

Different phosphoisoforms of RNA polymerase II engage the Rtt103 termination factor in a structurally analogous manner

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The carboxyl-terminal domain (CTD) of the largest subunit of RNA polymerase II (Pol II) orchestrates dynamic recruitment of specific cellular machines during different stages of transcription. Signature phosphorylation patterns of $Y_1S_2P_3T_4S_5P_6S_7$ heptapeptide repeats of the CTD engage specific "readers." Whereas phospho-Ser5 and phospho-Ser2 marks are ubiquitous, phospho-Thr4 is reported to only impact specific genes. Here, we identify a role for phospho-Thr4 in transcription termination at noncoding small nucleolar RNA (snoRNA) genes. Quantitative proteomics reveals an interactome of known readers as well as protein complexes that were not known to rely on Thr4 for association with Pol II. The data indicate a key role for Thr4 in engaging the machinery used for transcription elongation and termination. We focus on Rtt103, a protein that binds phospho-Ser2 and phospho-Thr4 marks and facilitates transcription termination at protein-coding genes. To elucidate how Rtt103 engages two distinct CTD modifications that are differentially enriched at noncoding genes, we relied on NMR analysis of Rtt103 in complex with phospho-Thr4- or phospho-Ser2-bearing CTD peptides. The structural data reveal that Rtt103 interacts with phospho-Thr4 in a manner analogous to its interaction with phospho-Ser2-modified CTD. The same set of hydrogen bonds involving either the oxygen on phospho-Thr4 and the hydroxyl on Ser2, or the phosphate on Ser2 and the Thr4 hydroxyl, can be formed by rotation of an arginine side chain, leaving the intermolecular interface otherwise unperturbed. This economy of design enables Rtt103 to engage Pol II at distinct sets of genes with differentially enriched CTD marks.

CTD code | CTD interactome | noncoding RNA | NMR | phosphothreonine

E ach stage of transcription relies on ordered recruitment and exchange of specific protein complexes that act on RNA polymerase II, its nascent transcripts, and the underlying chromatin. This dynamic process is orchestrated via patterned posttranslational modifications of the carboxyl-terminal domain (CTD). This unusual and essential domain of Rpb1, the largest component of the 12subunit polymerase, consists of repeating Y1S2P3T4S5P6S7 heptapeptides (26 repeats in budding yeast and 52 in humans) (1). The mechanistic consequences of phosphorylating Ser5 and Ser2 have been well documented (2-11). However, the role of Thr4 phosphorylation (pThr4), and even the necessity of Thr4 for cellular survival, appears to differ among closely related species and between growth conditions within a given species (12-15). Recent mass spectrometric analysis of an extensively engineered CTD revealed a paucity of pThr4, raising questions about the importance of this mark (16). In contrast, similar studies found pThr4 marks at levels comparable to or greater than the ubiquitously placed pSer2 mark in both yeast and human cells (17). Thus, much remains to be understood about the natural abundance and functional role of pThr4 marks on the endogenous, unmodified CTD.

Previous studies suggest that pThr4 has roles in transcriptional elongation, 3'-end processing, and termination (12-14, 18). Our data, as well as other recent studies, suggest that CTD bearing pThr4 is bound by Rtt103 (16, 18), a well-known component of the Rat1 exosome that is thought to play a role in transcription termination of protein-coding genes. However, the bulk of previous genetic, biochemical, and structural studies show Rtt103 binds pSer2 and may cooperatively recruit Pcf11 to the 3' ends of proteincoding genes (19, 20). Very few CTD-interacting proteins recognize multiple forms of phosphorylated CTD (3, 21-25), and the structures of even fewer are resolved. One intriguing example is Ssu72, which binds and dephosphorylates pSer5-CTD or pSer5-pThr4 peptides in the opposite orientation as pSer7-CTD peptides (22, 24, 26-28). How does Rtt103 discriminate between pSer2 and pThr4 CTD? Does Rtt103 use the same interface to engage pThr4 or pSer2 marks? Does pThr4-bound Rtt103 impact similar genes as Rtt103 recruited by pSer2? Here, we aim to answer these questions and further elucidate the role of pThr4 in Pol II function.

Significance

Stage-specific and gene-specific molecular machines are recruited to elongating RNA polymerase II (Pol II) through reversible phosphorylation of its carboxyl-terminal domain. This unusual domain is composed of a tandemly repeating $Y_1S_2P_3T_4S_5P_6S_7$ motif. Here, we identify a class of noncoding RNA that relies on phospho-Thr4 for effective termination. We also identify protein complexes that rely on Thr4 to associate with Pol II. Rtt103, one of the proteins that engages phospho-Thr4, also binds phospho-Ser2 and facilitates transcription termination of protein-coding genes. Using NMR, we show that Rtt103 binds with phospho-Thr4 in a nearly identical manner as phospho-Ser2. Our genomic, proteomic, and structural data suggest that phospho-Ser2 and phospho-Thr4 enable the recruitment of Rtt103 to different gene classes.

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Fig. 1. pThr4 levels correlate with snoRNA termination in *S. cerevisiae*. (A) Rpb1 with mutant CTDs are shown with black (WT), orange (T4A), and blue (T4E) CTD repeats. Plasmids bearing mutant Rpb1 and a LEU2 auxotrophic marker were transformed into yeast. Shown are serial dilutions of cells in the presence of 5-FOA, which counterselects against yeast retaining the URA3-bearing plasmid containing WT Rpb1. –LEU, agar plates lacked leucine; OD, optical density; –URA, agar plates lacked uracil. (*B*) T4A expression (from genome tiling microarrays) normalized to WT is plotted across the whole yeast genome with chromosomes (Chr.) listed below. Transcripts above the red line denote regions with $z \ge 5$ (*protein-coding genes). (*C*) RNA expression (*Top*) and Rpb3 ChIP analysis (*Bottom*) from WT (black), T4A (orange), or S2AWT (green) are shown at *SNR33*. Arrows denote the transcription start site, black bars indicate gene bodies, and red lines indicate cleavage and polyadenylation sites. Northern blot shows probing *SNR33* (*Right*). IP, immunoprecipiate. *SCR1* is a loading control. (*D*) Same as *C* at the *SNR40* locus.

We report that substituting all threonine 4 residues in the Saccharomyces cerevisiae CTD with alanines (T4A) results in defective transcription of a coherent set of small nucleolar RNA (snoRNA) genes. Genome-wide chromatin immunoprecipitation (ChIP) studies show high pThr4 and low pSer2 levels at snoRNA genes that display defective termination. Consistent with this observation, snoRNA that are unaffected by T4A substitution show high pSer2 and low pThr4 levels. Quantitative mass spectrometry with T4A bearing Pol II revealed dramatically reduced binding of discrete protein complexes involved in transcription elongation and termination. Among the many partners, Rtt103 displayed the greatest sensitivity to the removal of the Thr4 side chain. Remarkably, our NMR studies show that Rtt103 binds analogously to pThr4 and pSer2 peptides and that a key Arg (R108) within Rtt103 mediates key intermolecular interactions with both phosphorylated peptides. We propose that, whereas pSer2 functions as a generic termination mark at most Pol II-transcribed genes, pThr4 engages the transcriptional termination machinery in a gene class-specific manner.

Results

Thr4 Is Required for Transcription Termination. To further characterize the role of pThr4 in budding yeast, we examined the effect of Rpb1 alleles in which Thr4 was substituted with an alanine (T4A) or glutamate (T4E). Alanine substitution is commonly used to map binding interfaces because it is considered a benign perturbation, and this residue occurs in structured and unstructured protein folds (29). Moreover, alanine cannot be phosphorylated and does not replace the polar hydroxyl moiety of threonine with a hydrophobic methyl group (as is the case with a valine substitution). Alanine substitutions were made across all heptad repeats or in the proximal or distal halves of the CTD. We also created a strain in which Thr4 was substituted with the phosphomimic glutamate (T4E) in each repeat (Fig. 14, *Left*). Whereas the T4E strain was inviable, the T4A strain displayed a slow growth phenotype (Fig. 14, *Middle Right*).

Under exponential growth conditions in rich medium, the transcriptomes of WT and T4A cells displayed few global differences when examined with high-density genome tiling microarrays (30, 31). Avian histone transcripts require pThr4 to process 3' ends and are not polyadenylated (12); however, in budding yeast, histone genes are polyadenylated (32). Consistent with this species-specific difference, no defects in 3'-end processing of histone genes were observed in T4A-bearing yeast strains (*SI Appendix*, Fig. S1). Interestingly, we observed a minor but significant correlation between T4A and a strain containing a shortened CTD (33) (*SI Appendix*, Fig. S24 and Dataset S1).

Plotting the fold change in expression between T4A and WT, ~50 striking peaks were observed (Fig. 1*B*). Using a stringent *z*-score cutoff of \geq 5, we found significant enrichment at 3' ends of snoRNAs. In the T4A strain, 37 of 77 snoRNA transcripts extended well past the mature 3' end (Fig. 1*C*, *Top* and *SI Appendix*, Fig. S3). Whereas many formed chimeric transcripts that used the polyA signals of downstream protein-coding genes to terminate transcription (e.g., *SNR33*) (Fig. 1*C*) (34–36), others were read through and terminated even in the absence of a downstream gene (e.g., *SNR48*) (*SI Appendix*, Fig. S3). In contrast, 40 snoRNAs displayed no 3'-read-through defect (e.g., *SNR40*) (Fig. 1*D*, *Top* and *SI Appendix*, Fig. S3). We further performed genome-wide transcription analysis on a strain in which Ser2 was substituted with an alanine only in the first half of the CTD (S2A/WT) and observed no significant read-through at any snoRNA gene (Fig. 1 *C* and *D* and *SI Appendix*, Table S1).

Northern blots were performed to confirm that these 3' extensions were bona fide read-through events and not spurious initiation of new transcripts. RNA from WT, T4A, or S2A/WT cells was analyzed with a probe targeting *SNR33* or *SNR40* (Fig. 1 *C* and *D*, *Right*). A strong band at ~1.5 kb corresponding to the chimeric *SNR33-YCR015C* transcript was observed in T4A but not in WT. Evidence of minor read-through was observed in S2A/WT, consistent with previous results (37). As evident, no chimeric *SNR40*-*CAF40* transcript was observed in any strain tested (Fig. 1*D*, *Right*).

In addition to Northern blot analysis of specific transcripts, genome-wide chromatin immunoprecipitation of the Rpb3 subunit of Pol II was performed to further distinguish 3'-processing defects from bona fide transcription termination defects (Fig. 1 *C* and *D*, *Bottom*). Rpb3 was strongly retained downstream of *SNR33*, further suggesting a role for pThr4 in snoRNA termination.

Thr4-Dependent and -Independent snoRNAs. We quantified readthrough at each snoRNA, as previously described (35) (Fig. 2*B*), and identified 37 snoRNAs with a 3' extension (Fig. 2*A* and *SI Appendix*, Table S1). The Pol II read-through index (from ChIP of the Rpb3 subunit) in the region downstream of the 3' end of processed snoRNAs was significantly correlated (P < 0.00001) with the 3'-RNA extension index (from RNA expression) (Fig. 2*C*). However, no correlation was observed between the read-through defect and the length (P = 0.44) or the abundance of the snoRNA transcripts (P = 0.67) (*SI Appendix*, Fig. S2 *B* and *C*).

To determine why only a subset of snoRNA genes is read through in the T4A mutant, we examined each of the four classes of snoRNA processing sites. Of the 37 snoRNAs with significant readthrough, 36 occurred at terminal processing sites of monocistronic or polycistronic transcripts ($P = 3.3 \times 10^{-6}$, Fisher's exact test) (Fig. 2D and *SI Appendix, SI Materials and Methods*). We also observed that several snoRNAs, even those with minimal 3'-end extension, displayed high levels of Pol II downstream of the termination site when assayed by genome-wide ChIP analysis (Figs. 1D and 2C). This result indicates that certain snoRNA transcripts are more defective for transcription termination rather than 3'-transcript processing.

Previous work suggests a role for pSer2 in snoRNA termination (37–39). Therefore, we examined the levels of pSer2 as well as pThr4, normalized to Rpb3 levels, across snoRNA genes in a WT strain. Remarkably, *SNR33*, a transcript with a termination defect and high read-through in T4A, displayed significantly more pThr4 downstream of the 3' end than *SNR17A*, a snoRNA transcript that displays little, if any, read-through (Fig. 2*E*, *Left*). This pattern was observed across gene compilations of all snoRNAs (Fig. 2*E*, *Right*). Concomitant with an increased pThr4 level downstream of the 3' end, lower pSer2 levels were observed

downstream of snoRNA genes that displayed defective termination in the T4A strain. Consistent with this pattern, significantly higher levels of pSer2 were observed downstream of the compilation of snoRNAs that do not show a termination defect in T4A strain (Fig. 2*F*). Striking correlations between 3'-extension index, high pThr4, and low pSer2 levels implicate pThr4 in snoRNA termination. As α -pThr4 antibodies yield modest ChIP signals, we examined pThr4 signals in other genome-wide studies of this mark (15, 40). Similar reciprocal patterns of pThr4 and pSer2 profiles were evident in pThr4 ChIP data obtained with a different (6D7) α -pThr4 antibody (40, 41) (*SI Appendix*, Fig. S44).

Thr4 Is Required for Association of Elongation and Termination Factors. Previous efforts to identify factors that bind pThr4 used pThr4specific antibodies to affinity purify RNA polymerase II bearing this mark (18). Because pThr4-bound factors may be competitively displaced by the pThr4 antibody, we sought to identify binding partners in a CTD-antibody-independent manner. We affinity enriched HAtagged Pol II from strains bearing WT, T4A, and T4E CTDs. A control strain (Z26) lacking the HA-tagged Rpb1 was subjected to identical affinity enrichment procedure. Label-free quantitative proteomics (22, 42-44) was used to identify and quantify the relative abundance of affinity-enriched complexes (Fig. 3A). For Pol II enriched from T4A extracts, dramatic reduction was observed for multiple members of the Pol II termination complex (Rtt103, Rat1, Rai1, and Pcf11), cAMP-dependent PKA kinase holoenzyme and its regulatory subunit (Tpk2, Tpk3, and Tpk1, and Bcy1), PAF1c complex that bridges elongation and termination (Paf1, Ctr9, Cdc73, Rtf1, and Leo1), elongation factors (Set2, Spt4, Spt6, and Elf1), Pol II nuclear import proteins (Npa3, Iwr1, and Gpn3), as well as Fcp1, a phosphatase that acts on pThr4 (45) (Fig. 3B and Dataset S2). Many of these proteins were not known to rely on Thr4 for association with Pol II. Furthermore, the loss of all members of a given protein complex (e.g., Rtt103, PKA, and PAF1c) strongly supports the role of the Thr4 in association with the entire complex with Pol II, and it also serves as an internal control for label-free quantitative proteomic analysis.

Next, we charted the connectivity between Thr4-dependent partner proteins (Thr4 readers) using the publicly available STRING database of physical contacts (46). Substantial interconnectivity between Thr4 readers was evident, as were two groups of high- and low-density networks (Fig. 3C). Rtt103 termination complex, PKA, and nuclear localization factors form the outer set of interacting nodes, whereas PAF1c, elongation factors, Fcp1, and Pol II subunits form an intricate high-density network. The factors in the high-density network interact with Pol II subunits, suggesting that their interaction with Pol II may be less CTD dependent than the low-density network containing the Rtt103 termination complex and PKA. This interpretation is consistent with our unbiased and unsupervised quantitative mass spectrometry data that show greatest depletion of the Rtt103 and PKA complexes in the interactome of Pol II bearing the T4A substitution. Thus, our interactome data and the network analysis with known protein-protein interaction studies are internally consistent and mutually reinforcing.

Rtt103 Binds to CTD Phosphorylated by PLK3. Our mass spectrometrybased approach relies on quantifying differential binding to Rpb1 that lacks the Thr4 side chain. Thus, we cannot exclude the possibility that the factors identified in our study necessarily bind to phospho-Thr4. Therefore, we biochemically assayed for binding of these factors to pThr4. To examine binding to the CTD, we immunopurified (1C7) hypophosphorylated Pol II and phosphorylated the CTD with PLK3, a highly selective human Thr4 kinase (Fig. 3D, *Top*) (13). Significant phosphorylation of Thr4 was observed with negligible phosphorylation at other residues of the CTD (Fig. 3D, *Left*). Tandem affinity purification (TAP)-tagged Rtt103, a representative of the termination complex; Pa11-TAP of the eponymous PAF1 complex; Tpk1-TAP, the PKA subunit that



Fig. 2. Thr4-dependent and -independent snoRNAs. (*A*) The 3'-extension index (EI) for each snoRNA in T4A is plotted from highest to lowest. Orange bars indicate an El of >1.7. (*B*) Example gene (*SNR13*) and equation used to calculate EI, defined as fold change in the average expression 200 bp downstream of the 3' end of snoRNAs. (*C*) Comparison of RNA extension index to Rpb3 ChIP read-through, defined as fold change in the average Rpb3 ChIP occupancy in the 1,000-bp window downstream of the 3' end of snoRNAs. (*D*) Four snoRNA classes are illustrated with black boxes representing snoRNAs and purple representing exons. The fractions of snoRNAs in each class with 3' extensions in T4A are noted next to the red dotted line depicting the mature 3' end of respective snoRNAs. (*E*) pThr4 occupancy (normalized to Rpb3) centered at the 3' end of *SNR33* or *SNR17A* (*Left*) or a gene compilation of pThr4 occupancy across all snoRNAs read-through in T4A (dark) or no read-through (light). Shaded regions indicate 95% confidence interval. (*F*) Similar traces for pSer2.

associates with Pol II across the transcribed gene (47); and Set2-TAP, an H3K36 methyltransferase, were expressed and purified from budding yeast and assayed for binding to pThr4bearing Pol II. Rtt103 showed a dramatic increase in binding over the unphosphorylated Pol II, further suggesting an interaction between the termination factor and the pThr4-bearing CTD (Fig. 3D, *Right*). Purified Paf1, Tpk1, and Set2 did not bind appreciably to Thr4-phosphorylated CTD (Fig. 3D, *Right* and *SI Appendix*, Fig. S5B). To examine the role of pThr4 in recruiting Rtt103 in vivo, we performed genome-wide ChIP analysis of Rtt103 in WT and T4A cells. In T4A, we observed delayed recruitment of Rtt103 at read-through snoRNAs (~160 bp 3'), confirming the role of pThr4 levels in engaging the termination machinery in vivo (*SI Appendix*, Fig. S4*B*).

Rtt103 Forms a Structurally Analogous Interface with Different Phosphoisoforms of the CTD. To elucidate the molecular basis of pThr4-CTD binding to Rtt103, we performed NMR titrations with a diheptad phosphorylated at Thr4 on repeat "a" and repeat "b" (denoted as pThr4_{ab}-CTD) and compared it to a pSer2_{ab}-CTD diheptad peptide studied by us in the past (20) (Fig. 4.4). The first 131 residues of Rtt103, comprising the CTD interacting domain (CID), was expressed and purified in bacteria. Peptide binding to the CID of Rtt103 was examined using 2D ¹H-¹⁵N heteronuclear

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Fig. 3. Thr4 is required for association of elongation and termination factors. (A) Flowchart of the method to identify factors whose binding is compromised in T4A. (*B*) Volcano plot illustrating the fold change ln(T4A/WT) and the significance $-Log_{10}$ [false discovery rate (FDR)] of differential binding. Proteins depleted at least one natural log in T4A with an FDR of <0.05 are color coded and labeled as in C. (C) Physical interconnectivity (46) between the significantly depleted proteins identified in *B*. Gray background indicates low density connectivity. (*D*) Flowchart of the experimental design to assay binding of factors to human CTD (*Top*). Phosphorylated of HeLa extract, unphosphorylated human Rpb1, or Rpb1 phosphorylated with human PLK3 (*Left*). Enriched hypophosphorylated Pol II (Thr4-CTD) and PLK3-treated CTD (pThr4-CTD) was incubated with the indicated TAP-tagged proteins, expressed in yeast. Indicated proteins were assayed for binding to pThr4-bearing Rpb1 and probed with α -TAP (*Right*).

single quantum coherences (HSQCs) to monitor chemical shift perturbations (CSPs) of amide resonances upon addition of the respective peptide. The ¹H-¹⁵N HSQCs of the CID bound to the pThr4-marked peptide overlays well with the ¹H-¹⁵N HSQC of the CID bound to the pSer2-marked peptide, indicating that the CID forms a structurally analogous interface with both phosphoisoforms (Fig. 4B). By comparing the chemical shift perturbation upon addition of each peptide (Fig. 4C), we observed that the two peptides affect the chemical shifts of Rtt103-CID in the same way, even though they are phosphorylated at different positions. The chemical shift deviations from random coil for backbone carbons confirm that the secondary structure of Rtt103 is unchanged (Fig. 4D). The chemical shift differences between the two bound Rtt103-CIDs are very small, and the most significant changes occur in close proximity to the region where Ser2 and Thr4 are located in our structure of pSer2 bound to Rtt103 (Fig. 4E) (20).

This result is in good agreement with the previous finding by us and others that the structures of CIDs bound to CTD peptides generally overlap quite well. Specifically, the Rtt103-CID bound to pSer2-CTD (20) overlaps well with the other CIDs bound to CTD peptides with different phosphorylated states, with a 5.5-Å C α rmsd to Nrd1-CID bound to pSer5 (48), 5.2-Å C α rms deviation to the SCAF8-CID bound to pSer2 and pSer5 (49), 5.5-Å C α rms deviation to the SCAF8-CID bound to pSer2 and pSer7 (49), 3.9-Å C α rms deviation to Pcf11-CID bound to pSer2 (25), and 1.1-Å C α rmsd to the RPRD1A-CID bound to pSer7 (50) over the entire length of the polypeptide chain (*SI Appendix*, Fig. S8). Whereas CID is a widely conserved protein fold, it is important to note that other proteins can interact with the CTD using different folds, including Pin1 (WW domain) (51), Tdrd3 (Tudor domain) (52), and Ssu72 (PTP domain) (26–28).

We measured binding affinities for each peptide by fitting the chemical shift changes versus the ratio of added CTD peptide to CID protein for every resonance exhibiting significant changes upon peptide addition (Fig. 4*F*). pThr4_{ab}-CTD binds to Rtt103 CID fivefold weaker than pSer2_{ab}-CTD, but still 25-fold stronger than



Fig. 4. The CID domain of Rtt103 binds analogously to pThr4_{ab}-CTD and pSer2_{ab}-CTD peptides. (A) Schematic of diheptad CTD peptides used in NMR titrations. Each CTD repeat (arrow) is labeled as repeat "a" or repeat "b." Sequences of the peptides are shown on the *Top*. Phosphothreonines and phosphoserines are indicated by orange and green circles, respectively. (*B*) Overlay of ¹H-¹⁵N HSQC spectra of free Rtt103-CID (black) and of the same domain in complex with pThr4_{ab}-CTD (orange) and pSer2_{ab}-CTD (green). (*C*) Chemical shift perturbations (CSPs) of Rtt103-CID with diheptad peptides phosphorylated at Thr4 (orange) or Ser2 (green). Very similar trends are observed for both peptides, with generally smaller overall amplitude changes for pThr4_{ab}-CTD (*Top*). (*Bottom*) Chemical shift difference for Rtt103-CID in complex with the two peptides. Three residues V109, 1112, and K114 (red) have the largest chemical shift differences (>0.08 ppm) in the two complexes. (*D*) The secondary structure is unchanged for Rtt103-CID in complex with pThr4_{ab}-CTD. Chemical shift deviations from random coil shifts for ¹³Cα (*Top*), ¹³Cβ (*Middle*), and ¹³C' (*Bottom*) versus residue number are shown. (*E*) Model of Rtt103-CID in complex with pThr4_{ab}-CTD, generated from the existing structure of Rtt103-pSer2_{ab}-CTD, by substituting pSer2 with Ser2 and Thr4 with pThr4 and performing energy minimization within PyMOL. The three amino acids experiencing the largest changes between the two phosphorylated peptides, V109, 1112, and K114 (red), are located in the immediate proximity of the sites of phosphorylation (Ser2 or Thr4). (*F*) Affinities of Rtt103-CID for diheptad repeat CTD peptides, as measured by NMR titrations. Values are averaged over multiple amino acids (>10) used for extracting independent K_d values by fitting of chemical shift changes vs. the ratio of peptide to Rtt103. Data are represented as mean ± SD.

the unphosphorylated CTD peptide. Recent structural and mutational studies suggest that the functional CTD unit is not defined by the YSPTSPS heptad motif and may well extend to a longer sequence (53–55). Therefore, we examined binding of Rtt103-CID to longer CTD peptides (*SI Appendix*, Figs. S6 and S7). Consistent with the binding to two-repeat peptides, similar structural interface was observed between Rtt103 CID and a four-repeat CTD peptide bearing pThr4 residues at the internal two-repeats (*SI Appendix*,

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Figs. S6 and S7). These results provide a framework for understanding how two different phosphoisoforms interact with a common interaction surface of Rtt103.

Structural Basis for pThr4-Rtt103 Interaction. The solution-state NMR ensemble of structures of Rtt103 revealed a hydrogen bond between Arg108 of Rtt103 and the phosphate on pSer2 of the CTD (20). The data reveal that the Arg108 side chain can adopt many conformations when bound to the pSer2-CTD peptide (SI Appendix, Fig. S5D). To examine whether pThr4 could be accommodated at the same binding interface of Rtt103, we modeled the CTD with this mark on to our NMR structure and performed energy minimization in PyMOL (56) (Fig. 5A). The triad hydrogen bond/salt bridge interaction involving pSer2-Thr4-Arg108 in our previous structure of Rtt103 bound to the pSer2 peptide (20) can be replaced by a new triad Ser2-pThr4-Arg108 in pThr4_{ab}-CTD bound complex (Fig. 5 A and B) without altering the interface significantly, a result consistent with the conservative pattern of chemical shift changes (Fig. 4 B and C). Further, these modeling results are consistent with our biochemical experiments that show strong binding between Rtt103 and pThr4 marks (Fig. 3D).

To validate this model, we purified a mutant variant of Rtt103 in which the positively charged Arg108 was substituted with a neutral Asn (R108N). Previous studies confirm that this mutant is deficient in binding to pSer2 peptides (20), and we hypothesized that it would also be incapable of binding to pThr4. Unphosphorylated or Thr4-phosphorylated GST-CTD bearing 14 repeats was bound to glutathione beads and incubated with WT Rtt103 or the R108N mutant. GST-CTD was used to definitively show that Rtt103 directly binds to pThr4-CTD, because it was purified in bacteria that lack eukaryotic CTD-binding proteins. Consistent with the model, Rtt103 (WT) bound significantly better to pThr4 than to the unmodified CTD. No pThr4-dependent increase in binding was observed with the mutant Rtt103, confirming that R108 plays a crucial role in recognizing both pThr4 and pSer2 marks (Fig. 5C). Whereas we did not observe binding of Rtt103 to beads, both WT and mutant Rtt103 display low levels of binding to unphosphorylated CTD. This low level binding of Rtt103 to CTD is not observed with human Pol II (Fig. 3D). However, immunoprecipitated human Pol II is likely to co-enrich factors that bind unmodified CTD and prevent Rtt103 from interacting with unmodified CTD. On the other hand, the recombinant GST-CTD fusion does not contain eukaryotic CTD binding proteins and is therefore accessible for low levels of Rtt103 association.

Discussion

The CTD undergoes sequential waves of posttranslational modifications during the transcription cycle. These modifications facilitate engagement of cellular machines that process nascent transcripts, place posttranslational marks on histones, and facilitate each stage of the transcription cycle. Unlike other ubiquitous phosphorylation marks on the CTD, pThr4 plays a critical role in regulating transcription at specific gene classes. Convergence of results from independent experimental approaches have revealed (i) two classes of noncoding RNAs that rely on either pThr4 or pSer2 marks to recruit the transcriptional termination machinery, (ii) an interactome that includes protein complexes not known to rely on Thr4 to associate with the CTD, and (iii) the structural basis of the interaction between pThr4 and Rtt103. That Rtt103 binds analogously to pThr4-CTD and pSer2-CTD suggests the remarkable economy of design where two different CTD marks, which are enriched on different sets of genes, are recognized by the cellular machinery that affects a common function at both sets of genes.

More specifically, we find that substitution of Thr4 with Ala in all heptad repeats of the CTD results in defective transcription termination at a subset of snoRNAs in vivo (Figs. 1*C* and 2*A* and *SI Appendix*, Table S1). The genome-wide transcriptome data from the T4A strain reveal two classes of snoRNA genes with termi-



Fig. 5. Model of Rtt103 in complex with pThr4-CTD. (A) Two orthogonal views of the model of Rtt103 in complex with pThr4-CTD and (*B*) representative solution state structures of Rtt103 in complex with pSer2-CTD are shown. All structural figures were generated with PyMOL (56). (C) Biochemical validation of the model is shown. Purified Rtt103 (WT) or the mutant Rtt103 (R108N) was incubated with a bead only control, or beads bound to GST-CTD that were either in their unphosphorylated state (Thr4-CTD) or phosphorylated state (pThr4-CTD). Bar graph shows fold change in binding to the phosphorylated CTD vs. unphosphorylated (*P < 0.01).

nation at one class dependent primarily on pThr4 and termination at the other class primarily dependent on pSer2 marks. We hypothesize that a yet-to-be-identified yeast kinase may selectively place pThr4 marks on the CTD at this subset of snoRNAs. Remarkably, ChIP studies as well as single transcript analysis by Northern blots, show that in the absence of the pThr4 mark, Pol II transcribes through the *SNR33* termination site and continues to transcribe the entire downstream protein-coding gene, *YCR015C*, using the accruing pSer2 marks to terminate synthesis of the chimeric *SNR33-YCR015C* transcript (Fig. 1*C*).

Previous studies defined a role for pSer5 in recruiting the Nrd1-Nab3-Sen1 complex, which facilitates termination of many noncoding RNA transcripts (36, 57–59). This finding prompted us to ask whether the T4A substitution inherently prevents other CTD residues from being phosphorylated. Western blots of WT, S2A/ WT, and T4A cells confirm that pSer5 levels (H14 antibody) were equally abundant among all strains. Further, we only observed a partial reduction in pSer2 in the T4A strain (*SI Appendix*, Fig. S9). This partial reduction in pSer2 levels in the T4A strain may contribute to reduction in binding of known pSer2-CTD "readers" such as Set2 (60–63) and Paf1C (64, 65).

The genomic and proteomic data indicate that a balance between pThr4 and pSer2 levels leads to engagement of common components of the termination machinery at different genes. Rtt103 is known to bind pSer2-marked CTD and facilitate termination at protein-coding genes (19, 20); however, our results suggest that Rtt103 may also participate in termination at noncoding snoRNA genes. A minor reduction in pSer2 in the T4A strain (SI Appendix, Fig. S9) may explain the reduction in Rtt103 occupancy at snoR-NAs; there remains sufficient signal to efficiently terminate at genes that are not dependent on the Thr4 side chain (SI Appendix, Fig. S4B). Whereas the precise role of Rtt103 in termination is being actively investigated by several groups, the ability of termination factors, such as Pcf11, that function at both protein-coding and -noncoding genes has been well documented. Interestingly, NMR studies had previously shown that Rtt103 interacts with pSer2 marks and cooperatively engages Pcf11 at protein-coding genes (20). However, a strain harboring pcf11-13, an allele that disrupts interaction with the CTD, displays defective termination at a set of snoRNA genes (38). These pcf11-13-sensitive snoRNAs coincide with snoRNAs that display termination defects in our T4A strain (P = 0.011) (Fig. 2A and SI Appendix, Table S1). This functional overlap is consistent with our quantitative proteomics analysis that shows reduced association of Pcf11 with T4A-bearing Pol II (Fig. 3). It is interesting to note that structural studies of Pcf11 with pSer2 peptides do not show any direct interaction with this mark but may rely on structure of CTD stabilized by this modification (25, 66, 67). Whereas pThr4 may stabilize a similar conformation in the CTD repeats, it remains to be determined whether such repeat structure or cooperative assembly with direct pThr4 readers such as Rtt103 engages Pcf11 at noncoding snoRNA genes.

In addition to termination factors such as Pcf11 and Rat1 exosome, we also observe reduced association of all subunits of the PAF1 complex with T4A bearing Pol II (Fig. 3B). Consistent with the role of the PAF1 complex in snoRNA termination (35), the read-through defects in T4A strain overlapped remarkably well with those observed in *paf1* Δ cells (P = 0.00014) (SI Appendix, Table S1). In the context of the chromatin landscape, PAF1c plays a key role in snoRNA termination by recruiting the machinery that modifies histone H3 (35). In agreement with this finding, significantly lower levels of histone H3 modifications were observed across snoRNA genes that display termination defects in T4A strains (GSE72802) (SI Appendix, Fig. S4C). Further, a genetic link between Thr4 and chromatin remodeling through the histone variant Htz1 has been reported (15). Consistent with a termination defect, Htz1, a histone H2A variant that yields more labile nucleosomes (68, 69), was highly enriched downstream of snoRNAs that display read-through in T4A (70) (SI Appendix, Fig. S4D).

The exquisite dependence of Rtt103 on pThr4 led us to investigate the molecular basis for the ability of this factor to engage two distinct phosphoisoforms of the CTD. As noted above, pSer2 and pThr4 marks are reciprocally enriched at the two classes of snoRNA genes. Moreover, pSer2 plays a key role in engaging the transcription termination machinery at protein-coding genes. NMR-based structural models of the CTD interacting domain of Rtt103 indicate that the same surface of the protein interacts with the both pSer2- and pThr4-modified CTD peptides. A highly conserved arginine residue on Rtt103 can readily engage either phospho-CTD mark with minimal perturbation of the remaining interface. This structurally analogous mode of engagement contrasts with those observed for Ssu72 binding to pSer5- and pSer7modified CTD peptides. However, the preference for one modification over the other is also observed in the interactions of Rtt103 with pSer2 or pThr4 bearing CTD peptides. The NMR titration data indicate that Rtt103 prefers pSer2 over pThr4 by fivefold and lead us to hypothesize that Rtt103 might have a more direct role at pSer2-enriched genes, whereas it may function more cooperatively with other components of the transcriptional machinery at pThr4-enriched genes.

Taken together, the model that emerges from our results posits that pThr4 and pSer2 play overlapping roles in engaging the transcriptional termination machinery at distinct sets of genes. Unlike the near ubiquitous role for pSer2 in transcription termination, phosphorylation of Thr4 provides a ready means to selectively engage the termination machinery at specific genes. This model is consistent with variable levels of pThr4 marks observed at different gene classes and the differing sets of genes that display the dependence on pThr4 in response to different environmental and physiological cues.

Materials and Methods

Mutant CTD Construct and Plasmid Shuffle. Mutant Rpb1 CTDs were constructed and tested for viability in vivo as previously described with some modifications (71). Briefly, the CTD repeats were constructed by annealing and ligating 5'-phosphorylated oligonucleotides containing WT or mutant codons at position 4 (*SI Appendix*, Table S3). WT and mutant Rpb1 CTDs were cut with Aval and ligated into similarly cut pSB0, which was transformed into DH5 α and screened. pSB0 vectors were cut with KpnI and SnaBI and ligated WT Rpb1 gene. Transformed into Z26, which contains a URA3-linked WT Rpb1 gene. Transformants were plated on synthetic complete (SC) media lacking uracil and leucine. Single colonies were grown overnight in YPD, and plated on SC –Leu +5-FOA (1 mg/mL) (Toronto Research Chemicals) to counterselect against any colonies maintaining the WT Rpb1.

RNA Expression Analysis. Cell collection, RNA preparation, cDNA synthesis and labeling, array hybridization, and normalization were as previously described (72), using cyanine dyes (Flownamics) and SuperScript III (Life Technologies). Samples were hybridized to whole-genome tiled DNA microarrays (Roche NimbleGen). Biological triplicates were averaged, and significant read-through was quantified as the fold change (mutant/WT) in the region 200 bp downstream of the 3' end of processed snoRNAs as previously described (35). All expression data are deposited in the Gene Expression Omnibus (accession no. GSE95419). Additional details on Fisher's exact test used in Fig. 2D are described in SI Appendix, SI Materials and Methods.

Northern Blot Analysis. RNA was extracted from 25-mL cultures at OD₆₀₀ 0.5 using hot phenol. A total of 20 μ g of RNA was run on 1× MOPS [(3-(*N*-morpholino)propanesulfonic acid], 6% formaldehyde, and 1.5% agarose gel and transferred to Hybond-N membrane. ³²P-UTP was incorporated into probes (*SI Appendix*, Table S3) complementary to *SNR33*, *SNR40*, or *SCR1* and was hybridized overnight to the blots. Blots were exposed to a phosphorimaging screen and scanned on a Typhoon FLA9000 (GE Healthcare). Additional details are in *SI Appendix*, *SI Materials and Methods*.

Chromatin Immunoprecipitation. ChIP was done as previously described with minor modifications (9, 31). Chromatin was sonicated on a Misonix 4000 sonicator. Rpb3 (NeoClone, W0012, currently BioLegend), pThr4 (1G7), or the FLAG-tag on Rtt103 (M2, Sigma) was immunoprecipitated. Samples were labeled, hybridized, and processed as previously described (9, 31). All ChIP data are deposited in the Gene Expression Omnibus (accession no. GSE95419). Additional details are in *SI Appendix, SI Materials and Methods*.

Coimmunoprecipitation and Label-Free Quantitative Proteomics. WT, T4A, T4E (unshuffled), and Z26 were grown to OD₆₀₀ ~1.0 in 2 L of selective media. Cells were harvested and lysed via milling with a Retsch Mixer Mill MM 400. Lysed yeast was added to 1× NET-seq buffer (73) with protease and phosphatase inhibitors and 2,000 units DNase I (Sigma). Debris was pelleted, and Pol II was immunoprecipitated with α -HA Dynabeads (Life Technologies). Following washes, samples were eluted three times with 2 mg/mL HA peptide in TBS. Fractions were combined, and proteins were precipitated with TCA. Triplicate samples were analyzed by label-free quantitative proteomics analysis as previously described (42). Significant proteins were identified using QSpec (74). Raw data are available at the ProteomeXchange (accession no. PXD005967). Additional details are described in *SI Appendix, SI Materials and Methods*.

Validation of Binding to pThr4. Endogenous Pol II from HeLa whole-cell extract was immunoprecipitated with the antibody 1C7 (13) to enrich hypophosphorylated CTD (Pol IIA). Pol IIA was treated with PLK3 for 60 min at 28 °C in kinase buffer A [20 mM Tris·HCl (pH 7.4), 20 mM NaCl, 10 mM MgCl₂, 1 μ M

DTT, and 2 μ M ATP] to selectively phosphorylate Thr4. TAP-tagged proteins were expressed in yeast (*SI Appendix, SI Materials and Methods*) and incubated with phosphorylated CTD substrate for 60 min at 30 °C. Beads were washed, and proteins were eluted with Laemmli sample buffer before Western blot analysis using antibodies targeting CTD modifications, Pol II, or the TAP-tag.

NMR Experiments. The CID domain of Rtt103 (Rtt103-CID) used for NMR titration and chemical shift assignments was prepared as described previously (20) and ¹⁵N or ¹⁵N, ¹³C labeled. The diheptad peptides were purchased from AnaSpec or VCPBio Lab. All of the NMR experiments were recorded at 25 °C on a Bruker Avance 600 MHz spectrometer equipped with cry-ocooled ¹H/¹³C/¹⁵N triple-resonance probes and pulse-field gradients. NMR data were processed with NMRPipe (75) and analyzed within CcpNmr (76). The complex of Rtt103-CID with diheptad peptides was formed by titrating the peptide into the CID and monitoring the change of ¹⁵N-edited HSQC spectra until saturation was observed. The chemical shifts of backbone and side-chain resonances were assigned using standard triple resonance NMR experiments including 3D HNCO, HNCACB, and CBCA(CO)NH (77). The composite chemical shift changes for each HN

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cross-peak in ¹⁵N-edited HSQC are quantified using empirical equation $\Delta \delta = \sqrt{\Delta \delta H^2 + (0.15 \times \Delta \delta N)^2}$, where $\Delta \delta H$ and $\Delta \delta N$ are the chemical shift changes for ¹H and ¹⁵N, respectively. For binding affinities, a series of ¹⁵N-edited HSQC spectra were collected with increasing molar ratio of peptide to protein ranging from 0:1 to 3:1. The K_{dS} were calculated using the module "follow shift changes" within CcpNmr by fitting the fractional chemical shift changes against the ratios of ligand to protein.

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