A genetic screen for suppressors of hyper-repression of the fission yeast PHO regulon by Pol2 CTD mutation T4A implicates inositol 1-pyrophosphates as agonists of precocious IncRNA transcription termination

Angad Garg, Stewart Shuman, and Beate Schwer

1Molecular Biology Program, Sloan-Kettering Institute, New York, NY 10065, USA and 2Department of Microbiology and Immunology, Weill Cornell Medical College, New York, NY 10065, USA

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ABSTRACT

Fission yeast phosphate homeostasis genes are repressed in phosphate-rich medium by transcription of upstream IncRNAs that interferes with activation of the flanking mRNA promoters. IncRNA control of PHO gene expression is influenced by the Thr4 phospho-site in the RNA polymerase II CTD and the 3′ processing/termination factors CPF and Rhn1, mutations of which result in hyper-repression of the PHO regulon. Here, we performed a forward genetic screen for mutations that de-repress Pho1 acid phosphatase expression in CTD-T4A cells. Sequencing of 18 independent STF (Suppressor of Threonine Four) isolates revealed, in every case, a mutation in the C-terminal pyrophosphatase domain of Asp1, a bifunctional inositol pyrophosphate (IPP) kinase/pyrophosphatase that interconverts 5′-IP7 and 1,5-IP8. Focused characterization of two STF strains identified 51 coding genes coordinately upregulated vis-à-vis the parental T4A strain, including all three PHO regulon genes (pho1, pho84, tgp1). Whereas these STF alleles—asp1-386(Stop) and asp1-493(Stop)—were lethal in a wild-type CTD background, they were viable in combination with mutations in CPF and Rhn1, in which context Pho1 was also de-repressed. Our findings implicate Asp1 pyrophosphatase in constraining 1,5-IP8 or 1-IP7 synthesis by Asp1 kinase, without which 1-IPPs can accumulate to toxic levels that elicit precocious termination by CPF/Rhn1.

INTRODUCTION

The carboxyl-terminal domain (CTD) of the Rpb1 subunit of RNA polymerase II (Pol2), comprising tandemly repeated heptapeptides of consensus sequence Y1S2P3T4S5P6S7, serves as a scaffold to recruit proteins that regulate transcription initiation, elongation and termination, adjust chromatin structure, and catalyze or regulate mRNA capping, splicing and polyadenylation. The primary structure of the CTD, which is dynamically sculpted by serine, threonine, and tyrosine phosphorylation and by cis–trans proline isomerization, conveys information about the status of the transcription machinery—a CTD code—that is ‘read’ by CTD-interacting proteins and RNA processing assemblies (1–5).

Our aims have been to decipher the informational rules for the CTD code by genetically manipulating the composition and structure of the fission yeast Pol2 CTD; to understand how CTD coding ‘letters’ are assembled into ‘words’ (i.e. a vocabulary); and to elucidate how CTD coding cues govern specific cellular gene expression programs (6–10). The findings that fission yeast is viable when the CTD Tyr1, Ser2, Thr4 or Ser7 residues are uniformly replaced by a non-phosphorylatable side chain accords with transcriptome analyses showing that only a small fraction of fission yeast mRNAs is dysregulated by CTD phospho-site mutations. We hypothesized that the effects of mutating these phospho-sites are genetically buffered by other cellular factors that are functionally redundant to the phospho-mark or the side-chain hydroxyl. By identifying such functional redundancies (manifest as synthetic lethality), and gauging their specificity for a particular phospho-site mutation, we showed that the effects of mutating the inessential Tyr1, Ser2, and Thr4 CTD phospho-sites on fission yeast cell growth are buffered by subunits of the fission yeast cleavage and polyadenylation factor (CPF) complex (10).
Key insights emerged from studies of fission yeast phosphate homeostasis, a transcriptional response to phosphate availability that is governed by the CTD code and the 3′ processing/termination machinery (8–12). The *Schizosaccharomyces pombe* phosphate (PHO) regulon comprises three genes that specify, respectively, a cell surface acid phosphatase Pho1, an inorganic phosphate transporter Pho84, and a glycerophosphate transporter Tgp1 (13). Expression of the PHO genes *pho1*, *pho84*, and *tgp1* is actively repressed during growth in phosphate-rich medium by the transcription in cis of a long noncoding (lnc) RNA from the respective 5′ flanking genes *ptr*, *prt2*, and *nc-tgpl* (11–12,14–17). A *CTD*-S7A allele that prevents installation of the Ser7-PO4 mark in all heptads de-represses the PHO genes in phosphate-replete cells (8–10). By contrast, a *CTD-T4A* allele that precludes inscription of the Thr4-PO4 mark in all heptads de-represses the PHO genes under phosphate-rich conditions (8–10). Our model for the repressive arm of phosphate homeostasis is that: (i) transcription of the upstream lncRNA interferes with expression of the downstream mRNA genes by displacing the activating transcription factor Pho7 from its binding site(s) in the mRNA promoters that overlap the lncRNA transcription units (13,18–20); (ii) loss of the Ser7-PO4 mark leads to precocious termination of lncRNA transcription prior to the mRNA promoter; and (iii) loss of the Thr4-PO4 mark reduces termination and hence increased transcription across the mRNA promoter (10). This model is supported by RNA analyses, transcriptomics and epistatic effects of CTD mutations with mutations of CPB subunits and transcription termination factor Rhn1 (10).

A model for the transcriptional arm of the fission yeast phosphate starvation response is that phosphate deprivation triggers a signaling pathway that ultimately results in shut-off of transcription of the *ptr*, *prt2*, and *nc-tgpl* lncRNAs that—under phosphate replete conditions—interfere in cis with expression of the downstream *pho1*, *pho84*, and *tgp1* phosphate acquisition genes (11). Yet, the mechanism of phosphate sensing in fission yeast is obscure and it is not known how a starvation signal is transmitted to impact lncRNA transcription initiation by Pol2. Our work has defined a distinctive bipartite promoter—consisting of a HomolD box and a TATA box—that is essential for transcription of the three PHO-regulatory lncRNAs (11–12,14). The transcription factors that recognize the lncRNA promoters (particularly the HomolD box) and drive lncRNA synthesis are uncharted. The HomolD box [5′-CAGTCAC(A/G)] was identified initially as a Pol2 promoter signal in fission yeast genes encoding ribosomal proteins (21–23). Mutations in the lncRNA or in protein factors that diminish lncRNA 3′ processing/termination can delay, but do not eliminate, the upregulation of PHO gene transcription in response to phosphate starvation (9,11–12).

A notable outcome of our epistasis analyses, wherein we made pairwise combinations of the hyper-repressive *CTD*-T4A allele with various mutations that de-repress the PHO regulon in phosphate-replete cells, was that T4A repression generally ‘won out’ over de-repression (9,10). This signifies that the de-repressive mutations, which are posited to elicit precocious lncRNA termination, depend on Thr4-PO4 (or the Thr4 hydroxyl) to achieve their effect. Our aim in the present study was to query how the T4A mutant affects 3′ processing/termination, by performing a genetic screen in *CTD-T4A* cells for extragenic suppressors of the *pho1* hyper-repressive phenotype. We felt this was an auspicious scenario for suppressor genetics, because it is virtually impossible to obtain a revertant of the *rpb1-CTD-T4A* allele that has 27 different missense codon mutations that change Thr to Ala, and because the known mutations that de-repress *pho1* in a wild-type CTD background ought not to score as such in *T4A* cells. We envisioned two potential classes of T4A suppressor mutations: (i) those that reduce or shut off transcription of the *prt* lncRNA and thereby relieve *ptr* interference with the *pho1* promoter; and (ii) those that enhance the 3′ processing/termination efficiency of the T4A mutant Pol2 transcription complex engaged in *prt* lncRNA synthesis. With respect to the latter category, we can imagine several ways in which *T4A* might be suppressed. For example, by mutations in Pol2 itself (e.g. mutations that affect its elongation properties or CTD-independent interactions with elongation, processing and termination factors); by mutations in Pol2 elongation factors (that affect the temporal window to elicit termination of lncRNA synthesis); by hypomorphic mutations in essential 3′ processing/termination factors for which we presently have no genetic handles to establish their connections to the CTD code; by mutations that influence rNTP pools and thereby affect Pol2 elongation indirectly; or by mutations that affect the levels of signaling molecules that regulate 3′ processing/termination.

We describe here the isolation and characterization of a collection of *STF* (Suppressor of Threonine Four) mutants. Sequencing of 18 independent *STF* isolates revealed, in every case, missense or nonsense mutations in the C-terminal pyrophosphatase domain of Asp1—a bifunctional inositol pyrophosphatase (IPP) kinase/pyrophosphatase enzyme (24–28)—as responsible for the *STF* phenotype. Epistasis analysis vis-à-vis CPB and Rhn1 indicates that *STF* mutations promote 3′ processing/termination. Transcriptional profiling uncovered an ensemble of 51 coding genes that were cooperatively up-regulated in two different *STF* nonsense mutant strains; this set includes all three genes of the PHO regulon. Thus, our forward genetic screen fortifies the recent proposal (29) that fission yeast phosphate homeostasis is subject to metabolite control by inositol pyrophosphates, exerted via the 3′ processing/termination machinery and the Pol2 CTD.

**MATERIALS AND METHODS**

**Mutagenesis of rpb1-CTD-T4A cells and screening for increased pho1 expression**

Ethyl methanesulfonate (EMS) mutagenesis was carried out as described (30). In brief, *rpb1-CTD-T4A* cells were grown in minimal medium (PMG) at 30°C to an *A*₆₀₀ of 0.25. Cells were harvested by centrifugation and resuspended at ~1 × 10⁸ cells/ml in PMG medium. To one aliquot (2 ml), we added EMS (Sigma) to a final concentration of 2%, while a second aliquot was left untreated (to determine survival rate). The suspensions were placed on a nutator and incubated for 3.5 h at room temperature. Cells were washed thrice with NaCl (150 mM), diluted, and plated to YES
agar. After 5 days of incubation at 30°C, plates were overlaid with 1% agarose containing 0.015% α-naphthyl phosphate, 0.15% Fast Blue B Salt stain, and 0.1 M sodium acetate (pH 4.2) to assay acid phosphatase activity. Colonies that turned red were extracted from near confluent lawns of mutagenized cells and colony-purified by streaking or re-plating at low cell densities. Independently isolated candidate suppressor strains were back-crossed to rpb1-CTD-T4A cells of the opposite mating type, subjected to random spore analysis, and identified as red colonies using the overlay assay. After back-crossing individual suppressor strains for a second time, we measured acid phosphatase activity of cells grown in liquid culture (YES medium) at 30°C, as follows. Exponentially growing cultures were harvested, washed, and resuspended in water. Reaction mixtures (200 μl) containing 10 mM p-nitrophenylphosphate, 100 mM sodium acetate (pH 4.2), and serial dilution of cells (ranging from 0.01 to 0.1 A{sub 600} units) were incubated at 30°C for 5 min. To stop the reaction, 1 ml of 1 M sodium carbonate was added, the cells were removed by centrifugation, and the absorbance of the supernatant at 410 nm was measured. Acid phosphatase activity is expressed as the ratio of A{sub 410} (p-nitrophenol production) to A{sub 600} (cells). The data are averages (± SEM) of measurements from at least three independent cultures.

**Spot tests of fission yeast growth**

Cultures of *S. pombe* strains were grown in liquid YES (yeast extract with supplement) medium until A{sub 600} reached 0.5–0.8. The cultures were adjusted to an A{sub 600} of 0.1 and aliquots (3 μl) of serial 5-fold dilutions were spotted to YES agar. The plates were photographed after incubation for 2 days at 34°C, 2.5 days at 30°C and 37°C, 4 days at 25°C, and 6 days at 20°C.

**Whole-genome sequencing**

After PicoGreen quantification and quality control by Agilent BioAnalyzer, 500 ng aliquots of genomic DNA were sheared using a LE220-plus Focused-ultrasonicator (Covaris catalog # 500569) and sequencing libraries were prepared using the KAPA Hyper Prep Kit (Kapa Biosystems catalog # 500569) and sequencing libraries were subjected to size selection by mixture with 0.5 volume of aMPure XP beads (Beckman Coulter catalog # A63882) after post-ligation cleanup. Libraries were not amplified by polymerase chain reaction (PCR) and were pooled equivalent for sequencing. Samples were run on a NovaSeq 6000 in a 150/150 bp paired end run using the NovaSeq 6000 SBS v1 Kit and an S1 flow cell (Illumina). The average number of read pairs per sample was 10 million.

**Mapping STF mutations**

The FASTA file for the *S. pombe* genome was accessed from Pombase and modified by replacing the wild-type rpb1 locus with the rpb1-CTD-T4A allele sequence. To preserve chromosomal coordinates of all other genes, the natMX antibiotic-resistance gene and TPII terminator sequences placed 3′ of the rpb1-CTD-T4A ORF during strain construction were annotated separately in the genome FASTA file. The genome index was built from this modified FASTA file using Bowtie2 (31). The whole-genome sequencing data from the parental and STF mutant rpb1-CTD-T4A strains were aligned to the genome using Bowtie2 (31). The resulting SAM files were converted to BAM files using Samtools (32). Variants were identified by BCFTools (33) using the criteria of adjusted mapping quality = 40, minimum base quality = 20, and disabled probabilistic realignment for the computation of base alignment quality for considering variations or insertion-deletion events. The multi-allelic caller protocol was used for variant calling in BCFTools. Variants were annotated using SnvEff, with its in-built genome version for *S. pombe* (34). Variants were further filtered by removing all variations with an average mapping quality ≤ 25 (Phred scale). All variants present in the parental strain were excluded as non-causal mutations.

**Transcriptome profiling by RNA-seq**

RNA was isolated from *S. pombe* STF-6 and STF-9 cells and from parental CTD-T4A cells that were grown in liquid YES medium at 30°C to an A{sub 600} of 0.5–0.6. Cells were harvested by centrifugation and total RNA was extracted via the hot phenol method. The integrity of total RNA was gauged with an Agilent Technologies 2100 Bioanalyzer. The Illumina TruSeq stranded mRNA sample preparation kit was used to purify poly(A)+ RNA from 500 ng of total RNA and to carry out the subsequent steps of poly(A)^+ RNA fragmentation, strand-specific cDNA synthesis, indexing, and amplification. Indexed libraries were normalized and pooled for paired-end sequencing performed by using an Illumina NovaSeq 6000-S1 flow cell. FASTQ files bearing paired-end reads of length 51 bases were mapped to the *S. pombe* genome (Pombase) using HISAT2-2.1.0 with default parameters (35). The resulting SAM files were converted to BAM files using Samtools (32). Count files for individual replicates were generated with HTSeq-0.10.0 (36) using exon annotations from Pombase (GGF annotations, genome-version ASM294v2; source ‘ensembl’). RPKM analysis and pairwise correlations (Supplementary Figures S1 and S2) were performed as described previously (8). Differential gene expression and fold change analysis was performed in DESeq2 (37). Cut-off for further evaluation was set for genes that had an adjusted p-value (Benjamini–Hochberg corrected) of <0.05 and were up or down by at least 2-fold in STF-6 or STF-9 in comparison to CTD-T4A. Genes were further filtered on the following criteria: (i) ≥ 2-fold up and the average normalized read count for the mutant strain was ≥ 100; and (ii) ≥ 2-fold down and the average normalized read count for the parent strain was ≥ 100.

**RESULTS**

The STF screen

*rpb1-CTD-T4A* cells were mutagenized by treatment with 2% EMS for 3.5 h at 22°C to achieve a survival rate of ~40%. The cells were washed, plated on phosphate-replete YES agar medium, and incubated for 5 days at 30°C to allow formation of single colonies from individual mutagenized cells. To gauge acid phosphatase activity, the
plates were overlaid with 1% agarose containing 0.015% 6-naphthyl phosphate (a Pho1 substrate), 0.15% Fast Blue B Salt stain, and 0.1 M sodium acetate (pH 4.2). The cell surface Pho1 acid phosphatase causes formation of a red pigment and relative red color intensity of the colonies provides a semi-quantitative assay of Pho1 activity (8,38). T4A cells appear pale after this procedure. From an initial pool of about 390,000 EMS survivors, we picked 50 independent red colonies that we deemed candidate STF mutants. These were re-streaked for single colonies, then grown out in YES liquid medium, re-plated, and re-tested for red color. We then selected 36 of the most intensely red staining strains, back-crossed them to a T4A strain of the opposite mating type, and tested populations of post-sporulation haploid back-crossed them to a parental strain and the twelve independent twice back-crossed T4A progeny for Pho1 acid phosphatase activity via the overlay type, and tested populations of post-sporulation haploid back-crossed them to a parental strain and the twelve independent twice back-crossed T4A progeny for Pho1 acid phosphatase activity via the overlay assay. If the STF phenotype results from a mutation in a single gene, then we expect ~50% of the haploid progeny from the back-cross to stain red and ~50% to be pale. An example of such segregation is shown in Figure 1A. Twenty-four of the back-crossed putative single-gene STF mutants were back-crossed for a second time to the original parental T4A strain and 1:1 segregation of red/pale colony color was affirmed. Twelve independent twice back-crossed STF strains and the parental T4A strain were assayed quantitatively for Pho1 activity after growth at 30°C in phosphate-replete liquid medium. The STF strains expressed 23- to 49-fold higher Pho1 activity than the T4A parent (Figure 1B). All twelve of these STF strains grew as well as the T4A parent on YES agar at 30°C (Figure 1C).

Identification of STF mutations by whole genome sequencing

Paired-end Illumina sequencing of unamplified genomic DNA (average read length 150 bases) from the parental T4A strain and the twelve STF strains was performed by the MSKCC genomics core facility to achieve at least 100-fold coverage of each fission yeast genome. The striking finding was that 12/12 STF strains had acquired missense or nonsense mutations in the *asp1*+ gene that encodes a bifunctional kinase/pyrophosphatase enzyme involved in inositol pyrophosphate (IPP) metabolism (24–28) (Supplementary Figure S3). A list of missense, nonsense or frameshift mutations within other coding genes that were detected in the 12 STF strains vis-à-vis the parental T4A strain is shown in Supplementary Figure S3. In cases where the same mutation was found in multiple different STF strain genomes, it is most likely that they were acquired as polymorphisms from the T4A strain of opposite mating type that was employed during the two rounds of back-crossing prior to whole-genome sequencing. This is certainly the case for the *ade6*-(P489L) coding mutation found in 7/12 of the STF strains (Supplementary Figure S3), i.e. this *ade6* allele (known as *ade6-M210*) is an auxotrophic marker used to select diploids after mating.

Three pairs of STF strains had an identical nucleobase mutation in the *asp1*+ open reading frame: STF-6 and STF-10 (G1158A); STF-5 and STF-8 (G1928A); STF-3 and STF-11 (G2588A). Although STF-6 and STF-10 have the same *asp1*+ mutation, STF-10 has two additional changes (in the *dm1*+ and *yta12*+ genes) that are not found in STF-6 or any of the other STF strains (Supplementary Figure S3). Similarly, whereas STF-5 and STF-8 have the same *asp1*+ mutation, STF-8 has a mutation (in *mgl1*+) that is not found in STF-5 or any of the other STF strains (Supplementary Figure S3). And, though STF-3 and STF-11 have the same *asp1*+ mutation, STF-3 has 12 additional coding mutations that are not present in STF-11 or any of the other STF strains (Supplementary Figure S3). We surmise that these STF strains are independent isolates.

The 920-aa Asp1 protein, which was mutated in all of the STF strains, consists of two catalytic domains: an N-terminal IPP kinase module (aa 1–364) that converts 5-IP7 to 1,5-IP8; and a C-terminal IPP pyrophosphatase domain that converts 1,5-IP8 back to 5-IP7 (Figure 2B). Asp1 can also phosphorylate IP6 to yield 1-IP7 and dephosphorylate 1-IP7 back to IP6. Asp1 is inessential for fission yeast growth. The effect of an *asp1Δ* null allele or a kinase-defective *asp1* missense mutant is to eliminate intracellular IP8 and 1-IP7 and to increase the level of 5-IP7 (26,28). Conversely, a pyrophosphatase-defective *asp1* missense mutant increases the intracellular level of IP8, as does expression of a C-terminal truncation allele that retains the kinase domain but lacks the pyrophosphatase domain (26).

Recent studies show that failure to synthesize IP8 and 1-IP7 in Asp1 null or Asp1 kinase-defective strains results in hyper-repression of the *PHO* regulon under phosphate-replete conditions (29). Synthetic lethality of *asp1Δ* and *asp1* kinase-defective alleles with mutations of CPF subunits Ppn1, Swd22, and Ssu72 point to an important role for IP8 (or 1-IP7) in essential 3′ processing/termination events, albeit in a manner genetically redundant to CPF (29).

Pertinent to the present suppressor analysis is the observation that *asp1Δ* and the kinase-defective allele are synthetically lethal with *CTD-T4A* (B. Schwer, unpublished), i.e. the negative effect of T4A on 3′ processing/termination is exacerbated by the lack of IP8.

The salient finding here is that all of the STF mutations map to the C-terminal pyrophosphatase domain of Asp1 (Figure 2A). Asp1 pyrophosphatase is a member of the histidine acid phosphatase superfamily of phosphohydrolases that act via a covalent enzyme-(histidinyl-N)-phosphate intermediate (39). (Pho1 is a member of the same acid phosphatase superfamily.) The signature active site motif—390RHADR400 in the Asp1 pyrophosphatase—is located close to the N-terminal margin of the domain at residue 365. His397 in this motif is the Asp1 catalytic nucleophile. To aid in interpreting the STF mutations, we submitted the pyrophosphatase domain amino acid sequence to the Phyre2 structure modeling server (40), which returned a ‘top hit’ tertiary structure model templated on the 1.7 Å crystal structure of *Yersinia kristensenii* phytase (PDB ID: 4ARV; 39). The model starts at Trp386 and ends at Asp871 and is punctuated by gaps (aa 426–491, 641–659 and 699–738) comprising segments of the Asp1 pyrophosphatase that have no counterparts in *Yersinia* phytase. A stereo view of the Asp1 Phyre2 model is shown in Figure 2C with a phosphate anion in the active site derived from a superposition of the *Yersinia* phytase structure. The model places the His397-Nε nucleophile 3.3 Å from the phosphorus center and allows for a network of phosphate contacts to conserved arginine side chains Arg396, Arg400, and Arg536 (Figure 2C). Alanine mutations of Arg396,
Ala398Val is located within the RHADR400 active site motif and thereby affect its conformation. Thus, we infer that asp1-H686Y and asp1-A398V strains are defective for Asp1 pyrophosphatase activity. To our inspection, the model does not provide insights to any structural defects of the CTD-WT cells is several fold higher than the ‘hyper-repressed’ Pho1 state in CTD-T4A cells). With longer incubation under the agar overlay, tiny puncta of red staining were observed on the plates selective for CTD-WT progeny. Microscopic examination indicated that these puncta corresponded to progeny that had germinated and undergone only a few rounds of cell division. Our inference from these findings is that these two nonsense asp1-STF alleles are lethal in a wild-type rpbl-CTD background. Consistent with this interpretation, we genotyped the asp1 locus of 12 individual CTD-WT isolates from the STF-6 backcross and 12 individual CTD-WT isolates from the STF-9 backcross, and found that all of them were wild-type with respect to the Asp1 pyrophosphatase domain.

To affirm the results of the random spore analysis, we dissected nine individual four-spore asci from the STF-9 backcross and genotyped the viable haploid progeny for the rpbl-linked drug resistance markers and for Pho1 expression by agar overlay. Examples of the tetrad analysis are shown in Figure 3. The tetrads that yielded four viable progeny \( n = 4 \) were found to be parental ditypes (PD) comprising two CTD-T4A dark red progeny and two CTD-WT light red progeny. Tetrads that yielded two viable progeny \( n = 3 \) were non-parental ditypes (NPD) consist-
Figure 2. STF mutations map to the Asp1 IPP pyrophosphatase domain. (A) Whole-genome sequencing of 12 independent twice back-crossed STF strains revealed the indicated coding mutations in the C-terminal IPP pyrophosphatase domain of Asp1. (B) Asp1 is a bifunctional enzyme composed of an N-terminal IP7 kinase domain (gold) and a C-terminal IP8 pyrophosphatase domain (cyan). Asp1 kinase converts IP7 to IP8 and the Asp1 pyrophosphatase reverses this process, as shown. (C) Stereo view of the Phyre2 model of the Asp1 pyrophosphatase domain tertiary structure templated on the crystal structure of Yersinia kristensenii phytase, with a phosphate anion in the active site and surrounding active site side chains.

ing of two CTD-T4A pale progeny. Tetrads yielding three viable progeny \( (n = 2) \) were tetratypes (TT) comprising one CTD-T4A dark red haploid, one CTD-T4A pale haploid, and one CTD-WT light red haploid. Failure to recover the fourth CTD-WT dark red haploid indicated that the asp1-(W493-Stop) mutation, which eliminates the Asp1 pyrophosphatase domain while preserving the Asp1 kinase, is lethal in a wild-type rpb1-CTD background. We also dissected eight individual four-spore asc from the STF-6 backcross. Tetrad analysis revealed three parental ditypes with four viable progeny, three non-parental ditypes with two viable progeny, and two tetratypes with three viable progeny. Thus, the asp1-(W386-Stop) allele is also lethal in the context of a wild-type Pol2 CTD.

Other STF alleles are viable but sick in a wild-type rpb1-CTD background

We then proceeded to mate the other STF CTD-T4A strains with a differentially marked wild-type rpb1-CTD strain and performed both random spore analysis and tetrad dissections as described above. STF-1, STF-2, STF-4, and STF-12 matings to a wild-type rpb1-CTD strain yielded CTD-WT STF colonies at 30°C that stained deep red in the agar overlay assay. Red-staining CTD-WT STF-3, STF-5, and STF-7 haploids were also viable but grew out more slowly than the other strains after sporulation. Sequencing the asp1 pyrophosphatase domain ORF in these CTD-WT haploids affirmed the presence of the STF mutations. The CTD-WT STF strains were spot-tested for growth on YES agar medium. The STF-1, STF-2, STF-3, STF-5, STF-7, and STF-12 mutations conferred a severe cs growth defect at 20 and 25°C, a ts growth defect at 34 and 37°C, and slower growth at 30°C compared to the CTD-WT asp1+ control (Figure 4A). CTD-WT STF-4 cells displayed a cs growth defect, but grew well at 30°C and better than the other STF strains at 25, 34 and 37°C (Figure 4A).

The viable CTD-WT STF strains were grown in phosphate-replete medium at 30°C and assayed for acid phosphatase activity. The STF-1, STF-2, STF-3, STF-4,
CTD-T4A strains with dis2Δ, ctf1Δ, ssu72-C13S, ppn1Δ, swd22Δ, and rhn1Δ strains. A population of random spores was screened for the drug-resistance markers linked to the CTD-T4A locus and the CPF/rhn1 mutant loci. Haploid progeny bearing mutant CPF/rhn1 alleles and were wild-type with respect to the Pol2 CTD were then screened by agar overlay for Pho1 expression, which showed similar numbers of pale and red isolates in every genetic cross. The viable STF-6 CPF/rhn1 and STF-9 CPF/rhn1 double-mutants were spot-tested for growth on YES agar in parallel with the respective CPF/rhn1 single-mutants (Figure 5). Notable findings were that the STF-6 and STF-9 alleles alleviated the cold-sensitive growth defects of ppn1Δ and swd22Δ cells at 20°C and the slow-growth ts phenotype of rhn1Δ cells at 37°C. Also, the rhn1Δ STF-6 and rhn1Δ STF-9 double-mutants displayed a strong cs defect that was not seen in rhn1Δ per se (Figure 5). These genetic suppression data suggest that the lethality of the IPP pyrophosphatase-defective asp1-(W386-Stop) and asp1-(W493-Stop) mutations is exerted via 3′ processing/termination.

The STF CPF/rhn1 double mutants were assayed for Pho1 acid phosphatase activity during growth in phosphate-replete medium, in parallel with the respective CPF/rhn1 single mutants. In every case, the STF alleles elicited a strong de-repression of Pho1 expression, ranging in magnitude from 16- to 62-fold above the hyper-repressed level of Pho1 expression in the corresponding CPF/rhn1 single mutants (Figure 6).

Transcriptome profiling of the STF-6 and STF-9 strains

We performed RNA-seq on poly(A)+ RNA isolated from STF-6 and STF-9 cells and from the parental rpb1-CTD-T4A strain. cDNAs obtained from three biological replicates (using RNA from cells grown to mid-log phase in YES medium at 30°C) were sequenced for each strain. In the datasets, 90–96% of the reads were mapped to unique genomic loci (Supplementary Figure S1). Read densities (RPKM) for individual genes were highly reproducible between biological replicates (Pearson coefficients of 0.97–0.99; Supplementary Figure S2). A cutoff of ±2-fold change in normalized transcript read level and an adjusted P-value of ≤0.05 were the criteria applied to derive an initial list of differentially expressed annotated loci in the STF-6 and STF-9 mutants versus the parental CTD-T4A control. We then focused on differentially expressed genes with average normalized read counts ≥100 in either the STF or parental T4A strains in order to eliminate many (mostly non-coding) transcripts that were expressed at very low levels in vegetative cells. We thereby identified sets of 64 and 63 annotated protein-coding genes that were upregulated by these criteria in STF-6 and STF-9 cells, respectively, 51 of which were coordinately upregulated by ≥2-fold in both STF mutants (P < 6.795e-98) (Figure 7). The most highly upregulated sub-set (≥16-fold increase) includes all three phosphate homeostasis genes: pho1 (up 46- to 54-fold), tgp1 (up 19- to 26-fold), and pho84 (up 16- to 18-fold) (Figure 7). RNA-seq revealed no increase in STF-6 and STF-9 cells of the mRNA encoding Pho7, the transcription factor that drives pho1, pho84, and tgp1 mRNA synthesis. The ecl3

STF-5, STF-7, and STF-12 mutations increased Pho1 expression by 21-, 20-, 23-, 17-, 22-, 25-, and 21-fold, respectively, vis-à-vis the CTD-WT asp1+ control (Figure 4B). Moreover, Pho1 expression in STF-1, STF-2, STF-3, STF-4, STF-5, STF-7 and STF-12 cells was 4-, 4-, 3-, 4-, 3-, and 3-fold higher in the CTD-WT background than in the CTD-T4A background (Figure 4B versus 1B). Thus, Asp1 pyrophosphatase and CTD-Thr4 exert autonomous and opposing effects on Pho1 expression.

Lethality of asp1-(W386-Stop) and asp1-(W493-Stop) is suppressed by CPF and Rhn1 mutations

Fission yeast CPF is a 13-subunit protein assembly responsible for the co-transcriptional 3′ processing of Pol2 transcripts that precedes Pol2 transcription termination (41). Five of the CPF subunits (Ctf1, Ssu72, Dis2, Ppn1, and Swd22) are dispensable for growth. Rhn1 is an inessential CTD-binding Pol2 termination factor that recognizes the Thr4-PO4 CTD mark (42). Loss-of-function mutants of Rhn1 or any of the five inessential CPF subunits result in hyper-repression of pho1 RNA expression and Pho1 acid phosphatase activity in phosphate-replete cells (10). In light of the findings above that the STF-6 asp1-(W386-Stop) and STF-9 asp1-(W493-Stop) alleles were viable in combination with CTD-T4A (which is thought to exert a negative effect on 3′ processing/termination) but lethal in a wild-type rpb1-CTD background, we queried whether their lethality could be suppressed by non-CTD mutations in the 3′ processing/termination machinery. Accordingly, we performed pairwise matings of STF-6 CTD-T4A and STF-9

Figure 3. Tetrad dissection shows that STF-9 is lethal in the context of a wild-type Pol2 CTD. Individual spores (a–d) from five tetrads (1–5) resulting from a cross between asp1-STF-9 CTD-T4A and asp1-WT CTD-WT strains were arrayed on YES agar medium and incubated for 6 days at 30°C. Genotyping the viable progeny assigned the segregation patterns as non-parental ditype (NPD), parental ditype (PD), and tetratype (TT) as indicated.
Figure 4. STF mutants that are viable with a wild-type rpb1-CTD elicit conditional growth defects and lead to stronger pho1 de-repression than in the CTD-T4A background. (A) CTD-WT and CTD-WT STF strains (as specified on the left) were spot tested for growth at the indicated temperatures. (B) The CTD-WT and indicated CTD-WT STF strains were assayed for Pho1 acid phosphatase activity.

Figure 5. Lethality of asp1-(W386-Stop) and asp1-(W493-Stop) is suppressed by CPF and Rhn1 mutations. Serial 5-fold dilutions of the indicated strains were spot tested for growth at the indicated temperatures.

gene, upregulated by 28- to 39-fold, is located on chromosome II, adjacent to and in opposite orientation to the prt2 lncRNA gene of the phosphate-regulated prt2-pho84-prt-pho1 gene cluster (14). Two other genes involved in phosphate acquisition were also upregulated: SPBPA2B2.06c encoding an extracellular 5′ nucleotidase (up 28- to 48-fold) and SPBC1683.01 encoding a transmembrane phosphate transporter (up 3- to 4-fold) (Figure 7). It is suggested that ecl3 and SPBC1683.01, in common with the three PHO genes, might be subject to transcription interference by 5′ flanking lncRNAs (29).

The RNA-seq experiment identified 63 and 65 protein-coding genes that were downregulated by ≥2-fold in STF-6 and STF-9 cells, respectively, 59 of which were coordinately downregulated in both STF mutants (P < 2.152e-125) (Supplementary Figure S4). The set of downregulated transcripts includes those encoding proteins involved in fission yeast iron homeostasis: the siderophore Str1; sulfiredoxin Srx1; ferric reductase Frp1, iron oxidase-permease Fio1; iron permease Fip1; and ferric reductase Frp2. Expression of these genes is normally repressed during growth in rich medium by the iron-sensing DNA-binding GATA-family transcriptional repressor Fep1 (43,44). RNA-seq showed no effect of STF-6 or STF-9 on the level of fep1 mRNA compared to that in T4A cells, suggesting that the observed reduction in iron regulon transcripts is not caused by increased expression of the Fep1 repressor.
Identification of additional STF mutations by focused sequencing of the asp1 locus

In light of the whole-genome sequencing data showing that all of the initial set of 12 STF strains had Asp1 pyrophosphatase mutations, we analyzed a second set of six independent twice-backcrossed STF isolates (alleles 13–18) by focused PCR amplification of the Asp1 pyrophosphatase ORF and sequencing the PCR product. We thereby found that all six had mutations in the Asp1 pyrophosphatase domain, including one nonsense change (Trp642-Stop) and four missense changes—Arg396His, Arg400His (two independent isolates), Glu844Lys, and Gly863Asp (Figure 8B). The Asp1 Gly863Asp lesion in the STF-14 strain was identical to the missense mutations identified by whole genome sequencing in two independent STF mutants in the initial set (STF-3, STF-11). Moreover, the acid phosphatase activity of STF-14 cells (Figure 8A) was similar to that of STF-3 and STF-11 cells (Figure 1B). The other STF strains in this set expressed lower levels of Pho1, albeit still 9- to 10-fold higher than the parental T4A strain (Figure 8A). Reference to the Phyre2 model (Figure 2C) underscores that the sites of two STF mutations—Arg396 and Arg400—are catalytic residues of the pyrophosphatase active site. It is conceivable that single Arg-to-His changes at these positions do not completely eliminate IPP pyrophosphatase activity, insofar as histidine might fulfill some of the interactions made by arginine with the scissile phosphate.

Second-round STF mutants that are viable with a wild-type rpb1-CTD elicit stronger pho1 de-repression than in the CTD-T4A background

The STF-13, STF-15, STF-16, STF-17, and STF-18 strains from the second set of CTD-T4A STF mutants were mated to a wild-type rpb1-CTD strain, the resulting diploids were sporulated, and random haploid progeny were screened for the rpb1-linked drug-resistance markers and for Pho1 expression by agar overlay. In each case, we recovered viable CTD-WT dark red isolates at the frequency expected for random segregation, signifying that the asp1-E844K, asp1-W642Stop, asp1-R396H, and asp1-R400H alleles were not lethal in the context of a wild-type Pol2 CTD. We infer that these mutations are likely hypomorphs with respect to Asp1 pyrophosphatase function. The CTD-WT STF-13 and STF-15 strains grew well on YES agar at all temperatures (Figure 8C). The CTD-WT STF-16 and STF-17 strains grew well at 30–37°C but displayed a slow growth defect at low temperatures, as gauged by colony size (Figure 8C). Testing the CTD-WT STF strains for acid phosphatase activity (Figure 8D) was instructive in the following respects: (i) the STF-13, STF-15, STF-16, and STF-17 mutations increased Pho1 expression by 12-, 9-, 16-, and 16-fold, respectively, vis-à-vis the asp1-WT control; and (ii) Pho1 expression in cells bearing STF-13, STF-15, STF-16, and STF-17 alleles was 8-, 6-, 11-, and 10-fold higher in the CTD-WT background than in the CTD-T4A background (Figure 8D versus A). These results fortify the conclusion that the Asp1 pyrophosphatase and the CTD Thr4 mark exert opposing effects on phosphate homeostasis.

DISCUSSION

The results of the STF screen and ensuing genetic analyses engender two key conclusions: (i) ablating the Asp1 IPP pyrophosphatase domain while leaving the IPP kinase domain intact can be lethal; and (ii) the IPP pyrophosphatase domain requirement for viability in such circumstances is bypassed by the rpb1-CTD-T4A mutation and by loss-of-function mutations in CPF subunits and termination factor Rhn1. The inference we draw is that an important role of the Asp1 pyrophosphatase is to constrain the level of IP8 or 1-IP7 may accumulate to toxic levels that elicit precocious termination by CPF/Rhn1.

We had already proposed that too much IP8 is toxic to fission yeast based on our findings that a viable IPP pyrophosphatase-defective allele—asp1-H397A, constructed by the Fleig laboratory (24)—was lethal in the absence of a second fission yeast IPP pyrophosphatase enzyme Aps1 (29). Aps1 is a member of the Nudix hydro-
Figure 7. Transcription profiling identifies genes upregulated by STF-6 and STF-9 alleles. List of 51 annotated protein-coding genes that were upregulated at least 2-fold in STF-6 T4A and STF-9 T4A cells compared to the parental T4A strain. The log2 fold changes are shown. The 13 genes that were also upregulated in asp1−H397A cells are indicated in bold font.

| gene            | log2 change | protein / function                                      |
|-----------------|-------------|---------------------------------------------------------|
| pho1            | 5.75        | acid phosphatase                                        |
| SPBP2B2.06c     | 5.59        | extracellular 5′ nucleotidase                           |
| ecB             | 5.29        | extender of chronological lifespan                     |
| tdp1            | 4.69        | glycerophosphodiester transporter                      |
| pho84           | 4.18        | phosphate transporter                                   |
| SPCC11E10.01    | 2.67        | cystathionine beta-lyase                                |
| gpd3            | 2.55        | glyceraldehyde 3-phosphate dehydrogenase                |
| ngl1            | 2.34        | peptide N-glycanase                                    |
| SPBC1703.08c    | 2.31        | 5-formyltetrahydrofolate cyclo-ligase                   |
| isp4            | 2.18        | oligopeptide transmembrane transporter                  |
| SPBC1683.01     | 2.01        | phosphate transporter                                   |
| gep4            | 1.91        | phosphatidyglycerol phosphate phosphatase               |
| aoe1            | 1.84        | phenazine biosynthesis                                  |
| SPBC2582.08     | 1.78        | fettin related conserved fungal protein                 |
| rds1            | 1.72        | fettin related conserved fungal protein                 |
| SPAPB8E5.10     | 1.70        | fettin related conserved fungal protein                 |
| SPBC8E4.04      | 1.57        | aldol NAD+ 1-oxidoreductase                             |
| sdu1            | 1.49        | deubiquitinase/desumoylase                             |
| SPAC2H10.01     | 1.49        | Zn2-Cys6 transcription factor                          |
| qcr8            | 1.46        | ubiquinol-cytochrome-c reductase complex                |
| oga1            | 1.45        | Stm1 homolog                                           |
| SPAC1039.01     | 1.44        | amino acid transmembrane transporter                   |
| SPCC1235.01     | 1.42        | fettin related conserved fungal protein                 |
| mcbl            | 1.41        | MCM binding protein                                    |
| mae1            | 1.37        | malate/succinate:proton symporter                      |
| tpr1            | 1.37        | Paf1 complex subunit                                   |
| SPBPB7E8.01     | 1.37        | GPI anchor protein                                      |
| cyt1            | 1.32        | cytochrome c1                                          |
| tef10           | 1.30        | translation elongation factor EF-1 alpha                |
| mic26           | 1.29        | MICOS complex subunit                                  |
| pcm1            | 1.29        | RNA cap guanine-N7 methyltransferase                   |
| tef102          | 1.28        | translation elongation factor EF-1 alpha                |
| wdr13           | 1.27        | WD repeat protein                                      |
| qcr6            | 1.26        | ubiquinol-cytochrome-c reductase complex                |
| SPBC388.08      | 1.23        | fettin related conserved fungal protein                 |
| mae2            | 1.21        | malate dehydrogenase                                   |
| SPBC1703.13c    | 1.20        | mitochondrial phosphate transporter                     |
| abp2            | 1.19        | structural organization                                 |
| skl4            | 1.18        | structural organization                                 |
| mug2            | 1.13        | DUF1773 family protein                                 |
| set10           | 1.13        | lysine methyltransferase                               |
| mef1            | 1.11        | mitochondrial translation elongation factor G          |
| SPBC26H8.11c    | 1.10        | acyl-coenzyme A thioesterase                           |
| SPBPB21E7.09    | 1.09        | L-asparaginase                                         |
| aah2            | 1.09        | alpha-amylase                                          |
| leo1            | 1.09        | Paf1 complex subunit                                   |
| mic19           | 1.08        | MICOS complex subunit                                  |
| cox3            | 1.06        | cytochrome c oxidase subunit                           |
| cpn3            | 1.06        | kinetochore protein CENP-C                             |
| mfs3            | 1.06        | spermidine transporter                                 |
| coxl2           | 1.01        | cytochrome c oxidase subunit                           |
lase superfamily (45). Moreover, the synthetic lethality of asp1-H397A aps1Δ was suppressed by rpb1-CTD-T4A and by mutations of CPF subunits Ppn1, Swd22, Ssu72, and Ctf1 (29). These and other data, especially the effects of altered IP8 (or 1-IP7) levels on PHO gene expression, collectively pointed to IP8 (or 1-IP7) as an agonist of 3' processing/termination. Accordingly, absence of IP8/1-IP7 in asp1Δ cells (26) resulted in PHO gene hyper-repression (29); increased IP8 in asp1-H397A cells (26) resulted in de-repression of the PHO regulon under phosphate-replete conditions (29); and too much 1-IPPs in asp1-H397A aps1Δ cells elicited lethal precocious termination events affecting one or more essential fission yeast genes. Two key observations vis-à-vis the STF phenomenon are: (i) whereas the CTD-T4A mutation erased the strong de-repression of Pho1 by asp1-H397A, the low level of Pho1 expression in the phosphate-replete CTD-T4A asp1-H397A double-mutant was nonetheless 5-fold higher than that in hyper-repressed CTD-T4A single-mutant cells (29), i.e. in retrospect, asp1-H397A had a weak but detectable STF phenotype; and (ii) Pho1 de-repression ‘won out’ over T4A repression in the viable asp1-H397A aps1Δ CTD-T4A strain (29). These findings suggest that there is a threshold level of IP8 (or 1-IP7), above which precocious lncRNA termination is insensitive to loss of the CTD Thr4 mark.

A conundrum here is why the asp1-(W386-Stop) and asp1-(W493-Stop) nonsense alleles that emerged from the STF screen, which truncate all or most of the Asp1 pyrophosphatase domain, do not phenocopy the asp1-H397A allele used in prior studies (29), i.e. asp1-(W386-Stop) and asp1-(W493-Stop) fail to grow in a CTD-WT background whereas asp1-H397A CTD-WT cells grow well at all temperatures tested. Rather, asp1-(W386-Stop) and asp1-(W493-Stop) mimic the lethality and PHO de-repressive phenotypes observed for the asp1-H397A aps1Δ double pyrophosphatase-dead strain. A possible distinction is that the asp1-STF mutations are in the context of an otherwise unperturbed chromosomal asp1 locus, whereas the asp1-H397A strain was constructed by replacing the asp1+ gene with a marked mutant allele that is flanked by a kanamycin-resistance gene cassette (24). The asp1-H397A-kanMX junction in the asp1-H397A strain is such that the promoter element for the kanMX cassette was inserted 6 nt downstream of the asp1+ ORF stop codon, signifying that the native asp1+ poly(A) site and terminator were displaced during mutant construction. This raised the possibility that the level of asp1-H397A expression in the asp1-H397A-kanMX strain, and hence the level of unopposed Asp1 IPP kinase activity, might differ from that of an asp1-STF strain in which there is no 3' flanking marker. To address this point, we introduced a kanMX cassette 6 nt downstream of the ‘native’ asp1+ stop codon in the STF-6, STF-9, STF-2 and STF-12 mutant strains. The marked STF strains were mated to a wild-type rpb1-CTD strain and a large population of random haploid progeny was screened for the drug-resistance markers linked to the asp1-STF and rpb1-CTD-WT genes. Failure to recover any doubly drug-resistant STF-6 or STF-9 haploids signified that the lethal-
vity of the *asp1-STF-6* and *asp1-STF-9* alleles in a wild-type background could not be suppressed by the presence of a 3' flanking gene analogous to that of the Fleig lab's *asp1-H397A* strain. Doubly marked *STF-2 CTD-WT* and *STF-12 CTD-WT* haploids were recovered and they displayed growth defects comparable to those seen in Figure 4A for the *STF-2 CTD-WT* and *STF-12 CTD-WT* strain in which the *asp1-STF* alleles were unmarked.

An alternative conjecture is that binding of IP8 (or 1-IP7) to the active site of the C-terminal pyrophosphatase domain of Asp1 exerts a negative effect (presumably allosteric) on the activity of the N-terminal IPP kinase domain. This scenario resonates with hypotheses of interdomain allosteric communication for bifunctional mammalian IPP kinase/pyrophosphatase enzymes put forward by the Shears laboratory (46). Based on structures of histidine acid phosphatase enzymes, we envision that simple replacement of the histidine nucleophile by alanine, which obviously impairs catalysis, ought not to interdict the binding of the substrate to the Asp1 pyrophosphatase active site and hence not preclude the hypothetical allosteric effect on the kinase. By contrast, elimination of the pyrophosphatase domain (by a premature stop codon at or near the proximal margin of the pyrophosphatase domain) would certainly eliminate IP8 (or 1-IP7) binding and hence the potential for exerting a brake allosterically on the IPP kinase. A similar dampening of an allosteric effect might ensue from a missense mutation that locally distorts the pyrophosphatase active site and thereby compromises IP8 binding.

We posit that there is a narrow threshold around which the level of unopposed (or weakly opposed) Asp1 kinase activity becomes toxic or growth inhibitory, in which case *STF* mutations in the Asp1 pyrophosphatase domain might perturb the steady state levels or intracellular distribution of the Asp1 protein (even modestly), and thus the level of Asp1 IPP kinase activity, thereby leading to variable mutational effects on vegetative growth. There is an apparent trend whereby the *STF* pyrophosphatase mutations that most strongly suppress *CTD-T4A* with respect to Pho1 activity (Figure 1B) have more severe effects on vegetative growth in the *CTD-WT* background (Figures 3 and 4A) than do ‘second-round’ *STF* mutations that are relatively weaker *T4A* suppressors (Figure 8A and C).

The present transcriptome analysis of the *STF-6* and *STF-9 asp1* pyrophosphatase truncation mutants reveals a set of 51 coding genes that are coordinately upregulated compared to the parental *CTD-T4A* strain. We had previously identified 63 coding genes over-expressed in *asp1-H397A* cells versus wild-type asp1* (29). Of these, 32/63 are upregulated in *STF-6* cells and 16/63 are up-regulated in *STF-9* cells. Most pertinent is that 13 of the genes that were overexpressed in *asp1-H397A* were coordinately over-expressed in both *STF-6* and *STF-9*. These 13 are indicated in bold font in Figure 7 and include all three genes of the *PHO* regulon that are normally repressed by upstream flanking IncRNAs and are hyper-repressed by *CTD-T4A*. With respect to the 59 protein-coding genes that were down-regulated in *STF-6* and *STF-9* cells compared to the parental *T4A* strain, 11 of them were also down-regulated in *asp1-H397A* cells (indicated in bold font in Supplementary Figure S4), including four of the iron homeostasis genes.

We conclude that there is a coherent ensemble of 1-IPP-responsive genes dysregulated by inactivation of the Asp1 pyrophosphatase, be it by domain deletion or crippling of the active site and irrespective of CTD Thr4 status.

Finally, the successful implementation here of the *STF* screen rejuvenates and extends the power of classic forward genetics to illuminate the biology of the Pol2 CTD, which was initially practiced to great effect by the Young and Corden labs in the 1990s via their isolation of extragenic suppressors of budding yeast CTD truncations or S2A phospho-site mutations (47–50). The SRB (Suppressor of RNA polymerase B) screen was key to the identification via genetics of the subunits of the Pol2 holoenzyme (including components of the Mediator) that are essential for enhancer-dependent Pol2 transcription initiation. The present *STF* screen fortifies the connections between CTD Thr4 and IPP dynamics as governors of Pol2 termination in fission yeast.

### DATA DEPOSITION

The RNA-seq data in this publication have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE155609. ([https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE155609](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE155609)).

### SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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