CPSF30 and Wdr33 directly bind to AAUAAA in mammalian mRNA 3’ processing

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AAUAAA is the most highly conserved motif in eukaryotic mRNA polyadenylation sites and, in mammals, is specifically recognized by the multisubunit CPSF (cleavage and polyadenylation specificity factor) complex. Despite its critical functions in mRNA 3’ end formation, the molecular basis for CPSF–AAUAAA interaction remains poorly defined. The CPSF subunit CPSF160 has been implicated in AAUAAA recognition, but direct evidence has been lacking. Using in vitro and in vivo assays, we unexpectedly found that CPSF subunits CPSF30 and Wdr33 directly contact AAUAAA. Importantly, the CPSF30–RNA interaction is essential for mRNA 3’ processing and is primarily mediated by its zinc fingers 2 and 3, which are specifically targeted by the influenza protein NS1A to suppress host mRNA 3’ processing. Our data suggest that AAUAAA recognition in mammalian mRNA 3’ processing is more complex than previously thought and involves multiple protein–RNA interactions.

[Keywords: mRNA 3’ processing; cleavage and polyadenylation; polyadenylation signal]

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CPSF30, respectively, can be specifically cross-linked to AAUAAA-containing RNA substrates by UV irradiation (Keller et al. 1991). However, the identities of these proteins were not experimentally determined, and, more importantly, it was unclear whether either of these two proteins directly binds to AAUAAA. Second, the yeast homolog of CPSF160, Yhl1/Cft1p, has been shown to bind RNA (Dichtl et al. 2002). Third, recombinant human CPSF160 binds to RNAs in a pull-down assay and shows a preference for AAUAAA-containing RNAs [Murthy and Manley 1995]. However, CPSF160 has been shown to bind to an enhancer element located 76 nt upstream of the AAUAAA in HIV mRNA PASs [Gilmartin et al. 1995]. Additionally, a recent study that aimed to map protein–RNA interactions in vivo for mRNA 3' processing factors failed to detect specific CPSF160 binding at the AAUAAA hexamer [Martin et al. 2012]. Therefore, it remains unclear how CPSF recognizes AAUAAA in mammalian mRNA 3' processing.

The smallest subunit of the CPSF complex, CPSF30, contains five zinc finger (ZF) domains and, in metazoans, a zinc knuckle (ZK) domain [Barabino et al. 1997]. Recombinant CPSF30 has RNA-binding activity, and its ZK domain enhances this activity [Barabino et al. 1997]. Interestingly, the influenza virus protein NS1A specifically interacts with CPSF30 via its ZF2 and ZF3 domains, and this interaction is responsible for suppressing host mRNA 3' processing [Nemeroff et al. 1998; Twu et al. 2006]. However, the role of CPSF30–RNA interaction in mRNA 3’ processing remains poorly characterized, and it is unclear why CPSF30 is specifically targeted by the viral protein.

In an effort to define the molecular basis for CPSF–AAUAAA interaction, we unexpectedly found that CPSF30 and Wdr33 directly bind to the AAUAAA hexamer. We demonstrated that CPSF30–RNA interaction is essential for mRNA 3’ processing and is primarily mediated by its ZF2 and ZF3 domains. By combining these new results with previous findings, we propose a new model for AAUAAA recognition by CPSF that requires multiple protein–RNA interactions.

### Results

**Immunopurified CPSF specifically recognizes AAUAAA**

Several groups have previously purified CPSF from cultured mammalian cells or animal tissues using chromatography methods [Bienroth et al. 1991; Murthy and Manley 1992; Gilmartin et al. 1995]. However, the yield of CPSF was generally low, and the purity and activities of the isolated complexes tended to vary, making it difficult to carry out detailed functional studies. To overcome these technical limitations, we established HEK293 cell lines that stably express Flag-tagged CPSF73 near the endogenous levels [Shi et al. 2009]. To purify the CPSF complex, we prepared nuclear extract from these cells, carried out immunoprecipitation under stringent conditions using anti-Flag antibodies, and eluted immunoprecipitated proteins under native conditions using Flag peptides. Analyses of the eluted proteins by mass spectrometry and Western blotting suggested that the purified CPSF complex was intact, containing all known CPSF subunits plus an additional associated protein, symplekin (Table 1; Shi et al. 2009). It was noted that the purified CPSF complex contained two CPSF30 isoforms that are encoded by alternatively spliced mRNAs (Fig. 1A). The gel staining profile (Fig. 1A) and mass spectrometry analyses results (Table 1) suggest that all subunits are present in the purified CPSF complex at comparable levels. Furthermore, mass spectrometry analyses of biological replicates revealed that the molecular composition and apparent stoichiometry of the purified CPSF complex were highly consistent (Table 1). Therefore, these cell lines provide a reliable and efficient system for purifying CPSF.

Next we tested whether the immunopurified CPSF specifically recognizes the AAUAAA hexamer and functions in mRNA 3’ processing using multiple in vitro assays. As shown in Figure 1B, gel mobility shift assays demonstrated that a specific RNP complex was formed when CPSF was incubated with an AAUAAA-containing RNA substrate [WT: wild type] derived from the commonly used SV40 late [SVL] transcript PAS. A single-nucleotide substitution in the hexamer [AAGAAA] completely abolished this interaction [Fig. 1B, MT: mutant]. In addition, we also carried out in vitro polyadenylation assays with purified CPSF and recombinant poly[A] polymerase [PAP], which have been shown to be sufficient for reconstituting AAUAAA-dependent polyadenylation [Christofori and Keller 1988; Takagaki et al. 1988]. Consistent with the gel shift results [Fig. 1B], our in vitro polyadenylation assays showed that the AAUAAA-containing, but not the AAGAAA-containing, RNA substrate was specifically polyadenylated when incubated with CPSF and PAP [Fig. 1C]. Similar results were obtained with the adenovirus L3 PAS RNA, another commonly used RNA substrate for mRNA 3’ processing assays [Supplemental Fig. S1]. These results demonstrated that the immunopurified CPSF specifically recognizes the AAUAAA hexamer and is functional in mRNA polyadenylation and thus suitable for detailed characterizations of the CPSF–AAUAAA interaction.

### Table 1. Mass spectrometry analyses of the immunopurified CPSF complex

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<th>Protein name</th>
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<th>Coverage [replicate 2]</th>
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<tr>
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CPSF30 and Wdr33 directly bind to AAUAAA

To define the molecular basis of CPSF–AAUAAA interaction, a key first step was to determine which CPSF subunits directly bind to the AAUAAA hexamer. To this end, we synthesized an L3 RNA substrate containing the wild-type AAUAAA hexamer by splinted ligation. This RNA was radiolabeled at a single position between the third and fourth nucleotide within the AAUAAA hexamer (Fig. 2A, top panel). Another RNA (mutant) was synthesized that contained a mutant hexamer (AAUAC) but was otherwise identical in sequence and the position of the radiolabel (Fig. 2A, top panel). AAUAC was chosen such that the nucleotides immediately adjacent to the radiolabel phosphate were identical between the wild-type and mutant RNAs, thereby minimizing the possible bias in UV cross-linking efficiency caused by nucleotide context.

To identify AAUAAA-interacting proteins, purified CPSF was incubated with either the wild-type or mutant RNA substrates, UV-irradiated, and resolved by SDS-PAGE. Two major radiolabeled bands of ~200 kDa and ~40 kDa and a minor band of ~80 kDa, were specifically detected in AAUAAA-containing samples (Fig. 2A, bottom panel, lane 1), but no radiolabeled bands were observed when the mutant RNA was used (Fig. 2A, bottom panel, lane 4). Consistent with our gel mobility shift and in vitro polyadenylation assay results (Fig. 1B,C), the UV cross-linking pattern again demonstrated that the immunopurified CPSF can specifically bind to AAUAAA-containing RNAs and that a single-nucleotide substitution in the hexamer abolished this interaction (Fig. 2A). Additionally, the UV cross-linking results suggested that CPSF–RNA interactions are mediated primarily by two proteins. To determine which, if any, of these proteins directly binds to AAUAAA, we treated the UV cross-linked CPSF–RNA complexes with RNase T1 or RNase I. RNase T1 cleaves 3’ of unpaired G residues, while RNase I cleaves at all nucleotides in ssRNAs. RNA cleavage by these RNases was experimentally confirmed, and the presence of CPSF did not interfere with RNA digestion (Supplemental Fig. S2). After RNase treatment, only the proteins that are directly cross-linked to the hexamer region should remain radiolabeled. Under these conditions, the two major bands remained labeled, but their apparent molecular weights decreased to ~160 kDa and ~30 kDa, respectively (Fig. 2A, bottom panel, cf. lanes 1 and 2, 3). Such shifts were due to the trimming of the CPSF-associated RNA by the RNases (Supplemental Fig. S2). These data demonstrated that two CPSF subunits of ~160 kDa and ~30 kDa specifically and directly bind to the AAUAAA hexamer. It should be pointed out that the decreases in signal intensities of the two major bands following RNase treatments (Fig. 2A, cf. lanes 1 and 2, 3) indicate that these two proteins may also cross-link to sequences outside AAUAAA.

Based on the prevalent model for mammalian mRNA 3’ processing, we suspected that the 160-kDa species corresponds to CPSF160, which has been implicated in AAUAAA recognition [Murthy and Manley 1995; Dichtl et al. 2002]. Wdr33, a recently identified CPSF subunit with a molecular weight similar to that of CPSF160 [Shi et al. 2009], represents another candidate protein for the ~160-kDa species. The ~30-kDa species most likely corresponds to CPSF30 for several reasons. First, CPSF30 is the only CPSF subunit that has a molecular weight in the ~30-kDa range (Fig. 1A). Second, the ~30-kDa species appeared to contain a doublet [Fig. 2A, lanes 2, 3] that is reminiscent of the two CPSF30 isoforms present in the purified CPSF complex [Fig. 1A]. Finally, CPSF30 is a known RNA-binding protein [Barabino et al. 1997]. To determine the identities of the two proteins that directly bind to AAUAAA, we first treated the UV cross-linked CPSF–wild-type RNA complex with RNase I and then denatured the CPSF–RNA complex by boiling it for 3 min in the presence of 0.5% SDS. We then carried out immunoprecipitation using antibodies against candidate proteins with the denatured CPSF–RNA complexes (see the Materials and Methods for details). As shown in Figure 2B [top panel], CPSF30, CPSF160, and Wdr33 were all specifically immunoprecipitated by their respective antibodies, while other CPSF subunits were not coprecipitated, confirming the specificity of this procedure.
Using this procedure, we found that the 30-kDa radiolabeled species in the CPSF–RNA cross-linking product was specifically immunoprecipitated by anti-CPSF30 antibodies (Fig. 2B, bottom panel). Surprisingly, however, the 160-kDa radiolabeled band was specifically and efficiently immunoprecipitated by anti-Wdr33 antibodies but not by anti-CPSF160 antibodies (Fig. 2B, bottom panel). These results provided direct evidence that CPSF30 and Wdr33 directly bind to the AAUAAA hexamer. Under these conditions, we did not detect cross-linking between CPSF160 and the AAUAAA. The implications of these findings are discussed below.

Mapping the CPSF30 and Wdr33 footprints in the AAUAAA region

As both CPSF30 and Wdr33 bind directly to the AAUAAA hexamer, we next mapped their footprints in the AAUAAA hexamer region at a high resolution. To this end, we prepared a series of AAUAAA-containing RNAs that have sequences identical to those of the L3 wild-type RNA used in Figure 2A but were site-specifically radiolabeled at different sites at or near the AAUAAA hexamer, including 0, −13, −7, and +9 nt relative to the U in the AAUAAA hexamer (Fig. 3A, top panel). We then incubated these RNAs with CPSF and subjected them to UV cross-linking followed by RNase T1 digestion. Our results showed that both CPSF30 and Wdr33 cross-linked to 0 nt and −7 nt but weakly to −13 nt (Fig. 3A, bottom panel, lanes 6, 4, 2). In contrast, a prominent protein of ~70 kDa specifically cross-linked to −13 nt (Fig. 3A, lane 2). Using the same immunoprecipitation procedure as described above (Fig. 2B), we provided evidence that this ~70-kDa species most likely corresponds to Fip1 (Supplemental Fig. S3), a known CPSF subunit that has RNA-binding activity (Kaufmann et al. 2004; Lackford et al. 2014). At +9 nt, however, Wdr33 was the only major protein that cross-linked to RNA, and its cross-linking signal at this site was weaker compared with those at 0 and −7 nt (Fig. 3A, bottom panel, cf. lanes 8 and 4, 6), indicating weaker Wdr33–RNA interaction at +9 nt.

We also mapped the footprints of CPSF30 and Wdr33 on SVL RNA (Fig. 3B). To this end, four different SVL-derived RNAs were synthesized and 3’ radiolabeled. These RNAs contain a unique G residue that is positioned at different positions relative to the AAUAAA hexamer such that RNase T1 digestion of these RNAs will generate a series of different radiolabeled RNA fragments (Fig. 3B, top panel; Supplemental Fig. S4). These RNAs were incubated with CPSF, UV-irradiated, and then treated with RNase T1. Under this condition, only the proteins that directly cross-link to the 3’ fragments (underlined in Fig. 3B, top panel) will be radiolabeled. Our results showed that Wdr33 cross-linked to SVL1–3 but not to SVL4 (Fig. 3B, bottom panel), suggesting that Wdr33 binds to sequences upstream of +12 nt, including the AAUAAA region. In contrast, CPSF30...
bound to SVL1 but showed significantly lower interactions with SVL2 and SVL3, and no interaction was detected with SVL4 (Fig. 3B, bottom panel). These results provided evidence that CPSF30 binds directly to AAUAAA in SVL RNA. Together, our mapping results with two distinct RNA substrates suggest that both CPSF30 and Wdr33 bind specifically to a narrow region spanning the AAUAAA hexamer.

CPSF30 is essential for CPSF–RNA interaction

Mammalian CPSF30 contains five ZF domains and one ZK domain. We next set out to map the protein domains of CPSF30 that mediate its interaction with RNA. To this end, we expressed Flag-tagged full-length [CPSF30 isoform 2: NP_001075028.1] or truncated CPSF30 in which one of the ZF or ZK domains was deleted in HEK293T cells by transient transfection and then purified them by immunoprecipitation with anti-Flag antibodies. As shown in Figure 4B (top panel), the full-length and truncated CPSF30 were specifically purified. We next incubated the recombinant CPSF30 with a radiolabeled AAUAAA-containing SVL RNA substrate followed by UV irradiation. Our results showed that the full-length CPSF30 was efficiently cross-linked to RNA (Fig. 4B, bottom panel). Although small amounts of intact CPSF complex might be copurified in this procedure, it does not seem to contribute significantly to the observed RNA interaction, as no Wdr33 cross-linking signal was observed (Fig. 4B, bottom panel). In addition, recombinant Flag-CPSF30, which was expressed using the baculovirus–insect cell system, also cross-linked to RNA (Supplemental Fig. S5). These results suggest that CPSF30 directly binds to RNA. However, CPSF30 UV cross-linking signals were nearly absent when ZF2 or ZF3 was deleted (Fig. 4B, bottom panel). Additionally, the UV cross-linking efficiency decreased by >60% when the ZK domain was removed (Fig. 4B, bottom panel). These data suggest that CPSF30–RNA interactions are primarily mediated by its ZF2 and ZF3 domains, while the ZK domain also contributes to this interaction.

To determine the role of CPSF30 in the overall CPSF–RNA interaction, we wanted to generate intact CPSF complexes containing a mutant CPSF30 that is deficient of RNA binding. To this end, we established HEK293 cell lines that stably express both Flag-tagged CPSF73 and HA-tagged full-length or truncated CPSF30 in which ZF2 or ZF3 was deleted (ΔZF2/3). Interestingly, the expression of exogenous CPSF30 led to a significant decrease in the levels of the endogenous CPSF30 proteins (Supplemental Fig. S6), indicating a possible autoregulatory mechanism. A similar phenomenon has been described for a number of RNA-binding proteins, including splicing factors of the SR protein family (e.g., Wang et al. 1996). The low levels of the endogenous CPSF30 proteins in these cell lines...
allowed the purification of CPSF complexes that contain the full-length CPSF30, ΔZF2-CPSF30, or ΔZF3-CPSF30 by immunoprecipitation using anti-Flag antibodies. As shown in Figure 4C, Western blotting analyses detected similar levels of all tested CPSF subunits in these complexes, including the HA-tagged CPSF30. These results suggest that the deletion of either ZF2 or ZF3 did not interfere with the association between CPSF30 and the rest of the CPSF complex. We then incubated equivalent amounts of these complexes with radiolabeled AAUAAA-containing SVL RNAs followed by UV cross-linking. In line with our earlier results [Fig. 2A], Wdr33 and CPSF30 specifically cross-linked to the RNAs when CPSF complexes contained the full-length CPSF30 [Fig. 4D]. However, when CPSF complex containing CPSF30-ΔZF2 or CPSF30-ΔZF3 was used, the cross-linking signals of both CPSF30 and Wdr33 dramatically decreased (Fig. 4D). Consistent with these results, when CPSF complexes containing the full-length CPSF30 or ΔZF2/3 mutant were used for in vitro polyadenylation assays, significantly lower levels of RNA polyadenylation were observed for CPSF30-ΔZF2/3-containing CPSF (Fig. 4E). In contrast, when CPSF containing CPSF30-ΔZF2 was used in the same assay, we did not observe a significant difference in polyadenylation efficiency compared with the CPSF complex containing the full-length CPSF30 (Supplemental Fig. S7). These results suggest that the CPSF30-RNA interaction is primarily mediated by its ZF2 and ZF3 domains and that CPSF30 RNA-binding activity is required for the entire CPSF complex to interact with RNA.

Figure 4. Mapping the CPSF30 domains necessary for RNA binding. (A) A schematic of the CPSF30 domain structure. Each circle represents a putative domain. The five ZF domains are marked as ZF1–ZF5. (B) Full-length (FL) or truncated CPSF30 in which one of the ZF or ZK domains are deleted (ΔZF1–5/ZK) were expressed in 293T cells by transient transfection and purified by immunoprecipitation (IP) using anti-Flag antibodies. [Top panel] The purified proteins were resolved by SDS-PAGE and visualized by silver staining. These proteins were incubated with SVL RNA substrate [wild type [WT] in Fig. 1A], UV-irradiated, resolved by SDS-PAGE, and visualized by phosphorimaging. The mutant proteins that showed the most significant differences in UV cross-linking compared with the full-length CPSF30 are marked by white arrows. Cross-linking efficiency was calculated as UV cross-linking signal intensity/protein staining intensity and normalized by that of the full-length sample and is listed. (C) CPSF complexes were purified by immunoprecipitation with anti-Flag antibodies and lysates from HEK293 cell lines that express CPSF73-3xFlag and HA-tagged full-length CPSF30 or the ZF2 or ZF2 deletion mutant (ΔZF2 or ΔZF2). The purified complexes were analyzed by Western blotting with antibodies against specified proteins. (D) CPSF complexes containing the full length or ΔZF2 or ΔZF3 were incubated with SVL RNA substrate, UV-cross-linked, resolved by SDS-PAGE, and visualized by phosphorimaging. Wdr33 and CPSF30 signal intensities were normalized by those of the full-length sample and are listed at the bottom. (E) CPSF complexes containing the full length, ΔZF2, or ΔZF3 were incubated with SVL RNA substrate and recombinant PAP in the presence of ATP. The RNAs were resolved on an 8% urea gel and visualized by phosphorimaging. The percentage of poly(A)+ RNA in each sample was calculated and normalized by that of the full-length sample and is listed.
Transcriptome-wide analyses of CPSF–RNA interaction in vivo

In addition to the in vitro studies with selected RNA substrates as described above, we also characterized CPSF–RNA interactions in vivo through iCLIP (individual nucleotide resolution UV cross-linking and immunoprecipitation) analyses [Huppertz et al. 2014]. Since CPSF160 was believed to be the main RNA-binding subunit of CPSF, we initially wanted to perform iCLIP analysis of CPSF160. To this end, we used a stable HeLa cell line that expresses GFP–CPSF160 in a doxycycline (Dox)-dependent manner (Supplemental Fig. S8A). After GFP-CPSF160 was induced, we irradiated the cells with UV-C [wavelength 254 nm] to induce protein–RNA cross-links. After immunoprecipitation with anti-GFP antibodies, however, very low amounts of RNA were recovered (Supplemental Fig. S8B), most likely due to low cross-linking efficiency. To overcome this limitation, we increased cross-linking efficiency through incorporation of the photoactive ribonucleoside analog 4-thiouridine (4-SU) into nascent RNAs and then UV-A [wavelength 365 nm] irradiation [Hafner et al. 2010]. Significantly higher amounts of RNA were recovered from GFP-CPSF160 immunoprecipitation under this condition [Supplemental Fig. S8B]. To visualize the proteins that are cross-linked to RNAs, we next treated the cell lysates from UV cross-linked cells with RNase I, carried out immunoprecipitation using anti-GFP antibodies, radiolabeled the protein-associated RNAs, and resolved the immunoprecipitation sample by polyacrylamide gel electrophoresis. Strikingly, the most prominent radiolabeled species in the immunoprecipitation sample was not GFP-CPSF160 (~180 kDa) but an ~30-kDa band (Fig. 5A, lane 3). Based on our in vitro results, we suspected that this ~30-kDa species may correspond to CPSF30. To test this possibility, we subjected the anti-GFP immunoprecipitation sample to a second immunoprecipitation using anti-CPSF30 antibodies. Indeed, the ~30-kDa band was specifically immunoprecipitated by anti-CPSF30 antibodies (Fig. 5A, lane 6). These results point to CPSF30 as perhaps the major

Figure 5. Transcriptome-wide analyses of CPSF–RNA interaction by iCLIP. [A] A tet-inducible GFP-CPSF160 expression HeLa cell line was untreated (−) or treated (+) with Dox, incubated with 4-SU, and UV-irradiated. (Lanes 1–3) Cell lysates were treated with RNase I before being subjected to immunoprecipitation [IP] with control [Ctrl] or anti-GFP antibodies. For samples in lanes 4–6, the immunoprecipitation samples were subjected to a second immunoprecipitation with anti-CPSF30 antibodies. All immunoprecipitation samples were 5′ labeled with 32P, resolved on a polyacrylamide gel, and visualized by phosphorimaging. [B] CPSF iCLIP tracks on the Rps12 and Atf4 genes. CstF64 iCLIP tracks based on data from Yao et al. (2012) were included for comparison. Each peak represents a unique cross-linking site. Nucleotide sequences around the iCLIP peaks are shown above the tracks. [C] Transcriptome-wide distribution of CPSF cross-linking sites relative to the cleavage sites [ends of transcripts]. [D] Transcriptome-wide distribution of CPSF cross-linking sites relative to AAUAAA (the underlined A corresponds to position 0 on the X-axis).
RNA-binding subunit of the CPSF complex in vivo. Although weak protein–RNA cross-linking signals were observed at ~160 kDa, they were significantly lower than the signals from CPSF30–RNAs (Fig. 5A, lane 3), indicating that Wdr33–RNA cross-linking products were not efficiently recovered under these conditions.

To analyze the in vivo CPSF–RNA interactions at the transcriptome level, we constructed iCLIP libraries from RNAs recovered in the anti-GFP immunoprecipitation sample and subjected them to high-throughput sequencing using the Illumina HiSeq platform. Using these data, we generated a CPSF–RNA interaction map at single-nucleotide resolution. As shown earlier (Fig. 5A), CPSF30 seems to be the major mediator of these interactions. Consistent with our in vitro data, CPSF–RNA in vivo cross-linking sites are highly enriched near the AUAAA hexamer (Fig. 5B,C). As UV cross-linking only occurs at U residues in our procedure, CPSF cross-linking was often observed at the U runs nearest to the AUAAA hexamer, as shown in the Rps12 and Atf4 transcripts (Fig. 5B). CPSF iCLIP signals are distinct from those of CstF64 (Fig. 5B; Yao et al. 2012), demonstrating the specificity of our iCLIP analyses. At the transcriptome level, CPSF cross-linking sites showed two peaks at −26 and −7 nt, respectively, relative to the cleavage site (Fig. 5C). The valley between the two peaks coincides with the position of the AUAAA hexamer (Fig. 5C). To map CPSF-binding sites near AUAAA at a higher resolution, we plotted the distribution of CPSF iCLIP sites relative to the AUAAA itself. Strikingly, CPSF cross-linking sites displayed a bell-shaped peak that is only disrupted by a sharp valley near the center where AUAAA resides (Fig. 5D). As mentioned earlier, such a decrease in CPSF cross-linking was most likely due to the low U content in the AUAAA hexamer. As CPSF30 is the main mediator of CPSF–RNA interactions under the conditions used in this analysis (Fig. 5A), our iCLIP analyses provided in vivo evidence that CPSF30 binding encompasses the AUAAA hexamer on a transcriptome-wide level.

Discussion

A central question in the mRNA 3’ processing field has been how PASs are recognized and how PAS selection can be regulated. We provided critical and novel insights into the mechanism by which the AUAAA hexamer is recognized by CPSF. Based on data presented here and previous studies, we propose a new model for CPSF–RNA interaction that involves multiple protein–RNA interactions (Fig. 6). First, we provided direct evidence that CPSF30 and Wdr33 directly bind to AUAAA. These interactions anchor CPSF to RNA and provide specificity to mRNA 3’ processing. The individual contributions of CPSF30 and Wdr33 to CPSF RNA-binding specificity, however, remain to be determined. Second, previous in vitro and in vivo studies suggest that Fip1 binds to U-rich upstream sequences (Kaufmann et al. 2004; Martin et al. 2012; Lackford et al. 2014). Third, CPSF160 interacts with RNA over a broad region upstream of the cleavage sites, including the AUAAA area (Gilmartin et al. 1995; Murthy and Manley 1995, Martin et al. 2012). Among these factors, we studied CPSF30 in detail and demonstrated that CPSF30 is essential for the entire CPSF complex to interact with RNAs. Interestingly, we found that CPSF30–RNA interactions are primarily mediated by its ZF2 and ZF3, the exact same domains targeted by the influenza protein NS1A to suppress host mRNA 3’ processing (Nemerosoff et al. 1998; Tvu et al. 2006). Together with these earlier studies, our results indicate that the influenza virus has evolved to precisely target this essential protein–RNA interaction to suppress host mRNA 3’ processing (Fig. 6). Additionally, it has been shown in plants that CPSF30 plays an important role in PAS selection, and its activity is regulated by calmodulin [Delaney et al. 2006; Thomas et al. 2012]. Therefore, CPSF30–RNA interaction may serve as a hub for mRNA 3’ processing regulation.

We previously identified Wdr33 as a component of the mammalian mRNA 3’ processing complex and a bona fide subunit of the CPSF complex (Shi et al. 2009). The yeast Wdr33 homolog Pfs2p is a subunit of the CPF complex and an essential 3’ processing factor [Ohnacker et al. 2000]. Pfs2 inactivation in fission yeast also leads to defects in chromosome replication and segregation [Wang et al. 2005]. Wdr33/Pfs2 has been shown to regulate synapse and axon development in Caenorhabditis elegans [Van Epps et al. 2010]. Mutations in the plant Wdr33 homolog FY lead to aberrant mRNA alternative polyadenylation and defective flowering time control [Simpson et al. 2003]. These studies strongly suggest that Wdr33 plays an essential role in mRNA 3’ processing and in mediating cross-talk between Wdr33 and other UPF proteins.

Figure 6. A new model for CPSF–RNA interaction. Multiple proteins mediate CPSF–RNA interactions: CPSF30 and Wdr33 directly interact with AUAAA, and CPSF30 binds to AUAAA via its ZF2 and ZF3 (shown as two fingers). Fip1 binds to auxiliary upstream sequences. CPSF160 binds to RNA over a broad region upstream of the cleavage sites. CPSF30, CPSF100, and symplekin do not participate in PAS recognition. During influenza viral infection, the viral protein NS1A specifically binds to the ZF2 and ZF3 of CPSF30 within the CPSF complex. As CPSF30–RNA interactions are primarily mediated by its ZF2 and ZF3 and are essential for CPSF binding to RNA, NS1A–CPSF30 interaction blocks CPSF–AUAAA interaction and host mRNA 3’ processing.
mRNA 3’ processing and other cellular processes. However, the exact functions of Wdr33 in mRNA 3’ processing remained poorly understood. In this study, we provide in vitro evidence that Wdr33 directly binds to AAUAAA (Fig. 2). In addition, recent photoactivatable ribonucleoside-enhanced CLIP (PAR-CLIP) analyses suggest that Wdr33 binds to AAUAAA in vivo at the transcriptome level (see the accompanying study by Schönemann et al. 2014). Wdr33 contains a highly conserved WD40 repeat domain. This domain constitutes the majority of the yeast Pfs2p, while the metazoan Wdr33 proteins are significantly larger and contain additional domains (Chan et al. 2011; Xiang et al. 2014). Consisting of repeats of a 31-60-residue motif that forms β-propeller structures, WD40 domains are best known for mediating protein–protein interactions (Stirnimann et al. 2010). However, several studies suggest that this domain may also participate in RNA interaction. For example, the WD40 domain of Gemin5 in the SMN complex binds to snRNAs in a sequence-specific manner (Lau et al. 2009). Two recent proteomic analyses of poly[A]+ RNA-binding proteins identified 16 and 23 WD40-containing proteins in the mouse and human proteomes, respectively (Castello et al. 2012; Kwon et al. 2013). In future studies, it will be of great interest to characterize the Wdr33–RNA interactions through structural and biochemical analyses, which may provide important insights into the mechanisms of mRNA 3’ processing as well as WD40–RNA interactions in general.

CPSF160 has also long been implicated in RNA binding and AAUAAA recognition. Purified recombinant CPSF160 was found to bind AAUAAA with weak specificity in vitro (Murthy and Manley 1995), and the yeast CPSF160 homolog Yhh1/Cit1p avidly binds RNA around the PAS in vitro (Dichtl et al. 2002). While it is unclear why CPSF160 RNA binding was not detected in the present study or the accompanying one by Schönemann et al. (2014), our results suggest that the mechanism for AAUAAA recognition is more complex than previously thought. We propose several possible models to integrate our results with those from previous studies and help explain how these interactions could be accommodated and coordinated in the region around the AAUAAA. First, given the dynamic nature of mRNA 3’ processing (Chan et al. 2011), it is possible that CPSF160 and CPSF30/Wdr33 may bind to AAUAAA at different stages of the mRNA 3’ processing reaction. This may be similar to pre-mRNA splicing, in which the 5’ splice site is initially bound by the U1A protein and subsequently by U1 snRNA through base-pairing (Du and Roshbash 2002). Second, earlier biochemical studies suggested that the molecular weight of the CPSF complex ranged from ~350 kDa to ~500 kDa (Bienroth et al. 1991; Murthy and Manley 1992). More recent functional studies suggest that some CPSF subunits form stable subcomplexes (Sullivan et al. 2009; Schönemann et al. 2014). Therefore, there could be multiple alternative CPSF complexes or subcomplexes that rely on different protein–RNA interactions for AAUAAA recognition. Third, RNA sequences encompassing the AAUAAA hexamer could form secondary structures (Gilbert et al. 1992; Hans and Alwine 2000), thereby allowing CPSF subunits to contact RNA in multiple locations. This may explain the previous observations that CPSF contacts both the AAUAAA hexamer and an enhancer element >70 nt upstream (Gilbert et al. 1995) and that CPSF160 interacts with RNA in a broad region upstream of the cleavage sites (Martin et al. 2012). Finally, given the high degree of diversity in mammalian PAS sequences (Proudfoot 2011), CPSF160 and CPSF30/Wdr33 may be required for AAUAAA recognition in different PAS sequence contexts. This is reminiscent of the transcription factor TFIID, which relies on multiple protein–DNA interactions for the precise recognition of diverse promoter sequences (Cler et al. 2009). Interestingly, although CPSF160 shares little sequence homology with Wdr33, it was predicted to form a number of WD40-like β-propeller structures, including the putative conserved RNA-binding domain (Dichtl et al. 2002; Xiang et al. 2014). Therefore, both CPSF160 and Wdr33 may rely on β-propeller structures for their RNA-binding activities. In any event, our studies have provided evidence for a previously unappreciated complexity in AAUAAA recognition by CPSF.

Materials and methods

RNA substrates

SVL RNA and sequences used were as follows: SVL wild type, CUGCAUAUAACAGUGUA [Figs. 1B, C, 4; Supplemental Figs. S5, S7]; SVL mutant, CUGCAAGAAACAGUGUA [Figs. 1B, C, 4]; L3 wild type, AUGUACUAGGACACAAUAGCGAAU GUUUUAUUGUGUA [Figs. 2, 3A, Supplemental Figs. S1–S3]; and L3 mutant, AUGUACUAGGACACAAUACCGCAAU GUUUUAUUGUGUA [Fig. 2, Supplemental Fig. S1]. The SVL RNA substrates were chemically synthesized and 5’ radiolabeled using T7 RNA polymerase. The L3 RNA substrates were site-specifically radiolabeled by splinted ligation of two chemically synthesized RNAs. For mapping experiments in Figure 3B, the RNAs were synthesized by in vitro transcription with DNA oligo templates and T7 polymerase.

Protein purification

CPSF purification and mass spectrometry analyses were carried out as described before (Shi et al. 2009). To purify the CPSF complex shown in Figure 4, C–E, the HEK293-CPSF73-3xFlag cell line was retrovirally transduced to express HA-CPSF30 (full-length or truncated), and cell lysates were used for immunoprecipitation as described above. For purifying recombinant CPSF30 and its mutants, Flag-CPSF30 (full-length or truncated)-pCDNA3 plasmids were transiently transfected into HEK293T cells by using the calcium phosphate method. At 48 h post-transfection, whole-cell extracts were prepared from mock or transfected cells and used for immunoprecipitation.

In vitro assays

For UV cross-linking assays, ~1 fmol of RNA substrates and ~60 fmol of CPSF were incubated and UV-irradiated according to a previously published procedure (Keller et al. 1991). Afterward, the samples were immediately mixed with SDS sample buffer or incubated with 20 U of RNase T1 or 60 U of RNase I for 5 min at 37°C. The cross-linking samples were resolved by SDS-PAGE and visualized by phosphorimaging. Immunoprecipitation of UV cross-linked protein was carried out as previously described (Yuan et al. 1999) using the following antibodies: CPSF30 [Bethyl
For iCLIP experiments, cells were preincubated with 4-SU for 1 h
iCLIP streptomycin). hygromycin, 4 mM medium (DMEM) with 10% fetal bovine serum (FBS) supplemented
GFP-CPSF160 were maintained in Dulbecco’s modified Eagle medium (DMEM) with 10% fetal bovine serum (FBS) supplemented
Cell culture
HEK293, 293T, and HeLa FlpIn cell lines containing inducible GFP-CPSF160 were maintained in Dulbecco’s modified Eagle medium [DMEM] with 10% fetal bovine serum [FBS] supplemented with the appropriate antibiotics [CPSF73-3xFlag cell line: 200 μg/mL G418, HeLa FlpIn GFP-CPSF160 cell line: 100 μg/mL hygromycin, 4 μg/mL blasticidine, 10 U/mL penicillin, 10 μg/mL streptomycin].

iCLIP
For iCLIP experiments, cells were preincubated with 4-SU for 1 h and cross-linked with 2 × 400 ml/cm² and 365-nm UV light. Protein G Dynabeads [Life Technologies] were used for immunoprecipitation. iCLIP library construction and data analyses were performed as described (Huppertz et al. 2014). Sequencing data have been deposited to EMBL-EBI ArrayExpress Archive (accession no. E-MTAB-2939).

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References


Supplemental materials for:

CPSF30 and Wdr33 directly bind to AAUAAA in mammalian mRNA 3’ processing

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Content: eight supplemental figures
Figure S1. L3 RNAs containing a wild-type (WT) or mutant (MT) hexamer was incubated with immunopurified CPSF and recombinant PAP in the presence of ATP as described in Fig. 1C. The RNAs were resolved on an 8% urea gel and visualized by phosphorimaging. The pre-mRNA substrate and the polyadenylated (poly(A)+) RNA are marked.
Figure S2. RNAs were extracted from samples shown in Fig. 2A, resolved on a 20% urea gel, and visualized by phosphorimaging. An RNA ladder was loaded in the far left lane. The results showed that the RNases digested at the predicted sites and that the presence of CPSF did not interfere with the RNase digestion.
**Figure S3.** Immunopurified CPSF was incubated with RNA substrates and irradiated by UV as described in Fig. 2B. The sample was then denatured before subjected to IP using anti-Fip1 antibodies, similar to Fig. 2B. The IP samples were analyzed by Western blotting (top panel) or phosphorimaging (bottom panel). Note that Fip1 in the IP sample migrated slower than the input in both western blots and phosphorimage.
**Figure S4.** For samples in Fig. 3B, the RNAs were extracted after RNase T1 digestion and resolved on an 20% urea gel and visualized by phosphorimaging. RNase T1 digested at the predicted sites and the presence of CPSF did not affect RNase digestion.
Figure S5. (A) Flag-CPSF30 expressed in insect cells (Sf9) was purified by Flag IP and resolved by SDS-PAGE and visualized by silver staining. (B) Purified recombinant Flag-CPSF30 was incubated with SVL RNA substrate and UV crosslinked (right lane) as described in Fig. 4B. Purified CPSF complex was used for comparison (left lane).
Figure S6. Western blotting analyses of CPSF73-3xFlag & HA-CPSF30 cell lines. Cell lysates from HEK293 cell lines expressing CPSF73-3xFlag, or CPSF73-3xFlag plus HA-CPSF30 full-length (FL), zinc finger 2 deletion (ΔZF2), or zinc finger 3 deletion (ΔZF3). Note that HA-CPSF30 proteins are the predominant form of CPSF30 present in these cells.
Figure S7. (A) CPSF complexes were purified by Flag IP with and lysates from HEK293 cell lines that express CPSF73-3xFlag and HA-tagged full-length (FL) CPSF30 or zinc knuckle deletion mutant (ΔZK). The purified complexes were analyzed by Western blotting with antibodies against specified proteins. (B) CPSF complexes containing the FL or ΔZK CPSF30 were incubated with SVL RNA substrate and recombinant PAP in the presence of ATP. The RNAs were resolved on an 8% urea gel and visualized by phosphorimaging. Pre-mRNA and poly(A)+ RNA are marked. The percentages of poly(A)+ normalized to that of the FL samples and listed below the gel image.
Figure S8. (A) Western blotting analyses of GFP-CPSF160 cell line before (-) and after (+) Dox treatment. (B) Following GFP-CPSF160 induction, the cells were irradiated with UV-C or incubated with 4-thiouridine and irradiated with UV-A. The cell lysates were treated with RNase I and used for IP with anti-GFP antibodies. The IP samples were 5’ radiolabeled with Optikinase and γ-32P-ATP, resolved by SDS-PAGE and visualized by phosphorimaging.