Alterations of signaling pathways in response to chemical perturbations used to measure mRNA decay rates in yeast

NICHOLE ESHLEMAN, XIANGXIA LUO, ANDREW CAPALDI, and J. ROSS BUCHAN
Department of Molecular and Cellular Biology, University of Arizona, Tucson, Arizona 85721, USA

ABSTRACT
Assessing variations in mRNA stability typically involves inhibiting transcription either globally or in a gene-specific manner. Alternatively, mRNA pulse-labeling strategies offer a means to calculate mRNA stability without inhibiting transcription. However, key stress-responsive cell signaling pathways, which affect mRNA stability, may themselves be perturbed by the approaches used to measure mRNA stability, leading to artifactual results. Here, we have focused on common strategies to measure mRNA half-lives in yeast and determined that commonly used transcription inhibitors thiolutin and 1,10 phenanthroline inhibit TORC1 signaling, PKC signaling, and partially activate HOG signaling. Additionally, 4-thiouracil (4tU), a uracil analog used in mRNA pulse-labeling approaches, modestly induces P-bodies, mRNA–protein granules implicated in storage and decay of nontranslating mRNA. Thiolutin also induces P-bodies, whereas phenanthroline has no effect. Doxycycline, which controls “Tet On/Tet Off” regulatable promoters, shows no impact on the above signaling pathways or P-bodies. In summary, our data argues that broad-acting transcriptional inhibitors are problematic for determining mRNA half-life, particularly if studying the impacts of the TORC1, HOG, or PKC pathway on mRNA stability. Regulatable promoter systems are a preferred approach for individual mRNA half-life studies, with 4tU labeling representing a good approach to global mRNA half-life analysis, despite modestly inducing P-bodies.

Keywords: Hog1; P-bodies; PKC; TORC1; mRNA decay

INTRODUCTION
Exposure of cells to stress triggers compensatory changes in gene expression, of which changes in mRNA decay play a key role (Fan et al. 2002; Pérez-Ortín et al. 2007). Enhancing mRNA decay typically affords a more rapid means to reduce mRNA levels than transcriptional inhibition alone, and thus adjustments in both processes are often observed (García-Martínez et al. 2004; Molin et al. 2009). However, understanding how stress events are sensed and relayed mechanistically to alterations in mRNA stability remains poorly understood. Generally, stress-responsive signaling pathways are more closely studied for their effects on transcription and translation, though several are implicated in regulating mRNA decay in yeast and other species; these include the TORC1 pathway, HOG pathway, and PKC pathway. The TORC1 (target of rapamycin complex) pathway, conserved throughout eukaryotes, regulates cell growth and metabolism in response to environmental and nutrient cues, particularly carbon and nitrogen availability (Barbet et al. 1996; Thomas and Hall 1997; Loewith and Hall 2011). Inhibition of TORC1 activity during diauxic shift or treatment with rapamycin reduces levels of several reporter and endogenous mRNAs due to enhanced deadenylation or mRNA decapping (Albig and Decker 2001). Conversely, TORC1 inhibition stabilizes select stress-responsive genes (Talarek et al. 2013). Global analysis of yeast mRNA half-lives reveals that rapamycin treatment increased the average mRNA half-life, though also facilitated faster decay of numerous transcripts, including all ribosomal protein mRNAs (Munchel et al. 2011). Finally, TORC1 can directly phosphorylate Dcp2 (mRNA decapping enzyme), which correlates with the destabilization of autophagy mRNAs in C. neoformans (Hu et al. 2015).

The HOG (high osmolarity glycerol) pathway is a stress-activated protein kinase pathway that helps yeast cells sense and adapt to various environmental stresses, particularly osmotic stress (Brewster and Gustin 2014). During mild osmotic stress, most yeast housekeeping genes exhibit decreased mRNA stability, whereas stress-responsive genes increase in mRNA stability (Molin et al. 2009; Romero-Santacreu et al. 2009). However, in hog1Δ strains, destabilization of housekeeping mRNAs was impaired...
(Molin et al. 2009; Romero-Santacreu et al. 2009), with one study also suggesting stabilization of stress-responsive mRNAs was inhibited (Molin et al. 2009). Ste20, a kinase activator of HOG signaling, phosphorylates Dcp2 in response to various stresses, slowing the decay of select ribosomal protein mRNAs (Yoon et al. 2010). In human cells, p38 (Hog1 ortholog) activity is a key regulator of AU-rich element (ARE)-containing cytokine mRNAs (Yoon et al. 2010). During and following osmotic stress, HOG signaling has been linked to assembly and disassembly of P-bodies in yeast (Brengues et al. 2005; Romero-Santacreu et al. 2009). P-bodies are cytoplasmic assemblies of non-translating mRNA–protein complexes (mRNPs) that are implicated in regulation of mRNA repression and mRNA decay (Jain and Parker 2013). Thus, some effects of Hog1 on mRNA stability may also reflect altered P-body functions.

The PKC (protein kinase C) pathway in yeast is best studied for its role in sensing cell wall integrity. Under nutrient poor growth conditions, absence or conditional inactivation of PKC signaling pathway proteins (Pkh1, Pkh2, and Pkc1) impairs deadenylation and decay of various reporter and endogenous mRNAs, and causes synthetic growth defects in dhh1Δ null strains (Luo et al. 2011); Dhh1 is a P-body localizing protein that enhances mRNA decapping and translation repression. Pkh1/2 and Pkc1 also aid stress-induced increases in P-bodies under nutrient-poor conditions, though not under nutritionally rich conditions (Luo et al. 2011). Cell wall stress also triggers P-body induction in a PKC signaling-dependent manner, and relocalization of mRNAs that normally accumulate under cell wall stress to P-bodies, suggestive of possible alterations in their stability (García et al. 2019). Finally, as with p38, several studies in human cells indicate PKC signaling regulates stability of ARE-containing cytokine mRNAs (Schoenberg and Maquat 2012).

In this work, we examine how various chemical perturbations used to measure mRNA decay rates in yeast affect stress-responsive signaling pathways. Specifically, we examine at standard concentrations the effects of two general transcription inhibitory molecules, 1,10 phenanthroline (phenanthroline) and thiolutin, doxycycline (which controls Tet off/Tet on promoter systems) and 4-thiouracil (4tU), which is used in RNA pulse-labeling approaches. All approaches are commonly used to assess individual and global mRNA decay rates in yeast (Parker et al. 1991; Hughes et al. 2000; Grigull et al. 2004; Coller 2008; Munchel et al. 2011; Pérez-Ortínez et al. 2013; Wada et al. 2017; Chan et al. 2018), and yet often lead to highly divergent results. Our western blot and microscopy findings suggest part of the reason for this may be aberrant activation or inhibition of signaling pathways that regulate mRNA decay, particularly when using phenanthroline or thiolutin.

RESULTS AND DISCUSSION

Phenanthroline and thiolutin impair TORC1 signaling via the Sch9 branch

We first examined how 10 min of treatment with our transcription-regulating small molecules affected TORC1 signaling by examining Sch9 phosphorylation status in two common yeast strains (BY4741 and W303). Sch9 lies within one of two major downstream effector branches of the TORC1 signaling pathway (Fig. 1A; Loewith and Hall 2011; Hughes Hallett et al. 2014). Sch9 is directly phosphorylated by TORC1 under optimal growth conditions, but is rapidly dephosphorylated when TORC1 is inactivated, and thus is a commonly used indicator of TORC1 signaling activity. Rapamycin was used as a positive control of TORC1 inactivation, and after 10 min of rapamycin treatment, Sch9 is indeed strongly dephosphorylated (Fig. 1B, lanes 1,2; Urban et al. 2007; Hughes Hallett et al. 2014).

Strikingly, phenanthroline and thiolutin inhibit TORC1 to the same extent as rapamycin (Fig. 1B, lanes 3,4,7,8). In contrast, doxycycline, 4tU and β-estradiol have no effect on Sch9 phosphorylation status (Fig. 1B, lanes 5,6,9,10, Fig. 1E). These effects were consistent between both yeast strain backgrounds and were not a result of solvents (ethanol or DMSO) used to dissolve the small molecules (Fig. 1C). These data suggested that phenanthroline and thiolutin are inhibitors of TORC1 signaling.

Phenanthroline activates the PP2A TORC1 signaling branch

We next looked at the PP2A–Tap42 branch of TORC1 signaling by looking at Par32 phosphorylation, a downstream target and indicator of PP2A–Tap42 branch signaling activity (Fig. 1A; Hughes Hallett et al. 2014; Boeckstaens et al. 2015).

As before, the effect of our molecules was examined after 10 min of treatment with rapamycin serving as a control for TORC1 inhibition (which activates PP2A–Tap42 signaling); as expected, clear Par32 phosphorylation was observed (Fig. 1D, lanes 1,2). Interestingly, phenanthroline, but not thiolutin, caused a similar shift after 10 min of treatment (Fig. 1D, lanes 3,4 vs. 7,8). None of the other small molecules significantly affected Par32 phosphorylation status. Thus, phenanthroline leads to activation of the PP2A TOR signaling branch, whereas thiolutin does not. Joint inhibition of Sch9 and PP2A signaling branches by thiolutin is reminiscent of AMPK/Snf1-mediated effects on TORC1 during glucose deprivation stress (Hughes Hallett et al. 2014). In summary, phenanthroline and thiolutin differentially affect PP2A–Tap42 signaling, suggesting a

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different mode of action of these molecules on TORC1 signaling.

Dot6 and Sfp1 transcription factors relocalize in the presence of thiolutin and phenanthroline, consistent with TORC1 signaling inhibition

We next assessed the effect of a 10 min treatment with our small molecules on the localization of transcription factors Sfp1 and Dot6, whose nuclear or cytoplasmic localization is indicative of TORC1 signaling (Fig. 1A). Under optimal growth conditions, the transcription factor Sfp1 is directly phosphorylated by TORC1 and is localized in the nucleus where it facilitates ribosomal protein (RP) and ribosome biogenesis (Ribi) gene transcription (Lempiäinen et al. 2009). Conversely, cytoplasmic localization of RP/Ribi inhibitory transcription factors Dot6 and Tod6 is induced by Sch9 dependent phosphorylation. TORC1 inactivation,
and the resulting dephosphorylation of Sfp1, Dot6, and Tod6 reverses these localization phenotypes (Fig. 1A), causing Dot6 and Tod6-mediated transcriptional silencing of RP/Ribi genes (Huber et al. 2011).

As anticipated, Dot6 was strongly localized in the cytoplasm in our no-treatment control cells, with <6% of cells exhibiting nuclear-localized Dot6. Rapamycin, serving as a TOR-inactivating control, caused 19% of cells to display clear Dot6 nuclear localization within 10 min (Fig. 2A,D). Both phenanthroline (56%) and thiolutin (79%) induced more striking nuclear localization of Dot6 than rapamycin over the same time period (Fig. 2A,D); we hypothesize that additional inhibitory effects of thiolutin and phenanthroline on PKA signaling, which like TORC1 also promotes
cytoplasmic localization of Dot6 and Tod6 (Lippman and Broach 2009; Huber et al. 2011), may explain this result. In contrast, doxycycline and 4tU resembled control cells.

With Sfp1 localization, control, doxycycline and 4tU treated cells all showed >92% nuclear localization (Fig. 2B,E). Interestingly, whereas rapamycin and particularly thiolutin both caused strong cytoplasmic relocalization of Sfp1 (16% and 2% nuclear localization), phenanthroline treatment maintained 79% of cells with nuclear enrichment, albeit the intensity of Sfp1 nuclear signal was generally reduced compared to control cells (Fig. 2B). Inhibition of PKA signaling also promotes Sfp1 cytoplasmic relocalization (Jorgensen et al. 2004; Marion et al. 2004). Thus, if phenanthroline and thiolutin do in fact inhibit PKA signaling (as hypothesized above), an additional variable must explain the poorer phenanthroline-induced Sfp1 cytoplasmic relocalization relative to thiolutin and rapamycin.

In summary, despite thiolutin and phenanthroline exhibiting similar transcription repression effects in yeast (Grigull et al. 2004), there is a nuance to their effects. Thiolutin caused the strongest relocalization of Dot6 and Sfp1 to and from the nucleus, respectively, with phenanthroline also exhibiting a strong nuclear relocalization effect on Dot6 relocalization (more potent than rapamycin). Conversely, rapamycin exhibits a more notable effect on Sfp1 relocalization than phenanthroline. These results are consistent with both thiolutin and phenanthroline impairing TORC1 and PKA signaling, but hint that thiolutin may have a more severe inhibitory effect on RP/Ribi gene transcription.

P-bodies are induced by thiolutin and 4-thiouracil

We next examined how a 10 min treatment with our small molecules affected P-body and stress granule assembly (Buchan 2014). P-bodies and stress granules increase in size and number when the translation is inhibited, owing to a higher proportion of nontranslating mRNAs with which to nucleate them. P-bodies are present at low levels in yeast under normal growth conditions, whereas stress granules typically only form during stress.

In control cells, a modest amount of P-bodies was observed as expected (0.42 foci/cell), whereas thiolutin significantly induced P-bodies (1.80 foci/cell; Fig. 2C,F; Supplemental Table 2). 4tU, despite showing no evidence of impairing stress signaling pathways, also caused a modest but significant increase in P-body number (1.08 foci/cell). The dose tested here (0.2 mM) was used in prior labeling analyses (Munchel et al. 2011), and is fivefold lower than recent work (Chan et al. 2018). This compares to an average of 2.28 P-bodies/cell in cells subject to glucose deprivation, which served as a positive control of P-body induction (Fig. 2F and data not shown). Phenanthroline, rapamycin, and doxycycline did not induce P-bodies. Stress granule induction was not seen in any of our treatment conditions, except for glucose deprived cells, and in a tiny fraction (<1%) of thiolutin-treated cells. In summary, P-bodies are induced by thiolutin and to a lesser but still significant degree by 4tU.

Phenan throline and thiolutin induce HOG signaling

We next examined how our transcription-modifying small molecules affected HOG signaling. Hog1 is a mitogen-activated protein kinase (MAPK), an ortholog of p38 and JNK in human cells, and drives yeast transcriptional responses to hyperosmotic stress (Fig. 3A; Brewster and Gustin 2014).

The effect of our small molecules on Hog1 was assessed using Hog1 and phospho-Hog1 antibodies, to assess overall levels and activation status, respectively. Ten minutes of KCl stress, at a final concentration of 0.4 M, was used as a positive control of Hog1 activation; as expected, we observed the appearance of phosphorylated Hog1 under these conditions (Fig. 3B). Additionally, phenanthroline and thiolutin also induced Hog1 phosphorylation, albeit to a lower extent than 0.4 M KCl stress, which itself is a modest osmotic stress. No effects of any other small molecules examined on Hog1 levels or phosphorylation status were observed. In summary, phenanthroline and thiolutin both induce HOG signaling.

Phenan throline and thiolutin inhibit PKC signaling

Our final focus was on PKC signaling, a MAP Kinase cascade pathway which in yeast senses and maintains cell wall integrity; however additional roles in processes such as autophagy, apoptosis, endocytosis, cytokinesis, and actin cytoskeleton regulation are also known (Heinis ch and Rodicio 2018). Pkc1 activation ultimately leads to phosphorylation, activation and nuclear relocalization of Slt2 in yeast (Fig. 3C). Here, Slt2 phosphorylates various transcription factors involved in cell wall maintenance and other processes.

The effect of our small molecules on PKC signaling was assessed using an antibody recognizing phosphorylated Slt2; we were unable to identify an antibody capable of recognizing nonphosphorylated Slt2 but verified that our small molecules did not affect levels of GFP-tagged Slt2. GAPDH was also used as a loading control. As a positive control, 1 h of zymolase treatment was utilized to perturb cell wall integrity; a clear increase in phosphorylated Slt2 was observed over basal levels under normal growth (Fig. 3D). Interestingly, both phenanthroline and thiolutin reduced phosphorylated Slt2 levels, indicative of inhibition of PKC signaling. No effect on PKC signaling was observed with any other small molecules (Fig. 3D). In summary, phenanthroline and thiolutin both inhibit PKC signaling. The collective findings from our work are summarized in Figure 4.
Summary

Given the perturbing effects of thiolutin and phenanthroline on all signaling pathways assessed, and the potent induction of P-bodies caused by thiolutin, we recommend that neither of these reagents be used to obtain mRNA half-life data. Note, a temperature-sensitive subunit (rpb1-1) of RNA Polymerase II (RNAPII) is also often used in yeast to measure mRNA decay in the absence of transcription (Nonet et al. 1987). However, the rpb1-1 mutation causes a global decrease in transcription rates of 2.7-fold, and a global increase in mRNA stability (a presumed adaptation mechanism) at 30°C (Sun et al. 2012). Shifting to 37°C also induces heat-shock responses in yeast (Morano et al. 2012); hence we chose not to examine rpb1-1 yeast in our work and would suggest such a tool be used with caution.

Doxycycline and 4tU showed no effect on any of the stress signaling pathways examined. β-estradiol also had no effect on Sch9 phosphorylation status. Thus, should a yeast experimentalist's goal be the study of mRNA stability under differing stress conditions, our work suggests that regulatable promoters such as the Tet-off and β-estradiol promoter may represent good options. Finally, the modest induction of P-bodies by 4tU, and prior data on uracil analog effects on growth rate (Chan et al. 2018) and
rRNA processing (Burger et al. 2013) suggest that uracil analog incorporation can perturb RNA metabolic processes. To our knowledge, the effects of such incorporation on translation, secondary structure formation, and RNA modification, all of which may affect cytoplasmic mRNA regulation and thus mRNP granule assembly, remain unknown. A careful assessment of the effects on growth, and minimizing analog labeling concentrations and time periods is likely beneficial for the most accurate mRNA half-life data to be obtained, with the additional benefit that no promoter perturbations are required.

MATERIALS AND METHODS

Yeast strains and growth conditions

Untransformed yeast strains were cultured at 30°C with YPD medium. Strains transformed with plasmids were grown in SD medium with appropriate nutrients for plasmid selection. The transformation was performed by a standard LiAc method (Gietz and Woods 2006). The genetic background of strains used in this study are listed in Supplemental Table 1.

Small molecule modifiers of transcription and TORC1 signaling

The following molecules were incubated in yeast cultures for 10 min at the following concentrations. Rapamycin (TORC1 inhibitor, 200 ng/mL - Fisher Scientific BP2963-1), dissolved in DMSO (Amresco 0231); 1,10 Phenanthroline (RNAP I–III inhibitor, 100 µg/mL - Sigma-Aldrich 131377), in ethanol (Fisher Scientific BP28184); doxycycline (Antibiotic/regulator of Tet-Off/On promoters; 10 µg/mL - Sigma-Aldrich D-9891) in water, β-estradiol (regulator of β-estradiol promoter; 1 µM - Sigma-Aldrich E8875), in ethanol; thiolutin (RNAP I–III inhibitor, 3 µg/mL - WVR 89157-260), in DMSO; 4tU (Uracil analog, 25 µg/mL - Sigma-Aldrich 440736), in DMSO. All drug concentrations are based on standards in the field and/or the following studies (Grigull et al. 2004; Mnaimneh et al. 2004; Munchel et al. 2011). Ten minutes was chosen as a time period for analysis given that mRNA half-lives were recently reported to average 5 min (Chan et al. 2018), thus short-term effects are relevant for accurate half-life measurement.

Band shift assays

Sch9 and Par32 bandshift measurements were performed using a previously described protocol adapted from the Loewith laboratory (Urban et al. 2007; Hughes Hallett et al. 2014), with the following minor alterations. Forty-seven milliliters mid-log cultures harvested and incubated with 3 mL of 100% trichloroacetic acid (TCA) were incubated on ice for between 30 min and 2 h. During cleavage of carboxy-terminally HA-tagged Sch9 samples with 2-nitro-5-thiocyanatobenzoic acid (NTCB; aids separation of phospho-Sch9 fragments for gel analysis), incubation conditions were 12–16 h at 33°C in the dark (1 mM NTCB and 100 mM CHES, pH 10.5). Otherwise, all steps were as previously described (Hughes Hallett et al. 2014). HA (12CA5; Sigma-Aldrich, 11583816001) and C-Myc (Roche, M4439) antibodies were used for Sch9-HA and Par32-Myc detection, respectively (see Supplemental Table 1).

Western blotting

A 50 mL yeast culture was grown at 30°C in a shaking water bath until midlog (OD400, 0.5–0.6), split into 50 mL flasks with 5 mL of culture aliquoted to each, and treated for 10 min (30°C, shaking) with the previously mentioned small molecules, or KCl (0.4 M final concentration) as a phosphy-Hog1 inducing control, or zymolyase (0.26 U/mL final concentration) as a phosphy-Slt2 inducing control (Boorsma et al. 2004). An untreated flask was kept aside as a negative control. One milliliter of culture from each flask/treatment group was harvested, spun down at 13,300 rpm for 2 min. The supernatant was then discarded. Five-hundred microliters of media (YPD) and 0.7 M NaOH was added to the tubes, and then vortexed. Samples were incubated at room temperature for 3 min and then spun down at 5000 rpm for 2 min. Afterward, the supernatant was carefully removed and 50 µL of 2× SDS
sample buffer (4% SDS, 100 mM Tris pH 8.8, 100 mM DTT, 20% glycerol and bromophenol blue) was added to the samples. Samples were mixed with gentle pipetting and then cell extracts were heated to 95°C in SDS sample buffer for 10 min before being run on a 10% SDS-PAGE gel (10% acrylamide, 0.375 M Tris-HCl pH 8.8, 10% Glycerol, 0.1% SDS, 0.1% APS and TEMED). Samples were transferred to a nitrocellulose membrane and detected using Hog1 (D-3) (Santa Cruz Biotechnology; sc-165978), Phospho-p38 [Hog1] MAPK [Cell Signaling Technology, 9211S], or Phospho p44/42 (Stz2) MAPK (D13.14.4E; Cell Signaling Technology - 4370S) antibodies. GFP (Abcam, ab290) and GAPDH (Thermofisher Scientific, MA5-15738) antibodies were used as loading controls for Slt2 westerns. LiCOR secondary antibodies (IRDye 800CW Goat anti-Mouse IgG and IRDye 800CW Goat anti-Rabbit IgG) were used for final imaging. A precision plus dual color protein marker (Page Ruler Stained Protein Ladder, Fisher Scientific- 26617) was used for band size estimates.

Microscopy

Five milliliters of cultures were grown in 50 mL conical flasks shaking at 30°C until midlog phase (OD600 0.4–0.6) and treated with small molecules as described above. Glycein deprived yeast were washed twice in SD media lacking glucose. Cultures were placed back at 30°C to shake for 7 min. Next, 300 µL of culture was added onto an eight well Ibidi chamber slide (Ibidi 80821) treated with Concanavalin A (Cal Biochem - D00157580) to facilitate yeast adhesion, and left at room temperature for 3 min. Culture media in wells was gently aspirated to remove unbound yeast cells and replaced with 150 µL of identical culture media (± small molecule, ± glucose). Cells were then imaged on a DeltaVision Elite system as previously described (Buchan et al. ± small molecule, ± glucose). Cells were then imaged on a DeltaVision Elite system as previously described (Buchan et al. ± small molecule, ± glucose). Cells were then imaged on a DeltaVision Elite system as previously described (Buchan et al. ± small molecule, ± glucose). Cells were then imaged on a DeltaVision Elite system as previously described (Buchan et al. ± small molecule, ± glucose).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

ACKNOWLEDGMENTS

This work was funded in part by a grant to J.R.B. from the National Institute of General Medical Sciences (NIGMS, R01-GM1145664), and a T32 training grant (ST22GM008659-18) awarded to the MCB department at the University of Arizona.

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