A dynamic transcriptional network communicates growth potential to ribosome synthesis and critical cell size

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Cell-size homeostasis entails a fundamental balance between growth and division. The budding yeast Saccharomyces cerevisiae establishes this balance by enforcing growth to a critical cell size prior to cell cycle commitment (Start) in late G1 phase. Nutrients modulate the critical size threshold, such that cells are large in rich medium and small in poor medium. Here, we show that two potent negative regulators of Start, Sfp1 and Sch9, are activators of the ribosomal protein (R) and ribosome biogenesis (Rib) regulons, the transcriptional programs that dictate ribosome synthesis rate in accord with environmental and intracellular conditions. Sfp1 and Sch9 are required for carbon-source modulation of cell size and are regulated at the level of nuclear localization and abundance, respectively. Sfp1 nuclear concentration responds rapidly to nutrient and stress conditions and is regulated by the Ras/PKA and TOR signaling pathways. In turn, Sfp1 influences the nuclear localization of Fhl1 and Ihf1, which bind to R gene promoters. Starvation or the absence of Sfp1 causes Fhl1 and Ihf1 to localize to nucleolar regions, concomitant with reduced R gene transcription. These findings suggest that nutrient signals set the critical cell-size threshold via Sfp1 and Sch9-mediated control of ribosome biosynthetic rates.

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Cell-size homeostasis requires that proliferating cells coordinate growth and the cell cycle, such that each division is matched by a doubling of mass. Cell growth and the cell cycle are, however, separate processes that can be uncoupled, either by endogenous developmental mechanisms, for instance, during oogenesis and the rapid cleavage cycles following fertilization, or within the laboratory. In eukaryotic cells, doubling time is usually limited, not by the time required to duplicate and divide the genome, but rather by the time required to double cell mass. Consistently, cell cycle progression is typically dependent on cell growth. In contrast, mass accumulation continues unabated during most cell cycle arrests (Saucedo and Edgar 2002).

In the unicellular budding yeast, growth and the cell cycle are coupled at Start, the short interval during late G1 phase after which cells are committed to division (Hartwell et al. 1974). Passage through Start has several requirements as follows: (1) growth to a critical cell size (Johnston et al. 1977), (2) nutrient sufficiency (Hartwell et al. 1974), (3) attainment of a critical translation rate (Hartwell and Unger 1977; Moore 1988), and for haploids, (4) absence of mating pheromone (Hartwell et al. 1974). The first three conditions are likely to be interrelated. The critical size requirement and minimum translation rate explain why slowing growth rate increases the length of G1 phase, whereas the time required to transit the rest of the cell cycle is relatively constant (Hartwell and Unger 1977). The critical-size threshold maintains uniform cell size over many generations, and under minimal nutrient conditions forces cells to accumulate the energy stores required to complete the division cycle. At constant nutrient levels, the critical-size requirement couples growth and division only in daughter cells, as mother cells by definition have already attained critical size.

Start depends on activation of the SBF and MBF transcription factor complexes that bind the promoters of G1/S-regulated genes (Nasmyth 1996). SBF and MBF are composed of related DNA-binding proteins Swi4 and Mbp1, respectively, which interact with a common regulatory subunit Swi6 to drive expression of a massive suite of ~200 genes. Among these, the key transcripts are
the G1 cyclins *CLN1* and *CLN2* and the B-type cyclins *CLB5* and *CLB6*. Cln1 and Cln2 activate Cdc28, the cyclin-dependent kinase [CDK] that controls all cell cycle transitions in budding yeast. Cln1/2–Cdc28 phosphorylation events trigger bud emergence and inactivate Sic1 and Cdh1, two key inhibitors of B-type cyclin–Cdc28 activity. Once unleashed, Clb5/6–Cdc28 complexes initiate DNA replication [Nasmyth 1996]. Start can therefore be viewed as the short interval required to accumulate sufficient Cln1/2–Cdc28 activity to phosphorylate Cdh1 and Sic1, and perhaps other substrates.

The connection between cell size and SBF/MBF activation remains enigmatic. It has recently been discovered that SBF, and likely MFB, activation occurs upon dissociation of the Start-repressor Whi5 [Jorgensen et al. 2002; Costanzo et al. 2004, de Bruin et al. 2004]. Loss of the interaction between SBF/MBF and Whi5, which correlates with movement of Whi5 to the cytoplasm, is driven by phosphorylation of Whi5 by Cln3–Cdc28 [Costanzo et al. 2004, de Bruin et al. 2004]. Cln3 appears to act in parallel to Bck2, a poorly understood activator of SBF/MFB; as cells lacking *CLN3* and *BCK2* are inviable due to permanent G1 arrest [Wijnen and Futcher 1999]. Remarkably, deletion of WHI5 bypasses this arrest [Costanzo et al. 2004, de Bruin et al. 2004]. Because Start activation is highly sensitive to Cln3 dosage, it has been presumed that a critical concentration of Cln3–Cdc28 activity triggers Start. Although Cln3 protein and Cln3–Cdc28 kinase activity do not increase during G1 phase [Tyers et al. 1993; Melmerny et al. 1997], a size-dependent increase in Cln3 nuclear abundance may help activate Start [Futcher 1996; H. Wang et al. 2004].

Nutrients modulate the critical cell-size threshold in proportion to the proliferation rate [Johnston et al. 1979; Lorincz and Carter 1979; Tyson et al. 1979]. The effects of nutrients on critical cell size are conveyed rapidly, as cells are not committed to Start at a given size threshold until just before the threshold is reached [Lorincz and Carter 1979]. Thus, shifting cells from poor to rich medium temporarily increases the fraction of unbudded G1-phase cells as these cells grow to the new threshold [Johnston et al. 1979]. Nutrient effects are mediated in part by the Ras/PKA pathway; decreased Ras/PKA signaling, as in a *cdc25-1* strain, decreases critical cell size, whereas hyperactive Ras/PKA signaling, as in a *ras2va121* strain, increases critical cell size [Baroni et al. 1989]. Strains that exhibit constitutive PKA activity do not adjust cell size in response to carbon source quality [Tokiwa et al. 1994].

Ribosome biogenesis is a chief occupation of growing cells, accounting for >50% of total transcription in yeast and mammalian cells [Warner 1999; Moss and Stepanovsky 2002]. In yeast, the rate of ribosome synthesis is dictated by the rate of transcription of the RNA and protein subunits of the ribosome [Warner 1999]. tRNA and *RP* gene transcription is exquisitely sensitive to the growth potential of the cell and is rapidly repressed in response to a wide variety of internal and external stresses [Ju and Warner 1994; Warner 1999; Gasch et al. 2000]. The 137 *RP* genes, referred to here as the *RP* regulon, are tightly coregulated. The vast majority of genes in the *RP* regulon have promoter-binding sites for Rap1, whereas a few have sites for Abf1 [Warner 1999]. As Rap1 silences telomeric repeats but activates glycolytic and *RP* genes [Warner 1999], its activity depends on context, presumably generated by combinations of associated coregulators. The *RP*-specific transcription factors that cooperate with Rap1 have not yet been identified.

A large number (>200) of other genes, which we term the *Ribi* regulon, show nearly identical transcriptional responses as *RP* genes to environmental or genetic perturbations [Gasch et al. 2000; Hughes et al. 2000, Wade et al. 2001; Jorgensen et al. 2002; Miyoshi et al. 2003]. The promoters of these coexpressed genes are strongly enriched for the presence of two motifs, termed RRPE and PAC, and therefore appear to constitute a distinct regulon [Gasch et al. 2000; Hughes et al. 2000, Wade et al. 2001, Jorgensen et al. 2002]. Most of these genes encode proteins involved in ribosome biogenesis, a process involving more than 100 accessory factors that assemble and modify rRNA and RPs in the nucleolus [Hughes et al. 2000; Wade et al. 2001; Fatica and Torlervy 2002, Jorgensen et al. 2002]. A number of additional functional categories are present in this regulon, including subunits of RNA Polymerase I and III, enzymes involved in ribonucleotide metabolism, tRNA synthetases, and translation factors [Gasch et al. 2000; Wade et al. 2001; Jorgensen et al. 2002]. The *Ribi* regulon thus consists of non-RP genes that boost translational capacity. Two central nutrient-signaling conduits, the Ras/PKA and TOR signaling pathways, can activate tRNA, *RP*, and *Ribi* transcription [Klein and Struhl 1994; Neuman-Silberberg et al. 1995; Cardenas et al. 1999; Hardwick et al. 1999; Powers and Walter 1999; Y. Wang et al. 2004]. Of these, the control of rRNA synthesis is best understood, as it is known to depend largely on phosphorylation of the critical initiation factor TIF-1A/Rrn3 [Grummt 2003]. Both Rap1-binding sites and RRPE elements can render gene transcription sensitive to Ras/PKA signaling, although the presumed kinase targets have not been identified [Klein and Struhl 1994; Neuman-Silberberg et al. 1995; Y. Wang et al. 2004].

Systematic determination of cell-size distributions for all yeast deletion strains has recently identified many new potential Start regulators [Jorgensen et al. 2002; Zhang et al. 2002]. Many of the genes that encode potential Start repressors are implicated in ribosome biogenesis, suggesting not only a link between these two seemingly disparate systems, but also that cell-size mutants may identify new regulators of ribosome synthesis [Jorgensen et al. 2002]. Beginning with the most potent size regulators, Sfp1 and Sch9, we have elaborated a dynamic transcriptional network that dictates the ribosome synthesis rate and the critical cell-size threshold.

**Results**

*Sfp1* and *Sch9* strongly influence cell size

Of all nonessential genes, deletions in either *SFP1* or *SCH9* caused the greatest decrease in cell size [Whi phe-
notype]. These deletion strains also proliferate slowly and accumulate cells in G1 phase, suggesting SFP1 and SCH9 are important for mass accumulation [Fig. 1A] [Jorgensen et al. 2002]. Sfp1 is a putative transcription factor with a split zinc-finger domain at its C terminus that is essential for function [Fingerman et al. 2003]. Sch9 is an AGC family kinase that possesses a C2 lipid-binding domain and is the closest yeast homolog of the metazoan pro-survival kinase Akt/ PKB [Fabrizio et al. 2001; Jorgensen et al. 2002]. Sch9 kinase activity is required for its function [data not shown]. To conditionally overexpress Sch9, we constructed a strain bearing a galactose-inducible allele integrated at the endogenous locus [GAL1–SCH9]. Like SFP1, overexpression of SCH9 caused a large cell phenotype, but unlike SFP1, SCH9 overexpression was not toxic [Fig. 1A]. To enable conditional control of Sch9 activity at endogenous levels without confounding nutrient source shifts, we engineered an analog-sensitive allele of SCH9 at the endogenous locus [sch9αα]. Substitution of a bulky amino acid with either alanine or glycine in the ATP-binding pocket of kinases often permits highly specific inhibition by analogs of the cell-permeable kinase inhibitor PP1, such as C3-1'-naphthylmethyl PP1 (1NM-PP1) [Bishop et al. 2000]. Unexpectedly, the integrated sch9αα allele separated the proliferation and size functions of SCH9, conferring a nearly wild-type proliferation rate, but a strong Whi phenotype [Fig. 1B]. Sch9αα was almost fully inhibited in vivo by 100 nM 1NM-PP1, as determined by effects on doubling time and cell size [Fig. 1B]. At this concentration, 1NM-PP1 had no effect on wild-type cells [data not shown] [Bishop et al. 2000].

Sfp1 and Sch9 activate the RP and Ribi regulons

We have previously shown that Sfp1 activates the RP and Ribi regulons [Jorgensen et al. 2002; Fingerman et al. 2003]. Further expression profiles of GAL1–SFP1 and sfp1Δ/Δ strains elaborated the Ribi regulon to 236 genes [Fig. 1C; Supplementary Table 1]. We next asked whether Sch9 also controls these genes. Addition of 100 nM 1NM-PP1 resulted in a rapid repression [≤30 min] of the RP regulon [2.5-fold] and the Ribi regulon [1.5-fold] in an sch9αα strain, followed by a slight but reproducible

Figure 1. Sch9 and Sfp1 regulate cell size and RP and Ribi regulon transcription. [A] Galactose-regulated alleles of SFP1 and SCH9. [B] Inactivation of an analog-sensitive allele of SCH9 by 1NM-PP1. A total of 100 nM 1NM-PP1 or DMSO solvent was added at 0 min to log phase, glucose cultures of sch9αα and sch9Δ. Cell-size distributions were measured after 6 h and compared with reference distributions. Without 1NM-PP1, the sch9αα allele is hypomorphic, with nearly wild-type doubling time [tD], but a strong Whi phenotype [tD 96 min, median cell size: 30 fl, vs. 87 min, 42 fl for wild type and 153 min, 28 fl for sch9Δ]. [C] Sch9 and Sfp1 activate the RP and Ribi regulons. Expression profiles for GAL1–SFP1 and sfp1Δ/Δ were repetitions of experiments described previously [Jorgensen et al. 2002]. Expression profiles were determined for sch9αα and wild-type cultures harvested at indicated times after addition of 100 nM 1NM-PP1; expression ratios are relative to untreated cells from the same culture. The presence of RAP1, PAC, RRPE, or PAC + RRPE [P+R] promoter elements is indicated [Filpel et al. 2001; Jorgensen et al. 2002]. [D] Induction of representative genes in the Ribi [NOP1, NRSl] and RP [RPPO, RPLl18] regulons upon restoration of GAL1–SCH9 or GAL1–SFP1 expression. An ACT1 loading control and GAL7, SFP1, and SCH9 induction controls are shown. [E] Comparison of gene sets regulated by Sfp1 and Sch9. Expression profiles of sfp1Δ/Δ and inactivated sch9αα [1NM-PP1, 90 min] were plotted against one another. The number of genes in each functional group is indicated.
recovery of both regulons (Fig. 1C). These transcriptional effects preceded effects of 1NM-PP1 on proliferation rate and cell size [Fig. 1B]. Northern analysis confirmed that representative members of the RP [RPP0, RPL11B] and Ribi [NOP1, NSR1] regulons were induced upon expression of GAL1–SCH9 (Fig. 1D). Sfp1 and Sch9 are thus both required for maximal RP and Ribi expression. However, inhibition of sch9ΔΔ also induced 38 genes not altered in sfp1ΔΔ cells [Fig. 1E], almost all of which were strongly induced as cultures deplete glucose and enter stationary phase [DeRisi et al. 1997]. Many stationary phase genes are also repressed by Ras/PKA, consistent with the genetic placement of Sch9 parallel to Ras/PKA [Toda et al. 1988].

Sfp1 and Sch9 are negative regulators of Start

A number of observations suggested that Sfp1 and Sch9 might regulate the critical cell size threshold at Start. First, sfp1ΔΔ and sch9ΔΔ cells are much smaller than other deletion mutants with the same proliferation rate [Jorgensen et al. 2002]. Second, the hypomorph alleles sfp1ΔΔ [Jorgensen et al. 2002] and sch9ΔΔ caused marked decreases in cell size with little increase in doubling time, similar to mutations in canonical Start regulators [Fig. 1A,B]. Third, increasing the doubling time of wild-type cells to that of sfp1Δ or sch9Δ cells by treatment with sublethal doses of cycloheximide reduced the size of new-born daughter cells, but not average cell size [Fig. 2A]. We directly tested the role of Sfp1 and Sch9 at Start by synchronous release of G1-phase daughter cells obtained by centrifugal elutriation. The slow rate at which daughter cells lacking Sfp1 or Sch9 activity accumulated volume underscored the role of these gene products in cell growth [Fig. 2B]. All aspects of Start, including the activation of SBF- and MBF-dependent transcription, bud emergence, and DNA replication initiation were accelerated with respect to cell size in strains that lacked Sfp1 or Sch9 activity [Fig. 2C–G]. For example, whereas wild-type cells expressed the MBF-regulated gene RNR1 at a size of ~30 fL, cells that lack Sfp1 or Sch9 activity do so at sizes of ~16 fL and ~21 fL, respectively. These results indicate that Sfp1 and Sch9 lie upstream of the G1/S transcriptional machinery. In contrast, treatment of wild-type cells with a sublethal dose of cycloheximide that slowed growth to the same extent as loss of Sch9 actually delayed Start, as measured by all of the above parameters [Fig. 2B–D,H]. We note that the absence of Sfp1, and to a lesser extent Sch9, caused considerable heterogeneity in traversal through Start, as shown by the extended window of G1/S transcription and the slow accumulation of cells with buds and G2 DNA content. This effect may arise from a reduced translation rate, as cycloheximide-treated wild-type G1 phase cultures behaved in a similar manner. The heterogeneity in Start progression is not, however, caused by heterogeneous cell growth rate [Fig. 2D].

Figure 2. Sfp1 and Sch9 are negative regulators of Start. [A] Size distributions of sfp1ΔΔ and sch9ΔΔ strains. Doubling times were as follows: sfp1Δ Δ 220 ± 13 min, sch9ΔΔ 153 ± 3 min, wild-type [WT] 89 ± 2 min, wild-type [WT] + 200 nM cycloheximide 149 ± 4 min. [B–H] Determination of critical cell size. Small G1-phase daughter cells (>97% unbudded) were isolated from late log-phase cultures [3–4 × 10⁶ cells/mL, raffinose medium with no drugs] of wild-type, sfp1ΔΔ and sch9ΔΔ strains by centrifugal elutriation and reinoculated in glucose medium. A total of 100 nM 1NM-PP1 was added to the sch9ΔΔ culture upon reinoculation; 200 nM cycloheximide was added to a wild-type culture upon reinoculation. [B] Daughter cell size at various times after reinoculation for each culture. [C] Bud index as a function of cell size for each culture. [D] Cell-size distributions at the >25% budded time point for each culture. Passage through Start was monitored by cell size, bud index, and expression of SBF [CLN2]- and MBF [RNR1]-specific transcripts for wild-type cells [E], sch9ΔΔ cells in 100 nM 1NM-PP1 [F], sfp1ΔΔ cells [G], and wild-type cells in 200 nM cycloheximide [H]. The time point at which cultures were >25% budded is highlighted in pink. All data was reproduced in duplicate experiments. 1NM-PP1 [100 nM] had no effect on the critical cell size of a wild-type strain [data not shown].
Strains lacking SFP1 or SCH9 are impervious to carbon source effects on cell size

If Sfp1 and Sch9 mediate the effects of nutrients on the critical cell size threshold, strains lacking these gene products should be largely impervious to nutrient modulation of cell size. The size distributions of sfp1Δ/Δ and sch9Δ/Δ diploid cells were, in fact, barely reduced on poor carbon sources, unlike wild-type and whi5Δ/Δ cells [Fig. 3A,B]. sfp1Δ/Δ cells appeared to be at near minimal cell size, as overexpression of the hyperactive CLN3-1 allele had little impact on the mean size of these cells [Fig. 3B]. Importantly, sfp1Δ and sch9Δ strains proliferated much more slowly than wild type on all carbon sources examined [data not shown], unlike strains that lack the large ribosomal subunit RPL11B gene, which is only rate-limiting for growth when cells are in glucose medium [Zhao et al. 2003].

In contrast to sfp1Δ and sch9Δ cells, the size of cln3Δ bck2Δ whi5Δ cells was fully responsive to carbon source [Fig. 3C,D]. Introduction of sfp1Δ into the cln3Δ bck2Δ whi5Δ background rendered cells very small, but not as small as the sfp1Δ mutation alone [Fig. 3C,D]. Because sfp1Δ is not fully epistatic for cell size to the triple cln3Δ bck2Δ whi5Δ mutant or to any of the single mutants [data not shown], Whi5, Cln3, and Bck2 must still play a role in sfp1Δ cells. These results argue that nutrient signaling, via Sfp1 and Sch9, influences the activation of SBF and MBF by a mechanism independent of known upstream regulators.

Sch9 abundance and phosphorylation are altered by nutrient signals

As the RP and Ribi regulons respond dynamically to the environment, we examined Sfp1 and Sch9 abundance and localization under different nutrient and stress conditions using functional fusions to epitope tags and GFP variants. In log phase, GFP-Sch9 localized throughout the cell, but was unexpectedly enriched at the vacuolar membrane [Fig. 4A]. We note that a C-terminal Sch9-GFP fusion protein was not functional [data not shown]. GFP-Sch9 localization was dynamic, as the vacuolar membrane signal largely disappeared following carbon starvation [Fig. 4B,C]. GFP-Sch9 localization was not altered by different carbon sources or by rapamycin [data not shown]. The abundance of HA3-Sch9 was, however, dependent on TOR activity as all forms, and in particular HA3-Sch9 phosphoisoforms, decreased upon rapamycin treatment [Fig. 4D,E]. Under steady-state proliferation on different carbon sources, HA3-Sch9 abundance correlated with cell growth rate, RP/Ribi transcription, and cell size [Fig. 4D] [Gasch et al. 2000].

Sfp1 nuclear concentration is regulated by multiple nutrient and stress signals

As expected for a transcription factor, Sfp1-GFP was localized predominantly to the nucleus in growing cells [Fig. 5A]. Strikingly, Sfp1-GFP rapidly (~5 min) relocalized to the cytoplasm upon exposure to a broad range of stresses. Cytoplasmic relocalization was also observed with a GFP-Sfp1 fusion protein [data not shown]. Quantitation of the Sfp1-GFP nuclear:cytoplasmic [N:C] ratio revealed that under some conditions (carbon starvation, oxidative stress), Sfp1-GFP became uniformly distributed [i.e., an N:C ratio of unity], whereas in others [rapamycin, tunicamycin, nitrogen starvation], a greater fraction of Sfp1-GFP was retained in the nucleus [Fig. 5A,B]. Neither the abundance nor the electrophoretic mobility of Sfp1MYC13 was altered upon carbon starvation or rapa-

Figure 3. Cells lacking SFP1 or SCH9 fail to adjust cell size in response to carbon source. [A] Representative size distributions of log-phase cultures of the indicated strains in synthetic glucose [black], raffinose [blue], or glycerol [orange] medium. [B] Mean cell sizes of the indicated cultures (n = 6, range: 10–180 FL). Error bars extend one S.E. in each direction. The sfp1Δ/Δ GAL1–CLN3-1 [3-1] strain was propagated in synthetic galactose medium. [C] Nutrient and SFP1-dependent control of size in cells that lack known upstream regulators of SBF/MBF. Wild-type and cln3Δ bck2Δ whi5Δ strains were in rich glucose, raffinose, or glycerol medium. The sfp1Δ and sfp1Δ cln3Δ bck2Δ whi5Δ strains were in rich glucose medium. [D] Mean cell sizes of the indicated cultures (n ≥ 2, range: 10–180 FL). Error bars extend one S.E. in each direction.
cytochalasin D), and the partial nuclear retention of Spfp1 in the presence of amino acids suggested that Ste12, a kinase that is required for the TOR pathway, was involved in the control of Sfp1 localization. The TOR pathway plays a role in cellular growth and division by controlling the expression of genes involved in ribosome biogenesis, amino acid transport, and protein synthesis. However, the nuclear retention of Spfp1 after rapamycin treatment of wild-type cells suggests that repression of the TOR pathway does not suffice to explain the complete relocalization upon carbon starvation.

Importantly, Spfp1 localization was also linked to the TOR pathway. The rapamycin-resistant TOR1-1 allele blocked Spfp1 nuclear depletion in response to rapamycin (Fig. 5H). However, the partial nuclear retention of Spfp1 may suggest that rapamycin treatment of wild-type cells reduces TOR activity, which in turn affects Sfp1 localization.

Figure 4. Sch9 abundance and localization is modulated by nutrients. (A) Enrichment of GFP/Sch9 at the vacuolar membrane (vm). (B) Depletion of GFP/Sch9 from the vm after carbon starvation (37 min). (C) Kinetics of GFP/Sch9 depletion from the vm. (Closed squares) Glucose; (open diamonds) glucose. Error bars extend one S.E. in each direction. (D) HA-Sch9 abundance and phosphorylation status are altered by nutrient conditions. Cells that expressed HA-SCH9 from the SCH9 promoter were propagated in rich glucose, raffinose, or glycerol medium. Cells propagated in glucose medium were treated with rapamycin [rap, 200 ng/mL] or carbon starved. (E) HA-Sch9 was visualized by immuno blot with anti-HA antibody. GFP/Sch9 served as a no-tag control. Asterisk indicates nonspecific cross-reactive species. (E) Sch9 is a phosphoprotein. HA-Sch9 expressed at endogenous levels was immunoprecipitated with anti-HA antibody and either mock treated [M], treated with alkaline phosphatase [P], or treated with alkaline phosphatase in the presence of phosphatase inhibitors [P+I] and detected by anti-HA immunoblotting.

Cytochalasin D is a drug that disrupts actin filament assembly, and treatment with this compound led to a gradual decline in both nuclear Spfp1YFP and RP/Ribi transcription (Fig. 5B) [Miyoshi et al. 2003]. In log phase proliferation on different carbon sources, the extent of Spfp1YFP nuclear localization was also proportional to cell-growth rate, RP/Ribi expression, and cell size [Fig. 5E] [Gasch et al. 2000].

Importantly, Spfp1 localization responded quickly to improved nutrient conditions. Addition of glucose to stationary phase cells rapidly (∼5 min) increased the nuclear concentration of Spfp1YFP [Fig. 5I]. This nuclear import of Spfp1YFP occurred even in the presence of a lethal concentration of cycloheximide, and so did not require protein synthesis [Fig. 5F]. In fact, cycloheximide treatment of cells in suboptimal raffi nase medium rapidly increased the nuclear concentration of Spfp1YFP, but had no effect on Ste12YFP, an unrelated transcription factor [Fig. 5C; data not shown]. Spfp1 may thus effect the compensatory feedback loop that induces RP transcription within 15 min of cycloheximide treatment [Cárdenas et al. 1999].

Spfp1 localization also depended on TOR and Ras/PKA activity. The rapamycin-resistant TOR1-1 allele blocked Spfp1 nuclear depletion in response to rapamycin [Fig. 5H]. However, the partial nuclear retention of Spfp1 after rapamycin treatment of wild-type cells suggests that repression of the TOR pathway does not suffice to explain the complete relocalization upon carbon starvation [Fig. 5B]. A tpk1wimp strain, which has weak constitutive PKA activity [Cameron et al. 1988], had reduced nuclear Spfp1YFP [Fig. 5I], whereas overexpression of a hyperactive RAS2Val19 allele stimulated nuclear accumulation of Spfp1YFP [Fig. 5I] and RP/Ribi induction with the same kinetics [Y. Wang et al. 2004]. However, because a tpk1wimp strain still relocalized Spfp1 upon carbon starvation, diminished Ras/PKA signaling was also not sufficient to account for the totality of Spfp1 export [Fig. 5I]. Likewise, a high level of Ras/PKA activity caused by either of the hyperactive GAL10–RAS2Val19 or GPA2Aub73 alleles could not drive Spfp1 into the nucleus of carbon-starved cells [data not shown]. Deletion of Snf1, an AMP-activated kinase that signals many stress responses, also had no effect on Spfp1 relocalization upon carbon starvation [data not shown]. Multiple signals must therefore regulate nuclear localization of Spfp1. We note that Spfp1 cannot completely account for the pronounced effect of TOR and Ras/PKA signaling on RP transcription, as loss of Spfp1 activity causes an approximately threefold repression of RP genes as opposed to the >10-fold decreases caused by loss of TOR or PKA activity [Fig. 1C] [Neuman-Silberberg et al. 1995; Powers and Walter 1999].

Spfp1 and Sch9 act in parallel

Consistent with the above Spfp1 localization results, spfp1Δ strains are sensitive to cycloheximide [Fingerman et al. 2003] and to decreases in Ras/PKA and TOR signaling [Fig. 6A], arguing that these pathways regulate additional activators of RP and Ribi expression. The incomplete overlap in expression profiles [Fig. 1E], lack of
Figure 5. Sfp1 nuclear concentration responds rapidly to environmental stimuli and is regulated by the Ras/PKA and TOR signaling pathways. [(A) Sfp1 localization is regulated by nutrient signals. An SFP1^{YFP} SEC63^{CFP} strain was visualized, and cell and nuclear membranes were demarcated by Sec63^{CFP}. Carbon starvation (–C) was for 15 min, and rapamycin treatment [rap, 200 ng/mL] for 20 min. [(B) Sfp1 exits the nucleus in response to various stress conditions. An SFP1^{YFP} SEC63^{CFP} strain was depleted for carbon or nitrogen or treated with H_{2}O_{2} (0.30 mM), rapamycin (200 ng/mL), or tunicamycin [2 mg/mL]. The Sfp1^{YFP} nuclear:cytoplasmic ratio [N:C] of each cell at each time point was plotted (dashes, 30–100 cells), as was the average ratio (thick line). (C) Sfp1 abundance and electrophoretic mobility are not altered by carbon starvation (–C) or rapamycin [rap, 200 ng/mL] treatment. Cells were harvested after the indicated time [min]. Sfp1^{MYC13} in cell lysates was visualized by immunoblot with anti-MYC antibody. Fh1^{MYC13} served as a specificity control. (D) Sfp1 relocation correlates with repression of the RP and Ribi regulons. Genome-wide expression profiles comparing mRNA abundance before and after carbon starvation (–C) were obtained by de novo expression profiles for rapamycin addition [rap], oxidative stress [H_{2}O_{2}], and nitrogen starvation (–N) were derived from published data [Hardwick et al. 1999; Gasch et al. 2000]. Scale indicates fold change. (E) Nuclear localization of Sfp1 in different carbon sources. The average Sfp1^{YFP} N:C ratio was determined under steady-state proliferation in glucose, raffinose, and glycerol medium. Error bars extend one S.D. in each direction. (F) Sfp1 re-enters the nucleus rapidly in response to glucose. Stationary phase SFP1^{YFP} SEC63^{CFP} cells were re-fed with glucose in the presence or absence of cycloheximide [chx, 10 µM] and Sfp1^{YFP} N:C ratios determined. (G) Sfp1 may effect a feedback response to ribosome shortage. Sfp1^{YFP} N:C ratio was measured in SFP1^{YFP} SEC63^{CFP} cells proliferating in raffinose medium before and after addition of chx [10 µM]. (H) The rapamycin-resistant allele TOR1-I blocks Sfp1 relocation in response to rapamycin but not nitrogen starvation. A TOR1-1 SFP1^{YFP} strain was treated with rapamycin [red dotted line, 200 ng/mL] or starved for nitrogen [black, offset for visualization] for 30 min. (I) Compromised Ras activity lowers the nuclear concentration of Sfp1. The N:C ratio of Sfp1^{YFP} was quantitated before and after 30 min of carbon starvation in a tpk1^{wimp} strain [red dotted line, SFP1^{CFP} tpk1^{wimp} bcy1Δ tpk2Δ tpk3Δ] and in a control strain [black line, SFP1^{CFP}, offset for visualization]. At t = 0 min, the difference between the wild-type and wimp strain was significant (Student’s t-test, p = 4.4 × 10^{-11}). Due to higher cell autofluorescence, Sfp1^{CFP} N:C ratios are less than Sfp1^{YFP} N:C ratios. (J) Hyperactive Ras signaling drives Sfp1 into the nucleus. A GAL10–RAS2^{VP} SFP1^{CFP} strain [red dotted line] and a control SFP1^{CFP} strain [black, offset for visualization], proliferating in synthetic raffinose medium, were induced with galactose at t = 0, and visualized after 30 and 60 min. Both strains were deleted for GAL1 and are incapable of metabolizing galactose.

genetic epistasis [Fig. 6B], and the invariability of the sfp1Δ sch9Δ double mutant indicate that Sfp1 and Sch9 act in parallel pathways [Jorgensen et al. 2002]. Consistently, inhibition of sch9^{+} does not affect the localization of Sfp1^{CFP} (data not shown). However, because both sfp1Δ and sch9Δ strains have substantial growth defects, the invariability of the double mutant might simply reflect nonspecific additive growth defects. To precisely modulate Sfp1 function, we constructed an sfp1^{KR} allele in which the genomic SFP1 sequence is fused to the ligand-binding domain of the estrogen receptor [ER]. In the absence of the natural ligand β-estradiol, the ER domain is sequestered in an inactive complex with Hsp90 [Mattoni et al. 1994]. The activity of Sfp1^{KR} in vivo exhibits a smooth dose dependence on β-estradiol concentration [Fig. 6C]. To quantitatively assess genetic synergy, an sfp1^{KR} sch9^{+} double-mutant strain was challenged with increasing concentrations of β-estradiol and 1NM-PP1, in effect applying overlapping gradients of Sfp1 and Sch9 activity. Severe proliferation defects were observed at combined concentrations of β-estradiol and 1NM-PP1 that individually caused little defect, consistent with a parallel role for each in RP/Ribi transcription [Fig. 6D]. As Sfp1 and Sch9 both appear to be downstream of TOR signaling [Figs. 4D, 5A,H], their combined action may account for the TOR requirement in RP/Ribi gene expression.

Sfp1 appears to act at RP promoters via Fhl1 and Ifh1.

Genome-wide chromatin immunoprecipitation [ChiP] analysis of many potential transcriptional regulators in yeast, including Sfp1, has been recently reported [Lee et al. 2002]. Of the 211 intergenic regions most enriched in
Sfp1 complexes at p < 0.02, 42 corresponded to RP genes, a proportion not due to chance (p < 10^{-14}). But, the less than twofold promoter enrichment in the reported ChIP data for Sfp1 [Lee et al. 2002], and our own ChIP analysis [data not shown], suggests that the Sfp1–RP promoter interaction is at the limit of detection. Sfp1–Rbi1 promoter interactions have not yet been detected by ChIP [Lee et al. 2002; Fingerman et al. 2003; P. Jorgensen and M. Tyers, unpubl.]. To determine how Sfp1 and Sch9 might regulate RP promoters, we examined two novel transcription factors that also bind to these promoters, the forhead/FHA domain protein Fhl1 and the zinc-finger protein Rgm1 [Lee et al. 2002]. Fhl1 and a genetically interacting factor, Ihf1, are required for an unknown aspect of ribosome synthesis, suggesting that they may be activators of RP transcription [Hermann-Le Denmat et al. 1994; Cherel and Thuriaux 1995].

We detected several strong genetic interactions within this network of putative RP gene regulators. In light of the severe proliferation defects of sfp1Δ and fhl1Δ strains, we were surprised to find that a sfp1Δ fhl1Δ double-mutant strain proliferated as well as a fhl1Δ strain; in contrast, sfp1Δ caused additive defects with fhl1Δ [Fig. 7A]. However, deletion of FHL1 did not cause an obvious cell size phenotype [Fig. 7B]. The sfp1Δ size defect was epistatic to fhl1Δ as the sfp1Δ fhl1Δ double mutant was as small as the sfp1Δ single mutant [Fig. 7B]. The control of Sfp1 over cell size may thus be distinct from its control over the RP regulon. Conversely, the colony size of an sch9Δ, but not an sfp1Δ strain was severely compromised by an allele of IFH1 (ifh1^{100}) and by rgm1Δ, neither of which alone caused noticeable defects [Fig. 7C; data not shown].

These genetic interactions suggested that Sfp1 might influence RP transcription via Fhl1 and Ihf1. To investigate Fhl1 and Ihf1 regulation, we created various C-terminal epitope tag or GFP variant fusions by integration of fusion cassettes at the genomic FHL1 and IFH1 loci, all of which provided full function in vivo. We first recapitulated the association of IFH1Δ with RP promoters by ChIP analysis, as controls, Fhl1 did not locate to another highly active promoter regulated by Rap1 (PGK1) or to Rbi1 promoters [Fig. 7D]. We then determined that Ihf1^{MYC13} also bound specifically to RP promoters [Fig. 7D]. The association of Ihf1^{MYC13} and Fhl1Δ with RP promoters was reduced in sfp1Δ cells by approximately fourfold and approximately twofold, respectively [Fig. 7E]. Neither the sfp1Δ mutation nor carbon starvation detectably altered Ihf1^{MYC13} or Fhl1^{100}, abundance or electrophoretic mobility (data not shown).

Sfp1 dramatically influenced the localization of Ihf1 and Fhl1. In wild-type cells in glucose medium, Ihf1^{CFP} and Fhl1^{YPFP} were predominantly nuclear, but excluded from the nucleolus [Fig. 8A,B]. In sfp1Δ strains, however, a high percentage of cells showed striking enrichment of Ihf1^{CFP} and Fhl1^{YPFP} in a subnuclear focus that corresponded to the nucleolus, as shown by colocalization with Bud21^{YPFP} or Bud21^{CFP} [Fig. 8A,B]. These foci were not observed in strains lacking Sch9 activity or after treatment with a sublethal concentration of cycloheximide [Fig. 8C; data not shown], indicating that reduced growth or translation rate was not sufficient to induce nucleolar relocalization. We then examined whether Ihf1^{CFP} and Fhl1^{YPFP} changed localization in response to a nutrient stress that causes Sfp1 to exit the nucleus. Upon carbon starvation, Ihf1^{CFP} rapidly infiltrated the nucleolus of most cells [Fig. 8D,E]. Likewise, in most cells,
almost all Fhl1-YFP relocalized near the nucleolus after carbon-source depletion [Fig. 8F,G]. As controls, two unrelated transcription factors, Ste12-YFP and Rst1-YFP, did not form foci in spf1Δ cells or upon carbon starvation [data not shown]. Fhl1-YFP and Ifh1-CFP nucleolar localization thus inversely correlated with the nuclear concentration of Spf1. Surprisingly, in spite of the transcriptional repression of RP genes and Ifh1 and Fhl1 relocalization to the nucleolus, Ifh1 and Fhl1 were readily detected at RP promoters under conditions of carbon source limitation [Fig. 8H]. The activity of Fhl1/Ifh1 may therefore be dictated by the nuclear environment of promoter regions.

Discussion

An unexpected but fundamental connection has emerged between two ostensibly disparate systems, ribosome biogenesis and Start. Phenotypic identification of potential Start repressors isolated not only 15 ribosome biogenesis factors but, more significantly, upstream activators of the mRNA transcriptional program that sets the rate of ribosome synthesis, namely Sfp1 and Sch9 [Jorgensen et al. 2002]. We have elaborated these initial connections to uncover a dynamic transcription factor network that responds to nutrient signals in part through altered localization of critical regulators. Other uncharacterized potential Start repressors may well participate in this emerging network that controls RP and Ribi expression [Jorgensen et al. 2002].

A dynamic transcriptional network at RP promoters

Sfp1 is a primary element in RP/Ribi regulation. Given that the Ribi regulon responds more rapidly than the RP regulon in response to Sfp1 [Jorgensen et al. 2002], it is puzzling that Sfp1 binds weakly to RP promoters, but apparently not to Ribi promoters [Lee et al. 2002; Fingerman et al. 2003]. Nevertheless, Sfp1 activates Ribi transcription via the RRPE element [Fingerman et al. 2003]. Consistent with a primary regulatory role, Sfp1 nuclear concentration responds within minutes to environmental conditions and appears to dictate the nuclear localization of Fhl1 and Ifh1, two novel regulators of RP promoters [Fig. 9A]. Additional activators of the RP and Ribi regulons include the Sch9, TOR, and Ras/PKA pathways, all of which respond to nutrient signals. Sfp1 is downstream of TOR and Ras/PKA kinases, but the observed genetic redundancies indicate that these signaling pathways must modify multiple elements of the RP promoter network. Sfp1, Fhl1, and Rgm1 are likely to bind directly to RP promoter elements, such as the T-rich sequence or more recently described cis elements [Warner 1999; Pilpel et al. 2001; Beer and Tavazoie 2004], thereby placing the RP regulon under complex combinatorial control [Fig. 9A]. As Rap1-binding sites mediate nearly all transcriptional activation at RP promoters [Warner 1999], it seems likely that most other components will subserve Rap1, perhaps by switching Rap1 between activation and repression modes. In support of this model, overexpression of the N terminus of Ifh1 disrupts telomeric and mating-type silencing, both of which require Rap1 [Singer et al. 1998].

The dynamic nature of the RP/Ribi control network is manifest at several levels. First, the ability of the two regulons to respond quickly to environmental conditions rests on adept transcriptional responses coupled with mRNA instability [Warner 1999; Grigull et al. 2004]. Second, Sfp1 relocalization is remarkably responsive to the environment. As it is never completely excluded from the nucleus, the RP and Ribi promoters may be quite sensitive to the nuclear concentration of Sfp1, as might beft weak interactions of Sfp1 with RP promoters. Third, Fhl1 and Ifh1 are unexpectedly subject to

Figure 7. Sfp1 influences Fhl1 and Ifh1 interactions with RP promoters. [A] FHL1 deletion is epistatic to SFP1 deletion for colony size. Spore clones were imaged after 7 d [top] and 5 d [bottom]. [B] FHL1 deletion is not epistatic to SFP1 deletion for cell size. Strains recently derived from a tetradtype tetrad dissection (as in A) were sized in log phase in rich glucose medium. These representative distributions were highly reproducible. (C) Synthetic proliferation defects between sch9Δ and rgm1Δ and an allele of IFH1. Spore clones were imaged after 3 and 2 d, respectively. [D] Fhl1 and Ifh1 bind specifically to RP promoters. Real-time PCR was used to quantitate the efficiency with which the promoter regions of the indicated genes were captured in Fhl1Δ [F7 anti-HA antibody] or Ifh1Δ [MYC13, 9E10 anti-MYC antibody] complexes. Binding to Ribi promoters [URA7, RPA190, NSR1] and control promoters [PGK1, ACT1] was not observed. (E) Ifh1Δ [MYC13] and Fhl1Δ [HA1] bind poorly to RP promoters in spf1Δ cells. The spf1Δ/wild-type ratio of ChIP efficiency for ifh1Δ [MYC13] and Fhl1Δ [HA1] was calculated for two individual experiments. Error bars extend one S.E. in each direction.
nucleolar relocalization. Ifh1 and/or Fhl1 may potentially have dedicated nucleolar roles, such as repression of RNA Polymerase I and III, and thereby constitute an unsuspected line of communication between RP and rRNA transcription [Fig. 9B]. In an alternate, but not mutually exclusive model, the nucleolar focus of Ifh1 and Fhl1 in sfp1Δ and carbon-starved cells may signify a cluster of repressed RP genes, as Ifh1 and Fhl1 continue to bind to RP promoters in these cells [Fig. 9B]. Intriguingly, physical interactions have been found between Fhl1 and the rDNA-binding factor Hmo1 and between Ifh1 and the nucleolar protein Utp22 [data not shown], suggesting that Fhl1/Ifh1 may tether inactive RP genes to nucleolar structures. Spatial control at the level of gene compartmentalization is evident in nucleolar tRNA gene clusters in rapidly growing cells and in Rap1-dependent clustering of silenced loci at the nuclear periphery [Gasser 2001; Thompson et al. 2003]. Furthermore, the relocalization of transcription factors appears to be a common feature in ribosome biogenesis as the limiting regulatory factor for RNA transcription, Rrn3, as well as RNA PolI subunits, are released from the nucleolus upon rapamycin treatment [Tsang et al. 2003; Mayer et al. 2004].

Sch9 relays nutrient signals

An unexpected level of spatio-dynamic control has emerged with the localization of Sch9 to the vacuole. Despite its discovery as a high-copy suppressor of the Ras/PKA pathway in yeast [Toda et al. 1998], and its recently discovered roles in cell longevity and size [Fabrizio et al. 2001; Jorgensen et al. 2002], the presumed components that lie upstream and downstream of Sch9 have proven elusive. Budding yeast expresses two bona fide orthologs of the PDK1 kinase that activates Akt/PKB in metazoans, called Pkh1 and Pkh2, although these kinases are activated by sphingolipids, not phosphatidylinositol trisphosphate [Casamayor et al. 1999]. Pkh1 and Pkh2 activate Ypk1 and Ykr2, the yeast orthologs of the metazoan kinase SGK [Casamayor et al. 1999]. As its sequence contains the predicted activating phosphorylation sites in the activation loop and hydrophobic motif, Sch9 may also be a Pkh1/2 substrate. As the vacuole is an important reservoir of amino acids, phosphates, and other metabolites, Sch9 may communicate the status of these internal nutrient pools to RP/Ribi transcription and Start. Elaboration of the Sch9 pathway in yeast may
provide insights into the regulation of cell size by Akt/ PKB in metazoans.

Ribosome biogenesis and nutrient modulation of critical cell size

On the basis of the above observations, we propose a refined model for nutrient modulation of the critical cell-size threshold at Start. Sfp1, Sch9, and Ras/PKA function in a nonlinear network that dictates both critical cell size and expression of the Ribi and RP regulons. Critical cell size at Start is decreased when any of these components is crippled, whereas either constitutive activation [Ras/PKA] or inactivation [Sfp1, Sch9] renders cell size impervious to carbon source control [Baroni et al. 1989; Tokiwa et al. 1994]. Each component of this trio is needed for proper RP and Ribi gene expression [Klein and Struhl 1994; Neuman-Silberberg et al. 1995; Y. Wang et al. 2004]. Like Ras/PKA signaling, we have found that Sfp1 and Sch9 are sensitive to nutrient conditions, at the level of localization and abundance, respectively. In addition, strains deleted for numerous genes implicated in the actual events of ribosome biogenesis, as well as ribosome structural genes, are similarly, if less dramatically, uncoupled for growth and division [Jorgensen et al. 2002]. Finally, the cell size at which SBF/MBF activation, budding, and DNA replication initiation occur is diminished in cells proliferating in poor nutrients [Johnston et al. 1979; Lorincz and Carter 1979; Tyson et al. 1979; Stuart and Wittenberg 1995; Flick et al. 1998]. All of these observations can be unified by a model in which nutrient control of the critical cell-size threshold at Start is communicated by rates of ribosome production [Fig. 9C].

In this model, nutrient status influences the Start machinery via proximal events in ribosome biogenesis, rather than by downstream changes in protein synthetic rate. This configuration would effectively anticipate future changes in translation rate and adjust the cell-size threshold accordingly. In support of this idea, upon shift of cells from ethanol to glucose medium, rates of rRNA transcription reach maximal values within a few minutes, whereas protein synthetic rate does not achieve a maximum until nearly an hour after the shift [Kief and Warner 1981]. Concordantly, upon nutrient shifts, cells adjust their critical cell-size threshold very rapidly [Lorincz and Carter 1979].

This model begs the question of how ribosome biogenesis might impinge on the Start machinery. As ribosome biogenesis factors implicated in size control lie along the entire pre-60S assembly pathway, the cell may monitor flux through this branch [Fatica and Tollervey 2002]. Notably, inhibition of ribosome biogenesis by secretory defects also appears to proceed via the 60S branch [Zhao et al. 2003]. Dissection of the ribosome biogenesis signal is complicated by redundancy among the many components, the existence of feedback loops, and its essential role in the cell. As G1/S transcription is greatly accelerated in cells lacking SFP1 or SCH9, it is possible that SBF/MBF are a target of the presumed signal. Other less-direct models in which Ras/PKA, Sch9, and Sfp1 impinge

Figure 9. (A) Initial model for the transcription system at RP promoters. Sfp1 localization is controlled by various signals and in turn influences the localization of Ifh1 and Fhl1. As suggested by synthetic lethal interactions, Sch9 modulates RP transcription by a pathway parallel to Sfp1/Fhl1/Ifh1/Rgm1, perhaps by phosphorylating Rap1. Sfp1, Fhl1, and Rgm1 likely bind to the promoter directly via still uncharacterized sequence elements (gray box) [Warner 1999; Pilpel et al. 2001; Beer and Tavazoie 2004]. Ifh1 and Fhl1 may activate RP transcription by switching Rap1 [or Abf1] between intrinsic transcriptional activation [A] and repressive [R] functions. (B) Summary of transcription factor localization correlated with active and inactive (carbon starvation) RP transcription. Two possible nonmutually exclusive models explain the nucleolar relocalization of Ifh1 and Fhl1 upon carbon starvation. (Left) In the first model, unbound Ifh1 and Fhl1 are sequestered in or near the nucleolus (nucl), where they may have additional functions, such as repressing rRNA transcription. In the second model, Ifh1 and Fhl1 and repressed RP promoters are drawn into the nucleolar region upon carbon starvation. (C) A model of Start entry. Ribosome synthesis rates and ploidy establish the critical cell size threshold, which represses the SBF and MFB transcription factor complexes by an unknown mechanism. Cell size or a parameter that correlates with size, such as translation rate, signals to Cln3 and/or Bck2 to activate SBF and MFB. The model is complicated by the effects of ribosome synthesis on translation and by increased Cln3 abundance in rich nutrient conditions [dashed line].
on a second common target might also be considered. For instance, decreasing the overall rate of \( R_P \) and \( R_b i \) transcription in nutrient-limited cells might free up RNA Polymerase II and other core transcriptional regulators for recruitment by SBF/MBF, and presumably all other promoters [Thomas 2000].

Two mechanisms to explain nutrient modulation of critical cell size have been proposed previously, both of which invoke changes in the abundance of G1 cyclins [Baroni et al. 1994; Tokiwa et al. 1994; Polymenis and Schmidt 1997; Flick et al. 1998]. However, such models do not account for all of the data presented here and elsewhere. Carbon source control of Start occurs in part via transcriptional regulation of \( C L N \), because \( c l n^\Delta \) strains fail to appropriately increase critical cell size when shifted from poor to glucose medium, moreover, glucose and cAMP repress \( C L N \) even relative to other G1/S transcripts like \( C L N \) [Tokiwa et al. 1994; Flick et al. 1998]. However, in these studies, critical cell size at Start was inferred indirectly by bud emergence, rather than by measurement of SBF/MBF-dependent transcription. Because SBF/MBF activation is highly dependent on \( C L N \), but independent of \( C L N \) and \( C L N \) [Tycers et al. 1993; Dirick et al. 1995; Stuart and Wittenberg 1995], \( C l n^\Delta \) likely controls the length of the Start interval rather than the timing of Start entry.

A second model postulates that \( C l n^\Delta \) couples the critical cell size threshold to nutrients by virtue of \( C l n^\Delta \) translational control and protein instability [Polymenis and Schmidt 1997; Hall et al. 1998]. It is clear that carbon source controls critical cell size at the level of SBF/MBF activation, an effect that in principle might be transmitted through \( C l n^\Delta \) [Stuart and Wittenberg 1995]. As shown here, however, carbon source and \( S F P \) strongly impact cell size even in cells that lack the known upstream regulators of SBF/MBF, thereby obviating the \( C l n^\Delta \)-based model. Furthermore, the observation that in poor nutrients cells have very low levels of \( C l n^\Delta \) and translational capacity yet pass Start at a small size, is also at odds with this model [Tokiwa 1995; Hall et al. 1998]. That is, nutrient upshifts delay Start, despite increases in \( C l n^\Delta \) abundance [Johnston et al. 1977; Lorincz and Carter 1979; Tokiwa et al. 1994]. From this perspective, nutrient modulation of the critical cell-size threshold is quite remarkable, as not only must the yeast cell growing in poor nutrients pass Start with less translational capacity, but it must do so with much less \( C l n^\Delta \). That is, in poor nutrients, less \( C l n^\Delta \)-Cdc28 activity seems to be required to pass Start. Consistently, Start-defective \( c d c 28^\Delta \) mutants at a semipermissive temperature cycle threefold faster in pyruvate medium than in glucose medium [Shuster 1982].

To synthesize, a sharp distinction must be drawn between the critical cell-size threshold, which is altered by nutrients and ploidy, and the mechanism by which cells gage their size, perhaps through translation rate. A sizing role for translation rate is suggested by the increased critical size caused by sublethal doses of cycloheximide. Furthermore, even cells beyond the critical cell-size threshold require a minimum rate of protein synthesis before budding can occur [Moore 1988]. Given these observations, the critical cell-size threshold and the critical translation-rate requirement may be one and the same. As argued above, the mechanisms that set this threshold may be entirely distinct from the processes that measure size.

**Ribosome biogenesis and cell cycle control**

Several recent findings suggest that primordial links between ribosome biogenesis and the cell cycle may couple growth and division in metazoan systems [Saucedo and Edgar 2002; Ruggero and Pandolfi 2003]. As in yeast, disruption of ribosome biogenesis, but not the translational machinery, in flies causes a small cell-size phenotype [Montagne et al. 1999; Thomas 2000]. In mice, blocking the synthesis of new ribosomes allows hepatocytes to grow, but not enter the cell cycle [Volarevic et al. 2000]. Similarly, in human cells, overexpression of a dominant-negative version of a conserved ribosome biogenesis factor called Bop1 causes G1-phase arrest in a p53-dependent manner [Pestov et al. 2001]. Many additional connections between p53 and the nucleolus have been unearthed [Ruggero and Pandolfi 2003]. For example, the critical activator of p53, the Arf tumor suppressor, is a nucleolar protein and an inhibitor of ribosomal RNA processing [Sugimoto et al. 2003]. It has recently been proposed that all stresses that stabilize p53 do so by disrupting the nucleolus [Rubbi and Milner 2003].

As in yeast, mechanisms exist in mammalian cells to ensure that ribosomal content can be doubled with each cell cycle. Delineation of the signals that emanate from ribosome biogenesis to the cell cycle machinery in yeast should thus illuminate analogous processes in metazoans as well as the evolutionary history of size control processes. Like Sfp1, Myc is a direct modulator of RP transcription and cell size [Eisenman 2001], whereas Sch9 and Akt are highly related kinases that regulate ribosomal synthesis and cell size [Saucedo and Edgar 2002]. Understanding these connections has been lent new urgency by the discovery of numerous links between ribosome biogenesis and cancer [Ruggero and Pandolfi 2003], including the recent demonstration that RP genes are haploinsufficient tumor suppressors in zebrafish [Amsterdam et al. 2004].

**Materials and methods**

**Yeast strains, plasmids, and medium**

Extensive descriptions of all methods are provided as Supplemental Material. Yeast culture and size analysis was as described previously [Jorgensen et al. 2002]. All experiments were performed with log phase cells at O.D.\(_{600}\) < 0.5 and cell concentration <5 x 10\(^5\) cells/mL to minimize repression of the \( R_b i \) and \( R_P \) regulons by inadvertent nutrient depletion [Ju and Warner 1994; DeRisi et al. 1997]. Strains used in this study are listed in Supplementary Table 2 and were typically generated by genomic integration of standard C-terminal tagging cassettes, including GFP variant cassettes obtained from the Yeast Resource Center [University of Washington, Seattle]. Centrifugal elutria-
tion, Northern analysis, genome-wide expression profiles, chromatin immunoprecipitation, and as allele construction were carried out as described [Tyers et al. 1993; Bishop et al. 2000; Jorgensen et al. 2002; Costanzo et al. 2004].

Quantitative microscopy

Live-cell microscopy was performed with an Eclipse E600FN microscope [Nikon] and an Orca II CCD camera [Hamamatsu]. Cultures in synthetic medium were rapidly concentrated and immediately visualized to mitigate starvation effects. For quantification of Spf1 YFP or Spf1-3FP nuclear:cytoplasmic ratio, five serial sections were taken of each cell field. Sec63-CFP or Sec63-3FP was used to delineate the nuclear envelope. Metamorph [Universal Imaging] was used to capture and quantify the fluorescence signal from an identically sized region from the nucleus and the cytoplasm. To induce carbon starvation, cells were washed once and resuspended in synthetic medium with 0.02% glucose. For nitrogen starvation, cells were resuspended in synthetic glucose medium with 1/50th the standard concentration of amino acids and ammonium sulfate. Ifh1-1CFP and Fhl1-1CFP signals were visualized in the same manner, except that nucleolar regions were demarcated with Bud21-3CFP or Bud21-3FP. Because allowing cells to remain concentrated for even short periods of time caused redistribution of Spf1, Fhl1, and Ifh1, all results were confirmed with directly mounted dilute cultures.

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