Eukaryotic Release Factor 1 Phosphorylation by CK2 Protein Kinase Is Dynamic but Has Little Effect on the Efficiency of Translation Termination in Saccharomyces cerevisiae

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Protein synthesis requires a large commitment of cellular resources and is highly regulated. Previous studies have shown that a number of factors that mediate the initiation and elongation steps of translation are regulated by phosphorylation. In this report, we show that a factor involved in the termination step of protein synthesis is also subject to phosphorylation. Our results indicate that eukaryotic release factor 1 (eRF1) is phosphorylated in vivo at serine 421 and serine 432 by the CK2 protein kinase (previously casein kinase II) in the budding yeast Saccharomyces cerevisiae. Phosphorylation of eRF1 has little effect on the efficiency of stop codon recognition or nonsense-mediated mRNA decay. Also, phosphorylation is not required for eRF1 binding to the other translation termination factor, eRF3. In addition, we provide evidence that the putative phosphatase Sal6p does not dephosphorylate eRF1 and that the state of eRF1 phosphorylation does not influence the allosuppressor phenotype associated with a sal6Δ mutation. Finally, we show that phosphorylation of eRF1 is a dynamic process that is dependent upon carbon source availability. Since many other proteins involved in protein synthesis have a CK2 protein kinase motif near their extreme C termini, we propose that this represents a common regulatory mechanism that is shared by factors involved in all three stages of protein synthesis.

Protein synthesis is carried out in three distinct steps: initiation, elongation, and termination. While the first two steps have been extensively studied, our understanding of the termination process has lagged behind. Two classes of release factors mediate translation termination in the budding yeast Saccharomyces cerevisiae and other eukaryotes. Eukaryotic release factor 1 (eRF1) (encoded by the SUP45 gene) is a class I release factor that recognizes any of the three stop codons when they are located in the ribosomal A site (5, 10). Following stop codon recognition, eRF1 also induces polypeptide chain release by activating the peptidyl transferase center of the ribosome. eRF3 (encoded by the SUP35 gene) is a class II release factor that facilitates stop codon recognition and stimulates the termination reaction in a GTP-dependent manner (17, 37).

Protein synthesis requires a major commitment of cellular energy and resources. It is not surprising that translation is regulated by multiple mechanisms in response to external stimuli such as nutrient abundance or environmental stress. One mechanism of modulating translational efficiency is through the phosphorylation of various translation factors. Probably the most well-characterized example of this regulation is the phosphorylation of serine 51 of the α subunit of the mammalian translation initiation factor eukaryotic initiation factor 2 (eIF2α). When eIF2α is phosphorylated, it competitively inhibits GTP exchange by eIF2B (39). As a result, the ternary complex of eIF2-Met-tRNA

MATERIALS AND METHODS

Strains and growth conditions. The Saccharomyces cerevisiae strains used in this study are described in Table 1. Strains YDB447 and YDB640 were derived from YDB340 by using standard genetic techniques. Strain YDB640 was constructed using a one-step gene replacement strategy. The entire SAL6 open reading frame was removed by transforming yeast strain YDB340 and YDB447 with a PCR fragment containing the TRP1 gene from Candida glabrata.
flanking homology upstream and downstream of SAL6. The C. glabrata TRPI gene was amplified from pCGTRPI (27). The UFP1 gene was deleted in strain YDB641 by using a similar strategy. Candidate deletion strains were screened by PCR to verify the insertion. Construction of strains YDB830 and YDB447 have been described elsewhere previously (37). Strains YDH6 and YDH8 were kindly gifts from Claiborne V. C. Glover III, and their construction has been described previously (21).

Yeast extract-peptone-dextrose is a rich medium supplemented with 2% glucose; synthetic medium supplemented with 2% dextrose (glucose) and amino acids was also used. Wickerham’s minimal medium is a defined minimal medium (43). For [32P]orthophosphate labeling, the phosphate concentration in Wickerham’s minimal medium was reduced to 100 μM. Other specific nutritional supplements were added as needed.

**Plasmids.** The centromeric plasmid pDB800 (with a LEU2 selectable marker) carries the wild-type SUP45 gene with a 1,568 bp upstream of the AUG start codon and 1,036 bp downstream of the UAA stop codon. pDB800 was also used as a template for site-directed mutagenesis using the QuickChange mutagenesis kit (Stratagene). The S421A mutation was introduced into the SUP45 gene and verified by sequencing, and a SpeI-HindIII fragment was then subcloned back into pDB800 to generate pDB841. Using a similar approach, plasmids encoding the following SUP45 mutant alleles were constructed: S432A (pDB859), S421A/S432A (pDB902), S421D (pDB887), S432D (pDB860), and S421DS432D (pDB858). Similarly, the II22S mutation was introduced into the SUP45 gene and subcloned back into pDB800 using a Bpl-SpeI fragment to generate pDB970.

To make a construct to express the eRF1 mutant lacking the C-terminal 19 amino acids (eRF1-C19), an Spel site was generated immediately 5′ of the stop codon of SUP45. A Bpl-HindIII fragment was then subcloned back into pDB800, digested with Spel, and ligated together to precisely delete the last 19 codons. The resulting plasmid was named pDB943. The plasmid used for protein purification of N-terminal His6-tagged eRF1 and codons. The resulting plasmid was named pDB843. The plasmid used for protein purification of N-terminal His6-tagged eRF1-C19 was made by digesting plasmid pDB843 with Spel and HindIII and subcloned into SUP45-ePT-3A vector pDB696 to create pDB970. Similarly, plasmid pDB858 was digested with SpeII and HindIII and ligated into pDB696 to introduce the eRF1 S421D/S432D double mutant into the ePT-3A vector (pDB941).

**Metabolic labeling and Western blots.** Yeast strains were grown in Wickerham’s low-phosphate minimal medium at 30°C. Cells were grown to mid-log phase (0.5 to 1.4 A600 units/ml). Equal aliquots of cells were harvested and resuspended at 4 A600 units/ml in Wickerham’s low-phosphate medium with or without 2% glucose. After further incubation at 30°C for 2 h, 400 μCi/ml [32P]orthophosphate was added, and incubation was continued for 2 h. Cells were then harvested and processed as described above. For the chase experiment, cells were grown in Wickerham’s low-phosphate medium in the presence of 2% glucose to mid-log phase (0.5 to 1.4 A600 units/ml). Cells were then harvested and labeled at 4 A600 units/ml in fresh Wickerham’s low-phosphate medium containing 2% glucose. After 2 h, equal aliquots of cells were harvested and resuspended at 4 A600 units/ml in fresh Wickerham’s low-phosphate medium in the presence or absence of 2% glucose to initiate the chase period. Samples were taken at 0, 0.5, 1, 2, and 4 h after initiating the chase period and processed as described above.

To assess the role of CK2 protein kinase in eRF1 phosphorylation, strains YDH6 and YDH8 were grown to early log phase at 25°C in Wickerham’s low-phosphate medium. Equal aliquots of cells were then shifted to 37°C or maintained at 25°C with shaking for an additional 12 h (approximately three generations). The cells were then harvested and resuspended at 4 A600 units/ml in fresh medium, and aliquots were metabolically labeled as described above at either 25°C or 37°C for two more hours. The cells were then lysed, and samples were processed as described above.

**Analytical ultracentrifugation.** Purification of N-terminal His6-tagged eRF1 and His6-tagged eRF3 (residues 254 to 685) from the soluble fraction of Escherichia coli lysates has been described previously (37). Immediately following purification, proteins were dialyzed in phosphate-buffered saline (140 mM NaCl, 2 mM NaPO4, pH 7.4). Protein concentrations were determined using Beer’s law: A = εc, where A is the absorbance, ε is the molar absorptivity, c is the light path through the sample, and λ is the concentration. Protein concentrations for analysis were 29 μM for individual proteins or 58 μM total for the mixtures of eRF1 and eRF3. For complex formation, eRF1 and eRF3 were mixed in 1:1 molar ratios and incubated on ice for 20 min prior to centrifugation. Sedimentation values were obtained using a Beckman XL-A ultracentrifuge with a four-channel An-60 Ti rotor. Sedimentation profiles were subjected to the software program SEDFIT (available at http://www.analyticalultracentrifugation.com).

**Readthrough assays.** The stop codon readthrough assays were performed using a bicistronic reporter construct consisting of a Renilla luciferase gene followed by an in-frame firefly luciferase gene. Separating the two genes is either a tetranucleotide termination signal (e.g., UAA A) or a similar sequence containing a sense codon (e.g., CAA A). The construct is driven by the PGK1 promoter and has a CYC1 poly(A) addition signal. Cells were grown in synthetic medium with 2% dextrose and the appropriate amino acids. Percent readthrough was determined by taking the ratio of the firefly/Renilla activities obtained from the termination signal construct relative to the firefly/Renilla activities for the control (sense codon) construct. Samples were processed in quintuplicate, and each experiment was repeated at least twice (for further details, see reference 25).

**Table 1. Strains used in this study**

| Strain | Genotype | Reference or source |
|--------|----------|---------------------|
| YDB340 | MATα ura3-52 leu2-3,112 his3Δ200 trp1Δ401 his4Δ200 leu2-80 succ2-Δ901 [psi-] | 37 |
| YDB447 | MATα ura3-52 leu2-3,112 his3Δ200 trp1Δ401 his4Δ200 leu2-80 succ2-Δ901 sup45::His5[psi-] | 37 |
| YDB640 | MATα ura3-52 leu2-3,112 his3Δ200 trp1Δ401 his4Δ200 leu2-80 succ2-Δ901 sup45::His5[psi-] | This study |
| YDB641 | MATα ura3-52 leu2-3,112 his3Δ200 trp1Δ401 his4Δ200 leu2-80 succ2-Δ901 sup45::His5[psi-] | This study |
| YDH6  | MATα cka1Δ::His3 cka2Δ::TRPI1 cka2Δ::LEU2 | 21 |
| YDH8  | MATα cka1Δ::His3 cka2Δ::TRPI1 cka2Δ::LEU2 | 21 |
Northern blots. RNA was extracted from cells using a hot-phenol extraction method (38). Total RNA (25 μg/lane) from the indicated strains was resolved on a 1% agarose-formaldehyde gel and transferred onto a nitrocellulose membrane using a Posi-Blot pressure blower (Stratagene). Samples were baked in a vacuum oven at 80°C and then probed with a32P-labeled DNA probe for the CYH2 gene. After results were visualized using a PhosphorImager (GE Healthcare), the membrane was stripped by incubation in stripping solution (10 mM Tris-HCl, pH 7.5, 0.2% SDS) for 1.5 h at 75°C. The membrane was then probed with an ACT1 probe for the loading control, and results were again visualized using a PhosphorImager (GE Healthcare).

RESULTS
eRF1 is phosphorylated by CK2 protein kinase in vivo. Due to the frequent phosphorylation of translation initiation and elongation factors, we began this study by asking whether either one of the translation termination factors eRF1 and eRF3 is phosphorylated in the budding yeast Saccharomyces cerevisiae. Yeast cultures were metabolically labeled with [32P]orthophosphate or [35S]methionine for 2 h, and immunoprecipitation experiments were carried out using polyclonal antibodies to either eRF1 or eRF3. Figure 1 shows that while both eRF1 and eRF3 readily incorporated [35S]methionine, only eRF1 could be labeled with [32P]orthophosphate. These results demonstrate that eRF1 is a phosphoprotein in yeast cells.

Using the NetPhos 2.0 phosphorylation site predictor (http://www.cbs.dtu.dk/services/NetPhos), we found that 10 serine residues at positions 421 and 432 to alanine, both individually and together, and plasmids encoding the mutant forms of eRF1 were introduced into a sup45Δ strain expressing wild-type eRF1 from a plasmid carrying the URA3 gene as a selectable marker. Plasmid shuffling (6) was then carried out by plating the transformants onto medium supplemented with 5-fluoroorotic acid, a uracil analogue that allows the formation of colonies only from cells that have lost the wild-type SUP45 plasmid with the URA3 marker. Each strain was then tested for the ability to phosphorylate eRF1 (Fig. 3). The results show that the introduction of either the S421A or S432A mutation reduced the level of eRF1 phosphorylation, while both mutations together eliminated most (if not all) eRF1 phosphorylation. These results indicate that the serine residues at positions 421 and 432 are the major sites of eRF1 phosphorylation in vivo.

TABLE 2. Known CK2 phosphorylation sites near the C termini of translation factors

| Gene | Protein | Phosphorylation site | Function | Reference |
|------|---------|----------------------|----------|-----------|
| RPP2B | P2B | ...KEESDDDM... | Component of ribosomal stalk | 3 |
| RPP2A | P2Δ | ...KEESDDDM... | Component of ribosomal stalk | 3 |
| RPP1B | P1β | ...KEESDDDM... | Component of ribosomal stalk | 3 |
| RPP1A | P1α | ...KEESDDDM... | Component of ribosomal stalk | 3 |
| RPP0 | P0 | ...KEESDDDM... | Component of ribosomal stalk | 3 |
| TIF5 | elf5 | ...I1SEEE... | Promotes GTP hydrolysis by elf2 | 30 |
| SUI2 | elf2α | ...DS4DDEEEEEDE* | α subunit of elf2 | 16 |

* An asterisk (*) indicates the position of the stop codon.
The phosphorylation state of eRF1 has little effect on the efficiency of translation termination or nonsense-mediated mRNA decay (NMD). Having identified the two major sites of eRF1 phosphorylation, we used the phosphorylation mutants described above to characterize the role that eRF1 phosphorylation plays in the cell. The double mutant eRF1-S421A/S432A allowed us to examine how the loss of eRF1 phosphorylation affects the function of this protein. We also constructed another mutant, eRF1-S421D/S432D, in which the phosphorylated serine residues were changed to aspartic acid to mimic constitutive phosphorylation. We first used these mutants to determine whether phosphorylation plays a role in translation termination. To do this, we used a dual luciferase reporter system (19, 25, 37). Briefly, this system utilizes tandem Renilla and firefly luciferase genes that are separated by a single in-frame stop codon (Fig. 4). The activity of firefly luciferase, encoded by the distal open reading frame, provides a quantitative measure of the readthrough of the stop codon that separates the two open reading frames. The activity of Renilla luciferase, encoded by the proximal open reading frame, serves as an internal control for mRNA abundance and translation initiation rates. Since the efficiency of translation termination is influenced not only by the stop codon but also by the first nucleotide after a stop codon (together referred to as the tetranucleotide termination signal), we examined the effect of the eRF1 phosphorylation mutants on all possible tetranucleotide termination signals (Table 3). We found that the mutations that eliminated phosphorylation (eRF1-S421A/S4342A) caused a small (1.2- to 1.5-fold) increase in readthrough at the UAA A, UAA U, and UAG G termination signals. Similarly, the mutations that mimicked constitutive phosphorylation (eRF1-S421D/S432D) exhibited a small (1.3- to 1.6-fold) increase in readthrough at the UAA A, UAA U, UAG C, and UGA G termination signals but had little effect on stop codon recognition at other signals. These increases are significant ($P < 0.01$) but relatively small compared to an eRF1 mutant lacking the extreme C-terminal 19 amino acids (3.3- to 17.6-fold) (data not shown). Taken together, these results indicate that the phosphorylation state of eRF1 has little effect on the efficiency of stop codon recognition in vivo.

NMD rapidly degrades mRNAs that contain premature stop mutations in a process that is intimately linked with translation termination (1, 25). The induction of NMD in yeast requires a
Phosphorylation of eRF1 does not affect its ability to interact with eRF3. To test this hypothesis, we examined the association between eRF1 and eRF3 by performing sedimentation velocity experiments using analytical ultracentrifugation (Fig. 6). His-tagged forms of full-length eRF1 or a fragment of eRF1 consisting of amino acid residues 254 to 685 were purified from *E. coli* extracts using Ni²⁺-nitrilotriacetic acid affinity chromatography. Since *E. coli* does not contain protein kinases that recognize CK2 protein kinase sites, wild-type eRF1 purified from *E. coli* was assumed to be in the dephosphorylated form. As a negative control for eRF1-eRF3 complex formation, we used an eRF1 mutant lacking the last 19 amino acids (eRF1-CΔ19) that does not efficiently bind eRF3 (15). Preliminary experiments found that eRF1, eRF1-CΔ19, and eRF3 each have sedimentation coefficients ranging from 2.07S to 2.09S (Fig. 6). When wild-type (unphosphorylated) eRF1 and eRF3 were incubated together for 20 min at 4°C before centrifugation, we observed a new 3.06S complex that represented the eRF1-eRF3 complex. This indicated that a lack of phosphorylation did not prevent the formation of the eRF1-eRF3 complex. As expected, this 3.06S peak was not observed when eRF1-CΔ19 was incubated with eRF3, indicating that the deletion of the C-terminal 19 amino acids of eRF1 disrupted eRF1-eRF3 complex formation. Incubation of eRF3 with the eRF1 mutant that mimics constitutive phosphorylation (eRF1-S421D/S432D) resulted in the appearance of the 3.06S peak, indicating that complex formation still occurred when acidic residues were present at the sites of eRF1 phosphorylation. Taken together, these results indicate that the phosphorylation state of eRF1 does not play a significant role in eRF1-eRF3 heterodimer formation.

The state of eRF1 phosphorylation is not affected by the putative phoshatase encoded by *SLA6* and does not affect the allosuppressor phenotype associated with a *sal6Δ* mutant. Mutations in the *SUP45* and *SUP35* genes, encoding eRF1 and eRF3, respectively, were originally identified based on their ability to cause readthrough of stop mutations in various biosynthetic genes of *S. cerevisiae* (22). These mutations were
termed omnipotent suppressors because they cause readthrough at all three stop codons. Later, the \textit{SAL6} gene was identified in a screen for mutants that enhanced the readthrough phenotypes associated with omnipotent suppressor mutations (40). Mutations in the \textit{SAL6} gene alone did not result in a suppressor phenotype, but they exacerbated the readthrough associated with omnipotent suppressor mutations in the \textit{SUP35} or \textit{SUP45} gene. This enhanced suppression phenotype is referred to as an allosuppressor effect. The \textit{SAL6} gene was later sequenced and was found to encode a protein with significant homology to PP1-like protein phosphatases (11, 42). However, it is presently still not clear how the protein encoded by the \textit{SAL6} gene mediates the allosuppressor effect.

Given our finding that eRF1 is a phosphoprotein, we next tested the hypothesis that the allosuppressor phenotype associated with the \textit{sal6Δ} mutation is mediated through its effect on the phosphorylation state of eRF1. To do this, we used the dual luciferase reporter system to compare termination efficiencies in strains expressing the phosphorylation mutants in \textit{SAL6} and \textit{sal6Δ} backgrounds (Fig. 7). Since the allosuppressor effect requires the presence of an omnipotent suppressor mutation, the phosphorylation site mutations were combined with the eRF1-I222S mutation, which was originally identified as the \textit{sup45-2} omnipotent suppressor allele (41). When these eRF1 mutant proteins were expressed from a low-copy-number plasmid in a \textit{sup45/H9004} strain, we found that expression of the eRF1-I222S allele as the sole source of eRF1 resulted in a 3.3-fold increase in readthrough (Fig. 7A). When eRF1-I222S was expressed in the \textit{sal6Δ} strain, we observed a 6.6-fold increase in readthrough, indicating that the presence of the \textit{sal6Δ} mutation caused an additional twofold increase in readthrough. This observation recapitulated the previously observed allosuppressor effect associated with \textit{SAL6} mutations (40). However, we did not detect any significant change in readthrough when either the S421A/S432A or the S421D/S432D phosphorylation site mutation was combined with the eRF1-I222S mutation in the \textit{sal6Δ} strain. If Sal6p was the phosphatase for eRF1, the \textit{sal6Δ} mutation might increase the phosphorylation state of eRF1. This hypothesis predicts that the mutant protein that mimics constitutive phosphorylation, eRF1-S421D/S432D, will phenocopy the allosuppressor effect in a \textit{SAL6} strain, since the presence of the aspartic acid residues mimics the maximal phosphorylation state of eRF1. However, such an allosuppressor effect was not observed in the \textit{SAL6} strain expressing eRF1-I222S/S421D/S432D (Fig. 7A). These results suggest that the allosuppressor effect associated with the \textit{sal6Δ} mutation is not mediated by its effect on the phosphorylation state of eRF1.

The absolute level of eRF1 phosphorylation might also be expected to increase in a \textit{sal6Δ} strain if Sal6p is the phosphatase responsible for the dephosphorylation of eRF1 (assuming that it is not already fully phosphorylated under normal conditions). To test this hypothesis, we used our metabolic labeling assay to determine whether any change in the phosphorylation state of eRF1 could be detected in the \textit{sal6Δ} mu-
As shown in Fig. 7B, we did not observe any significant difference in the phosphorylation state of eRF1 between the wild-type and sal6Δ strains. We cannot rule out the possibility that eRF1 is completely phosphorylated under the growth conditions under which this experiment was carried out. If that were the case, the deletion of its phosphatase would not be expected to show a detectable increase in phosphorylation. However, when considered together with the allosuppressor readthrough data, these results argue that Sal6p is not the phosphatase responsible for the dephosphorylation of eRF1, and the state of eRF1 phosphorylation does not appear to be important for the allosuppressor effect associated with sal6 mutations.

eRF1 phosphorylation and dephosphorylation are dynamic processes that require cell growth. The results described above indicate that the phosphorylation state of eRF1 does not significantly influence its termination activity. Consistent with this finding, mutations that eliminated the CK2 phosphorylation sites in the five proteins that form the ribosomal stalk shown in Table 1 also did not affect either ribosome activity or the formation of the stalk structure (3, 36). Similarly, studies that altered the CK2 phosphorylation sites in eIF2α and eIF5 did not have any detectable effects on translation initiation (16, 30). However, the existence of CK2 phosphorylation sites in these and other proteins involved in translation suggests that this modification may carry out some common function. To gain further insight into the nature of eRF1 phosphorylation, we next asked whether this modification required active cell growth. Wild-type cells were grown to mid-log phase and harvested, and similar aliquots were resuspended in the same growth medium in the presence or absence of 2% glucose. The cultures were incubated for 2 h at 30°C to deplete any residual glucose in the glucose-depleted culture, and [32P]orthophosphate was then added to each culture. Incubation at 30°C was continued for an additional 2 h, and the cells were harvested. As expected, glucose depletion under these conditions effectively blocked cell growth (Fig. 8A). When we examined the phosphorylation state of eRF1, we found that growth inhibition by glucose depletion essentially eliminated the phosphor-
Cells were labeled for 2 h with [32P]orthophosphate and then possibilities, we performed a metabolic chase experiment (Fig. 9). Dephosphorylated form of eRF1. To distinguish between these cycles may arrest with cells that contain predominantly the growth is inhibited by glucose depletion, this phosphorylation dephosphorylated forms during normal growth. When cell alternatively after its synthesis. By subjecting the cells to glucose depletion, we were inhibiting de novo protein synthesis and eRF1 may only be phosphorylated once during (or immediately after) its synthesis. Previous studies have shown that several factors involved in the initiation and elongation steps of translation undergo post-translational modification by phosphorylation. In this study, we show that the translation termination factor eRF1 is also a phosphoprotein in yeast, which shows for the first time that all three steps in translation are modified in this manner. We found that eRF1 is phosphorylated in two sites near its C terminus by the CK2 protein kinase as previously shown for several other yeast proteins involved in translation, including all five proteins of the ribosomal stalk (3), and initiation factors eIF5 (30) and eIF2α (16) (Table 2). Surprisingly, we found that the introduction of mutations that either eliminated eRF1 phosphorylation or mimicked constitutive eRF1 phosphorylation had little effect on the efficiency of stop codon recognition in vivo (as measured by a translational readthrough assay). The level of eRF1 phosphorylation was also unaffected by the disruption of the SAL6 gene, which encodes a putative PP1-like phosphatase. Furthermore, the allosuppressor phenotype associated with the sal6Δ mutation was insensitive to the state of eRF1 phosphorylation. These results indicate that the phosphorylation of eRF1 does not play a significant role in its ability to mediate efficient translation termination under the growth conditions used in the current study.

We also found that the phosphorylation state of eRF1 did not influence its ability to bind eRF3, even though phosphorylation occurred within the portion of eRF1 that is responsible for eRF3 binding. In addition, our results indicate that phosphorylation of eRF1 does not influence nonsense-mediated mRNA decay. From these data, we can infer that the phosphorylation state of eRF1 probably does not affect Upf1p binding, since the association of Upf1p with the termination complex is required for optimal translation termination efficiency (12, 25).

Based on the results summarized above, the physiological role for CK2-mediated phosphorylation of eRF1 remains obscure. Previous studies of other proteins involved in translation that undergo CK2 phosphorylation near their C termini have also had difficulty establishing a function for this modification. A study of the P1β protein of the ribosomal stalk suggested that phosphorylation, in conjunction with an N-terminal signal, may stimulate protein degradation. This hypothesis is consistent with the previous observation that phosphorylation can function as a degradation signal for the vacuolar and protea-
some turnover pathways (23). However, it was subsequently shown that P1β was not degraded by either of these pathways. Instead, it was shown that a failure of the P1 dimer to associate with P2 causes a rapid turnover of P1 (33). Therefore, phosphorylation of P1 may simply play a role in its association with P2, with enhanced degradation occurring due to a lack of assembly. In another example, yeast eIF2α is phosphorylated by the CK2 protein kinase at three serine residues near its C terminus (amino acids 292, 294, and 301). Although mutation of these serine residues to alanine did not cause a detectable phenotype, it did exacerbate the growth defects observed when these mutations were combined with other mutations that reduced the efficiency of nucleotide exchange, suggesting that phosphorylation of the CK2 protein kinase sites in eIF2α may be required for optimum eIF2 activity (16). Similarly, we found that changes in the phosphorylation state of eRF1 resulted in only subtle effects on stop codon recognition that may help to fine-tune the termination process at certain termination signals (Table 3). Finally, it was shown that eIF5 is phosphorylated at multiple sites near its C terminus by the CK2 protein kinase (30), but a role for this modification was not found. Taken together, these results suggest that phosphorylation of various translation factors by the CK2 protein kinase does not play a strong role in their specific functions related to translation under steady-state growth conditions. Thus, the purpose of this posttranslational modification remains somewhat obscure.

The database searches carried out during this study also revealed that at least two other yeast proteins involved in translation, eIF2Be and eIF4γ, also have CK2 phosphorylation motifs near their extreme C termini. In addition, a number of other translation factors, including eIF1β, eIF1A, Tif53p, eIF2βγ, and eIF4B, have consensus CK2 phosphorylation motifs within 50 amino acids of their C termini. We hypothesize that these proteins are also phosphorylated by the CK2 protein kinase. Based on the results of the phosphorylation studies described above, the phosphorylation of these factors (if it occurs) may not play a dramatic role in regulating their activities. Instead, they may be phosphorylated by the CK2 protein kinase in response to favorable growth conditions in a manner similar to that of the five ribosomal stalk proteins, eIF2α, eIF5, and eRF1. Here, we have provided evidence that (i) the phosphorylation of eRF1 is dynamic and goes through a phosphorylation/dephosphorylation cycle rather than being constitutively phosphorylated following its synthesis, (ii) active cell growth is required for the phosphorylation/dephosphorylation cycle, and (iii) inhibition of this phosphorylation cycle by eliminating the sites of phosphorylation does not have adverse effects on translation or steady-state growth when cells are grown with glucose as a carbon source. It is possible that this phosphorylation pathway is important when the cells are grown under adverse conditions where subtle changes in the activity of the translational apparatus or the function of other related factors may be important. It is also possible that this conserved modification may act to down-regulate translation under conditions that are not favorable for cell growth.

A significant body of evidence also indicates that many factors have alternate cellular functions. For example, eIF1A not only plays a role in translation elongation but has also been shown to bind actin and influence the distribution of the actin cytoskeleton (20, 24, 32). Ccr4p, a subunit of the major cytoplasmic deadenylase complex, also acts as a transcription factor that becomes active in response to DNA damage (28). It is currently not known whether a posttranslational modification such as phosphorylation regulates any (or all) of the alternate functions of these factors. Further studies will be required to examine this possibility.

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REFERENCES

1. Amrani, N., R. Ganesan, S. Kervestin, D. A. Mangus, S. Ghosh, and A. Inamori. 2004. A faus 3′-UTR promotes aberrant termination and triggers nonsense-mediated mRNA decay. Nature 432:112–118.
2. Andjelkovic, N., S. Zolnierowicz, C. Van Hoof, J. Goris, and B. A. Hemmings. 1996. The catalytic subunit of protein phosphatase 2A associates with the translation termination factor eRF1. EMBO J. 15:7156–7167.
3. Ballesta, J. P., M. A. Rodriguez-Gabriel, G. Bou, E. Briones, R. Zambrano, and M. Remacha. 1999. Phosphorylation of the yeast ribosomal stalk. Functional effects and enzymes involved in the process. t7EFS Mol. Microbiol. 35:337–350.
4. Bandhakavi, S., R. O. McMann, D. E. Hanna, and C. V. Glover. 2003. Genetic interactions among ZDS1.2, CDC37, and protein kinase CK2 in Saccharomyces cerevisiae. FEBS Lett. 554:295–300.
5. Bertram, G. A., H. A. Bell, D. W. Rine, and L. Stansfield. 2000. Terminating eukaryote translation: domain I of release factor eRF1 functions in stop codon recognition. RNA 6:1236–1247.
6. Boeke, J. D., F. LaCroute, and G. R. Fink. 1984. A positive selection for mutants lacking orotidine-5′-phosphate decarboxylase activity in yeast: 5-fluoro-7-uracil acid resistance. Mol. Genet. 197:345–346.
7. Cardenas, M. E., Q. Dang, C. V. Glover, and S. M. Gasser. 1992. Casein kinase II phosphorylates the eukaryote-specific C-terminal domain of topoisomerase II in vivo. EMBO J. 11:1785–1791.
8. Carlberg, U., A. Nilsson, and O. Nygard. 1990. Functional properties of phosphorylated elongation factor 2. Eur. J. Biochem. 119:639–645.
9. Chang, Y. W., and J. A. Traup. 1998. Insulin stimulation of phosphorylation of elongation factor 1 (eEF-1) enhances elongation activity. Eur. J. Biochem. 218:1–207.
10. Chavatte, L., L. Frolova, L. Kisselev, and A. Favre. 2001. The polypeptide chain release factor eRF1 specifically contacts the s(4)UGA stop codon located in the A site of eukaryotic ribosomes. Eur. J. Biochem. 268:2986–2992.
11. Chen, M. X., Y. H. Chen, and P. T. Cohen. 1993. PPQ, a novel protein phosphate containing a Ser + Asn-rich amino-terminal domain, is involved in the regulation of protein synthesis. Eur. J. Biochem. 216:689–699.
12. Czarnowska, K., M. J. Butow, J. J. Guzarkiewicz, S. V. Pushklin, X. Han, Y. Weng, H. A. Perlick, H. C. Dietz, M. D. Ter-Avanesyan, and S. W. Peltz. 1998. The surveillance complex interacts with the translation release factors to enhance termination and degrade aberrant mRNAs. Genes Dev. 12:1665–1677.
13. Dahlseid, J. N., J. Puziss, R. L. Shirley, A. L. Atkin, P. Hieter, and M. R. Culbertson. 1998. Accumulation of mRNA coding for the efl3p kinetochore subunit of Saccharomyces cerevisiae depends on the same factors that promote rapid decay of nonsense mRNAs. Genetics 150:1019–1035.
14. Eibara, K., and Y. Nakamura. 1990. C-terminal interaction of translational release factors eRF1 and eRF3 of fission yeast: G-domain uncoupled binding and the role of conserved amino acids. RNA 6:579–730.
15. Eurlwaichitr, L., F. M. Graves, I. Stansfield, and M. F. Tuite. 2001. The polypeptide chain release factor eRF1- and ribosome-dependent guanosine triphosphatase. RNA 7:201–207.
16. Feng, L., H. Yoon, and T. F. Donahue. 1994. Casein kinase II mediates multiple phosphorylation of Saccharomyces cerevisiae eIF-2α (encoded by XU2), which is required for optimal eIF-2 function in S. cerevisiae. Mol. Cell. Biol. 14:5139–5153.
17. Frolova, L., X. Le Goff, G. Zhouravleva, E. Davydova, M. Philippe, and L. Kisselev. 1996. Eukaryotic polypeptide chain release factor eRF3 is an essential ribosome-bound guanine triphosphatase. RNA 2:334–341.
18. Gonzalez, C. I., W. Wang, and S. W. Peltz. 2001. Nonsense-mediated mRNA decay in Saccharomyces cerevisiae: a quality control mechanism that degrades transcripts harboring premature termination codons. Cold Spring Harb. Symp. Quant. Biol. 66:321–328.
19. Gretzmann, G., J. A. Ingram, P. J. Kelly, R. F. Gesteland, and J. F. Atkins. 1998. A dual-luciferase reporter system for studying recoding signals. RNA 4:479–486.
20. Gross, S. R., and T. G. Kinzy. 2005. Translation elongation factor 1A is essential for regulation of the cell actin cytoskeleton and cell morphology. Nat. Struct. Mol. Biol. 12:772–778.

21. Hanna, D. E., A. Rethinaswamy, and C. V. Glover. 1995. Casein kinase II is required for cell cycle progression during G1 and G2/M in Saccharomyces cerevisiae. J. Biol. Chem. 270:25905–25914.

22. Hawthorne, D. C., and U. Leupold. 1974. Suppressors in yeast. Curr. Top. Microbiol. Immunol. 64:1–47.

23. Hershko, A., and A. Ciechanover. 1998. The ubiquitin system. Annu. Rev. Biochem. 67:425–479.

24. Kandl, K. A., R. Munshi, P. A. Ortiz, G. R. Andersen, T. G. Kinzy, and A. E. Adams. 2002. Identification of a role for actin in translational fidelity in yeast. Mol. Genet. Genomics 268:10–18.

25. Keeling, K. M., J. Lanier, M. Du, J. Salas-Marco, L. Gao, A. Kaenjak-Angelletti, and D. M. Bedwell. 2004. Leaky termination at premature stop codons antagonizes nonsense-mediated mRNA decay in S. cerevisiae. RNA 10:991–703.

26. Kitada, K., H. J. Muller, H. E. Meyer, F. Marks, and M. Gochwendi. 1995. Protein kinase C delta-specific phosphorylation of the elongation factor eEF-alpha and an eEF-1 alpha peptide at threonine 431. J. Biol. Chem. 270:15616–6162.

27. Kielbassa, K., E. Yamaguchi, and M. Arisawa. 1995. Cloning of the Candida glabrata TRP1 and HIS3 genes, and construction of their disruptant strains by sequential integrative transformation. Gene 165:203–206.

28. Lenssen, E., N. James, I. Pedruzzi, F. Dubouloz, E. Cameroni, R. Bisig, L. Maillet, M. Werner, J. Roosen, K. Petrovic, J. Winderickx, M. A. Collart, and C. De Virgilio. 2005. The Ccr4-Not complex independently controls both Msn2-dependent transcriptional activation—via a newly identified Glc7/TFIIH promoter distribution. Mol. Cell. Biol. 25:488–498.

29. Lowry, O. H., N. J. Rosebrough, A. L. Farr, and R. J. Randall. 1951. Protein measurement with the folin phenol reagent. J. Biol. Chem. 193:265–275.

30. Maiti, T., A. Bandypadhyay, and U. Maitra. 2003. Casein kinase II phosphorylates translation initiation factor 5 (eIF5) in Saccharomyces cerevisiae. Yeast 20:97–108.

31. Meggio, F., and L. A. Pinna. 2003. One-thousand-and-one substrates of protein kinase CK2? FASEB J. 17:349–368.

32. Munshi, K. A., K. Kandl, A. Carr-Schmid, J. L. Whitacre, A. E. Adams, and T. G. Kinzy. 2001. Overexpression of translation elongation factor 1A affects the organization and function of the cell actin cytoskeleton in yeast. Genetics 157:1425–1436.

33. Nusspauer, G., M. Remacha, and J. P. Ballesta. 2000. Phosphorylation and N-terminal region of yeast ribosomal protein P1 mediate its degradation, which is prevented by protein P2. EMBO J. 19:6075–6084.

34. Paushkin, S. V., V. V. Kushnirov, V. N. Smirnov, and M. D. Ter-Avanesyan. 1997. Interaction between yeast Sup45p (eRF1) and Sup35p (eRF3) polypeptide chain release factors: implications for prion-dependent regulation. Mol. Cell. Biol. 17:2798–2805.

35. Poole, A., T. Poore, S. Bandhakavi, R. O. McCann, D. E. Hanna, and C. V. C. Glover. 2005. A global view of CK2 function and regulation. Mol. Cell. Biochem. 274:163–170.

36. Rodriguez-Gabriel, M. A., M. Remacha, and J. P. Ballesta. 1998. Phosphorylation of ribosomal protein P0 is not essential for ribosome function but can affect translation. Biochemistry 37:16620–16626.

37. Salas-Marco, J., and D. M. Bedwell. 2004. GTP hydrolysis by eRF3 facilitates stop codon decoding during eukaryotic translation termination. Mol. Cell. Biol. 24:7769–7778.

38. Schmitt, M. E., T. A. Brown, and B. L. Trumpower. 1990. A rapid and simple method for preparation of RNA from Saccharomyces cerevisiae. Nucleic Acids Res. 18:3091–3092.

39. Sonenberg, N., and T. E. Dever. 2003. Eukaryotic translation initiation factors and regulators. Curr. Opin. Struct. Biol. 13:56–63.

40. Song, J. M., and S. W. Liebman. 1987. Allosuppressors that enhance the efficiency of omnipotent suppressors in Saccharomyces cerevisiae. Genetics 115:451–460.

41. Stansfield, I., V. V. Kushnirov, K. M. Jones, and M. F. Tuite. 1997. A conditional-lethal translation termination defect in a sup45 mutant of the yeast Saccharomyces cerevisiae. Eur. J. Biochem. 245:557–563.

42. Vincent, A., G. Newnam, and S. W. Liebman. 1994. The yeast translational allosuppressor, SAl6: a new member of the PPI-like phosphatase family with a long serine-rich N-terminal extension. Genetics 138:697–708.

43. Wickerham, L. J. 1946. A critical evaluation of the nitrogen assimilation tests commonly used in classification of yeasts. J. Bacteriol. 52:293–301.

44. Wilson, L. K., N. Dhillon, J. Thorner, and G. S. Martin. 1995. Casein kinase II catalyzes tyrosine phosphorylation of the yeast nucleolar immunophilin Fpr3. J. Biol. Chem. 272:12961–12967.