Air2p is critical for the assembly and RNA-binding of the TRAMP complex and the KOW domain of Mtr4p is crucial for exosome activation

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ABSTRACT

Trf4/5p-Air1/2p-Mtr4p polyadenylation complex (TRAMP) is an essential component of nuclear RNA surveillance in yeast. It recognizes a variety of nuclear transcripts produced by all three RNA polymerases, adds short poly(A) tails to aberrant or unstable RNAs and activates the exosome for their degradation. Despite the advances in understanding the structural features of the isolated complex subunits or their fragments, the details of complex assembly, RNA recognition and exosome activation remain poorly understood. Here we provide the first understanding of the RNA binding mode of the complex. We show that Air2p is an RNA-binding subunit of TRAMP. We identify the zinc knuckles (ZnK) 2, 3 and 4 as the RNA-binding domains, and reveal the essentiality of ZnK4 for TRAMP4 polyadenylation activity. Furthermore, we identify Air2p as the key component of TRAMP4 assembly providing bridging between Mtr4p and Trf4p. The former is bound via the N-terminus of Air2p, while the latter is bound via ZnK5, the linker between ZnK4 and 5 and the C-terminus of the protein. Finally, we uncover the RNA binding part of the Mtr4p arch, the KOW domain, as the essential component for TRAMP-mediated exosome activation.

INTRODUCTION

Eukaryotic cells produce diverse types of protein-coding as well as noncoding RNAs, some of which have been recently shown to have direct regulatory role in the regulation of gene expression and chromatin structure [reviewed in (1)]. Most, if not all, RNAs are synthesized as a precursor molecule that needs to be posttranscriptionally processed and/or modified in order to form mature functional molecules. Several human genetic disorders relate to abnormalities in RNA processing and proper ribonucleoprotein assembly, such as cancer or neurodegenerative diseases (2). In yeast, nuclear RNA maturation and stability are under strict control from RNA surveillance carried out by the nuclear exosome and its cofactor the TRAMP4/5 polyadenylation complex (3–6).

The TRAMP recognizes a variety of nuclear transcripts produced by all three RNA polymerases, such as precursors of small nuclear and nucleolar RNAs (snRNAs and snoRNAs), ribosomal RNAs (rRNAs), tRNAs, as well as telomeric, and cryptic unstable transcripts (CUTs) (6–8). It adds short poly(A) tails to aberrant transcripts, creating a favorable substrate for the exosome (3–6,9), reviewed in (10). Among the complex subunits, Mtr4p provides helicase activity and Trf4p or Trf5p provide polyA-polymerase (PAP) activity, Air1p and Air2p are predicted to provide RNA-binding. Air proteins contain five tandemly arranged ZnK motifs of the CCHC-type (CX2CX4HX4C) that have previously been shown to...
bind nucleic acids and proteins in unrelated viral proteins (13). The ZnK regions in these proteins recognize exposed guanosines of RNA loops or in single-stranded RNA (13). The RNA helicase Mtr4p is not required for the polyadenylation activity of TRAMP; however, it regulates the length of a poly(A) tail added by Trf4p-Air2p dimer and modulates its polyadenylation rate in vitro (14).

Recently, three works reported on the structure of the full-length Mtr4p and a heterodimer of Trf4p-Air2p fragments (15–17). The structures of Mtr4p revealed a five-domain organization in which four domains constitute the DEh helicase core [RecA-1, RecA-2, winged helix (WH) and a helical bundle] (16,17). Compared to known structures of other DNA and RNA helicases, Mtr4p DEh core has an N-terminal β-hairpin extension located on the surface of the globular core. The most unique feature however is an insertion of 265 amino acids between the WH helices protruding from the globular core to form the arch domain with an arm-like structure and a fist (also termed the KOW domain) at the end. The KOW domain shares similarities with rRNA-binding proteins and contributes to RNA binding in vitro (17). The arch is dispensable for Mtr4 helicase and ATPase activities and for TRAMP integrity in vitro. A strain expressing an archless Mtr4p displays a slow growth phenotype that seems to result from an Rpr6p-dependent incomplete processing of 5.8S rRNA (16).

The study of the structure of the minimal Trf4p-Air2p heterodimer, covering the catalytic and central domains of Trf4p and only the last two ZnKs of Air2p, revealed a phylogenetically conserved surface on the central domain of Trf4p that contacts the last linker and ZnK region of Air2p (15). Nevertheless, this study still left many open questions such as what is the role of the other Air2p ZnK domains and the N- and C-terminal regions of Air2p and Trf4p. Finally, the RNA-binding mode of Trf4p-Air2p has been addressed only indirectly. The structural determination of the entire complex has proven to be difficult so far due to the instability of recombinant subunits. Moreover, it still remains a mystery as to how TRAMP interacts with and activates the downstream acting exosome. Thus despite the advances in our understanding of TRAMP function in vivo and of the individual structures, there is still very limited knowledge on the assembly of the complex and of the mechanistic details of its action.

Here, we have focused on two aspects of the TRAMP4 biochemistry. On one hand, we aimed to identify the code of recognition of structurally complicated substrates that need to be polyadenylated by TRAMP4 before exosome degradation, such as the hypomodified tRNA^{Met}. Second, to tackle the TRAMP4 complex assembly and molecular mechanisms of its function, we have performed detail analyses of Trf4p, Air2p and Mtr4p domain organization in vitro and in vivo. Furthermore, we succeeded to obtain the solution structure of the five zinc-knuckle region of Air2p by NMR spectroscopy. Overall, we uncovered regions required for cell viability, protein–protein interactions, RNA binding domains and regions responsible for exosome activation.

**MATERIALS AND METHODS**

**Yeast cultures and manipulation**

Yeast were cultured under standard conditions in media with selective markers corresponding to particular strains and vectors. Yeast transformations were performed by lithium acetate method. For growth test analyses, yeast were cultured overnight in minimal SD medium at 30°C, diluted with fresh SD media for OD_{600} 0.2 and grown at 30°C to OD_{600} 1. The cells were then serially diluted in decimal dilutions and incubated at 17°C, 25°C, 30°C and 37°C in SD media (or as indicated in each experiment). The yeast strains used in this work are listed in Supplementary Table S2.

**Purification of protein complexes from yeast**

The yeast culture was grown at 30°C in YPD medium to OD_{600} ~2.5. Cells were harvested, frozen in liquid nitrogen and manually grinded. Lysate was melted in D100 buffer [100 mM KCl, 50 mM Tris, 10% (v/v) glycerol, 0.02% (v/v) Nonidet P-40 (NP-40), 0.2 mM EDTA, pH 7.9] with protease inhibitors, soluble fraction was recovered by centrifugation and bound to IgG-sepharose beads for 3 h at 4°C. Beads were washed thoroughly with D150 (same as D100 but 150 mM KCl) or D1000 (high-salt wash, same as D100 but 1 M KCl) buffer, and proteins were eluted by TEV protease cleavage to D150 buffer. Elutions were snap frozen and kept at −80°C. Alternatively, elutions were merged and bound o/n to Ni-NTA agarose beads. The beads were washed with D150 buffer with 10 mM imidazole and eluted by 250 mM imidazole to D50 buffer. Elutions were snap frozen and kept at −80°C.

All immunoprecipitated samples that were subsequently subjected to western blot analysis were prepared in the presence of 100 μg ml⁻¹ of RNase A (added to the cell lysate). Samples that were used for enzymatic assays (polyadenylation, exosome degradation) were prepared in the absence of RNase in order to avoid interference of the residual RNase with the enzymatic analysis.

**RNA isolation and analysis**

For RNA analysis, cells were grown at 30°C o/n in liquid SD media, diluted to OD_{600} 0.25 and grown at 25°C to OD_{600} 1. RNA was isolated by the hot phenol extraction and stored at −80°C. Five micrograms of total RNA was denatured in 25% formamide, separated on 8% polyacrylamide gel and transferred to nylon membrane using semi-dry electro-transfer. RNA was cross-linked to the membrane by UV light (120 mJ cm⁻²) and hybridized with a probe in Ultra-Hyb buffer (Ambion) according to manufacturer’s instructions.

**Construction of yeast plasmids**

Yeast genomic DNA or WT full-length TRF4, AIR2 and MTR4 inserted in pET22b (5) were used to PCR-amplify the untranslated and coding regions of TRF4, AIR2 and MTR4 genes.
**TRF4 constructs.** To obtain plasmids containing truncated forms of the TRF4 gene, the desired region was amplified in a standard PCR reaction using primers listed in Supplementary Table S3 and inserted into pNOPPATA1L vector via restriction sites NdeI and Sall. To obtain point mutated forms, a standard PCR mediated site directed mutagenesis with pNOPPATA1LTRF4 (5) as a template was performed. All constructs contained N-terminal fusion protein A tag and a TEV protease recognition sequence. With the exception of C-terminally truncated forms (that is TRF4 ΔA544, CA499, CA463 and 182–463), the constructs also contained fusion C-terminal hexahistidine anchor. The His6 tag was not used during analyses and we observed no sign of its interference with the results. To express endogenous levels of the Trf4p, the genomic region spanning 500 nt upstream of the TRF4 initiation codon was amplified using primers TRF4 506 5′-UTR XbaI For and TRF4 506 5′-UTR ATG SpeI XhoI Rev and inserted into the pRS413 vector via XbaI and XhoI restriction sites. Into this construct, the TRF4 full-length gene and/or alleles D425A, DXD and/or N496 from pNOPPATA1L constructs including all fusion tags were inserted through SpeI site. For Escherichia coli expression, the TRF4 coding sequence was inserted into the expression vector pET30a(+) allowing for the N-terminal fusion protein A tag and a TEV expression, the TRF4 coding sequence was inserted into the expression vector pET30a(+) allowing for the N-terminal fusion protein A tag and a TEV fusion G-protein B1 tag (18) by using BamHI and XhoI restriction sites.

**AIR2 constructs.** Yeast genomic DNA or WT AIR2 gene inserted in pET22b (5) were used to PCR-ampify the untranslated and coding regions of AIR2. To prepare AIR2 constructs for phenotypic analyses in vivo, the coding sequence of AIR2 surrounded by 425-bp upstream of initiation codon and 479-bp downstream of stop codon was PCR amplified from Saccharomyces cerevisiae genomic DNA (BY4741 background) with primers AIR2 5′-UTR For1 and AIR2 3′-UTR Rev1. The fragment was inserted into the pRS415 vector via PstI and NotI restriction sites resulting in the plasmid V379. To prepare tagged versions of AIR2 for affinity purifications of TRAMP complexes, the full-length CDS of AIR2 was subcloned to a yeast expression vector [pNOPPATA1L, (5)] via NdeI and XhoI restriction enzymes, resulting in the plasmid V53. Single point mutations of AIR2 were prepared with WT AIR2 in pNOPPATA1L as a template by site-directed mutagenesis by using oligonucleotides listed in Supplementary Table S3. AIR2 truncation mutants were constructed by PCR amplification with specific primers listed in Supplementary Table S2. The PCR product was inserted into pNOPPATA1L vector using NdeI and Sall restriction sites. To generate mutants pRS415 derived constructs, we first introduced an NdeI restriction site at the position 3-bp upstream of AIR2 start codon resulting in the construct V394 WT (with primers AIR2 NdeI ATG For and AIR2 NdeI ATG Rev). Individual mutants prepared in pNOPPATA1L were then transferred via NdeI and BamHI sites to generate constructs listed in the Supplementary Table S4.

**MTR4 constructs.** Full-length MTR4 and all deletion variants were cloned into pNOPPATA1L vector. C-terminal His6 tag was added via PCR amplification with an extended reverse primer (MTR4 His Sall). The full-length MTR4 was amplified from genomic DNA (primers MTR4 NdeI and MTR4 His Sall); MTR4 archless was amplified from pAV674 vector (16), which was a kind gift from Dr Van Hoof. ΔKOW MTR4 version was prepared by ligation of two PCR products into pNOPPATA1L vector (using NdeI/Sall restriction sites). N-terminal partial (amplified with primers MTR4 KOW N) of MTR4 (aminoacids 1–665) was extended at the 3′-end by coding sequence for Gly-Ser-Gly-Ala-Ser and C-terminal part of MTR4 (aminoacids 819–1073, obtained with oligos MTR4 KOW C) was extended at 5′-end by coding sequence for Ala-Ser-Gly-Ser-Gly-Ser resulting in GSGSASGSGS bridge (connected via NheI restriction site in Ala-Ser coding sequence).

**Preparation of unmodified tRNAiMet for enzymatic and NMR binding experiments**

The plasmid ptRNAiMet for the in vitro transcription of yeast tRNAiMet (5) was linearized with BstNI. Unlabelled hypomodified tRNAiMet was produced by in vitro T7 polymerase run off transcription (19) using linearized plasmid as a template. The product was purified by anion exchange HPLC under denaturing conditions and desalted by a NAP column (Sephadex G-25). The hypomodified tRNAiMet was denatured at 95°C, pH 6.2 for 5 min and snap cooled on ice, to favor a monomeric conformation. The concentration of RNA was determined by UV-spectroscopy.

**Fluorescence anisotropy**

The equilibrium binding of Air2p ZnK1-5 (aminoacids 57–180) to different oligonucleotides was analyzed by fluorescence anisotropy. The 15-mer poly-A RNA and DNA oligonucleotides were 5′-labeled with N,N′,N′′,N′′′-tetramethyl-6-carboxyrhodamine (TAMRA) that was attached via a hexyl linker. The measurements were conducted on a FluoroMax-4 spectrofluorometer (Horiba Jobin-Yvon, USA). The instrument was equipped with a thermostatted cell holder with a Neslab RTE7 water bath (Thermo Scientific, USA). The system was operated using FluorEssence software (version 2.5.3.0, Horiba Jobin-Yvon, USA). The TAMRA fluorophore was excited at 561 nm, and its emission was collected at 581 nm. The widths of both excitation and emission monochromatic slits were 8 nm and integration time was set to 3 s. During the measurement 10 mM labeled oligonucleotide (volume 1.4 ml) was titrated with increasing amounts of the protein in 50 mM Tris buffer (pH 7.9), supplemented with 250 mM NaCl, 20 mM β-mercaptoethanol and 50 μM ZnSO4. The titrations were carried out at 20°C in a stirred 1.5 ml quartz cuvette. A fixed delay of 30 s was set between each aliquot addition and start of the measurement to allow the reaction to reach equilibrium. This delay was sufficient, as no further change in anisotropy was observable. Every data point is an average.
of three measurements. The data were analyzed in SigmaPlot 11 software (Systat Software, USA). The experimental isotherms were fit to a single-site binding model according to Heyduk and Lee (20) using nonlinear least squares regression. The data were normalized for visualization purposes.

Preparation of DNA probes for northern blot analysis
DNA probes for NEL025c and snR33 were amplified from *S. cerevisiae* BY4741 genomic DNA with the primers combinations (NEL025c For and Rev, and snR33 — mature + extension For and Rev, see Supplementary Table S5) and used as templates for Random Primed DNA Labeling Kit (Roche) according to manufacturer’s instructions. To detect U14 and snR13 3'-extended snoRNAs and tRNA Met, the oligonucleotides were 5'-phosphate labeled using T4 polynucleotide kinase (NEB) and [γ-32P]-ATP.

Whole-cell protein extract preparation for western blot analysis
The yeast culture was grown at 30°C in selective medium to OD600 ~1.0. The cells were resuspended in the D150 buffer with protease inhibitors (0.5 μg ml⁻¹ of leupeptin, 0.8 μg ml⁻¹ of pepstatin A, 0.6 mM PMSF and 0.6 mM DTT). The cell suspension was vortexed with glass beads at 4°C for 10 min. The beads and the cell debris were pelleted by centrifugation, a part of the cell-free extract was mixed with 1×SDS loading buffer for western blot analysis. The rest of the extract was used for protein concentration measurement by the method of Bradford.

Antibodies
The generation of specific antibodies against Trf4p, Air2p, Air1p have been described in (5,7). Mtr4p antibodies were kindly provided by Dr Patrick Linder (University of Geneva). The anti-proteinA antibodies and the anti-His6 antibodies were from Sigma. All antibodies were used at a 1:5000 dilution.

Expression and purification of recombinant proteins
Full-length Trf4p from *S. cerevisiae* was recombinantly expressed from pET30a vector with N-terminally fused G-proteinB1 solubility tag (18) in *E. coli* strain BL21-CodonPlus (DE3)-RIPL (Stratagene). The expression was induced by 0.4 mM IPTG at 37°C for 4 h in the presence of 0.5 mM ZnSO4. Cells were lysed in the lysis buffer (50 mM carbonate buffer, 300 mM NaCl, 5% glycerol, 400 μM ZnSO4, pH 10.5), and all proteins were purified in buffer containing 50 mM carbonate buffer, pH 10.5, 400 mM NaCl, 10% glycerol, 400 μM ZnSO4 using affinity Ni-NTA chromatography followed by gel filtration.

Expression and preparation of Air2 ZnK protein for NMR studies
The coding sequence of Air2 protein comprising residues 57–180 from *S. cerevisiae* was expressed with the N-terminal fusion Smt3 protein (21). The expression was induced by 100 μM IPTG in *E. coli* BL21-Codon Plus (DE3)-RIPL (Stratagene) strain overnight at 16°C in M9 minimal medium, supplemented with 50 μM ZnSO4. For isotope labeling, the medium was supplemented with 15NH4Cl and [U-13C6]glucose. Cells were harvested, resuspended in denaturing buffer (50 mM Tris, 500 mM NaCl, 8 M Urea, pH 8.0) and lysed by sonication, and proteins were purified using affinity Ni-NTA chromatography. The protein immobilized on beads was extensively washed and subsequently refolded in buffer containing 50 mM Tris pH 7.9, 500 mM NaCl, 20 mM β-mercaptoethanol, 5 mM imidazole, 50 μM ZnSO4, and protease inhibitors. Refolded protein was eluted and dialyzed against buffer containing 50 mM Tris pH 7.9, 500 mM NaCl, 20 mM β-mercaptoethanol and 50 μM ZnSO4. The Smt3 fusion protein was cleaved by Ulp1p protease and removed on Ni-NTA column. For NMR measurements the protein was concentrated using a Vivaspin 20 concentrator (Sartorius).

NMR spectroscopy and structure determination
The protein used in NMR spectroscopy was concentrated to ~150 μM. Due to a high salt buffer [50 mM sodium phosphate buffer, 500 mM NaCl, 10 mM β-mercaptoethanol pH 8.0 (90% H2O/10% D2O)], 3 mm or shaped NMR tubes were used. All NMR experiments were recorded at 20°C on a Bruker AVANCE 600 MHz and 900 MHz spectrometers equipped with a cryogenic triple resonance probe (Bruker BioSpin). The chemical shifts of backbone and side-chains were assigned using standard triple resonance experiments (22,23). All distance restraints were derived from the 3D 15N- and 13C-edited NOESYs (with mixing time of 80 ms) collected at 900 MHz spectrometer. The structure calculations were performed with the automated NOE assignment module implemented in the CYANA program (24). Initial structures were further refined with the CYANA-generated restraints along with the restraints for the tetrahedral zinc coordination with AMBER 10.0 software (25) as described previously (26,27). From 40 refined structures, the 20 conformers with the lowest AMBER energy were selected to form the final ensemble of structures.
Enzymatic assays

Polyadenylation assays were carried out in 15 μl reaction mixtures containing 5–50 ng of affinity purified complex, 50 fmol 5’-end labeled RNA, 0.5 mM ATP, 5 mM MgCl2, 25 mM Tris, pH 7.9, 20 mM KCl, 10% glycerol, 0.1 mg ml−1 BSA, 1 mM DT, 0.02% Nonidet P-40, 5U of RNasin Plus (Promega) with or without addition of 50 ng of affinity purified Rrp6-TAP eluate (5). Reaction mixture was incubated at 30°C, and 5 μl aliquots were collected at times indicated. The reaction was stopped by the addition of 25 mM EDTA and proteins removed by incubation with ProteinaseK (Promega) for 10 min at 37°C. Sample was mixed with one volume of formamide loading buffer and resolved on denaturing polyacrylamide gel.

Database searches and sequence analysis

BLAST searches were conducted by using the NCBI genome databases (http://www.ncbi.nlm.nih.gov/BLAST/) or other nonredundant or EST databases at the Swiss EMBl node (28). The multiple sequence alignment was performed by using ClustalW2 (29) at the http://www.ebi.ac.uk/Tools/services/web_clustalw2 interface. The Boxshade server (http://www.ch.embnet.org/software/BOX_form.html) was used to color the similarity in the alignment.

RESULTS

Identification of Air2p regions required for intermolecular contacts with Trf4p and Mtr4p

Trf4/5p, unlike canonical PAPs, possess no identifiable RNA-binding domain and thus are inactive on their own. We have previously demonstrated that Air2p activates Trf4p, likely through its RNA-binding properties (5). To identify Air2p regions that contact RNA, interact with Trf4p or Mtr4p, and are required for yeast viability, we have prepared Air2p mutants carrying point mutations in the second zinc-coordinating cysteine in each of the five ZnKs (ZnK1 to ZnK5) to disrupt the ZnK fold (Figure 1A and Supplementary Figure S1A). In order to tackle the importance of the surrounding N- and C-terminus, we generated deletion mutants lacking either the N- or C- terminus or both (Figure 1A). Mutants ZnK4, ZnK5, CA180 and ZnK1-5L were not able to rescue growth defect of the air1Δair2Δ strain at higher temperature and mutants in the first three ZnKs and NΔ57 showed reduced fitness (Figure 1B). The growth defects observed at 30°C were not caused by reduced expression or stability of Air2p variants nor other TRAMP subunits, as the western blot analysis of protein extracts prepared from cells grown at 30°C revealed comparable protein levels for the mutants tested (Supplementary Figure S1B and S1C). This pointed to the importance of the last two zinc knuckles and the Air2p C-terminus. Our previous yeast two-hybrid results indicated the minimal region in Air1 and Air2 proteins required for Trf4p interactions positioned across the zinc knuckles ZnK3 to ZnK5 (5). Accordingly, in NMR titration experiments, we observed that the fourth and the fifth ZnKs of Air2p associated with recombinant Trf4p (data not shown).

Furthermore, we observed chemical shift perturbation in the third (L3) and fourth linker (L4) regions upon Trf4p binding. It concerned mainly residues 117–120 (L3) and residues S138, I139, W140, R141 and Y143 (L4), most of them being highly conserved between Air homologs from distant species (Supplementary Figure S1A). The analysis of additional mutants in those regions revealed that double mutants in the last two ZnKs ZnK4+5 and in the fourth linker (W140G+R141E), respectively, exhibited poor fitness comparable to the WT strain at all temperatures tested (Figure 1B). The growth defects observed at 24°C correlated with deficiencies in TRAMP function in vivo as northern blot analyses revealed a strong accumulation of NEL025c CUT and snR33 pre-snoRNA in ZnK4+5, L4, CA180 and ZnK1-5L mutants cultivated at 24°C (Figure 1C). Cold sensitivity and to a certain extent RNA degradation defects seen at 24°C were observed also in all the other ZnK point mutants indicating that all ZnKs contribute to some degree to TRAMP activity in vivo (Figure 1B and C).

To address whether the in vivo phenotypes reflected defects in TRAMP assembly or catalytic activity, we isolated and analyzed yeast complexes containing the individual mutated Air2p versions. In agreement with our NMR studies, L4 and ZnK5 mutants failed to efficiently copurify Trf4p and Mtr4p, whereas the other point mutations did not affect TRAMP4 assembly (Figure 1D). Western blot analysis of the input fractions of individual Air2p mutants revealed comparable levels of all three TRAMP subunits in WT, ZnK and L4 mutant strains. Thus the lack of Trf4p and Mtr4p copurification with L4 and ZnK5 was not caused by their decreased stability or expression in these mutant strains. As the absence of Mtr4p does not disrupt the interaction with Trf4p (Figure 1D, HSW) (4,5), we conclude that the fourth linker and last zinc knuckle regions of Air2p are crucial for the binding to Trf4p. The lack of Trf4p and Mtr4p copurification in ZnK5 samples was reflected in the lack of PAP activity of the purified protein sample (Figure 2A). Interestingly, although the ZnK4 mutant was able to maintain the TRAMP integrity (Figure 1D), it was strongly defective in polyadenylation in vitro (Figure 2A). Accordingly, recombinant Air2 proteins mutated in ZnK4 or ZnK5 failed to activate yeast Trf4p in vitro (Figure 2B).

We have previously reported that purified TRAMP4 complex activates exosomes in vitro (5). In order to test whether some of the mutants may affect exosome activation, we performed coupled polyadenylation/ degradation assays. In this assay, the 5’-end labeled hypomodified tRNAiMet (a well-defined in vivo target of this pathway) is incubated in the presence of WT or mutant TRAMPs and yeast nuclear exosome purified via the Rrp6p subunit (5). Figure 2C shows Rrp6p-TAP purified fractions that were used in all subsequent degradation assays in this work compared to exosome obtained via the core subunit Rrp4p. The coupled polyadenylation/ degradation assays revealed that none of the polyadenylation-competent Air2p point mutants further affected exosome activation in vitro (Figure 2D).

The analysis of deletion mutants indicated the importance of the first 25 N-terminal amino acids for the
interaction with Mtr4p. We observed that both the N\(\Delta\)57 proteins did not co-purify Mtr4p (Figure 1D, lanes N\(\Delta\)57 and ZnK1-5L). Interestingly, the N\(\Delta\)57 protein pulled down significantly higher amounts of Trf4p than WT even in the presence of 1M KCl (HSW). At present, we do not know the reason for higher yields of Trf4p in this mutant. These results were not due to significantly altered levels of Mtr4p nor Trf4p in these mutants (Supplementary Figure S1C). In addition, the deletion of the last 139 amino acids weakened the interaction with both Mtr4p and Trf4p (Figure 1D, samples CA180 and ZnK1-5L).

Air2p is a genuine RNA-binding protein

To further investigate why TRAMP4 mutant in Air2p ZnK4 lacks PAP activity, we studied the central region of Air2p (ZnK1-5, residues 57-180) by NMR spectroscopy. The Air2p was purified under denaturing conditions and refolded on Ni-NTA column in the presence of N-terminal Smt3 fusion protein. We first confirmed, that this fragment is sufficient to activate the PAP activity of Trf4p (Figure 2B, first lane). For NMR experiments, a high-salt concentration of 500 mM NaCl was used to prevent the protein aggregation. Although this protein construct suffered from a very low solubility limit (150 \(\mu\)M), we were able to achieve \(1H, 13C\) and \(15N\) resonance assignments of ZnK1-5 and determine its three-dimensional structure (Figure 3A). The chemical shift deviations of C\(\alpha\) and H\(\alpha\) of the assigned backbone resonances of ZnK1-5 from the sequence-dependent random coil values show no pattern for the ZnK1-5 region. This agrees well with the absence of secondary structure elements in the...
The topology of zinc knuckles (13, 22). The histidines of the CCHC motif of ZnKs, namely H71, H90, H109, H131 and H172, are involved in coordination of zinc ions as the N-H correlations of these residues span the same region in the 2D $^1$H-$^15$N HSQC (Figure 3B) that is characteristic for zinc knuckles (22). Furthermore, the presence of folded zinc knuckles was confirmed by the addition of EDTA in excess of zinc ions that resulted in a loss of resonance dispersion in the 2D $^1$H-$^15$N HSQC (data not shown). The NOESY spectra suffered from a low signal-to-noise ratio as a result of low solubility of the ZnK1-5 protein and the presence of high-salt buffer. Therefore, the resulting NMR structure is of a low resolution (Supplementary Table S1, Supplementary Figure S2). The structure of this Air2p region consists of five independent zinc knuckles that all adopt a canonical zinc knuckle fold of the CCHC type. The domains are connected by flexible linkers, and no inter-domain contacts have been found (Figure 3A). The conserved IWRxY motif of linker 4 is also unstructured in the free form, whereas it forms a short $\alpha$-helix upon binding to Trf4p (15).

To investigate whether Air2p is a genuine RNA-binding protein, we performed a quantitative solution-binding assay.
Figure 3. Structure and RNA-binding of ZnK1-5 Air2p. (A) The lowest energy structure of the Air2p ZnK1-5. The protein is shown as a ribbon model (in red) with the zinc-coordinating residues (CCHC) shown in yellow. The zinc ions are shown in blue. (B) Close-up view of the $^1$H-$^{15}$N TROSY spectrum, showing the N-H correlations of histidines that are involved in the coordination of zinc ions. (C) Air2p ZnK1-5 binds A15 RNA. Binding isotherms for equilibrium binding of ZnK1-5 Air2p to fluorescently labeled A15 RNA, monitored by fluorescence anisotropy. (D) Air2p ZnK1-5 binding to dA15 DNA. The ionic strength and pH of the binding buffer was the same for both measurements. The dissociation constant ($K_d$) was calculated from the best fit of data using a single-site binding isotherm. (E) $^1$H-$^{15}$N TROSY spectra of Air2p ZnK1-5 alone (in blue) and in the presence of one equivalent of hypomodified tRNA$^{Met}$ (in red; $^1$H-$^{15}$N HSQC) at 20°C. (F) Summary of chemical shift perturbations and line broadening of Air2p ZnK1-5 upon binding to hypomodified tRNA$^{Met}$. Affected residues (qualitatively described by perturbation index: yes = 1, no = 0) are plotted against the amino-acid residue number. The assignments of residues indicated by asterisks could not be obtained. (G) Solvent-accessible surface representation of the RNA-binding ZnK of Air2p (ZnK2, ZnK3 and ZnK4) colored by electrostatic potential (blue, positive; red, negative) of the representative structure.
assay using fluorescence anisotropy (FA) experiments. We found that Air2p ZnK-5 binds unstructured A15 RNA with a $K_D$ in a low micromolar range (Figure 3C), whereas the binding affinity to dA15 DNA could not be detected in our experimental setup ($K_D >> 100 \mu M$) (Figure 3D). Next, we used NMR spectroscopy to reveal which ZnK domains of Air2p are used to bind to hypomethylated tRNA$^\text{Met}$, a substrate that is used for in vitro assays in this work. The chemical shift changes and resonance broadening observed in the $^1$H-$^{15}$N HSQC spectrum of this work. The chemical shift changes and resonance that are unstructured in the free form, become structured as well. This suggested that these highly conserved linkers that are unstructured in the free form, become structured upon RNA-binding akin to the linkers of viral nucleocapsid (NC) proteins (13). Our attempts to compare binding of Air2p to hypomethylated tRNA$^\text{Met}$ and native tRNA$^\text{Met}$ (the latter being a poor substrate for TRAMPs) failed, as we were not able to prepare sufficient quantities of native tRNA$^\text{Met}$ from yeast for NMR experiments.

**The minimal functional Trf4p**

The recent crystal structure of truncated Trf4p bound to a fragment of Air2p encompasses only the conserved catalytic (CAT) and central domains (CD) of Trf4p (residues 161-481) (15). We observed that these regions alone (fragments 182-463 or 161-481) are not functional in vivo as they were not able to rescue viability of the double deletion strain $^\text{trf4}_4/^\text{trf5}_5$ (Figure 4A and C and Supplementary Figure S3A and S3B). To search for the minimal functional Trf4p and to address the role of individual Trf4p domains, we prepared a set of deletion and point mutant constructs of $^\text{TRF4}$ (Figure 4B). Deletions were designed to remove poorly conserved, potentially unstructured parts of the protein. In addition to the catalytically inactive DXD mutant (D236A and D238A) (5), we introduced two point mutations: (i) D425A residing in a highly conserved aspartate residue in the central domain (30), which is not present in canonical PAPs, and (ii) K552A, which has been proposed to reside in the catalytic center for Trf4p dRP lyase activity (31).

Deletions of the entire N- (NΔ181) or C- (CΔ463) termini surrounding the CAT and CD parts of Trf4p did not support viability of the double trf4$^\Delta$/trf5$^\Delta$ strain (Figure 4C, Supplementary Figure S3A and S3B) and exhibited an accumulation of three well-characterized TRAMP substrates, NEL025c CUT and precursors of SnR13 and U14 snoRNAs when expressed in trf4Δ strain (Figure 4D). Although we were able to detect the expression of these fragments by western blot (Supplementary Figure S3C), our attempts to purify these mutants to assess the interaction with other TRAMP components and their activities failed. Thus, we hypothesize that the deletion resulted in protein misfolding and instability. The analyses of the rest of the deletion mutants indicated the minimal functional Trf4p region (spanning residues 97-499) containing CAT and CD of Trf4p surrounded by short extensions as NΔ96 and CΔ499 were sufficient for TRAMP4 assembly (Supplementary Figure S3D). In vivo, CΔ499 displayed cold sensitivity and impaired growth at 30°C when expressed in trf4$^\Delta$/trf5$^\Delta$ deletion background (Figure 4C, Supplementary Figure S3A and S3B). When expressed in the single trf4Δ strain, CΔ499 along with NΔ96 were able to suppress the RNA degradation/processing defects of the trf4Δ in vivo (Figure 4D). TRAMPs purified via proteinA-tagged NΔ96 and CΔ499 showed WT-like PAP activities (Figure 4E) and were able to activate tRNA$^\text{Met}$ degradation in the presence of Rrp6p-TAP exosomes in vitro (Figure 4F).

In agreement with previously published data, the catalytically inactive DXD TRF4 mutant suppressed only partially the NEL025C accumulation (6,7). This implies that TRAMP can also act in polyadenylation independent manner in vivo. In addition to DXD, we have identified D425A as the second point mutant with impaired PAP activity (Figure 4F). Compared to DXD, the D425A exhibited very weak, only above the background accumulation of TRAMP targets seen by northern blot analysis (Figure 4D). Moreover, it was able to support growth of trf4$^\Delta$/trf5$^\Delta$ strain (Figure 4C). The purified TRAMP4 containing the point mutation in D425 failed to polyadenylate hypomethylated tRNA$^\text{Met}$ in vitro under standard assay conditions and accordingly to activate the exosome (Figure 4E and F), although the complex integrity was not affected (Supplementary Figure S3D). We found that increasing ATP concentration resulted in a gradual increase of the PAP activity in the D425A mutant up to the 4mM concentration tested, whereas it did not change any further activity of the saturated WT protein (Figure 4G). This result indicated that the mutation reduced Trf4p affinity for ATP. We propose that lowered ATP substrate affinity of D425A slows down PAP activity of the complex, which is however sufficient for TRAMP4 function in vivo.

**The Mtr4p-arch domain is required for TRAMP4 complex-mediated exosome activation in vitro**

We have observed that in the absence of Air proteins the stability of the Trf4p-Mtr4p heterodimer is compromised even under low salt conditions (Figure 5A) implying that Air proteins mediate interaction between Trf4p and Mtr4p. This is in agreement with similar recent findings in *Schizosaccharomyces pombe* (32), suggesting a conserved mode of Mtr4p interaction within TRAMP in two distant yeast species. Based on sequence conservation and molecular surface scanning, several patches were proposed as possible binders for Trf4p-Air2p (17). To identify the interaction surface of Mtr4p with the rest of the TRAMP complex, we prepared a set of terminal and internal deletion variants N-terminally fused with cleavable 2×proteinA tag (Figure 5B). As expected, the mutants in the conserved core were not able to support growth in the absence of WT Mtr4p, while the removal of the first 143 amino acids (NΔ143) resulted in strong growth defect (Figure 5C). The archenal mutant shows reduced growth as reported previously (16).
Interestingly, the deletion of the KOW domain was sufficient to reproduce this phenotype (Figure 5D).

Using affinity purification from *S. cerevisiae* and subsequent western blotting, we confirmed the previous results from *in vitro* (17) showing that the arch domain is also dispensable for TRAMP formation *in vivo* (Figure 5E). Furthermore, we observed the full complex formation after deletion of the unusual RecA-1N-terminal β-hairpin extension (NΔ143) found in the structure (17), which is not conserved outside of the Ski2 protein family.
Deletion of any of the domains that form the helicase core, led to the loss of co-purification of TRAMP components (Figure 5E). Nevertheless, we cannot rule out the possibility of misfolding or cellular mislocalization. Since the base area of the helical core around the RNA exit tunnel was proposed to serve for exosome interaction
(16), we narrowed down the interaction site with the Trf4p/Air2p heterodimer to the side of the barrel formed by RecA-2, winged helix and helical bundle domains. Since Mtr4p is dispensable for TRAMP polyadenylation activity but is required for the activation of the exosome (4,5), we tested whether the loss of N-terminal extension or arch domains had any effect on the ability of the TRAMP complex to activate degradation in coupled in vitro polyadenylation and exosome assay. While the NA143 mutant did not show any deviation from WT, we observed almost complete loss of exosome activation in the presence of the archless mutant on two different structured substrates (Figure 5F, Supplementary Figure S4A and S4B). To investigate the importance of the KOW domain, the RNA binding element of the arch (17), we conducted the assay in the presence of Mtr4p variant lacking this domain. As shown in Figure 5G, removal of the KOW domain recapitulated the inability of the archless mutant to efficiently activate exosome in vitro. Since we observed apparently lowered polyadenylation in KOWless TRAMPs, we next compared the exosome activation in the presence of lowered amounts of WT Mtr4p. Despite the resulting reduction in observed polyadenylation activity, the WT Mtr4p TRAMP still activated Rtp6p-TAP, unlike KOWless mutant (Supplementary Figure S4C). This together with the slow growth phenotypes (Figure 5D) suggested that arch and particularly the KOW domains are crucial for exosome activation.

DISCUSSION

We have identified several key features of the assembly, RNA binding and exosome activation of the yeast TRAMP RNA surveillance complex. While we focused on the analysis of the TRAMP4 complex, due to the similar composition of TRAMP5 (33) and high level of sequence conservation between Trf5p and Trf4p, and between Air1p and Air2p (5), data presented here are likely valid also to TRAMP5.

Dual function of Air1/2p in the TRAMP complex

Here, we demonstrate that Air2p is a bona fide RNA-binding protein. The Air2p ZnKs 2, 3 and 4 bind aberrant tRNA (Figure 3E and F); however, only the fourth ZnK is fully essential for TRAMP PAP activity in vitro (Figure 2A and B). Although other study by Hamill et al. (15) proposed its involvement in RNA binding, our analyses have not revealed any specific role for ZnK1. It is possible that ZnK1 binds other RNAs, not used in our tests. Alternatively, it may be involved in the interaction with other TRAMP-associated proteins. Structural and sequence alignment analyses indicate that the RNA-binding ZnKs of Air2p use a different binding mode compared to the NC proteins. They lack the hydrophobic pocket, which accommodates guanine base in the NC proteins. Instead, they have large electropositive patches (Figure 3G) created out of lysines, arginines and serines. We speculate that these residues could be responsible for the recognition of a broad spectrum of RNA substrates based on a low-sequence specificity interactions with the sugar-phosphate backbone.

The analysis of deletion mutants indicated the importance of the first 25 N-terminal amino acids for the interaction with Mtr4p, as both the NAΔ57 (residues 57–344) and ZnK1-5L (residues 26–205) proteins did not co-purify with Mtr4p (Figure 1D, lanes NAΔ57 and ZnK1-5L). The N-terminus is poorly conserved among Air1/2p homologs. In fact, a metazoan Air1-like protein (also designated ZCCHC7) contains a long N-terminal extension (data not shown). Given the observation that the human Mtr4p (SKIV2L2) forms the NEXT (Nuclear EXosome Targeting) complex independently of PAPD5/ZCCHC7 (the human homologs of Trf4p/Air1/2p) (34), it is possible that the interaction between noncanonical PAP (Trf4p/Air2p) and Mtr4p helicase has diverged during evolution depending on the organism complexity.

On the other hand, the Trf4p interacting residues in Air2p identified by NMR titration experiments at positions 139–143, 141G and R141E reside in a stretch of amino acids forming the (P/T)xIWRxYxL motif conserved among Air-like proteins from distant species including humans (Supplementary Figure S1A). The protein pulldown experiments revealed strong defects in Trf4p and Mtr4p copurification with mutants in this conserved motif (L4 mutant) as well as in the mutant with the disrupted last fifth zinc knuckle fold (ZnK5) (Figure 1D). Mtr4p is not required for Trf4p–Air2p interaction. It is likely that dimerization of Air2p and Trf4p is needed for efficient assembly with Mtr4p, as the absence of either one results in loss of Mtr4p copurification (Figures 1D and 5A). Therefore, we believe that linker 4 and ZnK5 of Air2p are crucial regions responsible for interaction with Trf4p. The last ZnK also displays the highest degree of conservation among the five Air ZnKs (Supplementary Figure S1A). Likewise, the Trf4p surface interacting with the ZnK4–ZnK5 region is conserved among other Trf4-like proteins (15). This implies a phylogenetically conserved mode of interaction between Air1/2p homologs and Trf4p-like noncanonical PAPs. Although the human PAPD5 does not apparently require a cofactor for PAP activity in vitro (35), it does form a complex with ZCCHC7 and other putative RNA-binding proteins (34), suggesting that Air-like proteins mainly serve as RNA substrate specificity factors. During the preparation of this manuscript, Fasken and co-workers (36) reported similar results on Air1p as regard to the importance of the last two zinc knuckles for cell viability and IWRxY motif conservation, NEL025C degradation and Trf4p interaction (36). Our analysis goes beyond this work by distinguishing the different key roles of ZnK4 in RNA binding and Trf4p activation (see below) and ZnK5 in Trf4p interaction. Fasken and colleagues found that Air1p and Air2p mutants in ZnK5 are unstable at 37°C and to a lesser extend at 25°C, which is likely the reason of strong growth defects at high temperatures. Our western blot analysis of expression levels of the different Air2p variants shows only slight decrease in ZnK5 Air2p mutant at 25°C (Supplementary Figure S1B). Moreover, our in vitro reconstitution assays (Figure 2B) suggest that...
this mutant is unable to form functional Trf4p-Air2p PAP even in amounts comparable to WT. Thus we conclude that the absence of Trf4p in ZnK5 immunoprecipitates is not caused by lowered \textit{in vivo} expression of Trf4p, but due to weakened interactions between the two proteins as a response to the disruption of the fifth Air2p zinc knuckle. Furthermore, we show that Air2p mediates the interaction between Trf4 and Mtr4 proteins (Figure 5A). Our work extends to the analysis of the N- and C- termini of Air2p revealing crucial interactions with Mtr4p. The Air2p termini enhance Mtr4p binding, while the main surface for interaction with Trf4p is located in the fifth ZnK and the adjacent linker 4 regions (Figure 1D). Based on these mapping analyses and the prediction that the bottom of the Mtr4p helicase barrel serves for interaction with exosome (16), we propose a model of the quaternary interactions within the TRAMP complex. In this model, Trf4p/Air2p binds to the Mtr4p DEExH globular core, at the side of barrel formed by RecA-2, winged helix and helical bundle domains. We suggest that the RNA substrates are first screened by the Air2p ZnK2 – ZnK4 region that protrudes from Mtr4 surface and are consequently loaded to the Trf4p active site. We speculate that the RNA 3'-oligo(A) terminus leaves Trf4p in the direction toward the KOW-RecA-2 interface, where the RNA 3'-end is expected to enter the helicase core (17) (Figure 6).

TRF4 catalytical mechanism

We found that increasing ATP concentration resulted in a gradual increase of the PAP activity in the D425A mutant up to the 4 mM concentration tested, whereas it did not change any further activity of the saturated WT protein (Figure 4G). This result indicated that the mutation reduced Trf4p affinity for ATP.

By examining the available crystal structure (15), we observed that the D425 residue is situated approximately 20 Å from the active site. Close inspection of the mechanism of the homologous yeast canonical poly(A) polymerase Pap1p catalysis revealed that hydrogen bonding in this region is required for stabilization of conformational movements leading from an open to closed conformation of the protein (37). When the N189A (CAT domain) mutation is introduced in Pap1p, the equilibrium between the open and closed state is shifted, resulting in increased \( K_M \) for ATP (37). N189 forms a hydrogen bond with the main chain carbonyl of the Y307 (CD) in the closed conformation (Supplementary Figure S3E and S3F). Interestingly, the Y307 of Pap1p resides in both, sequentially (30) and spatially, identical position to D425 of Trf4p (Supplementary Figure S3G). Therefore, we suggest that the side chain of D425 may serve as a hydrogen bond acceptor analogous to Y307 main chain carbonyl. We have not identified any hydrogen bond formed by D425 in the published structure [(15) and Supplementary Figure S3G]. Notably, the structure was solved for the Trf4p apoenzyme; hence it is likely to represent the open state status. We, therefore, suggest that a conformational change analogous to Pap1p may occur in Trf4p upon RNA and ATP binding, thus indicating the conservation of the catalytic mechanism of both related proteins.

RNA binding KOW domain is involved in exosome activation

The role of the arch domain in exosome function \textit{in vivo} has been proposed previously by Jackson \textit{et al} (16) who observed the accumulation of 5.8S+30 rRNA and stabilization of 5'-ETS, both being specific substrates of Rrp6p activity (38). Here we report that the arch, particularly the KOW domain, of Mtr4p is necessary for the TRAMP-mediated exosome activation \textit{in vitro}. We show that TRAMPs lacking arch or KOW are able to add poly(A) tails, but they almost completely fail to activate exosomes to degrade structured RNAs \textit{in vitro} (Figure 1F and G; Supplementary Figure S4). Recently Jia \textit{et al} (14) showed that Mtr4p regulates the individual steps of TRAMP polyadenylation as well as the length of the poly(A) tails added. We do not exclude the possibility that arch/KOW is also involved in this regulation. However, in our assays, N\(\Delta\)143 and \(\Delta\)KOW TRAMPs showed comparable polyadenylation, but only N\(\Delta\)143 was fully able to initiate exosomal degradation, whereas \(\Delta\)KOW was highly defective. Therefore, we conclude that the observed differences in polyadenylation seen in KOW-less TRAMP cannot explain the lack of exosome activation, and we propose that the KOW domain is crucial for TRAMP-mediated exosome activation.

As neither the helicase nor ATPase activities of Mtr4p are affected by arch removal (16,17), we speculate that the RNA-binding properties of the arch might be necessary to handle the substrate between different activities of the TRAMP-exosome pathway. Both the close proximity of

![Figure 6. Model of TRAMP assembly and RNA binding. The Air2 protein is bridging the Mtr4p (interaction via N- and C-terminus) and Trf4p (interaction via L4, ZnK3 and C-terminus). The ZnKs 2, 3 and 4 are involved in RNA binding. The KOW domain of Mtr4p is required for exosome activation.](https://academic.oup.com/nar/article-abstract/40/12/5679/2414880)
the KOW to the helicase RNA entry site and the possible flexibility in the elbow of the arm (16) suggest that arch transports RNA between Trf4p-Air2p, the helicase core and potentially the Rrp6p. Alternatively, the arch is required for the displacement of other RNA-binding factors and shuttling the substrate to the helicase core. We also cannot exclude the possibility that the arch serves for protein–protein interactions with Rrp6p or other core exosome components. This scenario is supported by the presence of a similar structure recently uncovered in the cytoplasmic exosome cofactor Ski2p (16,17,39).

The nuclear exosomes contain two confirmed exoribonucleases—the Rrp6p and Rrp44p. Currently, it is still not fully understood how these two subunits interplay on divergent exosome substrates in vivo. However, it was shown that in the conditions used in our in vitro assays (5 mM Mg2+) the nuclease Rrp44p is inactive (12). Therefore, we propose that nuclease activity observed in our assays can be fully attributed to Rrp6p. Therefore, it will be interesting to address whether RNAs bound by KOW can also undergo processing by the core exosome or whether they are specifically triggered for the degradation by Rrp6p. The exact mechanistic details remain obscure and will be the subject of following studies.

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REFERENCES

DEsH helicase involved in nuclear RNA processing and surveillance. Proc Natl Acad Sci USA, 107, 12139–12144.