Yeast heat shock mRNAs are exported through a distinct pathway defined by Rip1p

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We reported previously that heat or ethanol shock in Saccharomyces cerevisiae leads to nuclear retention of most poly(A)+ RNA but heat shock mRNAs (encoding Hsp70 proteins Ssa1p and Ssa4p) are efficiently exported in a process that is independent of the small GTPase Ran/Gsp1p, which is essential for most nucleocytoplasmic transport. To gain further insights into proteins essential or nonessential for export of heat shock mRNAs, in situ hybridization analyses to detect mRNA and pulse-labeling of proteins were used to examine several yeast mutant strains for their ability to export heat shock mRNAs following stress. Rip1p is a 42-kD protein associated with nuclear pore complexes and contains nucleoporin-like repeat sequences. It is dispensable for growth of yeast cells under normal conditions, but we report that it is essential for the export of heat shock mRNAs following stress. When SSA4 mRNA was induced from a GAL promoter in the absence of stress, it was efficiently exported in a strain lacking RIP1, indicating that Rip1p is required for export of heat shock mRNAs only following stress. Npl3p, a key mediator of export of poly(A)+ RNA, was not required for heat shock mRNA export, whereas Rss1p/Gle1p, a NES-containing factor essential for poly(A)+ RNA export, was also required for export of heat shock mRNAs after stress. High-level expression of the HIV-1 Rev protein, but not of Rev mutants, led to a partial block in export of heat shock mRNAs following stress. The data suggest a model wherein the requirement for Npl3p defines the mRNA export pathway, the requirement for Rip1p defines a pathway used for export of heat shock mRNAs after stress, and additional factors, including Rss1p/Gle1p and several nucleoporins (Rat7p/Nup159p, Rat2p/Nup120p, and Nup145p/Rat10p), are required in both pathways.

Key Words: RNA export; heat shock; RIP1; hnRNP; RSS1/GLE1; Rev

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A distinguishing feature of eukaryotic cells is the nucleus, a distinct subcellular compartment separated from the cytoplasm by the double-membraned nuclear envelope. Embedded within the nuclear envelope are nuclear pore complexes (NPCs) that serve as the only known channels for transport between the nucleus and the cytoplasm (for review, see Davis 1995; Panté and Aebl 1996). Transport of macromolecules through NPCs is signal-mediated, saturable, and energy dependent. Considerable progress has been made in recent years in identifying (1) receptor molecules that recognize nuclear localization signals (NLSs) within karyophilic proteins and mediate interactions between these proteins and NPCs, (2) a small Ras-like GTPase (Ran in metazoan cells and Gsp1p/Gsp2p in Saccharomyces cerevisiae) and its accessory proteins, which play a central role in nuclear protein import, and (3) distinct components of NPCs required for nuclear protein import (for review, see Görlich and Mattaj 1996; Hicks and Raikhel 1996; Corbett and Silver 1997; Nigg 1997). The existence of permeabilized cell systems has allowed direct study of nuclear protein import in vitro and played an important role in elucidating the molecular details of this process (Adam et al. 1990; M oore and Blobel 1992; Schlenstedt et al. 1993).

In contrast, we know considerably less about the export of macromolecules from the nucleus. The same GTPase system (Ran/Gsp1p) is believed to be required for this process, as defects in RNA export occur rapidly when yeast or mammalian cells carrying temperature-sensitive alleles of Gsp1p/Ran, its GTPase activating protein (Rna1p), or its nucleotide exchange factor (Prp20p) are shifted to the nonpermissive temperature (Amberg et al. 1992, 1993; Forrester et al. 1992; Kadowaki et al. 1992; Shiokawa and Pogo 1974; Wong et al. 1997). Proteins that associate with nascent pre-mRNAs within the nucleus and package them into heterogeneous nuclear ribonucleoprotein (hnRNP) particles contain nuclear export signals (NESs) (Michael et al. 1995). For example, the yeast hnRNP protein Npl3p...
has been shown to be a key mediator of RNA export signals (Lee et al. 1996). In addition, several yeast nucleoporins (Nup82p, Nup84p, Nup85p, Nup120p, Nup133p, Nup145p, Nup159p, and Npl4p) have roles specifically in RNA export, as strains harboring mutant alleles of some or disruptions of others show temperature-dependent defects in mRNA export but not in nuclear protein import (Doye et al. 1994; Fabre et al. 1994; Wente and Blobel 1994; Aitchison et al. 1995; Gorsch et al. 1995; Heath et al. 1995; Hurwitz and Blobel 1995; Kraemer et al. 1995; Li et al. 1995; Dehoratius and Silver 1996; Goldstein et al. 1996; Murphy et al. 1996; Siniossoglu et al. 1996; Dockendorff et al. 1997).

Competition experiments indicate that export from the nucleus of distinct classes of RNA molecules requires class-specific factors (Jarmolowski et al. 1994). The existence of such factors creates the potential to regulate nuclear transport so that only certain types of macromolecules are transported under certain conditions or to enable the transport of RNA molecules that would otherwise be retained in the nucleus. For example, HIV-1 Rev protein facilitates the export of intron-containing mRNAs (Fischer et al. 1994). The export activity of Rev requires the presence of a leucine-rich NES in Rev (Malim et al. 1991) and appears to target Rev response element (RRE)-containing and intron-bearing HIV-1 mRNAs to an export pathway that shares components with those required for export of U snRNAs and SS RNA, but not mRNA, tRNA, or rRNA (Fischer et al. 1995; Fritz et al. 1995).

Yeast Rip1p is a 42-kD polypeptide identified as a two-hybrid interactor with HIV-1 Rev (Stutz et al. 1995). Similar two-hybrid screens led to the identification of the related mouse and human proteins, Rab/hRIP (Bogerd et al. 1995; Fritz et al. 1995). Yeast Rip1p contains a unique carboxy-terminal domain of ∼80 amino acids. The amino-terminal 75% of Rip1p contains many XFXF repeats and is most closely related to the repeat domain of the yeast nucleoporin Rat7p/Nup159p (Gorsch et al. 1995; Kraemer et al. 1995). This sequence similarity suggests that Rip1p may also be a nucleoporin, and nuclear rim staining was seen using antibodies to Rip1p (Stutz et al. 1995). Mammalian hRIP/Rab was detected primarily in the nucleoplasm (Fritz et al. 1995). It is not yet clear whether yeast Rip1p and hRIP/Rab are homologs or whether Rip1p is a component of NPCs or an NPC-associated protein.

RNA export is regulated following heat shock and other forms of stress in S. cerevisiae (Saavedra et al. 1996). Following stress, poly(A)⁺ RNA accumulates within yeast nuclei, but stress also induces transcription of heat shock genes. Synthesis of inducible heat shock proteins requires that these heat shock mRNAs be exported to the cytoplasm. We showed previously that SSA4 mRNA, encoding stress-inducible Hsp70, is efficiently exported following heat or ethanol shock and that its export is unaffected by mutations in components of the Ran/Gsp1 system (Saavedra et al. 1996).

Here we report the identification of factors required for export of heat shock mRNAs following stress in yeast. Rip1p is required, but no defect in export of mRNA was seen under normal growth conditions (23°C–37°C) in cells disrupted at RIP1. In the absence of stress in cells lacking Rip1p, export of SSA4 mRNA expressed from a GAL promoter occurred efficiently, indicating that export of SSA4 mRNA requires Rip1p only following stress. High-level expression in yeast cells of HIV-1 Rev, but not of a mutant form of Rev lacking a functional NES, partially prevented the nuclear export of heat shock mRNAs. Export of heat shock mRNAs after stress was not dependent on functional Npl3p but did require Gle1p/Rss1p, an NES-containing protein essential for nuclear export of poly(A)⁺ RNA (Del Priore et al. 1996; Murphy and Wente 1996) and also shown by two-hybrid analyses to interact with Rip1p (Murphy and Wente 1996). The data suggest a model wherein the requirement for Npl3p defines the mRNA export pathway, the requirement for Rip1p defines a distinct pathway used for export of heat shock mRNAs after stress, and additional factors, including Rss1p/Gle1p and several nucleoporins (including Rat7p/Nup159p, Rat2p/Nup120p, Nup145p), are required in both pathways.

Results

Rip1p is essential for nuclear export of heat shock mRNAs following cellular stress.

In a previous report, we showed that celler stress (heat or ethanol shock) causes accumulation of poly(A)⁺ RNA within yeast nuclei but that heat shock mRNAs are transcribed and exported efficiently (Saavedra et al. 1996). During the course of those studies we learned that yeast cells lacking the small NPC-associated protein Rip1p are partially compromised for Rev-dependent export of RRE-containing RNAs but showed no defects in cellular growth or nuclear processes at either 23°C or 37°C (Stutz et al. 1995). We tested whether Rip1p might be involved in the selective export of heat shock mRNA following stress by comparing the patterns of localization of poly(A)⁺ RNA and SSA4 mRNA (expressed from a high-copy plasmid and encoding stress-inducible Hsp70) in wild-type cells and in cells deleted for RIP1. Interestingly, SSA4 mRNA accumulated in nuclei of Δrip1 cells heat-shocked at 42°C for 1 hr (Fig. 1). Very little SSA4 mRNA was visible when these cells where grown at 23°C (Fig. 1G) or shifted to 37°C (Fig. 1H). Furthermore, in Δrip1 cells shifted to 42°C for 1 hr, the poly(A)⁺ RNA signal was exclusively nuclear (data not shown), whereas in wild-type cells shifted to 42°C, a faint signal for poly(A)⁺ RNA is detected in the cytoplasm following heat shock (Saavedra et al. 1996; data not shown) and most likely reflects the export of a variety of heat shock mRNAs produced following stress. We also examined the effect of ethanol shock on export of SSA4 mRNA at 23°C in Δrip1 cells (Fig. 2). Following ethanol shock, SSA4 mRNA accumulated within nuclei of Δrip1 cells, indicating that the requirement for Rip1p for export following stress does not merely reflect a high-temperature threshold for inactivation of the export function of NPCs. Together, these results provide evidence that
mRNAs present prior to heat shock turned over. Note particularly the dark band migrating just below Hsp104p in Δrip1 cells; this band was nearly undetectable in wild-type cells heat-shocked for 15 min but persisted during the heat shock period in Δrip1 cells.

Those bands that represent heat shock proteins produced constitutively continued to be translated during stress, reflecting their preferential translation. Heat shock proteins that reach high levels only following heat shock (e.g., Ssa4p and Hsp104p) were produced in Δrip1 cells at levels substantially lower than in wild-type cells and with different kinetics. Although the rates of synthesis of induced heat shock proteins in wild-type cells were declining within 60 min of a shift to 42°C, their levels were still increasing in Δrip1 cells. Although cells maintained at a very low density (e.g., OD₆₀₀ < 0.10) contain essentially no SSA4 mRNA (Stutz et al. 1997), low levels of SSA4 mRNA are produced as cells reach higher densities, and it is probable that most of the Ssa4p synthesized following heat shock in Δrip1 cells derives from enhanced translation of SSA4 mRNA produced prior to heat shock. However, the increase in the rate of synthesis of Ssa4p 60 min after heat shock in Δrip1 cells may also reflect some export of SSA4 mRNA late in the heat shock period. The in situ hybridization results (Fig. 1) clearly show that most SSA4 mRNA is prevented from leaving the nucleus in Δrip1 cells. The data shown in Figure 3A indicate that the production of several other classes of heat shock proteins was dramatically reduced in Δrip1 cells, suggesting that Rip1p is important for the export of multiple classes of heat shock mRNA.

We also examined SSA4 mRNA levels using a primer extension assay (Fig. 3B). Analysis of RNA levels in each sample (Fig. 3B) indicated that primer extension analyses were performed using approximately equal cell equivalents of RNA. Very low but detectable levels of SSA4 mRNA were present in both wild-type and Δrip1 cells.
Because Rip1p appears to be a nucleoporin whose repeat region is not essential, we wondered whether disruption of RIP1 caused any abnormalities in NPC distribution that could be involved in transport defects. We compared wild-type cells and cells disrupted for RIP1 by indirect immunofluorescence using either an anti-Rat7p/Nup159p antibody or the RL1 monoclonal antibody that recognizes multiple repeat-containing yeast nucleoporins and observed a normal nuclear rim-staining pattern (C. Heath and C. Cole, unpubl.).

The essential yeast hnRNP protein Npl3p is not required for export of heat shock mRNAs

Npl3p is an essential mediator of the export of poly(A)+ RNA in yeast (Lee et al. 1996). To determine whether Npl3p is required for export of heat shock mRNAs, we examined the localization of poly(A)+ RNA and SSA4 mRNA in cells carrying the npl3-1 and npl3-17 alleles and expressing SSA4 from its own promoter from a high-copy plasmid. As reported previously, poly(A)+ RNA accumulated in the nuclei of npl3-1 cells shifted to 37°C for 1 hr (Fig. 5A, h), but the block to export was incomplete as some cytoplasmic signal remained. Following a shift to 42°C, poly(A)+ RNA was detected primarily in nuclei, indicating that less poly(A)+ RNA was exported in npl3-1 cells following heat shock than at the nonpermissive temperature of 37°C. Little or no SSA4 mRNA was detected at 23°C (Fig. 5A, a) or 37°C (Fig. 5A, b), whereas it was primarily cytoplasmic following heat shock at 42°C (Fig. 5A, c). This indicates that Npl3p is not required for efficient export of SSA4 mRNA following stress. Essentially identical results were obtained with the npl3-17 allele (data not shown), even though this is a tighter allele than npl3-1, and the defect in poly(A)+ RNA export at 37°C in this strain was more complete than in npl3-1 cells. We also examined the pat-

grown at 23°C. SSA4 mRNA increased dramatically following the shift to 42°C, but to a lower extent in ∆rip1 cells than in wild-type cells. This may reflect a lower stability for the SSA4 mRNA that accumulates in the nuclei of ∆rip1 cells. As a control, we also analyzed LYS7 mRNA. This mRNA declined from similar levels and with essentially identical kinetics in wild-type and ∆rip1 cells (Fig. 3B).

To determine whether export of SSA4 mRNA was dependent on Rip1p under all conditions or only following stress, we examined the localization of SSA4 mRNA expressed from the GAL1 promoter in ∆rip1 cells. No SSA4 mRNA was detected when cells were grown on glucose (data not shown) or raffinose (Fig. 4A). When cells were incubated for 2 hr at 23°C in the presence of 2% galactose, SSA4 mRNA was present throughout the cell (Fig. 4B). This indicates that SSA4 mRNA can be exported from the nucleus by at least two pathways: at 23°C, Rip1p is not required for export of SSA4 mRNA, and this mRNA may be exported using the conventional poly(A)+ RNA export machinery; at 42°C, SSA4 mRNA export requires Rip1p (Fig. 1) and is also independent of the Ran/Gsp1p GTPase machinery (Saavedra et al. 1996).
tern of protein synthesis in heat-shocked npl3-17 cells (Fig. 5B) and found it to be indistinguishable from the wild-type pattern. We conclude that Npl3p is not required for the export of heat shock mRNAs following heat shock.

High-level expression of HIV-1 Rev protein interferes with efficient export of SSA4 mRNA following stress

HIV-1 Rev possesses a nuclear export signal required for the export of Rev itself and RRE-containing mRNAs (Malim et al. 1991; Fischer et al. 1994, 1995). Microinjection into Xenopus oocyte nuclei of a high level of the leucine-rich Rev NES conjugated to bovine serum albumin (BSA) inhibited the nucleocytoplasmic export of 5S RNA and U small nuclear RNAs (snRNAs) but did not affect export of rRNA, tRNA, or poly(A)+ RNA (Fischer et al. 1995). Because export of SSA4 mRNA following stress is dependent on the presence of Rip1p (Fig. 1) and Rev functions to facilitate the export of RRE-containing mRNAs even in yeast (Stutz and Rosbash 1994), we examined the effects of high-level expression of HIV-1 Rev on poly(A)+ and SSA4 mRNA export in wild-type yeast cells. Plasmids encoding either wild-type Rev or the M10 export-defective Rev mutant, under control of the GAL10 promoter, were introduced into yeast cells carrying a high-copy SSA4 plasmid. The M10 Rev allele contains two missense mutations within the Rev activation domain/NES and does not facilitate the export of RRE-containing RNAs in either yeast (Stutz et al. 1995) or mammalian cells (Malim et al. 1991). Although Rev interacts efficiently with Rip1p and hRIP/Rab in two-hybrid assays (Bogerd et al. 1995; Fritz et al. 1995; Stutz et al. 1995), the ability of mutant Rev activation domains to function in vivo closely correlates with their ability to interact with hRIP/Rab/Rip1p in two-hybrid assays (Bogerd et al. 1995; Stutz et al. 1996).

We induced production of wild-type or M10 Rev by addition of 2% galactose for 4 hr. As assayed by Western blotting using an anti-Rev antiserum, both Rev and M10 were induced to comparable levels by this treatment (Fig. 6A). Following induction at 23°C, cells were shifted to 42°C for 30 min to induce transcription from the SSA4 promoter. Expression of wild-type Rev caused partial inhibition of the export of SSA4 mRNA (Fig. 6B, cf. panel G with panel C), whereas SSA4 mRNA export was normal in heat-shocked cells maintained in raffinose (Fig. 6B, panel C). Expression of the M10 mutant did not interfere with the export of SSA4 mRNA (Fig. 6B, panel O). This suggests that wild-type Rev is able to titrate an important mediator of SSA4 mRNA export following stress. We note that only a minority (∼30%) of cells expressing wild-type Rev show clear nuclear accumulation of SSA4 mRNA. Most likely, there is a correlation between Rev levels and the degree of accumulation of SSA4 mRNA in nuclei. Natural cell-to-cell variation in copy numbers for both the Rev and SSA4 plasmids likely results in substantial differences in actual Rev and SSA4 mRNA levels.

We also examined the distribution of poly(A)+ RNA in cells (23°C) induced to express wild-type Rev or the M10 mutant from a GAL promoter. Induction of Rev and M10 expression for 4 hr caused no detectable alteration in the distribution of poly(A)+ RNA (Fig. 6C), demonstrating that the partial inhibition of SSA4 mRNA export in cells expressing a high-level of Rev and shifted to 42°C is not likely to be attributable to any general inhibition of RNA export by Rev.

Rss1p/Gle1p is required for export of both poly(A)+ RNA under normal conditions and heat shock mRNAs following stress

Previously, we identified Rss1p as a high copy suppressor of the rat7-1 mutation (Del Priore et al. 1996). Murphy and Wente (1996) also identified this protein (which they called Gle1p) and demonstrated that Rss1p/Gle1p con-
tains a NES essential for its function. Strong nuclear accumulation of poly(A)⁺ RNA was seen when cells were depleted for Rss1p/Gle1p (Del Priore et al. 1996) or in cells carrying temperature-sensitive alleles of RSS1/GLE1 shifted to the nonpermissive temperature (Murphy and Wente 1996). To understand further the role of Rss1p, we prepared a series of temperature-sensitive alleles of RSS1 (V. Del Priore and C.N. Cole, unpubl.).

We examined mutant cells carrying the rss1-37 allele for their ability to export SSA4 mRNA when heat-shocked at 42°C (Fig. 7). There was modest nuclear accumulation of poly(A)⁺ RNA in mutant cells grown at 23°C (Fig. 7A, a) and strong accumulation of poly(A)⁺ RNA in the nuclei of 100% of the cells when they were shifted to 37°C for 1 hr (panel b). In this strain, SSA4 mRNA accumulated in the nuclei of cells shifted to 42°C (panel c), and there was little or no increase in the rate of synthesis of those heat shock proteins whose increased production requires export of newly transcribed heat shock mRNAs (Fig. 7B, cf. Fig. 3). This indicates that Rss1p is required for efficient export of RNAs both through the conventional mRNA export pathway and through the distinct pathway used for export of heat shock mRNAs following stress. We also examined the effect of loss of functional Rss1p/Gle1p on NPC distribution by comparing wild-type cells and cells carrying the rss1-37 allele. No changes in NPC distribution were seen in cells carrying the rss1-37 mutant allele incubated at 23°C or shifted to 37°C or 42°C for 1 hr (data not shown).

**Discussion**

Cells respond to stress at multiple levels to enable them to adjust to rapid changes in their environment and to recover from stress when stress conditions have passed (for review, see Lindquist 1986; Jost and Lindquist 1988; Lindquist and Petersen 1990; Morimoto et al. 1994; Morimoto 1993; Panniers 1994; Sierra and Zapata 1994). The experiments presented in this paper indicate that stress-induced heat shock mRNAs are exported through a pathway that is independent of several but not all of the factors required for normal mRNA export. A model for these partially overlapping transport pathways is...
shown in Figure 8. In S. cerevisiae, export of heat shock mRNAs (Fig. 8, left), but not bulk poly(A)⁺ RNA (Fig. 8, right), depends on the nucleoporin-like NPC-associated Rip1p (Fig. 1) but does not require Npl3p (Fig. 5), a key mediator of poly(A)⁺ RNA export (Lee et al. 1996) or components of the Ran/Gsp1p GTPase system (Saavedra et al. 1996). The NES-containing protein Rss1p/Gle1p is required for the export of both poly(A)⁺ RNA under non-stress conditions (Del Priore et al. 1996; Murphy and Wente 1996) and heat shock mRNAs following stress (Fig. 7). This factor may also have roles in the export of other classes of RNA. There are likely to be additional factors required for export of both bulk poly(A)⁺ RNA under normal conditions and SSA4 mRNA following stress, and these as yet unidentified factors may be required for export of other classes of RNA as well. In addition, there is likely to be at least one factor (X in Fig. 8) which binds specific elements in heat shock mRNAs and has a role analogous to that of Npl3p for export of poly(A)⁺ RNA under normal conditions. The same subset of nucleoporins required for export of poly(A)⁺ mRNA is also required for export of heat shock mRNAs following stress and may also be needed for export of other classes of RNA.

Previous evidence for the existence of multiple RNA export pathways comes primarily from studies in Xenopus oocytes, where it was shown that radiolabeled RNAs microinjected into nuclei were exported efficiently and that their export could be competed by unlabeled RNAs of the same class but not by RNAs of other classes (Jarmolowski et al. 1994). This indicates that, at least in Xenopus, the limiting export factors are specific for single classes of RNAs. Other studies focused on the HIV-1 Rev protein, a small RNA-binding protein that shuttles in and out of nuclei (Meyer and Malim 1994) and is able to facilitate the export of intron-containing mRNAs that contain sequences to which Rev can bind (RRE) (Fischer et al. 1994). BSA coupled to the Rev NES was exported following microinjection into Xenopus oocyte nuclei and was able to inhibit the export of co-injected U snRNA and 5S RNA but not mRNA, rRNA or tRNA (Fischer et al. 1995). These and other studies suggest that Rev directs intron-containing HIV-1 mRNAs to an RNA export pathway normally used for export of none-poly(A)⁺ RNAs (Fridell et al. 1996; Fritz and Green 1996), most likely 5S RNA and U snRNAs. The machinery involved in Rev-dependent export appears to be conserved among distantly related organisms, as Rev functions in S. cerevisiae in a manner that is also dependent on RREs within RNAs and the presence of a functional NES within Rev (Stutz and Rosbash 1994).

In the studies reported here, we used two assays to monitor RNA export. The first involves in situ hybridization to detect poly(A)⁺ (Amberg et al. 1992) or specific mRNAs (Saavedra et al. 1996). In situ assays are quite sensitive for detection of partial defects in RNA export, as modest accumulation of RNA in nuclei is readily seen, even when cytoplasmic RNA levels remain high. However, this assay is insensitive for determining...
whether a block in RNA export is complete or partial, as nuclear signals for poly(A)+ RNA differ only modestly when there is a complete versus incomplete block in RNA export. In addition, it is difficult to distinguish between low concentrations of cytoplasmic RNA and background fluorescence in in situ hybridization (Fay et al. 1997).

The second assay uses pulse-labeling of proteins. This assay complements the in situ hybridization assay, as it readily detects increased production of heat shock proteins, even when there is a partial block in export of heat shock mRNAs. However, it provides no information about the extent to which heat shock mRNAs are accumulating in nuclei under various conditions. Using both assays to examine RNA distribution and protein synthesis in various mutant strains and under various environmental conditions has allowed us to determine whether various gene products are required for export of heat shock mRNAs following stress.

The radiolabeling assay also demonstrated that export of mRNAs encoding other classes of heat shock proteins (e.g., Hsp104p and Hsp82p) occurred preferentially following stress and was dependent on the same factors (Rip1p, Rss1p/Gle1p) and nucleoporins (Rat7p/Nup159p, Rat2p/Nup120p and Nup145p; Saavedra et al. 1996) as export of SSA4 and SSA1 mRNAs. Thus, heat shock mRNAs as a class appear to share an RNA export mechanism. Stutz and Rosbash observed a partial reduction in Rev-dependent export in yeast cells lacking Rip1p (Stutz and Rosbash 1994; Stutz et al. 1995), suggesting that Rev mediates export of RRE-containing RNAs through both Rip1p-dependent and Rip1p-independent pathways. Coupled with the observation that the Rev NES can inhibit export of U snRNAs and 5S RNAs in Xenopus, these results suggest that a Rip1p-dependent RNA export pathway is used for export of both heat shock mRNAs following stress and for other classes of RNA (perhaps 5S RNA) but that a separate pathway, independent of Rip1p, is available, at least under normal growth conditions, for export of those classes of RNA that use the Rip1p-dependent pathway during normal growth.

Inhibition of export of heat shock mRNAs by high-level expression of HIV-1 Rev

The partial inhibition of SSA4 mRNA export when Rev was expressed from a strong inducible promoter (Fig. 6) further strengthens the hypothesis that heat shock mRNAs exit the nucleus via a pathway that is distinct from that normally used for poly(A)+ RNA export. What might be the mechanism by which Rev expression partially inhibits export of heat shock mRNAs? Because wild-type Rev, but not the M10 mutant, interacts with Rip1p in the two-hybrid system, high-level expression of Rev could titrate Rip1p, thereby reducing export of heat shock mRNAs. However, it now appears that the interaction of Rev and Rip1p may be indirect, as two-hybrid interactions between Rev and Rip1p were eliminated by loss-of-function mutations in Crm1p (M. Neville et al. 1997). Most likely, Rev titrates one or more cellular NES-binding proteins. We tested whether overexpression of Rss1p/Gle1p, which contains a Leu-rich NES similar to Rev’s, would affect RNA export but saw no effect on heat shock mRNA export induced following a shift to 42°C or on poly(A)+ RNA export at 23°C or 37°C (C. Heath and C. Cole, unpubl.). Although Rev expression had no effect on export of poly(A)+ RNA within the first 4 hr of induction (Fig. 6C), cells containing either Rev or the M10 mutant under control of the GAL10 promoter failed to form colonies on galactose plates, indicating eventual growth inhibition by both Rev and M10 (C. Hammell and C. Cole, unpubl.). In contrast, little or no growth inhibition was seen when Rss1p was expressed from a 2μ plasmid, or when cells containing a galactose-inducible RSS1 plasmid were grown on galactose (Del Priore et al. 1996; C. Hammell, C. Heath, and C. Cole, unpubl.). Possibly Rss1p/Gle1p levels never reach those attained by Rev or M10. Alternatively, Rev may titrate a critical cellular factor that does not interact with Rss1p/Gle1p. Another possibility is that Rev interacts more strongly with a factor that also interacts with Rss1p/Gle1p.

Interestingly, Rss1p/Gle1p is required for both bulk poly(A)+ RNA export under normal growth conditions (Del Priore et al. 1996; Murphy and Wente 1996) and export of heat shock mRNAs after stress (Fig. 7). Because Rss1p/Gle1p contains an NES similar to that of HIV-1 Rev (Murphy and Wente 1996), this protein is most likely a soluble export factor that shuttles between the nucleus and the cytoplasm. Because ts mutations of RSS1 affect heat shock export dramatically (Fig. 7), we suspect that Rss1p/Gle1p plays a similar role in export of all mRNAs, and perhaps in export of other classes of RNA as well.

Following the observation of Stutz et al. (this issue), we found that rss1-37 was synthetically lethal with disruption of RIP1 (C. Heath and C. Cole, unpubl.). This means that in the presence of ts alleles of RSS1, Rip1p becomes essential for growth even at the permissive temperature (23°C). This could reflect a nonessential role for Rip1p during normal mRNA export. However, there are other possible explanations. We do not yet know whether Rss1p is important for export of other classes of RNA, and it could well be involved in all RNA export. Assays to examine export of other classes of RNA in S. cerevisiae have not yet been developed. Some RNAs could exit the nucleus using both Rip1p-dependent and Rip1p-independent pathways. If cells lack Rip1p, export solely through the other pathway may be insufficient to allow growth in the presence of mutant alleles of RSS1. Rip1p may be a structural component of NPCs, most likely anchored within NPCs through its unique carboxyl terminus. NPCs lacking Rip1p may be structurally altered so that export efficiency is compromised, perhaps through important binding sites on certain nucleoporins becoming less accessible. Under these conditions, mutations of Rss1p, or other factors important for RNA export, might readily lead to insufficient export under permissive conditions, and thus to the observed synthetic
lethality. The observation that the carboxy-terminal domain of Rip1p, lacking all nucleoporin repeats, suppresses the synthetic lethality between Δrip1p and temperature-sensitive mutations of RSS1 (Stutz et al. 1997) is consistent with this explanation. Murphy and Wente (1996) isolated GLE1/RSS1 because of its synthetic lethality with a disruption of NUP100, which encodes a nonessential nucleoporin. We tested Δnup100 cells for defects in export of heat shock mRNA after stress and saw no differences from wild-type; the same result was obtained with disruption of NUP2, another nonessential nucleoporin (C. Heath, C. Hammell, and C. Cole, unpubl.).

Several important questions remain regarding differential export of mRNA following stress. One concerns the mechanism by which export of most poly(A)+ RNAs (Saavedra et al. 1996) and import of some nuclear proteins (Liu et al. 1996; Saavedra et al. 1996) are blocked. In addition, we do not know whether the pathway used for export of heat shock mRNAs following stress contains any components that must be activated for the pathway to function. Although we have reported that the Gsp1p/Ran system is not required for export of heat shock mRNAs after stress (Saavedra et al. 1996), recent observations in our laboratory indicate that this system remains functional under these conditions, so its inactivation does not appear to be part of the mechanism for altering nucleocytoplasmic transport following stress (C. Saavedra, C. Hammell, and C. Cole, unpubl.). Another question concerns the nature of the factors that recognize heat shock mRNAs and presumably play a critical role in mediating the export of heat shock mRNAs. Finally, it will be necessary to determine the identity of the factors that play the same role for export of heat shock mRNA that the Ran/Gsp1p system performs for export of most classes of RNA.

Materials and methods

Strains, plasmids, and growth conditions

Yeast strains are listed in Table 1. Except where noted, cells were grown to early exponential phase at 23°C in YPD-rich medium or in synthetic complete medium lacking leucine (SC –Leu) or uracil (SC –Ura) (Rose et al. 1989). For induction of the heat shock response, cells in the exponential phase of growth (5 x 10^6 to 2 x 10^7 cells/ml) were transferred to water baths at the appropriate temperature. In some experiments, the stress response was also induced by the addition of ethanol to 10% (vol/vol). Strains FY23 and FY86, derived from S288C, were used as wild types (Winston et al. 1995). Yeast transformations were performed by electroporation using a Gene-Pulser (Bio-Rad Laboratories, Melville, NY); cells were allowed to recover in rich medium containing 1 M sorbitol at 23°C for at least 1 hr before placing. E. coli strain DH5α was used for most cloning procedures. Plasmids used in these studies are listed in Table 2. A strain containing an integrated copy of the npl3-17 allele was generously provided by Margaret Lee and Pamela Silver (Dana Farber Cancer Institute, Boston, MA).

Construction of a strain disrupted for RIP1

A PCR-based gene deletion approach (Baudin et al. 1993) was used to disrupt the RIP1 gene. We generated a HIS3 cassette by PCR amplification that was flanked at its ends by 45 nucleotides identical to sequences just upstream and downstream of the RIP1 open reading frame (ORF). The upstream oligonucleotide sequence was 5′-GTAATGTCAGCTTCCGGTACCCATTCACGTCGGGTGCTAAGCCTGGCCTCCTGCCTTCAGGTATATCAGGGTCGCGCGCCTCGTTCAGAATGGAGGTAT 3′ and the downstream oligonucleotide sequence was 5′-CAGGATATCATCAGGTCGCCGCGCTTCCAGTTTCCAGAATGGGGAGCTTCTATGCAACCAATGCAGGTGGTGTAATGTCAGCTTCCGGTACCC ATTCACGTCGGGTGCTAAGCCTGGCCTCCTGCCTTCAGGTATATCAGGGTCGCGCGCCTCGTTCAGAATGGAGGTAT 3′. For both oligonucleotides, the last 19 nucleotides are homologous to the HIS3 selectable marker. We used 20 ng of the HIS3-containing plasmid pBM2815 (obtained from P. Silver, Dana Farber Cancer Institute, Boston, MA) to generate the deletion construct in the PCR reaction. The 50 µL PCR reaction consisted of 1× PCR buffer (10 mM Tris-HCl, 15 mM MgCl₂, 500 µM dNTPs, and 2 units of Taq DNA polymerase (Boehringer Mannheim Biochemicals, Indianapolis, IN). The reaction was subjected to a 30-cycle amplification consisting of 1.5 min at 94°C, 2.0 min at 50°C, and 2.0 min at 72°C. The PCR product was transformed into a wild-type haploid strain FY86, containing the deletion allele his3Δ200. The flanking upstream and downstream sequences target the deletion construct to the RIP1 such that its ORF is replaced with HIS3 by homologous recombination. Recombinants were selected on SC –His plates, and recombination was verified by PCR.

Table 1. Yeast strains used in this study

| Strain | Genotype | Source |
|--------|----------|--------|
| FY23   | MATa trp1Δ63 leu2Δ1 ura3-52 | Winston et al. (1995) |
| FY86   | MATa his3Δ200 leu2Δ1 ura3-52 | Winston et al. (1995) |
| ACY1   | MATα/MATα his3Δ200/Δ200 leu2Δ1/Δ1 ura3-52/Δ52 trp1Δ63/TRP1 | Anita Corbett (Dana Farber Cancer Institute, Boston, MA) |
| CHY119 | MATα rip1::HIS3 his3Δ200 ura3-52 leu2Δ1 | this study |
| PSY361 | MATα npl3-1 trp1Δ901 ade2-101 his3Δ200 ura3-52 lys4-519 | Pamela Silver (Dana Farber Cancer Institute, Boston, MA) |
| PSY1026| MATα npl3-17 ade2-101 can1-100 his3Δ200 leu2Δ1 lys1-1 ade8 | Margaret Lee (Dana Farber Cancer Institute, Boston, MA) |
In situ hybridization assays.

Stains that grew at 23°C but not at 37°C were selected for further plating to SC−Leu plates and incubated at 23°C and 37°C. Mu-}

copy of containing medium to select for loss of the plasmid-borne wild-type up on the SC−Leu plates were replica plated onto 5-FOA con-

PCR fragment was transformed into VDPY111, and the trans-
gapped vector was gel-isolated. A mixture of the vector and the 2854 GENES & DEVELOPMENT

dgTP, 0.2 mM dATP, 1 mM dCTP, and 1 mM dTTP. These

resulting 350-bp DNA fragments were subcloned into pDAD-1 wild-type and M10 mutant Rev were amplified by PCR, and the

S Plasmids containing the ORFs of Rev and M10 mutant Rev, construction of plasmids for inducible expression of HIV-1

Gene between base pair 246 and 1966 was PCR amplified using the following

Construction of plasmids for inducible expression of HIV-1 Rev and M10 mutant Rev

Plasmids containing the ORFs of Rev and M10 mutant Rev, fused to glutathione S-transferase, were obtained from Michael Malim (University of Pennsylvania, Philadelphia). The ORFs of wild-type and M10 mutant Rev were amplified by PCR, and the resulting 350-bp DNA fragments were subcloned into pDAD-1 between the HindIII and EcoRI sites in its polylinker region, thereby placing Rev and M10 Rev under control of the yeast GAL10 promoter.

Construction of the RSS1 temperature-sensitive alleles

The RSS1 temperature-sensitive alleles were made using PCR mutagenesis (Cadwell and Joyce 1992). The RSS1 gene between base pair 246 and 246 was PCR amplified using the following primers: 5'-GGCCGTTGCTACAGGATATTC-3' and 5'-GGTATATAATTGATAACAAGAGAATCGTCG-

8

and 5

8

base pair 246 and 1966 was PCR amplified using the following

Gels were then dried and exposed to a Phosphor screen (Molecular Dynamics) and imaged using ImageQuant software (Molecular Dynamics).

Western analyses

Cells were grown to a total OD 600 of 0.75, pelleted by centrifugation at 2500 rpm for 2.5 min, and washed once with water. Two hundred micrograms of acid-washed glass beads and 200 µl of hot sample buffer (62.5 mM TrisCl at pH 6.8, 2% SDS, 10% glycerol, 8 mM urea, 0.72 mM 2-mercaptoethanol, 0.05% bromophenol blue) were added to the cell pellets. After brief vortexing, cells were then lyzed as follows: 10 sec vortexing and 50 sec incubation in the boiling water bath, repeated five times over 5 min. The supernatant was then transferred to a clean Eppendorf tube, and equal volumes of cell lysate were loaded onto a 10% polyacrylamide gel. Electrophoresis was carried out at 200 V for 50 min. Proteins in the gel were transferred to a PVDF membrane by electroblotting overnight at 100 mA in a cold room. The membrane was washed briefly with 1x phosphate-buffered saline (PBS), 0.1% Tween 20 solution (solution A), and then blocked for 1 hr with 1x PBS, 0.1% Tween 20, and 5% nonfat milk solution (solution B). The membrane was then incubated with anti-Rev antiserum (kindly provided by Dr. Michael Malim, Howard Hughes Medical Institute, University of Pennsylvania, Philadelphia) at a 1:5000 dilution in solution B for 2 hr followed by four washes of 10 min each with solution A. The membrane was then incubated with anti-rabbit antibody.

Cells were grown at 23°C in SC−Met and containing 0.85 grams/liter of yeast nitrogen base without amino acids and ammonium sulfate and 3.3 grams/liter of yeast nitrogen base with-
coupled to horseradish peroxidase (Amersham, Inc., Arlington Heights, IL), diluted 1:3000 in solution B for 1 hr. Washes were performed as described above. Signal was developed with the ECL kit (Amersham, Inc.) according to the manufacturer’s directions.

Analysis of SSA4 mRNA levels by primer extension

Both wild-type and Δprp1 cells were grown to an OD_{600} of 0.5. RNA extractions were performed as described by Pikielny and Rosbash (1985). The total RNA isolated from each sample was dissolved in 20 µl of distilled water. Identical aliquots from each sample were subjected to electrophoresis in a 1% agarose gel and stained with ethidium bromide to permit visualization of rRNAs. A primer with the sequence GTTGTGACCTAAATCAATACC was used to specifically measure SSA4 mRNA levels. LYS7 mRNA levels were measured with primer AGGGGC-TAGCTTCA. Primer extension assays were performed using 6 µl of each isolated RNA sample. Both primers were added to each RNA sample and primer extension performed as described (Pikielny and Rosbash 1985). Extension products were analyzed on a 12% denaturing polyacrylamide gel. The relative levels of extension products were quantified using a Molecular Dynamics PhosphorImager using ImageQuant software.

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