrie C. 1999. An RNA switch at the 5' splice site requires ATP and the DEAD box protein Prp28p. *Mol Cell* **3**: 55–64.
- Stevens SW, Barta I, Ge HY, Moore RE, Young MK, Lee TD, Abelson J. 2001. Biochemical and genetic analyses of the U5, U6, and U4/U6-U5 small nuclear ribonucleoproteins from *Saccharomyces cerevisiae*. *RNA* **7**: 1543–1553.
- Strauss EJ, Guthrie C. 1991. A cold-sensitive mRNA splicing mutant is a member of the RNA helicase gene family. *Genes Dev* **5**: 629–641.
- Strauss EJ, Guthrie C. 1994. PRP28, a “DEAD-box” protein, is required for the first step of mRNA splicing in vitro. *Nucleic Acids Res* **22**: 3187–3193.
- Teigelkamp S, Mundt C, Achsel T, Will CL, Lührmann R. 1997. The human U5 snRNP-specific 100-kD protein is an RS domain-containing, putative RNA helicase with significant homology to the yeast splicing factor Prp28p. *RNA* **3**: 1313–1326.
- Umen JG, Guthrie C. 1995. A novel role for a U5 snRNP protein in 3' splice site selection. *Genes Dev* **9**: 855–868.
- Umen JG, Guthrie C. 1996. Mutagenesis of the yeast gene *PRP8* reveals domains governing the specificity and fidelity of 3' splice site selection. *Genetics* **143**: 723–739.
- van Nues RW, Beggs JD. 2001. Functional contacts with a range of splicing proteins suggest a central role for Brr2p in the dynamic control of the order of events in spliceosomes of *Saccharomyces cerevisiae*. *Genetics* **157**: 1451–1467.
- Vijayraghavan U, Parker R, Tamm J, Iimura Y, Rossi J, Abelson J, Guthrie C. 1986. Mutations in conserved intron sequences affect multiple steps in the yeast splicing pathway, particularly assembly of the spliceosome. *EMBO J* **5**: 1683–1695.
- Wassarman DA, Steitz JA. 1992. Interactions of small nuclear RNA's with precursor messenger RNA during in vitro splicing. *Science* **257**: 1918–1925.
- Weber G, Cristão VF, De L, Alves F, Santos KF, Holton N, Rappsilber J, Beggs JD, Wahl MC. 2011. Mechanism for Aar2p function as a U5 snRNP assembly factor. *Genes Dev* **25**: 1601–1612.
- Will CL, Lührmann R. 2011. Spliceosome structure and function. *Cold Spring Harb Perspect Biol* **3**: a003707.
- Wyatt JR, Sontheimer EJ, Steitz JA. 1992. Site-specific cross-linking of mammalian U5 snRNP to the 5' splice site before the first step of pre-mRNA splicing. *Genes Dev* **6**: 2542–2553.
- Xu Y-Z, Newnham CM, Kameoka S, Huang T, Konarska MM, Query CC. 2004. Prp5 bridges U1 and U2 snRNPs and enables stable U2 snRNP association with intron RNA. *EMBO J* **23**: 376–385.
- Yang K, Zhang L, Xu T, Heroux A, Zhao R. 2008. Crystal structure of the B-finger domain of Prp8 reveals analogy to ribosomal proteins. *Proc Natl Acad Sci* **105**: 13817–13822.
- Yang F, Wang X-Y, Zhang Z-M, Pu J, Fan Y-J, Zhou J, Query CC, Xu Y-Z. 2013. Splicing proofreading at 5' splice sites by ATPase Prp28p. *Nucleic Acids Res* **41**: 4660–4670.
- Zhang M, Green MR. 2001. Identification and characterization of yUAP/Sub2p, a yeast homolog of the essential human pre-mRNA splicing factor hUAP56. *Genes Dev* **15**: 30–35.