Pho5p and Newly Identified Nucleotide Pyrophosphatases/Phosphodiesterases Regulate Extracellular Nucleotide Phosphate Metabolism in Saccharomyces cerevisiae

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Extracellular nucleotides play many biological roles, including intercellular communication and modulation of nucleotide receptor signaling, and are dependent on the phosphorylation state of the nucleotide. Regulation of nucleotide phosphorylation is necessary, and a specialized class of enzymes, nucleotide pyrophosphatases/phosphodiesterases (E-NPPs), has been identified in mammals to perform this function. Although the E-NPP class is conserved among complex eukaryotes, this system has not yet been identified in Saccharomyces cerevisiae. Using genetic and biochemical experiments, we show that two orthologs of the E-NPP family, referred to as Npp1p and Npp2p, exist in budding yeast and can perform nucleotide phosphate hydrolysis. This activity is enhanced during phosphate starvation, where hydrolyzed phosphates can be imported from extracellular sources and utilized to overcome phosphate starvation through the activity of the Pho5p acid phosphatase. The added compensatory effect by Pho5p is also a newly established role for Pho5p. This study demonstrates that extracellular nucleotide phosphate metabolism appears to be controlled by at least two independent regulatory mechanisms, uniting phosphate starvation with extracellular nucleotide regulation.

The phosphorylation state of extracellular nucleotides is critical for maintaining intercellular communication. Nucleotide phosphates are also involved in activation of many cell surface receptors for initiation of various intracellular processes. It is critical to regulate the phosphorylation state of extracellular nucleotides, as well as to control extracellular pyrophosphate concentrations, and it must be tightly synchronized with dynamic environmental conditions. A regulatory mechanism to control extracellular nucleotide hydrolysis consists of a multigene nucleotide pyrophosphatase/phosphodiesterase (E-NPP) family (14). Five members of this family have been identified in humans (NPP1 to -5), each with distinct functional roles (10, 13, 19, 22). NPP1 to -3 are found in nearly all human tissue types, and these enzymes are categorized to contain an alkaline ecto-nucleotide pyrophosphatase/phosphodiesterase type 1 domain (12). These enzymes catalyze hydrolysis of pyrophosphate and phosphodiester bonds from nucleotide sources. All three are involved in regulating nucleotide recycling. In addition, NPP1 and NPP2 may regulate pathologic mineralization, and NPP2 and NPP3 are involved in regulating cell motility (33, 43). NPP4 and NPP5 are additional members of this family; however, little is known about their activity. Several disorders have been found to correlate with misregulation or altered activity of E-NPP family members, including diabetes (5), chondrocalcinosis deposition disease (21), and tumor migration (48). Thus, although the NPP family performs considerable physiological roles in humans, the discovery of cellular roles of E-NPPs has not been previously established for Saccharomyces cerevisiae.

Another regulatory mechanism in yeast involves phosphate sensing and acquisition from the extracellular medium. Inorganic phosphate (P_i) is an essential nutrient for the biosynthesis of many cellular components. The availability of nutrients in the environment is critical in determining the cellular decision to undergo processes such as growth and proliferation. When phosphate supplies are limited, cells induce a response to acquire inorganic phosphate from multiple sources for cellular uptake (PHO response) (24). This includes expression of several genes, such as those encoding nonspecific scavenger phosphatases, to generate inorganic phosphates from various phosphate-containing molecules (6, 32). Free inorganic phosphate is subsequently transported and localized into the cytoplasm as a source for metabolic processes (47).

A hallmark of high-affinity phosphate transport pathway activation is the transcriptional upregulation of two classes of genes: acid phosphatases and phosphate symporters (2). The PHO response upregulates three genes (PHO5, PHO11, and PHO12) encoding repressible acid phosphatases (APases) when phosphate is scarce. These phosphatases are localized to either the periplasmic space or the cell wall and are responsible for phosphate scavenging by working in conjunction with high-affinity transporters to acquire phosphate when P_i concentrations in the environment are low (31, 41). Of the three phosphatases, Pho5p is responsible for >90% of APase activity (44). Whereas the APases are believed to utilize many different substrates for phosphate scavenging, it remains unclear whether they play a role in extracellular nucleotide recycling through phosphate hydrolysis.
The other key components of the PHO transport pathway are phosphate symporters. Two classes of symporters have been identified for phosphate uptake: a low-affinity and a high-affinity transport system (35). The low-affinity system is believed to be constitutively expressed, regardless of extracellular phosphate concentrations. It includes Pho87p, Pho90p, and Pho91p (2) and can import extracellular phosphate at a $K_m$ of 770 $\mu$M (45). The high-affinity phosphate regulatory system mediates specific uptake of inorganic phosphate from extracellular sources under phosphate starvation conditions when external phosphate concentrations fall below 100 $\mu$M (28). Under low-phosphate conditions, expression of the high-affinity transporters, Pho84p and Pho89p, is rapidly increased. This correlates with the decrease in expression of low-affinity phosphate transporters. These transporters have an approximately 10- to 100-fold-lower $K_m$ value for extracellular phosphates than their low-affinity counterparts (27, 34, 38).

In this report, we show that genes encoding putative E-NPPs (NPP1 and NPP2) can catalyze NPPase reactions and contribute to extracellular nucleotide-derived phosphate hydrolysis, demonstrating the existence and function of E-NPPs in budding yeast. Their levels of protein expression and enzymatic activity are enhanced during phosphate starvation. We demonstrate that nucleotide phosphate hydrolysis can also be performed by repressible Pho5p activity. Together, Npp1p, Npp2p, and Pho5p are the major contributors to NPPase activity. All three components are likely to work in conjunction with the high-affinity phosphate transporters to import scavenged phosphates once hydrolyzed from extracellular nucleotides. Regulation of extracellular nucleotide phosphatase activity in S. cerevisiae is distinct from that of multicellular eukaryotes, where E-NPPs and the PHO response share overlapping roles in yeast for phosphate acquisition and may implicate shared functions in extracellular nucleotide regulation.

### MATERIALS AND METHODS

**Reagents.** Adenosine-[α-32P]triphosphate, adenosine-[γ-32P]triphosphate, adenosine-[32P]monophosphate, and [2-3H]adenosine were obtained from Amersham Biosciences. Lactate dehydrogenase, phosphoethanolamine, NADH, morpholinepropanesulfonic acid (MOPS), MgCl2, Tris, and MgSO4 were obtained from Sigma Aldrich. Concentrated aqueous ammonia was purchased from EMD Chemicals. Pyruvate kinase was obtained from U.S. Biologicals. 4-Nitrophenyl phosphate was obtained from Fluka. Bacto-yeast extract and Bacto-peptone were purchased from Becton Dickinson.

**Strains and media.** The S. cerevisiae strains used in this study were derived from the Saccharomyces Genome Deletion Project (46) and are described in Table 1. All strains used in this study had comparable growth rates over a 24-h growth period. The strains were grown at 25°C unless otherwise specified, and a 24-h growth period, cultures were pelleted, washed three times with YPD, and resuspended in YPD. Each time point correlates with distinct stages of log-phase growth, where 8 h corresponds to early log phase ($OD_{600} = 0.2$ to 0.4), 16 h corresponds to mid-log phase ($OD_{600} = 0.6$ to 0.8), and 24 h corresponds to late log phase ($OD_{600} = 1.0$ to 1.3). The cells were broken with glass beads, and the supernatant was collected. The radioactivity of each of the fractions was determined by liquid scintillation counting (Beckman Coulter).

**Sequence homology for potential E-NPPs.** BLAST, PSI-BLAST, and RPS-BLAST searches were performed where each member of the human E-NPP family was independently queried against the S. cerevisiae genome. From these searches, two yeast genes, NPP1 (YCR026c) and NPP2 (YEL016c), were identified. Additionally, the active site amino acid sequences (see Fig. 2, where 17 residues are highlighted) from known human E-NPPs were queried by BLAST searches against the S. cerevisiae genome, and the same two genes were identified. Both genes are uncharacterized, hypothetical open reading frames with predicted type I phosphodiesterase/nucleotide pyrophosphatase motifs as determined by the Conserved Domains Database (26). These genes were also used to perform BLAST searches against the yeast Schizosaccharomyces pombe. Two putative, uncharacterized open reading frames were identified in the search that were also predicted to have type I phosphodiesterase/nucleotide pyrophosphatase motifs. Protein sequences for open reading frames were retrieved from GenBank (see Fig. 2); S. cerevisiae NPP1, AAC79270; S. cerevisiae NPP2, P25353; Homo sapiens NPP1, AAF63094; H. sapiens NPP2, BAA08260; H. sapiens NPP3, AAC75163; H. sapiens NPP4, BAA74902; H. sapiens NPP5, CAB56566; Rattus norvegicus NPP1, AF320854; R. norvegicus NPP3, BAA63063; and low-pvirus NPP, AAF43474.

**Assay of NPP activity.** Strains expressing tandem affinity purification (TAP)-tagged fusion proteins of Npp1p and Npp2p were grown overnight in YPD. Cells were lysed in 20 mM Tris-HCl (pH 7.5), 0.5 mM phenylmethylsulfonyl fluoride, 0.5 mM benzamide, 0.3 mM NaCl, and 0.2% Triton X-100. After centrifugation, the TAP tag fusion proteins were immunoprecipitated from the supernatant with TAP tag monoclonal antibodies (Open Biosystems) and protein A-agarose (Agarose (U. state)). Immunoprecipitates were washed five times, resuspended in 50 mM HEPES (pH 7.5), and assayed for enzymatic activity. Activity was measured at 30°C in 50 mM Tris HCl (pH 9.0) with 0.9 mM p-nitrophenyl dTMP as a substrate. The reaction was stopped by 10-fold dilution in 5% trichloroacetic acid (Sigma Aldrich). The amount of p-nitrophenyl formed was quantified at λ = 410 nm after the addition of a 0.1 volume equivalent of 5 M NaOH. One unit of NPPase activity was defined as the amount of enzyme required to hydrolyze 1 μmol of substrate/min under the assay conditions used.

**Isolation of the NPP-nucleotideylated intermediate.** The trapped intermediate was performed according to Blytt et al. (4) with minor changes (42). Purified Npp1p or Npp2p (1 U/ml) was incubated in 0.57 M imidazole-formate, pH 4.0, and 260 μM [α-32P]ATP. At specific time points, 0.3-mg/ml bovine serum albumin and 10% trichloroacetic acid were added. After 20 min on ice, the precipitated proteins were pelleted by centrifugation and washed three times in 50 mM Tris HCl, pH 6.8. The pellet was then boiled in 25 mM Tris HCl (pH 6.8), 2% sodium dodecyl sulfate (SDS), 1% glyceral, and 250 mM 2-mercaptoethanol (electrophoresis buffer). The trapped intermediates were visualized by autorad.
diography following SDS-polyacrylamide gel electrophoresis (PAGE) (7.5%). As a negative control, identical 60-min reactions were set up using [γ-32P]ATP as a substrate to verify that no radiolabeled trapped intermediates were generated under these conditions.

Isolation of the NPP phosphorylated intermediate. Phosphorylation was performed by incubating of 200-mU/ml of Npp1p or Npp2p in 50 mM HEPES, pH 7.5, with 50 μM [γ-32P]ATP. At specific time points, the reaction was stopped by the addition of electrophoresis buffer. The phosphorylated intermediates were visualized by autoradiography following SDS-PAGE (7.5%). As a negative control, an identical 60-min reaction containing [α-32P]ATP was run to verify that phosphorylated intermediates were not generated under these conditions.

TAP-tagged protein expression and Western blot immunodetection. Strains with chromosomally expressed genes of interest with either TAP fusions for Npp1p and Npp2p or green fluorescent protein (GFP) fusions for Pho5p were obtained (Open Biosystems). Cells from stationary-growth cultures were used to inoculate cultures in either YPD or low-phosphate medium with an initial OD600 of 0.05. Cultures were grown for various time points and normalized for 5 OD600 equivalents to be prepared per lysate. Cell lysates were prepared by bead beating in 1% SDS, 8 M urea, 10 mM MOPS, pH 6.8, and 10 mM EDTA. Normalized total cell lysates were loaded per lane on an 8% PAGE gel, electrophoresed, and transferred to a nitrocellulose membrane. Anti-TAP antibody or anti-GFP antibody was used as a primary antibody (Open Biosystems and Sigma, respectively). Horseradish peroxidase-conjugated antibodies against rabbit immunoglobulin G (Santa Cruz Biotechnology) were used as secondary antibodies. For protein loading controls, mouse anti-414 antibody (obtained from D. Forbes) was used to detect a constitutively expressed nuclear protein, Nap62p. Western blotting was performed using the ECL detection system protocol (Perkin-Elmer).

NPPase activity assays with ATP substrates. Cells grown to late log phase were used to inoculate either YPD or modified medium at an initial OD600 of 0.05. Cultures were grown for the times indicated, pelleted, washed three times with Tris-EDTA, and resuspended in 0.5 ml 50 mM Tris-HCl, pH 8.0. A kinetic activity assay was adapted to spectrophotometrically measure activity with ATP as a substrate (9). Cells were then combined with a reaction mixture containing 100 mM MOPS, pH 8.0, 10 mM MgCl2, 1 mM phosphoenol pyruvate, 12 U lactate dehydrogenase, 15 U pyruvate kinase, and 140 μM NADH. The absorbance was zeroed at λ = 340 nm, and then 10 μl of 100 mM ATP was added. ATP hydrolysis to ADP was monitored as a function of NADH oxidation over 1-min elapsed reaction time by measuring the decrease in absorbance at 340 nm, and then 10 μl of 100 mM ATP was added. ATP hydrolysis to ADP was monitored as a function of NADH oxidation over 1-min elapsed reaction time by measuring the decease in absorbance at 340 nm (Beckman-Coulter DU 640). NADH consumption (in micromoles per minute) was determined by the equation [\( \frac{A_{340} - A_{340} - 1 \text{ min}}{A_{340} - A_{340} - 1 \text{ min}} \) min].

RESULTS

Cellular uptake of extracellular nucleotide-derived phosphates is dependent on extracellular phosphate concentration. Ecto-nucleotide phosphatase activity had yet not been established to exist in yeast. To ascertain whether this activity was present in budding yeast, extracellular ATP hydrolysis was measured. If E-NPP activity occurs, hydrolyzed phosphates will be imported during cell growth to maintain proper intracellular phosphate concentrations. Cultures were grown in the presence of radiolabeled nucleotide substrates, and the level of imported radioactivity was monitored over time. This was measured in both low- and high-phosphate medium from early- to late-log-phase growth (Fig. 1). Nutrient-enriched YPD medium was utilized as high-phosphate conditions, whereas YPD medium with depleted inorganic phosphate provided low-phosphate conditions. Cells were grown in both conditions in the presence of 1 μCi [γ-32p]ATP. At each time point, cells were lysed and the amount of imported γ-phosphate was determined by comparison to that remaining in the medium. When grown in high-phosphate medium, <10% of the radioactivity was detected within the cellular lysate. However, when phosphate was depleted from the growth medium, >95% of the radiolabeled phosphate was detected in the cellular lysate. To demonstrate that the observed hydrolysis is a regulated cellular event, NPPase activity was monitored over time under normal and low-phosphate conditions, illustrating induced NPPase activity during phosphate starvation. Cells were grown for 24 h in the presence of radiolabeled compounds prior to uptake analysis. Tritiated adenosine uptake was evaluated as a control to establish that phosphates were hydrolyzed from the nucleotide base prior to cellular import.

Identification of two putative E-NPP genes, NPP1 and NPP2. In several eukaryotes, the E-NPP multigene family controls a complex mechanism of extracellular nucleotide regulation and metabolism. Despite their diversity, the family members share many physiological roles involving nucleotide phosphate hydrolysis. As demonstrated above, extracellular nucleotide-derived phosphate hydrolysis occurs in S. cerevisiae, yet the enzyme(s) involved in this activity has not been previously identified. Based upon sequence comparisons to this family of enzymes, two candidates for genes encoding E-NPPs were identified in S. cerevisiae, neither of which is individually required for viability. These genes share approximately 30% similarity and 15 to 20% identity to the E-NPP family within the
type I phosphodiesterase-nucleotide pyrophosphatase motif and have BLAST scores with E-values as significant as $10^{-79}$ compared to human homologs with demonstrated E-NPP activity (Fig. 2). Notably, in regions containing catalytically significant residues including the active site, nearly all residues were identical. Based on their sequence similarities and functional characterization (described below), we have named these genes \textit{NPP1} (\textit{YCR026c}) and \textit{NPP2} (\textit{YEL016c}), in accordance with the Saccharomyces Genome Database naming guidelines (7).

To ascertain whether these genes encode functional NPPs in yeast, Npp1p and Npp2p were purified and assayed with the same tests used for characterization of human NPPs. The primary trademark of the NPP family is to catalyze phosphate hydrolysis from nucleotide triphosphates, producing an intermediate where the nucleotide is covalently attached to the catalytic threonine residue during catalysis. Autoinhibition states were also monitored over 60 min as an indicator of autoinhibition. Npp2p was autoprophosphorylated at the earliest time point and persisted throughout the time course.
catalytic threonine (12). In the second step of this reaction, the nucleotide monophosphate is released. Previous characterization of other members of the NPP family illustrated that the nucleotidylated catalytic intermediate could be trapped under acidic conditions in the presence of imidazole using the substrate \([\alpha^{32}\text{P}]\text{ATP}\). Imidazole inhibits substrate release after nucleotide phosphate hydrolysis, leading to stabilization of the intermediate state. Npp1p and Npp2p were isolated by immunoprecipitation and verified by Western blotting analysis (Fig. 2C). The semipurified proteins were tested to determine whether they could perform the NPP reaction mechanism. Under these reaction conditions, the catalytic activity of Npp1p and Npp2p were measured over a 60-min time course (Fig. 2C). For both enzymes, the nucleotidylated intermediates were observed. As a negative control, the same experimental setup using \([\gamma^{32}\text{P}]\text{ATP}\) was performed to verify that nonspecific labeling did not occur under these conditions (data not shown). The difference in the kinetics of each intermediate trapping profile demonstrates that each enzyme has unique activity, and may be differentially regulated. A possible explanation for this observation is through autoinhibition of catalytic activity.

Some members of the NPP family can undergo autophosphorylation and autodephosphorylation of the catalytic threonine as a mechanism of catalytic regulation (42). To establish whether these two enzymes could undergo autophosphorylation, the phosphorylation state of each enzyme was measured over time (Fig. 2C). Npp1p and Npp2p were able to undergo autophosphorylation. Distinct profiles for autophosphorylation were observed, suggesting that their catalytic activities are differentially autoregulated. Both enzymes demonstrated the characteristic qualities of NPPases, signifying that NPPs are conserved in yeast.

**NPP1 and NPP2 are important for cellular uptake of extracellular nucleotide-derived phosphates.** To determine if these genes were responsible for the observed extracellular nucleotide phosphate hydrolysis, null mutants were tested for their ability to perform hydrolysis under low-phosphate conditions. Experiments were conducted parallel to those in Fig. 1 over a 24-h time course monitoring hydrolysis at early-, mid-, and late-log-phase growth. Using both \([\gamma^{32}\text{P}]\text{ATP}\) and \([\alpha^{32}\text{P}]\text{ATP}\), the mutant strains were assayed for levels of imported radiolabeled phosphate (Fig. 3A). When \(\gamma^{32}\text{P}\) uptake was monitored, strains lacking either NPP1 or NPP2 could import only 50% of the radiolabeled phosphate by 16 h. In the double mutant, the defect in phosphate hydrolysis was slightly more pronounced. These defects are in stark comparison to the wild type, which could import nearly the entire radiolabeled probe by this time point. Defects in these mutant strains were also seen when \(\alpha^{32}\text{P}\) uptake was measured, which correlates with complete phosphate hydrolysis from the nucleotide moiety (Fig. 3B). While nearly 60% of the radiolabel was imported in the npp2Δ mutant, only 5% of the radiolabel was imported in the npp1Δ mutant at 16 h. The decreased rate of \(\alpha\)-phosphate hydrolysis may simply be due to catalytic rates of hydrolysis, where more Npp2p was found to be in an autoinhibited, phosphorylated state than Npp1p. Yet this notable result also demonstrates that NPP1 may preferentially utilize substrates at the \(\alpha\)-phosphate position to perform phosphodiester hydrolysis. The hydrolysis defect was comparable in a double null mutant where about 10% of the radiolabel was detected in the cellular lysate. However, by 24 h, nearly all of the \(\alpha^{32}\text{P}\) and \(\gamma^{32}\text{P}\) could be imported in all three mutant strains, indicating that the absence of activity from these genes may be compensated from another source. Accordingly, Npp1p and Npp2p demonstrate conserved E-NPP activity in yeast and partially contribute to nucleotide-derived phosphate hydrolysis during phosphate starvation.

**E-NPPs can hydrolyze nucleotide-derived phosphates, particularly during phosphate starvation.** The absence of either or both genes encoding E-NPPs caused a substantial defect in measured phosphate uptake, indicating that both gene products can perform extracellular nucleotide hydrolysis. To establish that these gene products are directly utilizing nucleotide substrates for phosphate hydrolysis, mutant strains lacking NPP1 and/or NPP2 were tested for their ability to hydrolyze nucleotide phosphates in a direct NPPase assay using ATP as a substrate in a coupled enzymatic assay where ATP hydrolysis was detected as a function of NADH oxidation (Fig. 4A). Under normal phosphate conditions, a low level of NPPase...
activity was present in all strains at levels comparable to that of a wild-type strain. When grown under low-phosphate conditions, NPPase activity in the wild-type strain was greatly enhanced by 16 and 24 h. Strains lacking either NPP1 or NPP2 had no increased NPPase activity by 16 h but had a minor increase in activity by 24 h, although activity was <50% of that in the wild type. The double mutant was slightly worse, where NPPase activity levels were approximately 30% of the wild type. Thus, Npp1p and Npp2p can hydrolyze extracellular phosphate from ATP, and this activity is greatly enhanced when extracellular phosphate concentrations are low.

Since activity depends on phosphate starvation conditions, protein expression levels of Npp1p and Npp2p were monitored over a 24-h time course from early to late log phase under both normal and low-phosphate growth conditions (Fig. 4B). At each time point, cell lysates were prepared, and expression levels of TAP-tagged Npp1p and Npp2p were detected by immunoblotting. Protein expression levels were nearly undetectable at 0 h (stationary-growth phase), but levels were increased during active log-phase growth. In correlation with enhanced enzymatic activity, protein expression levels for both proteins were significantly augmented under low-phosphate conditions, particularly for Npp2p at later time points. These protein expression profiles may implicate Npp1p as an early responder for NPPase activity, and Npp2p may subsequently assist in sustaining the response over time by delayed expression. This may also demonstrate that under low-phosphate conditions, Npp1p and Npp2p play complementary roles for regulation of extracellular nucleotide phosphate hydrolysis during phosphate starvation.

Pho5p, a secreted acid phosphatase, partially mediates nucleotide-derived phosphate hydrolysis. While Npp1p and Npp2p appear to be highly activated during phosphate starvation, the possibility that an element of the PHO response may be involved in extracellular nucleotide phosphate hydrolysis was also investigated. The prospect seemed likely, since this activity is strongly phosphate dependent and only a 50% reduction in NPPase activity was seen in the absence of NPP1 and NPP2. When the PHO response system is activated, the repressible acid phosphatase encoded by PHO5 is upregulated and plays a major role in extracellular phosphate scavenging (47). Although two other acid phosphatases are also upregulated during this response (Pho11p and Pho12p), >90% of the detectable phosphatase activity was due to Pho5p. To determine if PHOS contributes to phosphate hydrolysis of extracellular nucleotides, a pho5Δ mutant was assayed for its ability to internalize phosphate. Both [α-32P]- and [γ-32P]ATP were tested as potential substrates (Fig. 5A and B). After 16 h of growth, the wild-type cells imported nearly all of the radiolabeled phosphate. However, the pho5Δ strain imported 60% of the γ-32P and only 20% of the α-32P at the same time point. By 24 h, the pho5Δ mutant approached the wild-type level of imported γ-32P but still lagged in importing α-32P. Although nucleotide phosphate hydrolysis still occurred in the pho5Δ mutant, the absence of Pho5p caused a significant decrease in the rate of hydrolysis. The defect was most pronounced in the instance with AMP as a substrate. In addition to Npp1p and Npp2p, Pho5p appears to play a significant role in phosphate-mediated uptake from extracellular nucleotide-derived phosphoryl substrates, and this function is distinct from its previously characterized activities.

Pho5p can utilize nucleotide-derived phosphates as substrates. Since the absence of PHOS caused a notable reduction in nucleotide-derived phosphate cleavage, this raised the possibility that Pho5p may also directly utilize extracellular ATP as a substrate. This model was tested using a coupled enzyme assay to monitor ATP hydrolysis (Fig. 5C). Both wild-type and pho5Δ strains were grown in low-phosphate medium, and APase activity was determined as a function of time. As cultures reached mid- and late-log-phase growth, the rate of ATP hydrolysis was significantly upregulated in the wild type. The pho5Δ mutant demonstrated no upregulation by early- or mid-log-phase growth. Even by late-log-phase growth, the level of ATP hydrolysis by the pho5Δ mutant only slightly increased, and this level of defective hydrolysis was comparable to the rate seen with the npp1Δ npp2Δ double mutant. The defect in the pho5Δ mutant strongly indicates that Pho5p plays a signif-
The observed loss of nucleotide phosphate hydrolysis in the absence of PHO5 should correlate with enhanced enzymatic activity. To establish that Pho5p expression is increased under these growth conditions, Pho5p expression levels were monitored over a 24-h time course under high- and low-phosphate conditions (Fig. 5D). Protein levels increased during log-phase growth and were enhanced under phosphate starvation conditions. Pho5p demonstrated the greatest increase of expression by 8 h, indicating that Pho5p is rapidly expressed alongside Npp1p. Thus, Pho5p and Npp1p may perform some redundancy in performing early NPPase activity, and this precedes the time-delayed response of Npp2p. This evidence links the increased nucleotide phosphate hydrolysis activity demonstrated in a wild-type strain through primary enhanced expression levels of Npp1p and Pho5p, followed by the secondary enhancement of Npp2p, to collectively perform NPPase activity during phosphate starvation.

NPP1, NPP2, and PHO5 are principal components of NPPase activity in S. cerevisiae. Three separate genes from two distinct gene classes were identified to perform NPP activity in yeast. All three genes appear to play a large role in overall activity, although their function appears to be redundant. It remains unknown whether there are any other genes that can additionally contribute to NPP activity and whether NPP activity is even a necessary cellular event. To address these questions, a strain lacking NPP1, NPP2, and PHO5 was generated. The strain was used to determine the extent of defectiveness in extracellular nucleotide phosphate hydrolysis. The rate of imported \(^{32}\text{P}\) from extracellular ATP was measured over 24 h (Fig. 6A). By the end of the time course, the triple-null strain could intracellularly import only about half of the total \(^{32}\text{P}\), compared to nearly 100% in either the npp1npp2 or pho5 double mutants. To determine whether the triple-null strain could fully dephosphorylate all three phosphate groups from ATP, \(^{32}\text{P}\) ATP was provided extracellularly. Phosphate hydrolysis and import were subsequently measured under low-phosphate conditions (Fig. 6B). In this case, only about 20% of the total \(^{32}\text{P}\) was hydrolyzed by 24 h. This is in stark contrast to the npp1npp2 or pho5 strains, where hydrolysis reached nearly 100% or 70%, respectively, throughout the same growth period. This clearly demonstrates that all three genes are major components of the NPPase activity.

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The NPPase activity of the triple-null strain was further tested by directly monitoring ATP hydrolysis over time (Fig. 6C). Using the coupled enzyme assay, ATP hydrolysis was measured at various time points after cells were grown in either YPD or phosphate-depleted medium. At each time point, the triple-null strain demonstrated diminished NPPase activity and was slightly more defective than the npp1Δ npp2Δ or pho5Δ strains alone. The level of NPPase activity measured was comparable to basal activity demonstrated under normal YPD growth conditions. This demonstrates that NPP1, NPP2, and PHO5 are collectively the major contributors to NPPase activity in yeast and interconnects the possibility that NPPase activity and the PHO response may share overlapping regulatory roles in S. cerevisiae.

FIG. 6. NPP1, NPP2, and PHO5 are the major contributors for NPPase activity. (A) Extracellular phosphate hydrolysis by the triple-null strain was monitored through measuring phosphate import over 24 h. Extracellular [γ-32P]ATP was utilized as a substrate. At the end of the time course, a 50% defect in phosphate uptake was measured in the triple-null strain. (B) Extracellular [α-32P]ATP was utilized to test phosphate hydrolysis at the proximal position. The absence of all three genes caused a nearly complete loss of phosphate hydrolysis at the α-position of ATP. (C) NPPase activity was measured in the triple-null strain using the coupled enzyme assay with ATP as a substrate. The triple-null strain demonstrated nominal NPPase activity throughout the entire time course.

DISCUSSION

Yeast homologs of the E-NPP family can hydrolyze nucleotide phosphates. In many eukaryotes, E-NPPs are directly responsible for phosphate hydrolysis from extracellular nucleotides. Based on sequence criteria, we identified two candidate E-NPPs in the S. cerevisiae genome. They were both evaluated for their roles during phosphate starvation, and both demonstrated the capability of performing extracellular phosphate hydrolysis from nucleotide substrates. We have demonstrated that both Npp1p and Npp2p are functional nucleotide pyrophosphatase/phosphodiesterases. Both enzymes catalyzed nucleotide phosphate hydrolysis to produce the trapped nucleotide monophosphate intermediate. These enzymes can subsequently catalyze hydrolysis of the monophosphate to produce the nucleotide base. Additionally, this is the first demonstration that NPP1 and NPP2 are upregulated via phosphate-regulated transcription. Distinct profiles of catalytic regulation for each enzyme parallels the functional diversity seen among NPPs specialized for particular functions in humans. This is the first documentation of E-NPP activity in yeast and highlights the notion that functional diversity is conserved among the NPP gene family.

E-NPPs may mediate intercellular signaling. In humans, the fundamental role of the E-NPP family is to regulate extracellular nucleotide metabolism and nucleotide-derived intercellular signaling. While budding yeast is a unicellular organism, intercellular signaling has been documented to coordinate events such as aggregation, differentiation, sexual reproduction, and sporulation (18). During sporulation, differentiating yeast cells cooperate to maintain high concentrations of extracellular purines and produce adenosine 5′-tetraphosphate and adenosine 5′-pentaphosphate to mark ascospore formation (17). Intriguingly, NPP2 expression is strongly induced during sporulation (8). Taken together, this may implicate a novel intercellular signaling role for E-NPPs by regulating extracellular nucleotide metabolism to mediate sporulation and possibly other intercellular signaling events.

Yeast can derive essential phosphate from extracellular nucleotides. Importing phosphate from extracellular sources during periods of phosphate starvation can be critical for cellular viability. If extracellular phosphate is scarce, sequestering phosphate from diverse chemical substrates would be advantageous for survival. Enzymes responsible for obtaining phosphates from polyinorganic phosphate chains and/or pyrophosphates have been identified in yeast (15, 23, 25, 29, 30, 39). In prin-
ciple, hydrolysis of phosphates from nucleotide sources should provide an additional nutrient source. The import of phosphate from extracellular nucleotide substrates demonstrates the diversity of substrates utilized to overcome phosphate starvation. This finding is also compelling in that it unifies E-NPP regulation with the phosphate starvation response. The PHO response is a powerful mechanism in many unicellular organisms that allows them to adapt to an ever-changing environment. It is intriguing that the repressible expression of several genes that specialize in acquisition and import of available phosphates can bridge its activity with extracellular nucleotide regulation.

A new function for Pho5p as a nucleotide phosphatase. Since extracellular nucleotide-derived phosphate hydrolysis was significantly upregulated during periods of phosphate starvation, several components of the PHO response were investigated for their contributions. When the PHO response is activated, the secreted acid phosphatase Pho5p is the most active acid phosphatase for scavenging and hydrolyzing phosphates. Our results demonstrate that nucleotide phosphate hydrolysis performed by Pho5p is intertwined with the PHO response during phosphate starvation. It is noteworthy that another component of the PHO response is high-affinity phosphate uptake. In conjunction with Pho5p, phosphate transporters are critical components for generating and importing extracellular free inorganic phosphate to maintain viability during periods of starvation. Additionally, nucleotide dephosphorylation has profound effects in cellular regulation of many mammalian systems. Furthermore, the fact that this activity coincides with the PHO response system may implicate an integrated scheme of regulation during phosphate starvation through multiple modes of cellular regulation.

A model of nucleotide-derived phosphate acquisition. Under high-phosphate conditions, low levels of nucleotide phosphates are hydrolyzed by constitutive E-NPP activity composed primarily of NPP1, NPP2, and PHO5. Such basal levels of inorganic phosphate stimuli can be detected by phosphate transporters to define an adequate level of available inorganic phosphate for consumption by various biological processes (3). If extracellular phosphate becomes depleted, then phosphate starvation is sensed. This leads to the upregulation of a robust, multicomponent system of proteins known as the PHO response. This response system involves dual upregulation of phosphatases to generate free inorganic phosphate and phosphate symporters to import the generated inorganic phosphate. Our results provide the first evidence that nucleotide-derived phosphates can be hydrolyzed extracellularly in budding yeast by Npp1p, Npp2p, and Pho5p during phosphate starvation. Once free inorganic phosphate has been generated extracellularly, the repressible phosphate symporters Pho84p and Pho89p are presumably the major contributors to phosphate import to maintain cellular viability (Fig. 7).

In summary, our results establish that the E-NPP family is conserved in yeast and can perform extracellular phosphate hydrolysis. Extracellular nucleotides, particularly adenosine, have been found to play many related regulatory roles in cell signaling in multicellular organisms, including platelet aggregation (36), differentiation (20), cell proliferation (11), and apoptosis (16, 37). NPP1 and NPP2 remain conserved in S. cerevisiae and possibly in several other diverse fungal species, raising the possibility that these genes may also perform critical biological roles in yeast. We also establish a new role for Pho5p in sharing overlapping activity with the E-NPP family. These studies establish a mechanism for extracellular nucleotide regulation that is distinct from multicellular organisms by uniting its function with the phosphate starvation response. It will be important to investigate whether the regulation and biological roles of extracellular nucleotides by the E-NPP family are conserved among unicellular and multicellular organisms or identify what divergent roles have been adapted over evolution. It will also be exciting to establish whether the PHO response plays a larger regulatory role in extracellular nucleotide regulation and further to ascertain how these distinct regulatory mechanisms complement the function of the other.

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