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The human PAF complex coordinates transcription with events downstream of RNA synthesis

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The yeast PAF (yPAF) complex interacts with RNA polymerase II and coordinates the setting of histone marks associated with active transcription. We report the isolation and functional characterization of the human PAF (hPAF) complex. hPAF shares four subunits with yPAF (hCtr9, hPaf1, hLeo1, and hDcd73), but contains a novel higher eukaryotic-specific subunit, hSki8. RNAi against hSki8 or hCtr9 reduces the cellular levels of other hPAF subunits and of mono- and trimethylated H3-Lys 4 and dimethylated H3-Lys 79. The hSki8 subunit is also a component of the human SKI (hSKI) complex. Yeast SKI complex is cytoplasmic and together with Exosome mediates 3′-5′ mRNA degradation. However, hSKI complex localizes to both nucleus and cytoplasm. Immunoprecipitation experiments revealed that hPAF and hSKI complexes interact, and ChIP experiments demonstrated that hSKI associates with transcriptionally active genes (Pokholok et al. 2002). Therefore, yPAF was postulated to be involved in both RNA polymerase II-mediated transcription initiation and elongation. Recently, the yPAF complex has been shown to be required for Rad6-mediated histone H2B-K123 monoubiquitination (Ng et al. 2003b; Wood et al. 2003), Set1-mediated histone H3-K4 methylation (Krogan et al. 2003; Ng et al. 2003a), and Dot1-mediated histone H3-K79 methylation (Krogan et al. 2003), placing PAF at the center of cotranscriptional histone modifications (for review, see Hampsey and Reinberg 2003). PAF subunits are also required for maintaining proper poly(A) tail length in yeast, suggesting a post-transcriptional role for PAF complex in mRNA processing and maturation (Mueller et al. 2004).

Cotranscriptional events include not only chromatin modifications but also RNA processing and surveillance. Cotranscriptional RNA processing events such as 5′-end capping, splicing, and polyadenylation are relatively well documented (for reviews, see Maniatis and Reed 2002; Neugebauer 2002; Orphanides and Reinberg 2002). In recent years, evidence for cotranscriptional RNA surveillance, particularly in higher eukaryotes, emerged as well. Higher eukaryotes developed the process of alternative splicing to generate multiple mRNAs from one gene. This requires RNA surveillance to ensure high fidelity of gene expression. In mammals, nonsense-mediated decay (NMD) occurs not only in the cytoplasm, but also in the nucleus (Humphries et al. 1984; Takeshita et al. 1984; Cheng and Maquat 1993). We found that hSKI localizes to transcriptionally active genes in an hPAF-dependent manner. This novel link between hPAF and hSKI supports the hypothesis that the hPAF complex coordinates events downstream of RNA synthesis such as RNA surveillance.

Results and Discussion

hPAF complex contains a novel higher-eukaryotic-specific subunit, hSki8

A cDNA encoding hPaf1 was originally isolated as PDX, a gene that is amplified and overexpressed in human pancreatic adenocarcinoma (S.K. Batra, unpubl.). We frac-

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The PAF complex was first identified in yeast (yPAF) as an RNA polymerase II-associated factor, consisting of five subunits, Pa1, Ctr9, Leo1, Rfl1, and Cdc73 (Shi et al. 1996; Krogan et al. 2002; Mueller and Jaehning 2002). The yPAF complex interacts with TBP (Stolinski et al. 1997), the elongation factors Spt4–Spt5, and FACT (Costa and Arndt 2000; Krogan et al. 2002; Squazzo et al. 2002). Chromatin immunoprecipitation (ChIP) experiments revealed the presence of the yPAF complex at both promoter and coding regions of transcriptionally active genes (Pokholok et al. 2002). Therefore, yPAF was postulated to be involved in both RNA polymerase II-mediated transcription initiation and elongation. Recently, the yPAF complex has been shown to be required for Rad6-mediated histone H2B-K123 monoubiquitination (Ng et al. 2003b; Wood et al. 2003), Set1-mediated histone H3-K4 methylation (Krogan et al. 2003; Ng et al. 2003a), and Dot1-mediated histone H3-K79 methylation (Krogan et al. 2003), placing PAF at the center of cotranscriptional histone modifications (for review, see Hampsey and Reinberg 2003). PAF subunits are also required for maintaining proper poly(A) tail length in yeast, suggesting a post-transcriptional role for PAF complex in mRNA processing and maturation (Mueller et al. 2004).

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tionated nuclear extracts derived from HeLa cells through conventional chromatography following the elution of hPaf1/Pd2 using antibody directed against Pd2 in Western analyses. The polypeptides coeluting with hPaf1 were identified by MALDI-TOF mass spectrometry. Besides hPaf1, three other human homologs of the yeast Rtf1 homolog exists in the database, which shares 21% sequence identity and 40% similarity with the yeast Rtf1 (Supplementary Fig. 2). We designated this protein as hRtf1 and established a stable cell line expressing Flag-hRtf1. Affinity-purified hRtf1 did not display any subunits of the hPAF complex (Fig. 1A; Supplementary Fig. 3A). These results demonstrate that unlike in yeast, hRtf1 is not a subunit of the hPAF core complex. Fractionation of affinity-purified hRtf1 on a gel filtration column suggested that hRtf1 has a native mass of 200 kDa and it did not coelute with any other polypeptide (Supplementary Fig. 3B).

Our results are consistent with the studies of Rozenblatt-Rosen et al. (2005) demonstrating the existence of hCtr9, hLeo1, hPaf1, and hCdc73 as hPAF subunits and the apparent loss of hRtf1 as an integral subunit to the hPAF complex. We believe the difference is likely due to the purification approaches. In their report, hPAF was affinity-purified with antibodies against hCdc73 and eluted with acidic glycine. This procedure inevitably elutes large amounts of IgG that likely masked the low-molecular-weight region of the SDS/PAGE gel. This would not impinge on the identification of subunits larger than IgG, but may have obscured the presence of hSki8, the only hPAF subunit smaller than the IgG heavy chain.

Since the yPAF complex includes Rtf1 and interacts with RNA polymerase II, we analyzed whether the hPAF complex and hRtf1 interact with RNA polymerase II. Immunoprecipitation using antibody against hPaf1 specifically enriched for the subunits of the hPAF core complex, such as hCtr9, hLeo1, and hCdc73 as well as substoichiometric amounts of RNA polymerase II and hRtf1. Similarly, antibodies against RNA polymerase II and hRtf1 coimmunoprecipitated substoichiometric amounts of each other as well as subunits of the hPAF core complex (Supplementary Fig. 4). These results establish interactions between RNA polymerase II, the hPAF core complex, and hRtf1.

**hSki8** is also a subunit of the hSKI complex, and hPAF interacts with hSKI in vivo

Little is known about hSki8; it is denoted after the yeast homolog Ski8. Yeast Ski8, together with Ski2 and Ski3, form a trimeric complex that is required for 3′-to-5′ mRNA decay (Ridley et al. 1984; Masison et al. 1995; Brown et al. 2000). To further analyze the association of hSki8 with the hPAF complex, we established a stable cell line expressing Flag-hSki8. Affinity purification from the stable Flag-hSki8 cell line revealed the presence of the entire hPAF complex, further confirming hSki8 as an integral subunit (Fig. 1C). Interestingly, distinct higher-molecular-weight polypeptides were observed, apparently specific to the Flag-hSki8 complex, as they were absent in the purified Flag-hPaf1 and Flag-hLeo1 complexes (Fig. 1A, C). MALDI-TOF mass spectrometry analysis identified all the hPAF subunits as well as the homologs of the yeast Ski2 and Ski3 proteins (for sequence alignments, see Supplementary Fig. 2). There-
Therefore, we designated the high-molecular-weight proteins as hSki2 and hSki3. Gel filtration of the affinity-purified hSki8 sample revealed two distinct complexes, one corresponding to the hPAF complex and the other exhibiting the coelution of three polypeptides: hSki2, hSki3, and hSki8 (Fig. 1D). We denoted this second complex as hSKI. Under stringent conditions (0.5 M KCl) and using antibodies against hPaf1, we were able to deplete the hPAF complex from the affinity-purified Flag-hSki8. A gel filtration column revealed that the hPAF-depleted material contained a trimeric complex composed of hSki2, hSki3, and hSki8 eluting with an apparent native mass of 400 kDa (Supplementary Fig. 3C).

The discovery of a novel subunit of the hPAF complex that is shared with the hSKI complex prompted us to investigate whether these two complexes interact. We performed immunoprecipitation experiments using nuclear extract derived from the Flag-hPaf1 stable cell line with antibodies against Flag. The results demonstrated substoichiometric amounts of hSki3 immunoprecipitated by anti-Flag antibody. Interestingly, whereas the extract (input) displayed apparently different isoforms of hSki3, the immunoprecipitated material contained only one isoform (Fig. 1E). Importantly, a reciprocal immunoprecipitation experiment using antibodies against hSki3 demonstrated immunoprecipitation of endogenous hPaf1 from nuclear extract (Fig. 1E). Thus, hPAF and hSKI complexes do interact.

**Figure 2.** hSKI complex localizes to both cytoplasm and nucleus of HeLa cells. The upper panel shows staining of DAPI, hSki3, and α-tubulin with corresponding merges. The lower panel shows the competition experiment with 10 µg of purified recombinant proteins Ski3 or Paf1.

**Figure 3.** RNAi targeting hSki8 or hCtr9 knocks down the protein levels of other hPAF subunits and reduces cellular levels of histone H3-K4 trimethylation, monomethylation, and H3-K79 dimethylation. (A) Western blots for various proteins and histone modifications with whole cell lysate from mock RNAi, hSki8 RNAi, and hCtr9 RNAi samples. (B) Quantification of the Western blots.
plasm, a significant amount of hSki3 signal was also localized within the nucleus; we also noticed that the hSki3 signal appears to be excluded from the nucleolus (Fig. 2). As a control we used antibodies against α-tubulin, which localizes exclusively to the cytoplasm (Fig. 2). Importantly, preincubation of the hSki3 antibodies with excess recombinant hSki3 protein, but not recombinant hPaf1 protein, inhibited the signal for hSki3, demonstrating that the hSki3 immunofluorescence is specific (Fig. 2, lower panels). In addition, hSki8 localized to both the cytoplasm and nucleus, while hPaf1 appears to be predominantly nuclear with a weak signal in the cytoplasm [Supplementary Fig. 5]. This suggested that the hSKI complex may associate with transcriptionally active genes [see below] and mediate RNA surveillance.

hSki8 is required for PAF-dependent mono- and trimethylation of histone H3-K4 and dimethylation of H3-K79 in vivo

To understand the functional importance of the hPAF complex in vivo, we used RNAi against two of the hPAF subunits, hCtr9 and hSki8. Interestingly, RNAi directed against either hCtr9 or hSki8 not only knocked down the levels of the hPAF complex such as hPaf1 and hLeo1. As controls, the levels of actin and the subunit of the TFIIF transcription factor Rap74 were unaffected [Fig. 3]. These results are consistent with another report showing that deletion of individual yPAF subunits leads to various degrees of reduction in the protein levels of the other subunits [Mueller et al. 2004].

We then investigated the histone methylation marks. In line with previous studies in yeast, RNAi against hCtr9 or hSki8 led to a significant reduction in the cellular levels of trimethylation of histone H3-K4 and dimethylation of histone H3-K79, but not in dimethylation of H3-K36. As controls, the total levels of histone H3 and methylation marks associated with repression (trimethylation of H3-K9 and H3-K27) were not affected. Interestingly, when the status of the different methylation marks for H3-K4 were analyzed, we found no significant reduction in dimethylation, but significant reductions in mono- and trimethylation [Fig. 3]. This is probably due to the existence of multiple histone H3-K4 methyltransferases in humans [for review, see Margueron et al. 2005] and the distinct roles for the H3-K4 methylation marks. Trimethylated H3-K4 is well documented to be associated with transcription (Santos-Rosa et al. 2002; Schneider et al. 2004), but little is known about H3-K4 monomethylation. Our results suggest that histone H3-K4 monomethylation may also be a mark associated with active transcription.

hSKI complex is present at transcriptionally active genes with recruitment dependent on hPAF complex

Given the nuclear localization of a fraction of the hSKI complex and the physical interaction between hSKI and hPAF complexes, we next analyzed if the hSKI complex, likely a part of the RNA decay machinery, may function as a transcription-related factor. ChIP experiments using primers directed to the promoter and the downstream regions of the RPB1 gene demonstrated that the hSKI complex subunits (hSki3 and hSki8) are present at both the promoter and coding regions, generating a pattern similar to that observed for the hPAF complex subunits (hPaf1, hLeo1, and hSki8) [Fig. 4A]. Importantly, while siRNA against the hPAF subunit hCtr9 did not affect hSki3 or hSki8 protein levels [Fig. 3], the ChIP signals for hSki1 and hPAF were both similarly reduced. On the other hand, TFIIB and RNA polymerase II (Rpb4) signals remained the same [Fig. 4A].

To further analyze the presence of the human SKI complex at transcriptionally active genes, the inducible MAGE-A1 gene was chosen. The expression of the MAGE family [melanoma antigen genes] is regulated by DNA methylation. These genes are heavily methylated and are transcriptionally inactive in most human tissues and cell lines, while the DNA demethylating agent 5-aza-deoxycytidine [5-aza-dC] induces the expression of MAGE genes in various cells [De Smet et al. 1996]. In the HEK293 cells used in this study, the MAGE-A1 gene is also known to be methylated and silent [Fig. 4B, B. Zhu and J.P. Jost, unpubl.]. Expression of the MAGE-A1 gene was observed after 48 h of 5-aza-dC treatment [Fig. 4B]. Like hPAF complex, hSki1 was absent at the methylated and transcriptionally inert MAGE-A1 gene. However, upon induction, hSki1 was recruited to the promoter and coding sequences of the MAGE-A1 gene, along with other factors of the transcription machinery. This hSki1 recruitment was again reduced after knock-down of the hPAF complex by siRNA targeting hCtr9 [Fig. 4B].

These data collectively suggest that hSki1, a protein complex that is required for 3′-to-5′ RNA decay in yeast [Maison et al. 1995; Brown et al. 2000], associates with transcriptionally active genes via its interaction with the hPAF complex in the case of human cells. Notably, Exosome, the other complex that is required for RNA 3′–5′ decay, was also reported to be associated with transcriptionally active genes via its interaction with the transcription elongation factor Spt6 [Andrulis et al. 2002].

Figure 4. hSki1 complex is present at transcriptionally active genes, and its recruitment is dependent on hPAF. [A] Human SKI complex is present at promoter and coding regions of the RPB1 gene and hPAF knock-down reduces the signal for hSki1 complex. [B] RT–PCR showing the induction of MAGEA1 expression upon 5-aza-dC treatment. [C] Human SKI complex is recruited to the MAGE-A1 gene upon its induction, together with the RNA polymerase II transcription machinery. hSki1 recruitment is hPAF-complex-dependent.

Characterization of the human PAF complex

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Interestingly, Exosome occupies both the promoter and coding regions as well (Andrulis et al. 2002). Taken together, the human SKI complex shares many common features with the Exosome complex as both likely compose the 3→5′ RNA decay machinery in higher eukaryotes, interact with transcription factors, and occupy transcriptionally active genes. Therefore, we suggest that the hSKI complex, together with the Exosome, likely regulate cotranscriptional events including mRNA quality control. That a portion of hSKI complex is present in the nucleus and that human-derived SKI and PAF complexes interact are distinct differences with the case in yeast. This may signify a novel additional function(s) for hPAF and hSKI complexes or that higher eukaryotes evolved to integrate transcription and surveillance more efficiently.

The PAF complex: elongation factor or modulator of RNA metabolism?

Several findings have demonstrated interactions between transcription elongation factors and either the yeast (Costa and Arndt 2000, Krogan et al. 2002, Squazzo et al. 2002) or the human PAF complexes [B. Zhu and D. Reinberg, unpubl.]. Yet, Jaehning and coworkers found that the loss of the PAF complex in yeast does not alter the distribution of RNA polymerase II or transcription elongation factors such as Spt5 and Spt16 on transcriptionally active genes [Mueller et al. 2004]. Moreover, although the PAF complex was detected in association with randomly selected active yeast genes [Pokholok et al. 2002], suggesting it has a general role in transcription, deletion of the PAF complex (or its subunits) in yeast and plants affected the expression of only a subset of genes [Porter et al. 2002, He et al. 2004; Oh et al. 2004]. These observations collectively suggest that the PAF complex does not function as a classical elongation factor, despite the fact that PAF is associated with the transcriptionally active RNA polymerase II [Mueller et al. 2004; Sims et al. 2004]. Our results and the previously published results with Exosome support the model that mRNA surveillance occurs “cotranscriptionally” rather than “post-transcriptionally.” We favor the hypothesis that the hPAF complex serves as a platform between RNA polymerase II and many other factors modulating RNA biogenesis, in essence modulating cotranscriptional events such as chromatin remodeling, chromatin modifications, mRNA maturation, and surveillance.

Materials and methods

GenBank accession numbers

The GenBank accession numbers are as follows: hPaf1, AJ401156; hLeo1, BC018147; hCdr9, BC058914; hCdc73, NM_024529; hSki8, AK024754; hRtf1, BC015052; hSki3, BC056893; and hSki2, Q15477.

Affinity purification of Rtf1 and the hPAF complex

M2 anti-Flag agarose (Sigma) was equilibrated with the same buffer used in nuclear extract preparation (20 mM Tris-HCl at pH 7.9, 1.5 mM MgCl2, 0.2 M NaCl, 0.2 mM PMSF) and then incubated with nuclear extract derived from various stable cell lines overnight at 4°C. The resin was washed with an excess amount of buffer C containing 0.5 M KCl plus 0.1% NP-40, and then eluted with buffer C containing 0.5 M KCl plus 0.1 mg/mL Flag peptide.

Coimmunoprecipitation

HeLa cell nuclear extracts were dialyzed against buffer C containing 0.1 M KCl, and then incubated with various antibodies cross-linked to protein A agarose. The resin was washed with an excess amount of buffer C containing 0.15 M KCl, and proteins were eluted with 0.1 M glycine-HCl at pH 2.5.

ChIP

ChIP was performed according to the protocol described by Upstate. For every assay, cells were derived from approximately one confluent 15-cm plate.

RNAi knock-down for the hPAF complex

siRNAs targeting hSki8 and hCdr9 were purchased from Dharmaco. siRNAs were delivered using RNAiFect from QIAGEN according to the user manual.

Immunofluorescence

HeLa cells along with monoclonal antibody B-5-1-2 against α-tubulin (Sigma) and affinity-purified rabbit polyclonal antibodies against hSki3, hSki8, and hPaf1 were used in the experiments.

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