Saccharomyces cerevisiae TORC1 Controls Histone Acetylation by Signaling Through the Sit4/PP6 Phosphatase to Regulate Sirtuin Deacetylase Nuclear Accumulation

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ABSTRACT The epigenome responds to changes in the extracellular environment, yet how this information is transmitted to the epigenetic regulatory machinery is unclear. Using a Saccharomyces cerevisiae yeast model, we demonstrate that target of rapamycin complex 1 (TORC1) signaling, which is activated by nitrogen metabolism and amino acid availability, promotes site-specific acetylation of histone H3 and H4 N-terminal tails by opposing the activity of the sirtuin deacetylases Hst3 and Hst4. TORC1 does so through suppression of the Tap42-regulated Sit4 (PP6) phosphatase complex, as sit4Δ rescues histone acetylation under TORC1-repressive conditions. We further demonstrate that TORC1 inhibition, and subsequent PP6 activation, causes a selective, rapid, nuclear accumulation of Hst4, which correlates with decreased histone acetylation. This increased Hst4 nuclear localization precedes an elevation in Hst4 protein expression, which is attributed to reduced protein turnover, suggesting that nutrient signaling through TORC1 may limit Hst4 nuclear accumulation to facilitate Hst4 degradation and maintain histone acetylation. This pathway is functionally relevant to TORC1 signaling since the stress sensitivity of a nonessential TORC1 mutant (tco89Δ) to hydroxyurea and arsenic can be reversed by combining tco89Δ with either hst3Δ, hst4Δ, or sit4Δ. Surprisingly, while hst3Δ or hst4Δ rescues the sensitivity tco89Δ has to low concentrations of the TORC1 inhibitor rapamycin, sit4Δ fails to do so. These results suggest Sit4 provides an additional function necessary for TORC1-dependent cell growth and proliferation. Collectively, this study defines a novel mechanism by which TORC1 suppresses a PP6-regulated sirtuin deacetylase pathway to couple nutrient signaling to epigenetic regulation.

KEYWORDS target of rapamycin; histone acetylation; Sit4; sirtuins; epigenetic

Dynamic chromatin regulation is key to the adaptive mechanisms eukaryotic cells employ to alter cellular phenotype in response to fluctuating environmental conditions, such as changes in nutrient availability and various forms of stress. Environmental regulation of the epigenome profoundly affects health and disease, but how the chromatin regulatory apparatus responds to such influences is not understood (Lu and Thompson 2012; Szyf 2015). A universally conserved signaling pathway regulated by the environment is the target of rapamycin (TOR) pathway (Loewith and Hall 2011; Laplante and Sabatini 2012). TOR consists of two distinct subpathways involving TOR complex 1 (TORC1) and TOR complex 2 (TORC2), of which only TORC1 is activated by environmental inputs (Loewith and Hall 2011; Laplante and Sabatini 2012). In single-celled organisms such as budding yeast, TORC1 is specifically stimulated by the quantity and quality of the available nitrogen source; however, while controversial, recent studies also have suggested potential crosstalk between glucose signaling, carbon metabolism, and TORC1 activation (Schmelzle et al. 2004; Soulard et al. 2010; Ramachandran and Herman 2011; Dechant et al. 2014). Metazoan TORC1 is also regulated by nutrients, specifically amino acids, but requires an additional mitogen/growth factor input for its activation (Laplante and Sabatini 2012). The TORC1 signaling pathway controls cell growth and proliferation by regulating nutrient and energy homeostasis in eukaryotes.
Yeast TORC1 consists of either the Tor1 or Tor2 kinase, Kog1, Lst8, and the nonessential subunit Tco89 (Loewith et al. 2002; Reinke et al. 2004). Loss of either Tor1 or Tco89 causes hypersensitivity to conditions that suppress TORC1 signaling, including nutrient deprivation and the naturally occurring TORC1 inhibitor rapamycin (Loewith and Hall 2011). In yeast, TORC1 resides predominantly on the vacuole’s surface where it is activated by luminal amino acid accumulation (Binda et al. 2009; Dechant et al. 2014). Active TORC1 then signals through at least two well-defined downstream effector pathways. The best characterized is the AGC kinase Sch9, which is directly phosphorylated and activated by TORC1 (Urban et al. 2007). Active Sch9 can then phosphorylate a number of distinct substrates, many of which function directly in ribosomal gene transcription and protein translation (Urban et al. 2007; Huber et al. 2009; Huber et al. 2011). A recent study identified the existence of a TORC1 effector pathway distinct from Sch9 involving the Ypk3 kinase, which phosphorylates ribosomal S6, yet this pathway still remains poorly characterized (Gonzalez et al. 2015).

In the presence of a high-quality nitrogen source, TORC1 also phosphorylates the essential factor Tap42 (Di Como and Arndt 1996; Jiang and Broach 1999; Yan et al. 2012). Phosphorylated Tap42 interacts with the evolutionarily conserved protein phosphatase 2A (PP2A) and PP2A-like phosphatases, which sequester these enzymes onto the vacuolar surface and restrict their access to client substrates (Di Como and Arndt 1996; Jiang and Broach 1999; Yan et al. 2012). Upon nitrogen starvation and reduction of vacuolar amino acids, Tap42-regulated phosphatase complexes are released that then dephosphorylate several key transcription factors, including Gln3 and Gat1 (Beck and Hall 1999; Kuruvilla et al. 2001; Crespo et al. 2002). Dephosphorylation triggers Gln3 and Gat1 dissociation from their cytoplasmic anchor Ure2 and their subsequent nuclear translocation. Nuclear Gln3 and Gat1 then regulate transcription of genes involved in the metabolism of nonpreferred nitrogen sources as part of the nitrogen catabolite repression (NCR) response (Cooper 2002). Each of the Tap42-regulated phosphatases consists of a defining catalytic subunit, as well as additional regulatory subunits that elicit distinct regulatory functions. For example, the catalytic subunit of the PP6 phosphatase complex, Sit4, forms heterodimers with one of four regulatory subunits (Sap4, Sap155, Sap185, or Sap190) that direct its enzymatic activity to specific substrates involved in cell-cycle regulation, transcription, and translation (Luke et al. 1996; Rohde et al. 2004).

Recently, TORC1 has also been implicated in the regulation of chromatin structure through control of histone post-translational modifications. For example, TORC1-dependent transcription of ribosomal protein gene expression was demonstrated to require the Esal histone acetyltransferase, which acetylates histone H4 (Rohde and Cardenas 2003). TORC1 suppression was also shown to cause rapid histone deacetylation of ribosomal DNA (rDNA) repeats and increased transcriptional silencing via recruitment of the sirtuin deacetylase Sir2 (Ha and Huh 2011). Our group recently demonstrated that TORC1 regulates global histone H3 lysine 56 acetylation (H3K56ac) to facilitate ribosomal RNA synthesis by RNA polymerase I through a pathway connected to the sirtuins Hst3 and Hst4 (Chen et al. 2012). However, the relationship between TORC1 signaling and these sirtuins was not defined in detail. A subsequent study demonstrated that TORC1 signaling causes rDNA repeat expansion, while decreased TORC1 activity reduces rDNA repeat number through a mechanism requiring Sir2, Hst3, and Hst4 (Jack et al. 2015). Sirtuin deacetylases are a family of class III NAD+–dependent deacetylases whose catalytic activities are inversely regulated by energy state (Imai and Guarente 2014). Interestingly, in metazoans, TORC1 signaling and sirtuin activity also are regulated in an opposing fashion (Ghosh et al. 2010; Guo et al. 2011; Csibi et al. 2013). Therefore, these studies suggest that TORC1 signaling and sirtuin activity may be coordinated in opposing ways to link control of epigenetic changes needed for anabolism with the availability of environmental nutrients to support these processes.

In this report, we expand on these concepts to demonstrate that nitrogen availability signals through TORC1 to regulate site-specific acetylation of select histone H3 and H4 N-terminal lysine residues besides H3K56ac. We demonstrate that TORC1 controls these modifications globally via repression of Tap42-regulated Sit4 phosphatase activity. TORC1 suppression promotes a decrease in histone acetylation, which requires the activation of Sit4 and the sirtuins Hst3 and Hst4. Specifically in the case of Hst4, reduced TORC1 activity increases Hst4 nuclear localization and reduces Hst4 protein turnover, suggesting that Hst4 nuclear localization may stabilize the enzyme to mediate histone deacetylation under nutrient-limiting conditions. This TORC1-dependent suppression of Sit4/PP6 and downstream negative regulation of sirtuins is biologically relevant as deletions of Sit4, Hst3, or Hst4 rescue some TORC1 mutant phenotypes.

Materials and Methods

Yeast plasmids, strains, and culture conditions

The strains and plasmids utilized are listed in Supplemental Material, Table S1 and Table S2, respectively. Gene deletion and epitope tagging procedures were conducted as described previously (Janke et al. 2004). Unless explicitly stated, all yeast cultures and plating assays were performed in 1% yeast extract/2% peptone/2% dextrose (YPD). Yeast drop-out
synthetic complete (SC) media were prepared as described previously (Laribee et al. 2015). All yeast culture media were purchased from US Biologicals and Research Products International, and antibiotics for selection were obtained from Invitrogen (Carlsbad, CA) or GoldBio. Cells were cultured either at 30°C or room temperature (for the tap42ts experiments) with shaking. For the spotting assays, equal cell numbers from overnight cultures were pelleted, washed, and serially diluted fivefold. Cells were then spotted to the appropriate plates, incubated at the indicated temperatures, and photographed daily. Full-length appropriate plates, incubated at the indicated temperatures, and containing DAPI were purchased from Vector Laboratories (Burlingame, CA).

Histone antibodies were purchased from Active Motif. goat anti-mouse and rabbit anti-rabbit secondary antibodies (Rockland). All antibodies were tested for specificity and sensitivity using Western blot, and appropriately diluted secondary antibodies were used for all immunodetection experiments. We used the following antibodies: 

- α-Myc A14 (Santa Cruz Biotechnology), 
- -FLAG (Stratagene, La Jolla, CA), 
- -RPS6 (Abcam), 
- -phosphoS6 (Cell Signaling), 
- -FLAG (Stratagene, La Jolla, CA), 
- -rabbit HRP-conjugated secondary (The Jackson Laboratory), 
- -HA and -Myc A14 (Santa Cruz Biotechnology), 
- -G6PDH (Sigma, St. Louis, MI), 
- -FLAG (ThermoFisher), and 
- -rabbit FITC-conjugated secondary (Rockland). All histone antibodies were purchased from Active Motif. For confocal microscopy, Vectashield mounting media containing DAPI were purchased from Vector Laboratories (Burlingame, CA).

**RT-quantitative PCR, statistical analyses, and immunoblot analyses**

Total RNA was extracted and 1 μg of DNase I digested RNA was used to synthesize randomly primed complementary DNA (cDNA) using the ImProm II reverse transcription system from Promega (Madison, WI). Gene-specific quantitative PCR (qPCR) with normalization to the SPT15 housekeeping gene was performed and analyzed as previously described (Laribee et al. 2015). Primer sequences are available upon request. All statistical analyses reported in this study were performed using Student’s t-test and the statistical suite available in Microsoft Excel. Whole-cell extracts were prepared and immunoblotting was performed as outlined previously (Laribee et al. 2015). To quantify histone immunoblot results, films were scanned and analyzed by ImageJ software. Histone acetylation states were always normalized to total histone H3 levels to account for any variability in histone levels.

**Indirect immunofluorescence confocal microscopy**

Small scale cultures (10 ml) were cultured to log phase and treated as indicated. The cells were then fixed with 37% formaldehyde, pelleted, and washed twice with 0.1 M potassium phosphate buffer (K₂HPO₄, pH = 6.5) and once with 4 ml P solution (1.2 M sorbitol, 0.1 M K₂HPO₄, pH = 6.5). Pellets were resuspended in 1 ml P solution, 15 μl LongLife Zymolase (15 mg/ml in P solution) and 5 μl β-mercaptoethanol, followed by a 25-min incubation with gentle mixing at room temperature. After zymolase treatment, cells were pelleted, resuspended in 150 μl of P solution, and distributed evenly across two hydrophobic regions of a polylsine-coated microscope slide. Cells were allowed to settle prior to blocking (8% bovine serum albumin, 0.5% Tween 20 in PBS for 20 min). Blocking solution was removed and primary antibody (1:100 α-Myc) was added for a humidified incubation overnight at 4°C. Slides were washed four times with blocking solution and then incubated with FITC-conjugated secondary antibody (1:100) for 60 min at room temperature. The slide washes were repeated four times with blocking solution, twice with PBS, and then a drop of DAPI-containing Vectashield mounting medium was added before coverslip addition and sealing.

**Image analysis in Zen 2 Blue**

Image quantification was performed using Zen 2 Blue Lite, version 2.0.0 software. Two borders were traced onto the image using the Spline Contour tool, one around the cell periphery and the other around the nucleus. After closing each border, values were obtained for the area encompassed by the border, as well as the mean intensity value for each channel inside that space. The area within the nuclear border was multiplied by the mean intensity value for the green channel (FITC-conjugated secondary fluorescence green). This gave a number for the total nuclear fluorescence intensity (TNFI):

\[
\text{TNFI} = \text{nuclear area} \times \text{nuclear mean intensity value}.
\]

This same calculation was repeated using the outer cell border values, providing us a measure of the total cellular fluorescence intensity (TCFI):

\[
\text{TCFI} = \text{cellular area} \times \text{cellular mean intensity value}.
\]

We then performed the following calculation to get the percentage of nuclear protein:

\[
\% \text{nuclear} = \frac{\text{TNFI}}{\text{TCFI}} \times 100.
\]

Cells were chosen for analysis at random, with ~20–40 quantified per condition, per biological replicate (four to six replicates). Only cells with clear nuclear DNA staining and detectable α-Myc signal were quantified. For the immunostaining, cells whose cellular mean intensity was <50 units were considered to have not been penetrated by secondary antibody and were excluded from calculations.

**Protein turnover analysis**

Myc-tagged Hst3 or Hst4 strains were cultured to log phase in 200 ml YPD media. The 50-ml samples were steriley retrieved from each flask prior to cycloheximide (CHX) addition (100 μg/ml). Following treatment, additional 50-ml
 aliquots were taken at the time points indicated. Whole-cell extracts were prepped and analyzed by SDS/PAGE and α-Myc immunoblot. Films were scanned and Hst3 or Hst4 levels were normalized to total protein (G6PDH) levels by ImageJ analysis.

Data availability
The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article.

Results

Metabolic signaling through TORC1 regulates site-specific histone H3 and H4 acetylation

Previously, we demonstrated that TORC1 signaling regulates global H3K56ac (Chen et al. 2012). We wanted to assess whether this effect was unique for H3K56ac, or if TORC1 also contributed to the regulation of other histone H3 and H4 acetyl modifications as well. To explore this possibility, we performed immunoblot analysis for site-specific histone H3/H4 N-terminal acetylation modifications. Duplicate wild-type (WT) cultures, as well as a tco89Δ (nonessential TORC1 subunit), were cultured to log phase and either mock treated or treated with 300 nM rapamycin for 60 min before preparing whole-cell extracts. The concentration and length of rapamycin treatment chosen was confirmed to inhibit TORC1 signaling via phosphoS6 analysis (Figure S1A). Immunoblot analysis identified H3K18ac, H3K23ac, and H4K12ac to be selectively decreased after TORC1 inhibition, whereas all other acetylation states were unaffected (Figure 1, A and B). Furthermore, this effect was not due to globally reduced transcription-coupled histone modifications, as histone H3 lysine 4 trimethylation (H3K4me3), which demarcates transcriptionally active genes, was unaffected (Figure 1A). These changes occurred rapidly as WT cells treated with 200 nM rapamycin for 20 min displayed profoundly reduced global H3K18ac (Figure S1B). Importantly, restoration of WT Tco89 expression in tco89Δ cells completely rescued histone acetylation, thereby demonstrating that the reduced acetylation were solely due to impaired TORC1 activity (Figure 1C).

Nitrogen metabolism and amino acid availability are known activators of TORC1 (Beck and Hall 1999; Cardenas et al. 1999; Hardwick et al. 1999; Crespo et al. 2002). Intriguingly, carbon metabolism has also recently been suggested to activate TORC1 through the vacuolar localized V-ATPase complex, while glycolysis is known to control global levels of many of the same histone acetylation sites we have identified to be regulated by TORC1 (Friis et al. 2009; Dechant et al. 2014). Therefore, we next determined the impact these distinct nutrient signals have on TORC1-regulated acetylation. Changing the carbon source from glucose to a nonfermentable carbon (glycerol) failed to decrease histone acetylation in WT cells, nor did it suppress the acetylation defect observed in the tco89Δ mutant (Figure 1D).

Because cells grow considerably slower in the presence of a nonfermentable carbon, these results also exclude the possibility that the difference in acetylation between WT and tco89Δ is caused by a reduced growth rate in tco89Δ. To assess the contribution of nitrogen metabolism, we cultured WT cells to log phase and mock treated or treated cells for 20 min with 2 mM L-methionine sulfoximine (MSX) to inhibit glutamine synthetase, thus mimicking a nitrogen starvation state. MSX rapidly decreased H3K18ac (Figure 1E, top panel, and F), which we confirmed also suppressed TORC1 activity (Figure 1E, bottom panel). These data demonstrate that nitrogen-, but not carbon-dependent TORC1 activation is responsible for regulating these site-specific histone acetylation states.

TORC1 signaling through the Tap42-Sit4/PP6 phosphatase is required for global site-specific histone acetylation

TORC1 activates multiple downstream effectors, including the Sch9 kinase, which is directly phosphorylated by TORC1 (Urban et al. 2007). To determine if TORC1 regulates histone acetylation through Sch9 activation, we utilized a series of previously described Sch9 plasmids including Sch9WT, Sch9SA, in which the TORC1 target sites are mutated to alanine to mimic the nonphosphorylated form, and Sch9D3E, in which the TORC1 target sites are mutated to acidic residues, rendering it active independent of TORC1 (Urban et al. 2007). Critically, we found that these various mutant forms had similar expression and stability (Figure S2A). Additionally, we confirmed that in sch9Δ, the Sch9D3E plasmid promoted growth on nonfermentable carbon and rapamycin as expected (Figure S2B) (Urban et al. 2007). We also confirmed that in a sch9Δ+pSch9D3E strain, Maf1 phosphorylation (a direct substrate of Sch9) was maintained after rapamycin inhibition (Figure S2C) (Huber et al. 2009; Wei and Zheng 2009), altogether demonstrating that Sch9D3E functioned as expected. However, tco89Δ cells expressing Sch9D3E failed to rescue histone acetylation (Figure 2A). Similarly, a sch9Δ strain did not reduce histone acetylation (Figure 2B), thus demonstrating that TORC1-dependent acetylation is regulated independently of Sch9.

Next, we asked whether TORC1 signaled through its other downstream effector, Tap42, to regulate histone acetylation. Tap42 is an essential gene, so we utilized tap42Δ cells containing a plasmid expressing either WT or temperature-sensitive (ts) tap42 alleles (Yorimitsu et al. 2009). We confirmed the temperature sensitivity of these strains and chose tap42-106 for further analysis because it exhibited the least growth inhibition at 30° (Figure 2C). As a consequence, it would be less likely to have nonspecific effects on acetylation due to loss of viability. WT and tap42-106 cells were cultured to log phase at permissive (25°) temperature before being shifted to 30° for 1 hr. Even at room temperature, where there is no evident growth phenotype (Figure 2C), the tap42-106 mutant exhibited significant H3K18ac reduction, thus implicating Tap42 in histone acetylation regulation (Figure 2, D
and E). Importantly, the decreased acetylation was specific for H3K18ac, as H3K9ac, which is not affected by TORC1 inhibition, was intact (Figure 1A and Figure 2D). At the more restrictive (30°) temperature, tap42-106 caused an even greater reduction in H3K18ac without affecting H3K9ac (Figure 2, D and E). Taken together, these results demonstrate that TORC1 signals through a Tap42-dependent phosphatase pathway to regulate site-specific histone acetylation.

TORC1-dependent Tap42 phosphorylation allows Tap42 to interact with PP2A and PP2A-like phosphatases (P’ase) complexes, which promotes their sequestration to the vacuolar surface and restricts their access to substrates (Di Como and Arndt 1996; Jiang and Broach 1999; Yan et al. 2012). To identify which candidate P’ase is required for TORC1-regulated acetylation, we screened deletion mutants of individual catalytic PP’ase subunits (PPG1/ppg1Δ, PP4/pph4Δ, and PP6/sit4Δ) for the loss of rapamycin-induced deacetylation. Since two redundant catalytic subunits exist for PP2A (Pph21 and Pph22), we instead examined a tpd3Δ, which eliminates the PP2A regulatory subunit and impairs PP2A activity (Hombauer et al. 2007). TORC1 inhibition suppressed H3K18ac, as well as H3K56ac, which we previously identified to be TORC1 dependent, in WT cells and all mutants examined except for sit4Δ (Figure 3A and data not shown).

To confirm the Sit4-catalyzed PP6 phosphatase complex was indeed responsible for TORC1-dependent acetylation, WT and sit4Δ cells were reconstituted with control vector or a vector expressing a Sit4-FLAG construct. In control vector containing sit4Δ cells, higher basal H3K18ac levels were detected that were insensitive to rapamycin, while sit4Δ expressing WT Sit4 reduced H3K18ac and restored rapamycin-dependent acetylation repression (Figure 3B). We also confirmed that sit4Δ alone increased basal H3K18ac while a tco89Δ sit4Δ restored H3K18 acetylation to near WT levels relative to tco89Δ (Figure 3C). These results suggest that Sit4 mediates active repression of histone acetylation under limited TORC1 signaling conditions. As Sit4 partners with one of four different Sap (Sit4-associated protein) regulatory subunits to target distinct substrates, we utilized the same approach from Figure 3A to evaluate the contributions of these ancillary subunits to acetylation regulation. Interestingly, we determined that no single SAP deletion is sufficient to restore WT H3K18 acetylation (compare to the effect of sit4Δ in Figure 3A), although the most significant rescue is observed in the sap4Δ and sap185Δ strains (Figure 3D). This suggests the possibility that these Saps have some functional redundancy as has been suggested previously (Rohde et al. 2004).

The best characterized Sit4 function is in the regulation of the NCR response (Beck and Hall 1999; Crespo et al. 2002). Because nitrogen signaling through TORC1 regulates histone acetylation in a Sit4-dependent fashion, we probed whether Sit4 mediated this effect through activation of the NCR gene expression program. Surprisingly, while Sit4 loss significantly upregulated basal H3K18ac levels as expected, we
determined that neither deletion of the transcription factors Gln3 and Gat1, nor their cytoplasmic anchor Ure2, resulted in a similar increase in acetylation (Figure 3E). These findings demonstrate that while the Sit4-catalyzed PP6 phosphatase complex specifically represses acetylation under TORC1 limiting conditions, it does so independently of the NCR pathway.

**TORC1 opposes sirtuin deacetylase function to promote histone acetylation in a site-specific fashion**

We next tested whether TORC1-dependent acetylation involved sirtuin deacetylases since we previously identified a connection between TORC1 and sirtuins in H3K56ac regulation (Chen et al. 2012). Initially, we cultured WT and tco89Δ cells to log phase and either mock treated or treated with the pan-sirtuin inhibitor nicotinamide before analyzing acetylation. Basal H3K18ac increased in nicotinamide-treated WT cells, suggesting a role for sirtuins in regulating global steady-state acetylation levels, while nicotinamide completely restored acetylation in tco89Δ (Figure 4A). Additionally, treating log phase WT and tco89Δ cultures with the inhibitor FK866, which inhibits nicotinamide phosphoribosyltransferase to suppress NAD⁺ biosynthesis and inactivate sirtuins, completely restored acetylation in tco89Δ (Figure 4B) (Hasmann and Schemainda 2003). TORC1 actively suppresses transcription of the **PNC1** gene, which encodes a nicotinamidase that degrades nicotinamide to nicotinic acid as part of the NAD⁺ salvage pathway (Medvedik et al. 2007). Therefore, one possible link between TORC1 suppression and reduced histone acetylation could be increased Pnc1 enzymatic activity, which would reduce nicotinamide levels to consequently activate sirtuins. Surprisingly, combining tco89Δ with a **pnc1Δ** failed to restore histone acetylation, demonstrating that TORC1 suppression does not reduce histone acetylation by upregulating **PNC1** gene expression (Figure 4C). Collectively, these results demonstrate that TORC1 suppression reduces histone acetylation through a Sit4 and sirtuin-dependent pathway, but it does so independently of changes to the NAD⁺ biosynthetic pathway known to activate sirtuins.

To test whether the decrease in acetylation downstream of TORC1 inhibition might additionally involve class I and II histone deacetylases, we combined tco89Δ with a deletion in either the major class I or class II deacetylase (**rpd3Δ** and **hda1Δ**, respectively) known to regulate transcription and then analyzed whether inactivating either of these enzymes restored histone acetylation (Rundlett et al. 1996). Although basal acetylation was increased at our TORC1-responsive sites (K18ac and K23ac) in both the rpd3Δ and hda1Δ cells, the tco89Δ rpd3Δ and tco89Δ hda1Δ still displayed diminished acetylation, thus confirming that TORC1 acts independently of major class I or class II deacetylases to control global histone acetylation states (Figure 4D). Five sirtuins (Hst1, Hst2, Hst3, Hst4, and Sir2) exist in yeast (Wierman and Smith 2014). To identify which specific sirtuin(s) are connected to the TORC1 pathway, we generated combinatorial mutants of tco89Δ paired with deletions of the individual
sirtuins and analyzed global histone acetylation. Each double mutant resulted in a unique array of acetylation changes, but only the $tco89^{-}$ $hst4^{-}$ restored H3K18ac, H3K23ac, and H4K12ac to near WT levels (Figure 4, E and F). These data are consistent with our previous finding that $hst3^{-}$ or $hst4^{-}$ restores H3K56ac in $tco89^{-}$ and suggest that TORC1-dependent regulation of site-specific H3/H4 N-terminal acetylation requires repression of PP6 activity and downstream suppression of sirtuin deacetylases, particularly Hst4.

**TORC1 regulates Hst4 cellular localization and stability**

Recently, inhibition of the mammalian TORC1 (mTORC1) complex was demonstrated to induce transcriptional upregulation of SIRT4 to promote glutamine anaplerosis (Csibi et al. 2013). We next assessed if the Hst4-dependent decrease in histone acetylation upon TORC1 inhibition could be explained by increased Hst4 expression. To address this, we integrated a 9xMyc epitope at the HST4 genomic locus in WT and $tco89^{-}$ backgrounds, and then cells were either mock treated or treated with MSX or rapamycin for the indicated times. No significant difference in Hst4 levels was detected in WT cells treated for 20 min with either MSX or rapamycin; however, both $tco89^{-}$ and WT cells treated with rapamycin for 60 min displayed increased Hst4 protein levels compared to the mock-treated WT control (Figure 5A and Figure S3). These results were surprising since we previously determined that 20 min of rapamycin or MSX treatment was sufficient to effectively decrease histone acetylation (Figure 1E and Figure S1B), yet Hst4 protein levels do not increase until much later after TORC1 suppression. We evaluated the other sirtuins in response to TORC1 inhibition and found that only Sir2 protein was similarly elevated (Figure S3). However, Sir2 is unlikely to contribute to the decrease in histone acetylation upon TORC1 limitation, since a $tco89^{-}$ sir2^{-} mutant failed to restore TORC1-responsive histone acetylation (Figure 4, E and F).

To ascertain whether Sit4 contributes to the regulation of Hst4 protein expression, we generated sit4^{-} and $tco89^{-}$ sit4^{-} strains in the Hst4-9xMyc background and examined Hst4 levels. While $tco89^{-}$ increases Hst4 expression, the sit4^{-} dramatically reduced Hst4 levels relative to WT (Figure 5B). Intriguingly, the sit4^{-} $tco89^{-}$ returned Hst4 expression to WT levels, suggesting that TORC1 acts to repress PP6 activity and limit Hst4 protein expression (Figure 5B). When the experiment was repeated in a sap4^{-} background, which is the regulatory subunit that most robustly restored histone acetylation in response to rapamycin (Figure 3D), we observed a comparably modest effect on Hst4 protein levels (Figure 5B). These findings are in line with the possibility that some functional...
redundancy may exist between these PP6 regulatory subunits in relation to Hst4 regulation. We then confirmed this Sit4-dependent, TORC1-responsive effect on Hst4 levels was not due to transcriptional or post-transcriptional changes in HST4 messenger RNA (mRNA) expression by analyzing HST4, as well as HST3 and SIR2 gene expression by qPCR. Expression of HST3 and HST4 was not increased in the tco89Δ background (Figure 5C), and while we do detect a minor, but statistically significant decrease in SIR2 mRNA expression in tco89Δ relative to WT, its significance is currently unclear.

Hst3 protein degradation has been previously shown to be cell-cycle regulated through Cdk1-dependent phosphorylation and subsequent SCF$	ext{Cdc4}$-dependent ubiquitination, leading to its proteasome-mediated degradation (Delgoshaie et al. 2014; Edenberg et al. 2014). Whether any phosphatase opposes this Cdk1-dependent phosphorylation remains unknown. Hst4 was also identified as a candidate SCF$	ext{Cdc4}$ substrate, although control of its turnover has not been examined in detail (Mark et al. 2014). We wondered whether TORC1-dependent regulation of Hst4 protein expression occurs through altered protein stability. WT and tco89Δ strains expressing Hst4-9xMyc were cultured to log phase and then treated with 50 μg/ml CHX to inhibit protein synthesis. As a comparison, we performed an identical experiment with Hst3-9xMyc-expressing cells. Intriguingly, we found that while Hst4 turnover was reduced in tco89Δ compared to WT, Hst3 degradation was accelerated in the TORC1 mutant (Figure 5, D and E). In addition we show that in a tco89Δ sit4Δ Hst4-9xMyc strain, which previously restored WT levels of Hst4 expression compared to a tco89Δ alone (Figure 5B), the rate of Hst4 turnover strongly resembles that detected in WT cells (Figure 5D). Together, these data demonstrate that TORC1 signaling via the Sit4 phosphatase complex regulates the proteolytic decay of the sirtuins Hst4 and Hst3 in an inverse fashion, such that TORC1 promotes Hst4 turnover while it stabilizes Hst3.

**Reduced TORC1 signaling and subsequent Sit4 activation promotes Hst4 nuclear accumulation, which precedes the increase in Hst4 protein levels**

A subset of metazoan sirtuins have been shown to actively shuttle between the nucleus and the cytoplasm as a function of cell-cycle progression or in response to cellular stress (Vaquero et al. 2006; Scher et al. 2007; Tanno et al. 2007; Hisahara et al. 2008). Hst4, which has been proposed to be the yeast SIRT3 ortholog, is normally distributed between the cytoplasm and nucleus. Upon biotin starvation, however, Hst4 accumulates at mitochondria to facilitate mitochondrial protein deacetylation (Madsen et al. 2015). As mentioned previously, the best-characterized role that TORC1 suppression, and consequent Sit4 activation, has is to induce nuclear localization of transcription factors controlling the NCR pathway (Cooper 2002). We therefore considered the possibility that reduced TORC1 signaling might also stabilize Hst4 by...
altering its cellular distribution. To test this, WT and tco89Δ cells expressing either Hst4-9xMyc or the other tagged sirtuins, were cultured to log phase before performing α-Myc immunostaining and quantitative confocal microscopy analysis. Random fields of cells were captured and the amount of nuclear sirtuin was manually quantified as described in Materials and Methods. Representative images for Hst4 are shown in Figure 6A and for all five sirtuins in Figure S4. Intriguingly, tco89Δ enhanced the Hst4 nuclear pool, even after accounting for the increase in Hst4 protein levels (Figure 6, A and B). A minor increase in Sir2 nuclear accumulation was also detected but none of the other sirtuins, including Hst3, exhibited a significant change in localization (Figure 6B and Figure S4). To specifically address whether the increase in Hst4 nuclear localization occurred before or after Hst4 protein levels were increased, we repeated the WT and tco89Δ Hst4-9xMyc immunostaining experiment, this time including an additional WT sample treated with 200 nM rapamycin for 20 min. We detected maximal Hst4 relocalization within 20 min after rapamycin addition, which was approximately equivalent to the amount of nuclear-localized Hst4 observed in tco89Δ (Figure 6C and Figure S4). This relocalization is far earlier than the observed increase in protein levels (Figure 5A), and it correlates with the decrease in histone acetylation (Figure S1B). Finally, we asked whether this relocalization was dependent on Sit4 activation downstream of TORC1 inhibition, since Sit4 regulates Hst4 protein levels (Figure 5, B and D). Strikingly, we found that in a tco89Δ sit4Δ Hst4-9xMyc strain, the cytoplasmic population of Hst4 is restored to near WT levels (Figure 6, A and C), supporting the hypothesis that decreased TORC1 signaling triggers Sit4/PP6 activation, which
promotes Hst4 nuclear localization and stabilization to mediate site-specific histone deacetylation.

**Hst4-regulated histone acetylation controls a subset of TORC1-dependent biological functions**

TORC1 was identified previously to regulate acetylation of the RP genes through the recruitment of the Esa1 acetyltransferase (Rohde and Cardenas 2003). The initial Tco89 characterization conducted by Reinke et al. (2004) determined tco89Δ has no effect on RP gene transcription. Given the observed acetylation defects in tco89Δ, we chose to independently determine whether reduced TORC1 signaling may alter RP gene expression through an Hst4-dependent mechanism. Similar to the findings reported by Reinke et al. (2004), we found that although TORC1 activity is reduced in tco89Δ (Figure S1A), no significant decrease in tco89Δ RP gene expression occurred (Figure 7A). Additionally, the hst4Δ had no effect on RP gene expression, and while tco89Δ hst4Δ modestly reduced expression of RPL23B, tco89Δ hst4Δ did not affect the expression of the other RP genes (Figure 7A). These data indicate that the residual TORC1 activity retained in tco89Δ is sufficient to maintain normal RP gene expression and that Hst4 does not significantly contribute to their regulation.

Previous studies demonstrated that exposure to nutrient starvation or rapamycin treatment causes tco89Δ cells to exit the cell cycle and enter into an irreversible, G0-like growth arrested state (Binda et al. 2009). We next asked whether restoration of TORC1-mediated histone acetylation could rescue the sensitivity of tco89Δ mutants to multiple conditions that impair TORC1 signaling (Hosiner et al. 2009; Kapitzky et al. 2010). We found this effect was reversed in
a tco89Δ hst4Δ mutant, and to a lesser extent in tco89Δ hst3Δ, under low (5 nM) rapamycin concentrations (Figure 7B). Surprisingly, at slightly higher (10 nM) rapamycin concentrations, neither double mutant grew significantly (Figure 7B). Intriguingly, the tco89Δ sit4Δ failed to restore growth in the presence of even the low rapamycin concentration; however, combining tco89Δ with either hst3Δ, hst4Δ, or sit4Δ completely restored growth on both hydroxyurea and arsenic (Figure 7B). These results demonstrate that sirtuins are important for maintaining the G0-like growth-arrested state that TORC1 mutants exhibit when exposed to environmental stressors that inhibit TORC1 activity. However, while Sit4 loss can rescue this growth arrest in the presence of some TORC1 inhibitors, it appears to have functions independent of its link to sirtuins that are required for TORC1 impaired cells to escape the growth arrest induced specifically by rapamycin.

Discussion

The epigenome of eukaryotic cells is responsive to environmental stimuli, including changes in nutrient availability. Previous studies have shown that TORC1 signaling affects epigenetic processes, including mediating Esa1 recruitment to RP genes to acetylate histone H4, as well as regulating global H3K56ac (Rohde and Cardenas 2003; Chen et al. 2012). In this report, we significantly expand our understanding of the mechanistic link between TORC1 signaling and its downstream effects on site-specific histone acetylation modifications. We demonstrate for the first time that in logarithmically growing cells, TORC1 inhibition results in a rapid, direct, and selective deacetylation of H3K18, H3K23, and H4K12 on the histone H3/H4 N termini. To date, studies examining TORC1-regulated acetylation have used the TORC1 inhibitor rapamycin to examine the functional consequences TORC1 activity has on chromatin (Rohde and Cardenas 2003; Ha and Huh 2011; Chen et al. 2012). And while we also utilized rapamycin for this purpose, we extended our analyses to include other methods to suppress TORC1, including subunit deletion (tco89Δ) and a pharmacological inhibitor of nitrogen metabolism (MSX) (Crespo et al. 2002). We provide clear evidence that either of these additional conditions are sufficient to decrease site-specific histone H3/H4 acetylation, thus implicating nitrogen-dependent TORC1 activation as a significant regulator of histone acetylation.

Previous studies demonstrated that stationary phase cells exhibit reduced histone acetylation at many of the same residues identified above (Friis et al. 2009). These acetylation states were rescued by glucose refeeding, suggesting that carbon metabolism is a significant regulator of these histone modifications. However, we find that culturing WT and tco89Δ cells in media containing either a preferred, fermentable (glucose) carbon or a nonpreferred, nonfermentable (glycerol) carbon source, resulted in no change to the spectrum of observed histone acetylation defects. Our data argue against the possibility that crosstalk between carbon metabolism and TORC1 regulation, as was previously suggested, is ultimately responsible for the chromatin effects we observe (Schmelzle et al. 2004; Dechant et al. 2014). Instead, our findings suggest that in actively growing cells, nitrogen signaling through TORC1 is the predominant regulator of these specific histone acetylation states. As a consequence, the vacuole likely serves a vital link in the transmission of environmental nutrient sufficiency to the epigenetic regulatory apparatus via TORC1 activation. Indeed, a previous genetic screen of the systematic yeast deletion collection identified several vacuole mutants, including many V-ATPase mutants, to be necessary for maintenance of global histone acetylation (Peng et al. 2008).

Our study also provides a more detailed mechanistic understanding of our previous work, which identified a functional relationship between TORC1, sirtuins, and the regulation of H3K56ac. We demonstrate by direct pharmacological inhibition, cofactor depletion, and individual sirtuin deletion that TORC1 signaling specifically opposes histone deacetylation mediated by several sirtuin family members. Importantly, we also demonstrate the important role of TORC1-mediated histone acetylation does not impact RP gene transcription but is involved in the DNA-damage response and cell-cycle progression. (A) WT, tco89Δ, hst4Δ, and tco89Δ hst4Δ cells were cultured to log phase, total RNA was extracted, cDNA was synthesized, and qPCR performed with the indicated primer sets. Data are the average and SD of three independent experiments and significance was determined by Student’s t-test. (B) WT, tco89Δ, hst3Δ, tco89Δ hst3Δ, hst4Δ, tco89Δ hst4Δ, sit4Δ, and tco89Δ sit4Δ were cultured overnight to saturation. Equal numbers of cells were serially diluted fivefold and spotted onto YPD, YPD 5 mM rapamycin, YPD 10 mM rapamycin, YPD 100 mM hydroxyurea (HU), and YPD 1 mM arsenic oxide (AsO3). Images were taken 4 days after spotting.
Hst4 was the only sirtuin that, when deleted in a \textit{tco89Δ} background, restored acetylation across all of the known TORC1-regulated residues. This suggests that while TORC1 may have a general role in opposing sirtuin activity, Hst4 is likely to be one of the predominant sirtuins whose function is negatively affected by signaling through this pathway. The intimate link between TORC1 and the sirtuins is interesting, considering that both are responsive to cellular energetics, albeit inversely. This relationship appears to be conserved at some level across all eukaryotes (Ha and Huh 2011; Chen et al. 2012; Csibi et al. 2013; Jack et al. 2015). Together, our data support a model whereby TORC1 and sirtuin activity are regulated in an opposing manner to coordinate both nitrogen and energy metabolism with epigenetic control to facilitate productive cell growth and proliferation.

The presented findings also implicate TORC1-dependent suppression of the Tap42-associated \textit{Sit4} containing PP6 phosphatase complex in the regulation of histone acetylation. We demonstrate that these effects are independent of changes in \textit{HST4} gene expression, and that \textit{Sit4}-regulated histone acetylation suppression is separable from \textit{Sit4}-dependent activation of the NCR pathway. Instead, we provide support for a mechanism in which \textit{Sit4} activation promotes both a rapid cytoplasmic-to-nuclear redistribution of Hst4, and an eventual increase in Hst4 protein levels due to reduced protein turnover. Considering that maximal Hst4 relocalization occurs within 20 min post-TORC1 inhibition, but Hst4 protein does not accumulate significantly until ~40 min later, Hst4 movement to the nucleus may function to shield a fraction of the enzyme from proteolytic turnover. These effects on Hst4 are unique, as identical experiments with a highly similar sirtuin, Hst3, revealed that lower TORC1 activity does not affect Hst3 nuclear localization but does decrease Hst3 protein stability. How TORC1-dependent suppression of the \textit{Sit4} phosphatase causes these opposing effects on Hst3 and Hst4 stability is the subject of ongoing investigation. A distinct possibility is that changes in nutrient sufficiency modify TORC1-regulated \textit{Sit4} activity to affect Hst3 and Hst4 phosphorylation status and regulate their SCP\textsuperscript{Calc} dependent turnover. Such a regulatory mechanism might better coordinate the cellular growth response with cell division timing, as all of these factors are also intimately connected to cell-cycle regulation.

Histone acetylation regulates gene expression, in part, by disrupting histone-DNA contacts to promote chromatin decondensation, as well as by serving as docking sites for bromodomain- or YEATS-domain-containing regulatory factors (Sanchez et al. 2014; Shanle et al. 2015). While TORC1 signaling is known to regulate the transcription of a large number of genes involved in cell growth and proliferation, the functional contributions of our newly characterized TORC1-regulated histone acetylation states have in relation to these processes are currently unclear. In agreement with a previous report (Reinke et al. 2004), we find that in the \textit{tco89Δ} TORC1 mutant, RP gene expression is largely unaffected. We believe this demonstrates that there is sufficient residual TORC1 activity present in these mutants to maintain basal Sch9 activation and RP gene expression. However, TORC1 transcriptional regulation extends beyond just RP genes, so it is likely that other genes will be preferentially sensitive to the reduced histone acetylation we have detected. Identifying which acetyl lysine binding proteins are affected by altered TORC1 signaling will be critical to completely define how TORC1-dependent histone acetylation contributes to transcriptional programs controlling growth and proliferation.

Cells deficient in TORC1, such as \textit{tco89Δ} mutants, are acutely sensitive to environmental exposures that suppress TORC1 signaling, including rapamycin, DNA replication inhibitors, metals, and a variety of other stress-inducing agents. And while these compounds are often grouped together as having a generalized inhibitory effect on TORC1, the downstream cellular responses they elicit can vary substantially (Tate and Cooper 2013; Hughes Hallett et al. 2014). We demonstrate that the TORC1–PP6–sirtuin pathway outlined above is functionally relevant for a subset of TORC1-regulated biological processes, as either \textit{hst3Δ} or \textit{hst4Δ} can suppress \textit{tco89Δ} sensitivity to arsenic, hydroxyurea, and low concentrations of rapamycin. We believe that these sirtuins may contribute to enforcing the cell-cycle exit that occurs in \textit{tco89Δ} exposed to TORC1 suppressive conditions, in part through their role in chromatin deacetylation. Loss of either sirtuin may facilitate cell-cycle reentry following TORC1 inhibition by maintaining a more basally hyperacetylated, transcriptionally permissive chromatin state, thus allowing for more facile induction of genes necessary to escape the arrested state. The recent identification of a glucose-regulated histone acetylation pathway necessary for transcription of the \textit{CLN3} cyclin, and subsequent entry into the cell cycle, provides support for this concept (Shi and Tu 2013).

Furthermore, we demonstrate that while \textit{sit4Δ} also rescues the sensitivity of \textit{tco89Δ} to both hydroxyurea and arsenic, it fails to promote growth on rapamycin. These results suggest the possibility that \textit{Sit4} has sirtuin-independent roles in regulating cell growth control that are necessary for TORC1-deficient cells to escape the rapamycin-induced growth arrest (Sutton et al. 1991; Fernandez-Sarabia et al. 1992). There is a precedent for Tap42-associated phosphatases regulating such dynamics as it was previously shown that the nitrogen-responsive \textit{Rim15}-dependent phosphorylation of PP2A phosphatase plays a critical role in cell-cycle entry, as well as exit from quiescence (Bontron et al. 2013). Another candidate explanation is that \textit{Sit4} association with distinct Sap regulatory factors may have context-specific effects. In support of this, a previous report determined that the function \textit{Sit4} has in response to TORC1 suppression depends on the identity of the associated Sap regulator (Rohde et al. 2004). Unraveling these differences, and their specific role in sirtuin regulation, will be the subject of future studies.
Overall, our data define a mechanism by which nitrogen signaling through TORC1 activation controls specific epigenetic modifications that are important for TORC1-regulated cell growth and proliferation. Because these pathways are conserved in mammals, we consider it highly likely that deregulation of nitrogen metabolism and aberrant mTORC1 activation, which occurs in many cancers, might alter the epigenome through similar mechanisms to promote disease pathogenesis.

Acknowledgments

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Saccharomyces cerevisiae TORC1 Controls Histone Acetylation by Signaling Through the Sit4/PP6 Phosphatase to Regulate Sirtuin Deacetylase Nuclear Accumulation

Jason J. Workman, Hongfeng Chen, and R. Nicholas Laribee
Figure S1
Figure S1. Optimization of rapamycin treatment conditions. (A) Wild-type (WT) and tco89Δ cells were cultured to log phase, and WT cells were then mock treated, or treated with varying rapamycin concentrations (25 nM or 300 nM) for either 20 minutes or 60 minutes. Whole-cell extracts were prepared and analyzed by immunoblot (IB) with the indicated antibodies. Short and long exposures for the phosphoS6 blots are presented for clarity. (B) WT cells were cultured to log phase and then mock treated or treated with 300 nM rapamycin for 20 minutes. Extracts were prepared and analyzed as in (A).
Figure S2. Control experiments for mutant Sch9 plasmids. (A) *sch9Δ* cells transformed with control vector (CV), wild-type Sch9 (pSch9<sup>WT</sup>), and mutant forms where Sch9 activity is no longer...
responsive to TORC1 (pSch9<sup>WT</sup> and pSch9<sup>2D3E</sup>) were grown to log phase in selective media. Whole-cell extracts were prepared and analyzed by immunoblot as indicated. (B) Strains from (A) were grown to saturation, five-fold serially diluted, and spotted onto the indicated plate media. (C) WT Maf1-9xMyc cells were transformed with CV, while the Maf1-9xMyc sch9Δ strain was transformed with CV, pSch9<sup>WT</sup> and pSch9<sup>2D3E</sup>. Cells were grown to log phase and then mock or 300nM rapamycin treated for 60 minutes. Extracts were prepared, resolved on 8% SDS-PAGE, and immunoblotted as shown.
Figure S3
Figure S3. Sirtuin protein levels in response to TORC1 inhibition. Representative immunoblots of no tag control, and WT or tco89Δ cells expressing the 9xMyc epitope-tagged sirtuins. Data shown is representative of at least 3 independent biological replicates.
**Figure S4.** Sirtuin subcellular localization in response to TORC1 inhibition. (A-E) No tag control, and WT or tco89Δ cells expressing the 9xMyc epitope-tagged sirtuins were cultured to log phase, treated as indicated, and imaged by indirect immunofluorescence as described in the methods. Images are representative of at least three independent biological replicates, and are a subset of the quantification shown in Figure 6B and 6C.
| Strain     | Genotype                        | Source               |
|------------|---------------------------------|----------------------|
| BY4741     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 | OPEN Bio             |
| tco89Δ     | BY4741; tco89Δ::KanMX          | OPEN Bio             |
| sch9Δ      | BY4741; sch9Δ::KanMX           | OPEN Bio             |
| pnc1Δ      | BY4741; pnc1Δ::KanMX           | OPEN Bio             |
| rpd3Δ      | BY4741; rpd3Δ::KanMX           | OPEN Bio             |
| hda1Δ      | BY4741; hda1Δ::KanMX           | OPEN Bio             |
| gln3Δ      | BY4741; gln3Δ::KanMX           | OPEN Bio             |
| gat1Δ      | BY4741; gat1Δ::KanMX           | OPEN Bio             |
| ure2Δ      | BY4741; ure2Δ::KanMX           | OPEN Bio             |
| tpd3Δ      | BY4741; tpd3Δ::KanMX           | OPEN Bio             |
| ppg1Δ      | BY4741; ppg1Δ::KanMX           | OPEN Bio             |
| pph3Δ      | BY4741; pph3Δ::KanMX           | OPEN Bio             |
| sit4Δ      | BY4741; sit4Δ::KanMX           | OPEN Bio             |
| sap4Δ      | BY4741; sap4Δ::KanMX           | OPEN Bio             |
| sap155Δ    | BY4741; sap155Δ::KanMX         | OPEN Bio             |
| sap185Δ    | BY4741; sap185Δ::KanMX         | OPEN Bio             |
| sap190Δ    | BY4741; sap190Δ::KanMX         | OPEN Bio             |
| YNL387     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::HphNT1 hst3Δ::NAT | (CHEN et al. 2013) |
| YNL389     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::KanMX hst4Δ::NAT | (CHEN et al. 2012) |
| Y3033      | W303-1A tap42Δ::HIS3 pRS414-TAP42 | (YORIMITSU et al. 2009) |
| Y3032      | W303-1A tap42Δ::HIS3 pRS415-tap42-11 | (YORIMITSU et al. 2009) |
| Y3035      | W303-1A tap42Δ::HIS3 pRS414-tap42-109 | (YORIMITSU et al. 2009) |
| Y3034      | W303-1A tap42Δ::HIS3 pRS414-tap42-106 | (YORIMITSU et al. 2009) |
| YNL541     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::HphNT1 hst3Δ::KanMX | This Study |
| YNL622     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT sit4Δ::KanMX | This Study |
| YNL487     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT | This Study |
| YNL516     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT hst1Δ::KanMX | This Study |
| YNL519     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT hst2Δ::KanMX | This Study |
| YNL517     | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT sir2Δ::KanMX | This Study |
| Strain  | Genotype                  | Marker  | Notes                  |
|---------|---------------------------|---------|------------------------|
| YNL502  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT rpd3Δ::KanMX | This Study |
| YNL670  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT hda1Δ::KanMX | This Study |
| YNL678  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst1-9xMyc::KanMX | This Study |
| YNL681  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT Hst1-9xMyc::KanMX | This Study |
| YNL685  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst2-9xMyc::KanMX | This Study |
| YNL700  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT Hst2-9xMyc::KanMX | This Study |
| YNL698  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst3-9xMyc::KanMX | This Study |
| YNL676  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT Hst3-9xMyc::KanMX | This Study |
| YNL612  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst4-9xMyc::KanMX | This Study |
| YNL614  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT Hst4-9xMyc::KanMX | This Study |
| YNL687  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Sir2-9xMyc::KanMX | This Study |
| YNL689  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 tco89Δ::NAT Sir2-9xMyc::KanMX | This Study |
| YNL716  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst4-9xMyc::HYGRO sit4Δ::KANMX | This Study |
| YNL718  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst4-9xMyc::HYGRO sit4Δ::KANMX tco89Δ::HphNT1 | This Study |
| YNL775  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst4-9xMyc::KANMX sap4Δ::HYGRO | This Study |
| YNL776  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Hst4-9xMyc::KANMX sap4Δ::HYGRO tco89Δ::NAT | This Study |
| YNL778  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Maf1-9xMyc::NAT | This Study |
| YNL780  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 BY4741 Maf1-9xMyc::NAT sch9Δ KANMX | This Study |
Table 2. Yeast plasmids.

| Parent Plasmid      | Plasmid Description                                      | Source                  |
|---------------------|----------------------------------------------------------|-------------------------|
| pADH TCO89-MYC      | $ADH1_{prom}TCO89-MYC; URA3; CEN6/ARS$                   | This study              |
| pADH SIT4-FLAG      | $ADH1_{prom}SIT4-FLAG; URA3; CEN6/ARS$                   | This study              |
| pRS416              | $CEN6; ARS4; URA3$                                       | (BRACHMANN et al. 1998) |
| pJU677              | pRS416; SCH9-6xHA                                        | (URBAN et al. 2007)     |
| pJU790              | pRS416; SCH9-6xHA (T723A, S726A, T737A, S758A, S765A)   | (URBAN et al. 2007)     |
| pJU855              | pRS416; SCH9-6xHA (T723D, S726D, T737E, S758E, S765E)   | (URBAN et al. 2007)     |