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The ‘scavenger’ m⁷GpppX pyrophosphatase activity of Dcs1 modulates nutrient-induced responses in yeast

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ABSTRACT

Dcs1, the m⁷GpppX pyrophosphatase of Saccharomyces cerevisiae, has been reported to ‘scavenge’ capped 5’ end fragments generated by 3’→5’ mRNA degradation. We now show that the absence of Dcs1, and the closely related Dcs2 protein, compromises cellular responses to glucose-deprivation stress as well as to step changes in glucose availability. Dcs1 and Dcs2 form homo- and heterodimers, with the heterodimer appearing as cells enter diauxic. Despite the previously observed increase in abundance of the mRNA encoding the neutral trehalase (Nth1) in the stationary phase, the total enzyme activity of Nth1 decreases in this phase of growth. Changes in trehalase activity are significant because the non-reducing disaccharide trehalose is thought to stabilize cellular components under stress conditions. In the dcs1Δ and dcs1Δdcs2Δ mutants, normal regulation of trehalase activity is lost. Nutrient stress induces DCS1 and DCS2 transcription via the cAMP-PKA signalling pathway. Dcs1 also becomes phosphorylated as the availability of glucose diminishes, and we test the role of this phosphorylation in the stress response. Further evidence indicates that Dcs1 plays a complementary role to the translation factor eIF4E in preventing capped 5’ fragments of mRNA from interfering with translation initiation. We conclude that Dcs1 function influences cellular responses to changes in nutrient availability, while Dcs2 seems to act as a modulator of Dcs1 function.

INTRODUCTION

In living cells, enzyme-catalysed degradation processes determine the lifetimes of different mRNAs and thereby influence their steady-state levels. mRNA decay therefore plays an important role in the post-transcriptional control of gene expression (1). The degradation of eukaryotic mRNA is performed largely by exonucleases, whereby access of these enzymes to the body of the mRNA requires removal of either the 3’ poly(A) tail, the 5’ cap, or of both (2–4). Decapping is an important rate-controlling step, since it exposes the 5’ end of mRNAs to attack by 5’→3’ exonucleases of the Xrn1 type (5). Work in the yeast Saccharomyces cerevisiae has identified two proteins, Dcp1 and Dcp2, as involved in m⁷GDP-generating decapping, and homologues of these proteins exist in higher eukaryotes (6,7). However, an m⁷GMP-generating decapping activity detected in mammalian cells has been identified [DcpS (2,8)], and related enzymes have been partially characterized in both S.cerevisiae [Dcs1 and Dcs2 (9)] and in Schizosaccharomyces pombe [Nhml1 (10)]. It has been proposed that the Dcs-type proteins act as ‘scavenging’ pyrophosphatases that help mop up short, capped mRNA fragments left over from 3’→5’ exonuclease decay (2,8).

Little is known about modulation of mRNA decay pathways under conditions of stress. In general, S.cerevisiae regulates gene expression in response to nutrient, temperature and osmotic stresses. This generally involves coordinate transcriptional regulation of different groups of genes. S.cerevisiae has both stress-specific stress factors and more general stress factors. Msn2 and Msn4 are two related factors of the latter type. They become active both during the diauxic shift and under a broad range of stress conditions (11,12). These factors may generally receive and integrate signals from different stress-signalling pathways. They bind to DNA cis-acting stress response elements (STREs) that control a large number of genes. Glucose and cAMP down-regulate MSN2/4 and consequently the STRE-controlled genes (13). Thus, Msn2/4 factors are implicated in PKA-dependent regulation of STRE-controlled gene expression (11,12,14). The Rim15 protein kinase is identified as a downstream target of PKA that acts as an activator of STRE-controlled gene expression (14). The cAMP-PKA pathway is also connected to the nutrient-regulated protein kinases Sch9 and Yak1/Sok1 (14). Protein kinases Rim15 and Yak1 are required for proper cell entry into the stationary phase and suppress the growth arrest caused by PKA depletion, as does deletion of MSN2/4 (15). Recently, Yak1, along with phosphoprotein Pop2, were proposed to function as part of a novel glucose-sensing system in yeast that is involved in growth control in response to glucose availability (16).

The cAMP-PKA pathway is linked to a complex network of signalling cascades. Other components of this network,
including the TOR (target of rapamycin), PKC MAPK (protein kinase C nitrogen-activated protein kinase) and HOG (high-osmolarity glycerol) MAPK pathways, also regulate gene expression in response to stress conditions (17–19). Recently, it was revealed that activation of the cAMP-PKA pathway suppresses TOR deficiency (20). Moreover, TOR can regulate PKA through control of the subcellular localization of both the PKA catalytic subunit Tpk1 and the kinase Yak1 (20). Other results suggest that the HOG pathway may regulate Msn2/4, possibly via cAMP/PKA (18).

In this paper, we characterize how nutrient stress influences the expression of yeast DCS1 and DCS2 at the transcriptional and post-transcriptional levels. Dcs1 function compromises the nutrient stress response and apparently leads to de-regulated synthesis of the stress-response disaccharide, trehalose. We discuss possible mechanisms underlying these regulatory phenomena and the basis for functional interactions between Dcs1 and the cap-binding complex eIF4F.

MATERIALS AND METHODS

Yeast growth conditions

Yeast strains were as shown in Table 1. Cells were grown in liquid YPD (1% yeast extract, 2% peptone, 2% glucose) and in minimal YNB medium (0.67% yeast nitrogen base, 2% glucose) supplemented with uracil and selected amino acids, or on plates containing YPD plus 2% agar. For growth under different conditions, cultures of exponentially growing cells were divided into smaller volumes, washed and resuspended in liquid media containing different carbon sources (glucose, glycerol, trehalose). Alternatively, cultures were serially diluted on YPD, YP, YP(gal), YP(glyc), YPD(H2O2) and YPD(NaCl) agar plates.

Plasmids

PCR fragments bearing the DCS1 and DCS2 ORFs plus flanking sequences were digested with XbaI and Bsp120I and cloned into pRS313 and pRS315 (21), yielding pRS313-DCS1 and pRS315-DCS2, respectively. These plasmids were used for complementation analyses of dcs1Δ and dcs2Δ strains. Plasmids pRS313-DCS1-178 (S60A) and pRS313-DCS1-196 (T66A) were constructed after PCR-mediated site-directed mutagenesis (performed according to (22)). Plasmids pURAGAL1-DCS1-HA, pURAGAL1-DCS1-MH, pURAGAL1-DCS2-HA and pURAGAL1-DCS2-MH were constructed using YcpSUPEX2 (23) by inserting genes DCS1 and DCS2 tagged with either haemagglutinin or Myc epitope. To construct a plasmid for recombiant Dcs1 protein expression, an Ndel–XbaI fragment encoding His8-tagged Dcs1 protein was PCR-amplified from S. cerevisiae genomic DNA and inserted into the pET5A vector (Novagen), yielding pET5A-DCS1. Plasmid pBS1539 (24) was used as the PCR template in the generation of TAP fusion strains PTC196 and PTC197.

Protein purification and sequencing

Dcs2 protein was purified both as the TAP fusion version, and as the non-tagged protein using a two-step purification technique procedure (24). Dcs2–TAP fusion protein was co-purified with the Dcs1 protein. Non-tagged Dcs2 protein was purified as part of the Dcs1–Dcs2 complex isolated via the Dcs1–TAP fusion protein. Of each purified protein complex, was purified as part of the Dcs1–Dcs2 complex isolated via the Dcs1–TAP fusion protein. Non-tagged Dcs2 protein was purified both as the TAP fusion version, and as the non-tagged protein using a two-step purification technique procedure (24). Dcs2–TAP fusion protein was

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Mass spectrometry

Sinapinic acid and calibration standards for mass spectrometry were obtained from Sigma. The MALDI matrix was 10 mg/ml sinapinic acid in 50% acetonitrile–aqueous 0.05% trifluoroacetic acid (TFA). BSA at a concentration of 10 μM (66.4 kDa) and 2 μM apomyoglobin (16.9 kDa) were used as calibrants,
mixed and applied in a 1:1 ratio with the matrix. The sample (Dcs1–TAP and Dcs2 in water) was mixed 1:1 with the matrix and applied to a stainless steel MALDI target and left to dry at room temperature. Positive ion linear MALDI-TOF MS spectra were acquired on an AXIMA-CFR (Kratos Analytical, Shimadzu Biotech, UK).

**Fluorescence microscopy**

Cells were grown in YPD to mid-log phase, fixed by incubation in 4% (v/v) formaldehyde for 30 min at 30°C, and converted to spheroplasts. The TAP fusion was detected using a rabbit anti-protein A antibody (diluted 1:200, Sigma) and a secondary FITC-conjugated goat anti-rabbit IgG (1:100, Vector laboratories, Inc). Haemagglutinin and Myc epitope-tagged proteins were allowed to react, respectively, with rabbit polyclonal anti-haemagglutinin antibodies (1:100, Sigma) followed by FITC-conjugated goat anti-rabbit IgGs (1:100), and with a mouse monoclonal anti-c-Myc antibody (1:50, Sigma) followed by Texas red-conjugated horse anti-mouse IgG (1:100, Vector laboratories, Inc). Nuclear DNA was stained with 4',6-diamidino-2-phenylindole (DAPI) that was included in the mounting medium (Vector Laboratories, Inc). Cells were analysed using a confocal microscope (Zeiss LSM510; objective: HC Plan APO OS 100x oil NA 1.4).

**Western blotting**

Crude extracts were prepared from cultures using glass beads. Equal amounts of protein were separated via SDS–PAGE and transferred to a PVDF membrane (Amersham). Rabbit anti-Dcs1 antibodies were prepared (CovalAb UK Ltd) against recombinant Dcs1–His<sub>6</sub> protein generated in *E. coli*. Dcs2 was detected in the form of Dcs2–TAP using PAP antibodies (Sigma). Immunoreactions on the membrane were visualized by chemiluminescence (ECL, Amersham).

**RNA extraction, northern hybridization and primer extension analysis**

RNA was extracted from cells at the stated time points and northern blots were performed as described elsewhere (25). Primer extension analysis was performed using AMV reverse transcriptase (Promega) and the 32P-labelled oligonucleotide primer (Promega) revealed the presence of stress-related human m7G(5′)pppN-pyrophosphatase, which has a known regulatory role in the transcriptional and translational start sites of *DCS2*. The 5′ end of the *DCS2* mRNA was characterized by means of primer extension (Figure 1A). Examination of the sequences upstream of this transcriptional start site revealed the presence of stress-response elements [STREs (12,28)] in the *DCS2* DNA sequence (Figure 1B). N-terminal protein sequencing also defined the correct position of the translation start codon, revealing that Dcs2 is 352 amino acids long (*M<sub>r</sub> = 39.9 kDa). This makes Dcs2 45 amino acids shorter than the protein that would be predicted using the ATG previously assumed to be the start codon (NCBI database, see upstream ATG highlighted in Figure 1B).

**RESULTS**

**DCS1 and DCS2 expression signals**

Examination of the *DCS1* (*YLR270w*) DNA sequence reveals the presence of a single open reading frame with a predicted product (*M<sub>r</sub> = 40.6 kDa) that corresponds to the electrophoretic migration behaviour of Dcs1 (as detected by western blotting in yeast—see below). No known regulatory motifs are evident in the proximity of the *DCS1* promoter region. In contrast, comparison of the DNA sequence of *DCS2* (*YOR173w*) with the previously estimated size of the encoded protein does not allow unequivocal identification of this gene’s reading frame. We therefore determined the transcriptional and translational start sites of *DCS2*. The 5′ end of the *DCS2* mRNA was characterized by means of primer extension (Figure 1A). Examination of the sequences upstream of this transcriptional start site revealed the presence of stress-response elements [STREs (12,28)] in the *DCS2* DNA sequence (Figure 1B). N-terminal protein sequencing also defined the correct position of the translation start codon, revealing that Dcs2 is 352 amino acids long (*M<sub>r</sub> = 39.9 kDa). This makes Dcs2 45 amino acids shorter than the protein that would be predicted using the ATG previously assumed to be the start codon (NCBI database, see upstream ATG highlighted in Figure 1B).

**Dimerization and subcellular localization of the Dcs proteins**

In order to identify complexes formed by the Dcs proteins, we used the TAP procedure (24) to pull down potential binding partners from cell lysates. Using either Dcs1-TAP or Dcs2-TAP, we observed the formation of both homo- and hetero-dimers (Figure 1C and D). The MALDI spectrum (Figure 1D) does not resolve the two CBD-tagged Dcs1 species evident in SDS–PAGE, which are evidently very similar in terms of molecular mass (Figure 1C; see also Figure 4). These data are consistent with the previous observations (2,8,29) that the related human m7G(5′)pppN-pyrophosphatase, which has a similar monomeric size, has an apparent *M<sub>r</sub>* of ~80 kDa as determined by sedimentation analysis and gel filtration.

We assessed the subcellular distribution of Dcs1 and Dcs2 using fusions carrying protein A, haemagglutinin or c-Myc tags. Fluorescence microscopy performed using Dcs1 and Dcs2 tagged with protein A revealed that both proteins are predominantly cytoplasmic (Figure 2), and thus present in the same compartment in which the degradation of mature mRNAs and the metabolism of carbohydrates takes place.
Very similar distributions were seen with the HA- and c-Myc-tagged derivatives (data not shown), thus confirming the cytoplasmic location of both proteins. These data are consistent with the previous result of human cell fractionation where the majority of the pyrophosphatase activity was found in the cytoplasmic extract, while $\%$ of the total activity was present in the nuclear fraction (8).

Dcs proteins support the cellular response to nutrient stress

We explored the potential role of Dcs proteins in the cellular response to nutrient starvation using disruption mutants constructed using a PCR strategy (30). Glucose is the preferred carbon source for budding yeast, which has an elaborate network of molecular systems for sensing changes in glucose availability and adjusting gene expression and cellular metabolism in response to such changes (31). We observed that the ability of yeast cells to respond to glucose deprivation is compromised when DCS1 is disrupted, and that this phenotype is exacerbated in the dcs1Δdcs2Δ double disruption (Figure 3A). The growth of the dcs2Δ strain was difficult to distinguish from that of the wild-type strain, and it is therefore uncertain whether this single disruption has a phenotype. Similar

Figure 1. Characterization of DCS2 mRNA 5’ ends and Dcs1/Dcs2 dimeric complexes. (A) Primer extension analysis was performed on DCS2 mRNA using total RNA isolated from S.cerevisiae BY4741 cells at mid-log phase (lane 1). In the control, total RNA was treated with RNase A before the primer extension reaction (lane 2). Each sequencing lane is labelled with the dideoxynucleotide used in the sequencing reaction. (B) Sequence of the upstream region of the DCS2 gene. The correct initiation codon leads off the N-terminal sequence confirmed by amino acid sequencing. The ATG upstream of this, which was originally thought to be the start codon of DCS2, is also shown in bold type. Potential STRE elements are in bold, while a typical promoter TATA motif is underlined. Arrowheads indicate the 5’ ends detected by the primer extension analysis. (C) A silver-stained, 10% SDS–PAGE gel shows the proteins found in the complexes purified using Dcs1–TAP and Dcs2–TAP. Lane 1, molecular mass markers (kDa); lane 2, TAP purification from strain BY4741, as negative control; lane 3, TAP purification from strain PTC197 (Dcs2–TAP); lane 4, TAP purification from strain PTC196 (Dcs1–TAP). The identities (confirmed by MALDI) of the respective proteins running in lanes 3 and 4 are indicated by labelling on the sides of the gel. Two forms of both Dcs1 and Dcs1CBD are evident (compare Figure 5A and B). CBD (calmodulin binding domain) indicates proteins that still carry part of the TAP tag. (D) Positive ion MALDI mass spectrum of the sample that was analysed in lane 4 of (C) The spectrum reveals the presence of Dcs1CBD and Dcs2 monomers, of homodimers of Dcs2 (a) and Dcs1CBD (c), as well as of the Dcs2–Dcs1CBD heterodimer (b). M, values are indicated next to the corresponding peaks.

Figure 2. Subcellular localization of Dcs1 and Dcs2 proteins in S.cerevisiae cells. Strains expressing Dcs1–TAP and Dcs2–TAP were examined by indirect immunofluorescence microscopy using the antiserum directed against protein A and FITC-conjugated anti-rabbit secondary antibodies (column 1). Nuclear DNA was stained with DAPI in the overlay images (column 2).
phenotypes for the respective strains were observed when glucose was replaced by glycerol, whereas the availability of galactose significantly reduced the impact of the disruptions. The DCS genes were found not to influence growth under conditions of oxidative or osmotic stress in this assay (Figure 3A).

We next tested whether the DCS mutations affect the adaptation of yeast to changes in the composition of the growth medium. Both the spot assays on agar plates (Figure 3B) and growth experiments in liquid media (Figure 3C) revealed marked reductions in the ability of yeast to adapt to a shift from minimal medium to YP medium or to YP medium containing glycerol. In control experiments, we found that the growth phenotypes could be suppressed by transformation of the mutant strains with the appropriate DCS1/2 expression plasmids (Figure 3B, lanes 3 and 5). Overall, we conclude from these experiments that both the nutrient stress response and the nutrient adaptation response in yeast are influenced by the DCS genes.

Regulation of DCS genes

We next investigated how changes in the available carbon source affect expression of the respective genes. Northern and western blotting were used to compare the levels of mRNA and protein, respectively, encoded by the DCS genes in the presence and absence of glucose (Figure 4A).

Removal of glucose from YPD medium switched on the synthesis of both DCS2 mRNA and Dcs2 protein. The observed increase in the mRNA signal correlates with the results of an earlier microarray study of glucose-related changes in the yeast transcriptome (32). DCS2 was also identified in a screen for cAMP-repressed genes whose expression responds to nutrient limitation (33). Induction of the synthesis of DCS2 mRNA and Dcs2 protein were also observed under conditions of osmotic, oxidative and heat stress, while cycloheximide treatment induced only DCS2 mRNA synthesis (data not shown). These results are consistent with the presence of...
trehalase activity is expressed as a ratio to the activity of control cells (OD600).

Trehalase activity was measured using the glucose oxidase/peroxidase method (26) in extracts from the wild-type strain (BY4741, filled circles), the dcs1Δ mutant Y12429 strain (closed triangles) and the dcs2Δ mutant Y12429 strain (open triangles) at different stages of growth. The increase in trehalase activity is expressed as a ratio to the activity of control cells (OD600 = 0.4), normalized to 1. The presented data represent averages of three independent experiments.

**Figure 4.** DCS1 and DCS2 mRNAs and proteins during diauxic shift and under carbon-source stress conditions. (A) Top two rows: western blots using antibodies against Dcs1 and Dcs2, respectively. Dcs2 was detected in the form of Dcs2–TAP using PAP antibodies (Sigma). Bottom two rows: northern blotting was performed using probes against DCS1 mRNA, DCS2 mRNA and 25S rRNA, respectively. (B) The samples used in (B) were taken from this culture of BY4741 at the points indicated. Cell density (filled circles) and glucose concentration (open circles) were measured on all samples. Growth was at 30°C in YPD medium. (D) Trehalase activity was measured using the glucose oxidase/peroxidase method (26) in extracts from the wild-type strain (BY4741, filled circles), the dcs1Δ strain (PTC194, open circles), the dcs2Δ strain (Y12429, closed triangles) and the dcs1Δdcs2Δ strain (PTC195, open triangles) at different stages of growth. The increase in trehalase activity is expressed as a ratio to the activity of control cells (OD600 = 0.4), normalized to 1. The presented data represent averages of three independent experiments.

STRE motifs upstream of the DCS2 transcriptional start site (Figure 1B), since STREs mediate activation of transcription in response to stress conditions. In the case of DCS1, glucose depletion resulted in weaker enhancement of mRNA levels while causing the appearance of a modified form of the Dcs1 protein that ran as a reduced mobility band on SDS–PAGE (Figure 4A and B). The addition of the alternative carbon sources trehalose and glycerol to culture medium lacking glucose (Figure 4A and B). The addition of the alternative carbon sources trehalose and glycerol to culture medium lacking glucose (Figure 4A and B). The addition of the alternative carbon sources trehalose and glycerol to culture medium lacking glucose (Figure 4A and B). The addition of the alternative carbon sources trehalose and glycerol to culture medium lacking glucose (Figure 4A and B). The addition of the alternative carbon sources trehalose and glycerol to culture medium lacking glucose (Figure 4A and B). 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orthophosphate metabolic labelling experiment using the wild-type protein treatment with \textit{dcs1} strain BY4741 (lanes 1 and 2) and the \textit{also eliminated the phosphoprotein. In further experiments, we ment of wild-type Dcs1 by the T66A form in a wild-type strain upper band intensity is lost. As shown in Figure 5D, replace-
in Figure 5C indicates that Yak1 plays a major role in the tested appropriate kinase mutants for their effects on Dcs1 tial phosphorylation sites, identifying S60 and T66. We then region of the Dcs1 protein sequence for the presence of poten-
slow running band seen in Figure 4A (Dcs1) does indeed contain a phosphorylated form of this protein (Figure 5A). Further evidence that the Dcs1 modification is a phosphate group was obtained by treating the modified protein with \(\lambda\) PPase. This treatment yielded only the faster migrating form of Dcs1 (Figure 5B, lane 3), thus confirming that Dcs1 is converted to a phosphorylated protein in response to glucose deprivation. Partial protein degradation revealed that the appearance of a slower migrating (phosphorylated) form of Dcs1 is eliminated if in the N-terminal 70 amino acids of the protein are removed (data not shown). We therefore examined this N-terminal region of the Dcs1 protein sequence for the presence of potential phosphorylation sites, identifying S60 and T66. We then tested appropriate kinase mutants for their effects on Dcs1 phosphorylation (Figure 5C). Comparison of lanes 1, 2 and 3 in Figure 5C indicates that Yak1 plays a major role in the phosphorylation of Dcs1, since in its absence >90% of the upper band intensity is lost. As shown in Figure 5D, replacement of wild-type Dcs1 by the T66A form in a wild-type strain also eliminated the phosphoprotein. In further experiments, we established that the T66A mutation does not cause either of the compromised stress-response phenotypes typical of \(dcs1\Delta\) or \(dcs1\Delta dcs2\Delta\) (data not shown). The retention of a small amount of the slower running Dcs1 band in the absence of Yak1 is likely to be attributable to a low level of phosphorylation of Dcs1 at other sites (catalysed by other kinases).

Both \textit{DCS} genes are regulatory targets of the cAMP-protein kinase \(\lambda\) pathway

Both a previous report (33) and the data presented in Figures 1 and 4 indicate that \textit{DCS2} expression is controlled by the cAMP-PKA pathway [for review, see (14)]. We also observed that \textit{DCS1} gene expression follows a similar pattern to that of \textit{DCS2} (Figure 4A and B). In order to investigate whether \textit{DCS1} gene expression is responsive to the cAMP-PKA pathway, we compared Dcs1 levels in strains containing single or combined disruptions of \textit{RIM15} and \textit{YAK1} (which encode protein kinases involved in growth regulation), \textit{TPK1-3} (which encode PKA catalytic subunits) and \textit{MSN2/MSN4} (which encode stress-inducible transcription activators that are negatively controlled by PKA). We found that Dcs1 abundance was drastically decreased in \textit{msn2/msn4} cells (Figure 5C, lane 5), while it was much increased in \textit{rim15yak1} (Figure 5C, compare lanes 4 and 3). Moreover, the strain lacking the Rim15 protein kinase, which has been described as a stimulator of STRE-controlled genes (14), showed a reduced level of Dcs1 protein (Figure 5C, lane 2). Taken together, these results indicate that expression of \textit{DCS1} is also at least partially responsive to the cAMP-PKA pathway.

Is Dcs1 a modulator of trehalase activity?

We next investigated the possibility that Dcs1 might influence the stress response by directly modulating Nth1 activity. If this were the case, we would expect purified Dcs1 protein to be capable of suppressing the enhanced Nth1 activity observed in extracts from a \(dcs1\Delta\) strain (see Figure 4D). Neither recombinant Dcs1 purified from \textit{E.coli} nor phosphorylated or non-phosphorylated forms of Dcs1 isolated from \textit{S.cerevisiae} showed any significant ability to down-regulate trehalase activity (Figure 6). This indicates that Dcs1 itself does not impose post-translational regulation on Nth1.

Functional overlap between Dcs1 and eIF4E

Since Dcs1 and the translation initiation factor eIF4E both bind to the mRNA cap, we wondered whether there is any functional interaction between the two \textit{in vivo}. This is relevant to the impact that changes in the level of 5′ m\textsuperscript{7}G-capped mRNA fragments may have on the cell. From previous work (40), we know that eIF4E is in considerable excess over eIF4G, the other core component of the cap-binding complex eIF4F, in \textit{S.cerevisiae}. This means that there is a considerable population of eIF4E potentially available in the cell that might be involved in activities other than translation. We decided to explore the possible relationship between the two proteins using a strain (YTH3) in which the eIF4E gene (\textit{CDC33}) has been placed under the control of a promoter that can be regulated via a tetracycline-responsive operator (40). This enabled us to vary the amount of eIF4E in the cell by adding doxycycline to different final concentrations (Figure 7). By means of genetic mating followed by sporulation, we generated a YTH3 \(dcs1\Delta\) strain that could be compared with the YTH3 isogenic parent. In the absence of functional Dcs1, we observed that yeast became highly

Figure 5. Dcs1 is phosphorylated in response to glucose depletion. (A) A \([^{32}P]\)orthophosphate metabolic labelling experiment using the wild-type strain BY4741 (lanes 1 and 2) and the \textit{dcs1}\Delta strain PTC194 (lane 3). \([^{32}P]\)orthophosphate was added to yeast previously grown for 45 min in phosphate-free medium in the presence (lane 1) or absence (lane 2) of glucose. The lower panel shows western blot analysis of immunoprecipitated Dcs1 proteins. The upper panel shows an autoradiograph of the same gel. (B) Dcs1 protein treatment with \(\lambda\) PPase. Lane 1, Dcs1 purified from yeast cells at stationary phase and used in lanes 2 and 3. Lane 2, Dcs1 after incubation with \(\lambda\) PPase and phosphatase inhibitor. Lane 3, Dcs1 after incubation with only \(\lambda\) PPase. (C) Western blot using anti-Dcs1 antibodies. Late log phase extracts from mutant strains were analysed in the following order: lane 1, MY1 (wild-type strain); lane 2, MY2872 (rim15\Delta strain); lane 3, MY3297 (rim15\Delta yak1\Delta strain); lane 4, MY3273 (rim15\Delta yak1\Delta tpk1-3\Delta strain); lane 5, MY2677 (mss2\Delta msn4\Delta strain); lane 6, BY4741 (wild-type strain); lane 7, PTC195 (dcs1\Delta dcs2\Delta strain). (D) Western blot using anti-Dcs1 antibodies. Late log phase extracts were prepared from PTC194 cells transformed with pRS313- DCS1 (lane 1), pRS313-DCS1-178 (lane 2) and pRS313-DCS1-196 (lane 3). Arrowheads indicate the positions of the phosphorylated form of Dcs1 protein in (A–D).
sensitive to reductions in the level of intracellular eIF4E. Indeed, at high doxycycline concentrations, the combination of the $P_{\text{tet}}\text{CDC33}$ and $dcs1\Delta$ alleles was synthetic lethal. In control experiments, we observed that the strain bearing $dcs1\Delta$ alone was not affected by doxycycline at either concentration (data not shown).

**CONCLUSIONS**

The results described in this paper indicate that the Dcs proteins influence the ability of yeast to respond to both nutrient deficiency and to nutrient availability. Most striking is the observation of two distinct modes of $DCS$ gene regulation as glucose becomes scarce during the diauxic shift. Both $DCS2$ and $DCS1$ are subject to STRE-dependent transcriptional up-regulation, while the Dcs1 protein is also subject to phosphorylation. The transcriptional regulation involves the cAMP-PKA pathway. Since we have observed that Dcs1 and Dcs2 can form a heterodimer, the induction of Dcs2 synthesis under conditions of glucose deprivation is likely to lead to the replacement of much of the homodimeric Dcs1 population by the Dcs1–Dcs2 heterodimer. We note that Dcs2 modulates the impact of Dcs1 on the nutrient stress response, whilst itself not being able to substitute for Dcs1. We have therefore identified a function for Dcs2, which was previously found not to participate in decapping (8).

Transcription of the $NTH1$ gene, which also features STREs in its promoter region, increases in response to diauxie (41,42). Yet Nth1 activity decreases under the same conditions (37),...
and it has been proposed that this is attributable to Dcs1-dependent inhibition at the protein level (39). In this context it is notable that Thevelein previously suggested that cAMP-dependent PKA-mediated phosphorylation is involved in the regulation of Nth1 activity (43), leading to the proposal that phosphorylation might be used to modulate an interaction between Dcs1 and Nth1 (39). However, we have found no evidence of direct regulation of trehalase activity by Dcs1, irrespective of the phosphorylation state of the latter protein.

An alternative interpretation is that increased levels of m7G-oligoribonucleotides, as present in dcs1 mutants, serve to trigger a stress response. Like other known stressors (37,44), they may act to induce increased Nth1 activity. The mechanism by which the cell senses enhanced levels of m7G-oligoribonucleotides is unclear, but one possibility is that the response is coupled to interactions of these competitive inhibitors with the translational apparatus (Figure 8). The observation that expression of a \( P_{NTH1}\)lacZ fusion is almost three times higher in a \( dcs1\Delta \) strain than that in an isogenic wild-type strain (39) indicates that this increased activity is at least partly due to transcriptional control, although the involvement of some element of post-transcriptional control cannot be ruled out.

The observation that reduced levels of the cap-binding protein eIF4E cause strong synthetic phenotypes in a \( dcs1\Delta \) mutant shows that a normal cellular content of eIF4E partially complements Dcs1 deficiency. The most likely explanation of this is that eIF4E can ‘buffer’ to some extent the effects of an excess of 5’ m7G-oligoribonucleotides on cellular processes (such as translation initiation). This suggests a model in which the inhibitory potential of m7G-oligoribonucleotides is modulated by the activity of (free) eIF4E (Figure 8). There seems to be an interplay between Dcs protein activity and the nutrient stress response. In the absence of pyrophosphatase activity capable of hydrolysing m7G-oligoribonucleotides, the prolonged longevity of these species causes an additionally stressed state under conditions of glucose depletion. This, in

![Figure 8. A testable model for the modulating role of Dcs1 in the yeast stress response. Increases in yeast growth rate, or stress conditions, can accelerate mRNA turnover, thus increasing the rate of generation of mRNA turnover products, including m7G-oligoribonucleotides, m7GDP and other m7G-derivatives. In the 5’→3’ pathway, decapping of deadenylated but capped mRNA yields m7GDP, which is a substrate of Dcs1. In the 3’→5’ pathway, m7G-oligoribonucleotides are the final products of exosome-catalysed degradation. Where there is a reduction in the activity of Dcs1 (and more markedly where Dcs2 activity is also reduced), m7G-oligoribonucleotides and m7GDP levels increase and compete with capped mRNA for eIF4F and thereby inhibit translation initiation. In our experiments, we studied the extreme case where \( DCS1 \) and/or \( DCS2 \) were disrupted. Due to this inhibition, or possibly via an alternative stimulatory route, cAMP-PKA-mediated activation of STRE-promoters occurs, inducing a stressed state. Dashed arrows represent pathways that generate strong inhibitors of translation. Dotted arrows indicate potential stimulatory routes for STRE-dependent transcriptional activation, with the cAMP-PKA pathway providing one such route. In the latter case, inhibition of cAMP-PKA activity has to be repressed in order for transcription to be stimulated.

turn, may induce enhanced trehalase activity in the stationary phase and compromise the cellular response to nutrient stress. It is not known whether the levels of mG-oligoribonucleotides can influence STRE-mediated regulation of stress genes via alternative routes (Figure 8). In conclusion, the Dcs1 and Dcs2 activities in yeast show the novel property of being able to modulate the nutrient stress response and this organism’s adaptation to nutrient availability. Future work will need to elucidate the relationship between mG-oligoribonucleotide turnover and parts of these responses, including the control of trehalase activity.

Finally, other organisms possess further members of the HIT motif superfamily which are closely related to _Saccharomyces cerevisiae_ Dcs1. Do these proteins constitute a subgroup whose members all manifest the same functional properties? Certainly the human DcpS protein manifests similar decapping properties to _Saccharomyces cerevisiae_ Dcs1 (9), although we do not yet know whether it too is required for the nutrient stress response. On the other hand, the cellular function of _Schizosaccharomyces pombe_ Nhm1, much of which is nuclear (10), remains unclear. We also note that despite being very similar in structure to Dcs1, _Saccharomyces cerevisiae_ Dcs2 is clearly functionally distinct. There may therefore be significant diversity of function within the family of Dcs-like HIT proteins.

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