Characterization of Schizosaccharomyces pombe RNA triphosphatase

Yi Pei, Beate Schwer1, Stéphane Hausmann and Stewart Shuman*

Molecular Biology Program, Sloan-Kettering Institute, New York, NY 10021, USA and 1Department of Microbiology and Immunology, Weill Medical College of Cornell University, New York, NY 10021, USA

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ABSTRACT

RNA triphosphatase catalyzes the first step in mRNA cap formation which entails the cleavage of the β-γ phosphoanhydride bond of triphosphate-terminated RNA to yield a diphosphate end that is then capped with GMP by RNA guanylyltransferase. Here we characterize a 303 amino acid RNA triphosphatase (Pct1p) encoded by the fission yeast Schizosaccharomyces pombe. Pct1p hydrolyzes the γ phosphate of triphosphate-terminated poly(A) in the presence of magnesium. Pct1p also hydrolyzes ATP to ADP and Pi in the presence of manganese or cobalt (K_m = 19 µM ATP; K_cat = 67 s^{-1}). Hydrolysis of 1 mM ATP is inhibited with increasing potency by inorganic phosphate (K_i = 1 mM), pyrophosphate (K_i = 0.4 mM) and tripolyphosphate (K_i = 30 µM). Velocity sedimentation indicates that Pct1p is a homodimer. Pct1p is biochemically and structurally similar to the catalytic domain of Saccharomyces cerevisiae RNA triphosphatase Cet1p. Mechanistic conservation between Pct1p and Cet1p is underscored by a mutational analysis of the putative metal-binding site of Pct1p. Pct1p is functional in vivo in S.cerevisiae in lieu of Cet1p, provided that it is coexpressed with the endogenous triphosphatase, guanylyltransferase or methyltransferase CaCet1p. Cet1p, guanylyltransferase (Ceg1p) and methyltransferase (Abd1p) gene products. The guanylyltransferase and methyltransferase components of the capping apparatus are structurally and mechanistically conserved between mammals and budding yeast. In contrast, the structures and catalytic mechanisms of the mammalian and yeast RNA triphosphatases are completely different (1).

The RNA triphosphatases of mammals and other metazoans belong to a superfamily of phosphatases that includes protein tyrosine phosphatases, dual specificity protein phosphatases and phosphoinositide phosphatases (2–5). Metazoan RNA triphosphatases catalyze γ phosphate cleavage via a two-step pathway. First, a cysteine thiolate nucleophile of the enzyme attacks the γ phosphorus to form a covalent protein-cysteiny1-S-phosphate intermediate and release diphosphate-terminated RNA. Then, the covalent intermediate is hydrolyzed to liberate inorganic phosphate. The mammalian RNA triphosphatases do not require a metal cofactor.

In contrast, the RNA triphosphatase of S.cerevisiae (Cet1p) is dependent on a divalent cation cofactor and there is no evidence of a covalent phosphoenzyme intermediate. Cet1p exemplifies a new family of metal-dependent phosphohydrolases that includes the RNA triphosphatases encoded by two groups of eukaryotic DNA viruses (poxviruses and baculoviruses) (6). The yeast/viral triphosphatase family is defined by two glutamate-rich peptide motifs (referred to as motifs A and C) that are essential for catalytic activity and comprise the metal binding site (6–9).

Genes encoding homologs of the three S.cerevisiae cap-forming enzymes have been identified in the pathogenic fungus Candida albicans (10–12). The C.albicans CaCET1 (triphosphatase), CaGT1 (guanylyltransferase) and CaCM1 (methyltransferase) genes can complement the growth of S.cerevisiae strains in which the endogenous triphosphatase, guanylyltransferase or methyltransferase genes are deleted. Indeed, the entire three-component capping system of S.cerevisiae can be replaced by the C.albicans capping system (13). The catalytic domain of the Candida RNA triphosphatase CaCet1p is similar in amino acid sequence to its S.cerevisiae counterpart Cet1p (Fig. 1). The active site of Cet1p is located within the hydrophilic core of a topologically closed eight-strand β barrel—the ‘triphosphate tunnel’ (14). The β strands comprising the tunnel are displayed over the Cet1p amino acid sequence in Figure 1. Most of the

*To whom correspondence should be addressed. Tel: +1 212 639 7145; Fax: +1 212 717 3623; Email: s-shuman@ski.mskcc.org
hydrophilic side chains of Cet1p that point into the tunnel are conserved in \textit{C. albicans} CaCet1p. Extensive mutational analyses of the tunnel in both Cet1p and CaCet1p suggest that the active site structure and catalytic mechanism are virtually identical in the \textit{Saccharomyces} and \textit{Candida} triphosphatases (6,15,16).

We consider the RNA triphosphatase to be an attractive target for antifungal drug development because: (i) the triphosphatase activity is essential for yeast cell growth; (ii) the enzyme is conserved among at least two species of fungi, including the bona fide human pathogen \textit{C. albicans} and; (iii) the proteomes of several metazoan species (nematode, arthropod and mammal) include no identifiable homologs of the known yeast RNA triphosphatases. Yet it remains possible that other species of fungi employ either a metazoan-like RNA triphosphatase or a new class of RNA triphosphatase.

We are therefore interested in tracking the stage in evolution at which the capping apparatus of lower and higher eukaryotes diverged with respect to triphosphatase structure and mechanism. Towards this end, we have initiated a molecular genetic and biochemical analysis of the cap-forming enzymes of the fission yeast \textit{Schizosaccharomyces pombe} (12,17). A wide evolutionary distance separates fission yeast from budding yeast. \textit{Schizosaccharomyces} is in many respects more closely related to metazoans in its genetic organization and pre-mRNA processing strategy than to \textit{Saccharomyces}. For example: (i) fission yeast has a much higher proportion of intron-containing genes (~45%) than budding yeast (~5%); (ii) the components of the pre-mRNA splicing machinery in fission yeast are more similar to their mammalian counterparts than to the corresponding budding yeast proteins; and (iii) several fission yeast splicing factors have homologs in humans, but are missing from the proteome of budding yeast (18).

We have already identified the genes encoding the guanylyltransferase (PCE1) and methyltransferase (PCM1) components of the \textit{S. pombe} capping apparatus (12,17). Neither gene contains an intron. Pce1p and Pcm1p are structurally similar to the \textit{S. cerevisiae} guanylyltransferase (Ceg1p) and methyltransferase (Abd1p), respectively, and expression of the \textit{S. pombe} guanylyltransferase or methyltransferase in \textit{S. cerevisiae} is sufficient for the growth of cells lacking the endogenous guanylyltransferase or methyltransferase proteins. The \textit{S. pombe} guanylyltransferase Pce1p displays extensive amino acid sequence similarity to the catalytic domains of Cet1p and CaCet1p (17), but lacks any counterpart of the N-terminal triphosphatase domain of Mce1p.

Here we show that the \textit{S. pombe} RNA triphosphatase is encoded by a separate gene, which we named \textit{PCT1} (17). \textit{PCT1} contains a 5′-proximal intron and encodes a 303 amino acid polypeptide with extensive structural similarity to the catalytic domains of Cet1p and CaCet1p. Purified recombinant Pct1p that were targeted for alanine substitution in the present study are also shaded.

\begin{figure}
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\includegraphics[width=\textwidth]{figure1.png}
\caption{\textbf{Schizosaccharomyces pombe} RNA triphosphatase is structurally related to the RNA triphosphatases of \textit{S. cerevisiae} and \textit{C. albicans}. The complete amino acid sequence of \textit{S. pombe} Pct1p is aligned to the sequences of the C-terminal catalytic domains of \textit{S. cerevisiae} Cet1p (residues 241–539) and \textit{C. albicans} CaCet1p (residues 202–520) to \textit{S. cerevisiae} Cth1p (residues 1–317). Gaps in the alignment are indicated by dashes. The β strands that comprise the triphosphate tunnel of Cet1p are denoted above the sequence. The peptide domain of Cet1p that mediates its interaction with the guanylyltransferase Ceg1p is conserved in CaCet1p and is shaded. This segment is not conserved in either Cet1p or Pct1p. Positions of side-chain identity or structural similarity in all four fungal RNA triphosphatases are denoted by dots. Conserved motifs A (β1) and C (β11) that define the metal-dependent RNA triphosphatase family are indicated below the sequence. Residues comprising the homodimer interface of Cet1p that are conserved in Pct1p are underlined below the Pct1p sequence. Residues Glu78, Glu80 and Glu260 of Pct1p that target for alanine substitution in the present study are also shaded.}
\end{figure}
protein has intrinsic metal-dependent RNA triphosphatase and nucleoside triphosphatase activities. Mechanistic conservation between fission and budding yeast triphosphatases is suggested by mutational analysis of the putative metal-binding motifs of Pct1p. Our results are consistent with an abrupt divergence of the RNA triphosphatase component of the capping apparatus during the transition from fungal to metazoan species. We find that the \textit{S. pombe} triphosphatase is functional in \textit{S. cerevisiae} in lieu of Cet1p when it is coexpressed with the fission yeast guanylyltransferase Pce1p. To our knowledge, this is the first example of species-specific genetic interactions between fungal cap-forming enzymes.

\textbf{MATERIALS AND METHODS}

\textbf{Yeast expression vectors for \textit{S. pombe} RNA triphosphatase}

The intron-containing gene SPAC644.04 which encodes a putative RNA triphosphatase (hereafter referred to as \textit{PCT1}, for ‘Pombe Capping enzyme Triphosphatase’) was amplified from \textit{S. pombe} genomic DNA using \textit{Pfu} DNA polymerase (Stratagene) and oligonucleotide primers designed to introduce an \textit{NdeI} restriction site at the translation start codon and a \textit{BamHI} restriction site immediately downstream of the stop codon. The amplified gene was inserted into the pCR-Blunt II-TOPO vector (Invitrogen) to generate the plasmid pCR-PCT1(+intron). The complete nucleotide sequence of the \textit{PCT1} coding sequence was PCR-amplified using an \textit{NdeI} coding sequence was PCR-amplified using an \textit{NdeI} antisense primer that changed the \textit{PCT1} stop codon to His and then inserted into the \textit{NdeI} site of pYX1-MCE1(211–597) (\textit{CEN TRP1}) (4) to yield the fusion gene \textit{PCT1–MCE1}(211–597). Expression of the chimeric gene is under the control of the \textit{TPP1} promoter.

\textbf{Expression and purification of recombinant \textit{S. pombe} RNA triphosphatase}

An \textit{NdeI–BamHI} restriction fragment containing the \textit{PCT1} cDNA was inserted into the bacterial expression plasmid pET16b. The resulting plasmid, pET-PCT1, was introduced into \textit{Escherichia coli} BL21(DE3). A 100 ml culture of BL21(DE3)/pET-PCT1 was grown at 37°C in Luria–Bertani medium containing 0.1 mg/ml ampicillin until the \textit{A}_{600} reached 0.5. The culture was placed on ice for 30 min and then adjusted to 0.4 mM IPTG and 2% (v/v) ethanol. Incubation was continued for 20 h at 18°C with constant shaking. Cells were harvested by centrifugation and the pellets were stored at –80°C. All subsequent procedures were performed at 4°C. Thawed bacteria were resuspended in 5 ml of lysis buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl, 10% sucrose). Lysozyme was added to a final concentration of 100 µg/ml and the suspension was incubated on ice for 15 min, adjusted to 0.1% Triton X-100 and then sonicated to reduce the viscosity of the lysate. Insoluble material was removed by centrifugation for 45 min at 17 000 r.p.m. in a Sorvall SS34 rotor. The soluble extract was applied to a 1 ml column of Ni\textsuperscript{2+}-NTA agarose that had been equilibrated with lysis buffer containing 0.1% Triton X-100. The column was washed with the same buffer and then eluted step-wise with buffer B (50 mM Tris–HCl pH 8.0, 100 mM NaCl, 10% glycerol, 0.05% Triton X-100) containing 50, 100, 200, 500 and 1000 mM imidazole. SDS–PAGE analysis showed that the recombinant Pct1p polypeptide was recovered predominantly in the 200 mM imidazole eluate fraction. The eluate was adjusted to 50 mM NaCl and 2 mM DTT by addition of an equal volume of dilution buffer (50 mM Tris–HCl pH 8.0, 4 mM DTT, 10% glycerol, 0.05% Triton X-100). This material was applied to a \textit{C} column of phosphocellulose that had been equilibrated with buffer C (50 mM Tris–HCl pH 8.0, 10% glycerol, 0.05% Triton X-100) containing 50 mM NaCl. The column was washed with the same buffer and then eluted step-wise with buffer C containing 0.1, 0.2, 0.5 and 1.0 M NaCl. Pct1p was recovered in the 0.2 and 0.5 M NaCl eluate fractions. The 0.5 M NaCl fraction was pooled and the protein concentration was determined using the Bio-Rad dye-binding method with bovine serum albumin (BSA) as the standard. The yield of recombinant Pct1p was ∼0.4 mg.

Alanine-substitution mutations at positions Glu78, Glu80 and Glu260 were introduced into the \textit{PCT1} gene by the two-stage overlap extension method (21). The mutated genes were digested with \textit{NdeI} and \textit{BamHI} and then inserted into the bacterial expression vector pET16b. The inserts were sequenced completely to confirm the desired alanine mutation and exclude the triphosphatase and guanylyltransferase transcription units arranged in a tail-to-tail configuration.

\textbf{Chimeric \textit{S. pombe/mouse} capping enzyme}

A gene encoding Pct1p fused to the guanylyltransferase domain of the mouse capping enzyme [Mec1(211–597)]p was constructed as follows. The \textit{PCT1} coding sequence was PCR-amplified using an antisense primer that changed the \textit{PCT1} stop codon to His and introduced an \textit{NdeI} restriction site at the C-terminus. The PCR product was digested with \textit{NdeI} and then inserted into the \textit{NdeI} site of pYY1-MCE1(211–597) (\textit{CEN TRP1}) (4) to yield the fusion gene \textit{PCT1–MCE1}(211–597). Expression of the chimeric gene is under the control of the \textit{TPP1} promoter.
the acquisition of unwanted changes during amplification or cloning. The pET-PCT1-Ala plasmids were transformed into E.coli BL21(DE3) and the recombinant mutant proteins were expressed and purified as described above for wild-type Pct1p.

RESULTS

PCT1: a candidate S.pombe RNA triphosphatase gene

We identified, on phylogenetic grounds, a candidate RNA triphosphatase gene by searching the S.pombe Genome Sequencing Project Database (Sanger Center) for proteins related to the S.cerevisiae RNA triphosphatase Cet1p. The initial search revealed a gene fragment on S.pombe chromosome I encoding a polypeptide with significant similarity to the catalytic domain of Cet1p extending from motif A in strand β1 to motif C in strand β11. Subsequent annotation of this portion of chromosome I by the Sanger Center indicated that the S.pombe gene fragment corresponds to the 3′ exon of an intron-containing gene SAC644.04 (GenBank accession no. AL355012). The predicted spliced mRNA product of this gene (which we have renamed PCT1) encodes a 303 amino acid polypeptide (Fig. 1). The 5′ exon of PCT1 encodes amino acids 1–78 and the 3′ exon encodes amino acids 79–303. Figure 1 shows a structure-based alignment of the sequence of Pct1p with the sequences of three known fungal RNA triphosphatases: S.cerevisiae Cet1p, C.albicans CaCet1p and S.cerevisiae Cth1p. Cth1p is a non-essential S.cerevisiae enzyme of unknown function (15,22). Positions of side-chain identity or conservation in all four proteins are denoted by dots. Reference to the crystal structure of Cet1p indicates that the identical and conserved residues are clustered in the β strands that comprise the walls of the triphosphate tunnel (Fig. 1) and include many of the hydrophilic amino acids that are essential for catalysis by Cet1p and CaCet1p. The 303 amino acid Pct1p protein is considerably smaller than Cet1p (549 amino acids) or CaCet1p (520 amino acids); Cet1p and CaCet1p contain non-essential N-terminal extensions that are missing from Pct1p. In the experiments presented below, we test biochemically and genetically whether Pct1p has the requisite activities of a cap-forming enzyme in vitro and in vivo.

Metal-dependent phosphohydrolase activity of Pct1p

A cDNA encoding the 303 amino acid S.pombe polypeptide was constructed by deleting the intron from a genomic DNA fragment containing the PCT1 gene. The cDNA was then cloned into a T7 RNA polymerase-based pET vector so as to place the ORF in-frame with an N-terminal leader encoding a 21 amino acid peptide with 10 tandem histidines. The expression plasmid was introduced into E.coli BL21(DE3), a strain that contains the T7 RNA polymerase gene under the control of the lac promoter. A new 38 kDa polypeptide corresponding to Pct1p was detectable by SDS–PAGE in extracts of IPTG-induced bacteria bearing the pET-PCT1 plasmid (not shown). Initial purification of the His-tagged fusion protein was achieved by adsorption to Ni-agarose and elution with 200 mM imidazole. Pct1p was purified further by adsorption to a column of phosphocellulose and step elution with NaCl. The phosphocellulose preparation was nearly homogenous with respect to the Pct1p polypeptide, as judged by SDS–PAGE (Fig. 2A). Further characterization of recombinant Pct1p was performed using the phosphocellulose fraction.

We found that purified recombinant Pct1p is indeed an RNA triphosphatase. Activity was assayed by the liberation of $^{32}$P from 2 µM γ-$^{32}$P-labeled triphosphate-terminated poly(A) in the presence of 1 mM magnesium chloride. The extent of γ phosphate hydrolysis during a 15 min incubation at 30°C was proportional to input protein (Fig. 2C). In the linear range of enzyme dependence, 180 fmol of $^{32}$P, was released per fmol of Pct1p. This value corresponds to a turnover number of ~0.2 s$^{-1}$, which is lower than the values reported for the hydrolysis of [γ-$^{32}$P]poly(A) by S.cerevisiae Cet1p (1 s$^{-1}$). C.albicans

Figure 2. RNA triphosphatase and ATPase activities of Pct1p. (A) Protein purification. Aliquots (4 µg) of the phosphocellulose fractions of wild-type Pct1p (WT) and mutants E78A, E80A and E260A were analyzed by electrophoresis in a 12% polyacrylamide gel containing 0.1% SDS. Polypeptides were visualized by staining with Coomassie blue dye. The positions and sizes (in kDa) of marker proteins are indicated to the left of the gel. (B) ATPase activity. Reaction mixtures (10 µl) containing 50 mM Tris–HCl pH 7.5, 5 mM DTT, 2 mM MnCl$_{2}$, 1 mM [γ-$^{32}$P]ATP and either wild-type (WT) or mutant proteins as specified were incubated for 15 min at 30°C. Reactions were quenched by adding 2.5 µl of 5 M formic acid. Aliquots of the mixtures were applied to a polyethyleneimine–cellulose thin-layer chromatography (TLC) plate, which was developed with 1 M formic acid, 0.5 M LiCl. $^{32}$P release was quantitated by scanning the chromatogram with a FUJIX phosphorimager and was plotted as a function of input protein. (C) RNA triphosphatase activity. Reaction mixtures (10 µl) containing 50 mM Tris–HCl pH 7.5, 5 mM DTT, 1 mM MgCl$_{2}$, 20 pmol (of triphosphate termini) of [γ-$^{32}$P]poly(A) and either WT or mutant proteins as specified were incubated for 15 min at 30°C. $^{32}$P release is plotted as a function of input protein.
CaCet1p (1.4 s⁻¹), baculovirus LEF4 (1 s⁻¹) and vaccinia D1 (0.8 s⁻¹) (6,8,16,23).

The signature feature of the fungal/viral triphosphatase family members is their ability to hydrolyze nucleoside triphosphates to nucleoside diphosphates and inorganic phosphate in the presence of manganese or cobalt (6,8,9,16). The divalent cation specificity of the NTPase is distinct from the RNA triphosphatase function, which is optimal in magnesium. We found that recombinant Pct1p catalyzed the release of ³²Pi from [γ-³²P]ATP in the presence of manganese and that the extent of ATP hydrolysis increased as a function of input enzyme (Fig. 2B). There was no detectable ATP hydrolysis in the absence of a divalent cation (Fig. 3A). Hydrolysis of 1 mM ATP was optimal at 1–2 mM MnCl₂ and declined slightly at 3–5 mM MnCl₂ (Fig. 3A). ATP hydrolysis with cobalt as the cofactor was optimal at 1.5–6 mM CoCl₂ (Fig. 3A). The titration curves were sigmoidal at manganese and cobalt concentrations below the level of input ATP. ATPase activity was tested with a battery of divalent actions added at 2 mM concentration: cobalt was ~70% as effective as manganese, whereas magnesium, calcium, copper and zinc were inactive (Fig. 3B). The manganese-dependent ATPase activity of Pct1p in 50 mM Tris–HCl buffer was optimal between pH 7.5 and 9.0; the activity at pH 6.0 was 33% of the activity at pH 7.5 (data not shown).

Kinetic analysis of ATP hydrolysis

The rate of release of ³²P from [γ-³²P]ATP was nearly identical to the rate of conversion of [α-³²P]ATP to [α-³²P]ADP in a parallel reaction mixture containing the same concentration of Pct1p (Fig. 4A). We detected no formation of [α-³²P]AMP during the reaction. Hence, we conclude that Pct1p catalyzes the hydrolysis of ATP to ADP and Pi. Pct1p catalyzed the conversion [α-³²P]GTP to [α-³²P]GDP at the same rate as it hydrolyzed ATP to ADP (data not shown). Pct1p also hydrolyzed [α-³²P]dATP to [α-³²P]dADP at approximately half the rate of ATP hydrolysis (not shown). Other nucleotides were not tested.

Kinetic parameters for Pct1p were determined by measuring the extent of ³²P formation during a 15 min reaction as a function of input [γ-³²P]ATP concentration in the range 2.5–60 μM. A double-reciprocal plot of the data fits well to a linear function (Fig. 4B). We calculated a $K_m$ of 19 μM ATP and a $k_{cat}$
of 67 s\(^{-1}\). The turnover number of Pct1p in ATP hydrolysis is higher than the values reported for Cet1p (25 s\(^{-1}\)), CaCet1p (17 s\(^{-1}\)), Chl1p (2 s\(^{-1}\)), D1 (10 s\(^{-1}\)) and LEF4 (30 s\(^{-1}\)) (6,8,15,16,24). The \(K_m\) of Pct1p for ATP (19 \(\mu\)M) falls in the middle of the range reported for other family members: Cet1p (3 \(\mu\)M), CaCet1p (9 \(\mu\)M), LEF4 (43 \(\mu\)M), Chl1p (75 \(\mu\)M) and D1 (800 \(\mu\)M).

**Pct1p activity is abolished by replacement of motif A and C glutamates with alanine**

Pct1p residues Glu78 and Glu80 in motif A and Glu260 in motif C (Fig. 1) were replaced individually by alanine. The E78A, E80A and E260A proteins were expressed as His-tagged fusions and purified from soluble bacterial lysates by Ni-agarose and phosphocellulose column chromatography. The purity of the E80A and E260A proteins was comparable to that of wild-type Pct1p (Fig. 2A). The E78A, E80A and E260A mutants were unable to hydrolyze \(\gamma\)^-32P-labeled poly(A) or ATP even when the levels of input enzyme were far in excess of the amount sufficient for maximal release of \(\gamma\)^-P, by wild-type Pct1p (Fig. 2B and C). Using these data, we calculated that the specific RNA triphosphatase and ATPase activities of E78A, E80A and E260A were <0.1% of the activity of wild-type enzyme. We conclude that both glutamates of motif A and the central glutamate of motif C are essential for the phosphohydrolase activity of Pct1p.

This trio of acidic amino acids is broadly conserved in the RNA triphosphatases encoded by *S.cerevisiae*, *C.albicans*, poxviruses, African swine fever virus and baculoviruses. The equivalent three glutamates in motif A and motif C directly coordinate manganese at the metal binding site of Cet1p and are essential for catalysis by Cet1p in *in vitro* and for Cet1p function in *in vivo* (6,14). The motif A and C glutamates are also essential for catalysis by CaCet1p, Chl1p, vaccinia D1 and baculovirus LEF4 (7,9,15,16,25). Motifs A and C are located within strands \(\beta\) and \(\beta1\) of Cet1p, which are widely separated in the primary structure, but closely approximated in the tertiary structure along the ‘floor’ of the tunnel (14). In motifs A and C of the budding yeast RNA triphosphatases, alternating charged side chains are interdigitated with alternating aliphatic/aromatic side chains (Fig. 1). This sequence pattern is reprised in motifs A and C of the *S.pombe* RNA triphosphatase, suggesting that the metal-binding residues of the fission yeast enzyme are also located within a pair of \(\beta\) strands.

**Sedimentation analysis**

The native size of recombinant Pct1p was analyzed by sedimentation of the protein through a 15–30% glycerol gradient containing buffer G. Marker proteins catalase, BSA and cytochrome c were included as internal standards in the same glycerol gradient. After centrifugation, the polypeptide compositions of the gradient fractions were analyzed by SDS-PAGE (Fig. 5, top). A plot of the S values of the three standards versus fraction number yielded a straight line (not shown). Pct1p (a 38 kDa polypeptide) was resolved into two discrete peaks: a predominant 4.7 S ‘light’ component sedimenting just ahead of BSA (a 68 kDa monomeric polypeptide) and a minor 12 S ‘heavy’ component sedimenting just ahead of catalase (a 248 kDa tetramer of a 62 kDa subunit). We conclude that the predominant form of Pct1p is a homodimer and the minor species is at least a hexamer, if not an octamer. The gradient fractions were then assayed for manganese-dependent ATPase. The bimodal ATPase activity profile coincided with the distribution of Pct1p polypeptide (Fig. 5, bottom). The same bimodal distribution of Pct1p protein and ATPase activity was observed when the enzyme was sedimented in the absence of internal standards (data not shown).

The homodimeric quaternary structure of *S.pombe* Pct1p is a feature that is shared with the RNA triphosphatase of *S.cerevisiae* (14,26). Indeed, many of the amino acids that comprise the mostly hydrophobic dimerization surface in the crystal structure of *S.cerevisiae* Cet1p (14) are conserved in the *S.pombe* protein (Fig. 1).

**Inhibition of Pct1p ATPase by phosphate, pyrophosphate and tripolyphosphate**

We tested the effects of various phosphate derivatives on the ability of 30 nM Pct1p to hydrolyze 1 mM ATP in the presence of 2 mM manganese (Fig. 6). Inorganic phosphate, which is a product of the phosphohydrolase reaction, inhibited activity in a dose-dependent manner. ATP hydrolysis was inhibited by 50% of the control value at 1 mM P\(_i\) (Fig. 6). Inorganic pyrophosphate was a better inhibitor than phosphate (50% inhibition at 0.4 mM P\(_i\)) while tripolyphosphate was >10-fold more potent than pyrophosphate (50% inhibition at 30 \(\mu\)M tripolyphosphate).

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**Figure 5.** Glycerol gradient sedimentation. Pct1p (40 \(\mu\)g) was mixed with catalase (40 \(\mu\)g), BSA (40 \(\mu\)g) and cytochrome c (40 \(\mu\)g) in 0.2 ml of buffer G (50 mM Tris–HCl pH 8.0, 100 mM NaCl, 1 mM EDTA, 2 mM DTT, 0.05% Triton X-100). The mixture was layered onto a 4.8 ml 15–30% glycerol gradient containing buffer G. The gradient was centrifuged in a Beckman SW50 rotor at 50 000 r.p.m. for 15 h at 4°C. Fractions (0.2 ml) were collected from the bottom of the tube. (Top) Aliquots (18 \(\mu\)l) of odd numbered fractions were analyzed by SDS–PAGE along with samples of the input protein mixture (L). Polypeptides were visualized by staining with Coomassie blue dye. The identities of the polypeptides are indicated at the left of the gel. (Bottom) ATPase activity profile. Reaction mixtures (20 \(\mu\)l) containing 50 mM Tris–HCl pH 7.5, 5 mM DTT, 2 mM MnCl\(_2\), 1 mM \([\gamma\)^-32P\]ATP and 2 \(\mu\)l of the indicated manganese gradient fractions were incubated for 15 min at 30°C. The peaks of the marker proteins are indicated by arrows.
Figure 6. Inhibition of ATP hydrolysis by phosphate, pyrophosphate and tripolyphosphate. Reaction mixtures (10 µl) containing 50 mM Tris–HCl pH 7.5, 5 mM DTT, 2 mM MnCl₂, 1 mM [γ-32P]ATP, 11 ng of Pct1p and either sodium phosphate, sodium pyrophosphate or sodium tripolyphosphate as specified were incubated for 15 min at 30 °C. The reactions were initiated by the addition of Pct1p. The extents of ATP hydrolysis were normalized to those of control reactions from which inhibitory phosphates were omitted (values of 2.5–3.5 nmol 32P released). The normalized activities (control value = 1.0) are specified were incubated for 15 min at 30 °C.

The finding that tripolyphosphate elicited 50% inhibition when present at a concentration 33-fold less than input ATP and 67-fold less than input manganese argues against the possibility that tripolyphosphate inhibits by acting as a chelator to compete with ATP (or enzyme) for the metal cofactor. A simple explanation for the potent inhibition by tripolyphosphate is that it binds more avidly than ATP to the triphosphate-binding pocket within the active site tunnel.

Figure 7. Genetic interaction between *S.pombe* RNA triphosphatase and *S.pombe* RNA guanylyltransferase. Yeast strain YBS20 (cet1Δ) was transformed with *CEN TRP1* plasmids containing either CET1, PCT1, PCT1 + PCE1 or PCT1 + CEG1 and with a 2µ plasmid containing PCT1. Trip+ isolates were streaked on an agar plate containing 0.75 mg/ml 5-FOA. The plate was photographed after incubation for 3 days at 30 °C.
and negative controls, respectively. Growth of YBS50 depends on maintenance of a CEN URA3 CET1 CEG1 plasmid (26). Transformation of YBS50 with MCE1, which encodes the bifunctional mammalian triphosphatase–guanylyltransferase, allowed growth of YBS50 on medium containing 5-FOA (Fig. 8). Transformants bearing MCE1(211–597), which encodes only the guanylyltransferase domain of mouse capping enzyme, failed to give rise to FOA-resistant colonies. This was expected because a functional triphosphatase on the CEN TRPI plasmid is needed to complement the cet1Δ ceg1Δ double deletion. The instructive finding was that YBS50 cells bearing the PCT1–MCE1(211–597) plasmid did grow on 5-FOA (Fig. 8). Moreover, FOA-selected PCT1–MCE1(211–597) cells grew as well as MCE1 cells on rich medium (YPD agar) at 30 and 37°C, as gauged by colony size (data not shown). We conclude from this experiment that S. pombe Pct1p can function in vivo in mRNA capping when it is appropriately targeted to the transcription complex.

Specific genetic interaction between S. pombe triphosphatase and guanylyltransferase

The fact that S. pombe guanylyltransferase Pce1p by itself can replace Ceg1p in vivo in S. cerevisiae (17) implies that S. pombe guanylyltransferase can interact with the S. cerevisiae triphosphatase. However, S. pombe triphosphatase by itself is not functional in budding yeast—possibly because it does not interact with Ceg1p. If Pct1p is a bona fide capping enzyme in S. pombe, then we postulate that it is targeted to the transcription complex via a chaperone specific to S. pombe. Given that an association of the triphosphatase and guanylyltransferase components of the capping apparatus is conserved in evolution (with the interaction being in trans in S. cerevisiae and in cis in metazoans, poxviruses and baculoviruses), the simplest model would be that S. pombe triphosphatase interacts with S. pombe guanylyltransferase Pce1p. To test this hypothesis, we asked whether the in vivo function of Pct1p in budding yeast could be restored by coexpressing Pce1p.

Yeast CEN TRPI expression plasmids were constructed that contained a PCT1 cDNA driven by the yeast GDP1 promoter plus either the S. pombe PCE1 gene or the S. cerevisiae CEG1 gene driven by the TP11 promoter. These dual expression vectors were transformed into the yeast cet1Δ strain and RNA triphosphatase activity of the plasmid-borne PCT1 allele was tested by plasmid shuffle (Fig. 7). The salient and instructive findings were that coexpression of Pct1p with S. pombe Pce1p completely revived the in vivo activity of Pct1p (as measured by growth on 5-FOA), whereas coexpression with the budding yeast guanylyltransferase Ceg1p had no salutary effect on Pct1p (Fig. 7). (Control experiments verified that the plasmid containing PCT1 plus CEG1 did complement the S. cerevisiae ceg1Δ mutant.) We subsequently found that budding yeast cells containing the triphosphatase and guanylyltransferase components of the fission yeast capping apparatus grew as well on rich medium (YPD agar) at 30 and 37°C as cells containing the wild-type Saccharomyces capping enzymes (not shown).

DISCUSSION

Our biochemical and genetic studies of the S. pombe RNA triphosphatase Pct1p provide new insights into the evolution and divergence of the mRNA capping apparatus in lower and higher eukaryotes. Given the sequence similarities between the S. pombe RNA triphosphatase and the catalytic domains of the S. cerevisiae and C. albicans RNA triphosphatases, the similar catalytic repertoires of these enzymes in metal-dependent hydrolysis of triphosphate-terminated RNA and free nucleoside triphosphates and the concordance of mutational effects in the metal-binding motifs A and C of all three proteins, we conclude that the active site architectures of fission yeast and budding yeast RNA triphosphatases are highly conserved. A reasonable extrapolation from the analysis of three fungal species is that all fungi are likely to encode the same basic capping machinery. This machinery is composed of separate triphosphatase, guanylyltransferase and methyltransferase gene products, with triphosphatase components from the metal-dependent family of Cet1p-like phosphohydrolases rather than the metal-independent cysteine-phosphatase family found in metazoans and plants. Thus, RNA triphosphatase retains its potential for being a target for mechanism-based antifungal drug discovery, insofar as an inhibitor that either occupies or occludes the active site of C. albicans RNA triphosphatase will probably be active against RNA triphosphatases from other pathogenic fungi. Tripolyphosphate is a relatively potent inhibitor of the Pct1p phosphohydrolase activity and a worthwhile platform on which to build and test new derivatives.

The guanylyltransferase and triphosphatase activities are linked in cis within a single polypeptide in all metazoan organisms examined to date [including the nematode Caenorhabditis elegans (2), the amphibian Xenopus laevis (33) and the arthropods Drosophila melanogaster and Artemia salina (34), as well as mice and humans] and in the plant Arabidopsis thaliana. The metazoan and plant RNA triphosphatases, members of the cysteine-phosphatase superfamily, are structurally and mechanistically unrelated to the fungal RNA triphosphatases. In all fungi examined to date, the triphosphatase and guanylyltransferase activities reside in separate polypeptides that interact in trans. The guanylyltransferase components are structurally and mechanistically similar in all eukaryotes. The mammalian guanylyltransferase domain can support yeast cell growth (albeit slower growth relative to wild-type yeast) with the yeast triphosphatase Cet1p present in trans (4). Moreover,
the mouse guanylyltransferase displays a vestigial low-affinity interaction in vitro with the Ceg1p-binding peptide domain of yeast Cet1p (31). These findings suggest that the guanylyltransferases of higher eukaryotes evolved from an ancestral enzyme that did interact in trans with a triphosphatase, but that selection for such an interaction was relaxed during the emergence of metazoa.

The enzymes that catalyze the basic nucleic acid transactions (DNA replication, DNA repair, RNA synthesis and RNA processing) are generally very well conserved in lower and higher eukaryotes. Yet, in the case of RNA triphosphatase, we see a complete divergence of structure and mechanism either at the time of metazoan evolution or prior to the branching of the metazoan phyla listed above. How might this have occurred? We envisage a gene rearrangement event early in metazoan evolution that transferred a cysteine-phosphatase domain into the same transcription unit as the guanylyltransferase transcription unit, leading to creation of the triphosphatase–guanylyltransferase fusion protein that we see today in higher eukaryotic species. Note that lower and higher eukaryotes encode a plethora of cysteine-phosphatase family members with diverse substrate specificity and one can easily imagine encoding a species-specific genetic interaction long after the proposed gene fusion. Note also that eukaryotes and eukaryotic viruses encode additional RNA 5′ phosphatases of the cysteine-phosphatase family. These proteins are not linked in cis to a guanylyltransferase and are of unknown function (35–37). RNA triphosphatase-like cysteine-phosphatases are not evident in the yeast proteome. Also, we cannot discern a homolog of RNA triphosphatase-like cysteine-phosphatases are not evident in the yeast proteome. Also, we cannot discern a homolog of RNA triphosphatases that exist in metazoans are those encoded by large DNA viruses such as poxviruses and baculoviruses.

Although the RNA triphosphatase components of the fungal capping systems have the same catalytic domain, the present study illustrates that the capping components are not always functionally interchangeable in vivo. There is an apparent hierarchy of cross-species complementation, whereby all of the known fungal guanylyltransferases can function in Schizosaccharomyces with the endogenous Cet1p triphosphatase (10,17), but the ability of heterologous triphosphatases to function with the Schizosaccharomyces guanylyltransferase is variable and correlates with the presence or absence of a conserved guanylyltransferase-binding domain on the surface of the heterologous RNA triphosphatase. Schizosaccharomyces pombe triphosphatase Pct1p, which lacks the surface domain and does not function on its own in Schizosaccharomyces, can support cell growth when S. pombe guanylyltransferase Pce1p is provided in trans. Thus, the fission yeast capping components display a species-specific genetic interaction in vivo. Insights into the physical basis of the Pct1p–Pce1p interaction will be dependent on crystallization of the S. pombe enzymes and fine mapping of their functional domains by mutagenesis.

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