

Pre-mRNA 3'-end processing complex assembly and function

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The 3'-ends of almost all eukaryotic mRNAs are formed in a two-step process, an endonucleolytic cleavage followed by polyadenylation (the addition of a poly-adenosine or poly(A) tail). These reactions take place in the pre-mRNA 3' processing complex, a macromolecular machinery that consists of more than 20 proteins. A general framework for how the pre-mRNA 3' processing complex assembles and functions has emerged from extensive studies over the past several decades using biochemical, genetic, computational, and structural approaches. In this article, we review what we have learned about this important cellular machine and discuss the remaining questions and future challenges. © 2010 John Wiley & Sons, Ltd. *WIREs RNA* 2011 2 321–335 DOI: 10.1002/wrna.54

INTRODUCTION

Pre-mRNA 3' processing is an essential step in eukaryotic mRNA biogenesis, and it directly impacts many other steps in the gene expression pathway, such as transcription termination, splicing, mRNA export, translation, and mRNA turnover.^{1–3} 3' processing is also a versatile mechanism for gene regulation.⁴ For example, the efficiency of 3' processing, the usage of alternative polyadenylation (APA) sites, and the length of poly(A) tails can all be modulated. Additionally, 3' processing-mediated gene regulation can be global or transcript-specific. Accumulating EST data revealed that a significant portion of eukaryotic genes produce alternatively polyadenylated mRNAs,⁵ indicating that APA may be involved in the regulation of a large set of genes.⁶ Indeed a series of recent genomic studies showed that APA regulation is widespread and plays important roles in the immune⁷ and the neural systems,⁸ oncogenesis,⁹ stem cell differentiation,¹⁰ and development.¹¹ Defects in pre-mRNA 3' processing have been associated with a wide spectrum of human diseases.¹² Therefore, it is critical to understand how the pre-mRNA 3' processing machinery functions and how its activity can be regulated.

Much progress has been made in understanding the basic molecular mechanisms and regulation of pre-mRNA 3' processing. The core *cis*-elements

and the majority of the basal *trans*-acting factors required for 3' processing have been identified.^{1,2,4,13} Biochemical, genetic, and structural analyses have provided insights into the functions of some 3' processing factors.¹³ A diverse array of mechanisms has been uncovered regarding how 3' processing can be regulated.^{2,4} More recently, purification of the functional human pre-mRNA 3' processing complex has allowed compositional characterization of the entire machinery and made it possible to study its structure and dynamics throughout the cleavage and polyadenylation processes.^{14,15} In this article, we provide a broad overview of our current knowledge of the assembly and function of the pre-mRNA 3' processing complex and highlight recent progress in the field. The main focus will be the mammalian pre-mRNA 3' processing complex, but comparisons with its counterparts in yeast and plants will be provided to illustrate both the conservation and divergence of the 3' processing system in eukaryotic evolution.

PRE-mRNA 3' PROCESSING COMPLEX: THE COMPONENTS

Assembly of the pre-mRNA 3' processing complex is initiated by the binding of *trans*-acting factors to their target *cis*-elements in the RNA. Unlike the spliceosome whose assembly is aided by a series of specific RNA–RNA base-pairing events,¹⁶ assembly of the 3' processing complex on the pre-mRNA is directed entirely by RNA–protein interactions.^{1,2,4,15} As will be discussed later, the *cis*-elements for 3' processing are

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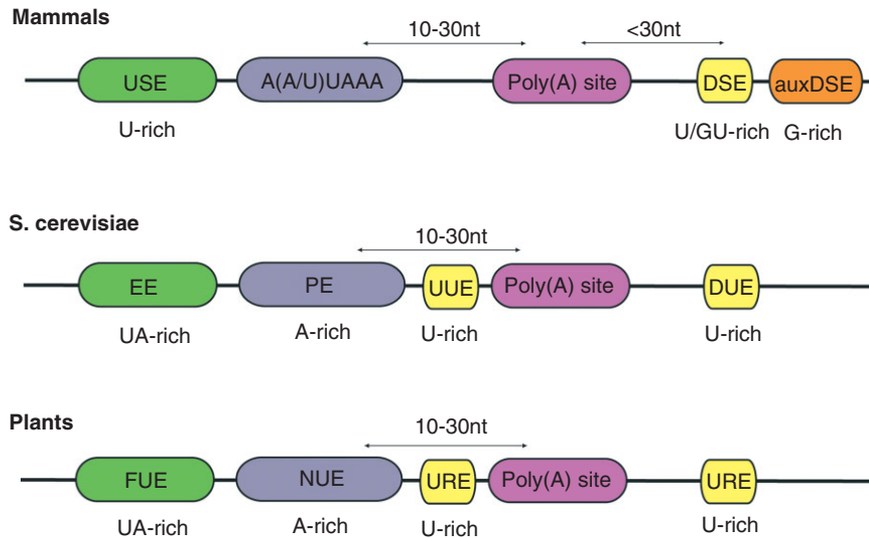


FIGURE 1 | Comparison of the poly(A) signals from animals (canonical) the budding yeast (*Saccharomyces cerevisiae*) and plants. USE: upstream element; DSE: downstream element; auxDSE: auxiliary downstream element; EE, efficiency element; PE, positioning element; UUE, upstream U-rich element; DUE, downstream U-rich element; FUE, far upstream element; NUE, near upstream element; URE, U-rich element. Same colors are used for similar *cis*-element to illustrate the conservation of the basic tripartite poly(A) signal across different phylogenetic groups. (Reprinted with permission from Ref 2. Copyright 1999 American Society for Microbiology)

generally short and/or degenerate (Figure 1) and the interactions of individual 3' processing factors with RNA are usually very weak. Therefore, an extensive network of RNA–protein and protein–protein interactions is required for the assembly of a stable 3' processing complex (Figure 2). Here we discuss the *cis*-elements and protein factors involved.

cis-Elements

Assembly of the pre-mRNA 3' processing complex is directed by multiple key *cis*-elements collectively called the polyadenylation signals (PAS).² Mammalian PASEs can be classified into two general types, the canonical and the noncanonical.^{2,4,17} The main *cis*-elements in a canonical PAS include a highly conserved A(A/U)UAAA element located 10–30 nucleotides upstream of the cleavage/polyadenylation site (poly(A) site) and a more variable U/GU-rich downstream element (DSE) ~30 nucleotides downstream of the poly(A) site (Figure 1). These two elements determine the general location of the poly(A) site, which is most frequently found after an adenosine residue.¹⁸ Genomic studies suggest that poly(A) site selection *in vivo* is fairly heterogeneous and, for any given PAS, the corresponding poly(A) sites are usually clustered in a ~20 nt window.^{5,19} In addition to the A(A/U)UAAA and the DSE, functionally important upstream sequence elements (USE) and auxiliary downstream elements (auxDSE) have also been identified in many viral and cellular mRNAs^{2,4,20} (Figure 1).

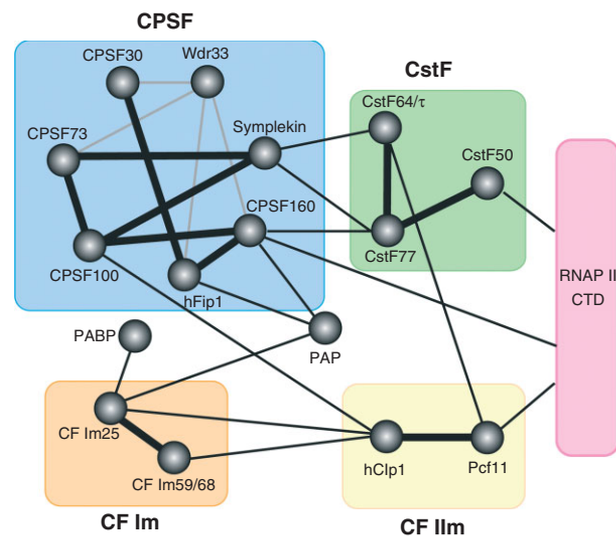


FIGURE 2 | The protein–protein interaction network of the mammalian pre-mRNA 3' processing complex. Circles represent individual proteins except for RNAP II CTD (pink rectangle). Colored rectangles represent subcomplexes. Experimentally verified interactions within subcomplexes are marked with thick black lines. Experimentally verified interactions among subcomplexes are marked with thin black line. Predicted interactions based on information on homologs from other species are marked with gray lines.

USES are generally U-rich while auxDSEs are mostly G-rich. These additional *cis*-elements help improve 3' processing efficiency by providing binding sites for the core 3' processing factors and/or regulatory factors.

Twenty to thirty percent of human PASEs lack the A(A/U)UAAA element and it remains poorly understood how these noncanonical PASEs are recognized by the 3' processing machinery or whether they are recognized by the same 3' processing factors that bind canonical PASEs.^{5,21} In some cases, UGUAN and related sequences have been identified as the key *cis*-element for directing A(A/U)UAAA-independent cleavage and polyadenylation.²² Recently another type of PAS was discovered that consists of only an A-rich USE and a potent DSE.²³ Currently, however, it is not clear whether the noncanonical PASEs share any common characteristics. Interestingly, noncanonical PASEs are often associated with APA and may play important regulatory roles.⁵

The PASEs of the budding yeast and plant pre-mRNAs are thought to be significantly different from the mammalian PASEs as they lack the highly conserved A(A/U)UAAA element.^{2,4} Instead, yeast and plant PASEs include a degenerate A-rich element which includes AAUAAA-like sequences (called the Positioning Element (PE) in budding yeast and the Near Upstream Element (NUE) in plants), a UA-rich USE (called the Efficiency Element (EE) in budding yeast and the Far Upstream Element (FUE) in plants), and a U-rich element (URE) upstream and/or downstream of the poly(A) site^{2,4} (Figure 1). Interestingly, these *cis*-elements are also shared by some mammalian non-canonical PASEs,^{22,23} indicating that PAS sequences might have been more conserved than previously thought.

Given the lack of strong, evolutionarily conserved consensus sequence in eukaryotic PASEs, RNA secondary structures have long been suspected to play a role in directing the assembly of the 3' processing complex.^{24,25} Multiple lines of evidence support this notion. First, an *in vitro* selection for 3' processing enhancers enriched motifs that are related in structure rather than sequence.²⁴ Secondly, functionally important secondary structures have been identified in the PASEs of a number of viral mRNAs.^{26,27} It remains to be seen, however, whether RNA secondary structure plays a general role in 3' processing and if so, what specific secondary structures are required and how they are recognized.

3' Processing Factors

Most, if not all, basal 3' processing factors have been identified^{4,13,15} (Table 1). Mammalian 3' processing factors include the poly(A) polymerase (PAP), the poly(A)-binding proteins (PABPs), the RNA polymerase II large subunit (RNAP II), and four multi-subunit protein complexes, CPSF, CstF, CF Im, and CF IIm.^{1,2,13} All of these factors except for PABPs are

required for cleavage, but CPSF and PAP are believed to be sufficient for the subsequent polyadenylation. PABPs are not required for either cleavage or polyadenylation, but function in activating PAP and in poly(A) tail length control.²⁸ Despite the significant divergence in the PAS sequences between yeast and mammals, most mammalian 3' processing factors have homologs in yeast and plants,^{2,13} indicating that the pre-mRNA 3' processing machinery and the biochemical mechanism have been conserved.

CPSF

CPSF is required for both cleavage and polyadenylation. It recognizes the A(A/U)UAAA element, helps to recruit other components of the 3' processing complex, and catalyzes cleavage.^{1,2,13} Originally it was shown that the purified CPSF consisted of CPSF160, 100, and 73,²⁹ and in some preparations also CPSF30.³⁰ Later studies have added symplekin, hFip1, and Wdr33 to the list of CPSF components.^{14,31,32} Next we will discuss the roles of the individual CPSF subunits in the context of the three aforementioned critical functions of CPSF.

CPSF in RNA Binding

CPSF recognizes the A(A/U)UAAA element with remarkable specificity as any single mutation within the hexamer strongly represses processing.^{33,34} Surprisingly, however, none of the CPSF subunits has a canonical RNA recognition motif (RRM). CPSF160 is believed to be the major CPSF subunit responsible for recognizing the A(A/U)UAAA element based on multiple lines of evidence. For example, CPSF160 can be UV crosslinked to AAUAAA-containing RNAs,^{33,35} and recombinant CPSF160 preferentially binds to RNAs containing AAUAAA than those with mutant hexamers.³⁶ Two fragments within the N-terminal half of CPSF160 bear limited similarities to the RRM domain sequence and were initially suspected to mediate RNA recognition.³⁶ However, a later study of Cft1p, the yeast CPSF160 homolog, attributed its RNA-binding activity to a central domain.³⁷ Although this region seems to be fairly well conserved from CFT1 to CPSF160, mutational analysis is needed to directly test the involvement of this segment in A(A/U)UAAA recognition by CPSF160.

At least two other subunits of the CPSF are involved in RNA binding, hFip1, and CPSF30.^{32,38} The C-terminal arginine-rich domain of hFip1 binds to U-rich sequences.³² The RNA-binding activity of CPSF30 is mediated by its zinc finger domain and it also prefers poly(U) sequences.³⁸ Since many USEs are U-rich sequences (Figure 1), hFip1 and CPSF30 may be involved in recognizing the USEs and other

TABLE 1 | Conservation of Polyadenylation Factors Between Yeast and Mammals

Mammalian Factor Name	Synonyms	Yeast Homolog	Function	Motifs
CPSF				
CPSF160	CPSF1	CFT1/YHH1	RNA-binding (AAUAAA)	SFT1, CPSF_A
Wdr33	WDC146	PFS2		WD40 repeat
Symplekin		PTA1	Scaffolding	
CPSF100	CPSF2	CFT2/YDH1		β -CASP
CPSF73	CPSF3	BRR5/YSH1	Endonuclease	β -CASP
hFip1	FIP1L1	FIP1	RNA-binding	
CPSF30	CPSF4	YTH1	RNA-binding	Zinc Finger
CstF				
CstF77	CSTF3	RNA14	Dimerization	HAT repeat
CstF64	CSTF2	RNA15	RNA-binding (DSE)	RRM
CstF64 τ	CSTF2T	PTI1?	RNA-binding (DSE)	RRM
CstF50	CSTF1			WD40 repeat
CF Im				
CF Im 68	CPSF6		RNA-binding	RRM
CF Im 59	CPSF7		RNA-binding	RRM
CF Im 25	CPSF5		Dimerization, RNA-binding	Nudix
CF IIm				
hPcf11		PCF11		CID
hClp1	HEAB	CLP1	RNA kinase	GTPase
Other Factors				
PAP	PAPOLA	PAP1	Poly(A) Polymerase	RNA-binding, NTP
Neo-PAP	PAPOLG		Poly(A) Polymerase	RNA-binding, NTP
Star-PAP	TUT1	CID1 (<i>S. pombe</i>)	Poly(A) Polymerase	RNA-binding, NTP, RS
PABPN1			Poly(A) tail length control	RRM
PABPC1		PAB1	Poly(A) binding	4 RRM
PABPC4		PAB1	Poly(A) binding	4 RRM
RNAP II large subunit		RPB1		CTD
PP1	PPP1CC	GLC7	Phosphatase	PP2Ac
Rbbp6	PACT	Mpel		DWNN, Zinc finger, RING
Homologues of Other Yeast Polyadenylation Factors				
Unknown		HRP1	RNA-binding	2 RRM
Unknown		REF2	RNA-binding	
Unknown		SYC1		CPSF73-100_C
Wdr82		SWD2		WD40 repeat
Ssu72		SSU72	CTD phosphatase	Ssu72

sequences, thereby providing additional RNA-protein contacts that may stabilize the specific interaction between CPSF160 and the A(A/U)UAAA element.

Protein-Protein Interactions of CPSF

CPSF subunits interact with one another extensively^{1,2,13} (Figure 2). CPSF100, 73, and symplekin

have recently been suggested to form the core of CPSF complex while other subunits join the complex as peripheral factors.³⁹ Furthermore, CPSF recruits other components of the 3' processing complex through direct physical interactions (Figure 2). For example, CPSF can be isolated in a pre-assembled complex with CstF, and this association is mediated

by multiple factors, including CPSF160, hFip1, and symplekin^{31,32,36} (Figure 2). Wdr33 may also be involved in bridging CPSF and CstF as its yeast homolog Pfs2p interacts with factors that are homologous to subunits of the CPSF and CstF complexes.⁴⁰ These interactions allow co-operative binding of CPSF and CstF to A(A/U)UAAA and the DSE.^{1,2,13} Following cleavage, CPSF remains bound to the A(A/U)UAAA element and anchors PAP to the RNA for polyadenylation through direct interactions between CPSF160, hFip1, and PAP.^{32,36}

CPSF also interacts with a variety of regulatory proteins to modulate 3' processing either globally or in a transcript-specific manner. For example, the influenza protein NS1 represses the 3' processing of host cellular pre-mRNAs by inhibiting CPSF activity via direct interaction with CPSF30.⁴¹ In another example, following stress, HSF1 not only activates transcription of heat shock protein (Hsp) genes, but also promotes the 3' processing of Hsp transcripts through its association with symplekin.⁴² Finally, the RRM-containing floral repressor FCA interacts with CPSF through FY, the plant homolog of Wdr33, to control flowering time at least in part by modulating the 3' processing of specific transcripts.⁴³

CPSF in the Catalysis of Cleavage

CPSF73 has been identified as the endonuclease responsible for the cleavage step based on several lines of evidence.^{44,45} First, CPSF73 can be UV crosslinked to the cleavage site in the RNA substrate.⁴⁴ Secondly, CPSF73 contains a metallo-beta-lactamase domain and a β -CASP domain.⁴⁵ Some members of the β -CASP protein family, such as Artemis, are known endonucleases.⁴⁶ Thirdly, recombinant CPSF73 shows zinc-dependent endonuclease activity.⁴⁵ It is noted that the endonuclease activity of the recombinant CPSF73 is very weak and stimulation by other 3' processing factor(s) may be necessary for efficient cleavage. Although CPSF100 is structurally highly similar to CPSF73, it lacks the zinc-binding motif and, as a result, does not possess endonuclease activity.⁴⁵ The functions of CPSF100 remain unclear.

The endonuclease activity of CPSF73 was also revealed by studies on metazoan histone pre-mRNA 3' processing.⁴⁷ Directed by unique *cis*-elements, the 3' processing of metazoan histone pre-mRNAs involves only a cleavage step without polyadenylation.⁴⁸ Recent studies suggest that some CPSF subunits, including CPSF73, 100 and symplekin, function in this process and CPSF73 functions as the endonuclease for cleavage.^{47,49} Intriguingly, CPSF73 has been suggested to play additional role in histone pre-mRNA 3' processing as the exonuclease for degradation of the downstream RNA.⁵⁰



FIGURE 3 | Domain structures of CstF 64 and CstF 64 τ . Rectangular boxes represent individual domains, which include RRM (RNA recognition motif), hinge domain, Pro/Gly (proline/glycine-rich domain), 12/8 \times MEARA/G (12/8 tandem copies of MEARA/G repeats), and CTD (C-terminal domain). (Reprinted with permission from Ref 1. Copyright 1997 Cold Spring Harbor Laboratory Press)

CstF

CstF recognizes the DSE and is specifically required for cleavage but apparently not for polyadenylation.^{1,2,13} CstF is comprised of three subunits, CstF77, CstF50 and either CstF64 or its paralog CstF64 τ .^{51,52} Recent studies suggest that CstF may function as a dimer in the 3' processing complex.^{53,54}

CstF in RNA Binding

The RNA-binding activity of CstF is mediated by CstF64/ τ .⁵¹ CstF64 contains an N-terminal RRM domain (Figure 3), which on its own binds with high affinity to GU-rich sequences similar to those found in natural DSEs.⁵⁵ The intact CstF complex, however, binds to PAS-containing pre-mRNAs very weakly as shown by both UV crosslinking and gel mobility shift assays.²⁹ Stable association of CstF to DSE requires co-operative binding of CPSF to the upstream A(A/U)UAAA element.²⁹ Although the exact mechanism is not well understood, this type of obligatory co-operative binding may ensure that 3' processing complex assembles only when both the A(A/U)UAAA and DSE are present and the spacing between the two elements is appropriate.

CstF64 τ shares the same domain structure as CstF64 (Figure 3), but seems to have different RNA-binding specificity.⁵⁶ CstF64 τ is highly expressed in the testis and has been proposed to mediate testis-specific poly(A) site choice.⁵² CstF64 τ knockout mice showed specific defects in spermatogenesis and male infertility,⁵⁷ suggesting that the CstF64 τ -mediated pre-mRNA 3' processing plays important roles in the testis. Therefore, the relative abundance of CstF64 and related proteins may be involved in regulating tissue-specific APA. Both CstF64 and CstF64 τ seem to be evolutionarily conserved from yeast to human¹⁴ (Table 1).

Protein–Protein Interactions of CstF

CstF77 is the scaffold holding the CstF complex together as it interacts with both CstF64 and CstF50

via its proline-rich domain, while CstF64 and CstF50 do not interact with each other^{31,58} (Figure 2). In addition, CstF77 also contains a so-called Half a TPR (HAT) domain. Crystal structures revealed that the CstF77 HAT domain forms a homodimer.^{53,54} This is consistent with earlier observations that both CstF77 and its yeast homolog RNA14p can self-associate.^{31,59} In addition, CstF50 may also dimerize.³¹ Together, these results provided strong evidence that CstF functions as a dimer in the 3' processing complex.

In addition to the aforementioned RRM domain, CstF64/ τ contains a 'hinge' domain, a Pro/Gly-rich region, and a highly conserved C-terminal domain (CTD) (Figure 3). Embedded within the Pro/Gly-rich region is 12 tandem copies of MEARA/G (mouse and human) or 11 copies of LEPRG (chicken) pentapeptide repeats, but the functions of these repeats as well as the entire Pro/Gly-rich domain remain unclear.^{1,2,13} The CstF64 'hinge' domain mediates its interactions with CstF77 and its CTD binds to Pcf11 and the transcription co-activator PC4.^{60–62} As mentioned earlier, both CstF77 and 64 interact with CPSF subunits to facilitate the co-operative binding of CPSF and CstF to pre-mRNAs (Figure 2).

As mentioned above, CstF50 can self-associate and this interaction is mediated by its N-terminal region.³¹ Furthermore, CstF50 contains seven WD-40 repeats, a domain known for mediating protein–protein interactions.^{1,2,13} CstF50 binds to RNAP II CTD and this interaction has been proposed to help recruit CstF to the transcription elongation complex.⁶³ Following UV-induced DNA damage, CstF50 becomes associated with the BRCA1–BARD1 complex and other factors and these interactions are responsible for transiently repressing pre-mRNA 3' processing.⁶⁴

CF Im

Similar to CstF, CF Im is involved in RNA-binding and is required only for the cleavage step.^{1,2,13} CF Im consists of CF Im25 and one of two structurally related proteins, CF Im 59 and CF Im68. Again similar to CstF, CF Im has recently been shown to function as a dimer.^{65,66} There are no clear homologs of CF Im in yeast (Table 1). But it has been proposed that CF Im may be the functional equivalent of the yeast 3' processing factor Hrp1,²² a RNA-binding factor that does not have a clear mammalian homolog.

CF Im in RNA Binding

All CF Im subunits can be UV crosslinked to RNA,⁶⁷ suggesting that they are all involved in RNA recognition. Recombinant CFIm25–68 binds specifically to UGUAN motif and this interaction has

recently been shown to be mediated by CF Im25.^{66,68} CF Im25 contains a Nudix domain and belongs to the Nudix phosphohydrolase superfamily.⁶⁵ However CF Im25 differs from other Nudix hydrolases at several key residues and lacks enzymatic activity.⁶⁵ Instead the CF Im25 Nudix domain functions as a RNA-binding domain that specifically recognizes the UGUAN motif.⁶⁶ CF Im59 and 68 both have a RRM domain, but only bind to RNA in the presence of CF Im25.⁶⁹ Currently it is unclear what RNA sequences CF Im59 and 68 recognize and what the functional significance of their RNA-binding activities is. CF Im–RNA interactions help improve the 3' processing efficiency at canonical PASEs.^{22,70} For noncanonical PASEs that lack the A(A/U)UAAA element but contain UGUAN motifs, CF Im can function as the primary RNA-binding factor for 3' processing complex assembly and recruit other 3' processing factors such as CPSF and PAP through direct interactions.²² In keeping with the important roles of CF Im25 in PAS recognition, depletion of CF Im25 by RNAi causes significant changes in the APA profile for many transcripts.⁷¹

Protein–Protein Interactions of CF Im

In addition to self-association, CF Im25 also interacts with PAP and PABPN1.¹³ As mentioned earlier, both CF Im 59 and 68 have an RRM domain. Interestingly, however, the RRM of CF Im68 is required for its interaction with CF Im25.⁶⁹ CF Im 59 and 68 also contain a C-terminal RS domain that is rich in arginine/serine dipeptide repeats. The RS domain is commonly found in the SR protein family, a large group of proteins best known as splicing regulators.⁷² The RS domain has been shown to mediate RNA-binding as well as protein–protein interactions, including interactions with other SR proteins.⁷² Indeed, CF Im 59 and 68 binds to a number of SR and SR-like proteins,^{69,73} and the CF Im complex has been identified as a component of the spliceosome.⁷⁴ The interactions between CF Im and splicing factors play important roles in coupling splicing and 3' processing.

CF IIm

CF IIm is required only for the cleavage step,^{1,2,13} but its exact functions remain unclear. CF IIm contains at least two subunits, Pcf11 and hClp1, both conserved from yeast to human. Pcf11 homologs in yeast and *Drosophila* are required for both pre-mRNA 3' processing and transcription termination.⁷⁵ Pcf11 binds to the RNAP II CTD in a phosphorylation-dependent manner via its CTD-interacting domain (CID domain).⁷⁵ It has been proposed that RNAP II-bound Pcf11 promotes transcription termination by dismantling the elongation complex.⁷⁶ Additionally,

the yeast Pcf11p has been shown to recruit the mRNA export adaptor Yra1p to the mRNA, thereby linking pre-mRNA 3' processing to mRNA export.⁷⁷

Although its role in pre-mRNA 3' processing remains unknown, hClp1 has recently been identified as an RNA 5'-kinase and it functions as a component of the endonuclease complex in tRNA splicing.^{78,79} The kinase activity of hClp1 is required for maintaining the phosphorylation of the 5'-end of the 3' exon for the subsequent ligation.⁷⁸ It is still unclear, however, if the kinase activity of hClp1 is necessary for pre-mRNA 3' processing. But the recombinant yeast Clp1 protein has no detectable kinase activity,⁸⁰ indicating that the kinase activity may be unique to Clp1 homologs in higher eukaryotes and this activity may not be required for pre-mRNA 3' processing itself.

Poly(A) Polymerases

PAP is perhaps the best-characterized 3' processing factor so far.^{1,2,13} A nucleotidyltransferase catalytic domain occupies the N-terminal half of PAP and is highly conserved (Figure 4). A RNA-binding domain is located near the middle of the protein. RNA binding by PAP alone, however, is not sequence-specific and its recruitment to 3' processing complex requires interactions with other 3' processing factors including CPSF and CF Im.^{1,2,13} The C-terminal domain (CTD) of PAP contains a bipartite nuclear localization signal (NLS) and is rich in serine and threonine residues. The CTD is a hot spot for post-translational modifications and plays important regulatory roles. First, PAP CTD

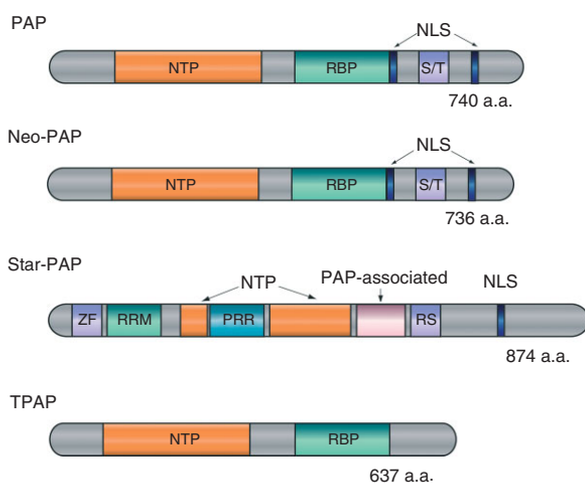


FIGURE 4 | Domain structures of the three nuclear mammalian PAPs. Rectangular boxes represent individual domains, which include NTP (Nucleotidyltransferase domain), RBP (PAP RNA-binding domain), NLS (nuclear localization signal), S/T (serine/threonine-rich domain), ZF (zinc finger domain), RRM (RNA recognition motif), PRR (proline-rich domain), and RS (RS domain). (Reprinted with permission from Ref 85. Copyright 2008 Nature Publishing Group)

is hyperphosphorylated during mitosis by the mitosis promoting factor (MPS, p34^{cdc2}/cyclin B) and, as a result, PAP activity is strongly inhibited.⁸¹ Repression of PAP activity is part of the cellular mechanism for blocking protein synthesis during mitosis. Secondly, PAP CTD binds to 14-3-3 ϵ in a phosphorylation-dependent manner and this association inhibits PAP activity and redistributes PAP to the cytoplasm.⁸² Thirdly, PAP CTD is acetylated by the CREB-binding protein (CBP) and acetylation inhibits its association with CF Im25 and its nuclear localization.⁸³ Finally, PAP CTD is sumoylated at multiple sites, and sumoylation stabilizes PAP, promotes its nuclear localization, but inhibits its enzymatic activity *in vitro*.⁸⁴

In addition to the canonical PAP, metazoans have at least three additional nuclear poly(A) polymerases, neo-PAP, Star-PAP and TPAP (Figure 4). Encoded by an intronless gene, TPAP is specifically expressed in the testis and can be found in both nucleus and the cytoplasm.⁸⁶ The functions of TPAP remain poorly understood. Neo-PAP shares the same overall domain organization with PAP and its *in vitro* polyadenylation activity is indistinguishable from that of the canonical PAP.^{87,88} Neo-PAP can be incorporated into functional human 3' processing complex *in vitro*,¹⁴ but its *in vivo* functions have not been characterized.

Star-PAP has a very different domain structure (Figure 4). It contains a zinc finger domain and a RRM near its N-terminus, a split catalytic domain in the middle, and a PAP-associated domain as well as a RS domain in the C-terminal one third.⁸⁵ Star-PAP was originally identified as a terminal uridylyl transferase (TUTase) specific for U6 snRNA.⁸⁹ The putative homolog of Star-PAP in fission yeast, Cid1, also has TUTase activity.⁹⁰ A later study, however, demonstrated that Star-PAP preferentially uses ATP as substrate and mainly functions as a poly(A) polymerase.⁸⁵ Star-PAP functions in a complex with the type I phosphatidylinositol 4-phosphate 5-kinases (PIP1 α) as well as subunits of the CPSF to control the 3' processing of a subset of transcripts.⁸⁵ Interestingly, the activity of Star-PAP is directly regulated by phosphatidylinositol-4,5-bisphosphate (PtdIns4,5P).⁸⁵ Compare to *Saccharomyces cerevisiae* which only has one nuclear PAP, the presence of multiple nuclear PAPs in metazoans may allow for more elaborate regulation of pre-mRNA 3' processing under different conditions.

Poly(A)-Binding Proteins

There are at least five PABP proteins in humans, one nuclear form (PABPN1) and four cytoplasmic ones (PABPC1, 3–5)^{28,91} (Figure 5). The nuclear and the

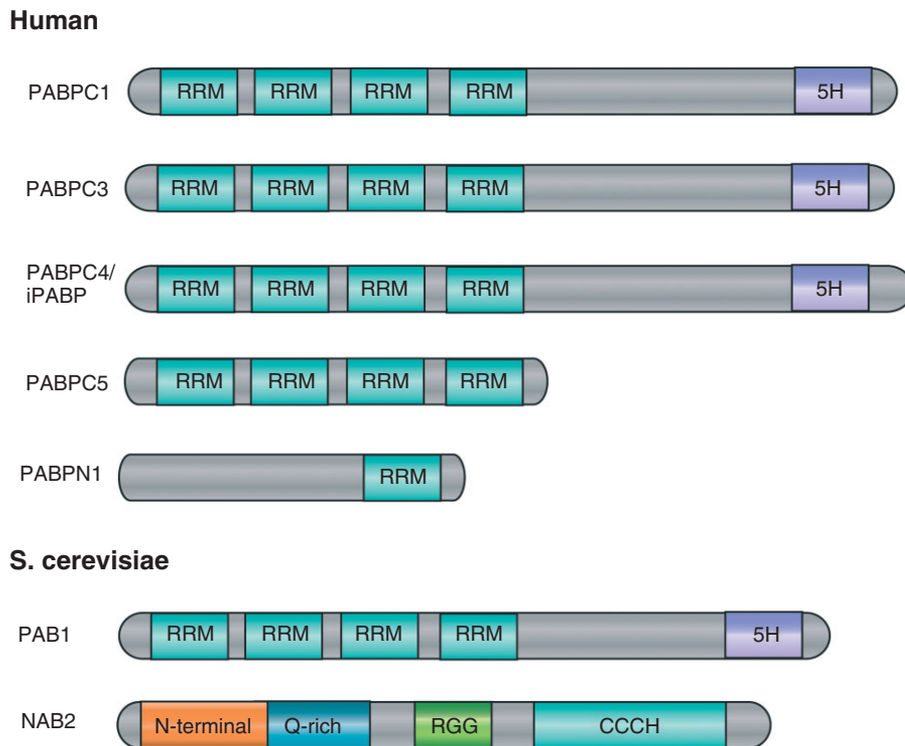


FIGURE 5 | Domain structure of yeast and human PABPs. Rectangular boxes represent individual domains, which include RRM (RNA recognition motif), 5H (a unique 5 conserved helices domain), RGG (RGG box involved in RNA binding), N-terminal (N-terminal domain), Q-rich (Q-rich domain) and CCCH (zinc finger domain). (Reprinted with permission from Ref 28. Copyright 2003 BioMed Central)

cytoplasmic PABPs have distinct domain structures. It has been shown that PABPN1 stimulates PAP activity and plays an important role in poly(A) tail length control.⁹² Following cleavage, PAP, anchored to the pre-mRNA by CPSF, initiates polyadenylation in a slow and distributive reaction.⁹² Binding of PABPN1 to the newly synthesized poly(A) tail stabilizes the polyadenylation complex and dramatically stimulates PAP activity. As a result, polyadenylation switches to a fast and processive phase. PABPN1 coats the entire length of poly(A) tail as it emerges from PAP. When the poly(A) tail reaches ~250 nt in length, polyadenylation is terminated by a poorly understood but PABPN1-dependent mechanism.⁹² PABPs have also been implicated in promoting mRNA export.^{28,91} In the cytoplasm, PABPs play critical roles in translation by promoting 5' → 3' interaction of the mRNA and stimulating initiation.^{28,91}

The budding yeast has only one PABP gene (PAB1) (Figure 5), which is the ortholog of the mammalian cytoplasmic PABPs.^{28,91} The budding yeast does express a structurally distinct nuclear poly(A) binding protein Nab2p (Figure 5), which is probably the functional equivalent of PABPN1. Both Nab2p and Pab1p have been implicated in poly(A) tail length control and mRNA export.^{28,91}

RNA Polymerase II C-Terminal Domain

In addition to transcribing genes, RNAP II also plays critical roles in coordinating co-transcriptional pre-mRNA processing.^{93–95} This function is mediated by its CTD,⁶³ a unique domain consisting of multiple repeats of the conserved heptapeptide YSPTSPS. The number of the repeat ranges from 26 in yeast to 52 in mammals. *In vitro*, RNAP II CTD is required for cleavage in the absence of transcription.⁹⁶ Although the exact functions of CTD in pre-mRNA 3' processing are unclear, it may promote the assembly of a stable 3' processing complex through its extensive interactions with other 3' processing factors, including CPSF, CstF, and CF IIm.^{63,93–95,97}

PRE-mRNA 3' PROCESSING COMPLEX: ASSEMBLY AND DYNAMICS

In vivo, pre-mRNA 3' processing factors are intimately connected with the transcription machinery and the 3' processing complex assembles on pre-mRNAs co-transcriptionally.⁹³ CPSF joins the transcription machinery as early as in the preinitiation complex stage through interactions with TFIID.⁹⁸ CPSF, CstF, and CF IIm are all associated with RNAP II

PRE-mRNA 3' PROCESSING COMPLEX: REGULATION

Like other steps of the gene expression pathway, pre-mRNA 3' processing is subject to regulation.^{1,2,4} Many examples of 3' processing regulation have been reported and the different modes of regulation can be classified into the following three general mechanisms.

Regulation of the Basal 3' Processing Factors

Changes in the protein levels, accessibility, or the activities of the basal 3' processing factors can influence pre-mRNA 3' processing and their effects can be global or transcript-specific.

First, changes in the protein levels of basal 3' processing factors can regulate 3' processing. For example, CstF64 is specifically up-regulated when primary B cells are induced to differentiate.^{106,107} The limited CstF complexes in primary B cells preferentially bind to the high affinity distal poly(A) site of the IgM pre-mRNA, resulting in the production of the membrane-bound form of IgM. However, in activated B cells in which CstF is more abundant, 3' processing shifts to a 'first-come, first-serve' mode and a lower affinity proximal poly(A) site is now preferentially recognized, leading to synthesis of the secreted form of IgM.^{106,107} It is highly likely that many more transcripts are regulated in a similar fashion by the changing levels of CstF64. Additionally the protein levels of CF Im subunits show developmental stage-specific variations and may cause changes in 3' processing efficiency as well as APA patterns.¹⁰⁸

Secondly, changes in the accessibility of the basal 3' processing factors can regulate 3' processing. For example, regulatory viral or cellular proteins could sequester the basal 3' processing factors and prevent them from participating in 3' processing. As mentioned earlier, the influenza virus NS1 protein specifically interacts with CPSF30, thereby inhibiting CPSF-RNA interaction and cleavage/polyadenylation of cellular pre-mRNAs.⁴¹ As mentioned earlier, CstF50 becomes associated with the BARD1/BRCA1 complex following UV-induced DNA damage and this association leads to transient inhibition of pre-mRNA 3' processing.⁶⁴

Finally, changes in the activity of the basal 3' processing factors can regulate 3' processing. Multiple post-translational modifications have been shown to modulate the activities of 3' processing factors. As mentioned earlier, hyperphosphorylation of PAP during mitosis represses PAP activity and leads to general inhibition of polyadenylation.⁸¹ Other modifications could have more profound effects. For example, sumoylation affects the stability, subcellular

localization, and activities of PAP.⁸⁴ CPSF73 and symplekin are also sumoylated and sumoylation stimulates the assembly of the 3' processing complexes.¹⁰⁹

Regulation of 3' Processing Complex Assembly by Local RNA Context and Regulatory Factors

First, 3' processing at a specific PAS can be affected by regulatory factors bound to adjacent *cis*-elements. For example, 3' processing is activated by an upstream intron.¹¹⁰ This stimulation is mediated by multiple interactions between 3' processing factors and splicing factors, such as the CF Im-U2AF and CPSF-U2 snRNP interactions, and their co-operative binding to the PAS and the upstream splicing *cis*-elements.^{73,111} Conversely, 3' processing is strongly inhibited by a 5' splice site located upstream of a PAS.¹¹² This is due to inhibition of PAP activity by U1-70K and U1A of the U1 snRNP bound to the upstream 5' splice site.¹¹² Another example of this type of regulation is provided by the 3' processing of the HIV pre-mRNA at the 3' LTR.¹¹³ The SR protein 9G8, in a complex with CDK11 and eIF3f, binds to a specific sequence upstream of the AAUAAA hexamer. Through its interaction with CF Im bound to an adjacent UGUAN motif, 9G8 stimulates 3' processing at the downstream PAS. Interestingly, overexpression of eIF3f or just its N-terminal 91 amino acid fragment disrupts the eIF3f-CDK11-9G8 complex and prevents efficient 3' processing at the 3' LTR, leading to reduced viral replication.¹¹³

Secondly, some regulatory factors could directly compete with basal 3' processing factors for binding to specific *cis*-elements within the PAS, thereby inhibiting 3' processing. For example, both the mammalian PTB (polypyrimidine tract-binding protein) and the *Drosophila* Sxl (sex-lethal) proteins can compete with CstF64 for binding to the DSE.^{114,115} The outcome of this competition is determined by the affinities of these factors for the DSE and their relative abundance.

Regulation of Pre-mRNA 3' Processing After Complex Assembly

Although regulation of pre-mRNA 3' processing most often occurs at the earliest step, i.e., the 3' processing complex assembly, it should be pointed out that later steps are also subject to regulation. For example, U1A protein regulates its own expression through modulating the 3' processing of its own transcripts.¹¹⁶ U1A proteins bind to a specific RNA structure in the 3' UTR of its own transcripts just upstream of the PAS. The RNA-bound U1A specifically inhibits PAP activity

and therefore prevents polyadenylation without affecting 3' processing complex assembly or cleavage. This forms a negative feedback loop that helps maintain the steady state levels of U1A protein.¹¹⁶

CONCLUSION

Despite the tremendous progress in the past decades, many key questions remain regarding the basic mechanisms of pre-mRNA 3' processing complex assembly and function as well as its regulation. First, what are the defining features of a PAS? Although a number of key *cis*-elements have been identified, one or more of these elements are missing in many PASEs. In the coming years, the key elements in the noncanonical PASEs will undoubtedly become better defined and the mechanisms for their recognition by the 3' processing machinery will be revealed. It will be of great interest to see whether there are any unifying features for all PASEs. Secondly, what are the functions

of each 3' processing factor? Systematic structure-function analyses of individual 3' processing factors as well as the entire 3' processing complex will provide new insights into the molecular mechanisms of pre-mRNA 3' processing, including RNA recognition, complex assembly, catalysis, and complex disassembly. Finally, as shown by recent studies,^{7–11} APA is a widespread phenomenon in higher eukaryotes. It will be critical to understand how 3' processing complex assembly is regulated at alternative poly(A) sites and what is the impact of APA under physiological as well as pathological conditions. Instead of focusing on one transcript or one 3' processing regulator at a time, it is becoming increasingly important to take a systems approach to investigate how 3' processing is regulated at the transcriptome level. Coupling high-throughput sequencing- and microarray-based genomic analyses with biochemical characterization will be critical in this effort.

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