A Transcription Factor Cascade Involving Fep1 and the CCAAT-Binding Factor Php4 Regulates Gene Expression in Response to Iron Deficiency in the Fission Yeast *Schizosaccharomyces pombe*

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We have identified genes encoding candidate proteins involved in iron storage (pel1+), the tricarboxylic acid cycle (sdh4+), and iron-sulfur cluster assembly (isa1+) that are negatively regulated in response to iron deprivation. Promoter deletion and site-directed mutagenesis permitted identification of a new cis-regulatory element in the promoter region of the pel1+ gene. This cis-acting regulatory sequence containing the pentanucleotide sequence CCAAT is responsible for transcriptional repression of pel1+ under low iron supply conditions. In *Schizosaccharomyces pombe*, the CCAAT-binding factor is a heteromeric DNA-binding complex that contains three subunits, designated Php2, Php3, and Php5. Inactivation of the php2+ locus negatively affects the transcriptional competency of pel1+. A fourth subunit, designated Php4, is not essential for the transcriptional activation of target genes under basal and iron-replete conditions. We demonstrate that, in response to iron-limiting conditions, Php4 is required for down-regulation of pel1+, sdh4+, and isa1+ mRNA levels. In vivo RNase protection studies reveal that the expression of php4+ is negatively regulated by iron and that this regulated expression requires a functional fep1+ gene. The results of these studies reveal that Fep1 represses php4+ expression in response to iron. In contrast, when iron is scarce, Fep1 becomes inactive and php4+ is expressed to act as a regulatory subunit of the CCAAT-binding factor that is required to block pel1+, sdh4+, and isa1+ gene transcription.

Iron is a biological constituent required by most organisms (22). As a cofactor in a wide range of biochemical reactions, iron is important (46). For example, iron is present in the active site of enzymes involved in metabolic processes such as the tricarboxylic acid (TCA) cycle, the cellular respiratory chain, oxygen transport, and photosynthesis. Iron’s value resides in the reactivity of the Fe2+/Fe3+ redox couple, which enables it to donate or accept an electron to catalyze enzymatic reactions. Paradoxically, the same redox-active nature of iron renders it biotoxic. This is due to its ability to react with oxygen species to generate the hydroxyl radical (19). Production of this free reactive oxygen species can damage cellular components such as DNA, proteins, and membrane lipids (20). Therefore, organisms need efficient mechanisms to acquire iron but also to keep its reactivity in check.

Iron presents an additional problem in that it is found under aerobic conditions as insoluble ferric hydroxides, making it biologically unavailable (10, 14). Consequently, organisms have evolved different iron-scavenging systems for solubilizing iron and transporting it into cells. Many bacteria and fungi synthesize and excrete low-molecular-weight iron chelators known as siderophores (15, 18, 24, 58). Subsequent to excretion of the siderophores, iron-siderophore complexes are efficiently recaptured by microorganisms via specific cell surface receptors (28, 46). Other iron assimilation mechanisms include reduction via a cell surface reductase and subsequent transport of the iron across the membrane by iron uptake proteins (2, 11, 12, 17, 29, 32, 54), proteolytic degradation of the host iron-binding proteins (49, 53), and surface receptors for mammalian iron carriers such as transferrin, lactoferrin, and heme (20).

In the model organism *Schizosaccharomyces pombe*, two types of iron uptake systems have been found (3, 48, 52). The first system consists of two genes, *sib1* (SPAC23G3.02c) and *sib2* (SPAC23G3.03), which are involved in the acquisition of iron from the hydroxamate-type siderophore ferrichrome (52). The ability of fission yeast cells to produce and excrete ferrichrome is consistent with the existence of *S. pombe* Str1, which confers ferrichrome uptake when ectopically expressed in a *Saccharomyces cerevisiae* fet3Δ arm1-4Δ mutant strain (44). Two other proteins found in the *S. pombe* proteome exhibit sequence similarity to Str1. These two homologs were designated Str2 and Str3 (44). Although Str2 and Str3 may participate in the mobilization of iron bound to siderophores, their substrate specificity has not been determined. Under low environmental iron conditions, a second system is used by *S. pombe* to accumulate iron. It is divided into multiple steps. A first step is the reduction of ferric iron by a cell surface electron transporter encoded by the *frp1*+ gene (48). Then, subsequent to reduction, ferrous iron is captured by an oxidase-permease complex formed by Fio1 and Fip1 (3). Fio1 converts ferrous iron to ferric iron, which is then transported across the membrane by the iron permease Fip1 (3).

Under iron-deficient conditions, the extent of transcription of *frp1*+, *fio1*+, *fip1*+, *str1*+, *str2*+, and *str3*+ increases several times (~8- to 18-fold), whereas when environmental iron is abundant, expression of these genes is extinguished (3, 43, 44, 89).
Conserved cis-acting elements in the fpr1+, fio1+, fip1+, str1+, str2+, and str3+ promoters with the sequence 5′-(A/T)GATAA-3′ are required for iron-mediated repression of gene expression (43, 44). Fep1 is a GATA transcription factor that has been shown to negatively regulate iron uptake by repressing fpr1+, fio1+, fip1+, str1+, str2+, and str3+ gene expression under iron-replete conditions (43, 44). Deletion of fep1+ (fep1Δ) results in constitutive expression of these iron transport genes. Based on the ability of the iron chelator BPS to preclude its binding, Fep1 has been shown to associate with DNA in an iron-dependent manner (43). The N-terminal 241 amino acids of Fep1 contain its DNA-binding domain. At its C terminus, Fep1 contains a leucine zipper domain that mediates Fep1 dimerization and increases its potency as a transcriptional repressor (45).

In the fission yeast S. pombe, following glucose depletion, a protein complex comprising the Php2, Php3, and Php5 proteins positively regulates many of the genes involved in oxidative phosphorylation, including cycl1+, which encodes the cyochrome c oxidase protein (38, 42, 60). The S. pombe Php2, Php3, and Php5 proteins are orthologous to Saccharomyces cerevisiae Hap2, Hap3, and Hap5, respectively (37, 39). Consistently, expression of S. pombe php2+, php3+, and php5+ functionally complements S. cerevisiae hap2Δ, hap3Δ, and hap5Δ mutants, respectively, for growth on a nonfermentable carbon source (38, 41, 60). The S. pombe Php2, Php3, and Php5 proteins contain short regions of homology with their S. cerevisiae counterparts that correspond to evolutionarily conserved core domains (38). In S. cerevisiae, the DNA-binding capability of the CCAAT-binding complex is conferred by the Hap2/Hap3/Hap5 subunits (39). In addition, a fourth subunit, denoted Hap4, enhances the Hap2/Hap3/Hap5 complex activity via its trans-activation domain (16). The S. cerevisiae heteromeric CCAAT-binding complex is modulated via the expression of Hap4, which is repressed in the presence of glucose and induced in media containing nonfermentable carbon sources such as lactate, ethanol, and glycerol. This suggests that Hap4 is a key regulator of CCAAT complex activity in response to carbon source in S. cerevisiae. Until recently, Hap4 homologs had not been identified in other fungi. A recent in silico study has revealed that such homologs exist in the genomes of Saccharomyces species closely related to S. cerevisiae, as well as two more distantly related Saccharomyces species (55). Furthermore, functional complementation studies revealed that Hap4 homologs exist in Kluyveromyces lactis (8) and Hansenula polymorpha (56).

The strict requirement for appropriate intracellular iron concentrations suggests that there may be regulatory mechanisms that prevent futile expression of the genes encoding iron-requiring proteins when iron is limiting. In Escherichia coli, a small noncoding regulatory RNA, termed RyhB, is induced under conditions of iron starvation (35). It hybridizes with its target mRNAs and causes their degradation (34). RyhB targets encode iron storage and iron-containing proteins (36). Transcription of the ryhB gene is repressed by the iron-Fur complex (33). A similar process has been described for the action of two small RNAs, called prrF1 and prrF2, in the control of iron-regulatagable genes in Pseudomonas aeruginosa (57). In S. cerevisiae, the RNA-binding protein Cth2 has been shown to mediate the degradation of transcripts encoding iron-containing proteins in response to iron deprivation (47). Under iron-adequate conditions, CTH2 expression is down-regulated. In contrast, increased CTH2 gene expression occurs during iron scarcity and depends on the iron-responsive transcription factors Aft1 and Aft2 (47).

In this study, we have identified a set of genes (pcl1+, sdh4+, and isa1+) whose expression is down-regulated under iron starvation conditions. Analysis of regions in the pcl1+ promoter demonstrated that a CCAAT-type regulatory sequence is necessary for its transcriptional regulation. Consistently, inactivation of the php2+ locus that encodes a subunit of the heteromeric CCAAT-binding factor negatively affects the transcriptional competency of pcl1+. To examine the function of the CCAAT-binding factor in S. pombe, we cloned php4+ and generated a php4Δ mutant strain. We found that in contrast to its activator function in S. cerevisiae, the S. pombe Php4 factor appeared to function as a repressor of genes encoding iron- using and iron storage proteins. We determined that php4+ is regulated at the level of gene transcription; it is induced under conditions of iron deprivation and turned off under conditions of iron repletion. Furthermore, its iron-dependent regulated expression requires a functional fep1+ gene. Taken together, these results reveal the existence of a transcription factor cascade composed of Fep1 and the heteromeric CCAAT-binding factor to regulate gene expression in response to iron deficiency in fission yeast.

**MATERIALS AND METHODS**

**Strains and growth conditions.** S. pombe strains used in this work were the wild-type strain FY435 (b′ his7-366 leu1-32 ada6-d18 ade6-M1000 (6) and the pcl1Δ (isogenic to FY435 plus pcl1Δ::Kan), php2Δ (isogenic to FY435 plus php2Δ::Kan), php3Δ (isogenic to FY435 plus php3Δ::Kan), php4Δ (isogenic to FY435 plus php4Δ::Kan), php5Δ (isogenic to FY435 plus php5Δ::Kan), and zfs1Δ (isogenic to FY435 plus zfs1Δ::Kan) disruption strains. The gene disruptions were created by replacing the coding region of pcl1+, php2+, php3+, php4+, php5+, or zfs1+ with a loxP-KANMX-loxP cassette through homologous recombination (21). An isogenic fep1Δ disruption strain (b′ his7-366 leu1-32 ada6-d18 ade6-M1000 fep1Δ::ura4+) was also used as described previously (43). Yeast strains were cultivated at 30°C in yeast extract plus supplements or in selective Edinburgh minimal medium lacking specific nutrients required for plasmid selection (1). For iron starvation or iron repletion experiments, S. pombe strains were grown to mid-logarithmic phase (A600 of ~1.0) and then incubated for 90 min in the presence of 250 μM 2,2′-dipyridyl (Dip) or 100 μM FeCl3, respectively. After treatments, 15-ml samples were withdrawn from the cultures for subsequent steady-state mRNA analysis.

**Analysis of gene expression.** Total RNA was extracted by the hot phenol method as described previously (4). RNAs were quantified spectrophotometrically, and 15 μg of RNA per sample was used in the RNase protection protocol, which was carried out as described previously (5). 32P-labeled antisense pcl1-, sdh4-, isa1+, php2-, php3-, php5-, fio1+, and zfs1+ RNAs were produced from the BamHII-linearized plasmids pSKccl1+, pSKdhd4+, pSKa1+, pSKphp2+, pSKphp3+, pSKphp5+, pSKfio1+, and pSKzfs1+, respectively. 32P-labeled antisense lacZ RNA was generated from the HindIII-linearized plasmid pSKlacZ (26). The riboprobe derived from the plasmid pSKa1+ (25) was used to probe act1+ mRNA as an internal control for normalization during quantification of the RNA protection products.

**Plasmids.** Plasmid pSKccl1+ was created by inserting a 170-bp BamHI-EcoRI fragment from the pcl1+ gene into the same sites of pBlueScript SK (Stratagene, La Jolla, CA). The antisense RNA hybridizes to the region between +18 and +188 downstream from the initiator codon of pcl1+. Plasmid pSKdhd4+ was constructed by ligating a 178-bp BamHI-EcoRI fragment from the sdh4+ gene into the BamHI-EcoRI sites of pBlueScript SK. This fragment corresponds to the region between +3 and +181 down to the first base of the translational start codon of sdh4+. To generate pSKa1+, a 188-bp fragment from the isa1+ gene (corresponding to the coding region between +3 and +191) was amplified and cloned into the BamHI-EcoRI sites of pBlueScript SK. pSKphp2+ was made by
ligating a PCR product containing a 174-bp fragment from the php2\(^+\) gene (corresponding to the coding region between +51 and +225) into the BamHI-EcoRI sites of pBluescript SK. Plasmid pSkphp3\(^+\) was created by inserting a 183-bp BamHI-EcoRI fragment from the php2\(^+\) gene (matching to the coding region between +54 and +237) into the corresponding sites of pBluescript SK. Plasmid pSkphp4\(^+\) was constructed by inserting a 182-bp BamHI-EcoRI frag-
ment of the php4\(^+\) gene into the same sites of pBluescript SK. The antisense RNA pairs to the region between +536 and +518 downstream from the A of the start codon of php4\(^+\). pSkphp5\(^+\) was generated by ligating a 196-bp fragment from the php5\(^+\) gene into the BamHI-EcoRI sites of pBluescript SK. The latter plasmid was used to produce an antisense RNA probe that hybridizes specifically to php5\(^+\) mRNA (positions +307 to +503). Plasmids pSkchi1\(^+\) was generated by inserting a 184-bp fragment from the zfi1\(^+\) gene (corresponding to the coding region between +11 and +194) into pBluescript SK using the BamHI and EcoRI sites. Plasmids pSlaucZ, pSkXhol\(^-\), and pSacchi1\(^-\), for determining lacZ, fox1\(^+\), and act1\(^+\) mRNA levels, respectively, were described elsewhere (25, 43).

To generate the pSpIpc12l, a 2,472-bp Xhol-BamHI PCR-amplified DNA segment containing the S. pombe pell1 locus starting at +1060 from the translational start codon up to +683 after the stop codon was inserted into the Xhol and BamHI sites of pSP1 (9). The plasmid pSPIpc1l-1398lacZ was constructed by introducing a BamHI-Asp718 PCR-amplified fragment from the pell1\(^+\) promoter containing 1,398 bp of the 5' noncoding region and the first 10 codons of the pell1\(^+\) gene into the BamHI-Asp718-cut YEplac195 vector (40). Once generated, the pell1\(^+\) promoter region was isolated from YEplac195 and used for various experiments. The resulting plasmid, pell1\(^+\) promoter and the pell1\(^+\) promoter, was used to generate the equivalent DNA restriction fragment in pSPIpc1l-1398lacZ (43) (to generate pSPIpc1l-1398lacZ). Six plasmids (pSPIpc1l-1398lacZ, pSPIpc1l-1478lacZ, pSPIpc1l-1364lacZ, pSPIpc1l-1247lacZ, pSPIpc1l-1151lacZ, and pSPIpc1l-1089lacZ) were obtained. Each PCR product obtained was purified, digested with BamHI and XhoI, and used to replace the equivalent DNA restriction fragment in pSPIpc1l-1398lacZ. Plasmids pSPIpc1l-21lacZ and pSPIpc1l-21lacZ were used to introduce mutations in the CCAAT box (positions +265 to +291 with respect to the A of the ATG codon of pell1\(^+\)).

RESULTS

Preparation of S. pombe extracts and EMSAs. S. pombe cells were grown to an A\(_{600}\) of ~1.0 at 30°C. Cells were broken with glass beads in extraction buffer (200 mM Tris-HCl (pH 8.0), 400 mM (NH\(_4\))\(_2\)SO\(_4\), 10 mM MgCl\(_2\), 1 mM EDTA, 10% glycerol, 7 mM β-mercaptoethanol, 1 mM phenylmethylsulfonyl fluoride) in the presence of a protease inhibitor cocktail (P-8340; Sigma) followed by centrifugation at 10,000 rpm at 4°C for 5 min. Typically, binding reactions contained 35 to 40 μg of whole-cell extract in 1× binding buffer [50 mM Tris-HCl (pH 7.9), 100 mM (NH\(_4\))\(_2\)SO\(_4\), 2.5 mM MgCl\(_2\), 1.2 mM EDTA, 50 mM NaCl], 1 μg of poly(dI-dC), and ~1 ng of radiolabeled probe in a final reaction volume of 20 μl. Reactions were incubated at room temperature for 30 min, and the protein-DNA complexes were resolved by gel electrophoresis (5 h at 40 V) on 4% polyacrylamide gels (acylamide/ bisacylamide ratio, 37.5:1) in 0.5× Tris-borate at 4°C. After electrophoresis, the gels were fixed, dried, and subjected to PhosphorImager analysis. For reaction mixtures containing a purified recombinant Php2, the conditions were identical, except that the protein was present at ~200 ng. For reaction mixtures containing affinity-purified MBP-Fep1 (200 ng), expression of the MBP-Fep1 fusion protein in E. coli, preparation of extracts, and electrophoretic mobility shift assays (EMSAs) were carried out as described previously (45). Furthermore, purified proteins were analyzed by SDS-polyacrylamide gel electrophoresis and immunoblotting as described previously (43). Oligonucleotides 5′-AGCTTAAATCCCCTTGACATATTAAACCGACCCG-3′ or 5′-CGCGGATATCCGTTAGAAACGGCGCA-3′ were used to generate a CCAAT box of 8-mer, identical to the one present in S. pombe. Preceding studies have revealed that genes encoding components of the reductive (e.g., fpl1\(^+\), fox1\(^+\), and fpl1\(^-\)) and nonreductive (e.g., str2\(^+\), str2\(^-\), and str3\(^+\)) iron transport systems are induced by iron starvation and repressed by iron repletion (43, 44). However, little is known of the expression profiles of genes involved in storage and utilization of iron. To investigate the latter aspect, we characterized the regulation of genes encoding proteins that may play a role in iron detoxification (pell\(^+\)) in S. pombe that may play a role in iron detoxification (pell\(^+\)) in S. pombe that may play a role in iron detoxification (pell\(^+\)) in S. pombe.
These two regions encompass five putative transmembrane domains (called the DUF125 module). Pcl1 and Ccc1 are the only known proteins of *S. pombe* and *S. cerevisiae*, respectively, that belong to the DUF125 transmembrane protein family. Importantly, as observed for the intracellular iron transporter encoded by the *CCC1* gene in *S. cerevisiae*, deletion of the *pcl1* gene in *S. pombe* renders cells sensitive to iron compared with the wild-type strain (A. Mercier, B. Pelletier, and S. Labbé, unpublished data). Clearly, this result reveals that Pcl1 plays a role in protecting cells against iron toxicity. Examination of the *S. pombe* genome database suggests that the open reading frame SPBP23A10.16 encodes a putative ortholog of *S. cerevisiae* Sdh4. Both proteins are predicted to be structurally

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**FIG. 1.** *pcl1*<sup>+</sup>, *sdh4*<sup>+</sup>, and *isa1*<sup>+</sup> gene expression is down-regulated under conditions of iron depletion. (A) Wild-type strain FY435 was grown to mid-logarithmic phase in yeast extract plus supplements. Total RNA from Dip (250 μM), control (−), or FeCl<sub>3</sub> (Fe) (100 μM) cultures was isolated. Shown is a representative RNase protection assay of *pcl1*<sup>+</sup>, *sdh4*<sup>+</sup>, *isa1*<sup>+</sup>, and *act1*<sup>+</sup> (as control) mRNA steady-state levels. Results shown are representative of three independent experiments. (B) Quantification of *lacZ* levels after treatments shown in panel A. Values are the averages of triplicate determinations ± standard deviations. (C) Representative RNase protection assay of *foi1*<sup>+</sup> (used as a control gene known to be induced under conditions of iron starvation) and *pcl1*<sup>+</sup> mRNA steady-state levels. (Bottom) *act1*<sup>+</sup> mRNA as an internal control. Total RNA was extracted from aliquots of cultures incubated in the absence (−) or presence of 250 μM Dip or 100 μM FeCl<sub>3</sub> (Fe) for 90 min at 30°C. (D) Graphic representation of quantification of three independent RNase protection assays, including the experiment shown in panel C. The values are the means of three replicates ± standard deviations.
related, with 35% sequence homology with each other. Based on that observation, it was inferred from homology that the *S. pombe* Sdh4 protein represents the mitochondrial membrane anchor subunit of the succinate dehydrogenase, which is a key iron-sulfur protein of the TCA cycle. Excluding the mitochondrial targeting sequence, the *S. pombe* Isa1 protein shows 71% and 64% sequence similarity to *S. cerevisiae* Isa1 and *E. coli* IscA, respectively. Furthermore, it has been demonstrated that *S. pombe* Isa1 is implicated in iron-sulfur cluster assembly paths (59). We found that in the presence of the iron chelator Dip, pcl1+, sdh4+, and isa1+ remained approximately equal to those observed in untreated cells (Fig. 1A and B). As expected, fio1+ mRNA levels (assayed as a control) were up- and down-regulated after treatment with Dip and iron, respectively (Fig. 1C and D). These results reveal that the pcl1+, sdh4+, and isa1+ genes, encoding proteins that may function in sequestration and utilization of iron, are regulated by iron-limiting conditions in a direction opposite that of the iron transport genes.

**Identification of the cis-acting element necessary for activation of pcl1+ gene expression under basal and iron-replete conditions.** In order to determine the regulatory sequence responsible for the iron-replete-mediated expression of pcl1+, sdh4+, and isa1+, we examined several regions of the pcl1+
promoter as well as the 3’ untranslated region of the pcl1’ gene. To ascertain whether the pcl1’ promoter was involved in pcl1’ regulation in response to changes in iron levels, a series of pcl1’ promoter deletion mutants were fused upstream of and in-frame to the lacZ gene in pSP1-lacZ (43). Gene expression from these plasmids was analyzed by RNase protection experiments. As shown in Fig. 2, removal of DNA sequences between –680 and –211 of the pcl1’ promoter had no effect on the iron-regulatable expression of pcl1’-lacZ fusions. In the presence of Dip, plasmids pSP1 pcl1’-680lacZ, pSP1 pcl1’-364lacZ, pSP1 pcl1’-247lacZ, and pSP1 pcl1’-211lacZ were still repressed by approximately sevenfold compared with the basal level of expression observed during testing under both untreated and iron-replete conditions (Fig. 2). Furthermore, for all of the above-mentioned pcl1’-lacZ promoter derivatives, basal and elevated iron concentrations resulted in high constitutive levels of pcl1’-lacZ mRNA expression (Fig. 2). When the pcl1’ promoter was deleted to position –173, iron-regulatable expression was completely abolished. Furthermore, truncation to position –173 also abolished the highly expressed steady-state level of pcl1’-lacZ mRNA in the presence of iron, lowering its expression to a minimal threshold (Fig. 2).

On the other hand, we also analyzed wild-type pcl1’ containing its own 3’ untranslated region and pcl1’ coding sequence with its 3’ untranslated region replaced by the CYC1 3’ region. Both constructs were expressed under the control of the wild-type pcl1’ promoter and both were down-regulated (approximately sevenfold) under low-iron conditions (Mercier et al., unpublished). Although we have identified a putative S. pombe ortholog to Cth2, designated Zfs1 (encoded by the locus SPBC1718.07c), we determined that zfs1’ mRNA is not regulated by iron addition or depletion (Mercier et al., unpublished). Furthermore, we ascertained that iron deprivation-repressed pcl1’ expression is independent of zfs1’ (Mercier et al., unpublished). Total RNA isolated from an isogenic pair of strains, one carrying an inactivated copy of pcl1’ and the other harboring a wild-type copy of the gene, showed no apparent effect on pcl1’ transcript levels or its regulation by iron availability. Consistently, inspection of the pcl1’ 3’ untranslated region did not allow the identification of putative AU-rich elements (47).

Because of the observation that the presence of the promoter region between –211 and –173 was required for governing iron-replete-mediated expression of the pcl1’-lacZ fusion gene, we examined whether a pcl1’ promoter segment including this region could contain a cis-acting element responsible for such iron-regulatable gene expression. Interestingly, the pcl1’ promoter region between –205 and –201 harbors a copy of the sequence CCAAT that is identical to the binding site for the CCAAT-binding transcription factor (30). To examine whether this CCAAT sequence could mediate gene expression as a function of iron availability, we inserted multiple point mutations that mimic changes known to abolish binding of the CCAAT-binding factor to CCAAT sequences in both elements within the sdh4+ and isa1+ promoters, a low and constitutive basal level of expression was observed. In fact, there was a complete lack of either down- or up-regulation of the sdh4+-CYC1-lacZ and isa1+-CYC1-lacZ fusions. Collectively, these results indicate that CCAAT sequences in pcl1’, sdh4’, and isa1’ genes are required to confer iron-regulated pcl1’, sdh4’, and isa1’ gene expression. pcl1’, sdh4’, and isa1’ genes require the Php2 subunit of the heteromeric CCAAT-binding factor for expression under conditions of iron availability. Based on the gene expression data we obtained, we predicted that the CCAAT-binding complex of S. pombe, which contains products of the php2’, php3’, and php5’ genes, may be required to regulate the expression of pcl1’. To test this hypothesis, and because Php2, Php3, and Php5 are essential for the formation of the CCAAT-binding complex in S. pombe (38), we first deleted the php2’ gene
php2Δ, pcl1+ mRNA was barely detected in php2Δ cells and not regulated by iron deprivation (Fig. 5). Importantly, iron limitation-dependent down-regulation of pcl1+ gene expression was corrected by integrating a php2+ allele expressed from its own promoter (Fig. 5). As a control, under conditions of iron deficiency, pcl1+ expression was found to be reduced (approximately sevenfold) compared with the basal level of pcl1+ transcript detected in untreated wild-type cells (php2Δ) (Fig. 5). Cells harboring an inactivated php2+ gene (php2Δ) also failed to mediate iron starvation-dependent repression of sdh4+ and isa1+. Interestingly, in php2Δ cells the magnitude of the steady-state basal level of sdh4+ or isa1+ mRNA was greater than that observed for pcl1+. This suggests the presence of additional transcription factors for basal gene expression of sdh4+ and isa1+. As shown in Fig. 5, the analysis of sdh4+ and isa1+ mRNA levels in a strain expressing php2+ showed complete repression of both sdh4+ and isa1+ expression in the presence of the iron chelator Dip. In php2Δ cells, sdh4+ or isa1+ gene expression was not completely repressed after treatment with 250 μM Dip. This observation clearly suggests a requirement for the Php2/Php3/Php5 complex to repress gene expression during iron deficiency. Consistently, mutations in the php3+ - and php5+ -encoded CCAAT-binding proteins were phenocopies of php2Δ (Mercier et al., unpublished). Taken together, these data establish a requirement for the presence of the CCAAT-binding complex to control the
level of \textit{pcl1}, \textit{sdh4}, and \textit{isa1} expression as a function of iron availability.

The \textit{pcl1} promoter CCAAT element is recognized by the CCAAT-binding complex. To further investigate the function of the CCAAT-binding factor in this regulatory pathway, an oligomer spanning the CCAAT box derived from the \textit{pcl1} promoter (Fig. 6A) was end labeled and used in an EMSA with extracts derived from \textit{php2} wild-type and \textit{php2} deletion strains. As shown in Fig. 6B, the \textit{pcl1} promoter CCAAT element formed a DNA-protein complex when extracts were prepared from \textit{php2} wild-type cells, whereas the extracts prepared from the \textit{php2} deletion strain showed loss of the complex. When extracts were prepared from \textit{php2} deletion cells in which a wild-type copy of \textit{php2} was returned by integration, a DNA-protein complex was formed and had an electrophoretic mobility similar to that observed with extracts derived from wild-type cells. Together, the results suggest that Php2 is necessary for the assembly and DNA-binding activity of the DNA-protein complex.

To determine whether CCAAT-binding activity could be reconstituted by the addition of recombinant Php2 to \textit{php2} extracts, we expressed a conserved core domain of Php2 (residues 2 to 70) fused to MBP in \textit{E. coli} cells. The polypeptide was purified to near homogeneity by two rounds of one-step affinity chromatography based on MBP affinity for maltose. The fusion protein was used in a mobility shift assay with extracts prepared from the \textit{php2Δ} deletion strain. As shown in Fig. 6B, the addition of MBP-\textsuperscript{2}Php2\textsuperscript{70}
fusion protein, but not MBP alone, reconstituted DNA-binding activity. To determine the specificity of the DNA-protein complex formation, we carried out competition experiments with unlabeled oligomers using either wild-type CCAAT or CCAAT with multiple point mutations within the 33-bp DNA fragment specified in Fig. 6A. Formation of the DNA-protein complex was inhibited by incubation with excess wild-type oligomer but not by the mutant competitor, indicating that the complex was formed by sequence-specific interactions (Fig. 6C). Taken together, these data indicate that the Php2 N-terminal 70 amino acid residues contain CCAAT recognition and subunit association domains and, therefore, are likely required for CCAAT-dependent transcription of pcl1\(^+\).

**Php4 is required for down-regulation of pcl1\(^+\), sdh4\(^+\), and isa1\(^+\) mRNA levels under low-iron conditions.** In *S. cerevisiae*, a feature of the CCAAT-binding factor is the presence of a fourth subunit in addition to subunits 2, 3, and 5. This fourth subunit of the complex, denoted Hap4, is down-regulated in the presence of glucose and induced in its absence, while the expression of *HAP2* (subunit 2), *HAP3* (subunit 3), and *HAP5* (subunit 5) is constitutive (13). Although it has been observed that Hap4-like proteins exist in other fungi, the sequence homology between these proteins is very limited (37, 56). Analysis of genomic DNA sequence from the *S. pombe* Genome Project revealed one locus (*SPBC16E9.01c*) that encodes a putative Hap4 homolog, which we have termed Php4. Although Php4 exhibits only 8.5% identity on the amino acid level with the Hap4 sequence from *S. cerevisiae*, one conserved motif found in Php4 (residues 26 to 41), which corresponds to residues 64 to 79 of Hap4, is strongly conserved (see Fig. 8A). This 16-amino-acid domain is known in *S. cerevisiae* to be essential for the interaction of Hap4 with the CCAAT-binding complex that is composed of Hap2, Hap3, and Hap5 (37). Interestingly, using the cross-linking agent EGS, we found that the Php4 protein associates with the Php2/Php3/Php5 complex (Mercier et al., unpublished). To investigate the role of Php4 in fission yeast, we deleted the *php4\(^+\)* gene (*php4\(^-\)*). Inactivation of *php4\(^+\)* gave rise to a sustained level of *pcl1\(^+\)*, *sdh4\(^+\)*, and *isa1\(^+\)* gene expression without any change in response to iron starvation (Fig. 7). On the contrary, *php4\(^-\)* cells in which a wild-type *php4\(^+\)* allele was reintegrated regained the capacity to

![Diagram](image-url)
down-regulate *pcl1*, *sdh4*, and *isa1* gene expression when cells were grown during iron starvation (Fig. 7). These data strongly suggest that repression of *S. pombe* iron metabolic genes during iron deficiency depends on the presence of Php4.

Given these results, we analyzed steady-state mRNA levels of *php2*, *php3*, *php5*, and *php4* as a function of iron availability in the presence of glucose. *php2*, *php3*, and *php5* steady-state mRNA levels were found to be constitutive.

FIG. 6. The *php2* gene product is required for the DNA-binding activity of the CCAAT-binding factor. (A) Sequences of the synthetic oligomers used. The box indicates the wild-type element CCAAT of the *pcl1* promoter, whereas the asterisks indicate that the element contains five substitutions. The nucleotide numbers refer to the position relative to the A of the initiator codon of the *pcl1* open reading frame. (B) DNA mobility shift assays were carried out with DNA-binding reaction mixtures containing the radiolabeled wild-type CCAAT box-containing probe (as shown in panel A) incubated with *S. pombe* extracts prepared from the wild-type (WT) strain or a *php2* mutant strain. An empty vector (−) or a wild-type copy of the *php2* gene was reintegrated (php2+) and tested for the ability to restore DNA-binding activity to *php2* extracts. Purified recombinant MBP or MBP-Php2 was included in the DNA-binding reaction mixtures prepared with *php2* extracts. The positions of the CCAAT-binding complexes are indicated on the left (B). (C) Electrophoretic mobility-shifted gel of a representative competition experiment using whole-cell extracts prepared from strain FY435 (*php2*). Competition was performed with double-stranded DNA unlabeled oligomers corresponding to wild-type (WT) and mutant elements. The amount of competitor used in each reaction is shown over the lanes, and the probe concentration was 1 ng/reaction. B, bound probe DNA; F, free probe DNA.
and unresponsive to cellular iron status (Mercier et al., unpublished). To determine whether the \( \text{php4}^- \) gene has a similar pattern of expression, we carried out RNase protection experiments to detect \( \text{php4}^- \) mRNA from untreated cells or from \( S. \text{pombe} \) cells incubated in the presence of 250 \( \mu \text{M Dip} \) or 100 \( \mu \text{M FeCl}_3 \). As shown in Fig. 8B, \( \text{php4}^- \) transcript levels were barely detectable from iron-replete cells. In contrast, in cells treated with Dip, \( \text{php4}^- \) steady-state mRNA levels were increased approximately fivefold over basal levels detected in untreated cells. Consistently, \( \text{php4}^- \) mRNA was absent in \( \text{php4}^- \) cells. Because \( \text{php4}^- \) was repressed in the presence of exogenous iron, we carried out RNase protection assays with wild-type and\( fep1\Delta \) cells to ascertain whether Fep1 plays a role in \( \text{php4}^- \) gene regulation. In wild-type cells, \( \text{php4}^+ \) mRNA levels were clearly derepressed in the presence of 250 \( \mu \text{M Dip} \). However, in the presence of 100 \( \mu \text{M FeCl}_3 \), the steady-state levels of \( \text{php4}^+ \) were down-regulated (Fig. 8B and C). Interestingly, in a \( fep1\Delta \) strain, \( \text{php4}^- \) mRNA was highly expressed and unresponsive to iron for repression (Fig. 8B and C). Taken together, these data reveal that Fep1 is required for iron-mediated repression of \( \text{php4}^+ \) gene expression, which itself is involved in repressing \( \text{pcl1}^+ \), \( \text{sdh4}^+ \), and \( \text{isa1}^+ \) gene expression under conditions of iron deprivation.

**Iron-induced Fep1 binding to the \( \text{php4}^+ \) GATA elements.**

Based on the iron-dependent regulation of \( \text{php4}^+ \) gene expression we obtained (Fig. 8 and Fig. S2 in the supplemental material), we predicted that Fep1 interacts with the sequences \(-186\text{AGATAT}-183\) and \(-165\text{TGATAA}-160\) found in the \( \text{php4}^+ \) promoter. To determine whether such interaction occurs, we produced the N-terminal 241 amino acids of Fep1 fused to MBP in \( E. \text{coli} \) cells grown in the presence of either 100 \( \mu \text{M FeCl}_3 \) or 250 \( \mu \text{M Dip} \). Subsequent to its purification, the recombinant Fep1 fusion protein was used for binding analyses. As shown in Fig. 9 by a representative EMSA, the wild-type 32\(^{P}\)-end-labeled 51-bp \( \text{php4}^+ \) promoter fragment, which contains the above-mentioned GATA sequences, forms a DNA-protein complex in the presence of metallated Fep1. To ascertain the specificity of this complex formation, we carried out competition experiments with unlabeled oligomers using either wild-type GATA or GATA with multiple point mutations.
within the 51-bp DNA fragment (Fig. 9A). Formation of the DNA-protein complexes was inhibited by incubation with excess wild-type oligomer but not by the mutant competitor, indicating that the complexes were formed by sequence-specific interactions (Fig. 9B). To test whether Fep1 binds to GATA elements derived from the *php4*/*H11001* promoter in an iron-dependent manner, EMSAs were conducted using iron-and Dip-treated purified protein preparations. As shown in Fig. 9C, the 32P-end-labeled double-stranded *php4*/*H11001* oligomer formed a complex with MBP-Fep1 when the fusion protein was purified from iron-treated *E. coli* extracts. In contrast, no complex was formed when the purified fusion protein was isolated from Dip-treated *E. coli* extracts (Fig. 9C). Furthermore, a reconstitution experiment was carried out to determine the ability of iron to restore MBP-Fep1 GATA-binding activity. Dip-treated MBP-Fep1 peptide was incubated with 10 μM FeCl₃ for 30 min. A 20-μl portion of the treated sample was assayed by gel shift retardation assay. As shown in Fig. 9C (last
lane). Fep1 GATA-binding activity was restored by iron. The fusion proteins from iron- and Dip-treated cells were probed with monoclonal antibodies against MBP to confirm their presence in the chromatographic fractions (Fig. 9D). Taken together, these results indicate that the N-terminal 241 amino acids of Fep1 associate with GATA promoter elements from \(php4^+\) in an iron-mediated manner.

**DISCUSSION**

In this paper, we present a novel homeostatic mechanism whereby the fission yeast \(S. pombe\) regulates gene expression in response to iron deficiency. We identified three genes encoding putative iron-containing proteins (\(pcl1^+\), \(sdh4^+\), and \(isa1^+\)) that are down-regulated at the mRNA level in response to iron depletion. Our results showed that the \(pcl1^+\), \(sdh4^+\), and \(isa1^+\) promoters require a CCAAT-type regulatory sequence for expression and regulation by iron starvation. Given the fact that multiple point mutations within the \(pcl1^+\), \(sdh4^+\), and \(isa1^+\) promoters CCAAT-type sequences abolished iron starvation-dependent repression of \(pcl1^+\), \(sdh4^+\), and \(isa1^+\), we tested the possibility that the CCAAT-binding complex of \(S. pombe\) plays a role in the regulation of \(pcl1^+\), \(sdh4^+\), and \(isa1^+\) gene expression. Indeed, we demonstrated that cells bearing a disruption of \(php2^+\), which encodes one subunit of the CCAAT-binding factor, are defective in the regulation of \(pcl1^+\), \(sdh4^+\), and \(isa1^+\) mRNA levels. To further understand how the \(S. pombe\) CCAAT-binding factor functions to regulate target gene expression in response to iron deprivation, we looked for clues about a potential mechanism. First, we found similar levels of \(php2^+\), \(php3^+\), and \(php5^+\) transcripts in low- and high-iron-grown cells of \(S. pombe\), indicating that transcription of \(php2^+\), \(php3^+\), and \(php5^+\) is independent of cellular iron status. Second, we observed no variation of \(Php2\), \(Php3\), and \(Php5\) protein steady-state levels in cells grown in the absence or presence of iron (Mercier et al., unpublished). Third, because EMSAs using extracts from Dip-treated, control, or iron-treated cells revealed the presence of a specific CCAAT-protein complex (Mercier et al., unpublished), we concluded that the DNA-binding activity of the \(S. pombe\) CCAAT-binding factor was insensitive to iron depletion. Thus, the question of how lack of iron within cells affects the CCAAT-binding complex was still an enigma. From there, we hypothesized that this might occur through an upstream component that could produce an alteration in activity of the \(php2/Php3/Php5\) heterotrimer. Based on a report that a putative homologue of \(S. cerevisiae\) \(Hap4\) had been identified in the genome of \(S. pombe\) (56), we tested the possibility that \(php4^+\) plays a role in the regulation of \(pcl1^+\), \(sdh4^+\), and \(isa1^+\) gene expression during iron deficiency. \(php4^+\) encodes a protein of 295 amino acids harboring a conserved 16-amino-acid region, \(^{26}RSVKQWV\) \(PPRPKPG\). The conserved motif that is present in \(Hap4\) is known to be required for its association with \(Hap2/Hap3/Hap5\). Although \(Php4\) bears this N-terminal 16-amino-acid motif found within \(S. cerevisiae\) \(Hap4\) as well as other \(Hap4\) homologues (55, 56), the rest of \(Php4\) exhibits a limited overall sequence homology with \(Hap4\) (8.5% identity, 12.8% similarity). Furthermore, four notable differences exist between the \(S. pombe\) \(Php4\) and \(S. cerevisiae\) \(Hap4\) products. First, under glucose conditions, \(php4^+\) mRNA is readily detectable in fission yeast. Although \(php4^+\) mRNA was unregulated by cellular iron status, under glucose conditions and in the absence of Fep1, its expression was clearly observed. This contrasts with the situation in \(S. cerevisiae\), where the \(HAP4\) gene has been shown to be repressed in the presence of glucose and derepressed when cells are grown on nonfermentable carbon sources (16). Second, in \(S. pombe\), inactivation of \(php4^+\) (\(php4\)) does not affect the ability of cells to grow on nonfermentable carbon sources (Mercier et al., unpublished), while \(S. cerevisiae\) \(hap4\) deletion cells are defective in growth on nonfermentable carbon sources (16). Third, in \(S. pombe\), we observed that induction of \(ycl1^+\) expression by a shift to a nonfermentable carbon source is independent of \(Php4\) (Mercier et al., unpublished). In \(S. pombe\) cells containing a \(hap4\) null allele, genes involved in mitochondrial electron transport such as \(CYC1\) are not induced when cells are shifted from glucose to a nonfermentable carbon source (16). Fourth, the \(S. pombe\) \(Php4\) protein is predicted at neutral pH to have a net charge of +9 (pI = 9.9) without any indication of an acidic cluster region. As opposed to this situation, the \(S. cerevisiae\) \(Hap4\) protein is quite hydrophilic, with a predicted isoelectric point of 5.2. Furthermore, two regions capable of stimulating transcription have been mapped within \(Hap4\), one between residues 359 and 476 and the other between residues 124 and 329 (55). Taken together, these differences led us to hypothesize that the \(Php4\) protein in \(S. pombe\) may play a regulatory function different from that of the \(S. cerevisiae\) \(Hap4\) protein.

Consistent with a role for \(Php4\) as a regulatory protein in iron-regulated gene expression, we determined that its expression is under the control of the iron-regulatory transcription factor Fep1. We identified two GATA-like elements (positions −188 to −183 and −165 to −160) in the \(php4^+\) promoter that can be specifically bound by Fep1. Furthermore, as previously observed for the interaction between Fep1 and \(fio1^+\) promoter GATA sequences (43), Fep1 interacts with the \(php4^+\) promoter GATA elements in an iron-dependent manner. Based on our previous (43) and current observations, we propose a model for the regulation of genes encoding iron-containing proteins during iron deficiency (Fig. 10). In the absence of iron, an inactivated Fep1 fails to bind GATA elements in the \(php4^+\) promoter. The \(Php4\) protein is synthesized and associates with the \(Php2/Php3/Php5\) heterotrimer already present on the promoters of genes required for iron utilization. As a result, the \(Php2/Php3/Php5/Php4\) complex blocks target gene expression, presumably to avoid a futile expenditure of energy in producing iron-using proteins that lack the necessary cofactor (iron) to function. On the other hand, under conditions of iron excess, Fep1 interacts with GATA elements in the \(php4^+\) promoter, repressing \(php4^+\) gene expression. Lack of \(Php4\) allows the CCAAT-binding \(Php2/Php3/Php5\) heterotrimer to activate gene expression. Recently, using DNA microarrays, Lan et al. (27) have demonstrated that the \(Candida albicans\) \(HAP43\) gene (orf19.8298/orf9.681), which encodes a putative transcriptional regulator orthologous to \(S. pombe\) \(Php4\), is regulated in response to iron status in the same manner as \(php4^+\). Furthermore, it has been shown that the iron-responsive transcriptional control of \(HAP43\) is mediated by the \(C. albicans\) Fep1 ortholog, Sfu1 (27). These findings with \(C. albicans\) support our results that \(Php4\) is an iron-regulated protein. Furthermore, this also suggests that \(C. albicans\) may have a closer
relationship to *S. pombe* than to *S. cerevisiae* with respect to the ability to establish and maintain normal iron homeostasis.

All organisms share a requirement for the regulation of iron metabolism during iron deficiency. In bacteria, a posttranscriptional down-regulation of iron-using pathways, which depends on small noncoding regulatory RNAs, has been uncovered (35, 57). In *S. cerevisiae*, two transcription factors, Aft1 and Aft2, are iron sensors that activate genes encoding components of iron transport systems in response to iron deprivation (7, 50, 51, 61, 62). Cth2, an RNA-binding protein, is also synthesized.
when the availability of iron is limited. Induction of Cth2 is mediated by increased transcript levels, and the Aft1 and Aft2 transcription factors play an essential role in this process (47). In response to low environmental iron levels, Puig et al. (47) have shown that Cth2 binds to specific mRNAs and triggers them for degradation. The AU-rich elements through which Cth2 effects its function are found in the 3′ untranslated regions of targeted mRNAs, which encode several iron-containing components, including enzymes involved in the TCA cycle, Fe-S cluster assembly, and heme biosynthesis (47). Although we have found a potential S. pombe ortholog to Cth2, named Zfs1, its deletion had no apparent effect on pcl1+′, sdh4+′, and isa1+′ transcript levels or their regulation by iron availability. Thus, this suggested that S. pombe has a distinct regulator for iron limitation-dependent down-regulation of gene expression.

Optimizing the utilization of limited available iron represents a critical challenge for all organisms, and it is therefore not surprising that various solutions have evolved. The results presented here describe a novel regulatory mechanism that governs cellular metabolism during iron deficiency in S. pombe. The question of whether a similar basic mechanism exists in other organisms represents a pertinent area for future investigation.

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