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Mechanism for Aar2p function as a U5 snRNP assembly factor

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Little is known about how particle-specific proteins are assembled on spliceosomal small nuclear ribonucleoproteins (snRNPs). Brr2p is a U5 snRNP-specific RNA helicase required for spliceosome catalytic activation and disassembly. In yeast, the Aar2 protein is part of a cytoplasmic precursor U5 snRNP that lacks Brr2p and is replaced by Brr2p in the nucleus. Here we show that Aar2p and Brr2p bind to different domains in the C-terminal region of Prp8p; Aar2p interacts with the RNaseH domain, whereas Brr2p interacts with the Jab1/MPN domain. These domains are connected by a long, flexible linker, but the Aar2p–RNaseH complex sequesters the Jab1/MPN domain, thereby preventing binding by Brr2p. Aar2p is phosphorylated in vivo, and a phospho-mimetic S253E mutation in Aar2p leads to disruption of the Aar2p–Prp8p complex in favor of the Brr2p–Prp8p complex. We propose a model in which Aar2p acts as a phosphorylation-controlled U5 snRNP assembly factor that regulates the incorporation of the particle-specific Brr2p. The purpose of this regulation may be to safeguard against nonspecific RNA binding to Prp8p and/or premature activation of Brr2p activity.

[Keywords: pre-mRNA splicing, protein interaction, protein phosphorylation, protein structure, spliceosome, yeast]
et al. 2002). Thus, snRNP assembly, restructuring, and reassembly are intimately tied to the splicing process itself.

Two distinct forms of U5 snRNP have been characterized in yeast, distinguished by the presence or absence of the Aar2 protein [Gottschalk et al. 2001; Boon et al. 2007]. Aar2p was discovered as a factor involved in the splicing of some pre-mRNAs in yeast [Nakazawa et al. 1991]. While Aar2p is not required for splicing per se, removal of the protein blocked repeated rounds of splicing in vitro (Gottschalk et al. 2001), suggesting that it could be involved in U5 snRNP or U4/U6–U5 tri-snRNP [re]assembly. More recently, it was shown that the Aar2p-U5 snRNP has a cytoplasmic phase (Boon et al. 2007). In addition to Aar2p and the Sm proteins, it contains the U5-specific Prp8 and Snu114 proteins, but it lacks the essential Br2 helicase [Gottschalk et al. 2001; Boon et al. 2007] that is required for spliceosome activation [Staley and Guthrie 1998] and disassembly [Small et al. 2006]. In the nucleus, Aar2p is replaced by Br2p, which, along with the remaining U5-specific proteins, gives rise to mature U5 snRNP (Boon et al. 2007).

Here, we investigated the structural basis of Aar2p’s association with Prp8p and the mechanism by which it regulates U5 snRNP biogenesis in Saccharomyces cerevisiae. We found that Aar2p forms a stable complex with the RNaseH-like domain of Prp8p, and the C terminus of Aar2p sequesters the C-terminal Jab1/MPN domain of Prp8p, which is a major interaction site of Br2p. Thus, Aar2p directly competes with Br2p for binding to Prp8p. Aar2p is found to be phosphorylated in vivo, including residue S253. Introduction of a phospho-mimetic mutation in Aar2p at position 253 disrupts the Aar2p–Prp8p interaction and allows Br2p entry. Our data suggest that Aar2p acts as a phosphorylation-controlled U5 snRNP assembly factor that regulates the interaction of Br2p, possibly to avoid nonspecific RNA binding by Prp8p and/or premature activation of Br2p activity.

Results

**Aar2p stably interacts with the RNaseH-like domain in the C-terminal region of Prp8p**

Previous work suggested that Aar2p interacts with a C-terminal region of Prp8p [Boon et al. 2006, 2007], although a direct contact between the proteins was not demonstrated. We tested interaction of full-length Aar2p fused to the LexA DNA-binding domain (in pBTM116) with HA-tagged C-terminal Prp8p fragments fused to the Gal4 activation domain (in pACTIII). Prp8pE1 (residues 1649–2413) includes the RNaseH and Jab1/MPN domains, Prp8pE3 (residues 2010–2413) includes the Jab1/ MPN domain but lacks most of the RNaseH domain, and Prp8pE3H (residues 2010–2413) carries two point mutations (Y2037H, I2051T) that increase its interaction with Br2p (Fig. 1A; van Nues and Beggs 2001). As the Aar2-LexA fusion protein alone activated transcription in yeast two-hybrid assays, we tested interactions among the proteins using pull-down assays instead. Aar2-LexAp coprecipitated the Prp8pE1 fusion protein but not Prp8pE3 or Prp8pE3H (Fig. 1B), indicating that Aar2p binds the C-terminal region of Prp8p and that this interaction requires sequence in the E1 fragment between residues 1649 and 2010.

We next used analytical gel filtration analysis to assess the interaction between Aar2p-His6p and fragments of Prp8p, both recombinantly produced in Escherichia coli. Upon mixing individually purified proteins, Aar2p-His6p coeluted with Prp8p1836–2413 [covering the RNaseH and Jab1/MPN domains, herein referred to as Prp8pCTF] at the expected size of a 1:1 complex (Fig. 1C–E). The two proteins could also be coproduced and copurified via three chromatographic steps (Supplemental Fig. S1A), confirming a stable and direct interaction of Aar2p-His6p and Prp8pCTF. Similarly, Aar2p-His6p stably bound to Prp8p1836–2092 [Prp8pRNaseH]. While Aar2p-His6p also comigrated with Prp8p2112–2413 [Prp8pJab1/MPN], the elution volume was the same as for the individual proteins, showing that Aar2p-His6p does not stably interact with Prp8pJab1/MPN [Fig. 1F–H]. The latter finding was confirmed in a pull-down assay (Supplemental Fig. S1B). Thus, Aar2p binds directly to the C-terminal region of Prp8p, and the RNaseH domain of Prp8p is the primary interaction module for Aar2p.

Crystal structure analysis of Prp8pCTF*, Aar2p, and an Aar2p-Prp8pRNaseH complex

To reveal the molecular basis of the Aar2p-Prp8p interaction, we attempted to crystallize the Aar2p-His6p-Prp8pCTF complex but failed. However, we could elucidate the crystal structure of Prp8p1836–2397 (called Prp8pCTF”) alone, using all data collected up to 3.3 Å resolution in the refinement (Fig. 2A; Supplemental Table S1; Supplemental Fig. S2A). Prp8pCTF” lacked 16 amino acids at the C terminus compared with Prp8pCTF. In that structure, a 60-residue linker between the domains lacked electron density, demonstrating its intrinsic flexibility (Fig. 2A). In the crystal structure, the unstructured linker could bridge various pairs of crystallographically equivalent domains [e.g., a closely interacting pair compared with a pair of domains in an open conformation] (Fig. 2A) and we could not unequivocally attribute a particular pair of domains to the Prp8pCTF” molecule in our crystals. However, individually produced RNaseH and Jab1/MPN domains ran as separate entities in gel filtration [Fig. 1C,F]. In addition, protease cleavage in the linker of Prp8pCTF” dissected RNaseH and Jab1/MPN domains that again migrated as individual proteins [Fig. 2B]. These data suggest that in solution, Prp8pCTF” adopts an open conformation in which the two domains are disconnected.

As the Aar2p-His6p-Prp8pCTF complex also failed to crystallize, we suspected additional flexible elements in Aar2p-His6p and subjected the protein to limited proteolysis. Subtilisin cleaved Aar2p-His6p into two fragments of ~18 kDa each, comprising residues 1–159 and 168–324 [Supplemental Fig. S1C]. Thus, subtilisin removed an internal loop [residues 160–167] and the C-terminal 31
amino acids plus the tag of the protein. Gel filtration analysis indicated that the two fragments of subtilisin-treated Aar2p (Aar2pSub) remained stably associated. Like full-length Aar2-His6p, Aar2pSub bound stably to Prp8pRNaseH (Fig. 1I) but failed to interact with Prp8p Jab1/MPN (data not shown).

Aar2pSub crystallized alone and in complex with Prp8pRNaseH. We first solved the structure of the complex by molecular replacement using the structure coordinates of isolated Prp8pRNaseH (Pena et al. 2008; PDB IDs: 3E9O and 3E9P). The Aar2pSub structure was subsequently solved using the coordinates of the Aar2pSub portion of the complex structure. Both structures were refined to low R/Rfree factors with good stereochemistry, using all diffraction data collected to 2.1 Å [Aar2pSub] and 1.8 Å [Aar2pSub–Prp8pRNaseH complex] resolution (Supplemental Table S1; Supplemental Fig. S2B,C).

Prp8pRNaseH grasps Aar2pSub between its thumb and fingers

Aar2pSub is organized in two globular domains, connected via a flexible loop between residues 153 and 171, part of which was cleaved during subtilisin treatment (Figs. 3A; Supplemental Figs. S2D, S3). The two domains interact closely via an extensive hydrophobic interface (53% nonpolar atoms) that buries 1504 Å² of combined surface area (Supplemental Fig. S2E), explaining why they remained stably associated after subtilisin treatment. The N-terminal domain (NTD) contains a 10-stranded, mixed β sandwich with three helices surrounding the upper rim.
The C-terminal domain (CTD) shows an all-helical fold with nine α helices, a π helix, and three 3_10 (η) helices. Residues 319–324 were not seen in the electron density of either Aar2p/Sub alone or in the complex with Prp8pRNaseH.

As previously seen in the isolated protein (Pena et al. 2008; Ritchie et al. 2008; Yang et al. 2008), the Prp8p portion in the complex adopts a central mixed α/β fold reminiscent of RNaseH-like enzymes, with a long β-hairpin insertion [residues 1859–1875] and an additional C-terminal α-helical domain (Fig. 3A). Globally, Prp8pRNaseH resembles a mitten with the RNaseH-like core representing the palm, the β-hairpin insertion representing the thumb, and the helical appendage corresponding to the fingers. The fingers and thumb contact Aar2p/Sub through two discontinuous interfaces [I and II] with a large gap above the palm region in between [Fig. 3A]. At interface I, the tip of the Prp8pRNaseH fingers [formed by the C terminus of helix H6 of Prp8pRNaseH] interacts with both Aar2p/Sub domains primarily via hydrogen bonds and ionic interactions [Fig. 3C, left]. At interface II, the thumb of Prp8pRNaseH latches onto one side of the Aar2p/Sub CTD, running along a cleft between helices α10 and α13 with polar interactions dominating the upper part and hydrophobic interactions prevailing in the lower part of the contact [Fig. 3C, right].

Of the combined surface area, 772 Å² is buried upon complex formation. An extended electronegative surface patch on Aar2p/Sub at interface II and across the gap faces a similarly extensive electropositive surface area on the thumb and palm of Prp8pRNaseH [Fig. 3B]. The association of Aar2p/Sub with Prp8pRNaseH has hallmarks of a facultative interaction, since large parts of the interfaces are hydrophilic [30% nonpolar atoms] and thus compatible with exposure to the aqueous environment. Indeed, the surface areas of the proteins facing each other across the gap are blanketed with water molecules [Fig. 3D].

The structure of Aar2p/Sub in isolation is very similar to the Prp8pRNaseH-bound structure [root-mean-square deviation (RMSD) of 0.92 Å for 289 Ca atoms] (Supplemental Fig. S2F, left), with significant adjustments only in the loop between strands β5 and β6 [NTD] and in the neighborhood of helices η3 and α11 [CTD]. While the β5–β6 loop undergoes an approximate rigid body movement, the region between helices η3 and α11 is structurally rearranged upon complex formation (Supplemental Fig. S2G). Similarly, there are only limited changes in the Prp8pRNaseH structure upon interaction with Aar2p/Sub, which entail small rigid body movements of the thumb and the tip of the fingers toward each other to grasp Aar2p/Sub in between [Supplemental Fig. S2F, right].

The structure of Aar2p/Sub alone or in the complex with Prp8pRNaseH is reminiscent of RNaseH-like enzymes, with a long β-hairpin insertion [residues 1859–1875] and an additional C-terminal α-helical domain (Fig. 3C, right). Phobic interactions prevailing in the lower part of the complex (Fig. 3A) resemble cargo binding by vesicular transport adaptors

Comparison of the Aar2p/Sub structure to known structures in the Protein Data Bank [http://www.pdb.org] revealed the NTD as a novel fold. Although it resembles diverse proteins with similar β sandwiches, no other protein has analogous interspersed helices. The CTD of Aar2p/Sub is most similar to the domain of Pcf11p that binds Ser2-phosphorylated heptad repeats of the C-terminal tail of RNA polymerase II (Z-score = 7.4, RMSD 3.6 Å for 112 Ca atoms) and to the related families of VHS and ENTH domains known from Golgi-ER transport adaptors and endocytic adaptor proteins, respectively [Z-score = 7.0 to the VHS domains of human GGA1 and GGA3; RMSD 3.7 Å for 111 Ca atoms; Fig. 3E]. The helical stack of the CTD also resembles armadillo repeat proteins such as β-catenin or importin-β, posing the question of whether Aar2p may act as a transport adaptor for pre-US snRNP during nucleo-cytoplasmic shuttling. However, Prp8p has its own nuclear localization signal (Boon et al. 2007).

VHS domain proteins bind acidic cluster–dileucine motifs of their cargo proteins between two helices in an extended conformation [Fig. 3F; Misra et al. 2002; Shiba et al. 2002]. This mode of cargo binding resembles the binding of the thumb region of Prp8pRNaseH to the CTD of Aar2p/Sub, although the atomic contacts differ in detail; the Aar2p/Sub–Prp8pRNaseH interaction displays a different arrangement of polar and hydrophobic interactions, and amino acids from both strands of the Prp8pRNaseH β hairpin bind to Aar2p/Sub. In contrast, Pcf11p binds Ser2-phosphorylated RNA polymerase II heptad repeats on the
opposite side of its VHS-like domain (Meinhart and Cramer 2004).

**Aar2p directly competes with Brr2p binding at the Prp8p C-terminal region by sequestering the Jab1/MPN domain**

A number of studies have shown that Brr2p also interacts with the C-terminal region of Prp8p (van Nues and Beggs 2001; Liu et al. 2006; Pena et al. 2007), and it was reported that more Aar2p associates with Prp8p in cells producing less Brr2p (Boon et al. 2007). We therefore investigated whether Aar2p and Brr2p bind competitively to Prp8p.

Gel filtration analysis revealed that Brr2p formed a stable complex with Prp8pCTF (Fig. 4A,B). However, unlike Aar2p, Brr2p interacted stably with Prp8pRNaseH but not with Prp8pRNaseHCTF (Fig. 4C).

Despite the preferred open structure of isolated Prp8pCTF, which might permit concomitant binding of both Aar2p and Brr2p, we could not assemble a ternary Aar2p-His₈p–Prp8pCTF–Brr2p complex. Mixing the three proteins stoichiometrically produced primarily a binary Aar2p-His₈p–Prp8pCTF complex and free Brr2p (Fig. 4D).

Increasing amounts of Brr2p led to formation of a mixture of binary Brr2p–Prp8pCTF and Aar2p–Prp8pRNaseH complexes but were not able to displace Aar2p quantitatively (Supplemental Fig. S4).

These findings could be explained if a preformed Aar2p–Prp8pRNaseH complex sequestered the Jab1/MPN domain, making it unavailable for Brr2p. This idea is consistent with the observation that a transposon insertion in the C-terminal Jab1/MPN domain of Prp8p (at residue 2173) interfered with Aar2p binding to U5 snRNP (Boon et al. 2006), suggesting an involvement of the Jab1/MPN domain in Aar2p binding to Prp8p. Indeed, we observed that addition of Prp8pRNaseHCTF to a preformed Aar2p–Prp8pRNaseH complex assembled a ternary complex, as revealed in gel filtration (Fig. 4F) and by GST pull-down assays (Fig. 5A, lanes 22, 23). Unlike full-length Aar2p, a Aar2pSub–Prp8pRNaseH complex did not bind Prp8pRNaseHCTF (Fig. 4, cf. F and G).

The very C terminus of Aar2p, not contained in our crystal structures, is well conserved among Aar2p orthologs (Supplemental Fig. S3). To test the importance of this C-terminal tail for binding of Prp8pRNaseHCTF, we generated Aar2pΔC-His₈p lacking only the last 23 amino acids, but, in
contrast to Aar2pSub, retaining the flexible loop between NTD and CTD. While Aar2AC•His6p stably bound to Prp8pRNaseH, the preformed Aar2AC•His6p–Prp8pRNaseH complex failed to bind Prp8pJab1/MPN [Fig. 4H]. Furthermore, unlike full-length Aar2-His6p, Aar2AC•His6p allowed formation of a ternary complex Aar2AC•His6p-Brr2p, showing that in the Aar2AC•His6p–Prp8pCTF complex, the Jab1/MPN domain remains available for binding to Brr2p [Fig. 4E]. These data support a model in which Aar2p bound to the RNaseH domain provides a binding platform for the Jab1/MPN domain. The C-terminal 23 amino acids of Aar2p are required for stable sequestration of Prp8pJab1/MPN and competition with Brr2p.

Prp8p residues involved in interface I with Aar2p [Fig. 3C, left] belong to a region [residues 2033–2067] in which
Figure 5. Aar2p is phosphorylated. (A) SDS PAGE analysis of Aar2p–Prp8p interactions by GST pull-down. All experiments were analyzed on the same gel. Numbers on the left indicate the molecular mass of standard proteins [M] in kilodaltons. [I] Mixture added to the glutathione sepharose, [II] pulled-down (bead) fraction. Proteins or protein mixtures added to the beads are identified above the gel. Red double arrows connect pull-downs of wild-type Aar2-His6p compared with Aar2 S253E-His6p. GST-Prp8pRNaseH brought down significantly reduced amounts of Aar2 S253E-His6p (lane 21) compared with wild-type Aar2-His6p (lane 19). Similarly, significantly less Prp8pJab1/MPN was brought down by GST-Prp8pRNaseH in the presence of Aar2 S253E-His6p (lane 25) compared with wild-type Aar2-His6p (lane 23). [B] Phosphatase treatment indicates that Aar2p is phosphorylated. AGY8 cells were grown in different concentrations of galactose to induce Aar2-His10p production to increasing levels. After incubation of extracts with Ni²⁺-NTA beads, the precipitated Aar2-His10p was treated with phosphatase (+) or not (−) as recommended by the manufacturer [New England Biolabs], fractionated by SDS PAGE, blotted, and probed with anti-Aar2 antibody (R5725). [C, D] Aar2p has phosphorylated serine(s) and tyrosine(s) in vivo. Aar2-His10p produced as in A was analyzed by SDS PAGE, blotted, and probed with anti-phospho-serine (mAb p5747, Sigma) [C] or anti-phospho-tyrosine (mAb 42H4; New England Biolabs) [D] antibodies. [E, F] The S253E variant of Aar2p does not interact with Prp8p. [E] Extracts from yeast cells producing Prp8 E1-HAp and wild-type, S253A, or S253E variants of Aar2-LexA fusion protein or with only LexAp (−) were immunoprecipitated with anti-LexA antibodies. Western blot with anti-HA and anti-Prp8 (a8.6) [Boon et al. 2006] antibodies shows that Aar2p S253E pulls down neither full-length Prp8 nor Prp8 E1-HAp. The blot was reprobed with anti-LexA antibody, showing efficient pull-down of Aar2-LexAp. [F] Extracts from yeast cells producing Prp8 E1-HAp or with Gal4AD-HAp [pACT] control as well as wild-type, S253A, or S253E variants of the Aar2-LexA fusion protein were immunoprecipitated with anti-HA antibodies, blotted, and probed with anti-LexA, indicating that Prp8 E1-HAp interacts with Aar2-LexAp wild type and with S253A, but not with S253E. Reprobing with anti-HA antibody verified that Prp8 E1-HAp was efficiently immunoprecipitated in the three extracts. [G, H] Gel filtration analysis of Aar2 S253E-His6p–Prp8p interactions. Details and labeling are as in Figure 1, C–I. Aar2 S253E-His6p does not form a stable complex with Prp8pRNaseH [G], but addition of Prp8p lab1/MPN rescues the ternary complex in gel filtration [H].
amino acid substitutions led to enhanced interaction with Brp2p [van Nues and Beggs 2001]. In light of the present Aar2pSub–Prp8pRNaseH structure, the direct competition between Aar2p and Brp2p for Prp8p suggests that this phenotype may be explained in part by reduced binding of Aar2p and thus easier access of Brp2p to Prp8p.

**Aar2p is phosphorylated in vivo**

In search of a mechanism that would allow replacement of Aar2p by Brp2p to form functional U5 snRNP, we speculated that post-translational modification of Aar2p might regulate the Aar2p–Prp8p interaction. To test whether Aar2p is phosphorylated in vivo, extracts from AGY8 yeast cells [PGAL1:AAR2-HIS10] were treated with λ phosphatase, which dephosphorylates phospho-serine (pSer), phospho-threonine (pThr), and phospho-tyrosine (pTyr) residues, the most frequently phosphorylated amino acids in eukaryotes (Zhuo et al. 1993). λ Phosphatase treatment caused Aar2p to migrate faster than nontreated Aar2p in an SDS denaturing gel (Fig. 5B), indicative of the removal of one or more phosphate groups. In addition, antibodies specific for pSer or pTyr detected Aar2p by Western blotting, with the signal increasing with the amount of Aar2p in the extracts (Fig. 5C,D). However, there was no signal with anti-pThr antibodies.

In order to identify the phosphorylated amino acids in Aar2pHis10p, the protein was purified from AGY8 cell extracts, digested with proteases, and analyzed by mass spectrometry [MS]. Five phosphorylated residues were found (S253, T274, Y328, S331, and T345) (Supplemental Fig. S3), confirming the immunodetection of pSer and additional detection of pThr residues, possibly due to superior sensitivity of the MS approach.

A multiple sequence alignment of Aar2p orthologs (Supplemental Fig. S3) showed the overall similarity in the sequences (23.0% identity and 38.3% similarity between the budding yeast and human proteins) and the locations of the five phosphorylated amino acids in the C-terminal third of Aar2p. Among those, only S253 is conserved in most organisms, including humans, but not in Arabidopsis thaliana, Caenorhabditis elegans, or Xenopus laevis. Thus, phosphorylation of S253 might be important for a widely conserved function of Aar2p.

**The Aar2pS253E phospho-mimetic mutation inhibits binding of Aar2p to Prp8p**

To test possible effects of Aar2p phosphorylation on the ability of Aar2p to bind Prp8p and compete with Brp2p, we mutated each of these phosphorylated residues to glutamate (mimicking phosphorylation) or to alanine (preventing phosphorylation) in the Aar2p-LexA fusion protein, and tested the effect of these mutations in pull-down assays. Strikingly, the wild-type Aar2p-LexA fusion protein and all of the mutant variants, except S253E, brought down endogenous Prp8p and the Prp8E1 fusion protein, whereas the S253E mutant protein reproducibly did not coprecipitate either protein [Figs. 5E, Supplemental Fig. S5A,B]. Likewise, Prp8pE1 pulled down the S253A mutant and wild-type Aar2p-LexA fusions but not the S253E variant [Fig. 5F]. Furthermore, the smaller S253D phospho-mimetic substitution behaved like S253E [data not shown]. These results show that the S253E or S253D phospho-mimetic substitutions inhibit interaction of Aar2p with Prp8p.

We also investigated the growth phenotypes of yeast strains overproducing the interacting proteins. Coproduction of the Aar2pLexA fusion protein and Prp8pE3 was slightly detrimental to growth at low temperature, and the coproduction of Aar2pLexAp and Prp8pE3H was even more so [Supplemental Fig. S5C]. These results suggest that the C-terminal E3 and E3H fragments of Prp8p, containing the Jab1/MPN domain, might sequester Brp2p, preventing it from functioning normally. As the growth inhibition depended on the coproduction of the Aar2pLexA fusion [data not shown], the effect of sequestering Brp2p might be exacerbated by the excess Aar2p competing with endogenous Brp2p for binding to Prp8p. Mutating the S253 residue in the Aar2pLexA fusion protein to alanine or glutamate gave opposite effects, with S253A slightly exacerbating and S253E slightly suppressing the growth defect [Supplemental Fig. S5C]. Thus, the inability of Aar2p with the S253E substitution to interact with Prp8p in vivo alleviates the growth inhibition caused by overproduction of the Prp8E3 or Prp8E3H proteins, supporting a model in which Aar2p competes with Brp2p for binding to Prp8p, unless residue S253 is phosphorylated.

We also tested whether the S253E mutation has a direct effect on binding of Aar2p to Prp8p in vitro. In GST pull-down assays, a strongly reduced amount of Aar2pS253E-His6p was brought down by GST-Prp8pRNaseH compared with wild-type Aar2p-His6p [Fig. 5A, lanes 18–21], and the fraction of Prp8pJab1/MPN pulled down with GST-Prp8pRNaseH in the presence of Aar2pS253E-His6p was similarly reduced compared with wild-type Aar2p-His6p [Fig. 5A, lanes 22–25]. These data show that binding of Aar2pS253E-His6p to Prp8p is strongly reduced compared with wild-type Aar2p-His6p, and that this effect is exerted primarily through a reduced interaction with the Prp8p RNaseH domain. In analytical gel filtration, Aar2pS253E-His6p separated from Prp8pRNaseH [Fig. 5G], but in the gentler conditions of this assay, addition of the Prp8pJab1/MPN fragment rescued the ternary complex [Fig. 5H]. Similarly, in the reductionist in vitro system, Brp2p was not able to quantitatively titrate Prp8pCTF from Aar2pS253E-His6p. However, reproducibly more Prp8pCTF was associated with Brp2p in the presence of Aar2pS253E-His6p compared with wild-type Aar2p-His6p [Supplemental Fig. S5D,E]. These results qualitatively agree with the in vivo pull-down data but suggest that additional factors may be involved in regulating the Aar2p–Prp8p interaction in vivo.

**The Aar2pS253E phospho-mimetic mutation allows binding of Brp2p to Prp8p and leads to conformational changes in Aar2p**

In order to investigate the effect of the Aar2p mutants on the ability of Prp8p to bind to Brp2p in vivo, extracts from cells producing wild-type or mutant Aar2p-LexA fusions were incubated with anti-Prp8 antibodies. Endogenous
Phosphorylation can affect binding of Prp8p RNaseH, which conformational changes may explain how S253 phosphorylation of Aar2p at S253 (mimicked by Aar2pS253E) reduces Aar2p affinity for Prp8p, allowing Brr2p to interact with Prp8p.

S253 lies directly C-terminal of helix α9 in the Aar2p CTD (Fig. 6C). Its side chain hydroxyl engages in a direct hydrogen bond to the backbone carbonyl of L249. Clearly, conformational adjustments would be required to accommodate a phosphate group on S253 and similar conformational rearrangements are expected upon replacement of S253 with the larger, negatively charged glutamate or aspartate. In the present conformation, the larger side chains would lead to steric clashes and the negatively charged moieties would face a hydrophobic pocket formed by F183, L205, and L249. The structural changes are expected to affect the relative positioning of the first segment of the VHS-like domain, encompassing helices α6, α7, and π1, which lie on top of S253 (Fig. 6C). Such conformational changes may explain how S253 phosphorylation can affect binding of Prp8pRNaseH, which occurs remote from the S253 position (Fig. 6C).

Indeed, when subjected to comparative limited proteolysis as a measure of conformational differences, Aar2S253E-His6p yielded different digestion patterns compared with wild-type Aar2-His6p with a number of proteases tested. For example, upon treatment with trypsin, fragments of ~24-kDa, 20-kDa, and 18-kDa as well as several smaller fragments were observed with both mutant and wild-type proteins, but these fragments appeared at lower trypsin concentrations with the mutant protein, which also gave rise to a novel band at ~22 kDa (Supplemental Fig. S6A).

Circular dichroism (CD) spectra of Aar2-His6p and Aar2S253E-His6p reproducibly showed small differences (Supplemental Fig. S6B), indicating some minor changes in the secondary structure contents (calculated from CD: 35.3/33.2% helix, 18.3/17.8% β strand for wild-type/S253E Aar2-His6p, crystal structure of Aar2pSub relative to full-length: 33.5% helix, 15.2% β strand). CD melting profiles showed that both proteins exhibit cooperative unfolding with a similar melting temperature (46.1°C–46.5°C) for the main transition (Supplemental Fig. S6C). Reproducibly, wild-type Aar2-His6p showed a second minor transition above 60°C. Qualitatively, these results are in line with differences in the protease digestion pattern and suggest that upon phosphorylation at S253, Aar2p would undergo a limited conformational change, which most likely affects the VHS-like domain and possibly the C-terminal tail, and that this structural change leads to reduced affinity for Prp8p.

Discussion

Assembly of snRNPs depends on chaperones and recycling factors

Assembly of functional macromolecular complexes is a challenging task in the crowded cellular environment and is frequently assisted by trans-acting factors (for review, see Charle et al., 2010). Among such factors, assembly chaperones associate with subunits or partial complexes to promote the formation of native-like sub-assemblies, prevent the formation of unproductive interactions, and/or avoid premature contacts. Frequently, assembly chaperones impose kinetic or thermodynamic traps that have to be resolved by the action of additional factors and recycling.
assembly factors. Furthermore, different assembly steps often take place in different cellular or subcellular compartments. These principles are nicely illustrated in the biogenesis of Sm core RNPs in metazoans (Char and Fischer 2010). However, it is presently unknown to what extent and how the de novo assembly of snRNP-specific proteins is also regulated.

Continued pre-mRNA splicing requires snRNP recycling and reassembly, since several snRNPs are compositionally and structurally remodeled during the splicing process (Wahl et al. 2009). As the Sm core RNPs apparently remain intact during the splicing cycle, the main task in recycling of snRNPs may be the reorganization of snRNAs and reassociation of snRNP-specific proteins. While Prp24p and the U6-associated Lsm2–8 protein complex indeed assist reassembly of U4 and U6 snRNAs after each round of splicing (Raghunathan and Guthrie 1998; Verdone et al. 2004), no recycling factors for snRNP-specific proteins have so far been characterized.

Aar2p—an assembly factor for a U5 snRNP-specific protein

Like the U4/U6 di-snRNP, U5 snRNP is also profoundly remodeled during splicing (Makarov et al. 2002). Based on the documented precursor–product relationship between Aar2p-U5 snRNP and mature U5 snRNP in yeast (Boon et al. 2007), and since Aar2p depletion interferes with repeated rounds of splicing in vitro (Gottschalk et al. 2001), we investigated Aar2p as a candidate assembly and recycling factor for U5 snRNP-specific proteins.

We show that Aar2p forms a stable complex with the C-terminal region of the U5-specific Prp8p, binding its RNaseH domain and sequestering its Jab1/MPN domain via Aar2p’s C terminus. It thereby hinders Br2p from interacting with the Jab1/MPN domain. Our in vitro binding analyses demonstrate that Br2p alone is inefficient in displacing Aar2p from Prp8p (Supplemental Fig. S4), suggesting that Aar2p imposes a kinetic or thermodynamic trap on U5 snRNP assembly, possibly functioning as an assembly chaperone. Phosphorylation mimicry at residue S253 of Aar2p reduces the affinity of the protein for Prp8p, and our data indicate structural changes in Aar2p[S253E] compared with wild-type Aar2p. Similar structural changes can be expected upon phosphorylation at position 253. Therefore, a presently unidentified Ser/Thr protein kinase may act as another U5 snRNP assembly factor that overcomes the block imposed by Aar2p. Taken together, our results demonstrate that Aar2p is a veritable U5 snRNP assembly factor, which, unlike previously characterized snRNP assembly factors, is required for the ordered binding of a U5 snRNP-specific protein, Br2p (Fig. 7). Further experiments will be required to determine whether Aar2p actually functions as a molecular chaperone.

Figure 7. Model for the U5 snRNP maturation. New elements added to the U5 snRNP maturation model proposed by Boon et al. (2007). Aar2p-U5 snRNP is assembled in the cytoplasm and transported to the nucleus. In the nucleus, Aar2p is phosphorylated by an unidentified kinase (Kinase X). Phosphorylated Aar2p exhibits reduced affinity for Prp8p, leaves the pre-U5 snRNP, and allows Br2p entry. In view of our finding that the RNaseH and Jab1/MPN domains, which are joined by a flexible linker region, do not interact, we propose that the C terminus of Prp8p takes on a different conformation when not bound by Aar2p. Additional U5 snRNP proteins lacking from Aar2p-U5 snRNP are presumably assembled concomitantly with Br2p (Boon et al. 2007). Phosphorylated Aar2p has a possible active role in recruiting Br2p (not shown).
configuration of the Prp8pCTF* fragment (Fig. 2A, left) may still represent a biologically relevant interaction between these domains, which is too weak to survive gel filtration or pull-down assays with the separated domains. The surface of the RNaseH domain that interacts with Aar2p is still accessible in the close conformation of Prp8pCTF*. Furthermore, the C-terminal — 35 residues of the Prp8p Jab1/MPN domain, which comprise a putative Brr2-binding region [Pena et al. 2007], are also surface-exposed in the close conformation of Prp8pCTF*. However, since in a combined model the Jab1/MPN C terminus is remote from the Aar2p binding site on the RNaseH domain [data not shown], additional structural and biochemical studies are required to clarify whether the close conformation of Prp8pCTF* is relevant for the sequestering of the Jab1/MPN domain by Aar2p.

Functional significance of the Aar2p-assisted U5 snRNP assembly

What could be the benefit of the complex Aar2p-mediated U5 snRNP assembly mechanism? Prp8p is an important scaffolding protein at the heart of the spliceosome and interacts with all functional elements of the pre-mRNA, several snRNAs, and other key protein splicing factors (for review, see Grainger and Beggs 2005). Structural and functional analyses of the C-terminal region of Prp8p have suggested that its RNaseH-like domain provides a platform for the handover of the pre-mRNA 5'-splice site from U1 snRNA to U6 snRNA and may be involved in stabilizing the catalytic RNA network [Pena et al. 2008; Ritchie et al. 2008; Yang et al. 2008]. Consistent with a role during spliceosome catalytic activation, the C-terminal region of Prp8p encompassing its RNaseH and Jab1/MPN domains directly modulates the activity of Brr2p [Maeder et al. 2009; Pena et al. 2009; Zhang et al. 2009].

The C-terminal region of Prp8p directly binds the U4/U6 snRNA duplex [Zhang et al. 2009], and the Prp8p RNaseH-like domain also interacts with other RNAs, including a putative mimic of the catalytic RNA core of the spliceosome [Ritchie et al. 2008]. Modeling studies have suggested that the groove between the thumb and fingers of the RNaseH-like domain serves as an RNA-binding site [Pena et al. 2008]. Thus, Aar2p binding at the same site, as shown in the present Aar2pSub–Prp8pRNaseH cocrystal structure, may prevent premature binding of the U4/U6 duplex or binding of nonspecific RNAs at this domain during U5 snRNP assembly.

Upon U4/U6–U5 tri-snRNP formation, Brr2p must be regulated to avoid unwinding U4/U6 di-snRNA prematurely. We suggest that Aar2p regulates the incorporation of Brr2p into U5 snRNP in a manner that silences the Brr2p enzymatic activity during U4/U6–U5 tri-snRNP formation. This suggested role of Aar2p is supported by the observation that the D281N mutant of yeast AAR2 acts as a suppressor of the temperature-sensitive prp38-I allele [Pandit et al. 2006] that causes slow release of U1 and U4 snRNAs during catalytic activation [Xie et al. 1998]. D281 of Aar2p forms a salt bridge with K2066 of Prp8p in interface I (Fig. 3C, left), an interaction that is expected to be weakened upon mutation of D281 to an asparagine.

As in vitro experiments have indicated a role for Aar2p in recycling snRNPs during extended splicing reactions [Gottschalk et al. 2001], Aar2p may have a reciprocal function, displacing Brr2p from Prp8p in a post-splicing complex, to facilitate the regeneration of functional U5 snRNPs. Thus, cycles of phosphorylation and dephosphorylation of Aar2p may regulate U5 snRNP assembly and recycling, and identifying the relevant kinase and phosphatase is the next important step in further elucidating the U5 snRNP [re]assembly mechanism.

Materials and methods

Yeast work

Details of yeast strains and plasmids are provided in Supplemental Tables S2 and S3. For MS analysis, yeast extract was prepared from AGY8 [pCAI1::AAR2-His10] cells grown in 2% (w/v) galactose, and Aar2p-His [p355+pCAI1::AAR2-His10] was affinity-purified using Ni2+-NTA beads followed by SDS-PAGE and analyzed as described in the Supplemental Material.

Analysis of recombinant proteins

All proteins for biochemical, biophysical, and structural studies were purified from yeast and, except Brr2p, were produced in E. coli and purified to near homogeneity by chromatographic techniques. Brr2p was produced in insect cell culture. Structural integrity was checked by CD spectroscopy and CD melting analyses. Targets were crystallized by the sitting drop vapor diffusion method, and diffraction data were collected at beamline 14.2 of the BESSY storage ring (Berlin, Germany). The structures were solved by molecular replacement and refined by standard strategies. Details are given in the Supplemental Material.

Database deposition

Structure coordinates and diffraction data were deposited with the Protein Data Bank [http://www.pdb.org] under accession codes 3SBG (Prp8pCTF*), 3SBS (Aar2pSub), and 3SBT (Aar2pSub–Prp8pRNaseH) and will be released upon publication.

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