Negative regulation of Gcn4 and Msn2 transcription factors by Srb10 cyclin-dependent kinase

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The budding yeast transcriptional activator Gcn4 is rapidly degraded in an SCF-Cdc4-dependent manner in vivo. Upon fractionation of yeast extracts to identify factors that mediate Gcn4 ubiquitination, we found that Srb10 phosphorylates Gcn4 and thereby marks it for recognition by SCF-Cdc4 ubiquitin ligase. Srb10 is a physiological regulator of Gcn4 stability because both phosphorylation and turnover of Gcn4 are diminished in srb10 mutants. Gcn4 is almost completely stabilized in srb10Δ pho85Δ cells, or upon mutation of all Srb10 phosphorylation sites within Gcn4, suggesting that the Pho85 and Srb10 cyclin-dependent kinases (CDKs) conspire to limit the accumulation of Gcn4. The multistress response transcriptional regulator Msn2 is also a substrate for Srb10 and is hyperphosphorylated in an Srb10-dependent manner upon heat-stress-induced translocation into the nucleus. Whereas Msn2 is cytoplasmic in resting wild-type cells, its nuclear exclusion is partially compromised in srb10 mutant cells. Srb10 has been shown to repress a subset of genes in vivo, and has been proposed to inhibit transcription via phosphorylation of the C-terminal domain of RNA polymerase II. We propose that Srb10 also inhibits gene expression by promoting the rapid degradation or nuclear export of specific transcription factors. Simultaneous down-regulation of both transcriptional regulatory proteins and RNA polymerase may enhance the potency and specificity of transcriptional inhibition by Srb10.

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Precise modulation of intracellular protein concentration is an important means by which diverse cellular processes are regulated. One way cells effectively achieve this is through proteolysis of key regulatory proteins. The ubiquitin system is the major cytoplasmic pathway by which proteins are degraded. Ubiquitin-mediated degradation of cellular proteins involves attachment of ubiquitin chains to substrate proteins, which are subsequently targeted for degradation by the 26S proteasome (Ciechanover et al. 2000). Assembly of a multiubiquitin chain upon a substrate typically requires three classes of enzymes: the ubiquitin-activating enzyme [E1], a ubiquitin-conjugating enzyme [E2], and a ubiquitin ligase [E3]. The E1 enzyme activates ubiquitin in an ATP-dependent manner by linking to its C terminus via a thioester bond. The E2 enzyme accepts the thioesterified ubiquitin and transfers it to a lysine residue of the target protein. This step usually requires assistance from an E3 enzyme. There are potentially 13 E2-like proteins in budding yeast. In contrast, there are two distinct classes of E3s: the HECT domain family and the RING domain family (Deshaies 1999; Seol et al. 1999; Jackson et al. 2000). The RING domain family of E3s includes multimeric complexes such as SCF and anaphase-promoting complex/cyclosome (APC/C), as well as single subunit ligases like Ubr1. Whereas HECT E3s act as a direct intermediary by accepting ubiquitin from E2 and transferring it to substrate, RING-based E3s are thought to catalyze ubiquitination by enabling the direct transfer of ubiquitin from E2 to substrate (Seol et al. 1999). Regardless of their mechanism of action, E3s typically bind directly to E2 and to substrate, suggesting that they provide substrate specificity in cellular ubiquitination reactions.

One salient example of how proteolysis can provide a regulatory switch stemmed from genetic analysis of the G1/S transition in budding yeast (Schwob et al. 1994). Cells harboring temperature-sensitive mutations in SKP1, CDC53, and CDC4 arrest in G1 phase at the nonpermissive temperature because they fail to degrade...
the S-phase cyclin/cyclin-dependent kinase (CDK) inhibitor Sic1 (Schwob et al. 1994; Bai et al. 1996). Subsequent in vitro reconstitution of Sic1 ubiquitination led to the identification of SCF\textsuperscript{Cdc4}, the prototype of the SCF (for Skp, Cdc53/cullin, F-box receptor) family of ubiquitin ligases [Feldman et al. 1997; Skowrya et al. 1997; Verma et al. 1997c]. Recently, Hrt1 [also known as Roc1 and Rbx1], an essential fourth subunit of the SCF complex, was identified (for review, see Deshaias 1999). The SCF family of ubiquitin ligases is potentially large given that the yeast genome encodes at least 17 potential F-box receptor subunits [Patton et al. 1998b], and at least two other SCF complexes—SCF\textsuperscript{Cin1} and SCF\textsuperscript{Mec30}—have been identified in budding yeast [Patton et al. 1998a]. Cdc34 appears to be the primary E2 enzyme that interacts with SCF complexes and catalyzes ubiquitination of their substrates in budding yeast.

Besides Sic1, the CDK inhibitor Far1 [Henchoz et al. 1997] and the replication initiation protein Cdc6 [Drury et al. 1997; Elsasser et al. 1999] have been shown to be substrates of SCF\textsuperscript{Cdc4}. A common feature in the ubiquitination of SCF\textsuperscript{Cdc4} substrates is that they must be phosphorylated by the major cell cycle CDK, Cdc28 [Henchoz et al. 1997; Verma et al. 1997c; Elsasser et al. 1999]. Phosphorylation appears to serve as a general signal that promotes binding of the F-box receptor Cdc4 to the substrates [Feldman et al. 1997; Skowrya et al. 1997]. To investigate the generality of the Cdc34/SCF\textsuperscript{Cdc4} pathway, we initiated biochemical analysis of the roles of these proteins in Gen4 ubiquitination. Gen4, a transcription activator involved in the regulation of amino acid and purine biosynthetic genes [Hinnebusch 1992], is very unstable, and its degradation is dependent on Cdc34 and proteasome function [Kornitzer et al. 1994]. Very recently, it was shown that Gen4 is stabilized in cdc4, cdc53, and skp1 temperature-sensitive mutants, and in pho85?Δ cells [Meimoun et al. 2000]. This suggests that SCF\textsuperscript{Cdc4} contributes to the rapid degradation of Gen4 in vivo and that a CDK other than Cdc28 is involved in Gen4 degradation. However, there is no biochemical evidence to date that either SCF\textsuperscript{Cdc4} or Pho85 directly promotes ubiquitination of Gen4.

Here we provide evidence that the Srb10 CDK of the SRB/mediator complex phosphorylates both Gen4 and the multistress response transcription factor Msn2. Whereas Srb10 targets Gen4 for SCF\textsuperscript{Cdc4}-dependent degradation, it helps enforce the nuclear exclusion of Msn2. It has been proposed that Srb10 negatively regulates transcription of certain genes by binding and phosphorylation of the C-terminal domain (CTD) of the largest subunit of RNA polymerase II [Hengartner et al. 1998]. Our results suggest that Srb10 can also repress the transcription of specific genes by directly antagonizing transcriptional activators.

**Results**

**Ubiquitination of Gen4 in yeast extracts**

Our in vitro studies on Gen4 ubiquitination were prompted by the observation that Gen4 turnover in vivo depends on Cdc34 [Kornitzer et al. 1994]. As a first step toward understanding the mechanism and regulation of Gen4 turnover, we set out to reconstitute Gen4 ubiquitination in vitro. Ubiquitination of \textsuperscript{35}S-methionine-labeled Gen4 was evaluated in G1-cyclin-depleted whole-cell yeast extracts as described for Sic1 [Verma et al. 1997c]. Although ubiquitination of Sic1 required the addition of GST–Cln2 to supply Cdc28 kinase activity [Fig. 1A, lanes 6–7], ubiquitination of Gen4 did not (lanes 2,3). Multiubiquitination of both substrates was confirmed by the addition of methyl-ubiquitin [lanes 4,8], a chain-terminating derivative of ubiquitin. Concomitantly with ubiquitination, Gen4 exhibited a characteristic molecular weight up-shift upon incubation in yeast extract (cf. Fig. 1A, lanes 1,2). To test if this modification was caused by phosphorylation, we first incubated Gen4\textsuperscript{HA} with a yeast extract fraction enriched for this activity, and then immunoprecipitated Gen4\textsuperscript{HA} and treated it with calf intestinal alkaline phosphatase (CIAP). As shown in Figure 1B, CIAP treatment reversed the molecular weight shift of Gen4, indicating that it arises from phosphorylation.

The results above suggested that Gen4 might be targeted for ubiquitination by a novel pathway. Alternatively, a protein kinase besides Cdc28 may be able to direct Gen4 to the SCF pathway. To distinguish between these possibilities, we fractionated whole cell extract from mutant cdc4ts cells into a protein kinase fraction and an SCF-containing fraction [the SCF fraction lacks the E2 Cdc34; Verma et al. 1997c]. As shown in Figure 1C, multiubiquitination of Gen4 occurred only when both yeast fractions and exogenous Cdc34 and Cdc4 were added. Since ubiquitination of Sic1 did not require the kinase fraction [data not shown], we concluded that Gen4 must be phosphorylated by an unknown protein kinase before it can be ubiquitinated by the SCF\textsuperscript{Cdc4}/Cdc34 pathway.

**Identification of the Srb10 subunit of RNA polymerase II holoenzyme as a Gen4 kinase**

To identify the activity required for Gen4 ubiquitination in vitro, we first attempted to purify this activity by conventional column chromatography. We used both Gen4 kinase assays and ubiquitination assays to monitor this activity during purification. Unfortunately, we did not purify enough activator to obtain protein sequence. However, upon partial purification [Fig. 2A], we identified a high-molecular-weight Gen4 kinase activity that cofractionated with the ubiquitination-promoting activity [Fig. 2B]. Note that although there are two contaminating low-molecular-weight Gen4 kinases that peak in fractions 11 and 15, neither of these fractions can sustain ubiquitination of Gen4 by SCF\textsuperscript{Cdc4}. We suspected the kinase activity was a CDK because the Cdc28 CDK is required for SCF\textsuperscript{Cdc4}-dependent ubiquitination of Sic1, Far1, and Cdc6. Given that Kin28 and Srb10, both of which are components of the RNA polymerase II holoenzyme [Myer and Young 1998], are the only CDKs
known to reside in a high-molecular-weight complex, we tested whether either of these kinases was responsible for the purified activity. Extract fractions enriched for the high-molecular-weight Gcn4 kinase activity were prepared from SRB10 and srb10/H9004 cells and tested for their ability to support Gcn4 ubiquitination by Cdc34 and purified recombinant SCF\textsubscript{Cdc4} complex. As shown in Figure 2C, the kinase fraction from SRB10 but not from srb10Δ cells supported Gcn4 ubiquitination. In contrast, we did not observe any defect when the kinase fraction was prepared from a kin28Δ strain [data not shown]. Taken together, these observations suggest that Gcn4 is directly ubiquitinated by Cdc34/SCF\textsubscript{Cdc4}, and that the Srb10 subunit of RNA polymerase II holoenzyme is the predominant protein kinase in yeast extract that is able to specify ubiquitination of Gcn4 by this pathway.

Srb10, but not Kin28, phosphorylates Gcn4 in vitro and targets it for SCF-dependent ubiquitination

Because both Srb10 and Kin28 associate with the RNA polymerase II holoenzyme complex [Myer and Young 1998], we next tested whether either Srb10 or Kin28 could directly phosphorylate Gcn4 by immunoprecipitating each of the kinases from yeast whole-cell extracts and performing kinase assays using purified Gcn4 expressed in Escherichia coli. Anti-\textit{myc} immunoprecipitates from SRB10\textit{myc} cells exhibited potent Gcn4 kinase activity, whereas those from \textit{KIN28\textit{myc}} cells had little or no activity [Fig. 3A, lanes 2–4]. As expected [Hengartner et al. 1998], GST–CTD (C-terminal domain) was phosphorylated by both Srb10\textit{myc} and Kin28\textit{myc} immunoprecipitates [lanes 6–8]. GST–CTD kinase activity observed in the Kin28\textit{myc} immunoprecipitate was not caused by contaminating Srb10 because similar activity was recovered from an srb10Δ strain [lane 8]. These data demonstrate that the high-molecular-weight Gcn4 kinase activity that we enriched for is owing to Srb10, not to Kin28. To further confirm the specificity of the kinase activity present in Srbl0\textit{myc} immunoprecipitates, we evaluated the Gcn4 kinase activity of immunoprecipitates prepared from an srb10-3\textit{myc} strain, which harbors an active-site mutant form of the Srb10 kinase that is incorporated into the RNA polymerase II holoenzyme [Liao et al. 1995]. Although similar amounts of Srb protein were present in Srbl0\textit{myc} and Srb10-3\textit{myc} immunoprecipitates [Fig. 3B, lower panel], only the former phosphorylated Gcn4 [Fig. 3B, upper panel].

Figure 1. Gcn4 ubiquitination in yeast extracts requires SCF components and a protein kinase activity other than Cdc28. (A) [\textsuperscript{35}S]Met-thionine-labeled Gcn4 (lanes 1–4) or Scl1 (lanes 5–8) synthesized by in vitro translation were incubated with G1-cyclin-depleted yeast whole cell extracts from RJD885 strain in the absence [lanes 2, 6] or the presence [lanes 3, 7] of purified GST–Cln2. Uba1, Cdc34, and SCF activities were supplied by the yeast extracts. Lanes 1 and 5 contained only input substrates. Methylated ubiquitin [me-ubiquitin] was used instead of wild-type ubiquitin in lanes 4 and 8. Encircled P refers to phosphorylated forms of Scl1 and Gcn4. All samples in panels A–C were evaluated by SDS-PAGE followed by autoradiography. (B) [\textsuperscript{35}S]Met-thionine-labeled Gcn4\textsubscript{4\textit{Ath}} translation product was phosphorylated by the DEAE flow-through fraction of RJD885 whole cell extract (see Materials and Methods). An aliquot of the reaction was immunoprecipitated using 12CA5 antibody and mock-treated [lane 2] or treated with calf intestinal alkaline phosphatase (CIAP) in the absence [lane 3] or the presence [lane 4] of phosphate inhibitor as described [Verma et al. 1997c]. Immunoprecipitated input Gcn4\textsubscript{4\textit{Ath}} is shown in lane 1. (C) Ubiquitination of Gcn4 in fractionated yeast extracts from a cdc4\textsuperscript{ts} strain [RJD893]. Extract fractions lacking Cdc34 but supplying either SCF components or a kinase activity were used [see Materials and Methods]. A complete reaction [lane 6] included a kinase fraction [\textsim 10 \mu g], a ubiquitination fraction [\textsim 15 \mu g], Cdc34 [100 ng], and insect cell lysate containing baculovirus-expressed Cdc4 [\textsim 4 \mu g total protein; Verma et al. 1997c]. In lanes 1–5, one or more of the above components was omitted, as indicated. Insect cell lysate containing baculovirus-expressed Cdc28 was used as a specificity control [lane 7] for the Cdc4 lysate.
In our initial experiments to test the ability of Srb10 to phosphorylate Gcn4, we used a strain (Z689) that expressed HA-tagged Srb11, the cyclin partner of Srb10. Anti-HA immunoprecipitates from Z689 cells contained Gcn4 kinase activity similar to that shown in Figure 3A (data not shown). [35S]methionine-labeled Gcn4, after preincubation with an Srb11 HA immunoprecipitate, was ubiquitinated in the presence of purified Uba1, Cdc34, and SCF<sub>Cdc4</sub> (Fig. 3C, lane 4). This reaction was specific because it was dependent on all of the above components and ubiquitin (lanes 2, 5–8). Ubiquitination of Gcn4 by the Cdc34/SCF<sub>Cdc4</sub> pathway was also reconstituted with Gcn4 produced in <i>E. coli</i> (Fig. 3A) and purified Uba1, Cdc34, and SCF<sub>Cdc4</sub> (Fig. 3D). The greater reaction efficiency observed in this experiment was most likely owing to the use of SCF preparations containing Hrt1. Taken together, these results indicate that phosphorylation of Gcn4 by Srb10 promoted its recognition and ubiquitination by SCF<sub>Cdc4</sub>.

**Srb10 phosphorylates Gcn4 and regulates its stability in vivo**

To investigate the physiological relevance of our biochemical data, we first tested whether Srb10 influences the phosphorylation state of Gcn4 in vivo by evaluating the electrophoretic mobility of Gcn4 and its phosphorylation state in pulse-radiolabeled wild-type and <i>srb10</i> mutant cells. Gcn4 was phosphorylated in vivo in an Srb10-dependent manner, as evidenced by a phosphatase-sensitive (Fig. 4A, right panel) molecular weight (MW) up-shift that was diminished in <i>srb10</i> mutant cells (Fig. 4A, left panel). This combined with our in vitro observations, strongly suggests that the Srb10 CDK is a physiological Gcn4 kinase. If Srb10 targets Gcn4 for turnover by the SCF<sub>Cdc4</sub> pathway, we predicted that Gcn4 should be stabilized in <i>srb10</i> mutants. To test this possibility, we carried out pulse-chase experiments to measure the turnover rate of Gcn4<sub>myc9</sub> in wild-type and <i>srb10</i> mutants. It is important to note that the tagged Gcn4 was expressed from the endogenous <i>GCN4</i> locus, under the control of the native promoter. Consistent with previously published results (Kornitzer et al. 1994; Meimon et al. 2000), Gcn4<sub>myc9</sub> was rapidly degraded in wild-type cells, with a half-life of 2.5–5 min (Figs. 4B, 5B). In contrast, Gcn4<sub>myc9</sub> was moderately stabilized in both <i>srb10-3</i> and <i>srb10A</i> mutants with a half-life of 10–12 min (Figs. 4B, 5B). Consistent with this difference in half-life, the steady-state level of Gcn4<sub>myc9</sub> is about two-fold higher in <i>srb10</i> mutant cells compared to wild-type (Fig. 4C). Careful examination of autoradiograms revealed that the phosphorylation-dependent up-shifted form of Gcn4<sub>myc9</sub> was strongly reduced, but not eliminated, in <i>srb10</i> cells (Figs. 4B, 5A, 5B). Furthermore, Gcn4<sub>myc9</sub> was more effectively stabilized in <i>cdc34</i> cells at the restrictive temperature (<i>t</i><sub>1/2</sub> > 20 min; Fig. 4D). We thus inferred that there might be additional protein kinases that can target Gcn4 for degradation via the Cdc34/SCF<sub>Cdc4</sub> pathway.

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parent rate of Gcn4 degradation. To test this, we directly compared the rates of Gcn4\textsuperscript{myc9} degradation in strains that expressed Gcn4\textsuperscript{myc9} from a GAL promoter with the strains that expressed Gcn4\textsuperscript{myc9} from its own promoter. Indeed, when Gcn4\textsuperscript{myc9} was overproduced, its rate of degradation appeared very similar in wild-type and srb10\textalpha cells (Fig. 5A, top panels). At normal levels of expression, however, Gcn4 was clearly stabilized in an srb10\textalpha mutant (Fig. 5A, bottom panels). Immunoblotting confirmed that the steady-state levels of Gcn4\textsuperscript{myc9} were at least fivefold higher in GAL-driven Gcn4\textsuperscript{myc9} strains compared with native-promoter-driven Gcn4\textsuperscript{myc9} strains (data not shown). Interestingly, we also noticed a progressive, SBR10-independent post-translational modification of Gcn4\textsuperscript{myc9} when it was overproduced (Fig. 5A, arrowhead; note that Gcn4 expressed from its own promoter does not show the same extent of progressive modification in srb10\alpha cells), suggesting that the overexpressed protein is more susceptible to a protein kinase that normally does not act (or acts poorly) upon endogenously expressed Gcn4.

We evaluated Gcn4\textsuperscript{myc9} stability in several other CDK mutants in an attempt to address whether multiple CDKs play a redundant role in Gcn4 turnover. Gcn4\textsuperscript{myc9} degradation was unaffected in ctk1\Delta and α-factor-arrested cells, suggesting that neither Ctk1 (a CDK homolog that phosphorylates CTD) nor Cdc28 contributed to turnover (data not shown). However, we found that Gcn4\textsuperscript{myc9} was moderately stabilized in a pho85\Delta mutant with a half-life of about 20 min [Fig. 5B], similar to what was reported by Meimoun et al. (2000). To test whether Srb10 and Pho85 contribute independently to Gcn4 turnover, we tested Gcn4\textsuperscript{myc9} stability in an srb10\Delta pho85\Delta double mutant strain. As shown in Figure 5B, Gcn4\textsuperscript{myc9} was further stabilized in srb10\Delta pho85\Delta cells with a half-life >40 min, similar to what was observed in cdc34\textalpha cells at 37°C [Fig. 4D]. Because srb10 and pho85 mutations had an additive effect, we conclude that at least two CDKs (Srb10 and Pho85) contribute to rapid Gcn4 turnover in vivo.

Srb10-dependent degradation of Gcn4 is not regulated by amino acid starvation

Because stabilization of Gcn4 under amino acid starvation conditions is achieved at least in part by the down-regulation of Pho85 activity [Meimoun et al. 2000], we wanted to test whether Srb10-dependent degradation of Gcn4 was regulated in a similar fashion. To test this, we
imposed amino acid starvation conditions by culturing cells in minimal medium lacking a specific amino acid for which the cells were auxotrophic and then measured 

Gcn4 stability under starved and unstarved conditions in wild-type, srb10Δ, and pho85Δ cells. Although Gcn4myc9 was significantly stabilized upon amino acid starvation with a half-life of about 20 min (Fig. 6A), it is further stabilized in srb10Δ cells with a half-life of about 40 min (Fig. 6B). Furthermore, Srb10-dependent phosphorylation of Gcn4myc9 (as indicated by the molecular weight shift) was not diminished upon starvation (Fig. 6A,B), suggesting that it is not regulated by amino acid starvation. Whereas amino acid starvation significantly stabilized Gcn4myc9 further in srb10Δ cells, it only increased Gcn4myc9 half-life slightly in pho85Δ cells (Fig. 6C). Taken together, these results suggest that Srb10 is not involved in the starvation-regulated aspect of Gcn4 turnover, and down-regulation of Pho85 is the primary means of stabilizing Gcn4 upon amino acid starvation.

CDK phosphorylation is required for rapid Gcn4 turnover in vivo

The protein sequence of Gcn4 reveals five putative CDK phosphorylation sites (S/TP): S17, T61, T105, T165, and S218. Our analyses of Gcn4 ubiquitination in vitro and stability in vivo suggest that these sites may be important for rapid Gcn4 turnover. To test this possibility, we phosphorylated purified Gcn4 using Srb10myc9 immunoprecipitates and subjected the in vitro phosphorylated Gcn4 to mass spectrometry analysis. All phosphopeptides that were recovered contained one of the five consensus S/TP sites. Moreover, direct sequencing confirmed that four of the consensus sites (S17, T61, T105, and S218) were phosphorylated. The exact site of phosphorylation on T165-containing peptides could not be determined, but was presumed to be T165. To analyze whether any of these sites is important for Gcn4 turnover, we first mutated each of the five sites by site-directed mutagenesis, and produced mutant forms of Gcn4...
protein by in vitro translation. We then tested the ability of the five single and a quintuple (3T2S) phosphorylation-site mutants of Gcn4 to serve as substrates for SCF^{Cdc4}-dependent ubiquitination in vitro. Although we did not observe a marked defect for any of the single-site mutants, ubiquitination of the 3T2S quintuple mutant was greatly diminished (Y. Chi and R. Deshaies, unpubl.). Based on our in vitro data, we constructed a strain in which the wild-type GCN4 locus was replaced with gcn4–3T2Smyc9, and the stability of the encoded mutant protein was tested by pulse-chase analysis. As shown in Figure 7B, Gcn4–3T2Smyc9 was very stable in vivo, and there was no appreciable degradation during the 40-min chase period. Noticeably, the mutant protein did not exhibit the characteristic molecular weight shift caused by Srb10-dependent phosphorylation.

The gcn4–3T2Smyc9 strain did not show any obvious morphology or growth defect compared with wild type. Since we used a C-terminally myc9-tagged version of GCN4 for all our in vivo studies, we were concerned that the myc9 tag might render Gcn4 inactive. To test the activities of various Gcn4 derivatives in our strains, we employed a starvation plate assay similar to that described by Natarajan et al. [1999]. Cells were streaked on
a control plate with normal leucine supplement and a plate with excess leucine, which causes isoleucine (Ile) and valine (Val) starvation (Niederberger et al. 1981). This assay monitors Gcn4-dependent activation of Ile and Val biosynthetic genes. As shown in Figure 7A, all strains [tagged or untagged GCN4] except gcn4 deletion mutants grew on −Ile, −Val plates equally well as on control plates, suggesting that the myc9 tag did not diminish the function of Gcn4. Furthermore, normal growth of gcn4−3T2Smyc9 cells on the starvation plate suggests that CDK phosphorylation is not essential for Gcn4 function.

**Srb10 phosphorylates Msn2 and influences its localization**

Microarray analysis of genomewide gene expression in the srb10-3 mutant reveals 173 genes whose expressions are up twofold or more than wild type, and a significant fraction of these genes are involved in stress response and diauxic shift (Holstege et al. 1998). Msn2 and Msn4 are two partially redundant zinc-finger transcription factors that activate expression of a suite of stress-response genes (Martinez-Pastor et al. 1996; Schmitt and McEntee 1996; Boy-Marcotte et al. 1998), including those induced in srb10-3 cells. Based on our analysis of Gcn4 regulation, we hypothesized that Srb10 might also directly repress the activities of transcription factors, including Msn2 and Msn4, by targeting them for degradation by the SCF pathway. To this end, we tested the ability of Msn2 to serve as a substrate for Srb10 in vitro. Using an assay similar to that shown in Figure 3A, we found Srb10 to be a potent kinase for recombinant GST–Msn2 (Fig. 8A, lane 6). Furthermore, phosphorylated GST–Msn2 was efficiently ubiquitinated by SCFCdc4 [data not shown]. However, pulse-chase analysis revealed that Msn2myc9 was a stable protein in wild-type cells with a half-life of more than 1 h [data not shown]. Interestingly, when pulsed-labeled at 27°C and chased under heat stress conditions (37°C), Msn2 was rapidly (within 5 min) phosphorylated in an Srb10-dependent manner [Fig. 8B]. Taken together, these data suggest that Srb10 phosphorylates Msn2 in vivo in response to stress. However, Srb10-dependent phosphorylation did not appear to target Msn2 for degradation via the Cdc34/SCF^Cdc4^ pathway.

Recent evidence suggests that Msn2 and Msn4 are regulated by shuttling between the cytoplasm and the nucleus in response to environmental conditions. Under normal conditions, Msn2 and Msn4 are retained in the cytoplasm, presumably by a combination of active nuclear export and cytoplasmic anchorage (Gorner et al. 1998; Beck and Hall 1999). To test whether Srb10 influences the localization of Msn2/Msn4, we monitored Msn2 localization in Msn2myc9 strains by indirect im-
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Figure 8. Msn2 is phosphorylated in an Srb10-dependent manner in vitro and in vivo, and its localization is influenced by Srb10. (A) Kinase assays were carried out with purified GST (lanes 1, 2), GST–Gcn4 (lanes 3, 4), and GST–Msn2 (lanes 5, 6) using immunoprecipitates from SRRB10 myc9 strain (YC17, lanes 2, 4, 6) and sb10-3 myc9 strain (YC7, lanes 1, 3, 5) as described in Figure 3A. (B) Cells from MSN2 myc9 (YC84, lanes 2–4) and MSN2 myc9 sb10-3 (YC124, lanes 6–8) strains grown at 27°C were pulse-labeled with Tran35S label at 27°C for 5 min, and chased at 37°C for 5 min. Aliquots (1/3 before the heat shock and 2/3 after the heat shock) of the cultures were subjected to immunoprecipitation with 9E10 antibody. Immunoprecipitates from the heat shock samples were divided and either mock-treated (lane 3, 7) or treated (lanes 4, 8) with 2 U CIAP for 30 min at 37°C. Untagged strains Z719 (lane 1) and Z690 (lane 5) were used as controls. (C) Localization of Msn2 myc9 in wild-type (YC84, a), msn5Δ (YC107, b), and sb10-3 (YC124, c) cells was evaluated under resting conditions by indirect immunofluorescence with 9E10 antibody. DAPI was used to stain cell nuclei. An sb10-3 strain containing untagged MSN2 (Z690) was used as control (d).

munofluorescence. As expected, Msn2 myc9 was mostly cytoplasmic in wild-type cells under unstressed conditions [Fig. 8C, panel a, Gorner et al. 1998]. Interestingly, Msn2 myc9 was localized to the nucleus in a subpopulation (15%–30%) of unstressed sb10-3 cells [Fig. 8C, panel c] and sb10Δ cells [data not shown]. Although we do not understand why Msn2 was mislocalized in only a fraction of sb10Δ cells, our results are consistent with the elevated expression of MSN2/MSN4-dependent genes in sb10-3 cells as determined by microarray analysis of a population of cells [Holstege et al. 1998].

Because Msn5 has been shown to be a nuclear exporter for Pho4 that has been phosphorylated by Pho85 [Kaffman et al. 1998], we next tested whether Msn5 was also required for the export of Msn2. Strikingly, Msn2 myc9 was concentrated in the nucleus in >90% of msn5Δ cells under normal conditions [Fig. 8C, panel b], suggesting that Msn5 is the primary export receptor for Msn2.

Discussion

SCF ubiquitin ligases have been implicated in the regulation of multiple transcription factors, including β-catenin, Gcn4, and Met4 [Deshaies 1999]. Here, we provide four major lines of evidence that the Srb10 CDK complex of the SRB/mediator module of the RNA polymerase II holoenzyme contributes to Gcn4 instability by phosphorylating Gcn4 and thereby targeting it to SCFCdc4. First, a high-molecular-weight Srb10-containing complex was purified as the most prominent activity in yeast extract that sustains SCFCdc4-dependent ubiquitination of Gcn4. Second, immunopurified Srb10 phosphorylated recombinant Gcn4, thereby rendering it a substrate for recombinant SCFCdc4. Third, Gcn4 was phosphorylated in an SRRB10-dependent manner in vivo. Fourth, Gcn4 was partially stabilized in sb10 mutants. Taken together, our observations indicate that Gcn4 is a physiological target of Srb10, and raise the intriguing possibility that recruitment of SRB/mediator may limit how long a transcriptional regulatory protein can occupy a promoter.

Regulation of Gcn4 by Srb10 and Pho85

During the course of our work, Meimoun et al. (2000) reported that Pho85 is required for rapid turnover of Gcn4. Although they show that Gcn4 is phosphorylated in vitro by recombinant Pcl1/Pho85, there is no direct evidence that this modification triggers its ubiquitination in vitro by recombinant Pcl1/Pho85. Meimoun et al. (2000) degradation through an indirect effect on SCFCdc4 activity. Nevertheless, we favor the hypothesis that both Srb10 and Pho85 directly promote Gcn4 turnover, and that the kinases act independently to target Gcn4 degradation, based on the observation that Gcn4 was modestly stabilized in pho85Δ and sb10Δ cells, but strongly stabilized in pho85Δ sb10Δ double mutants. Our observations coupled with those of Meimoun et al. suggest that it may be possible to generate more sophisticated regulatory controls by making the SCF-dependent turnover of a substrate require its phosphorylation by two or more kinases. Because distinct protein kinases can target different substrates to a single type of SCF complex (e.g., G1/CDK target Sic1, whereas either Srb10 or Pho85 targets Gcn4, to the SCFCdc4 complex), and because there are potentially 17 different types of SCF complexes and 120 protein kinases in yeast, combinatorial interactions between these elements can potentially generate a staggering array of regulatory controls.
Recent evidence suggests that amino acid starvation stabilizes Gcn4 through the negative regulation of Pho85-dependent phosphorylation (Meimoun et al. 2000). Interestingly, we found that Gcn4 was only moderately stabilized upon starvation and was further stabilized in srb10Δ cells. Moreover, Srb10-dependent phosphorylation of Gcn4 remained intact in amino acid-starved cells [Fig. 6A,B]. This indicates that whereas Pho85 is down-regulated upon starvation, Srb10-dependent degradation of Gcn4 persists. It appears that Srb10 and Pho85 contribute to SCFCdc4-dependent degradation of Gcn4 in different ways, either by targeting different pools of Gcn4 or by responding to different cellular signals.

If Srb10 negatively regulates Gcn4 by promoting its turnover, one would expect to see increased expression of Gcn4 target genes in srb10 mutants. Although microarray analysis of genomewide gene expression in the srb10-3 mutant reveals 173 genes whose expressions are up twofold or more compared to wild type [Holstege et al. 1998], only two of these are potential Gcn4 targets. This suggests that srb10 mutation does not lead to global derepression of Gcn4 target genes under normal growth conditions. We provide the following possible reasons to explain the microarray data. First, stabilization of Gcn4 in srb10 mutants only leads to an ~1.5-2-fold increase in the steady-state level of Gcn4 [Fig. 4C]. Compensatory regulatory mechanisms, such as redundant kinases that promote Gcn4 turnover and the well-characterized Gcn4 translational controls, may serve to limit the accumulation of Gcn4 in srb10 cells. The former possibility is supported by the observation that the 3T2S mutations have a greater effect on Gcn4 stability than does the pho85 srb10 double disruption [cf. Figs. 5B, 7B]. Second, Srb10 may have self-canceling positive and negative effects on Gcn4. It is possible that phosphorylation of Gcn4 by Srb10 stimulates its activity while also promoting its turnover. If so, the loss of Srb10-dependent phosphorylation of Gcn4 may not only stabilize the protein but also decrease its potency as a transcriptional activator. Third, pleiotropic effects of srb10 mutation, such as derepression of the stress response genes [Holstege et al. 1998] and slow growth, may indirectly alter the expression profile of Gcn4 target genes. Despite the absence of major change in the transcriptional output of Gcn4 in srb10 mutants, it is evident that Srb10 negatively regulates Gcn4 by promoting its rapid turnover.

Regulation of Msn2 and Msn4 by Srb10

The transcriptional activities of the partially redundant Msn2 and Msn4 proteins are tightly regulated. In resting cells these factors are inactive, whereas in stressed cells they promote transcription of a large suite of genes [Martinez-Pastor et al. 1996; Schmitt and McEntee 1996; Boy-Marcotte et al. 1998]. Srb10 is implicated in the regulation of Msn2/Msn4, in that multiple transcriptional targets of these proteins are increased in srb10-3 cells [Holstege et al. 1998]. Msn2 was phosphorylated by Srb10 in vitro, and the phosphorylation state of Msn2 was modulated by Srb10 in vivo [Fig. 8A,B]. In contrast to Gcn4, however, Msn2 was stable in resting or stressed cells, regardless of whether it was localized to the cytoplasm or to the nucleus (data not shown).

Activation of Msn2 in stressed cells is linked to its translocation from the cytoplasm to the nucleus [Gorner et al. 1998]. Whereas Msn2 is exclusively cytoplasmic in resting wild-type cells, Msn2 was primarily nuclear in 15%–30% of srb10 cells [Fig. 8C]. Although our data show that SRB10 can influence Msn2 localization, other factors, including the TOR and cAMP-dependent protein kinases, govern the nucleocytoplasmic shuttling of Msn2 in stressed cells [Gorner et al. 1998; Beck and Hall 1999], and regulated trafficking of Msn2 still occurred properly in the majority of srb10 cells [data not shown]. The underlying basis for the heterogeneity of srb10 cell populations is unclear. Perhaps the majority of srb10Δ cells adapt to chronic Msn activity in the nucleus by inducing factors that partially bypass Srb10’s role in Msn2 export. Regardless of the exact relationship between stress, Msn2/Msn4, and Srb10, the most parsimonious hypothesis is that the global induction of Msn target genes in srb10-3 cells [Holstege et al. 1998] results, at least in part, from failure to properly phosphorylate and thereby promote Msn5-dependent export of Msn2/Msn4 from the nucleus. Mapping and mutation of Srb10 phosphorylation sites in Msn2 will be needed to vigorously establish a direct role for Srb10 in the nuclear export of Msn2.

Implications for negative regulation of transcription by SRB/mediator: beyond CTD phosphorylation

The Srb10/Srb11 CDK/cyclin pair is part of a heterotrimeric module, which associates with the multiprotein complex termed SRB/mediator (Myer and Young 1998). Current models suggest that the Srb10 module functions through CTD of RNA polymerase II, in part by controlling its phosphorylation [Hengartner et al. 1998]. Both genetic and microarray analyses indicate that Srb10 serves as a negative effector of specific transcriptional programs [Carlson 1997; Holstege et al. 1998]. These observations pose a key question, namely, how does Srb10 activity selectively repress the expression of a small subset of the genome? Biochemical studies revealed that phosphorylation of CTD by Srb10 blocked the recruitment of RNA polymerase holoenzyme to promoter DNA [Hengartner et al. 1998]. It was postulated that gene-specific control by Srb10 might be mediated by promoter-bound factors that influence whether or not Srb10/Srb11 phosphorylates CTD before a promoter-bound preinitiation complex is formed.

An expanded view for how Srb10 negatively regulates transcription of selected genes is supported by our work. Specifically, we propose that Srb10 directly phosphorylates transcription factors, thereby altering their activity.
This provides a simple and logical explanation for how Srb10 mediates gene-specific transcriptional regulation. Interestingly, the exact mechanism of regulation appears to differ for different transcription factors. Whereas Srb10 targets Gcn4 for SCF<sub>Cdc4</sub>-dependent degradation, it appears to down-regulate Msn2 activity by promoting nuclear exclusion [although the exact manner in which this occurs remains to be determined]. In support of a direct role for Srb10 in the regulation of transcription factors, phosphorylation of the S699 residue of Gal4 by Srb10 was recently shown to be essential for Gal4-dependent induction of<sup>1</sup> GAL genes (Hirsch et al. 1999). Coupled with phosphorylation of the CTD, direct phosphorylation of specific transcription factors by Srb10 might potentiate its inhibitory effect on transcription and help focus its repressive activity upon specific promoters.

An interesting idea that emerges from our and other studies is that direct negative regulation of transcriptional activators by Srb10 may be confined to the vicinity of promoters to limit the lifetime of transcription factors that are actively engaged in promoting transcription. We propose that Srb10 comprises a timer that constitutively targets promoter-bound Gcn4 for degradation to limit the number of transcripts that a DNA-bound molecule of Gcn4 can promote. Intriguingly, analysis of a set of synthetic transcriptional activators revealed an inverse relationship between their potency and their stability, suggesting that the degradation of transcription factors may be mechanistically coupled to their ability to promote assembly of active transcriptional complexes on promoter DNA (Molinari et al. 1999). Thus, spatially restricted activation of transcription factor proteolysis may be a general regulatory theme. Note that a related argument can also be advanced for Msn2 and Msn4, whereby Srb10-dependent phosphorylation initiates the nuclear export of Msn molecules located at promoter elements. Although our studies highlight roles for Srb10 in the control of Gcn4 stability and Msn2 export, it is possible that Srb10 modulates other aspects of transcription factor activity in a spatially restricted manner. It will be interesting to see if the mammalian homolog of Srb10 likewise mobilizes diverse regulatory strategies to enable tight regulation of gene expression.

Materials and methods

Yeast strains

The yeast strains used in this study are listed in Table 1. All YC strains were derived from Z719, Z690, and Z687 [Liao et al.
Yeast cells were cultured in standard rich or synthetic medium supplemented with 2% dextrose, raffinose, or galactose as described (Sherman 1991). The GCN4 locus was modified to encode Gcn4 tagged with three copies of the HA epitope (HA3) at its C terminus as described (Schneider et al. 1995), and a gcn4 deletion strain (gcn4::HA3) was generated using a similar scheme. The GCN4, SRB10, srb10-3, KIN28, and MSN2 loci were modified to encode proteins tagged with nine copies of the myc epitope (myc9) at their C termini as described (Seol et al. 1999). GAL–GCN4 alleles were generated by replacement of the entire GCN4 promoter region with a PCR fragment containing URA3 and GAL10 promoter sequences. *cddc34* strains were constructed by homologous recombination using a plasmid carrying *cddc34-2* (Kornitzer et al. 1994). The *gcn4*, *pho85*, and *msn5* deletion strains were generated by PCR-based one-step replacement. Deletion alleles were confirmed by scoring mutant phenotypes and by PCR. To generate the *gcn4–ST2* strain, sequences spanning the five ST/TP sites within GCN4 were first replaced by URA3. The resulting GCN4::URA3 strain was subsequently transformed with a *gcn4–ST2* fragment that also contained adjacent 5′ untranslated sequences. Transformants were cultured inYPD for 6–10 h prior to plating on 5-fluoroorotic acid [5-FOA] medium. 5-FOA-resistant transformants were screened by PCR and confirmed by sequencing. All strains derived from Z719, Z690, or Z687 were constructed either by direct transformation using lithium acetate method or by genetic cross. Details of the strain constructions or oligonucleotide sequences are available upon request.

**Fractionation of yeast extracts**

Whole cell or DEAE Sepharose FF [Amersham Pharmacia] fractions of yeast extracts from G1 cyclin-depleted RJD885 or RJD893 were prepared as described (Verma et al. 1997b) with the following modifications. After the frozen cell powder was thawed in Buffer93 [B93], saturated [100%] ammonium sulfate solution was added slowly to a final concentration of 10% to allow further extraction of proteins. To concentrate fractionated proteins prior to assay, the flow-through and 0.15–0.25 M NaCl eluate from the 0%–35% ammonium sulfate cut were concentrated (to ~30 mg/mL) using an Ultrafree-15 centrifugal filter (Biomax-10K, Millipore) and dialyzed against B93.

To resolve Gcn4 kinase and ubiquitination activities, RJD893 whole-cell extract (WCE) was first fractionated by ammonium sulfate precipitation. The 0%–35% and the 45%–60% cuts were resuspended in CBW (25 mM HEPEs at pH 7.6, 25 mM NaCl, and 1 mM DTT) and applied to a DEAE Sepharose column. The 0.25 M eluate from each DEAE fractionation was concentrated to 10–15 mg/mL and dialyzed against B93. The 0.25 M DEAE eluate from the 0%–35% ammonium sulfate cut contained Gcn4 kinase activity, whereas that from the 45%–60% cut contained Gcn4 ubiquitination activity (when supplemented with the kinase fraction plus Cdc34). For assays shown in Figure 2C, the dialyzed 0%–35% ammonium sulfate cut was used directly without further purification.

To partially purify the Gcn4 kinase, we grew RJD481 cells in YPD to late log phase (optical density at 600 nm, OD600 = 2.5–3) and harvested and lysed them as described (Verma et al. 1997b). Whole cell extract (~1000 mg protein) was first fractionated by ammonium sulfate precipitation, and the 0%–40% cut (~150 mg protein) was resuspended in CBW and loaded onto a 10-mL SP Sepharose FF [Amersham Pharmacia] column. The column was washed with 30 mL CBW and eluted with a 100–500 mM NaCl gradient. Fractions containing the peak activity (as measured by kinase and ubiquitination assays, see below) were pooled (~7 mg protein), diluted in CBW, and applied to 2 mL of glutathione agarose (Sigma) previously loaded with ~4 mg GST–Gcn4. The affinity column was washed with 5 mL of CBW + 100 mM NaCl and eluted with 4 mL of CBW + 400 mM NaCl. The eluate was concentrated to 200 µL (~300 µg protein), and 50 µL was loaded onto a 2.4-mL Superdex200 column [Amersham Pharmacia] equilibrated with CBW + 500 mM NaCl. Fractions of volume 50 µL were collected, and 2 µL of each fraction was used in the activity assays.

**Preparation of assay substrates**

GCN4 and SIC1 were transcribed from EcoRI-linearized YEpl8–GCN4 and RBD445 by SP6 and T7 polymerase, respectively. A GCN4–HA transcription template was generated by PCR using a 5′ oligonucleotide containing a T7 RNA polymerase promoter [Verma et al. 1997b] and a 3′ oligonucleotide containing sequences that encode the HA epitope. All mRNAs were translated in rabbit reticulocyte lysates according to manufacturer’s instructions [Promega]. For ubiquitination assays, in vitro-translated Gcn4 was partially purified by batch chromatography on DEAE resin as described for Sic1 [Verma et al. 1997c]. The 100–400 mM NaCl DEAE eluate containing Gcn4 was heated at 65°C for 5 min to destroy a nonspecific Gcn4 kinase activity present in reticulocyte lysates (brief heating at up to 90°C did not significantly compromise the ability of Gcn4 to serve as a ubiquitination substrate). The eluate was then centrifuged at 10,000g for 5 min, and the supernatant was exchanged into 30 mM Tris·HCl at pH 7.5, 50 mM potassium acetate, and 1 mM DTT by three cycles of dilution and concentration to the original volume in a Centricon-10 [Amicon]. Recombinant Gcn4 was also expressed in *E. coli* and purified as described [Kim et al. 1994]. GST, GST–Gcn4, and GST–Msn2 were expressed in BL21(DE3) containing pLyS, and were purified by glutathione affinity chromatography. GST–CTD was expressed and purified as described [Thompson et al. 1993].

**Immunoprecipitations, kinase assays, and immunobLOTS**

Active protein kinases were isolated from yeast strains in which the relevant genes were modified to encode proteins with a myc9 epitope at the C terminus. Cells were grown to late log phase (OD600 = 2.5–3), and whole cell extracts were prepared as described above. All purification procedures were performed on ice or at 4°C. Typically, cell extracts (and supernatants from k 30–40 mg/mL) were diluted three-fold with Buffer B (30 mM Tris·HCl at pH 7.5, 6 mM ammonium sulfate, 1 mM EDTA, 1 mM DTT, 0.02% Triton X-100, and 0.5 mM PMSF), and preincubated with 15 µL protein A beads for 1 h. After brief centrifugation, the supernatant was incubated with 1 µL of anti-*myc* monoclonal antibody in 9E10 ascites fluid for 2 h and then with 10–15 µL of protein A beads for an additional 1–2 h. The protein A beads were washed three times with Buffer B and once with kinase assay buffer (KAB: 30 mM Tris·HCl at pH 7.5, 10 mM MgCl2, 10 mM NaCl, 1 mM DTT, and 0.02% Triton X-100).

32P kinase assays were performed by incubating immobilized kinase with 1 µg substrate [*Gcn4, GST–CTD or GST–Msn2*] in a 10-µL reaction containing KAB, 0.5 µL γ-32P-ATP [4500 Ci/m mole], and 50 µM ATP at 22°C for 1 h with occasional mixing. The reaction was briefly centrifuged, and the supernatant was either used immediately as substrate for the ubiquitination reactions [Figs. 3D] or mixed with Laemmli sample buffer, boiled for 3 min, resolved by SDS-PAGE, and visualized by autoradiography. For [%35S]methionine-labeled...
Gcn4 substrates, kinase assays (Figs. 1B and 3C) were carried out for 30 min at 25°C in KAB supplemented with ATP regenerating system. Immunoblotting was performed with either monoclonal 9E10 ascites fluid (1:2000 dilution) or affinity-purified anti-Cdc28 polyclonal antibody. Blots were visualized using horseradish peroxidase-conjugated secondary antibodies and a chemiluminescence (ECL+ kit; Amersham). For the experiment illustrated in Figure 4C, blots were scanned and quantitated using the STORM system [Molecular Dynamics].

Ubiquitination assays

Ubiquitination reactions in yeast extracts [Verma et al. 1997c] or with purified components (Feldman et al. 1997; Seol et al. 1999) were carried out essentially as described. Two forms of SCF(Cdc4) were used during the course of this study: a triple infection complex (Skp1/Cdc53PyHA/Cdc4PyHA) and a quadruple infection complex (Hrt1/Skp1/Cdc53/Cdc4PyHA), both of which were expressed in baculovirus-infected Sf9 insect cells, purified by binding to anti-polyoma [α-Py]-conjugated protein A beads, and eluted using Py peptide as described [Seol et al. 1999]. Approximately 100–150 ng of SCF(Cdc4) as estimated by Coomassie-stained gels was used in each reaction. Reactions were incubated at 25°C for 1 h, terminated by addition of Laemmli sample buffer, and evaluated by SDS-PAGE and autoradiography.

Pulse-chase experiments

GCN4myc9, MSN2myc9, and untagged control cells were grown overnight in SD-leucine-histidine-uracil at 30°C to mid-log phase (OD$_{600} = 0.5$). Approximately 3 × 10$^8$ cells were harvested and concentrated by centrifugation into 3 mL of the same medium. Cells were pulse-labeled for 5 min with 500 µCi of $[^35]$S label (1175 Ci/mmole, ICN), then chased in the same medium. Cells were washed three times with Triton buffer (30 mM Tris-HCl at pH 7.5, 0.5% SDS, 6% ammonium sulfate, 0.1 M NaF, and 0.1% NaN$_3$). Cells were centrifuged, and cell pellets were frozen in liquid nitrogen and stored at −80°C if necessary. The cell pellets were then supplemented with 100 µL of 0.5 mm glass beads and 100 µL of lysis buffer (30 mM Tris-HCl at pH 7.5, 0.5% SDS, 6% ammonium sulfate, 1 mM EDTA, 1 mM DTT, and 0.5 mM PMSF), vortexed for 2 sec, and boiled for 3 min. The samples were cooled briefly, vortexed for 2 min, boiled again for 2 min, and vortexed again for 1 min. Samples were then centrifuged at 10,000g for 10 min, and the supernatant was diluted fivefold with Triton buffer (30 mM Tris-HCl at pH 7.5, 1% Triton X-100, 1 mM DTT, and 1 mM PMSF) and immunoprecipitated with 9E10 antibody and protein A beads. The protein A beads were washed three times with Buffer B, aspirated to dryness, and mixed with 2× Laemmli buffer. Samples were boiled for 3 min prior to being evaluated by SDS-PAGE, followed by autoradiography and PhosphorImager (Molecular Dynamics) analysis.

ESMS Analysis of phosphorylated Gcn4

Phospho-Gen4 (70 pmol) was digested with modified trypsin (Promega) for 6 h at 37°C and then subjected to multidimensional phosphopeptide analysis by ESMS [Verma et al. 1997a]. Briefly, phosphopeptides were detected by on-line LC-ESMS. The flow from the column was split, with 95% going to a fraction collector and 5% to the mass spectrometer. The mass spectrometer is optimized to produce and detect CID-generated m/z 79 (PO$_3^-$) product ions, which are highly specific for phosphorylated peptides [Huddleston et al. 1993]. The MS is operated in a single ion-monitoring mode for enhanced sensitivity. Phosphopeptide-containing HPLC fractions (containing m/z 79 ions) were then analyzed by negative-ion nanoelectrospray, using precursor ion scans to distinguish phosphopeptides from unmodified peptides and determine their mass [Carr et al. 1996]. Candidate phosphopeptides were sequenced by tandem mass spectrometry using nanoES [Carr et al. 1996] or on LC-ES MS/MS [Zhang et al. 1998]. Phosphorylation site stoichiometry was determined by measuring the ratio of phosphorylated to nonphosphorylated peptide in the positive ion ES spectrum. Positive ion spectra for each phosphorylated and nonphosphorylated peptide were extracted and summed from a full scan LC-MS analysis of 18 pmoles of Gcn4 tryptic digest.

Immunofluorescence

Cells were grown overnight in 30 mL YPD at 27°C to mid-log phase (OD$_{600} = 0.5$). 4 mL of cells were then removed and fixed with 4.5% formaldehyde at room temperature for 1 h. Indirect immunofluorescence was carried out essentially as described [Pringle et al. 1991], using 9E10 (1:3000) as primary antibodies and fluorescein-conjugated-goat-anti-mouse antibodies (1:3000) as secondary antibodies. Fujichrome Provia 400 slide film was used to record data images on a Zeiss Axioskop microscope.

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