Distinct roles of Pcf11 zinc-binding domains in pre-mRNA 3′-end processing

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ABSTRACT

New transcripts generated by RNA polymerase II (RNAPII) are generally processed in order to form mature mRNAs. Two key processing steps include a precise cleavage within the 3′ end of the pre-mRNA, and the subsequent polymerization of adenosines to produce the poly(A) tail. In yeast, these two functions are performed by a large multi-subunit complex that includes the Cleavage Factor IA (CF IA). The four proteins Pcf11, Clp1, Rna14 and Rna15 constitute the yeast CF IA, and of these, Pcf11 is structurally the least characterized. Here, we provide evidence for the binding of two Zn2+ atoms to Pcf11, bound to separate zinc-binding domains located on each side of the Clp1 recognition region. Additional structural characterization of the second zinc-binding domain shows that it forms an unusual zinc finger fold. We further demonstrate that the two domains are not mandatory for CF IA assembly nor RNA polymerase II transcription termination, but are rather involved to different extents in the pre-mRNA 3′-end processing mechanism. Our data thus contribute to a more complete understanding of the architecture and function of Pcf11 and its role within the yeast CF IA complex.

INTRODUCTION

Polyadenylation of mRNA 3′-ends is a co-transcriptional event that occurs for nearly all eukaryotic mRNA with the exception of histone mRNAs (1). This process relies on a two-step mechanism starting with cleavage of the pre-mRNA in the 3′-untranslated region (3′-UTR) (2–4). This first step is immediately followed by the polymerization of adenosines at the 3′-hydroxyl end of the upstream cleavage product, while the downstream fragment is degraded by Rat1 (5). In yeast, the polyadenylation machinery comprises two large multi-protein complexes: the Cleavage/polyadenylation Factor IA (CF IA) (6,7) and the Cleavage and Polyadenylation Factor (CPF). Additional factors are also required for the in vitro process such as Hrp1/Nab4 (6,8) and the poly(A)-binding proteins Pab1 and Nab2 (7,9–11).

The CF IA complex consists of the four proteins Rna14, Rna15, Pcf11 and Clp1. Each of these proteins consists of distinct domains that contribute to the function of CF IA, including assembly of the complex and interaction with other components of the processing machinery or target pre-mRNA. Investigation of these structured domains has directly led to insight into CF IA architecture and function. Solution studies and crystal structures of the Kluyveromyces lactis Rna14, as well as the orthologue CstF-77 from Encyphalitozoon cuniculi and Mus musculus, have shown that Rna14 and CstF-77 self-associate into tight homodimers (12–15). The C-terminal region of Rna14 also forms a heteromeric complex with Rna15 as revealed by the solution structure of the Rna14–Rna15 minimal heterodimer (16) and the structure of the Rna14–Rna15 sub-complex from K. lactis (15). A three-helix bundle at the C-terminus of Rna15 has been shown to contact Pcf11 (17), and a short region in Pcf11 mediates association with Clp1 as observed in the structure of a Clp1–Pcf11 minimal complex (18,19). These contacts contribute to the correct assembly of CF IA, although they also dictate a more complicated stoichiometry versus a simple heterotetramer (20).

Recent studies have proposed an overall model for CF IA (21), with a stoichiometry of 2:2:1:1 or 2:1:1:1 (21,22) respectively for Rna14:Rna15:Clp1:Pcf11, but the functional basis for such an asymmetry has not yet been described.
Protein domains within CF IA also enable interaction with the target pre-mRNA and components of the transcription and processing machinery. Molecular characterization of the RNA recognition motif (RRM) domain of Rna15, and the orthologue CstF-64, reveal aspects of pre-mRNA binding (23). Increased RNA sequence specificity has also been demonstrated with structural details of the auxiliary subunit Hrp1 (also known as CF IB or Nab4), and in particular within a Hrp1–Rna15–RNA ternary complex (24–26). A connection to the largest subunit of RNA polymerase II, Rpb1 is mediated by the C-terminal interacting domain (CID) of Pcf11 that binds to the Rpb1 C-terminal YSPTPS heptamer repeats with phosphorylation on the serine in position 2 (27,28).

Despite the characterization of the CF IA domains described above, additional folded domains remain to be studied. This is most evident for Pcf11, for which only the N-terminal CID, the short Clp1-interacting region and recently, a zinc-binding domain (29) are described at the atomic level. In this study, we have analysed additional regions of Pcf11 and demonstrated that Pcf11 has two zinc-binding domains. Interestingly, mutations of these domains affect Pcf11 function to different extents. Mutation of the C-terminal zinc-binding motif has a dramatic impact on both yeast viability and pre-mRNA 3′-end processing, whereas mutations in the upstream zinc-binding domain has a small but noticeable effect on cell growth and 3′-end formation, suggesting different functions of these potentially similar motifs. The entire zinc finger region of Pcf11 is also capable of a non-specific interaction with RNA but only in the absence of Clp1. The C-terminal zinc-binding domain, which we find essential for pre-mRNA 3′-end processing, was also produced in isolation and was shown to contain flexible residues around an unusual zinc-binding fold. Altogether, this study provides a refined description of Pcf11 organization and gives novel insights into CF IA in mRNAs 3′-end maturation.

**MATERIALS AND METHODS**

**Cloning and rCF IA complex expression**

Each full-length protein was obtained by PCR on yeast *Saccharomyces cerevisiae* genome and cloned into modified versions of pET-15b, pET28b, pCDF and pLysS bacterial expression plasmids (30,31). Integrity of each clone was verified by sequencing. For co-expression, BL21(DE3) (Novagen) were transformed with the relevant combinations of plasmids and plated on Petri dishes with half of the recombinant antibiotic concentration. Mini-cultures were grown allowing the production of an N-terminally His<sub>6</sub>-GST-tagged proteins. The plasmids were transformed into *E. coli* BL21 (DE3). For large-scale protein purification, 1 L of TB was inoculated with the relevant overnight pre-culture and grown up to an OD of 1.5. After cell harvesting, the cell pellet was resuspended in a buffer containing 25 mM Tris–HCl pH 7.5, 150 mM NaCl and lysed by three passes into an EmulsiFlex C3 (Avestin). The crude extract was centrifuged at 50 000 g for 1 h at 4°C. The supernatant was incubated with His-affinity resin (Sigma) and loaded onto an AK 16 column (GE Healthcare). Contaminants were removed by extensive wash with buffer A containing 25 mM Tris–HCl pH 7.5, 150 mM NaCl. Proteins were eluted from the column with a gradient of buffer A supplemented with 250 mM imidazole. The eluted proteins were then loaded onto a HP-Heparin 5 ml (GE Healthcare) and separated by a gradient of NaCl up to 500 mM in 25 mM Tris–HCl pH 7.5. The pool of fractions was concentrated and loaded onto a Superdex-200pg column (GE Healthcare) equilibrated in 25 mM Tris–HCl pH 7.5, 250 mM NaCl, 1 mM DTT.

**Zinc-binding domains purification**

The DNA sequence coding for residues 530–626 or 403–626 of Pcf11 were cloned into a modified pET-15b plasmid allowing the production of an N-terminally His<sub>6</sub>-GST-tagged proteins. The plasmids were transformed into *E. coli* BL21 (DE3). After cell harvesting, the cell pellet was resuspended in a buffer containing 25 mM Tris–HCl pH 7.5, 150 mM NaCl and lysed by three passes into an EmulsiFlex C3 (Avestin). The crude extract was centrifuged at 50 000 g for 1 h at 4°C. The supernatant was incubated with His-affinity resin (Sigma) and loaded onto an AK 16 column (GE Healthcare). Contaminants were removed by extensive washes with a buffer A containing 25 mM Tris–HCl pH 7.5, 250 mM NaCl. Proteins were eluted from the column with a gradient of imidazole up to 250 mM supplemented in buffer A. The eluted proteins were cleaved with Tew protease to remove the His-GST tag overnight at 4°C. Then a Hi-prep 26/10 Desalting column was used to exchange the protein buffer into the Buffer A. The protein was

**Pull-down assays**

After overnight induction of 10 ml culture with 0.1 mM IPTG, harvested cells resuspended in lysis buffer (1.5x PBS, 1 mM Mg(OAc)<sub>2</sub>, 0.1% NP-40, 20 mM Imidazole, 10% (w/v) glycerol) were lysed by 10 s sonication at 3 W. Crude extract was then centrifuged and soluble fraction incubated 30 min with 30 μl of His-Affinity Cobalt resin. After three washing steps with 1 ml of lysis buffer, beads were eluted with 3 volumes of lysis buffer supplemented with 250 mM imidazole and one-third of the sample was loaded on a SDS-polyacrylamide gel.
then loaded on a His-affinity column to eliminate the His- 
GST tag and Tev protease. The zinc-finger domain was fur- 
ther purified using a High-resolution Superdex S-75pg gel- 
filtration column (GE Healthcare) equilibrated in 50 mM Tris (pH 7.5), 150 mM NaCl supplemented with 50 μM Zn(OAc)2. The purified protein samples were concentrated to 2.6 mg ml⁻¹, frozen and stored at –80°C. The concentra-
tion was estimated from absorbance at 280 nm assuming 
absorption of 1.54 cm⁻¹ for a 1 mg ml⁻¹ solution as calcu-
lated from the construct sequence.

NMR spectroscopy

In addition to protein expressed in LB media at natu-
ral abundance, GST-tagged Pcf11 (530–626) was also ex-
pressed using Rosetta2 (DE3) cells in M9 minimal me-
da supplemented with 1 g/l [15N]-ammonium chloride, 2 
g/l [13C]-glucose or with 2 g/l [10%-13C]-glucose and puri-
fied with as with the natural abundance protein. Purified sam-
ple contained 100–360 μM untagged protein in 50 mM Tris 
(pH 7.5), 150 mM NaCl, 50 μM Zn(OAc)2; with either 10% or 99% (v/v) 2H2O. NMR spectra were acquired at 20°C on a 700 MHz or 800 MHz Bruker Avance III spectrometer, equipped with a room temperature or cryo-
genic triple resonance gradient probe, respectively. Data ac-
quisition used TopSpin versions 2.1 and 3.2. Processing used 
NMRPipe/NMRDraw (32) and data analysis used 
Sparky (T.D. Goddard & D. G. Kneller, University of 
California, San Francisco, USA). Chemical shift assignment 
of the backbone 1H, 13C and 15N nuclei of Pcf11 for residues 548–603 relied on 2D 1H,15N-HSQC, 3D HNCO, 3D HNCACB and 3D HNHA spectra. Aliphatic sidechain assign-
ment used 2D 13C-HSQC, 3D (H)CCH-TOCSY (14 ms mixing time), H(C)CH-TOCSY (14 ms mixing time) 
(H)CCH-TOCSY (14 ms mixing time). Stereospecific as-
signment of the ψ1/ψ2 methyl groups of valine and the 
δ1/δ2 methyl groups of leucine used the sample prepared 
with [10%-13C]-glucose (33). Aromatic sidechain assign-
ment used 2D 1H,13C-HSQC and 1H,1H-NOESY (150 
ms mixing time) spectra, with an additional long-range 
1H,15N-HMBC spectrum to determine the protonation 
state of the δ1 and ε2 nitrogen atoms of the two histidine 
residues. The 1H, 13C and 15N chemical shift assignments 
have been deposited in the Biomolecular Magnetic Reso-
nance Data Bank (http://bmrb.wisc.edu) under accession 
code 34061.

Structure calculation

The backbone 1H, N, 1H,α, 13Cα, 13Cβ, 15N, 13C and 15N chemical shift values of Pcf11 (530–626) provided Ψ and φ angu-
lar restraints for 51 residues by using TALOS-N (34). In addi-
tion, 30 χ1 angular restraints were obtained. Two addi-
tion χ angle restraints for Val552 and Ile588 were based on analysis of 13C side chain resonances and the program 
Sider (35,36). Distance restraints were obtained from 2D 
1H,1H-NOESY (150 ms mixing time), 3D 1H,1H-NOESY-
13C-HMQC (150 ms mixing time) and 3D 1H,15N-HSQC– 
1H-NOESY (150 ms mixing time) with an initial set of 
976, 701 and 338 crosspeaks, respectively. The final list of 
unambiguous restraints included 353 intraresidue, 160 se-
quential, 54 medium and 332 long range distances, with an 
additional list of 271 ambiguous distance restraints. The four distance and 13 dihedral zinc-binding geometry con-
straints were excluded from the initial structure calculation, 
and were only incorporated in the final annealing step per-
formed in explicit water. The final ensemble contains the 15 
lowest energy structures calculated by using CNS1.1 (37) 
implemented within ARIA1.2 (38). The structure ensem-
ble has been deposited in the Protein Data Bank (http: / / www.ebi.ac.uk/pdbe) with PDB ID 5M9Z.

Atomic absorption spectroscopy

Pcf11 (530–626) constructs and Clp1-Pcf11 (403-626) sub-
complexes were first buffer exchanged against a 250 mM ammonium acetate (NH4Ac) solution at pH 7.5 using gel filtration (Zeba 0.5 ml, Thermo Scientific, Rockford, IL, USA) and ultrafiltration spin columns (Vivaspin 500 MWCO 10 000, Sartorius, Göttingen, DE), respectively. Protein concentration was determined spectrophotometri-
cally. NanoESI-MS characterization of Pcf11 variants was 
next generated on an electrospray TOF mass spectrometer (LCT, Waters, Manchester, UK) and on a Q-TOF (Synapt G2 HDMS, Waters, Manchester, UK), both equipped with an automated chip-based nanoESI source (Triversa Nanomate, Advion Biosciences, Ithaca, NY, USA) operating in the positive ion mode. External calibrations were gener-
ated from the multiple charged ions produced by a 2 μM horse heart myoglobin solution diluted in a 1:1 (v/v) water: acetonitrile mixture acidified with 1% (v/v) formic acid, and caesium iodide clusters diluted to 2 g/l in a 1:1 (v/v) water: isopropanol solution. Intact masses of indi-
vidual Pcf11 constructs were first determined in denatur-
ing conditions by diluting proteins to 2 μM in a 1:1 (v/v) water/acetonitrile mixture acidified with 1% (v/v) formic acid. Non-covalent complexes analysis in non-denaturing conditions was then performed by diluting samples to 5 μM 
in 250 mM NH4Ac buffer at pH 7.5. Fine-tuning of instru-
mental parameters was notably applied to preserve non-
covalently bound species in the gas phase and ensure effi-
cient ion transmission. Particularly, LCT and Synapt G2 
source back pressures were respectively raised to 6.7 and 7.7 mbar using a throttling valve, while acceleration volt-
age applied on the sample cone were set to 120 and 80 V. 
Data processing was finally carried out on MassLynx 4.0 
(Waters, Manchester, UK).

Mass spectrometry experiments

Yeast extracts were prepared following the spheroplast pro-
cedure and tandem-affinity purification (TAP) of the fac-
tors was performed as described previously (10). The im-
munoblot analysis of 3′ processing complexes was done
with equal amounts of purified factors run on 12% polyacrylamide-SDS gels and probed after transfer with rabbit polyclonal antibodies to Rna14 (1:3000), Rna15 (1:10000), Pcf11 (1:2000) and Clp1 (1:1000).

**In vitro pre-mRNA 3′-end processing assays**

Pre-mRNA 3′-end processing assays were performed with purified factors as described previously on the CYC1 or CYC1 pre-cleaved (CYC1pre) precursors radiolabeled *in vitro* with α-32P-UTP (10,39). Native CF IA (nCF IA) and nCPF were purified from yeast cells according to the TAP procedure (40) from strains expressing N-terminal TAP-tagged RNA15 and TAP-tagged Flpl, respectively (see Table 3 for the strains). The purification of nCFP was performed in a nra15-1 mutant background (inactive in this mutant form (39)) to ensure that no active nCF IA co-purifies and obscures the complementation assays with CF IA coming from native or recombinant origins (Figures 2 and 7). Hrp1/Nav4 (50 ng per assay) and Pab1 (140 ng per assay) were obtained from recombinant sources (10). Recombinant CF IA (rCF IA-ΔQ20) was obtained as described above. Reaction products of processing assays were analyzed on 6% polyacrylamide–8.3 M urea gels and visualized by phosphorimaging.

**Yeast strains and plasmids**

Strain YLM232 containing the pcf11Δ::HIS3 allele was constructed by replacing the TRP1 marker disrupting the PCF11 gene in strain NA50 (41) by the HIS3 marker. All of the yeast strains used in this study are listed in Table 3. pNA39 (URA3-ARS/CEN-AmpR (41)) was used to complement the pcf11Δ::HIS3 null allele in YLM233 strain. The PCF11 gene was also amplified by PCR with *S. cerevisiae* genomic DNA as template, from position –458 to +2566 (nucleotide +1 corresponding to the A of the start codon of PCF11 open reading frame) and cloned into the pFL36ΔSX plasmid (LEU2-ARS/CEN-AmpR); this plasmid was constructed as described in (41). Mutations of the two zinc-binding domains were obtained by site-directed mutagenesis and re-introduced in pFL36 or pFL36ΔSX by gap-repair in yeast (for pcf11-18) or by direct cloning (for pcf11-19), respectively. Plasmid pFL36ΔSX-pcf11-ΔQ20 was constructed by gap-repair. All of the constructs were verified by sequencing. Haploid strains expressing wild-type (YLM244) or mutant (YJG23, YJG24) plasmid-borne PCF11 alleles together with the TAP-tagged RNA15 gene were obtained by crossing YLM237, YJG10 and YJG12 with YLM221, respectively (see Table 3), and analysis of their progeny after meiosis. Mutant pcf11-20 expressing the Pcf11 protein mutated in the two zinc-binding domains was cloned in pFL36ΔSX and used to transform the diploid strain YLM232. Phenotypic analysis of the progeny after meiosis is described in Figure 1C.

**S1 nuclease protection assay**

Plasmids used for the S1 nuclease protection assays were made as follows: a PCR fragment encompassing the sequence ranging from the *Xho I* site located at nucleotide +45 (according to the start of mature snR13 RNA) to nucleotide +93 in the TRS31 gene (+1 being the start of TRS31 ORF) was amplified from yeast genomic DNA and cloned into the pGEM4 vector between the *EcoR I* and *BamH I* restriction sites (Promega). This plasmid was digested with *Xho I* and *Pvu II*, the released fragment was purified on an agarose gel and 3′-end labeled with the Klenow enzyme and [α-32P]-TTP (25 μCi/3000 Ci/mmol). Approximately 30 ng of the labeled DNA probe were hybridized with 30 μg of total RNAs extracted from each strain or 2 μg of control yeast tRNA and digested with 150 units of S1 nuclease (42). The protected fragments were resolved on 8% polyacrylamide–8.3 M urea gels and visualized by phosphorimaging. The positive control RNA was obtained from a PCR fragment covering the whole snR13 mature transcription unit down to nucleotide +291 cloned between the *EcoR I* and *HinD III* restriction sites of pGEM4. The resulting plasmid was cut with *EcoR I* and served as template for the generation of a synthetic RNA by run-off transcription with T7 RNA polymerase.

**UV-crosslinking assay**

UV cross-linking reactions were performed in microtiter plates and contained (in 10 μl) 100 fmol of 32P-labeled CYC1 in 75 mM potassium acetate, 2 mM magnesium acetate, 0.01% NP-40, 1 mM DTT, 0.1 mg/ml purified BSA, 2 μM tRNA (0.5 μg), 5 μg total yeast RNA (when applicable) and 100 ng of recombinant proteins. The proteins were first incubated for 15 min at 30°C in the reaction mixture either alone or in the presence of 5 μg of total yeast RNA. The *in vitro*-labeled CYC1 transcript was then added and the incubation was further continued for 15 min at 30°C. The samples were irradiated twice at 500 mJ on ice in a UV Stratalinker, and then digested with RNase A (100 ng per reaction) for 30 min at 37°C. The proteins were mixed with 3 μl of SDS-loading buffer and separated on SDS–12% polyacrylamide gels. The gels were fixed, dried and visualized by phosphorimaging. To control that equal amounts of the different versions of the recombinant Pcf11[403–626] polypeptides were used for UV cross-linking, ~600 ng of each recombinant proteins was loaded on a SDS–12% polyacrylamide gel and the proteins were compared after Coomassie staining (see Figure 9A).

**RESULTS**

**Putative zinc-binding domains in Pcf11 are required for cell viability**

Sequence analyses of the yeast Pcf11 protein have previously revealed the presence of two zinc-finger motifs of the C2H2 and CCHC types neighboring the Clp1 interaction domain (Figure 1A; Supplementary Figure S1) (29,43,44). As a step towards experimental confirmation, we analysed whether these putative zinc-binding residues are important for cell viability. To do so, the *PCF11* gene under its natural promoter was amplified by PCR and cloned into pFL36 and pFL38 centromeric plasmids. We next introduced point mutations in the *PCF11* gene to target two potential zinc-binding residues within each region. Specifically, the conserved C421 and C424 residues within the first
region, and independently the conserved C564 and C567 amino acids located in the second region, were mutated into serine and the resulting mutants, named pcf11-18 and pcf11-19, respectively, were assayed for growth at different temperatures (Figure 1B). Mutant pcf11-18 did not show any strong growth defect at physiological temperatures of 24°C or 30°C. In contrast, cell growth was significantly affected at 37°C and addition of 2% formamide in the medium exacerbated this phenotype at 24°C in comparison to the wild-type strain. Unlike the pcf11-18, pcf11-19 exhibited a strong growth defect at both 24°C and 30°C and turned out to be lethal at 37°C or in the presence of formamide at 24°C. We next constructed an allele that combined the mutations present in pcf11-18 and pcf11-19 such that both putative zinc-binding motifs were mutated. The resulting allele, pcf11-20, was cloned into a LEU2-marked plasmid and used to transform the heterozygous pcf11Δ::HIS3/PCF11 diploid strain YLM232 (see Materials and Methods and Table 3). After meiosis, only a few spores were observed showing a very strong slow-growth phenotype (Figure 1C; the spores are indicated with an arrow). These spores corresponded to the pcf11-20 mutants since they grew on a synthetic medium lacking histidine and leucine. This mutant therefore has a much stronger growth defect in comparison to the pcf11-19 allele. Despite the centromeric nature of the plasmid carrying the pcf11-20 allele, only a few spores corresponding to the mutant were able to grow (spores 2B, 4C, 5C, 8B from the first diploid, and spores 5A, 8B from the second diploid, Figure 1C). After inspection under the mi-
Figure 2. Recombinant CF IA works efficiently in mRNA 3’-end formation. (A) CF IA was reconstituted by co-expression of the four CF IA subunits in E. coli. Each subunit was alternatively fused at its N-terminus with a hexahistidine tag allowing for protein purification. Pull-down experiments were performed from 10 ml cell culture after immobilization on affinity-resin. The bound proteins were eluted with imidazole and analyzed by a 10% SDS-PAGE. Proteins were revealed by Coomassie-blue staining. The asterisk (*) refers to the His-tagged subunit of CF IA. (B) In vitro cleavage assay with the CYC1 precursor was performed in the presence of native CPF (nCPF) and wild-type native CF IA (nCF IAWT) purified by TAP-tagging (lane 8) or recombinant CF IA (rCF IA-D1Q20; lane 9). Hrp1 was included in the reaction to avoid cleavage at cryptic sites. Lanes 1 and 7: unreacted CYC1 precursor; lanes 1–6: negative controls showing the absence of cleavage activity of the different factors on their own. Lane 5 shows however the intrinsic and normal polyadenylation activity of Pap1-containing CPF. (C) In vitro polyadenylation assay of the CYC1 precleaved precursor (CYC1/pre) with nCF IAWT (lane 5) or rCF IA-D1Q20 (lane 6) and nCPF, Hrp1 and Pab1. The latter is required to control poly(A) tail elongation. Lanes 1 and 4: unreacted CYC1/pre precursor; Lane 2: nCPF alone can polyadenylate unspecifically CYC1/pre; lane 3: Hrp1 and Pab1 are inactive on their own.

croscope, many of the spores did not go beyond one or two cell divisions. Therefore, the pcf11-20 mutant conferred a strong germinative defect and a near-lethal phenotype for those spores which were able to escape germination.

In conclusion, the mutations of putative Pcf11 zinc-binding cysteines designed to abolish metal chelation in both the C2H2 and CCHC regions led to a clear growth defect, and thus supports a general need for Zn2+ binding in Pcf11 function.

Recombinant expression of CF IA

Prompted by the cell growth defects in the mutations described above and in order to circumvent native CF IA purification from these alleles, we first set-up a procedure for reconstituting S. cerevisiae CF IA from bacterially expressed proteins as previously described (22). This approach allows for the study of CF IA containing deletions of essential domains (e.g. Pcf11 CID) or point mutations that cannot be studied in vivo due to their lethal phenotype, and
thus enable studies on CF IA architecture and pre-mRNA 3′-end processing. Briefly, each subunit was cloned into a plasmid carrying a resistance to a different antibiotic allowing for the introduction of up to four plasmids at a time into a single bacterial cell (30,31,45). Due to limited levels of protein expression of full-length Pcf11 (data not shown), we opted for a slightly altered version of Pcf11 in which a 20-aminoacid stretch of glutamine residues has been deleted (residues 234–253). This shorter version of Pcf11 is viable and has no detectable phenotype in yeast (Supplementary Figure S2), and will be referred to as Pcf11-ΔQ20.

The expression of individual proteins was confirmed (Figure 2A). Subsequently, a four-subunit co-expression assay was attempted by swapping the His-tag alternatively to the N-terminus of each subunit. Each combination successfully retrieved the entire recombinant CF IA (rCF IA), however the best combination was found when Rna15 is Histagged (Figure 2A). A robust purification procedure was then set-up in order to retrieve pure rCF IA-ΔQ20 complex for further experiments (see Materials and Methods).

rCF IA retains in vitro 3′-end processing activity

CF IA is essential in pre-mRNA 3′-end processing through its role in the positioning of the catalytically active CPF complex onto the pre-mRNA. We used an in vitro reconstituted cleavage and polyadenylation assay to test whether rCF IA is able to substitute for the yeast native complex (referred to here as nCF IA). As expected, nCF IA in combination with TAP-purified nCPF readily processed the CYCI precursor in a cleavage-only reaction (where polyadenylation is inhibited with EDTA and Hrp1 added to control cleavage site selection (8); Figure 2B, lane 8), as well as in a polyadenylation reaction in which Hrp1 and Pap1 were also added (Figure 2C, lane 5). Note that none of the different factors used in the assays exhibit any processing activity on their own (Figure 2B, lanes 1–4, 6 and 2C, lane 3) except for CPF which is capable of non-specific polyadenylation activity on any RNA due to the presence of the poly(A) polymerase Pap1 as one of its intrinsic subunits as previously described in the literature (e.g. (40,46)) (Figure 2B, lane 5 and 2C, lane 2). When the recombinant complex, the cleavage and polyadenylation reactions appeared as efficient when rCF IA-ΔQ20 replaced native nCF IA in the assays (Figure 2B, lane 9, and 2C, lane 6, respectively). Therefore, Pcf11-ΔQ20 can efficiently work in recombinant rCF IA compared to native nCF IA, as far as in vitro 3′-end processing is concerned.

Pcf11 contains two Zn²⁺ atoms

rCF IA provides an ideal source to quantify the Zn²⁺ content in the complex by atomic absorption, as well as recombinant sub-complexes of Rna14–Rna15 and Pcf11–Clp1 heterodimers. Samples were analysed at 213.9 nm using the dialysis buffer as the baseline, and a titration curve ranging from 1.25 to 20 μM of ZnSO₄ solution was also prepared in the dialysis buffer. According to these measurements, only trace amounts of Zn²⁺ were detected in the Rna14–Rna15, whereas the rCF IA and Pcf11–Clp1 samples displayed significant Zn²⁺ content (Table 1). These data demonstrate the presence of natively bound Zn²⁺ atoms within rCF IA and most specifically within Pcf11, since the structure of Clp1 did not reveal any bound Zn²⁺ atoms (18,19).

To further identify an exact stoichiometry and to better define the zinc-binding regions in Pcf11, we used mass spectrometry analysis in non-denaturing conditions. We first designed a construct from residue 403 to the C-terminus (residue 626) that spans the two-potential zinc-binding regions. The protein was expressed in E. coli and purified to homogeneity and the accurate Zn²⁺ stoichiometry was determined by comparing the mass measured under non-denaturing or denaturing conditions (Figure 3). The mass spectrometry data obtained under non-denaturing conditions fit the theoretical value expected for Pcf11 (403–626) of 25 742.5 Da, which includes a G–H–M tripeptide at the N-terminus. Mass spectrometry measurement under native conditions shows an additional molecular mass of 130.5 Da corresponding to two bound Zn²⁺ atoms (Figure 3A). As a final test, the second C-terminal zinc-binding region of Pcf11 (residues 530–626) was expressed in isolation. The mass under denaturing conditions of 11 149.3 ± 0.1 Da again fits the theoretical value. Comparison to the denaturing conditions shows an additional molecular mass of
Table 1. Zn\(^{2+}\) content analysis in recombinant CF IA

| Protein       | [Protein] (µM) | [Zn\(^{2+}\)] (µM) | Ratio |
|---------------|----------------|--------------------|-------|
| Rna15–Rna14   | 3.14           | 0.5                | 0.16  |
| CF IA–ΔQ20    | 6.15           | 11                 | 1.79  |
| Pcf11–ΔQ20–Clp1| 17             | 23.5               | 1.38  |

65.7 Da, attributed to a single bound Zn\(^{2+}\) atom (Figure 3B). Attempts to produce the other isolated zinc-binding region from residue 403–454 failed due to excessive instability. Nevertheless, the results demonstrate that Pcf11 harbors two zinc-binding regions: one Zn\(^{2+}\) binds at the C-terminus (from 530–626), and another is likely bound by the conserved amino acids within 403–454. Further insight into the architecture of these domains necessitated a structural approach.

**The Pcf11 C-terminal zinc-binding domain has a novel fold**

To obtain atomic information on both zinc-binding domains from Pcf11, we first attempted to determine a structure of the larger complex between Pcf11 (403–626) bound to full-length Clp1. Even though crystals diffracting to a limited resolution (<4 Å) were obtained, no electron density could be attributed to Pcf11 apart from the Clp1 interacting domain, as has been previously described (18,19).

Due to the inability to produce the first zinc-binding domain in isolation, we focused on determining the structure of the second zinc-binding domain of Pcf11 (530–626). By using NMR spectroscopy, we found that the Pcf11 (530–626) construct indeed contains a well-folded domain, as well as a significant number of residues that fall within the region typical of a conformationally disordered peptide (Figure 4A). It was also apparent that Zn\(^{2+}\) binding was required to maintain the overall fold, since the protein was unstable in the absence of added Zn\(^{2+}\) during purification. Although it was not possible to assign the chemical shift values of the N- and C-terminal residues, a central region from G548 to T603 was amenable to detailed structural analysis by NMR-derived dihedral and distance restraints (Figure 4B, Table 2 and Materials and Methods). To prevent bias of the Zn\(^{2+}\) binding residues during structure calculation, the protein fold was first determined without any Zn\(^{2+}\)-based restraints. This calculated ensemble presented a consistent and favorable Zn\(^{2+}\) binding geometry formed by the side chains of C564, C567, H596 and C599. Therefore, a final refinement step in explicit water was performed, with the addition of a Zn\(^{2+}\) ion and a limited set of distance and dihedral restraints derived from an analysis of Zn\(^{2+}\) coordination in proteins (47). The overall structure is composed of seven β-strand elements and a single C-terminal α-helix (Figure 4C). The relatively large zinc-binding domain and high content of β-strands suggested an unusual zinc-binding fold. A recent structure of a similar region of Pcf11 from residues 538 to 608 (PDB ID 2NAX; (29)) displays the same overall fold, with a global RMSD between the two ensembles of 0.7 and 1.3 Å for the backbone and all heavy atoms, respectively, calculated by using SuperPose (48).

A search of the available structures by using DALI (49) failed to find a close structural homologue, but instead identified local homology (Z-score > 2) to one of the two zinc-binding sites within the RING finger fold (Figure 4D). For example, comparison to the recent structure from RING finger protein 165 (RFP165; PDB ID 5D0I; (50)) reveals a common β-turn placement of the first two cysteine residues in a C-x-I-C motif (C564 and C567 in Pcf11; C294 and C297 in RFP165), as well as a helical location of the remaining...
Table 2. Statistics for the ensemble of 15 lowest energy structures of Pcf11 residues 548–603

| NMR distance and dihedral restraints |  |
|-----------------------------------|---|
| Distance restraints               |  |
| Total                             | 1170 |
| Intraresidue                      | 353  |
| Sequential (ii-j = 1)             | 160  |
| Medium range (1<i-j<5)            | 54   |
| Long range (i-j>4)               | 327  |
| Ambiguous                         | 271  |
| Dihedral angle restraints         |  |
| \(\phi\)                          | 51   |
| \(\psi\)                          | 51   |
| \(\chi\)                          | 32   |
| Zn\(^{2+}\) coordination restraints |  |
| Distance                          | 4    |
| Dihedral                          | 13   |
| Structure statistics              |  |
| Violations (mean and SD)          |  |
| Distance restraints (Å)           | 0.010 ± 0.001 |
| Dihedral angle restraints (°)     | 0.76 ± 0.09 |
| Deviations from idealized geometry |  |
| Bond lengths (Å)                  | 0.001 ± 0.000 |
| Bond angles (°)                   | 0.300 ± 0.005 |
| Improperss (°)                    | 0.22 ± 0.01 |
| Ramachandran plot (%a,b          |  |
| Most favored regions              | 91.0 (95.0)c |
| Generally allowed regions         | 0.4 (0.0) |
| Disallowed regions                | 0.8 (0.0) |
| Average pairwise rmsd (Å)a       | 0.6 (0.4) |
| Backbone                          | 1.0 (0.9) |
| Heavy                             |  |

\(^a\) Using the Protein Structure Validation Software suite (71).
\(^b\) PROCHECK (72).
\(^c\) Number in parentheses for ordered regions: 549–555, 561–602.

A sequence alignment was then used to identify putative functional residues within the second zinc-binding domain (Figure 5A). Outside of the CCHC residues, only W582 is identical in all 13 species. This residue is found to be >40% solvent exposed in the structural ensemble by using the NACCESS program (52) and is thus able to contribute to biomolecular interactions. By extending the alignment analysis to residues that share sidechain properties in all 13 species, several residues around the zinc-binding site are identified (Figure 5B) which may contribute directly to Zn\(^{2+}\) binding properties (e.g. E569, N585, E602). Opposite to the zinc-binding residues, the aromatic amino acid (Y575) and two acidic residues (E577 and E578) cluster with W582 (Figure 5B). This conserved surface therefore presents a region of four exposed sidechains anchored in the triple-stranded \(\beta\)-sheet, specifically within strands \(\beta4\) and \(\beta5\) as well as the turn that connects them. This area may provide a binding surface for interaction with CPF subunits. Comparison of this region to the surface charge properties (Figure 5C) emphasizes the acidic and exposed nature of this region.

Zinc-binding mutations have little effect on CF IA assembly

Subunit–subunit interactions in CF IA have been reported through pairwise assays either with the two-hybrid analysis or by pairwise association/expression (13,16,17,20,41,43,53) giving rise to a refined interaction map in CF IA (26). However, interactions between two proteins that require a third stabilizing protein may not be detected, as for example with the PYM protein interaction with Mago and Y14 (54). We thus used our CF IA reconstitution assay to refine the subunit-subunit interaction map with respect to Pcf11, and to test whether the zinc-binding domains are necessary for CF IA assembly. We first tested the influence of double point mutations C421S/C424S (Pcf11-18) or C564S/C567S (Pcf11-19) in a pairwise coexpression assay between Pcf11 and Clp1 (Figure 6A, left panel). As observed, no significant decrease in binding between Pcf11 (331–626) and Clp1 was detected. We next addressed the effect of these double mutations on CF IA architecture (Figure 6A, right panel). The Pcf11 \(\Delta Q_{20}\) protein carrying the double mutations C421S/C424S (Pcf11-18) or C564S/C567S (Pcf11-19) did not alter significantly the presence of any CF IA subunit in comparison to the wild-type in pull-down assays.

We also tested the impact of the double cysteine mutants on the assembly of the native nCF IA complex (nCF IA). The mutant nCF IA factors were purified by the TAP-tagging method from strains YIG23 and YIG24, where Pcf11 was expressed from the pcf11-18 and pcf11-19 alleles, respectively (Figure 1 and Table 3). It should be noted that purifications of nCF IA from the mutant strains reproducibly retrieved lower amounts of factors compared to the wild-type strain for unknown reasons but likely related to the relative effects of the mutations on cell growth and viability. Nevertheless, all four subunits were present in nCF IA purified from the wild-type and mutant backgrounds (Figure 6B, lanes 1–3), but the bands corresponding to Pcf11 appeared slightly less abundant in the zinc-binding domain mutant alleles. Based on this silver-stained protein gels, equal amounts of factors were loaded on SDS-polyacrylamide gels and subjected to Western blot analyses with antibodies to all four subunits separately. The blots showed that, in comparison to equivalent signals detected with anti-Rna14 and anti-Rna15, the association of Pcf11 and also Clp1 was slightly impaired in the mutant factors (Figure 6C). With regard to nCF IA purified from the most affected mutant, the pcf11-19 allele, the Pcf11 mutant protein appeared to be more prone to degradation and was revealed as double bands yet both capable to associate with nCF IA (Figure 6B, lane 3 and Figure 6C, lane 3, panel \(\alpha\)-Pcf11). Nevertheless, the consequence of cysteine mutations in the zinc-binding motifs on nCF IA assembly is not dramatic enough to explain their impact on pcf11-18 and particularly pcf11-19 growth phenotypes.
Figure 5. Surface residue conservation separate from the zinc-binding site. (A) Sequence alignment of the second zinc-binding domain of Pcf11 from 13 species. Boundaries of the construct used in NMR studies (residues 530–626) and of the calculated structure (residues 548–603) are indicated, as are the locations of β-strands (arrows) and the α-helix (cylinder). Solvent accessibility for sidechains are calculated by using NACCESS (52) and indicated with varying shades of blue squares from solid blue (>80% accessible) to white (<20% accessible). Residues with complete sequence identity or similarity is highlighted in black and orange, respectively. Residues with similarity to yeast Pcf11 in >75% of the sequences are highlighted in light orange. (B) Surface representation of Pcf11 (548–603) with residues coloured by conservation as in panel A. The conserved patch formed by residues Y575, E577, E578 and W582 as well as the conserved residues around the Zn2+ atom (E569, V583, N586, E602) are indicated. (C) Surface charge properties of Pcf11 (548-603) with the negatively charge acidic regions in red, and the positively charged basic regions in blue.

Table 3. Yeast strains used in this study

| Strain   | Genotype                                      |
|----------|-----------------------------------------------|
| YLM232   | MATα/α, pcf11Δ::HIS3/PCF11, ade2-1/ade-2 leu-3,112/leu3-1 ura3-1/ura3-1 trp1-1/trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM233   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11, ade2-1/ade-2 leu-3,112 ura3-1 trp1 his3-11,15 cam1-100 GAL+ |
| YLM237   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM221   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11, [pNOP1]-TAP::RNA15-TRP1-Kl, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM244   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11, [pNOP1]-TAP::RNA15-TRP1-Kl, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM239   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11ΔQ20, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM244   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11ΔQ20, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM239   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11ΔQ20, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |
| YLM244   | MATα, pcf11Δ::HIS3, pFL36Δ-SX-PCF11ΔQ20, ade2-1/ade-2 leu-3,112 ura3-1 trp1Δ his3-11,15 cam1-100 GAL+ |

Mutations of the zinc-binding motifs impair mRNA 3′-end formation to different extents

We next examined the involvement of Pcf11 zinc-binding domains in mRNA 3′-end maturation in vitro. We assayed nCF IA factors purified from the wild-type and the mutant pcf11-18 and pcf11-19 strains in either cleavage or polyadenylation reactions at 30°C. As controls, no cleavage or polyadenylation activity was found with the purified factors on their own (Supplementary Figure S3). We took into account the lower amounts of nCF IA complexes retrieved from the mutant strains after TAP purification compared to wild-type as judged by the differences observed on protein gels (see Figure 6B and C) to carry out the assays. Moreover, to circumvent the variabilities observed in the Pcf11–Clp1 protein contents and their possible impact on 3′ processing activities, we performed the cleavage assays with increasing amounts of the different nCF IA factors. Efficient and very similar endonucleolytic cleavage of the CYC1 full-length precursor was obtained when nCF IAWT (Figure 7A, lanes 3–5) or nCF IApcf11-18 (Figure 7A, lanes 6–8) was used in the assay. Conversely, cleavage activity was undetectable with nCF IApcf11-19 no matter the amount of factor employed (Figure 7A, lanes 9–11). The possibility that the recombinant wild-type domain of Pcf11 could ameliorate the cleavage activity of mutant and wild-type nCF IA was tested. Starting from the same reaction conditions used in
Figure 6. The Pcf11 zinc-binding domains do not participate in CF IA assembly. (A) Pull-down assays of recombinant Pcf11–Clp1 and CF IA. The C421S/C424S point mutations and C564S/C567S point mutations were introduced into the ORF of Pcf11 (311–626) or Pcf11 ΔQ20. Pcf11:Clp1 or CF IA was pulled-down via a His-tag at the N-terminus of Pcf11. Proteins were eluted from the beads with imidazole and analysed by SDS-PAGE followed by Coomassie-Blue staining. Equivalent amounts of native wild-type and mutant TAP-purified CF IA were loaded on SDS–12% polyacrylamide gels and subjected to silver staining (B) or Western blot (C) analyses. Lanes 1: 5 μl of purified nCF IAWT; lanes 2: 20 μl of nCF IA ΔQ20; lanes 3: 20 μl of nCF IA ΔQ20.

lanes 4, 7 and 10 of Figure 7A, we complemented the assays with 100 ng of either recombinant Pcf11 [403-626], Clp1 or Pcf11 [403-626] alone. No significant improvement was observed with nCF IA WT or nCF IA ΔQ20 with Pcf11 [403-626]. Clp1 dimer but a slight inhibition could be noticed after addition of Pcf11 [403-626] alone (Figure 7B, lanes 2–4 and 5–7). Cleavage assays with nCF IA [ΔQ20] were not dramatically changed with either proteins (Figure 7B, lanes 8–10). Therefore, reconstitution of an active factor or stimulation of the activity was not possible by simply complementing the assay with wild-type domains alone.

We then performed polyadenylation assays with the same wild-type and mutant factors. In these conditions, less efficient polyadenylation of the CYC1 pre-cleaved precursor was reproducibly obtained with nCF IA WT compared to the wild-type factor (Figure 7C, lanes 4 and 3). In contrast, the absence of characteristic polyadenylated product was observed with the nCF IA [ΔQ20] mutant factor (Figure 7C, lane 5). Therefore, the impact on pre-mRNA 3′-end processing in vitro may likely explain the growth defects seen when Pcf11 zinc-binding motifs are mutated. Noticeably, these deficiencies also parallel the amplitude of the growth defects observed with the two mutants (Figure 1B).

RNA polymerase II transcription termination in Pcf11 zinc-binding mutants

The C-terminal position of the two Pcf11 zinc-binding motifs would predict that they are not important for RNA polymerase II (RNAPII) transcription termination as it has been demonstrated that only mutations in the 266 N-terminal amino acids of the protein affect termination (43). To test this assumption, an S1 nuclease protection assay was developed to map the 3′ ends of an RNAPII-transcribed gene coding for a stable snoRNA, snR13 (see Materials and Methods). Briefly, we designed a probe capable to hybridize with RNAs transcribed from the snr13-TRS31 region of the genome (Figure 8A). This 32P-5′-end-labeled probe (Figure 8B, lane 1) is sensitive to extensive degradation when subjected to S1 nuclease treatment when alone or in the pre-
Figure 7. Mutations in the zinc-binding motifs have distinct effects on 3′-end processing. (A) Pre-mRNA cleavage assay on the CYC1 precursor with TAP-purified factors from wild-type (nCF IA^WT) and mutant (nCF IA^p11-18) and nCF IA^p11-19) cells. Lane 1: unreacted CYC1 precursor, lane 2: nCPF and Hrp1 were assayed as negative controls for the lack of cleavage activity in the absence of CF IA. Increasing amounts of nCF IA^WT (0.5 μl, 1 μl and 2 μl; lanes 3–5) or nCF IA^p11-18 or nCF IA^p11-19 (2 μl, 4 μl and 8 μl; lanes 6–8 and 9–11, respectively) were added to nCPF and Hrp1 to carry out the cleavage reaction. (B) Complementation of cleavage assays. Cleavage reactions were run as in (A) with 1 μl of nCF IA^WT or 4 μl of nCF IA^p11-18 or nCF IA^p11-19. 100 ng of recombinant Pcf11[403-626]-Clp1 (lanes 3, 6, 9) or Pcf11[403-626] (lanes 4, 7, 10) were added to the reaction as indicated. Lane 1: unreacted CYC1 precursor. Lane 2: nCPF+Hrp1 were incubated with CYC1 as a negative control. (C) Polyadenylation assays performed on the CYC1 pre-cleaved precursor (CYC1/pre) with the same amounts of nCF IA^WT, nCF IA^p11-18 and nCF IA^p11-19 as in (B). Pab1 (140 ng) was added to the reaction to control de novo synthesis of poly(A) tails.
The zinc-binding domains do not participate in RNA polymerase II transcription termination at snoRNA genes. (A) Schematic representation of the S1 nuclease protection assay to map transcription termination at snoR13 transcription unit. (B) Detection of snoR13 read-through products by S1 nuclease protection. The radiolabeled probe (P, lane 1) was incubated with S1 nuclease either alone (lane 2) or in the presence of non-specific RNAs (tRNA; lane 3), a specific RNA designed to protect a 142-nt fragment (lane 4), or total RNAs extracted from wild-type or mutant cells grown at 24°C or after a shift to 37°C during 60 min (lanes 5–12). The protected fragments were separated by electrophoresis on 8% polyacrylamide–8.3 M urea gels. The bands on the gels were indicated as diagrammed in (C). The labeled molecular weight markers (in number of nucleotides) are the Msp I-digested pBR322 (M1) and the 100-bp DNA ladder (Promega; M2).

Pcf11 zinc-finger domains can bind RNA in the absence of Clp1 with different avidities

The presence of zinc-binding motifs in polypeptides is sometimes associated with an ability to bind nucleic acids. This is often the case with proteins involved in RNA processing. Even though the structure of the Pcf11 C-terminal zinc-binding domain has no overall structural homology with classical zinc fingers, we tested whether the region of Pcf11 containing the two zinc-binding motifs can associate with RNA. After confirming that equal amounts of the Pcf11 zinc-binding domains were used with a Coomassie-stained gel (Figure 9A), we performed UV cross-linking experiments with uniformly 32P-labeled CYC1 substrate, using recombinant Rna15 as a positive control and testing recombinant Pcf11[403–626], Clp1 (Figure 9B, lanes 1 and 2). As expected, a strong signal corresponding to Rna15 binding to the RNA was observed after SDS-PAGE and autoradiography of the gel (Figure 9B, lane 1). This interaction is due to the presence of a canonical RNP-type RNA-binding domain at the Rna15 N-terminus. Conversely, no signal was obtained in the same assay conditions with Pcf11[403–626]–Clp1 heterodimer (Figure 9B, lane 2). It has been previously shown that the region of Pcf11 encompassing the Clp1 binding site flanked by the two zinc finger domains binds not only Clp1 but also the RNA-binding protein and export factor Yra1 in a mutually exclusive mode (55). In competition experiments, the interaction between Yra1 and Pcf11 zinc-binding regions was lost after addition of Clp1 or RNA, despite the fact that the Yra1 RRM is not involved in the interaction between the proteins (56). To test whether the association of Pcf11[403–626] with Clp1 could mask an intrinsic capacity of the zinc finger domains to

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**Figure 8.** The zinc-binding domains do not participate in RNA polymerase II transcription termination at snoRNA genes. (A) Schematic representation of the S1 nuclease protection assay to map transcription termination at snoR13 transcription unit. (B) Detection of snoR13 read-through products by S1 nuclease protection. The radiolabeled probe (P, lane 1) was incubated with S1 nuclease either alone (lane 2) or in the presence of non-specific RNAs (tRNA; lane 3), a specific RNA designed to protect a 142-nt fragment (lane 4), or total RNAs extracted from wild-type or mutant cells grown at 24°C or after a shift to 37°C during 60 min (lanes 5–12). The protected fragments were separated by electrophoresis on 8% polyacrylamide–8.3 M urea gels. The bands on the gels were indicated as diagrammed in (C). The labeled molecular weight markers (in number of nucleotides) are the Msp I-digested pBR322 (M1) and the 100-bp DNA ladder (Promega; M2).
interact with RNA, we extended the UV cross-linking experiments to the Pcf11(403–626) region of the wild-type protein alone and of the corresponding mutant polypeptides Pcf11-18(403–626) and Pcf11-19(403–626) (Figure 9A, lanes 3–5). As hypothesized before, the absence of Clp1 revealed that the zinc-binding region of Pcf11 was able to effectively interact with CYC1 (Figure 9B, lane 3). Strikingly, the mutant Pcf11-18(403–626) polypeptide exhibited a strong signal comparable to that of Rna15 in the same assay conditions (Figure 9B, compare lane 4 and 1). Pcf11-19(403–626) was also prone to cross-link to the labeled RNA but in a manner similar to the wild-type protein (Figure 9B, lane 5 and 3). To evaluate the strength and/or specificity of Pcf11(403–626)–RNA interaction, we challenged CYC1 binding by incubating the proteins with an excess (5 µg) of total yeast RNA before adding the labeled transcript (see Materials and Methods). The right panel of Figure 9B showed that albeit diminished, Rna15 still exhibited a significant interaction with RNA (lane 6) but CYC1 interaction was lost for all of the Pcf11(403–626) versions (lanes 7–10). These experiments showed that Pcf11 zinc-binding domains have the capacity to bind non-specifically to RNA but only in the absence of another protein or RNA interactor.

DISCUSSION

In this article, we report the biochemical, biophysical and biological characterization of the two zinc-binding domains of Pcf11. The identified zinc-binding domains surround the Clp1 binding region (residues 454–530) and are located between residues 403–454 and 530–626, belonging to the C2H2 and CCHC class respectively (57). The zinc-binding domain spanning aminoacids 403–454 was the first to be identified (44) while both domains were more carefully depicted from the sequence of Pcf11 (43). Of note, the structural and functional analysis of the second zinc-binding domain was recently published (29). Mass spectrometry measurements demonstrate that each domain binds a single Zn2+ atom, and each zinc-binding domain displays a thermo-sensitive phenotype when two out of the four Zn2+ binding residues are mutated into serine. Simultaneous mutation of the two zinc-binding domains is close to lethality in yeast. We further demonstrate that these Pcf11 domains are not required for transcription termination in yeast and are involved to different degrees in pre-mRNA 3′-end processing in vitro.

CF IA bearing an altered version of Pcf11 in its first zinc-binding domain can efficiently cleave a pre-mRNA substrate and has only slightly impaired polyadenylation activity as compared with the wild-type factor. In contrast, the equivalent mutations in the second zinc-binding motif have a dramatic effect on both cell growth and 3′ processing. The overall assembly of the factor is mostly unchanged from the wildtype which is different than mutations in the Clp1 protein that abolished the association of Clp1 and Pcf11 with Rna14–Rna15 (53), or mutations in Rna14 or Rna15 which were described to impair CF IA subunit organization (16).

Along with this study, we also demonstrate that the poorly-conserved poly-glutamine extension of Pcf11 (residues 234–253), connecting the N-terminal CID to the Rna14–Rna15 binding domain, is not necessary for CF IA assembly. The poly-glutamine linker is also not required for mRNA 3′ end maturation, which likely explains the viability of pcf11 ΔQ20 mutant. As suggested by Amrani and colleagues (9), the poly-glutamine extension could confer a degree of freedom to the N-terminal and C-terminal domains, allowing them to fulfill independent functions. This possibility is reinforced by the observation that the region between residues 253 and 331 of Pcf11 is not otherwise necessary to assemble CF IA and may mainly function to extend the
linker region by more than 75 residues. This type of organization is also observed for Nab2, implicated in the regulation of the poly(A) tail length (58,59). Nab2 also exhibits a repetition of 12 glutamine residues linking the N-terminal domain related to mRNA export function, and the RNA interacting region located at the central and C-terminal regions. Interestingly, deletion of the Q-rich region of Nab2 also has no impact on pre-mRNA 3′-end processing either (10).

We have also reported the solution structure of the second zinc-binding domain of Pcf11. Similar to the findings by the Varani and Moore labs (29) the overall fold does not exhibit similarity to other reported zinc-binding domains. So far only CCCH-type zinc knuckles have been characterized among the proteins involved in mRNA 3′-end maturation. This includes Mpe1, an essential protein required for both cleavage and polyadenylation, that contains a central zinc knuckle (60), Nab2 (58,59,61) and Yth1 that contains five zinc knuckle domains (62,63). It should be noted, however, that mutant analysis for zinc-binding residues in these domains, as performed for Pcf11, have not been described. Previously reported PCF11 alleles, such as pcf11-2 (E232G, D280G, C424R, S538G, F562S, S579P) or pcf11-9 (A66D, S190P, R198P, R227G, E354V, K435V) exhibit only a thermo-sensitive phenotype (9,43). Cysteine mutations within Mpe1 or Nab2 lead to a cold sensitive phenotype and no detectable effect on growth at 37°C (58,60). The phenotype difference reported in this article may also illustrate the functional independence of each zinc-binding domain. Indeed, for the five CCCH domains of Yth1 there is variation in function: ZF2 and ZF4 interact with RNA, ZF4 and ZF5 interact with Fip1, ZF1 and ZF4 interact with the endonuclease Ysh1/Br5 and ZF1 and ZF2 are required for pre-mRNA cleavage (63).

Finally, Pcf11 is a key player in connecting RNAPII transcription termination, via its N-terminal CID, to pre-mRNA 3′-end cleavage, through its association to Clp1 and Rna14:Rna15 heterodimer. In this scheme, the Pcf11 zinc-binding domains appear to ensure the connection to pre-mRNA in the absence of CF IA and Hrp1. In addition to the contacts within CF IA subunits, Pcf11 exhibits direct contacts to CF IA subunits and other proteins. This is the case for Ysh1, Ydh1 and Yhh1 (64), Mpe1 (65), Ssu72 (66), Pt1 (67,68), Swd2 (69). Considering these interactions, the zinc-finger domains of Pcf11 could be seen as protein-protein interactors with specific CFIA subunits. As Swd2 is not required for the cleavage and polyadenylation steps in vitro (69,70), yet can affect 3′-end formation of some mRNAs and snRNAs in vivo, it is unlikely that a disruption of its interaction with Pcf11 could be responsible for the observed phenotypes.

Alternatively, but not in opposition, a putative role in RNA binding of the zinc-finger domain of Pcf11 has been suggested in our study. We showed that the region containing these motifs flanking the Clp1 interaction domain of Pcf11 can interact with RNA only in the absence of Clp1. This situation is reminiscent of Yra1 binding to the same region of Pcf11 as reported by David L. Bentley and coworkers (56). Tethering of Pcf11 to the pre-mRNA enhances binding of Yra1 to the C-terminal ZF1–Clp1-ID–ZF2 region of Pcf11 before transcription termination happens. Sub2 replaces Pcf11 and interacts with Yra1 which will cause the mRNA to be competent for export. In place of Yra1, Clp1 interacts with Pcf11 within the CF IA complex at the time of transcription termination (56). We propose that during these different exchange of interactions, a transient but efficient contact of Pcf11 with the elongating pre-mRNA occurs through the C-terminal ZF1–Clp1-ID–ZF2 domain and before Clp1 recruitment to maintain Pcf11 and associated proteins such as Rna14 and Rna15 on the pre-mRNA. It is also possible that Yra1 could act as a chaperone - as also proposed by Bentley and coworkers (56) - to induce a conformational change thus unmasking the Clp1-binding site of Pcf11.

As a result, the role of the C-terminal zinc-finger domain in pre-mRNA 3′-end processing is very likely direct and mediated through protein-protein contacts made with CFIA subunits that are still to be uncovered. The RNA-binding capacity of the zinc-finger mutants is probably not responsible for the observed phenotypes on growth as the purified mutant factors both showed that assembly of the complexes is maintained. We cannot however rule out that a proportion of the mutant Pcf11-19 protein is not associated to the CF IA complex in vivo and thus may perturb RNA processing and export, leading to the strong growth phenotype associated with the pcf11-19 mutant.

In summary, we provide evidence that Pcf11 has two zinc-binding domains mandatory for Pcf11 function in vivo. Even though the exact role of each zinc-binding domain remains to be elucidated, both impact mRNA 3′-end formation to very different intensities when mutated. In comparison, transcription termination at snRNA genes happens normally, and CF IA assembly is not affected. We propose a direct role in the interaction with specific CFIA subunits or a possible function through RNA recognition during the process of coordinating 3′-end processing and mRNP export.

ACCESSION NUMBER

The structure ensemble has been deposited in the Protein Data Bank (http://www. ebi.ac.uk/pdbe) with PDB ID 5M9Z.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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