Increased dosage of a transcriptional activator gene enhances iron-limited growth of \textit{Saccharomyces cerevisiae}

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We have selected for genes that, when present in multiple copies, enhance growth of wild-type cells of \textit{Saccharomyces cerevisiae} in an iron-limiting medium. A gene designated \textit{FUP1}, for ‘ferric utilization proficient’, was isolated by this approach. Increased dosage of \textit{FUP1} reduces the concentration of iron in the medium required for efficient growth and confers elevated levels of iron uptake activity in iron-limited cells. Disruption of the \textit{FUP1} locus reduces wild-type iron uptake rates by 2-fold in cells grown on raffinose medium but has no effect on glucose-grown cells. DNA sequencing showed that \textit{FUP1} encodes a hydrophilic 43 kDa protein identical to \textit{MSN1}, a gene encoding a transcriptional activator implicated in carbon source regulation. Our results suggest that \textit{FUP1/MSN1} also regulates synthesis of gene products involved in iron uptake.

Introduction

Iron is a critically important component of cellular biochemistry. Its two stable oxidation states, Fe$^{3+}$ and Fe$^{2+}$, allow the metal to participate in a wide variety of redox reactions involved in such diverse processes as the synthesis of DNA, lipids and amino acids, and in respiratory electron transport. Although abundant, iron is often unavailable for cell growth because the oxidized form, Fe$^{3+}$, is extremely insoluble at neutral pH. Therefore, organisms have evolved mechanisms to efficiently obtain iron from their environment (for a comprehensive review, see Winkelmann \textit{et al.}, 1987). For example, many bacteria and fungi secrete iron-binding compounds known as siderophores that bind extracellular iron; this complex is then brought into the cell via a receptor-mediated process.

The iron transport systems of many organisms are regulated in response to iron limitation and metabolic demand for iron (Winkelmann \textit{et al.}, 1987). In bacteria and fungi, the synthesis, secretion and internalization of siderophores are increased during growth in low iron environments.

The yeast \textit{Saccharomyces cerevisiae} apparently does not produce any siderophores (Schwyn & Neilands, 1987). The major mechanism of iron accumulation by \textit{S. cerevisiae} involves an iron reductase, located in the plasma membrane, that reduces Fe$^{3+}$ to Fe$^{2+}$ (Lesuisse \textit{et al.}, 1987; Lesuisse & Labbe, 1989; Dancis \textit{et al.}, 1990). Studies indicate that the reductase is probably encoded by the \textit{FRE1} gene (Dancis \textit{et al.}, 1990). Reduced iron is then transported into the cell by a carrier or channel located in the plasma membrane. The activity of the membrane iron reductase is regulated in response to iron availability, i.e. the reductase is more active as iron becomes increasingly limited (Dancis \textit{et al.}, 1990). This regulation is probably the result of transcriptional control because steady-state levels of \textit{FRE1} mRNA correspond to the level of reductase activity (Dancis \textit{et al.}, 1990).

The long-range goal of our research is to understand the process of iron uptake in yeast and determine how cells adapt to iron limitation. This communication describes a genetic strategy that we have used to identify genes involved in iron metabolism. We have screened a multicopy plasmid library for genes that, when present in several copies per cell, enhance growth of wild type cells in iron-limiting conditions. Using this approach we have identified the \textit{FUP1} gene. DNA sequencing revealed that \textit{FUP1} is identical to \textit{MSN1}, a transcriptional activator involved in carbon source regulation (Estruch & Carlson, 1990). Increased gene dosage of \textit{FUP1}/\textit{MSN1} reduces the amount of iron required for optimum cell growth.
This lower iron requirement is because *FUP1/MSNI* stimulates iron uptake during growth in low iron medium.

**Methods**

**Preparation of low-iron medium (LIM).** LIM is based on the Wickerham nitrogen base recipe (Wickerham, 1946) with two modifications essential to control iron availability. First, 1 mM-EDTA is added to provide a buffering capacity for the concentrations of multivalent metal cations. Second, the medium is pH-buffered at 4.2 with 20 mM-citrate to prevent pH changes that could greatly alter the metal binding ability of EDTA. This is an important modification because growing yeast can acidify an unbuffered medium to below pH 3.

The stock solutions from which LIM was made are described in Table 1. These were prepared from reagent grade chemicals and deionized-distilled water (DDW). The pH values of stocks 1 and 7 were adjusted to 8.0 and 4.2, respectively, with HCl. Stock 10 and the FeCl₃ stocks were prepared in 0.1 M-HCl. Solutions were filter-sterilized with 0.22-μm cellulose nitrate filters (Costar) and stored in polycarbonate bottles. To prepare the medium, the stock solutions were added in numerical order to 438.5 ml of DDW, filter-sterilized, and stored in deionized-distilled water (DDW). The pH values of stocks 1 and 7 were 9.0 and 5.0, respectively.

Contamination from culture inocula was minimized by inoculating from cultures that had negligible iron concentrations. Such cultures are not iron starved, because growing yeast can acidify an unbuffered medium to below pH 3.

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were tested for a growth enhancement effect by transforming the plasmids into DBY2063 and measuring cell growth rate in LIM(0-25). The 2-3 kb SalI–EcoRI fragment from pDE82 was used to replace the wild-type gene in DBY2063 by gene transplacement (Rothstein, 1991) to generate the strain DEYll94. Transplacement was confirmed by Southern blot analysis.

To sequence the FUP1 gene, we inserted the 6 kb SalI–Smal fragment from pDE1 into YEplp5 to yield pDE2. The 3-8 kb EcoRV–SalI fragment from pDE2 was then cloned into M13mp18 and the pDE2 3-8 kb EcoRV–HindIII fragment was cloned into M13mp19 to produce mpDE1 and mpDE2, respectively. Deletions spanning the mpDE1 and mpDE2 inserts were generated using the method of Henikoff (1984). Overlapping deletions were sequenced with the Sequenase DNA Sequencing Kit from United States Biochemicals.

**Iron uptake and reductase assays.** Exponentially growing cells were chilled on ice for 20 min, centrifuged at 1000 g for 5 min at 4 °C, washed twice in ice-cold LIM – EDTA (i.e. LIM lacking EDTA), and resuspended in approximately 1/100th the original culture volume in LIM – EDTA. Cell suspensions were kept on ice prior to use. Uptake assay solutions were prepared by diluting 59FeCl3 (Amersham) into chilled LIM – EDTA at the appropriate concentrations. In assays in which iron was supplied as Fe2+, 1 mM-sodium ascorbate was included in the assay solution. To begin the uptake assay, 50 μl of cell suspension was added to 450 μl of assay solution and transferred to 30 °C. After 10 min, the assay samples were chilled on ice, vacuum filtered through Whatman GF/C filters and washed in 10 ml ice-cold SSW (1 mM-EDTA, 20 mM-trisodium citrate pH 4.2, 1 mM-KH2PO4, 1 mM-CaCl2, 5 mM-MgSO4, 1 mM-NaCl). Non-specific uptake due to surface adsorption was determined by preparing parallel assays that were held on ice for 10 min before filtration and washing. These background levels of cell-associated 59Fe were subtracted out before uptake rates were calculated. 59Fe levels were measured with a Packard Minaxi γ Autogamma 5000 gamma counter. Specific activity, rate of decay and cell number were used to calculate the uptake rates.

Assays of the iron reductase were performed in a manner similar to that used for uptake assays. Aliquots (50 μl) of the cell suspensions were inoculated into 950 μl ice-cold LIM – EDTA supplemented with 10 μM-FeCl3 and 1 mM-bathophenanthroline disulphonate (BPS) (Landers & Zak, 1958). Samples were incubated at 30 °C for 10 min and then chilled on ice. Cells were removed by spinning for 20 s at 10000 g, and the absorbance at 520 nm was measured. The concentration of iron reduced was determined by comparison to a standard curve. Background levels were determined using parallel samples that contained no cells; these values were then subtracted from the values obtained with the cell-containing samples before calculating the reduction rate.

**Results**

**LIM is iron-limiting**

Media capable of controlling cell growth by iron limitation can be difficult to prepare because iron is a major contaminant of reagents, glassware, etc., and is required by yeast in only trace amounts. One common approach to preparing an iron-limited medium is to extract the iron from the media (Nicholas, 1957; Hewitt, 1966). Small amounts of iron can then be added back to the medium to restore cell growth. A second method, which we used in this work, takes advantage of the fact that chelating agents, such as EDTA, bind metal ions and make them unavailable to the cell. This approach was suggested by the work of F. M. M. Morel and co-workers (Morel et al., 1979; Anderson & Morel, 1982) on iron limitation of phytoplankton. Iron limitation with chelators has several advantages over iron extraction methods. The resulting medium is less sensitive to contamination by iron, and precipitation of constituents is prevented.

A wild-type yeast strain, DBY2063, could not grow on LIM without added iron. Addition of 10 μM-FeCl3 was sufficient to promote cell growth, whereas 10 μM supplements of Zn2+, Cu2+, Mn2+, Mg2+ and Ca2+ failed to stimulate growth. This observation precludes the possibility that the added FeCl3 was affecting the availability of another metal by titrating out the EDTA in the medium. To determine how different amounts of added iron affect cell growth, we measured the growth rate of DBY2063 on LIM supplemented with 0 to 25 μM-FeCl3 (Fig. 1a). The specific growth rate (μ) of DBY2063 increased as iron was added to the medium up to a concentration of 10 μM. The linearity of these data when displayed on a Lineweaver–Burk reciprocal plot (Fig. 1b) demonstrated that this response followed the relationship of Monod (\(\mu = \mu_{\text{max}} \times S/[K_p + S]\)), where \(\mu\) = specific growth rate, \(\mu_{\text{max}}\) = maximum specific growth rate, \(S\) = substrate concentration and \(K_p\) = concentration...
Table 2. Effect of EDTA on iron uptake and reduction

Wild-type DBY2063 cells were grown to the exponential growth phase in LIM(10). These cells were collected and assayed for uptake of iron supplied as $^{59}$Fe$^{3+}$ or $^{59}$Fe$^{2+}$ and for reduction activity in LIM – EDTA and LIM. All assays were performed in 10 $\mu$M-Fe. The mean values of two experiments, each performed in duplicate, is shown; the numbers in parentheses are the standard deviations.

<table>
<thead>
<tr>
<th>Assay medium</th>
<th>Iron uptake rate [fmol min$^{-1}$ (10$^6$ cells)$^{-1}$] with iron supplied as:</th>
<th>Iron reduction rate [nmol min$^{-1}$ (10$^6$ cells)$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>– EDTA</td>
<td>Fe$^{3+}$ 822 (± 75) Fe$^{2+}$ 894 (± 33)</td>
<td>18 (± 2)</td>
</tr>
<tr>
<td>+ EDTA</td>
<td>Fe$^{3+}$ 32 (± 15) Fe$^{2+}$ 26 (± 4)</td>
<td>19 (± 1)</td>
</tr>
</tbody>
</table>

EDTA inhibits uptake but not reduction

LIM prepared without EDTA supported cell growth without additional iron. It was clear from this result that the EDTA in the medium limits the availability of iron to the cells. EDTA may exert this effect by either preventing the reductase from reducing iron or by blocking uptake of Fe$^{2+}$. To test these hypotheses, we determined how EDTA affects iron reduction and uptake when supplied to cells as either Fe$^{3+}$ or Fe$^{2+}$. DBY2063 cells were grown to the mid-exponential phase in LIM(10) and assayed in LIM or LIM prepared without EDTA (LIM – EDTA) for iron reductase activity and uptake of $^{59}$Fe$^{3+}$ or $^{59}$Fe$^{2+}$ (Table 2).

Uptake of iron, supplied as Fe$^{3+}$, was inhibited by EDTA. This inhibition was not due to a block in reduction because this rate was not affected by the chelator. Because the vast majority of iron in reductase assays containing EDTA would be bound to the chelator ($K_a = 10^{23}$), we conclude that the Fe$^{3+}$–EDTA complex is an acceptable substrate for the reductase. Iron can be reduced in the uptake assay from Fe$^{3+}$ to Fe$^{2+}$ by adding 1 mM-ascorbate. Uptake of Fe$^{2+}$, as measured in the presence of ascorbate, was inhibited by EDTA as much as when iron was supplied as Fe$^{3+}$. Therefore, the iron-limiting properties of EDTA are the result of its ability to chelate Fe$^{2+}$ ($K_a = 10^{14}$) and make it unavailable for uptake.

Isolation of genes that enhance iron-limited growth

Our ability to limit the iron available to cells allowed us to select for genes that lower the iron concentration in the medium required for efficient growth. We screened a yeast genomic library in a multicopy plasmid for genes that enhanced cell growth in LIM supplemented with 0.25 $\mu$M-FeCl$_3$ [i.e. LIM(0.25)]. DBY2063 was transformed with a genomic plasmid library, pooled, and grown for several passages on LIM(0.25) until library-transformed cultures grew better than a vector-only control. Individual cells from these cultures were then cloned and tested for growth on LIM(0.25). Two plasmids that enhance the growth rate of DBY2063 in iron-limited conditions were isolated from 15 000 independent transformants. Restriction mapping showed that these plasmids contained overlapping but non-identical inserts. The plasmid pDE1, which contained the smaller of the two inserts (approximately 6 kb), was used in subsequent experiments.

Effect of pDE1 on cell growth and iron uptake

To ascertain the effect of pDE1 on the iron requirement of cells, we determined the relationship between the specific growth rate, $\mu$, and the concentration of iron in the medium ([Fe]). DBY2063 bearing pDE1 or the parental vector, YEp24, was grown in LIM containing a range of iron concentrations, and growth rates were measured (Fig. 2). The $K_s$ value of the pDE1-bearing strain, 0.2 $\mu$M-FeCl$_3$, was approximately one-fourth the $K_s$ of the parent strain (0.8 $\mu$M, Fig. 1) and the YEp24 control (0.7 $\mu$M, Fig. 2). No difference in growth rate was observed for these strains when iron was not limiting.
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**Fig. 3.** Restriction maps of the yeast genomic fragments in pDE1 and its derivatives. The plasmid pDE1 was digested with the restriction enzymes *SnaBI* (A), *BamHI* (B), *EcoRI* (R), *EcoRV* (V), *MhuI* (M), *SmaI* (S), *SalI* (L), *XhoI* (H) and *XbaI* (X), and the products were analysed by agarose gel electrophoresis. The map shows the positions of restriction sites within the pDE1 insert (thick line) and flanking vector (dashed line) DNA. The open box labelled FUP1 indicates the location of the open reading frame as determined by DNA sequencing and the arrow indicates the direction of transcription. The lines below the map represent the fragments that are present in the subclones pDE103, pDE104, pDE105, pDE106 and pDE107. The UR43 gene was inserted into the deletion interval in pDE107. The five resulting plasmids were transformed into DBY2063 and assayed for their ability to enhance cell growth in iron-limited medium by determining the growth of the transformant in LIM(0.25). To the right of each line is the result of these growth tests. The region sequenced is also indicated.

**Table 3. Effect of multiple FUP1 copies on iron uptake**

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Iron uptake rate [fmol min⁻¹ (10⁶ cells)⁻¹]</th>
</tr>
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<tbody>
<tr>
<td>None</td>
<td>97 (± 6)</td>
</tr>
<tr>
<td>YEp24</td>
<td>78 (± 2)</td>
</tr>
<tr>
<td>pDE1</td>
<td>506 (± 2)</td>
</tr>
</tbody>
</table>

Thus, the insert in pDE1 enhanced the ability of cells to grow under iron-limiting conditions.

One possible mechanism by which the plasmid enhanced cell growth was by stimulating the iron uptake apparatus. To test this hypothesis, we grew DBY2063 and DBY2063(YEp24) and DBY2063(pDE1) were grown exponentially in LIM(0-25) and assayed for uptake in 2 μM-iron supplied as ⁵⁸Fe⁺. The mean values of two experiments, each performed in duplicate, are shown; the numbers in parentheses are the standard deviations.

**Analysis of the FUP1 gene**

To determine the location of the gene responsible for this effect within the 6 kb insert of pDE1, deletions were generated in this plasmid and tested for their ability to enhance growth of DBY2063 on LIM(0-25) (Fig. 3). The results indicated that the gene is located between the *SnaBI* and *EcoRV* sites of the pDE1 insert. The sequence of this region was determined and a single large open reading frame (ORF) of 1146 bases was found. This ORF was found to be identical to the *MSN1* gene [see Estruch and Carlson (1990) for the sequence]. We refer to this gene as *FUP1* for ‘ferric utilization proficient’. The *FUP1* ORF was affected by all deletions that disrupted the growth effect of the plasmids. The gene encodes a protein of 382 amino acids with a predicted molecular mass of 43 kDa. A search of the NBRF protein database found no proteins similar to the FUP1 protein (Lipman & Pearson, 1985). The codon adaptation index, a measure of codon bias, of *FUP1* was calculated to be 0.104, suggesting that the gene is not highly expressed (Sharp & Li, 1987). The protein is rich in serine and threonine (17%) and asparagine plus glutamine (18%). A hydropathy plot of the FUP1 amino acid sequence generated by the method of Kyte & Doolittle (1982) shows a largely hydrophilic profile. The longest region of hydrophobicity (residues 85 to 105) has an average...
hydropathy index of +0.4. We have noted some specific features of the FUPI amino acid sequence (Fig. 4). The C-terminus is basic, having a net charge of +13 over 78 residues. There is a potential leucine zipper, a series of four heptad leucine repeats, spanning residues 12 to 40, with one gap at position 33 (Landschulz et al., 1988). We have also noted a consensus target sequence for phosphorylation of the protein by the cAMP-dependent protein kinase, RRXS, at amino acids 28 to 31 (Kemp et al., 1988).

**Effect of a FUPI deletion allele on cell growth and iron uptake**

A deletion/substitution mutation in which the entire FUPI gene had been removed was constructed by deleting the SnaBI–XhoI interval and inserting the URA3 gene (pDE107, Fig. 3). When this allele, designated fupl-1::URA3, was tested on a yeast multicopy plasmid vector, it did not enhance cell growth in LIM(O.25). A fragment containing the fupl-1::URA3 allele was introduced into DBY2063, and we selected for URA+ prototrophs. A URA+ transformant, DEY1194, was analysed by Southern blotting and determined to contain the disruption allele at the FUPI locus (data not shown).

No defect in growth was observed when DEY1194 was cultured on LIM(O-75), a medium in which cells are iron-limited. Thus, a strain completely lacking the FUPI product was wild-type for growth in iron-limiting conditions. Furthermore, no growth defect was observed when this strain was tested for growth on rich media containing either glucose or lactate as the carbon source at either 30 or 37°C.

However, we did note that DEY1194 grew at approximately 75% of the wild-type rate in media containing raffinose as the carbon source. We tested if the growth defect on raffinose was associated with a defect in iron uptake. Wild-type and fupl-1::URA3 cells were grown on glucose and raffinose and assayed for their iron uptake rate when iron was supplied as Fe3+ or Fe2+ (Table 4). The uptake rates of glucose- and raffinose-grown wild-type and fupl-1::URA3 cells were very low when assayed with Fe3+, but elevated approximately 10-fold when assayed with Fe2+. This result was consistent with our observation that the iron reductase was almost undetectable in these cells (data not shown). Therefore, the iron reductase is rate-limiting for uptake in cells grown in these media. Glucose-grown wild-type and mutant cells showed no difference in uptake rate. However, in raffinose-grown cells, the uptake rate observed for the fupl-1::URA3 strain was reduced to 50 to 60% of wild-type rates.

**Table 4. Effect of FUPI disruption on iron uptake**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Glucose</th>
<th>Raffinose</th>
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<tbody>
<tr>
<td></td>
<td>Fe3+</td>
<td>Fe2+</td>
</tr>
<tr>
<td>Wild-type</td>
<td>12.3 (±3)</td>
<td>113.9 (±18)</td>
</tr>
<tr>
<td>fupl-1::URA3</td>
<td>154.4 (±2)</td>
<td>130.3 (±13)</td>
</tr>
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DEY1296 (wild-type) and DEY1298 (fupl-1::URA3) were grown to exponential phase in SD-glucose or SD-raffinose. These cells were assayed for iron uptake with 2 μM-iron supplied as 59Fe3+ or 59Fe2+. The numbers in parentheses are the standard deviations of two experiments, each performed in duplicate.

*Discussion*

We have used a genetic strategy to isolate genes that affect iron-limited growth. Our prediction was that if the product of a gene was itself limiting in low iron, overproduction of that protein would enhance cell growth. The higher gene dosage present in a strain containing a multicopy plasmid often results in the accumulation of higher levels of the product of a plasmid-borne gene relative to the level produced by the chromosomal locus alone (Rine, 1991). Vectors of the type used in this study are present in 25 to 100 copies per cell (Clark-Walker & Miklos, 1974).

The FUPI gene was isolated because of its ability to enhance iron-limited growth. We demonstrated that the FUPI gene, when present in multiple copies, lowers the uptake...
$K_g$ for growth of a wild-type strain by fourfold without affecting the growth rate when iron was in excess. Our prediction is that this effect is due to overproduction of the FUP1 gene product. We have demonstrated that higher levels of FUP1 mRNA accumulate in pDE1-bearing cells than in wild-type cells (data not shown).

It seems likely that FUP1 is involved in the uptake of iron. We observed that strains bearing multiple copies have elevated levels of iron uptake. Yet a fup1 deletion mutant grows normally in iron-limiting media and exhibits wild-type iron uptake rates when grown on glucose and approximately 50% of wild-type when grown on raffinose. One explanation for this apparent paradox is that this effect is due to overproduction of a lexA-FUP1/MSN1 fusion protein could promote transcription from a reporter gene containing a lexA operator as an upstream activation sequence. These results, along with the effects of overexpression on SUC2 expression, suggested that the FUP1/MSN1 gene encodes a transcriptional activator that plays a role in gene regulation in response to carbon source.

There is a remarkable similarity between the effects of altered FUP1/MSN1 copy number on SUC2 expression and iron uptake activity. What is the connection between glucose repression and iron metabolism? In a fup1/msnl mutant, we have observed a defect in iron uptake with raffinose-grown cells but not with glucose-grown cells. Lesuisse et al. (1987) have shown that iron uptake is induced by growth on non-fermentable carbon sources. This induction is presumably because iron in the form of haem is required to metabolize these nutrients. These observations suggest that, to some extent, carbon source utilization and iron uptake are coregulated processes. FUP1/MSN1 may play a role in this regulation by activating transcription of genes involved in both of these responses.

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Estruch & Carlson (1990) presented several lines of evidence to prove that FUP1/MSN1 is a transcriptional activator. A FUP1/MSN1 β-galactosidase fusion protein was nuclear localized, and can bind DNA. Furthermore, a lexA-FUP1/MSN1 fusion protein could promote transcription from a reporter gene containing a lexA operator as an upstream activation sequence. These results, along with the effects of overexpression on SUC2 expression, suggested that the FUP1/MSN1 gene encodes a transcriptional activator that plays a role in gene regulation in response to carbon source.

Estruch & Carlson (1990) concurrently isolated FUP1, which they have called MSNL, as a multicopy suppressor of temperature sensitive mutations in the SNFL locus. SNFL encodes a protein kinase that is required for transcription of glucose-repressible genes when cells are grown on sugars other than glucose, such as raffinose, sucrose and galactose, or on non-fermentable carbon sources, such as lactate or ethanol. When present on a multicopy plasmid, the FUP1/MSN1 gene restores growth and regulated expression of at least one glucose-repressible gene, SUC2. A fup1/msnl disruption, however, caused only a 3- to 4-fold reduction in derepressed levels of invertase, the product of the SUC2 locus, when cells were grown on raffinose.

References


