Fission yeast RNA triphosphatase reads an Spt5 CTD code

SELOM K. DOAMEKPOR,¹ BEATE SCHWER,² ANA M. SANCHEZ,² STEWART SHUMAN,³ and CHRISTOPHER D. LIMA¹,⁴
¹Structural Biology Program, Sloan-Kettering Institute, New York, New York 10065, USA
²Microbiology and Immunology Department, Weill Cornell Medical College, New York, New York 10065, USA
³Molecular Biology Program, Sloan-Kettering Institute, New York, New York 10065, USA
⁴Howard Hughes Medical Institute, Structural Biology Program, Sloan-Kettering Institute, New York, New York 10065, USA

ABSTRACT

mRNA capping enzymes are directed to nascent RNA polymerase II (Pol2) transcripts via interactions with the carboxy-terminal domains (CTDs) of Pol2 and transcription elongation factor Spt5. Fission yeast RNA triphosphatase binds to the Spt5 CTD, comprising a tandem repeat of nonapeptide motif TPAYWSNGSK. Here we report the crystal structure of a Pct1·Spt5-CTD complex, which revealed two CTD docking sites on the Pct1 homodimer that engage TPAYN/S segments of the motif. Each Spt5 CTD interface, composed of elements from both subunits of the homodimer, is dominated by van der Waals contacts from Pct1 to the tryptophan of the CTD. The bound CTD adopts a distinctive conformation in which the peptide backbone makes a tight U-turn so that the proline stacks over the tryptophan. We show that Pct1 binding to Spt5 CTD is antagonized by threonine phosphorylation. Our results fortify an emerging concept of an “Spt5 CTD code” in which (i) the Spt5 CTD is structurally plastic and can adopt different conformations that are templated by particular cellular Spt5 CTD receptor proteins; and (ii) threonine phosphorylation of the Spt5 CTD repeat inscribes a binary on–off switch that is read by diverse CTD receptors, each in its own distinctive manner.

Keywords: transcription; elongation; mRNA processing; mRNA capping; X-ray crystallography

INTRODUCTION

The m⁷GpppN cap of eukaryal mRNA is formed by three enzymatic reactions: (i) RNA triphosphatase hydrolyzes the RNA ⁵′ triphosphate end (pppRNA) to a diphosphate (ppRNA); (ii) GTP:RNA guanylyltransferase converts ppRNA to GpppRNA via a covalent enzyme-(lysyl-N³)-GMP intermediate; and (iii) AdoMet:RNA(guanine-N7)-methyltransferase converts GpppRNA to m⁷GpppRNA (Ghosh and Lima 2010). Capping is targeted to nascent RNA polymerase II (Pol2) transcripts via physical interactions of one or more of the capping enzymes with the carboxy-terminal domain (CTD) of the Pol2 Rpb1 subunit. The Pol2 CTD consists of tandemly repeated heptapeptides of consensus sequence Y₁S₂P₃T₄S₅P₆S₇. Phosphorylation and dephosphorylation of the Tyr1, Ser2, Thr4, Ser5, and Ser7 residues and cis–trans isomerization of Pro3 and Pro6 inscribe a complex “CTD code” read by diverse receptor proteins that control transcription, modify chromatin structure, and catalyze or regulate mRNA capping, splicing, and polyadenylation (Buratowski 2009; Corden 2013; Eick and Geyer 2013; Jeronimo et al. 2013).

The Ser5-PO₄ “letter” of the CTD code plays a vital role in recruiting RNA guanylyltransferase (GTase) to the Pol2 elongation complex (Ho and Shuman 1999; Pei et al. 2001a; Schwer and Shuman 2011; Schwer et al. 2012). Indeed, the direct binding of GTase to the Ser5-PO₄ form of Pol2 CTD is a conserved theme among diverse eukaryal taxa (Ho and Shuman 1999; Pei et al. 2001a; Fabrega et al. 2003; Ghosh et al. 2011). Comparisons of the crystal structures of Candida albicans GTase (Cgt1), mammalian GTase (Mce1), and Schizosaccharomyces pombe GTase (Pce1) bound to Ser5-phosphorylated Pol2-CTD ligands illuminated how eukarya take divergent structural routes to achieve GTase-CTD interaction (Fabrega et al. 2003; Ghosh et al. 2011; Doamekpor et al. 2014). Whereas the CTD docking sites are dominated in each case by interactions of GTase with Ser5-PO₄ and Tyr1, the CTD segments bound to the GTases differ in their length and in the phase of the heptad sequence bound. For example, fission yeast GTase recognizes an 8-amino acid CTD segment (S₅PSYSPTS₅P) bracketed by two Ser5-PO₄ marks; mammalian GTase captures a 6-amino acid segment (S₆PYSYP); and Candida GTase has two distinct CTD-docking sites that recognize Ser5-PO₄ containing heptads in differently phased registers (TS₅PYSYP and PSYSPTS₅P).
Moreover, the atomic contacts to CTD differ among the three GTases and few of the CTD-interacting side chains are conserved. Thus, capping enzymes from different taxa have evolved unique strategies to read the same Pol2 CTD code.

The RNA triphosphatase (TPase) components of the mammalian and budding yeast capping apparatus are recruited passively to the Pol2 CTD, by virtue of their physical association with the GTase: in cis as a covalently fused TPase-GTase enzyme Mce1 in mammals (Ho and Shuman 1999) or in trans as separately encoded subunits of a TPase-GTase complex in budding yeast (Ho et al. 1999; Gu et al. 2010). The fission yeast S. pombe has a distinctive strategy for targeting cap formation to Pol2 transcripts, whereby the TPase (Pct1) and GTase (Pce1) enzymes are not associated physically, but instead bind independently to the Ser5-phosphorylated Pol2 CTD (Doamekpor et al. 2001a; Doamekpor et al. 2014).

Mammalian and fungal capping enzymes can also access nascent Pol2 transcripts via physical interactions with transcription elongation factor Spt5 (Wen and Shatkin 1999; Pei and Shuman 2002; Doamekpor et al. 2014). Spt5 is a large polypeptide, composed of multiple domain modules, that associates with the Pol2 transcription complex shortly after initiation and can exert negative and positive effects on transcription elongation (Hartzog and Fu 2013). Fission yeast Spt5 has a distinctive C-terminal repeat domain (the “Spt5 CTD”), composed of 18 repeats of a nonapeptide motif (T^3P^3A^3W^3N^3S^3G^3S^3K^3), that (i) binds the RNA capping enzymes Pct1 and Pce1 (Pei and Shuman 2002) and (ii) is targeted for threonine phosphorylation by the Cdk9 kinase (Pei and Shuman 2002, 2003; Viladevall et al. 2009). The CTDs of fission yeast Pol2 and Spt5 play overlapping roles in recruiting the capping enzymes in vivo (Schneider et al. 2010). Unlike the capping enzyme-Pol2-CTD interactions, which stringently depend on the Ser5-PO4 mark, Thr1 phosphorylation of the Spt5 CTD is not required for binding to fission yeast Pct1 or Pce1 (Pei et al. 2001a; Pei and Shuman 2002) and Pce1 binding to a Spt5 CTD is antagonized by Thr1 phosphorylation (Doamekpor et al. 2014).

Key insights to the interaction of the capping apparatus with Spt5 were gained via the crystal structure of a fission yeast Pce1-Spt5-CTD complex, which revealed a docking site in the OB domain of the GTase enzyme that captures the Trp4 residue of the fission yeast Spt5 nonamer repeat (Doamekpor et al. 2014). We found that a disruptive mutation in the Spt5-CTD binding site of Pce1 was synthetically lethal with mutations in the Pol2-CTD binding site (Doamekpor et al. 2014), signifying that the Spt5 and Pol2 CTDs cooperate to recruit guanylyltransferase to the transcription elongation complex in vivo.

Here we analyze the fission yeast RNA triphosphatase Pct1 and its interactions with the Spt5 CTD. We report crystal structures of the Pct1 apoenzyme and a Pct1-Spt5-CTD complex and show that Pct1 binding to Spt5 CTD is antagonized by threonine phosphorylation. We establish by structure-guided mutagenesis the relevance of the CTD interface to CTD binding in vitro and Pct1 function in vivo. Our results fortify an emerging definition of an “Spt5 CTD code.”

RESULTS

Crystallization of Pct1 and a Pct1-Spt5-CTD complex

Crystallization trials were conducted with S. pombe RNA triphosphatase Pct1 that had been premixed with (i) manganese and the high-affinity competitive inhibitor tripolyphosphate (Pei et al. 2001b), in an effort to mimic a substrate complex, or (ii) an 18-mer Spt5-CTD peptide TPAGNSGTAPWNGSK composed of two nonamer repeats corresponding to Spt5-(845–862). Crystals of the Pct1, tripolyphosphate, manganese mixture were grown by vapor diffusion against a precipitant solution containing 2.8 M sodium chloride, 10% PEG6000. Crystals were in space group P21. The structure was solved by molecular replacement, using the S. cerevisiae Cet1 homodimer (pdb 1D81) as the search model, and ultimately refined at 2.6 Å resolution with R/Rfree of 0.199/0.245 (Supplemental Table S1). No electron density was observed for tripolyphosphate or manganese; therefore we refer to this structure as Pct1 apoenzyme. The Pct1 apoenzyme crystal contained four protomers in the asymmetric unit organized as two homodimers, A/B and C/D. Ordered amino acids in A/B include 46–161, 166–303/40–161, 166–303 and in C/D include 46–108, 112–160, 166–303/40–108, 112–160, 165–303.

Crystals of the mixture of Pct1 and Spt5-CTD peptide were grown by vapor diffusion against 3 M sodium formate as the precipitant. The Pct1-Spt5-CTD crystals were in space group P21212. The structure was solved by molecular replacement (Hertog and Fu 2013). Fission yeast Spt5 has a distinctive C-terminal repeat domain (the “Spt5 CTD”), composed of 18 repeats of a nonapeptide motif (T^3P^3A^3W^3N^3S^3G^3S^3K^3), that (i) binds the RNA capping enzymes Pct1 and Pce1 (Pei and Shuman 2002) and (ii) is targeted for threonine phosphorylation by the Cdk9 kinase (Pei and Shuman 2002, 2003; Viladevall et al. 2009). The CTDs of fission yeast Pol2 and Spt5 play overlapping roles in recruiting the capping enzymes in vivo (Schneider et al. 2010). Unlike the capping enzyme-Pol2-CTD interactions, which stringently depend on the Ser5-PO4 mark, Thr1 phosphorylation of the Spt5 CTD is not required for binding to fission yeast Pct1 or Pce1 (Pei et al. 2001a; Pei and Shuman 2002) and Pce1 binding to a Spt5 CTD is antagonized by Thr1 phosphorylation (Doamekpor et al. 2014).

Overview of the Pct1 structure

The fold of Pct1 is shown in Figure 1A, with the A and B protomers colored green and cyan, respectively; the two Spt5 CTD peptides are depicted as stick models. The Pct1 homodimer consists of parallel triphosphate tunnels, each comprising a topologically closed 8-strand antiparallel β-barrel, resting on a predominantly α-helical globular pedestal. The pedestal forms the dimer interface at a central twofold symmetry axis, running perpendicular to the page in the view in Figure 1A. A surface model of the homodimer in Figure 1B, in the same orientation, highlights the through-and-through tunnel aperture. Rotating the surface model 90° around the
x-axis in Figure 1C underscores the concave entrance to the triphosphate tunnel.

A DALI search of the Protein Database with the Pct1 structure recovered S. cerevisiae RNA triphosphatase Cet1 (pdb 1D8I; Z score 26.8; 2.1 Å RMSD at 247 Ca positions) and the RNA triphosphatase domain of mimivirus capping enzyme (pdb 2QY2; Z score 14.0; 3.8 Å RMSD at 176 Ca positions) as the most closely related homologs (Lima et al. 1999; Benarroch et al. 2008). These enzymes belong to, and define, the triphosphate tunnel metalloenzyme superfamily (Gong et al. 2006; Keppetipola et al. 2007). They share a tunnel architecture and a mechanism of metal-dependent hydrolysis of the γ phosphate of nucleoside triphosphates and RNA 5′-triphosphate ends.

A horizontally offset superposition of the Pct1 and Cet1 protomers is shown in Supplemental Figure S1A. The loop segment from amino acids 43 to 54 preceding the first helix of the pedestal appears to be dangling free; however, as shown in Figure 1C, this segment actually extends across the twofold axis to interdigitate with the pedestal domain of the partner protomer. This feature of the structure accords with an earlier finding (Hausmann et al. 2003) that Pct1 homodimerization depends on the peptide segment 41VPKIEMNFLN50.

Deletion of this peptide converts Pct1 into a catalytically active monomer, which is defective in vivo in S. pombe and hypersensitive to thermal inactivation in vitro. In contrast, deleting the N-terminal 40 amino acids of Pct1, for which there was no interpretable electron density in our Pct1 crystal structures, had no effect on Pct1 homodimerization, activity in vivo, or thermal stability in vitro (Hausmann et al. 2003).

The structure of the homologous budding yeast Cet1 protein extends from amino acids 265 to 539. The Pct1 structure includes two interstrand loops that were disordered in the Cet1 structure (Supplemental Fig. S1A). The N-terminal 240 amino acids of Cet1 are dispensable for its RNA triphosphatase function in vivo and in vitro (Lehman et al. 1999). The Cet1 segment from amino acid 246–261 is essential for Cet1 function in vivo, but not for TPase catalytic activity in vitro, because this segment mediates tight association of Cet1 with the budding yeast GTase Ceg1 (Ho et al. 1999; Lehman et al. 1999; Gu et al. 2010). S. pombe Pct1 lacks a counterpart of this GTase-binding motif, consistent with the fact that Pct1 does not interact with the fission yeast GTase Pce1 (Pei et al. 2001a).

### Pct1 active site

The TPase active site within the tunnel aperture is shown in stereo in Supplemental Figure S1B. The aligned Cet1 active site, which is depicted in Supplemental Figure S1C, features a manganese ion in an octahedral coordination complex that includes a sulfate anion as a putative mimetic of the substrate γ phosphate (Lima et al. 1999). Virtually all of the catalytic residues of the Cet1 active site are conserved in Pct1 (Supplemental Fig. S1B,C). From the superposition, we infer that Pct1 acidic side chains Glu78, Glu80, Asp240, Glu260, and Glu262—projecting up from the tunnel floor—bind the essential metal cofactor; this inference is consistent with prior findings that mutating Glu78, Glu80, or Glu260 to alanine abolished Pct1 triphosphatase activity (Pei et al. 2001b). An ensemble of basic side chains emanating from the lateral walls and roof of the Pct1 tunnel (Arg169, Lys185, Lys227, Arg229) is implicated, by analogy to Cet1, in binding the γ phosphate and stabilizing the transition state of the phosphohydrolase reaction.

### Pct1·Spt5-CTD complex

An Spt5 CTD peptide T1P2A3W4N5 is docked on the lateral portion of pedestal domain of each protomer of the Pct1 homodimer (Fig. 1A; Supplemental Fig. S1A). (The main chain and Cβ atoms of the Arg9 residue of the preceding non-amer of the input 18-mer CTD peptide were placed into electron density, but the rest of the Arg side chain was disordered.) The N-termini of the two Spt5 CTD peptides are oriented toward each other and are separated by ∼40 Å across the protein surface, underscoring that each CTD docking
site on the homodimer is occupied by a different copy of the 18-mer CTD. We estimate that 11–12 amino acids would be required to bridge the 40 Å gap between TPAWN segments. As such, a Spt5 CTD composed of TPAWN-SGSKT PAWNSGSK-TPAWN with one intervening repeat could be sufficient to engage both sites. While it is not known if both binding sites are required for function in vivo, a S. pombe mutant strain with a Spt5 CTD containing just three repeats was sufficient for normal growth (Schneider et al. 2010). The surface view in Figure 1C highlights the proximity of the Spt5 CTD to the homodimer interface. The close-up stereo view of the CTD docking site (Fig. 2A) affirms that elements of both Pct1 protomers contribute to the Spt5 CTD binding site.

The atomic contacts of the Spt5 CTD amino acids with Pct1, proceeding from N to C termini of the bound peptide, are as follows: Arg9-Cβ makes van der Waals contacts with Phe109; the Arg9 main chain carbonyl is within hydrogen bonding distance of Asn106-Nδ; Pro2 makes van der Waals contact with Glu108 and with Met100 of the other protomer; Trp4 makes van der Waals contacts with Val60, Phe64, and His67, and with Met100 of the other protomer; Asn5-Oδ is within hydrogen bonding distance of Asn63-Nδ, while Asn5-Nδ is within hydrogen bonding distance of Asn197-Oδ of the other protomer (Fig. 2A). A noteworthy feature of the CTD conformation is that the peptide backbone makes a tight U-turn about Trp4 so that Pro2 stacks over the Trp4 indole (Fig. 2A,B). The centrality of Trp4 to the crystallographic Pct1·Spt5-CTD interface rationalizes previous findings that Pct1 failed to bind in vitro to an Spt5 CTD nonamer repeat array in which every Trp4 was replaced by alanine (Schneider et al. 2010). The Spt5 CTD Thr1 side chain makes no contacts with Pct1 in the crystal (Fig. 2A), consistent with an earlier report that mutating every Thr1 to alanine did not affect Pct1 binding to an Spt5 nonamer array (Schneider et al. 2010).

The fission yeast Pct1106NPEF109 motif that engages the Spt5 CTD is not conserved in budding yeast Cet1; indeed, the PEF109 tripeptide is an inserted element at the tip of an interstrand loop (denoted by the arrow in Supplemental Fig. S1A, left panel) that is not even present in Cet1, where the corresponding shorter loop is displaced compared with its position in Pct1 (see arrow in Supplemental Fig. S1A, right panel). This motif acquisition in Pct1 likely contributes to its distinctive interactions with the Spt5 CTD.

Alignment of the structures of the Pct1-Spt5-CTD complex and the Pct1 apoenzyme reveals an Spt5-induced local conformational change, depicted in Figure 3, where the apoenzyme fold and pertinent side chain carbons are colored beige and the Spt5-bound Pct1 fold and side chain carbons are colored green. The positions of most of the Spt5-interacting amino acids (e.g., Val60, Asn63, Phe64, His67, Met100, Asn106, and Asn197) are unchanged in the two structures. In contrast, residues 108 to 112 undergo a significant rearrangement. In the apoenzyme, this segment adopts a 310 helix.
Pct1 homodimer interface

The homodimer is stabilized by two distinct Pct1-Pct1 interfaces, each repeated twice across the twofold axis. The pertinent cross-protomer contacts are depicted in Figure 2C. The first interface is between the N-terminal leader peptide and the pedestal of the other protomer. Specifically, the Glu45 carboxylate makes a bidentate salt bridge to Arg143 and is within hydrogen bonding distance of His234; the Glu45 main chain carbonyl and amide nitrogen hydrogen bond with Gln235 Nε and Oε; the Phe48 main chain carbonyl and amide nitrogen hydrogen bond with the Arg143 and Asn294 side chains, respectively; and the Val54 main chain carbonyl receives hydrogen bonds from the Arg300 guanidinium nitrogens (Fig. 2C). The second dimer interface abuts the Spt5 CTD (shown as a thin stick model in Fig. 2C) and involves several of the amino acids that contact the CTD. This dimer interface entails the following: van der Waals interaction of Asn106 with Met100 and of Phe109 with Pro98; hydrogen bonds between Ile104 and Asn106 main chain atoms in one protomer and the Asn101 side chain and Met100 and Pro98 main chain atoms in the other protomer; and a hydrogen bond between Asp56 and Asn197 side chains (Fig. 2C). Many of the charged and polar amino acids at the Pct1 homodimer interface are conserved in Cet1. For example, Pct1 Asp56 is the counterpart of Cet1 Asp280, mutation of which to alanine converts the budding yeast triphosphatase into a catalytically active monomer that is unable to function in vivo (Hausmann et al. 2003).

Effect of threonine phosphorylation of the Spt5 CTD on interaction with Pct1

We implemented a fluorescence polarization assay to study the binding of Pct1 to purified recombinant Spt5-(801–898) protein, comprising 10 nonamer repeats (Pei and Shuman 2002), that was labeled at the N-terminus with BODIPY-FL. Fluorescence anisotropy increased with Pct1 concentration, with an apparent $K_d$ of 0.68 ± 0.10 µM (Fig. 4A). We also gauged Pct1 binding to a fluorescein-labeled 22-amino acid synthetic Spt5 CTD peptide SGSKTPAWNS GSKTPAWNSGSK that contained two complete nonamer repeats. The apparent $K_d$ of 124 ± 10 µM (Fig. 4B) was 180-fold higher than the $K_d$ for Pct1 for the longer Spt5-(801–898) ligand. This acute dependence on the number of nonamer repeats for high-affinity binding of Pct1 to Spt5 CTD in the fluorescence assay accords with results obtained using a yeast two-hybrid interaction assay, which showed that Pct1 binding to the Spt5 CTD in vivo required a long tandem nonamer repeat array (Pei and Shuman 2002). The Pct1-Spt5-CTD structure suggests how a longer nonamer array would have high avidity, based on simultaneous occupancy of both CTD docking sites on the Pct1 homodimer.

Threonine-1 of the S. pombe Spt5 nonamer motif is targeted for phosphorylation by Cdk9. Here and in previous studies, we found that Pct1 binds to the unphosphorylated Spt5 CTD. However, it is unclear whether and how the Pct1·Spt5 interaction is affected by phosphorylation. Although the Thr1 side chain does not contact the Pct1 interface in our crystal structure of Pct1 bound to the peptide TPAWN (Fig. 2A), the addition of a phosphate group at Thr1-Oγ could clash with the tight hairpin turn conformation of the Pct1-bound CTD peptide if the Thr1 side chain rotamer is maintained after phosphorylation. While a clash could be alleviated if the Thr1 side chain adopted an alternative rotamer, the structure engenders a prediction that Thr1 phosphorylation might be inimical to Spt5 CTD binding to Pct1.

To evaluate the impact of Thr1 phosphorylation, we tested the ability of unphosphorylated and Thr1-phosphorylated 22-amino acid synthetic peptides SGSKTPAWNSGSKTPA WNSGSK and SGSK(Tp)PAWNSGSK(Tp)PAWNSGSK to compete with a fluorescein-tagged 22-mer Spt5 CTD peptide for binding to Pct1 in a fluorescence polarization assay. Whereas unphosphorylated 22-mer CTD displaced the fluorescent CTD with a $K_i$ of 79 ± 9 µM, the Thr1-phosphorylated CTD did so with a $K_i$ of 2600 ± 280 µM (Fig. 4C). Thus, Thr1
phosphorylation elicited a 33-fold decrement in affinity of the fission yeast TPase for the Spt5 CTD.

**Functional probing of the Pct1-Spt5-CTD interface**

Mutations of Pct1 amino acids comprising the Spt5 CTD docking site, and a three amino acid internal deletion of the Pct1-specific loop segment at the docking site (Δ^{106}NPE^{108}), were introduced by marked allelic replacement into one chromosomal pct1 locus of a diploid *S. pombe* strain. The diploids were sporulated and viable haploids were obtained and genotyped to verify the allelic replacement. Serial dilutions of the wild-type and mutant strains containing equal numbers of cells were spot tested for growth on YES agar medium at 20°C, 25°C, 30°C, 34°C and 37°C. The M100A, N197A, and N197R mutants grew as well as wild-type cells at all temperatures (as gauged by colony size). The F64A, H67A, and Δ^{106}NPE^{108} mutants thrived at 20°C–25°C, but displayed severe temperature-sensitive (ts) growth defects at 30°C–37°C. The M100R strain grew slowly at 20°C–30°C and failed to thrive at 37°C (Fig. 5A). Because of the overlap between the Spt5 CTD and homodimer interfaces of Pct1, and prior studies indicating that weakening the homodimer interface of Cet1 elicits a ts growth defect and renders the protein thermolabile in vitro (Hausmann et al. 2003), we surveyed the mutant strains for their steady-state levels of Pct1 protein by Western blotting of whole-cell extracts of equal numbers of cells grown in liquid medium at 30°C. Probing the blot with anti-Spt5 antibody verified comparable sample loading. However, probing with anti-Pct1 antibody showed that the steady-state levels of Pct1 were decreased in F64A, H67A, and Δ^{106}NPE^{108} cells, likely accounting for the growth defects of these strains (Fig. 5B). In contrast, the Pct1 level in M100A and M100R cells were comparable to wild-type cells at comparable sample loading. However, probing with antibodies to fission yeast Spt5 and Pct1. The positions and sizes (kDa) of polypeptide size markers are denoted on the left. (C) Purified recombinant wild-type Pct1 and mutants M100A and M100R were assayed for manganese-dependent ATP hydrolysis as described under Materials and Methods. The extents of phosphate release (from 1 nmol of input ATP) are plotted as a function of reaction time. (D) Mutations that disrupt binding to the Spt5 CTD do not affect binding to the phosphorylated Pol2 CTD. Wild-type Pct1 and the indicated mutants were assayed for binding to BODIPY-FL-Spt5 CTD-(801–898) and BODIPY-FL-Pol2 CTD-Ser5-PO4 ligands. The Kd values are shown.
PO4 was also dependent on phosphorylation as interactions (Supplemental Fig. S2B). Interaction with BODIPY-Pol2-CTD-PO4 with an unlabeled CTD ligand (Supplemental Fig. S2A), interactions that were reduced to 90.2 ± 5.2 (Supplemental Fig. S2A), interactions that were reduced to 90.2 ± 5.2 (Supplemental Fig. S2A) by recombinant wild-type Pct1, arguing against a catalytic defect as the cause of the M100R growth phenotype (Fig. 5C).

We proceeded to test the binding of recombinant M100A and M100R proteins to the BODIPY-FL-Spt5-(801–898) ligand by Pct1 titration in the fluorescence polarization assay. The $K_d$ values were 1.4 ± 0.15 µM for M100A and 21.4 ± 2.4 µM for M100R, versus 0.68 ± 0.10 µM for wild-type Pct1. Thus, M100A, which had no apparent effect on fission yeast growth, had a twofold effect on Sp5 CTD binding (which presumably reflects loss of the van der Waals contact of Met100 with the CTD), whereas M100R, which compromised yeast growth, elicited a 31-fold decrement in Sp5 CTD binding, likely via steric hindrance at the CTD docking site (Fig. 2A). To confirm that defects associated with M100 mutations were specific to Sp5 we proceeded to test the binding of Pct1 to a phosphorylated 28 amino acid Pol2 CTD ligand containing four YSPTSPS consensus motifs. Pct1 bound to BODIPY-Pol2-CTD-PO4 with a $K_d$ value of 90.2 ± 5.2 (Supplemental Fig. S2A), interactions that were competed effectively with an unlabeled CTD ligand (Supplemental Fig. S2B). Interaction with BODIPY-Pol2-CTD-PO4 was also dependent on phosphorylation as interactions with the same ligand were decreased by at least an order of magnitude after treatment with calf-intestinal alkaline phosphatase (Supplemental Fig. S2A). No defects were observed in binding for recombinant M100A and M100R proteins to BODIPY-Pol2-CTD-PO4 (Fig. 5D). We infer that the M100R allele reflects the contribution of Sp5 CTD interaction to Pct1 activity in vivo.

**DISCUSSION**

**Structural plasticity of the Sp5 CTD**

The fission yeast Sp5 CTD has a relatively uniform primary structure consisting of tandem repeats of a nonapeptide motif that extend to the very C-terminus of the 990-amino acid Sp5 protein (Pei and Shuman 2002). Metazoan Sp5 proteins have an analogous repeat domain, often referred to as the CTR (C-terminal repeat), located near but not at the C-terminus (Hartzog and Fu 2013). Whereas the repeated elements of fission yeast Sp5 CTD and metazoan Sp5 CTRs characteristically include a Thr-Pro or Ser-Pro dipeptide (targeted for threonine or serine phosphorylation by a cyclin-dependent kinase), the spacer intervals between the dipeptide repeats differ among metazoan species, and they also differ from the regular spacing in *S. pombe* Sp5. The metazoan repeats do not adhere to the S. pombe consensus sequence TPAWNSGSK. In particular, whereas the residue located two positions downstream from the Thr-Pro dipeptide in Sp5 is typically a tryptophan, it is never a tryptophan in the human, nematode, or zebrafish proteins; instead, this position is usually occupied by tyrosine or histidine (Pei and Shuman 2002). The consensus view is that the Sp5 CTD/CTR functions to recruit cellular proteins to elongating Pol2 complexes with which Sp5 is engaged.

The present structural snapshot of the RNA capping enzyme Pct1 bound to Sp5 CTD sequence TPAWN extends insights from the recently reported crystal structure of a human Sp5 CTR binding protein (Wier et al. 2013): the Ptt3 domain of Rtf1 bound to a phosphorylated CTR peptide SGRSTPMYGSQ. Figure 6 shows an alignment of the Sp5 ligands from the two structures, superimposed at their respective Thr-Pro dipeptides. The proline is in the trans conformation in both structures. In the case of Rtf1, the Sp5 interface is dominated by hydrogen bonding contacts to the phosphate group of the Sp5 ligand, consistent with the fact that threonine phosphorylation is required for Rtf1 binding to Sp5-CTR (Wier et al. 2013). In contrast, Pct1 makes no contacts with the Sp5 threonine side chain and its binding to Sp5 is antagonized by threonine phosphorylation. The phosphotheorneine in the Rtf1 complex adopts a different rotamer configuration than does threonine in the Pct1 complex. Although our structure did not visualize the SGRST peptide segment preceding Thr1, it is noteworthy that the amino acid sequence of this segment of *S. pombe* Sp5 is identical to the threonine-flanking peptide segment present in the Rtf1 structure (Fig. 6). The two Sp5 CTD conformations diverge sharply after the Ca of the amino acid following the proline. In Rtf1, the bound Sp5 segment YGSQ adopts a $\beta\alpha\beta$ helix-like conformation that bears no resemblance to the tight hairpin turn made by the WN dipeptide of the Pct1-bound Sp5. Indeed, the positions of the aromatic tyrosine and tryptophan side chains of the human and *S. pombe* Sp5 are different. Whereas the tyrosine is a minor contributor to the interface with Rtf1 (i.e., compared with phosphotheorneine),
the fission yeast tryptophan is the dominant player at the interface with Pct1 (Fig. 2A). The comparison of the fission yeast and human Spt5 CTD/CTR complexes highlights the theme that the Spt5 CTD is structurally plastic and can adopt different conformations that are templated by particular cellular Spt5 CTD receptor proteins. In these respects, the Spt5 CTD resembles the Pol2 CTD.

An Spt5 CTD code

Spt5 binds to and travels with elongating Pol2, helping to coordinate elongation kinetics with RNA processing and chromatin modification (Hartzog and Fu 2013). Several of the key effector functions of Spt5 rely on its CTD/CTR as a platform to recruit cellular proteins to the Pol2 elongation complex. Available evidence suggests that threonine phosphorylation of Spt5 CTD repeats inscribes a binary code (on-off) that is read by diverse CTD receptors, each in its own distinctive manner. Whereas the Paf1 complex subunit Rtf1 recognizes only the phosphorylated Spt5 CTR (Wier et al. 2013), the RNA guanylyltransferase Pct1 (Doamekpor et al. 2014) and the RNA triphosphatase Pct1 (present study) specifically bind to the unmodified form of the CTD. Moreover, the positive elongation function of Spt5 depends on phosphorylation of the CTD/CTR by Cdk9 (Yamada et al. 2006).

The organization of the Spt5 CTD/CTR as a tandem array of Thr/Ser-Pro-containing motifs allows (in principle) for simultaneous engagement of multiple cellular CTD receptors that respond to different CTD cues. Taking into account just the on-off phosphorylation state of the repeated Thr-Pro and Ser-Pro motifs, the fission yeast Spt5 CTD can adopt up to 262,144 different primary structures. Cdk9 is a principal Spt5 CTD kinase in fission yeast (Pei and Shuman 2003; Viladevall et al. 2009), but it is conceivable that other kinases may also contribute to the writing of an Spt5 CTD code. How the code is edited by Spt5 CTD phosphatases is still uncharted territory, at least in fission yeast. Finally, by analogy to the Pol2 CTD, it is conceivable that the complexity of the Spt5 CTD code is amplified by cis-trans isomerization of the defining proline residue of the repeated Spt5 motif.

Two key themes highlighted by structural, biochemical, and genetic analyses of the reading of the Pol2 and Spt5 CTD codes by the fission yeast capping apparatus are (i) that CTD phosphorylation has opposite effects on the interaction of fission yeast guanylyltransferase and triphosphatase with Pol2 (Ser5-PO4 is required for binding) versus Spt5 (Thr1-PO4 inhibits binding) and (ii) the two CTD codes function in parallel to aid cotranscriptional capping.

MATERIALS AND METHODS

Purification of S. pombe Pct1

The pct1 open reading frame was inserted into pET-based bacterial expression plasmid pSMT3 so as to encode Pct1 fused to an N-terminal His6Smt3 domain. Mutations were introduced into the pSMT3-Pct1 plasmids with a QuickChange kit (Stratagene). The mutated pct1 ORFs were sequenced to verify the absence of unwanted coding changes. The expression plasmids were transformed into E. coli BL21(DE3) codon plus RIL (Stratagene). Cultures of E. coli containing the expression plasmids were grown in super broth in baffled flasks at 37°C to an A600 of 1.5, after which the cells were cold-shocked by placing the flasks on ice for 30 min. The cultures were adjusted to 0.5 mM isopropyl-β-D-thiogalactoside (IPTG) and then incubated overnight at 18°C with constant shaking. Cells were harvested by centrifugation at 16,000g and resuspended in buffer containing 50 mM Tris–HCl pH 8.0, and 20% (w/v) sucrose. All subsequent procedures were performed at 4°C.

For purification of the His6Smt3-tagged Pct1 proteins, cells were resuspended in lysis buffer containing 20 mM Tris–HCl, pH 8.0, 500 mM NaCl, 20 mM imidazole, 1 mM β-mercaptoethanol (BME), 0.1% IGEPAL (Fluka), 1 mM phenylmethanesulfonylfluoride (PMSF) and then disrupted by sonication. Lysates were clarified by centrifugation at 45,000g and the resulting supernatants were applied to a Ni-NTA superflow resin (Qiagen) that had been equilibrated with 50 mM imidazole buffer (Pct1). The Ni-NTA resin was washed with a 20-column-volume linear gradient of 50 mM to 1 M NaCl in 20 mM Tris-HCl, pH 8.0, 1 M NaCl, 1 mM BME) containing 20 mM imidazole. Bound proteins were eluted with 250 mM imidazole in buffer A. The His6Smt3 tags were removed by incubating the 250 mM imidazole eluate fractions with Ulp1, a Smt3-specific protease (Mossessova and Lima 2000), at a protein/Ulp1 ratio of 1000:1 (w/w) for 6–18 h at 4°C. The tag-free Pct1 proteins were separated from His6Smt3 by gel-filtration through a Superdex 200 column equilibrated in buffer A. The peak Pct1 fractions were pooled, dialyzed against buffer B (20 mM Tris–HCl, pH 8.0, 50 mM NaCl, 1 mM BME) and concentrated by centrifugal ultrafiltration. For crystallography experiments, wild-type Pct1 was further purified by cation exchange chromatography. The Superdex fraction was adsorbed to a MonoS 10/10 column (GE Healthcare) and eluted with a 20-column-volume linear gradient of 50 mM to 1 M NaCl in 20 mM Tris–HCl, pH 8.0, 1 mM BME. Pct1 was then concentrated to 10 mg/mL in 20 mM Tris–HCl, pH 8.0, 50 mM NaCl, 1 mM TCEP.

Pct1 crystallization and structure determinations

Pct1 apoenzyme

Aliquots of a mixture of 140 µM Pct1, 2 mM triplyorthophosphate, 5 mM manganese chloride were mixed on a cover slip with an equal volume of precipitant solution containing 2.8 M sodium chloride, 10% (w/v) PEG6000. Crystals were grown at 18°C by hanging drop vapor diffusion against the precipitant solution, then flash-frozen in liquid nitrogen. Diffraction data from a single crystal to 2.6 Å resolution were collected on a Rigaku RAXIS-4 image plate detector with CuKα radiation from a Rigaku RU200 rotating anode generator. The data were indexed, integrated and scaled using HKL2000 (Otwinowski and Minor 1997). The crystal was in space group P21. Initial phases were obtained by molecular replacement in PHASER (McCoy et al. 2007) using the coordinates of the Cet1 homodimer (pdb 1D8I) as the search model. Iterative rounds of refinement and adjustment in CNS (Brünger et al. 1998), PHENIX (Adams et al. 2010), and COOT (Emsley and Cowtan 2004) yielded a 2.6 Å Pct1 model with R/RFree of 0.199/0.245 and good geometry (97.9%, 99.9%, and 0.1% in favored, 1.0% in allowed, and 0.8% in outlier parameters).
allowed, and disallowed regions of Ramachandran space, respectively) (Chen et al. 2010). The model contained four Pct1 protomers in the asymmetric unit organized as two homodimers (A/B and C/D). No electron density was observed for tripolyphosphate or manganese.

**Pct1-Spt5-CTD complex**

A mixture of 140 μM Pct1 and 340 μM 18-mer Spt5-CTD peptide T1P2A3W4N5S6G7S8R9T1P2A3W4N5S6G7S8K9 was incubated for 1 h at 4°C, then mixed on a cover slip with an equal volume of 3 M sodium formate. Crystals were grown at 18°C by hanging drop vapor diffusion against 3 M sodium formate and cryoprotected with 3.2 M sodium formate, 20% (v/v) ethylene glycol. Diffraction data from a single crystal to 2.8 Å resolution were collected at APS NE-CAT beamline 24-IDE equipped with an ADSC Quantum 315 detector. Data were indexed, integrated and scaled using HKL2000. The crystal was in space group P2₁2₁2₁ with R/ Rfree of 0.191/0.248 and good geometry (96.0%, 100%, and 0% in most favored, allowed, and disallowed Ramachandran regions, respectively). The model contained four Pct1 protomers in the asymmetric unit organized as two homodimers (A/B and C/D). Each homodimer bound two copies of a Spt5-CTD peptide, of which five amino acids corresponding to TP A W N were ordered in electron density.

**Assays of Pct1-CTD interactions by fluorescence polarization**

Recombinant N(Cys)-Spt5-(801–898) protein was produced in *E. coli*, purified, and labeled with BODIPY-FL at the N-terminus using maleimide chemistry as described previously (Doamekpor et al. 2014). A fluorescein-labeled 22-mer synthetic Spt5-CTD peptide SGSKTPAWNNSGSKTPAWNNSGSK was purchased from New England Peptides. A 28-mer synthetic Pol2 CTD-Ser5-PO₄ peptide (YSPTS₅-PYSPYSPTS₃-PYSPYSPTS₃-PYSPYSPTS₃) was labeled with BODIPY-FL at the N terminus via N-hydroxy succinimimidyl ester chemistry as previously described (Doamekpor et al. 2014).

Binding reaction mixtures (18 μL) containing 20 mM Tris–HCl, pH 8.0, 50 mM NaCl, 4 mM dithiothreitol (DTT), either 300 nM BODIPY-FL-N(Cys)-Spt5-(801–898), or 25 nM fluorescein-labeled 22-mer synthetic Spt5-CTD peptide and increasing concentrations of Pct1 polypeptide as specified were incubated on ice for 15 min. For binding reactions of Pct1 with the Pol2-CTD, 400 nM untreated labeled BODIPY-FL-Pol2 CTD-Ser5-PO₄ or labeled BODIPY-FL-Pol2 CTD-Ser5-PO₄ which had previously been incubated with 0.5 units alkaline phosphatase (CIP, New England Biolabs) at 37°C for 2 h were used. Aliquots (15 μL) were transferred to wells of a 384-well microplate and fluorescence polarization was measured at 23°C with a SpectraMax M5 microplate reader (Molecular Devices) using an excitation wavelength of 485 nm, emission wavelength of 525 nm, and a cutoff of 515 nm. Experiments were performed in triplicate except for the reaction involving the CIP treated Pol2-CTD which was performed in duplicate. For the experiments using a 22-mer amino acid Spt5-CTD ligand or a 28-mer amino acid Pol2-CTD, in which the Pct1 concentration that achieved 50% saturation was at least three times greater than the concentration of the labeled ligand, the Kᵦ value was derived by fitting the data in Prism to a single-site binding model (Lundblad et al. 1996). For experiments with Spt5-(801–898), in which the Pct1 concentration that achieved 50% saturation was similar to the concentration of labeled ligand, the Kᵦ value was derived by fitting the data in Prism to a single-site binding model that accounts for receptor depletion (Lundblad et al. 1996). For competition experiments, Pct1 was incubated with serial dilutions of the unlabeled competitor for 15 min on ice. Fluorescent-labeled Spt5-CTD or Pol2-CTD-PO₄ was then added and incubation was continued for another 15 min on ice. Aliquots (15 μL) were transferred to the wells of a 384-well microplate, and measurements were obtained as described above. In assays with the 22-mer Spt5-CTD, the fluorescent ligand and Pct1 concentrations were 25 nM 22-mer fluorescein-Spt5-CTD and 120 μM Pct1 polypeptide. In the assays with the Pol2-CTD, the fluorescent ligand and Pct1 concentrations were 400 nM BODIPY-FL-Pol2 CTD-Ser5-PO₄ and 40 μM Pct1 polypeptide. Experiments were performed in triplicate, and data were fit in Prism to a one-site binding model for competitor.

**Effects of CTD interfacial mutations on Pct1 function in vivo**

Alleric replacement at chromosomal pct1⁺ locus was performed as follows. We constructed series of integration cassette plasmids containing, in the 5′–3′ direction, (i) a 1.4-kbp segment encompassing the pct1 cDNA (wild-type or mutated) and 484-bp of 5′ flanking chromosomal DNA; (ii) a 268-bp DNA segment harboring the nmt1 transcription termination signal; (iii) a hygMX cassette conferring resistance to hygromycin; and (iv) a 498-bp segment of 3′ flanking pct1 chromosomal DNA. The 5′-pct1–hygMX-pct1-3′ integration cassettes were excised from the plasmids and transfected into *S. pombe* diploids. Transformants selected for growth on YES agar containing 0.3 mg/mL hygromycin were genotyped by Southern blotting to confirm correct integration of hygMX at one of the pct1 loci. The pct1–hygMX allele was then PCR-amplified and sequenced to confirm the presence of the desired mutation(s). The heterozygous pct1⁺ pctl–hygMX strains were sporulated. A random population of haploid progeny (~2500, as gauged by plating on YES agar) was plated on YES-hygromycin agar. Viable pct1–hygMX haploid strains formed colonies on selective hygromycin agar at frequencies consistent with random segregation. To gauge the effect of the pct1 mutations on vegetative growth, cultures of haploid *S. pombe* pct1–hygMX strains were grown in liquid medium at 30°C until A₆₀₀ reached 0.6–0.9. The cultures were adjusted to a final A₆₀₀ of 0.1 and aliquots (3 μL) of serial fivefold dilutions were spotted on YES agar. The plates were incubated at 20°C, 25°C, 30°C, 34°C, and 37°C.

**ATPase activity**

Reaction mixtures (100 μL) containing 20 mM Tris–HCl, pH 8.0, 50 mM NaCl, 0.5 mM DTT, 2 mM MnCl₂, 0.2 mM ATP, and 40 nM of
the Superdex preparations of wild-type or mutant Pct1 polypeptide were incubated at 30°C. At the times specified, aliquots (5 µL) were removed and mixed with 100 µL Biomol green reagent (Enzo) for 30 min at 22°C before measuring the A_{570}. The molar amount of phosphate released was determined by interpolation to a phosphate standard curve.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

ACKNOWLEDGMENTS

We thank NE-CAT beamlines (Advanced Photon Source) supported by RR-15301 (NIH NCCR). APS is supported by the US Department of Energy, Office of Basic Energy Sciences, under Contract No. DE-AC02-06CH11357. Research reported in this publication was supported by the National Institute of General Medical Sciences of the National Institutes of Health under award numbers GM061906 (C.D.L.) and GM052470 (B.S. and S.S.). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. S. S. is an American Cancer Society Research Professor. C.D.L. is an Investigator of the Howard Hughes Medical Institute.

Received September 19, 2014; accepted October 24, 2014.

REFERENCES


Fission yeast RNA triphosphatase reads an Spt5 CTD code

Selom K. Doamekpor, Beate Schwer, Ana M. Sanchez, et al.

RNA published online November 20, 2014