Spliceosome disassembly catalyzed by Prp43 and its associated components Ntr1 and Ntr2

Rong-Tzong Tsai,1,4 Ru-Huei Fu,1,4 Fu-Lung Yeh,1,4 Chi-Kang Tseng,1,3 Yu-Chieh Lin,1,2 Yu-hsin Huang,1 and Soo-Chen Cheng1,2,3,5

1Institute of Molecular Biology, and 2Taiwan International Graduate Program, Molecular and Cellular Biology, Academia Sinica, Nankang, Taipei, Taiwan 115, Republic of China; 3Institute of Microbiology and Immunology, National Yang-Ming University, Shih-Pai, Taipei, Taiwan 112, Republic of China

Two novel yeast splicing factors required for spliceosome disassembly have been identified. Ntr1 and Ntr2 (NineTeen complex-Related proteins) were identified for their weak association with components of the Prp19-associated complex. Unlike other Prp19-associated components, these two proteins were primarily associated with the intron-containing spliceosome during the splicing reaction. Extracts depleted of Ntr1 or Ntr2 exhibited full splicing activity, but accumulated large amounts of lariat-intron in the spliceosome after splicing, indicating that the normal function of the Prp19-associated complex in spliceosome activation was not affected, but spliceosome disassembly was hindered. Immunoprecipitation analysis revealed that Ntr1 and Ntr2 formed a stable complex with DExD/H-box RNA helicase Prp43 in the splicing extract. Ntr1 interacted with Prp43 through the N-terminal G-patch domain, with Ntr2 through a middle region, and with itself through the carboxyl half of the protein. The affinity-purified Ntr1–Ntr2–Prp43 complex could catalyze disassembly of the spliceosome in an ATP-dependent manner, separating U2, U5, U6, NTC (NineTeen Complex), and lariat-intron. This is the first demonstration of physical disassembly of the spliceosome, catalyzed by a complex containing a DExD/H-box RNA helicase and two accessory factors, which might function in targeting the helicase to the correct substrate.

[Keywords: Spliceosome disassembly; NTR complex; Prp43; Ntr1; Ntr2]

Received September 21, 2005; revised version accepted October 28, 2005.

Introns are removed from precursor messenger RNA (pre-mRNA) via a two-step transesterification reaction catalyzed by a large ribonucleoprotein particle called the spliceosome (Brody and Abelson 1985; Frendewey and Keller 1985; Grabowski et al. 1985). The spliceosome consists of U1, U2, U4/U6, and U5 small nuclear ribonucleoprotein particles (snRNPs) as well as non-snRNP protein factors that bind to the pre-mRNA in a sequential manner (for reviews, see Burge et al. 1999; Brow 2002). Following binding of all four snRNPs, the spliceosome undergoes a large structural rearrangement, releasing U1 and U4 and forming new base pairings between U2 and U6 and between U6 and pre-mRNA. This leads to the activation of the spliceosome, on which catalytic reactions can take place.

During spliceosome activation, a protein complex as-
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2001, 2002), encoded by genes not essential for cellular growth. These proteins are associated with the spliceosome at the same time during spliceosome assembly in a step after the dissociation of U4, suggesting that they might function as an integral complex. Interestingly, all of these proteins were also found in the yeast penta-snRNP complex, which is proposed to be a functional particle of the preassembled spliceosome capable of splicing added pre-mRNA (Stevens et al. 2002).

Homologs of several NTC components, Prp19, Ntc90, Ntc85, Ntc77, Ntc30, and Ntc25, are also found in the 35S complex identified in mammalian splicing extracts by gradient sedimentation (Makarov et al. 2002). The 35S complex contains U5 snRNP with an incomplete set of protein components and a subset of proteins found in the 45S activated spliceosome. It has been proposed that this complex might represent a disassembly intermediate generated from the activated spliceosome post-splicing reaction (Makarov et al. 2002).

Disassembly of the spliceosome after completion of the splicing reaction is necessary for recycling of splicing factors to promote efficient splicing. The molecular mechanism of the disassembly has not been extensively studied. The yeast Prp22 protein, a DExD/H-box RNA helicase, is required for the dissociation of the mature message from the spliceosome (Company et al. 1991), but also plays a role in the second catalytic step (Schwer and Gross 1998). Another protein of the DExD/H-box family Prp43 has been shown to be involved in spliceosome disassembly in the release of lariat-intron (Arenas and Abelson 1997; Martin et al. 2002). Both Prp22 and Prp43 confer RNA-dependent ATPase activities, which are required for the release of mRNA and lariat-intron, respectively (Schwer and Gross 1998; Martin et al. 2002). Nevertheless, the mechanistic details of the disassembly process remain unclear.

The availability of genome sequences has facilitated global studies of gene functions. Several yeast proteomic databases provide valuable information for analyzing constituents of complex assemblies involved in various biological pathways (Zhou et al. 2002; Hazbun et al. 2003; Krogan et al. 2004). Two yeast ORFs, YLR424w and YKR022c, of unknown function are found to associate with proteins that include five NTC components and other known splicing factors (Hazbun et al. 2003). On testing whether these two gene products were components of NTC, we found that these two proteins, being only weakly associated with NTC, were not required for either the assembly of the spliceosome or catalytic reactions, but were required for disassembly of the spliceosome after the splicing reaction is complete. Immunoprecipitation analysis revealed that these two proteins formed a stable complex with Prp43. The affinity-purified complex could catalyze disassembly of the spliceosome in an ATP-dependent manner, separating U2, U5, U6, NTC, and lariat-intron. Our results not only demonstrated physical disassembly of the spliceosome, but also showed essential function of Ntr1 and Ntr2 as accessory factors of Prp43 in catalyzing spliceosome disassembly.

Results

Coprecipitation of Ntr1 and Ntr2 with NTC components

Three uncharacterized ORFs were suggested to be splicing factors in a recent study of yeast proteomes (Hazbun et al. 2003). Two of these proteins, YLR424w and YKR022c, were purified together with five identified NTC components as well as several other splicing factors. To verify whether these two proteins are components of NTC, YLR424w and YKR022c were tagged with the HA-epitope for immunoprecipitation with the anti-HA antibody, and proteins coprecipitated with Prp19 and with YLR424w or YKR022c were compared. Extracts prepared from PRP19-HA, YLR424w-HA, YKR022c-HA, and nontagged strains were precipitated with the anti-HA antibody followed by Western blotting as shown in Figure 1A. Due to differences in the accessibility of the HA-epitope on each protein, different amounts of the antibody were used for precipitation of different extracts as indicated in the legend to Figure 1. Precipitated proteins were first probed with the anti-HA antibody to reveal Prp19-HA, YLR424w-HA, and YKR022c-HA. The amounts of precipitated materials loaded on the gel were adjusted accordingly to contain approximately the same amount of HA-tagged proteins in each lane. Association of NTC components with Prp19-HA, YLR424w-HA, and YKR022c-HA was compared by Western blotting probed with antibodies against NTC components. Figure 1A shows that all the identified NTC components were coprecipitated with YLR424w and YKR022c, but at much lower levels than those coprecipitated with Prp19, indicating that both YLR424w and YKR022c were either weakly associated with NTC or only a small fraction was associated with NTC. YLR424w and YKR022c were unlikely true NTC components, and were therefore named Ntr1 and Ntr2, respectively, for NTC-related proteins. The protein sequences of Ntr1 and Ntr2 are shown in Figure 1B.

Ntr1 and Ntr2 are associated with the intron-containing spliceosome

Eight proteins have previously been identified to be components of NTC (Chen et al. 1998, 2001, 2002; Tsai et al. 1999). All of these proteins bind to the spliceosome at the same time during spliceosome assembly in a step after dissociation of U4 but prior to catalytic reactions. To determine whether Ntr1 and Ntr2 bind to the spliceosome in a similar way to NTC components, the spliceosomes formed in HA-tagged Ntr1 or Ntr2 extracts at various ATP concentrations were immunoprecipitated with anti-HA and anti-Ntc20 antibodies as shown in Figure 2. In Ntr1-HA extracts, although no splicing products were detected at 0.1 mM ATP [Fig. 2A, lane 1], a small amount of pre-mRNA was precipitated by the anti-Ntc20 antibody [Fig. 2A, lane 2], indicating formation of a small amount of the active spliceosome under this condition. Nevertheless, the anti-HA antibody did not precipitate any pre-mRNA. At 0.5 or 2 mM ATP, large
amounts of pre-mRNA, splicing intermediates, and products were precipitated by the anti-Ntc20 antibody (Fig. 2A, lanes 6,10), but only the lariat-intron was precipitated by the anti-HA antibody (Fig. 2A, lanes 8,12). Similar results were obtained with Ntr2-HA extracts (Fig. 2B), except that a trace amount of splicing activity was detected in this extract at 0.1 mM of ATP. These results suggest that Ntr1 and Ntr2 behave differently from NTC in their binding to the spliceosome during spliceosome assembly in that they are present primarily on the intron-containing spliceosome.

To rule out the possibility that failure in coprecipitating pre-mRNA or splicing intermediates with Ntr1 or Ntr2 was due to inaccessibility of the HA-epitope to the antibody during early steps of spliceosome assembly, proteins present on the total spliceosome and the active spliceosome were examined by precipitation with streptavidin agarose of spliceosomes formed on biotinylated wild-type and Ac/Cla actin pre-mRNA. The Ac/Cla pre-mRNA is actin-sequence-truncated beyond the branch point and thus, although unable to undergo catalytic reactions, allows formation of the active spliceosome and binding of NTC (Cheng 1994). As shown in Figure 2C, Ntr1 and Ntr2 were found on the spliceosome formed with wild-type pre-mRNA [lane 2], but only present in trace amounts with Ac/Cla pre-mRNA [lane 4], in contrast to NTC components, Prp19, Ntc85, and Ntc30, which were effectively associated with both pre-mRNA. It is speculated that binding of low levels of Ntr1 and Ntr2 to Ac/Cla pre-mRNA might be through their association with NTC. Nevertheless, this result is consistent with that of the immunoprecipitation experiment, in which Ntr1 or Ntr2 co-precipitated predominantly lariat-intron, and suggests that these two proteins become associated with the spliceosome primarily during or after catalytic steps.

Ntr1 and Ntr2 are required for the release of lariat-intron from the spliceosome

To examine the function of Ntr1 and Ntr2 in the splicing reaction, Ntr1 and Ntr2 were individually depleted from Ntr1-HA or Ntr2-HA extracts, with the anti-HA antibody for splicing assays. As shown in Figure 3A, while depletion of Prp19 abolished the splicing activity (lane 2), depletion of Ntr1 or Ntr2 resulted in accumulation of lariat-intron after splicing (lanes 4,6). This indicates that neither Ntr1 nor Ntr2 was essential for the splicing reaction, but both proteins were required either for disassembly of the spliceosome in release of either mature mRNA or lariat-intron, or required for the degradation of lariat-intron RNA.

To see whether Ntr1 or Ntr2 was required for the release of mature mRNA from the spliceosome, spliceosome was prepared by precipitation with the anti-HA antibody for splicing assays. As shown in Figure 3B, depletion of Prp19 abolished the splicing activity [lane 2], depletion of Ntr1 or Ntr2 resulted in accumulation of lariat-intron RNA. This suggests that neither Ntr1 nor Ntr2 was essential for the splicing reaction, but both proteins were required either for disassembly of the spliceosome in release of either mature mRNA or lariat-intron, or required for the degradation of lariat-intron RNA.

Accumulation of pre-mRNA and lariat-intron in Ntr2-depleted cells

Tetrad analysis had revealed that both NTR1 and NTR2 are essential for cell viability [data not shown]. To see
whether this reflected their essentiality in splicing in vivo, yeast strains were constructed such that NTR1 or NTR2 was placed under the control of a GAL-promoter for their metabolic depletion. In glucose-based media, the GAL-NTR1 strain grew normally for unknown reasons (data not shown), but the growth of GAL-NTR2 was suppressed on prolonged incubation, as shown in Figure 4A. RNA was isolated from GAL-NTR2 cells grown in glucose for 0 or 28 h, and subject to primer extension analysis using an 5' end-labeled primer, R13, complementary to a region in the second exon of the U3 gene. Figure 4B shows that two extension products representing pre-mRNA of U3A and U3B accumulated at 28 h (lane 5), but not at 0 h (lane 4) or in the wild-type control

Figure 2. Ntr1 and Ntr2 associate with lariat-intron during the splicing reaction. (A, B) The splicing reaction was performed at 2 mM, 0.5 mM, and 0.1 mM of ATP in Ntr1-HA (A) and Ntr2-HA (B) extracts, respectively, and reaction mixtures were subjected to immunoprecipitation with anti-HA and anti-Ntc20 antibodies. (C) The splicing reaction was carried out with biotinylated actin or Ac/Cla pre-mRNA and the spliceosome isolated by precipitation with streptavidin Sepharose. Components of the precipitated spliceosome were analyzed by Western blotting using antibodies against Ntr1, Ntr2, Prp19, Ntc85, and Ntc30.

Figure 3. Failure of lariat-intron release in Ntr1- or Ntr2-depleted extracts. (A) Splicing was carried out in Prp19-, Ntr1-, or Ntr2-depleted extracts, depleted from Prp19-HA, Ntr1-HA, and Ntr2-HA extracts, respectively, with the anti-HA antibody. (B) Splicing reactions carried out in mock-depleted (top panel) or Ntr1-depleted (bottom panel) extracts were subjected to sedimentation on 10%–30% glycerol gradients. RNA was extracted from fractions collected from each gradient and analyzed on 8% acrylamide/8 M urea gels.

Figure 4A. RNA was isolated from GAL-NTR2 cells grown in glucose for 0 or 28 h, and subject to primer extension analysis using an 5' end-labeled primer, R13, complementary to a region in the second exon of the U3 gene. Figure 4B shows that two extension products representing pre-mRNA of U3A and U3B accumulated at 28 h [lane 5], but not at 0 h [lane 4] or in the wild-type control
Similar extension products were also seen in prp2 and prp22 temperature-sensitive mutants after they were grown at restrictive temperatures for 2 h (Fig. 4B, lanes 2,3). This indicates that in vivo depletion of Ntr2 gave rise to a defect in pre-mRNA splicing. Considering the role of Ntr2 in spliceosome disassembly, depletion of Ntr2 should cause accumulation of lariat-intron, which cannot be detected by extension with R13. A second primer R14, located between the branch point and the 5'/H11032 splice site of the U3A intron, was used for primer extension to detect the lariat-intron. As shown in Figure 4C, one strong extension stop at the 5' end of pre-U3A was seen in prp2 [lane 2] and prp22 [lane 3] mutants and in GAL-NTR2 cells grown in glucose medium for 12 and 24 h [lanes 7,8], but was barely detectable in wild type or in GAL-NTR2 grown in galactose medium. This is consistent with the result using primer R13 except that only U3A was detected with primer R14. A minor extension product, representing pre-U3A terminated at the 5'/H11032 splice site, was also seen in prp22 [Fig. 4C, lane 1] or prp2 [Fig. 4C, lane 2]. This indicates that in vivo intron is accumulated when Ntr2 or Prp22 is not functional, corroborating a post-catalytic role for Ntr2 in pre-mRNA splicing.

**Association of Ntr1, Ntr2, and Prp43 in a functional complex**

Since in vitro depletion of either Ntr1 or Ntr2 resulted in accumulation of lariat-intron during the splicing reaction, these two proteins are either independently required for or coordinately function to promote disassembly of the spliceosome. To determine whether Ntr1 and Ntr2 are associated as a functional unit, Ntr1 was affinity-purified from Ntr1-HA extracts with the anti-HA antibody conjugated to protein A-Sepharose and used for complementation of Ntr1- or Ntr2-depleted extracts. As shown in Figure 5A, affinity-purified Ntr1 did indeed complement the intron-accumulation phenotype of both Ntr1- and Ntr2-depleted extracts [lanes 4,8], suggesting functional association of Ntr1 and Ntr2.

The yeast DExD/H-box family of RNA helicase Prp43 has previously been implicated in spliceosome disassembly [Arenas and Abelson 1997]. Prp43 was shown to confer RNA-dependent ATPase activity and to be required for release of the lariat-intron from the spliceosome after completion of the splicing reaction [Martin et al. 2002]. Therefore, Prp43 could potentially work cooperatively with Ntr1 and Ntr2 in mediating disassembly of the spliceosome. To test whether Prp43 is functionally associated with Ntr1 and Ntr2, Prp43 was tagged with the V5-epitope and depleted from Prp43-V5 extracts using the anti-V5 antibody. Accumulation of lariat-intron was seen in Prp43-depleted extracts (Fig. 5A, lane 11), but was complemented by affinity-purified Ntr1 (Fig. 5A, lane 12). This suggests that Prp43 is likely associated with Ntr1 and Ntr2 in a functional complex.

Physical association of Ntr1, Ntr2, and Prp43 was confirmed by immunoprecipitation analysis using extracts prepared from double-tagged strains PRP43-V5/NTR1-HA and PRP43-V5/NTR2-HA. Splicing extracts from these strains were precipitated with anti-HA or anti-V5 antibodies followed by Western blotting and probed with...
polyclonal anti-Ntr1 and anti-Ntr2 antibodies and anti-V5 antibody for Prp43-V5. As shown in Figure 5B, precipitation with the anti-V5 antibody for Prp43 coprecipitated Ntr1 and Ntr2 [lanes 1,3], or anti-HA [lanes 2,4] antibody followed by Western blotting probed with anti-Ntr1, anti-Ntr2, anti-V5, and anti-Prp19 antibodies. [Lane 5] Prp19-HA extracts were also precipitated with the anti-HA antibody as a control. [C] Western blotting of Ntr1-, Ntr2-, and Prp43-depleted extracts was probed with anti-Ntr1, anti-Ntr2, anti-Prp19 polyclonal, and anti-V5 monoclonal antibodies. Ntr1 was depleted from Prp43-V5/Ntr1-HA extracts with the anti-HA antibody [lanes 1,2], Ntr2 depleted from Prp43-V5/Ntr2-HA extracts with the anti-HA antibody [lanes 3,4], and Prp43 depleted from Prp43-V5/Ntr1-HA extracts with the anti-V5 antibody [lanes 5,6], respectively. [D] The affinity-purified NTR complex was subjected to sedimentation on 10%–30% glycerol gradients, and proteins from each fraction were analyzed by Western blotting probed for Ntr1, Ntr2, and Prp43. The molecular weight markers are indicated on the top.

Figure 5. Association of Ntr1, Ntr2, and Prp43 in a functional complex. (A) Ntr1-HA, Ntr2-HA, and Prp43-V5 extracts were depleted of Ntr1, or Ntr2, or Prp43 with anti-HA or anti-V5 antibody, respectively, and used for splicing and complementation assays with purified Ntr1 complex. [B] Prp43-V5/Ntr1-HA [lanes 1,2] and Prp43-V5/Ntr2-HA [lanes 3,4] extracts were immunoprecipitated with anti-V5 [lanes 1,3] or anti-HA [lanes 2,4] antibody followed by Western blotting probed with anti-Ntr1, anti-Ntr2, anti-V5, and anti-Prp19 antibodies. [Lane 5] Prp19-HA extracts were also precipitated with the anti-HA antibody as a control. [C] Western blotting of Ntr1-, Ntr2-, and Prp43-depleted extracts was probed with anti-Ntr1, anti-Ntr2, anti-Prp19 polyclonal, and anti-V5 monoclonal antibodies. Ntr1 was depleted from Prp43-V5/Ntr1-HA extracts with the anti-HA antibody [lanes 1,2], Ntr2 depleted from Prp43-V5/Ntr2-HA extracts with the anti-HA antibody [lanes 3,4], and Prp43 depleted from Prp43-V5/Ntr1-HA extracts with the anti-V5 antibody [lanes 5,6], respectively. [D] The affinity-purified NTR complex was subjected to sedimentation on 10%–30% glycerol gradients, and proteins from each fraction were analyzed by Western blotting probed for Ntr1, Ntr2, and Prp43. The molecular weight markers are indicated on the top.
cosedimented primarily in fractions 12–15, corresponding to a size of 150–200 kDa, consistent with the calculated size of the trimeric complex. The minor form of Ntr1–Ntr2 dimer is presumed to copurify with the trimer, but was not clearly distinguished.

**Interactions between Ntr1, Ntr2, and Prp43**

Large-scale genomic studies have revealed physical interactions between Ntr1 and Ntr2 and between Ntr1 and Prp43 (Gavin et al. 2002; Hazbun et al. 2003). Genome-wide two-hybrid analyses have also identified interactions between Ntr1 and Ntr2 (Ito et al. 2001; Hazbun et al. 2003). To confirm and further characterize interactions between these three proteins in the complex, interactions between each pair of proteins were analyzed by two-hybrid assays. Ntr1, Ntr2, and Prp43 were individually fused to the LexA-DNA-binding domain and GAL4-activation domain, and interactions were assayed by measuring β-galactosidase activity. Figure 6A shows that Ntr1 interacted with Ntr2, Prp43, and itself, whereas Ntr2 and Prp43 interacted only with Ntr1. Thus, formation of the NTR complex might be mediated through interactions of Ntr1 with Ntr2 and Prp43.

A recent study has found that the DExD/H-box RNA helicase Prp2, which is required for the first catalytic reaction, binds to the spliceosome through an interaction between the G-patch domain containing protein Spp2 (Roy et al. 1995; Silverman et al. 2004). Such interaction requires the G-patch domain of Spp2 and a region at the C-terminal end of the Prp2 sequence (Silverman et al. 2004). It has been proposed that Spp2 serves as a Prp2 accessory factor for targeting Prp2 to the spliceosome (Silverman et al. 2004). Interestingly, Ntr1 also contains a G-patch domain near its N terminus (Fig. 1). We therefore examined whether the G-patch domain or other regions of Ntr1 was responsible for the interaction between Ntr1 and Prp43 by two-hybrid assays. A series of deletion mutants spanning the entire Ntr1 sequence were generated, and the position of the deleted region in each deletion mutant is shown in Figure 6B. Interactions of these deletion mutants fused to the LexA-DNA-binding domain, with Prp43, Ntr2, and Ntr1 fused to the GAL4-activation domain, were assayed by measuring β-galactosidase activity. As shown in Figure 6C, deletion of the G-patch domain alone, or in combination with the upstream ΔNG or downstream region ΔA from Ntr1 abolished the interaction between Ntr1 and Prp43, suggesting the requirement of the G-patch domain for Ntr1’s interaction with Prp43. Moreover, a G-patch domain containing an N-terminal fragment with amino acid residues 1–122 (NG) was sufficient for interactions between Ntr1 and Prp43. The G-patch domain was not required for the interaction between Ntr1 and Ntr2 or for Ntr1 self-interaction, since neither ΔG nor

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**Figure 6.** Two-hybrid analysis of interactions between Ntr1, Ntr2, and Prp43. (A) Ntr1, Ntr2, and Prp43 were fused to the LexA-DNA-binding domain and Ntr1 and Prp43 fused to the GAL4-activation domain, and interactions were assayed by measuring β-galactosidase activity. (B) The map of Ntr1 and the positions of deletion mutants constructed for two-hybrid assays. Fragment NG, shown in the solid box, contains amino acid residues 1–122. Regions of deletion, shown in the open boxes, are ΔNG, 1–109; ΔG, 63–109; ΔA, 62–295; ΔB, 122–263; ΔC, 123–344; ΔD, 345–427; ΔE, 428–581; ΔF, 649–708. Regions of Ntr1 interact with Prp43; Ntr2 and Ntr1 self-interaction are indicated on the bottom. (C) Deletion mutants of Ntr1 were fused to the LexA-DNA-binding domain and their interactions with Prp43, Ntr1, and Ntr2, fused to the GAL4-activation domain, were assayed by measuring β-galactosidase activity. Far left panel (AD) shows the background generated from each deletion mutant when fused to the DNA-binding domain.
ΔNG had any effect on their interactions. Further analysis of deletion mutants revealed that while ΔC completely abolished interactions between Ntr1 and Ntr2, ΔD moderately and ΔB greatly weakened the interaction. The interaction in the ΔA mutant appeared to be at a level similar to the control of the activation domain (Fig. 6C, far left panel). These results suggest that the maximal region required for interactions between Ntr1 and Ntr2 was between amino acid residues 123 and 427. In contrast, self-interaction of Ntr1 involved the carboxyl half of the protein since ΔC, ΔD, and ΔE completely abolished the interaction, but ΔB had no effect. The interaction in ΔF was also weakened in view of the high basal level in this mutant.

**NTC is associated with the Ntr1- and Ntr2-containing spliceosome**

Although the lariat-intron coprecipitated with Ntr1 and Ntr2 during the splicing reaction, the amount precipitated was much lower than that coprecipitated with Ntc20 (Fig. 2). This could reflect a differential efficiency of precipitation by different antibodies or that Ntr1 and Ntr2 might be only transiently associated with the intron-containing spliceosome either while NTC was still on the spliceosome or after NTC was dissociated. To test whether Ntr1 and Ntr2 coexist with NTC on the spliceosome, we performed a double immunoprecipitation experiment. The spliceosome formed in Ntr1-HA extracts was immunoprecipitated with anti-HA antibody and eluted with the HA-peptide. The eluted spliceosome was then immunoprecipitated with an anti-Ntc20 antibody to examine the presence of NTC. As shown in Figure 7, the spliceosome coprecipitated with Ntr1 predominantly contained lariat-introns [lane 3] that, upon elution from beads with the HA-peptide, could be precipitated by the anti-Ntc20 antibody [lane 4], indicating that NTC was still present on the spliceosome when Ntr1 joined. This also suggested the possibility of isolating NTC-containing spliceosome formed in Ntr1-depleted extracts for functional assay of NTR complex.

**Disassembly of the spliceosome mediated by NTR complex**

To assay the function of NTR complex, the splicing reaction was carried out in Ntr1-depleted extracts and the intron-accumulated spliceosome isolated by precipitation with the anti-Ntc20 antibody as shown in Figure 8A [lane 2]. When the affinity-purified NTR complex and ATP were added to the precipitated spliceosome, ~70% of the lariat-intron was released from beads following incubation [Fig. 8A, lanes 9, 10]. Incubation with NTR complex or ATP alone failed to release lariat-intron [Fig. 8A, lanes 5–8]. Hydrolysis of ATP was required for lariat-intron release as revealed by the low efficiency of intron release with γS-ATP [Fig. 8A, lanes 11, 12]. This suggests that NTR complex promotes dissociation of NTC from spliced lariat-intron RNA in an ATP-dependent manner. To see whether NTR complex also promoted dissociation of snRNPs from lariat-intron, a similar experiment was performed except that the intron-containing spliceosome was precipitated with the anti-Smd1 antibody. As shown in Figure 8B, ~50% of the lariat-intron precipitated by the anti-Smd1 antibody was released from beads upon incubation with ATP and NTR complex [lanes 5, 6], indicating dissociation of the intron from snRNPs. Together, these results show that NTR complex catalyzed the dissociation of lariat-intron from NTC as well as from snRNPs, possibly as free RNA. To confirm that lariat-intron was released as free RNA, the dissociated fraction was treated with proteinase K and subjected to gradient sedimentation, as shown in Figure 8C. Gradient I shows the total splicing reaction performed in Ntr1-depleted extracts; the spliceosome sedimented primarily in fractions 5–7 and mature mRNA in fractions 11 and 12 [see also Fig. 3B]. Gradient II shows that the dissociated lariat-intron sedimented near the top of the gradient in a position [fractions 13 and 14] lighter than mature mRNA. Nearly an identical profile was seen when this fraction was deproteinized prior to sedimentation, as shown in gradient III, indicating that the lariat-intron had been released from the beads as naked RNA.

To see whether snRNPs were separated from each other accompanying intron dissociation, fractions released from anti-Ntc20 antibody precipitated spliceosome were subject to sedimentation on glycerol gradient, and RNA was extracted from each fraction for Northern blotting probed with five snRNAs. As shown in Figure 9, released U2, U5, and U6 sedimented at different positions [top panel], indicating separation of these three snRNPs. For comparison, total splicing extracts [Fig. 9, bottom panel] and deproteinized extracts [Fig. 9, middle panel] were also subject to sedimentation to reveal the positions of individual snRNPs and naked snRNAs. In-
Interestingly, while the released U5 sedimented like that in the extract, U2 was dissociated as a lighter particle than that in the extract, but not as naked RNA. This suggests that some components of U2 snRNP might have been dissociated during spliceosome assembly or disassembly or U2 might be dissociated in an altered conformation. Furthermore, U6 was released as naked RNA, consistent with our previous observation that Lsm proteins were dissociated from U6 during spliceosome activation (Chan et al. 2003). Altogether, these results demonstrate that NTR complex was functional in catalyzing disassembly of the spliceosome, resulting in total separation of U2, U5, U6, NTC, and lariat-intron.

Discussion

In this study, we have identified two novel splicing factors involved in disassembly of the spliceosome. Ntr1 and Ntr2 were identified through their association with NTC components. Nevertheless, only a small fraction of Ntr1 and Ntr2 was associated with NTC since depletion of NTC from the splicing extract did not affect the amount of Ntr1 or Ntr2 to a large extent (data not shown), and vice versa, only a small fraction of NTC components was associated with Ntr1 and Ntr2, as depletion of Ntr1 or Ntr2 did not affect the function of NTC in spliceosome disassembly. It is not clear why small amounts of Ntr1, Ntr2, and NTC are associated with each other. Since nuclease treatment of splicing extracts did not abolish such association (data not shown), the interaction was unlikely to be mediated through binding of RNA, excluding the possibility of their being the endogenous spliceosome. The functional significance of such association remains unknown.

Both NTR1 and NTR2 are essential for cellular growth. Metabolic depletion of Ntr2 resulted in accumulation of pre-mRNA and lariat-intron, indicating requirement of Ntr2 for pre-mRNA splicing in vivo. Although the in vivo role of Ntr1 in splicing was not demonstrated, it is likely that Ntr1 is also essential for pre-mRNA splicing in vivo, considering that Ntr1 was tightly associated with Ntr2 and the two proteins functioned in a coordinative manner. Primer extension analysis revealed that despite the NTR complex not being required for splicing catalysis or prior steps of spliceosome assembly, pre-mRNA accumulated at much higher levels than lariat-intron upon depletion of Ntr2. It is conceivable that block of spliceosome disassembly due to a faulty NTR complex will result in accumulation of the spliceosome and failure in recycling splicosomal components. As a consequence, a large part of pre-mRNA cannot be spliced and accumulates in the cell. This can be demonstrated in vitro using a splicing recycling assay described previously (Supplementary Fig. S1; Raghunathan and Guthrie 1998). Consistently, cells carrying mutations in the PRP22 gene, defective in release of mature mRNA, also accumulated much more pre-mRNA than lariat-intron.

We have demonstrated that Ntr1 and Ntr2 were associated with Prp43 to form a trimeric complex. While Ntr1 and Ntr2 were always associated with each other,
Figure 9. Dissociation of snRNPs catalyzed by the NTR complex. The released fraction as in Figure 8A (top), total extracts (bottom), and deproteinized extracts (middle), were subjected to sedimentation on 10%–30% glycerol gradient, and RNA was extracted from each fraction for Northern blotting, probed with U1, U2, U4, U5, and U6. The positions of U5, U6, U4/U6 di-snRNP, and U4/U6.U5 tri-snRNPs are marked.

Prp43 appeared to exist in great excess, and a large fraction of Prp43 was not associated with either Ntr1 or Ntr2. Furthermore, ~30% of Ntr1 and Ntr2 were not associated with Prp43. Thus, these three proteins existed in at least three different forms, uncomplexed Prp43, Ntr1–Ntr2 heterodimer, and Ntr1–Ntr2–Prp43 heterotrimer. The Ntr1–Ntr2–Prp43 trimeric complex was the functional form that mediated spliceosome disassembly. Uncomplexed Prp43 did not function by itself since depletion of Ntr1 or Ntr2 without changing the level of Prp43 to any great extent resulted in the accumulation of lariat-intron in the splicing reaction. Depletion of Prp43 from splicing extracts also led to intron accumulation despite a fraction of Ntr1 and Ntr2 being still present, suggesting that the Ntr1–Ntr2 dimeric complex was not functional without Prp43.

Prp43 belongs to the family of DExD/H-box RNA helicases, which play important roles in pre-mRNA splicing throughout the whole splicing process. Eight DExD/H-box RNA helicases are involved in different steps of the splicing reaction [Staley and Guthrie 1998]. All DExD/H-box family proteins confer RNA-dependent ATPase activity, and some have also demonstrated RNA unwinding activity [Laggerbauer et al. 1998]. These proteins use the energy gained from ATP hydrolysis to unwind RNA duplexes or to drive conformational changes in RNA molecules or ribonucleoprotein complexes. Nevertheless, DExD/H-box RNA helicases show very little substrate specificity for unwinding of RNA duplexes in vitro. It is speculated that these helicases might require extrinsic factors for targeting to specific substrates to execute their normal functions. Factors that interact with RNA helicases and possibly play roles in regulating their functions have been documented [Silverman et al. 2003]. The pre-mRNA splicing factor Spp2, initially identified as a high-copy suppressor of the prp2-1 temperature-sensitive mutant [Last et al. 1987], has been shown to be required for the activity of Prp2 in promoting step one reaction [Silverman et al. 2004]. Interactions of Prp2 and Spp2 can be demonstrated by twohybrid and GST pull-down assays [Roy et al. 1995; Silverman et al. 2004]. Spp2 also contains a G-patch motif important for its interaction with Prp2 [Silverman et al. 2004]. It has been proposed that the interaction of Prp2 with Spp2 is essential for the function of Prp2, and that Spp2 might act as a cofactor of Prp2 in determining its spliceosome specificity [Silverman et al. 2004]. Similarly, Prp43 interacts with Ntr1 through the G-patch domain, and the association of Prp43 with Ntr1 and Ntr2 is required for spliceosome disassembly. Ntr1 and Ntr2 together might function as coordinators to target Prp43 to the spliceosome to mediate spliceosome disassembly. Whether the G-patch motif represents a general DExD/H RNA helicase-interacting motif for directing the helicase to specific substrate remains to be seen.

Both NTR1 and NTR2 are essential for vegetative yeast growth and at least Ntr2 is essential for efficient splicing in vivo. Nevertheless, proteins homologous to either Ntr1 or Ntr2 have not been identified in other organisms despite the fact that Prp43 is evolutionarily conserved [Fouraux et al. 2002]. It is possible that the metazoan Prp43 is evolved to carry extra domains for targeting to its substrate, substituting the function of trans-acting regulators. Indeed, the metazoan Prp43 has a highly charged N-terminal domain not present in yeast or Caenorhabditis elegans. Determining whether this N-terminal region is functional for interaction with other spliceosomal components awaits further study.

Although Prp43 required Ntr1 and Ntr2 for its targeting to the spliceosome, only a small fraction of Prp43 was associated with Ntr1 and Ntr2. It is not clear whether the remaining protein exists in a free form or is associated with other macromolecules. The human ortholog of Prp43, DDX15 or hPrp43, has been demonstrated to associate with human L1 autoantigen and accumulate in the nucleoli of Hep-2 cells [Fouraux et al. 2002], suggesting a possible function of hPrp43 other than splicing. Association of hPrp43 with U1 and U2 snRNAs was also shown by immunoprecipitation with antibodies against hPrp43 [Fouraux et al. 2002]. Furthermore, hPrp43 was found associated with the affinity-pu-
rified 17S U2 snRNP (Will et al. 2002). These results suggest that hPrp43 might also function at the early stage of spliceosome assembly or even in facilitating the assembly of U1 or U2 snRNP. Nevertheless, association of yeast Prp43 with U1 or U2 was not detected by immunoprecipitation analysis (data not shown). Neither were early steps of the splicing reaction affected in the absence of Prp43. This suggests that the yeast Prp43 could be deficient in these extra functions that hPrp43 might have.

The fact that Ntr1 and Ntr2 coprecipitated a small amount of lariat-intron from the splicing reaction suggests that these two proteins might be only transiently associated with the spliceosome after completion of the splicing reaction. It is conceivable that the spliceosome rapidly undergoes disassembly upon binding of NTR complex and, consequently, this results in the detection of only a low level of Ntr1–Ntr2-associated spliceosome. In contrast, in normal splicing reactions, Prp43 was not detected to stably associate with the spliceosome in any amount by immunoprecipitation analysis using either anti-V5 antibody for Prp43-V5 or a polyclonal antibody against Prp43 [data not shown], suggesting that Prp43 did not act concordantly with Ntr1 and Ntr2 in binding to the spliceosome. It is possible that Prp43, while binding to the spliceosome as a component of NTR complex, might be immediately dissociated from the spliceosome upon hydrolysis of ATP, triggering disintegration of the spliceosome that still contains Ntr1 and Ntr2. The detected Ntr1- and Ntr2-containing splicing complex might represent a disassembly intermediate, which is either on its way to disassembly or defective for further disassembly. In this case, following disassembly of the spliceosome, Ntr1 and Ntr2 would need to reassociate with Prp43 to form the functional complex for subsequent rounds of the reaction. Alternatively, Prp43 might be associated with Ntr1 and Ntr2 in a dynamic manner. Ntr1–Ntr2 dimeric complex could be recruited to the spliceosome and form a relatively more stable complex, as detected by immunoprecipitation of Ntr1 or Ntr2. The Ntr1–Ntr2-bound spliceosome may further recruit spliceosome and form a relatively more stable complex, Ntr2 dimeric complex could be recruited to the Ntr1–be associated with Ntr1 and Ntr2 in a dynamic manner.

Materials and methods

**Yeast strains**

The following yeast strains were used: BJ2168, Mata prc1 prb1 pep4 leu2 trp1 ura3; YSCC1, Mata prc1 prb1 pep4 leu2 trp1 ura3 PRP19HA; YSCC131, Mata prc1 prb1 pep4 leu2 trp1 ura3 NTR1HA; YSCC132, Mata prc1 prb1 pep4 leu2 trp1 ura3 NTR2HA; YSCC133, Mata prc1 prb1 pep4 leu2 trp1 ura3 PRP43V5; YSCC141, Mata prc1 prb1 pep4 leu2 trp1 ura3 NTR1HA PRP43V5; YSCC142, Mata prc1 prb1 pep4 leu2 trp1 ura3 NTR2HA PRP43V5; YSCC152, Mata prc1 prb1 pep4 leu2 trp1 ura3 GAL-NTR2, J401, Mata his3 his7 ade3 ura3 prp2-1, prp22, Mata ade2 his3 ura3 tyr1 prp22-1.

**Oligonucleotides**

The following oligonucleotides were used: R1, GGGCGGAC GTCCCGAGCTAGCCTGTTAGAAGTACCCTATAAAAAGG; R2, GGCGGACGTCGTAGTGGAAGCTGAAGGCGGCGG; R3, GGCGGATCTTACATCGGAGATCG; R4, CCCGGTCAAGTGAATGCTAATGATA; R5, GCC GCAGCTCCACAGACTCGGGTTAAGCTTAAAGGCGG; R6, GGCGGAGCATCGTATGGGTAGTACAAATGAG CTTATTTTAGG; R7, GCCGCGATCCATTGTTGAGAC CTGAGG; R8, CCCCGGCTCAGGCTATGCGAAGAGATGCC; R9, GCCGCGATCCGTATCCATAAGTGCT G; R10, CCG GAAGCTTTCTTGGATCTGGTAC; R11, CCGGACTAG TTAGCATTTTGTGTA; R12, CCCGCTGAGCTCCTCC GACAAAGGTA; R13, GAGTTGACGATTCCTATAG; R14, AACCCGTCCGCAAAATG.

**Antibodies and reagents**

The anti-V5 antibody was purchased from Invitrogen Inc. The anti-HA monoclonal antibody 8G5F was produced by immunizing mice with a KLH-conjugated HA-peptide (R.-T. Tsai and S.-C. Cheng, unpubl.), and anti-Ntr2 polyclonal antibody was produced by immunizing rabbits with the Escherichia coli expressed full-length protein. Anti-Ntr1 and anti-Prp43 antibodies were produced by immunizing rabbits with the E. coli expressed Ntr1 C-terminal fragment of amino acid residues 563–709 and Prp43 N-terminal fragment of amino acid residues 1–101, respectively. Protein A-Sepharose was from Amersham Inc., Streptavidin Sepharose from Sigma Inc., and SuperScript II from Life Technology.

**Construction of NTR1-HA-tagged, NTR2-HA-tagged, and PRP43-V5-tagged strains**

For construction of the NTR1-HA-tagged strain, two DNA fragments, A and B, were generated by polymerase chain reaction (PCR) using primers R1 and R4, and R2 and R3, respectively. Following digestion of fragment A, which contains 436 base pairs (bp) of the 3' end of the NTR1 ORF and the 5' half of the HA-epitope, with BamH and AatII, and digestion of fragment B, which contains 609 bp of the 3' untranslated region (UTR) and the 3' half of the HA-epitope, with AatII and XhoI, fragments A and B were ligated into plasmid vector pRS406 digested with BamHI and XhoI. The resulting plasmid was linearized with BclI for transformation into yeast strain BJ2168 to displace the wild-type allele with the HA-tagged allele by the pop-in and pop-out gene displacement method (Winston et al. 1983). For construction of the NTR2-HA-tagged strain, DNA fragments C and D were generated by PCR using primers R5 and R8 and R6 and R7, respectively. Following digestion of fragment C, which contains 471 bp of the 3' end of the NTR2 ORF and the 5' half of the HA-epitope, with BamHI and AatII, and digestion of fragment D, which contains 502 bp of the 3' UTR and the 3' half of the HA-epitope, with AatII and XhoI, fragments C and D were ligated with prRS406 digested with BamHI and XhoI. The resulting plasmid was linearized with BglII for transformation into yeast strain BJ2168 to displace the wild-type allele with the HA-tagged allele by the pop-in and pop-out gene displacement method. For construction of the PRP43-V5-tagged strain, DNA fragments E and F were generated by PCR using primers R9 and R10 and R11 and R12, respectively. Fragment E, which contains 510 bp of the 3' end of the PRP43 ORF, was digested with BamHI and HindIII and ligated with plasmid vector pDK85 digested with BamHI and HindIII. The resulting plasmid was digested with SpeI and XhoI, and ligated with fragment F, which contains 599 bp of the 3' UTR, and had previously been digested with SpeI and XhoI. The resulting plasmid was linear-
ized with MscI for transformation into yeast strain BJ2168 to displace the wild-type allele with the V5-tagged allele.

**Primer extension**

Primer extension was performed with SuperScript II reverse transcriptase based on the method of Chan et al. [2003] using primer R13 or R14. Extension products were analyzed by electrophoresis on 8% polyacrylamide/8 M urea gels.

**Preparation of splicing extracts and substrates**

Yeast whole-cell extracts were prepared according to Cheng et al. [1990]. Actin precursors were synthesized in vitro, using SP6 RNA polymerase according to Cheng and Abelson [1987], and biotinylated pre-mRNA was synthesized following the procedure described by Chan et al. [2003].

**Splicing assays, immunoprecipitation, immunodepletion, and precipitation of the spliceosome by streptavidin agarose**

Splicing assays were carried out according to Cheng and Abelson [1987]. Immunoprecipitation was performed as described by Tarn et al. [1993a] with the anti-HA or anti-Ntc20 antibody. Immunodepletion of Ntr1, Ntr2, and Prp43 was performed by incubation of 0.1 mL each of NTR1-HA, NTR2-HA, and PRP43-V5 splicing extract with 150 ng of anti-HA antibody or 15 µL of anti-V5 antibody, respectively, conjugated to 50 µL of protein A-Sepharose. Precipitation of the spliceosome with streptavidin agarose was carried out according to Chan et al. [2003].

**Purification of NTR complex**

NTR1-HA strain was grown in 10 L of YPD for preparation of splicing extracts and the 40% saturated ammonium sulfate pellet fraction (40P) of the extract according to Cheng et al. [1990]. The anti-HA antibody was conjugated to protein A-Sepharose at a final concentration of 2 mg/mL and cross-linked with dimethylpimelimidate. Approximately 150–200 mg of 40P was added to KPO₄ [pH 7.0] to a final concentration 60 mM, and applied to a final concentration 60 mM, and applied to buffer containing 20 mM HEPES [pH 7.9], 100 mM NaCl, and 0.2 mM EDTA. Gradients were centrifuged in SW60 at 50,000 rpm for 3 h at 4°C and collected in 0.25-mL fractions.

**Gradient sedimentation**

Splicing reactions or released intron fractions were subjected to sedimentation analysis on 10%–30% glycerol gradients in a buffer containing 20 mM HEPES [pH 7.9], 100 mM NaCl, and 0.2 mM EDTA. Gradients were centrifuged in SW60 at 50,000 rpm for 3 h at 4°C and collected in 0.25-mL fractions.

**Acknowledgments**

We thank W.-Y. Tarn and P. Lin for critical reading of the manuscript and K.J. Deen for English editing. This work was supported by the Academia Sinica and by the National Science Council [Taiwan] NSC92-2321-B-001-017.

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Spliceosome disassembly catalyzed by Prp43 and its associated components Ntr1 and Ntr2

Rong-Tzong Tsai, Ru-Huei Fu, Fu-Lung Yeh, et al.

*Genes Dev.* 2005, 19:
Access the most recent version at doi:10.1101/gad.1377405

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