Ubiquitin–proteasome-mediated cyclin C degradation promotes cell survival following nitrogen starvation

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ABSTRACT Environmental stress elicits well-orchestrated programs that either restore cellular homeostasis or induce cell death depending on the insult. Nutrient starvation triggers the autophagic pathway that requires the induction of several Autophagy (ATG) genes. Cyclin C–cyclin-dependent kinase (Cdk8) is a component of the RNA polymerase II Mediator complex that predominantly represses the transcription of stress-responsive genes in yeast. To relieve this repression following oxidative stress, cyclin C translocates to the mitochondria where it induces organelle fragmentation and promotes cell death prior to its destruction by the ubiquitin–proteasome system (UPS). Here we report that cyclin C–Cdk8, together with the Ume6-Rpd3 histone deacetylase complex, represses the essential autophagy gene ATG8. Similar to oxidative stress, cyclin C is destroyed by the UPS following nitrogen starvation. Removing this repression is important as deleting CNC1 allows enhanced cell growth under mild starvation. However, unlike oxidative stress, cyclin C is destroyed prior to its cytoplasmic translocation. This is important as targeting cyclin C to the mitochondria induces both mitochondrial fragmentation and cell death following nitrogen starvation. These results indicate that cyclin C destruction pathways are fine tuned depending on the stress and that its terminal subcellular address influences the decision between initiating cell death or cell survival pathways.

INTRODUCTION

In response to environmental cues, cells activate well-orchestrated processes to either restore cellular homeostasis or commit to cell death programs. Incorrectly deciding between these cell fates is detrimental to the cell and linked to the etiology of many diseases. The cellular response to adverse environmental cues can be divided into three stages. First, the stress damage is recognized and a signal transduced to the nucleus. Second, transcription programs are altered to repress progrowth programs while inducing genes necessary for detoxification and damage repair. Finally, the decision made in the nucleus has to be communicated to the organelles to provide a unified cellular response. Importantly, mitochondria are a key regulatory node for proper response to cellular damage with stress-induced fragmentation being an initial step in mitochon-drion-dependent cell death pathways (Tait and Green, 2013; Kasahara and Scorrano, 2014).

The budding yeast Saccharomyces cerevisiae executes a regulated cell death (RCD) response to various stresses (oxidative stress, acetic acid, fungicides) (Madeo et al., 1999; Ludovico et al., 2002; Wissing et al., 2004; Buttner et al., 2007; Pereira et al., 2007) that is coupled with a fragmented mitochondrial phenotype (Braun and Westermann, 2011; Cooper et al., 2014). For example, although excessive hydrogen peroxide (H$_2$O$_2$) exposure initially induces both prosurvival (e.g., chaperones) and antioxidant (e.g., catalase) encoding genes (Levin, 2005), RCD inevitably ensues (Madeo et al., 1999). In contrast, nitrogen limitation induces macroautophagy pathways (hereafter referred to as autophagy) that enhance recycling of
findings that the Rpd3–Sin3–Ume6–Ume1 complex represses early meiotic genes whose transcription is also inhibited in nutrient-rich conditions (Vidal et al., 1991; Strich et al., 1994; Mallory and Strich, 2003). How repression is relieved upon nitrogen depletion is less well understood. For example, Rim15, the great wall kinase, induces ATG8 transcription (Bartholomew et al., 2012) but it remains unknown if this kinase works by inhibiting Ume6 or other HDAC members or by other mechanisms.

In addition to this HDAC complex, another repressor system was discovered that represses early meiotic genes (Strich et al., 1989; Cooper et al., 1997). This system is composed of the conserved Cdk8 kinase module (CKM) of the mediator complex (see Figure 1A). The CKM is composed of cyclin C, Cdk8, Med12, and Med13 and associates with the RNA polymerase II mediator complex and predominantly represses genes induced by environmental stress (Cooper et al., 1997; Holstege et al., 1998; Bjorklund and Gustafsson, 2005; van de Peppel et al., 2005; Allen and Taatjes, 2015) and entry into meiosis (Cooper et al., 1997; Strich et al., 1989; Surosky et al., 1994). In response to stress, cyclin C repression is relieved by its translocation to the cytoplasm (Cooper et al., 2014) following the destruction of its nuclear anchor Med13 by the SCF^cIts^ ubiquitin (Ub) ligase-mediated Ub–proteasome system (UPS) (Khakhina et al., 2014; Stieg et al., 2018). On translocation to the cytoplasm, cyclin C associates with mitochondria by directly binding to the fission machinery and inducing stress-mediated mitochondrial fragmentation (Cooper et al., 2014; Strich and Cooper, 2014; Ganesan et al., 2019; Jezek et al., 2019b). In addition, cyclin C is required for promoting RCD (Krasley et al., 2006; Cooper et al., 2014). Finally, cyclin C is destroyed following mitochondrial fragmentation by the UPS (Cooper et al., 1997, 2012), which attenuates the stress response. Taken together, these studies revealed that cyclin C connects the nuclear and mitochondrial oxidative stress response through its transcriptional repressor and mitochondrial morphology regulatory roles.

These studies are consistent with the emerging theme that proteins can have two very different functions, coined “day and night jobs” (Shamas-Din et al., 2013; Gross and Katz, 2017), which can be induced by different external or intrinsic stimuli. Importantly, our studies have shown that both the day and night jobs of cyclin C are conserved (Wang et al., 2015; Stieg et al., 2019). More recently, we have shown that in mammalian cells, different domains of cyclin C directly interact with the mitochondrial fission GTPase Drp1 to direct stress-induced mitochondrial fission (Ganesan et al., 2019). The cyclin C–Bax interaction is required for normal Bax activation and its efficient mitochondrial localization (Jezek et al., 2019a). Further studies in both yeast and mammals revealed that the aberrant release of cyclin C does not kill cells but does make them hypersensitive to oxidative or chemotherapeutic damage.

**FIGURE 1:** The UPS is required for cyclin C destruction following nitrogen starvation. (A) Model showing the two roles coined “day job” and “night job” that cyclin C plays to regulate the oxidative stress response (see text for details and reviewed in Jezek et al., 2019b; Strich and Cooper, 2014). Importantly, the night job is independent of Cdk8 and both roles are conserved (Wang et al., 2015). (B) Western blot analyses of extracts prepared from mid-log cultures with the indicated genotypes expressing cyclin C-myc resuspended nitrogen starvation medium (SD-N) for the indicated times. (C) Degradation kinetics of cyclin C signals obtained in B. Error bars indicate SD, N = 3. (D) Western blot analyses of extracts prepared from a wild-type mid-log culture were treated with 200 ng/ml rapamycin for the indicated times. Pgk1 levels were used as a loading control for all Western blot studies.
(Khakhina et al., 2014; Jezek et al., 2019a). Taken together, these results demonstrate that cyclin C translocation to the mitochondria is an early step in the RCD pathway (Jezek et al., 2019a). As anticipated from these results, both yeast and mammalian cell lines devoid of cyclin C are less sensitive to stress (Cooper et al., 2014; Wang et al., 2015) and acts as a tumor suppressor in both solid and hematological cancers (Li et al., 2014; Jezek et al., 2019c).

Given cyclin C’s bipartite role following cell death stimuli, we investigated whether it has a regulatory role when cells induce the autophagic survival response following nitrogen starvation. Similar to oxidative stress, we discovered that cyclin C represses the transcription of a gene (ATG8) required for autophagy. In addition, genetic studies revealed that cyclin C-Cdk8 function in the same pathway as the Ume6-Rpd3 HDAC system. Cyclin C-Cdk8-dependent repression in nitrogen-starved cells is also relieved by cyclin C proteolysis via the UPS. However, several differences from oxidatively stressed cells were observed. First, no detectable cytoplasmic cyclin C was seen and the mitochondria remained reticular. This difference is important as targeting cyclin C to the mitochondria changed nitrogen starvation from evoking a survival pathway to one inducing cell death. These findings reveal a complex system in which the subcellular address of cyclin C plays a key role in cell fate decisions by regulating gene expression and mitochondrial-dependent RCD.

RESULTS
Cyclin C degradation following nitrogen starvation requires 26S proteasome activity

We previously reported that cyclin C relocalization and subsequent destruction play key roles in directing cell fate following oxidative stress (Cooper et al., 2012, 2014; Jin et al., 2014a). Therefore, we asked whether cyclin C is similarly regulated upon nitrogen starvation. Exponentially growing wild-type cells expressing a functional cyclin C-myc reporter (Cooper et al., 1997) were starved for nitrogen (SD-N) and cyclin C levels were monitored by Western blot analysis. The results showed that cyclin C levels were reduced following nitrogen starvation with an apparent half-life of 45 min (Figure 1B, quantified in Figure 1C, half-life in Supplemental Figure S1A). Rapamycin also inhibits TORC1 (Cardenas et al., 1999) by a mechanism not physiologically identical to nitrogen starvation (Tate and Cooper, 2013). Therefore, we repeated these experiments in replete medium containing rapamycin at 200 ng/ml. Western blot analysis produced results similar to those obtained with nitrogen starvation (Figure 1D, quantified in Supplemental Figure S1A) with the decay rate being slightly slower (half-life of 75 min). The same rate of decay was also obtained when 50 ng/ml rapamycin was used (Supplemental Figure S1, A and B). Treatment with rapamycin, but not its solvent (Tween/ethanol), also triggered cyclin C destruction when tagged with YFP (Supplemental Figure S1C). We previously demonstrated that CNC1 mRNA levels remain constant following 4 h nitrogen starvation (Cooper et al., 1999) and found that the half-life of cyclin C is ~3 h in unstarved cultures using cycloheximide translation inhibition assays (Supplemental Figure S1, D and E). Taken together, these results suggest that cyclin C is targeted for degradation following nitrogen starvation or rapamycin stress. Next, we addressed whether cyclin C destruction required either the vacuolar or the UPS pathways. First, cyclin C-myc levels were monitored in nitrogen-starved cultures harboring mutations in two major vacuolar proteases, Pep4 and Prb1 (Takeshige et al., 1992; Van Den Hazel et al., 1996). The results showed that cyclin C was degraded with similar kinetics to wild type (Figure 1B, quantified in Figure 1C). In contrast, repeating this experiment in a mutant deficient for normal 20S proteasome assembly (ump1Δ) (Ramos et al., 1998; Li et al., 2007; Czabotar et al., 2013; Uekusa et al., 2014), significantly stabilized cyclin C (Figure 1B, quantified in Figure 1C). These results indicate that the UPS degrades cyclin C in response to nitrogen starvation.

Cyclin C degradation following nitrogen starvation requires 19S proteasome activity

It has been reported that the 20S proteasome can degrade ubiquitylated proteins independent of 19S function (Wang et al., 2010). To address if 19S function is required for cyclin C degradation in response to nitrogen starvation, a sem1Δ mutant was used. Sem1 is required for efficient lid assembly, catalyzing the incorporation of subunits Rpn3 and Rpn7 into the 19S regulatory complex (Jantti et al., 1999; Tomko and Hochstrasser, 2014). Cyclin C was significantly stabilized in sem1Δ compared to wild-type cells (Figure 1B, quantified in Figure 1C). Taken together, these results confirm the above conclusions that the UPS is required for cyclin C degradation and that the cap region is required for this activity.

Cyclin C destruction following nitrogen starvation uses the same E2s but different E3 enzymes to oxidative stress

UPS-mediated destruction is executed by proteins being tagged with Ub chains by a sophisticated three-step enzymatic cascade utilizing E1 Ub-activating, E2 Ub-conjugating and a variety of E3 Ub-ligating enzymes (Pickart, 2001; Tsujiya et al., 2017). Ub chains are in turn recognized by intrinsic or extrinsic (or a combination of both) receptor proteins that deliver and/or capture tagged proteins to the 19S cap of the proteasome (Schauber et al., 1998; Hartmann-Petersen et al., 2003; Verma et al., 2004; Lander et al., 2012; Rosenzweig et al., 2012; Shi et al., 2016). As H2O2-mediated destruction of cyclin C requires the action of the functionally redundant UBC4 and UBC5 E2 enzymes (Cooper et al., 1999, 2012), we investigated whether they performed the same role in nitrogen-starved cells. Cyclin C was significantly stabilized in ubc4Δ ubc5Δ cells compared with the isogenic wild type under these conditions (Figure 2A, quantified in Figure 2B). Next, the role of Ub receptor proteins in cyclin C proteolysis was tested. Cyclin C was still subjected to at least partial proteolysis in strains harboring single, double, and triple deletions in the extrinsic receptor proteins Ddi1, Dsk2, and Rad23 (Supplemental Figure S2A). Likewise, partial proteolysis was observed in strains harboring single and double deletions in Rpn10 and Rpn13, the intrinsic receptor proteins (Supplemental Figure S2B). Intriguingly, compared with its isogenic wild type, partial proteolysis of cyclin C was also observed in the intrinsic receptor triple mutant strain (Figure 2C) in which Rpn1, Rpn10, and Rpn13 each harbor mutations that prevent Ub binding (Shi et al., 2016). However, cyclin C was significantly stabilized in the “quintΔ” receptor mutant strain (Figure 2C, quantified in Figure 2B), in which the only functional Ub receptor is Rpn1 (Shi et al., 2016). Therefore, similar to other UPS substrates (Zhang et al., 2009; Shi et al., 2016; Saeki, 2017), the extrinsic and intrinsic Ub receptors are functionally redundant with regard to cyclin C degradation following nitrogen starvation. A caveat to this interpretation is that Sem1 is a potential Ub receptor, having such a role in fission yeast. However, here Sem1 does not recognize Ub when assembled in the complete proteasome (Paraskevopoulos et al., 2014).

We next asked if Not4, the E3 ligase that mediates cyclin C destruction following oxidative stress (Cooper et al., 2012, 2014) also plays this role in nitrogen-starved cells. However, cyclin C was still destroyed after 4 h nitrogen depletion in not4Δ cells (Figure 2E) indicating that a different E3 ligase fulfills this role. As Ubc4/5 can interact with both HECT domain and RING E3 ligases (Stoll et al., 2011), we decided to hunt for the E3 ligase mediating cyclin C degradation.
by initially testing candidate genes. Rsp5 was first tested as it both interacts with RNA polymerase II and is involved in a variety of stress responses (Huibregtse et al., 1997; Wang et al., 1999; Hiraishi et al., 2009). A doxycycline-inducible N-end rule approach (Bachmair et al., 1997; Wang et al., 1999; Hiraishi et al., 2009) forbidding the use of Ub receptor mutant (YSS781a, dsk2Δ, rad23Δ, ddi1Δ rpn13-pru rpn10-uum) and the triple intrinsic receptor mutant (YSS786a, rpn13-pru rpn10-uum rpn1-ARR). (D) Quantification of the results obtained in C. N = 2. (E) Cyclin C-YFP following wild blot analysis following nitrogen starvation in wild type (MHY414), ubc4/5Δ (MHY508), and not4Δ. (F) The Rsp5-HA strain (RSY2301) harboring the Tet operator plasmid (pCM1888) and cyclin C-myc were grown to mid-log phase and a sample removed for Western analysis to visualize Rsp5-HA (far right, top panel). The remaining culture was treated with doxycycline for 5 h before being subjected to nitrogen starvation. Thereafter, cyclin C-myc was monitored by Western blot analysis. For all blots, Pgk1 levels were used as loading controls.

FIGURE 2: The UPS is required for cyclin C degradation following nitrogen starvation. (A) Western blot analyses of extracts prepared from mid-log ubc4/5Δ (MHY508) and wild-type (MHY414) cultures expressing cyclin C-myc resuspended in nitrogen starvation medium (SD-N) for the indicated times. (B) Quantification of the results obtained in A. N = 3. (C) As in A except that cyclin C levels were monitored in wild type (SUB62), the “quant、“Ub receptor mutant (YSS781a, dsk2Δ, rad23Δ, ddi1Δ rpn13-pru rpn10-uum), and the triple intrinsic receptor mutant (YSS786a, rpn13-pru rpn10-uum rpn1-ARR). (D) Quantification of the results obtained in C. N = 2. (E) Cyclin C-YFP was monitored by Western blot analysis following nitrogen starvation in wild type (MHY414), ubc4/5Δ (MHY508), and not4Δ. (F) The Rsp5-HA strain (RSY2301) harboring the Tet operator plasmid (pCM1888) and cyclin C-myc were grown to mid-log phase and a sample removed for Western analysis to visualize Rsp5-HA (far right, top panel). The remaining culture was treated with doxycycline for 5 h before being subjected to nitrogen starvation. Thereafter, cyclin C-myc was monitored by Western blot analysis. For all blots, Pgk1 levels were used as loading controls.

Cyclin C does not translocate to the mitochondria following nitrogen starvation

Cyclin C is nuclear during normal growth conditions but forms mitochondrial-associated foci required for organellar fragmentation in response to oxidative stress (Cooper et al., 2014; Jin et al., 2014a) (see Figure 3A, top and middle panels). Thereafter cyclin C is destroyed by the UPS in the cytoplasm (Cooper et al., 1986, 1997, 2012). To determine whether cyclin C responds similarly following TORC1 inhibition, subcellular localization of cyclin C-YFP and mitochondrial morphology were monitored by fluorescence microscopy following 200 ng/ml rapamycin treatment. These experiments revealed that cyclin C-YFP was below the limits of detection following rapamycin treatment using standard exposure times (Figure 3A, bottom panel). No observable cyclin C-YFP cytoplasmic presence was observed even when longer exposures were used in cells harboring an NLS-NAB-mCherry fusion protein (Malinovska et al., 2012) to mark the nucleus (Supplemental Figure S3, A and B). Likewise, no observable cyclin C-YFP cytoplasmic presence was observed even at earlier timepoints in cells harboring a Nup49-mCherry fusion protein that marks the nuclear pore complex (NPC; Bucci and Wente, 1997) (Figure 3B). Last, consistent with the model that vacuolar hydrolysis is not needed to degrade cyclin C, we did not observe cyclin C-YFP in vacuoles marked by Vph1-mCherry in WT (Figure 3C) or pep4Δ prb1Δ-1 strains (Figure 3D). Likewise, cyclin C-YFP was still detectable in ump1Δ cells after rapamycin treatment with the caveat that we also observed some unexplained foci that were predominantly nuclear in both unstressed and stressed cells (Supplemental Figure S3D).

Consistent with cyclin C not having an observable cytoplasmic presence, we observed that the mitochondria remain predominantly reticular following 200 ng/ml rapamycin stress (Figure 3, A and C). These findings match early reports in mammalian cells that mitochondrial fragmentation is infrequent during starvation (Gomes et al., 2011; Rambold et al., 2011a,b). As nitrogen starvation triggers a slightly different stress response (Tate and Cooper, 2013), we next examined cyclin C-YFP localization and mitochondrial morphology during nitrogen depletions. Figure 4A shows that identical to rapamycin treatment, cyclin C-YFP was below the limits of detection after 3 h in nitrogen starvation media. Furthermore, these cells also harbored Nup49-mCherry and no observable cytoplasmic presence was detected at earlier timepoints (1 and 2 h). Importantly, even after longer exposure times (4 h) to nitrogen starvation media, the mitochondria remained predominantly reticular (Figure 4B) and cyclin C was only detectable on overexposure (Supplemental Figure S3D). Taken together, these data indicate that similar to oxidative stress, cyclin C is destroyed by the UPS following TORC1

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FIGURE 3: Cyclin C does not induce mitochondrial fragmentation following rapamycin stress. (A) Wild-type cells (RSY10) harboring expression plasmids for Mito-TFP (mitochondrial marker) and cyclin C-YFP were untreated (top panels) treated with 0.8 mM H$_2$O$_2$ (middle panel) or rapamycin (200 ng/ml) for the times indicated. Merged panels of cyclin C-YFP and Mito-TFP are shown. The stippled box indicates enlarged panels. (B) Wild-type cells harboring cyclin C-YFP and Nup49-mCherry grown to mid-log were treated with rapamycin (200 ng/ml) as indicated. Merged and enlarged panels are indicated. (C) As in A except that the cells also expressed Vph1-mCherry to mark the vacuoles were then treated with either 200 ng/ml or 2.5 ng/ml rapamycin for 3 h. (D) Mutant pep4Δ prb1Δ-1 (BJ5459) cells harboring cyclin C-YFP and Vph1-mCherry were treated with 200 ng/ml rapamycin for 4 h. Representative fluorescence microscopy images are shown.
FIGURE 4: Cyclin C does not induce mitochondrial fragmentation following rapamycin stress. (A) Wild-type cells (RSY10) harboring expression plasmids for cyclin C-YFP and Nup49-mCherry were washed and resuspended in SD-N. Representative fluorescence microscopy images of the results are shown. (B) As in A except that the cells also express Mt-Ds-Red to mark the mitochondria. For all panels, Nom. indicates Nomarski imaging. (C) Degradation kinetics of cyclin C after exposure to 1.2 and 0.4 mM H$_2$O$_2$ or SD-N as indicated. The data were obtained from previously published experiments (Jin et al., 2015) and Figure 1C. (D) Mitochondrial morphology of cells 2 h after the stress. The data for the H$_2$O$_2$ experiments were obtained from (Jin et al., 2015). (E) cnc1Δ cells (RSY1696) harboring a wild (pKC337) or cyclin C$^{A110V}$-myc expression plasmid (pKC354) were grown to mid-log phase and then washed and resuspended in nitrogen starvation medium (SD-N) for the indicated times. Cyclin C-myc levels were monitored by Western blot analysis. Pgp1 levels were used as a loading control.
inhibition triggered by either rapamycin treatment or nitrogen starvation. However, an important difference between the two stressors is that cyclin C destruction occurs prior to an observable cytoplasmic presence or activation of the mitochondrial fission machinery in response to starvation (see Discussion). Although the degradation kinetics of cyclin C during H2O2 stress (Jin et al., 2014a, 2015) and nitrogen starvation are similar (compared in Figure 4C), a significant increase in mitochondrial fragmentation is only observed following H2O2 stress (Jin et al., 2015) (compared with Figure 4D). Moreover, previous studies demonstrated that mitochondrial fragmentation was not observed with a cis mutant (CNC1A110V) that does not permit nuclear translocation of cyclin C after H2O2 stress (Cooper et al., 2014; Jin et al., 2014a). However, this mutant did not stabilize cyclin C following exposure to nitrogen starvation, indicating that a different signaling system likely triggers cyclin C destruction (Figure 4E). Taken together, these results support the model that nitrogen starvation stimulates cyclin C degradation prior to its detectable presence in the cytoplasm. Consistent with this model, the mitochondria do not exhibit significant stress-induced fragmentation.

Cyclin C-Cdk8 negatively regulates ATG8 expression within the Ume6-Rpd3 HDAC pathway

Our previous studies identified cyclin C as a repressor of both meiotic and stress response genes (SRG; Cooper et al., 1997, 2012). In yeast, ATG8 is induced shortly following autophagy stimulation (Kirisako et al., 1999) and encodes a protein that is incorporated into the phagophore membrane to regulate both autophagosome size as well as the binding of selective cargo receptor proteins (Ichimura et al., 2000; Xie et al., 2008). To determine if cyclin C-Cdk8 represses ATG8 transcription, RT-qPCR analysis was performed on total RNA samples prepared from unstressed yeast cnc1Δ and cdk8Δ strains. These studies revealed that ATG8 mRNA levels were increased ~two-fold compared with the wild-type control (Figure 5A). Induced levels following 1 h rapamycin treatment were similar in wild-type and mutants cells, indicating that loss of cyclin C-Cdk8 activity was not additive to the normal induction pathway (Figure 5A). GFP-Atg8 protein levels also mirrored these results (Figure 5B, example blot in Supplemental Figure S4A). Furthermore, GFP-Atg8 levels were similarly upregulated in the unstressed cdk8 kinase dead strain (cdk8S290A) (Surosky et al., 1994) (Figure 5C, compare 0 h time-points). Taken together, these results indicate that cyclin C-Cdk8 kinase activity represses ATG8 transcription and are consistent with the model that cyclin C destruction is a mechanism to relieve this repression.

To test whether cyclin C-Cdk8 is part of the previously established Ume6-Rpd3 HDAC repression pathway, a series of epistasis experiments was performed. As previously reported (Backues et al., 2012; Bartholomew et al., 2012), RT-qPCR analysis revealed that ATG8 mRNA levels were elevated ~fourfold in the ume6Δ mutant compared with wild type (compare Figure 4D to Figure 4A).

**FIGURE 5:** Cyclin C-Cdk8 negatively regulates ATG8 mRNA expression. (A) RT-qPCR assays probing for ATG8 mRNA expression in the mutant shown before and after 1 h 200 ng/ml rapamycin treatment. Transcript levels are given relative to the internal ACT1 mRNA control. (B) Fold increase in GFP-Atg8 levels in strains shown before and after 6 h 200 ng/ml rapamycin treatment. (C) GFP-Atg8 cleavage assays before and after 200 ng/ml rapamycin in wild-type (YC7) and cdk8 kinase dead (YC17) strains. The asterisk indicates a previously reported background band (Huang et al., 2014). (D) As in A for the ume6Δ (RSY431) and ume6 cdk8Δ (RSY2128) strains as indicated. (E, F) RT-qPCR assays probing for ATG7 and ATG14 mRNA expression respectively in wild-type (RSY10) and cdk8Δ (RSY1726) unstressed cells. Transcript levels are given relative to the internal ACT1 mRNA control. For all RT-qPCR assays, the error bars indicate the SD from the mean of two technical replicates from three independent cultures. ***p < 0.001 and NS represents no significance.
However, no increase in ATG8 mRNA levels was observed in the um6Δ cdk8Δ double mutant compared with um6Δ alone. In addition, rapamycin treatment induced a similar increase in ATG8 mRNA levels in um6Δ and um6Δ cdk8Δ double mutants (Figure 5D). Taken together, these data argue that cyclin C-Cdk8 acts within the Ume6-Rpd3 HDAC pathway to repress ATG8 transcription. Consistent with this, the mRNA levels of two genes ATG7 or ATG14, which are used to follow autophagy induction and nuclear translocation, respectively (Bernard et al., 2015), but not repressed by Ume6 (Jin et al., 2014b), were also not changed in unstressed cdk8Δ cells compared with wild type (Figure 5, E and F). Although this study is not exhaustive, it suggests that Ume6-Rpd3 HDAC and the CKM work together by an unknown mechanism to repress ATG8 transcription.

Cyclin C-Cdk8 repression restricts cell growth in response to partial TORC1 inhibition

Our epistasis experiments place cyclin C-Cdk8 and Ume6 in the same pathway with regard to ATG8 repression in replete media. Next, we addressed whether deleting CDK8 increased autophagic activity as observed in um6Δ mutants (Bartholomew et al., 2012). One method, termed autophagic flux, is based on the finding that GFP is protected from rapid degradation in the vacuole compared with its fusion partner. Therefore, comparing the amount of free GFP cleaved from the non-vacuolar GFP-Atg8 can be used as a measure of autophagic activity (Suzuki et al., 2001; Loos et al., 2014; Delorme-Axford et al., 2015). Quantifying the results described in Figure 5B and Supplemental Figure S4A revealed no significant difference in autophagic flux among wild-type, cdk8Δ, and cnc1Δ strains (Figure 6A). Another established method of assessing earlier autophagic activity is to determine the amount of substrate being processed through the system as a function of autophagosome formation (Torggler et al., 2017). Using GFP-Atg8 foci as a measure of autophagosome production, we found a significant increase in foci in unstressed cdk8Δ and cnc1Δ yeast strains compared with wild type (Figure 6B). Taken together, these results suggest that although GFP-Atg8 foci formation increases in cdk8Δ and cnc1Δ, deletion of the CKM is not sufficient to activate autophagic pathways in the absence of stress.

To further explore the physiological role of cyclin C destruction in the response to nitrogen starvation, we asked whether loss of this repressor enhanced cell survival under nitrogen-limiting conditions. One model consistent with the results from Figure 6, A and B is that the increased number of GFP-Atg8 foci seen in cnc1Δ or cdk8Δ cells primes them to better survive once autophagy pathways are initiated. Most protocols examining autophagic induction use either total nitrogen starvation or high rapamycin concentrations (e.g., 200 ng/ml) which drives cells into quiescence (Gray et al., 2004; Axford et al., 2014). However, more physiological conditions are found when cells are grown to mid log in SD-complete medium and 10-fold dilutions plated on SD-complete medium containing 0, 200, or 2.5 ng/ml rapamycin. (D) Identical exposures of GFP-Atg8 cleavage assays in wild-type and cdk8Δ cells after treatment with 2.5 ng/ml rapamycin for the timepoints indicated. The asterisk indicates a previously reported background band (Huang et al., 2014). (E) Wild-type cells (RSY10) harboring expression plasmids for cyclin C-YFP, Vph1-mCherry, and MT.

**FIGURE 6:** Deletion of either CDK8 or CNC1 promotes cell survival following low-dose rapamycin treatment. (A) Quantification of GFP-Atg8 flux depicted in Supplemental Figure S4A for wild-type (RSY10), cdk8Δ (RSY1726) and cnc1Δ (RSY1696) cultures. N = 3. (B) Representative Nomarski and fluorescence microscopy images of GFP-Atg8 foci are shown for unstressed wild-type, cnc1Δ, and cdk8Δ cells as in A. Quantification of GFP-Atg8 foci depicted as a percentage of population displaying 0, 1, 2, or 3 foci (percentage of mean ± SD) grown under unstressed conditions. N = 3, for all assays: **p < 0.01, ***p < 0.001, ****p < 0.0001. (C) Wild-type, cdk8Δ, and cnc1Δ cells were grown to mid log in SD-complete medium and 10-fold dilutions plated on SD-complete medium containing 0, 200, or 2.5 ng/ml rapamycin. (D) Identical exposures of GFP-Atg8 cleavage assays in wild-type and cdk8Δ cells after treatment with 2.5 ng/ml rapamycin for the timepoints indicated. The asterisk indicates a previously reported background band (Huang et al., 2014). (E) Wild-type cells (RSY10) harboring expression plasmids for cyclin C-YFP, Vph1-mCherry, and MT. CFP were washed and resuspended in 2.5 ng/ml rapamycin for 3 h. Representative fluorescence microscopy images of the results are shown. Arrows point to detectable cyclin C-YFP after 3 h rapamycin stress. (F) Doubling times of wild type (RSY10), cnc1Δ (RSY1696) and atg8Δ (RSY2144) cells in YPDA with and without 2.5 ng/ml rapamycin. N = 3. (E) For all blots, Pgk1 was used as a loading control.
As anticipated from these studies, wild-type, cnc1\(\Delta\), and cdk8\(\Delta\) cells were unable to grow on plates containing 200 ng/ml rapamycin (Figure 6C, middle panel). We thus sought to find the lowest rapamycin concentration that still evoked an autophagic response. We found that 2.5 ng/ml rapamycin induced autophagy using the GFP-Atg8 cleavage assay with kinetics similar to either higher drug concentrations or nitrogen starvation (Figure 6D). Fluorescence microscopy and Western blot analysis showed that cyclin C is also degraded in wild-type cells under these conditions though not to the extent seen at cell arresting concentrations (200 ng/ml—compare Figure 6E to Figure 3C and see Supplemental Figure S1C). Survival assays on medium containing 2.5 ng/ml rapamycin revealed that the plating efficiency increased in cnc1\(\Delta\) and cdk8\(\Delta\) mutants compared with wild-type controls (Figure 6C, bottom panel). Moreover, a growth rate advantage was also observed in the liquid cnc1\(\Delta\) cultures grown in 2.5 ng/ml rapamycin (doubling time of 2.3 h) compared with wild type grown in 2.5 ng/ml rapamycin (doubling time of 7.2 h, Figure 6F). These findings were most likely due to elevated autophagic activity as atg8\(\Delta\) mutants displayed a significant growth reduction (Figure 6F). These experiments indicate that 2.5 ng/ml rapamycin is sufficient to reduce TORC1 activity to a level that induces autophagy but still permits cell growth, suggesting that this amount of drug triggers a nutrient-limiting, but not eliminated, signal. These experiments also indicate that precocious derepression of the autophagic pathway through CNC1 or CDK8 deletion provides cells with a growth advantage under these partial starvation conditions. However, aberrant up-regulation of ATG8 does not accelerate the timing of autophagic induction as seen by the similar appearance of cleaved GFP-Atg8 in both wild type and the cdk8\(\Delta\) mutant (Figure 6D). Rather, consistent with the GFP-Atg8 foci formation assay (Figure 6A) once autophagy is initiated, the amount of cleaved product increases in the mutant arguing that more substrate is moving through the pathway.

**Autophagy induction protects cells from cyclin C-induced mitochondrial fission and cell death following H\(\text{2O}_2\) stress**

In controlling the cellular response to oxidative stress, yeast cyclin C has two roles (see Figure 1A). First, along with Cdk8, it represses SRGs in the absence of cell damage. Second, in response to stress, cyclin C performs its Cdk8-independent job by promoting stress-induced mitochondrial fission and RCD. These studies also show that on reaching the cytoplasm, cyclin C mediates mitochondrial fission, even in the absence of stress. This is illustrated by our studies in mammalian cells where *Escherichia coli*-purified yeast GST-cyclin C, but not GST alone, induces mitochondrial fission when added to digitonin-treated mouse embryonic fibroblasts (Wang et al., 2015). In response to nitrogen starvation, cyclin C is still destroyed by the UPS, thereby relieving the repression of ATG8. However, our data indicate that cyclin C destruction prevents cytoplasmic accumulation, thus preventing mitochondrial fission following nitrogen starvation. Therefore, one model consistent with these results is that cyclin C proteolysis protects mitochondrial integrity and increases cell survival. Congruous with this hypothesis, amino acid or nitrogen starvation in mammalian cells induces a hyperperfused mitochondrial network that promotes cell survival (Rambold et al., 2011b; Gomes and Scorrono, 2013; Morita et al., 2017). To test this model in yeast, wild-type cells were treated with 50 ng/ml rapamycin that induces quiescence similar to the 200 ng/ml concentration described above and destroys cyclin C with similar kinetics (Supplemental Figure S1, A and B). Cyclin C is also degraded using this concentration of rapamycin (Supplemental Figure S1, A and B). Following 0, 2, 3, or 4 h, the cells were subsequently treated with 0.8 mM H\(\text{2O}_2\) for an additional hour. Thereafter, the percentage of cells exhibiting mitochondrial fission and two or more cyclin C-YFP cytoplasmic foci was quantified. The results show that as previously reported (Cooper et al., 2014; Jin et al., 2014a), the addition of H\(\text{2O}_2\) induces cyclin C nuclear translocation and mitochondrial fragmentation (Figure 7A). As anticipated from results presented in Figures 3 and 4, the mitochondria remained reticular even after 4 h of rapamycin stress and cyclin C-YFP was below the limits of detection (Figure 7B). However, increasing the rapamycin exposure time before adding the H\(\text{2O}_2\) reduced both H\(\text{2O}_2\)-induced mitochondrial fission and cyclin C-YFP cytoplasmic foci (Figure 7A, quantified in C and D). The addition of H\(\text{2O}_2\) after rapamycin treatment did not affect the appearance of free GFP from GFP-Atg8 in either wild-type or cdk8\(\Delta\) cells (Figure 7E, right panel). Furthermore, similar to a previous report (Perez-Perez et al., 2014), we observed that H\(\text{2O}_2\) stress induces GFP-Atg8 cleavage (Figure 7E, middle panel). This is dependent on autophagy as no cleavage occurs in atg1\(\Delta\) or atg1\(\Delta\) cnc1\(\Delta\) cells (Supplemental Figure S4B). Autophagy, however, is not required for the cytoplasmic translocation of cyclin C following H\(\text{2O}_2\) stress and the mitochondria are able to execute fission in atg1\(\Delta\) cells (Supplemental Figure S4C). Taken together, these results suggest that once autophagy is established, cells are prevented from executing mitochondrial fragmentation. These results are consistent with the model that cyclin C destruction prior to cytoplasmic translocation prevents mitochondrial fragmentation.

To further test the model that rapamycin induced autophagy observed in Figure 7E protects cells from H\(\text{2O}_2\)-induced cell death we executed two additional assays. Thereafter either 2 mM H\(\text{2O}_2\) or water was added to the rapamycin treated cells for an additional 2 h. Cell viability was then monitored by plating serial dilutions of the cells on rich medium (YPDA). The results clearly show that pretreatment of the cells with 50 ng/ml rapamycin protects them from 2 mM H\(\text{2O}_2\)-mediated cell death. The experiment was repeated by measuring live cells by fluorescein diacetate (FDA) staining and fluorescence activated cell (FACs) analysis. Even after longer exposure times to H\(\text{2O}_2\) (20 h) 4 h pretreatment in rapamycin was protective of cell death with 80% of the H\(\text{2O}_2\)-treated cells dying compared with 10% in the pretreated group. These results clearly show that rapamycin protects cells from H\(\text{2O}_2\)-mediated cell death.

**Mitochondrial cyclin C localization triggers cell death following autophagy induction**

The experiments described above are consistent with a model that preventing cyclin C–mitochondrial interaction increases cell survival in response to starvation conditions. Previously, we have shown that deleting MED13, the gene encoding the nuclear anchor for cyclin C, allows constitutive cyclin C cytoplasmic localization and mitochondrial fragmentation (Khakhina et al., 2014). Importantly, these results, as well as those obtained in mammalian cells (Jezek et al., 2019a), showed that simply placing cyclin C at the mitochondria does not induce RCD in the absence of stress but does make the cells hypersensitive to cell damage. In other words, yeast and mammalian cells undergo RCD more efficiently when cyclin C is located at the mitochondria before the stress. To test the possibility that preventing cyclin C from reaching the mitochondria may help increase survival upon starvation conditions, we constructed a chimeric gene fusing the outer mitochondrial membrane (OMM)-associating domain of Fis1 to the C terminus of cyclin C-YFP (called cyclin C-YFP-Fis). Fis1 is the receptor for the GTPase Dnm1-Mdv1 complex required for mitochondrial fission (Tieu and Nunnari, 2000; Karren et al., 2005; Bhar et al., 2006). As expected, fluorescence
microscopy revealed that cyclin C-YFP-Fis localized to the OMM and this fusion protein is sufficient to induce mitochondrial fragmentation in the absence of a stress signal (Figure 8A, quantified in B and C).

Next, we addressed whether redirecting cyclin C to the mitochondria affects cell survival in response to rapamycin stress. The survival assay described above (Figure 6C) was repeated using cnc1Δ cells expressing either cyclin C-YFP or cyclin C-YFP-Fis. Compared to the cyclin C-YFP control, cells harboring the cyclin C-YFP-Fis construct exhibited approximately a 10-fold decrease in viability on 2.5 ng/ml rapamycin and thereafter cell viability determined by growth (10-fold dilutions) on rich medium (YPDA). (G) Mid-log wild-type cells were treated as described in F except that except that the % of live cells was monitored by FDA staining and FACS analysis after 24 h. The % of dead cells (FDA negative) was graphed for each condition. N = 3. ****p < 0.0001.

FIGURE 7: Rapamycin treatment protects mitochondria from H2O2-mediated fragmentation. (A) Wild-type cells harboring the plasmids shown were grown to mid-log and treated with 50 ng/ml rapamycin and thereafter 0.8 mM H2O2 for the times indicated. Representative fluorescence microscopy images of the results are shown. Bar = 5 µm. (B) Representative images of cells from A treated with 50 ng/ml rapamycin. Bar = 5 µm. (C, D) The percentage of cells from A that had cytoplasmic cyclin C and fragmented mitochondria respectively. N = 3. (E) GFP-Atg8 cleavage assays following treatment with 50 ng/ml rapamycin (left panel) or 0.8 mM H2O2 (middle panel) for the timepoints indicated. In the right-hand panel, H2O2 was added for 1 h after rapamycin addition. (F) Mid-log wild-type cells were subjected to the conditions shown (50 ng/ml rapamycin and 2 mM H2O2) and thereafter cell viability determined by growth (10-fold dilutions) on rich medium (YPDA). (G) Mid-log wild-type cells were treated as described in F except that except that the % of live cells was monitored by FDA staining and FACS analysis after 24 h. The % of dead cells (FDA negative) was graphed for each condition. N = 3. ****p < 0.0001.
phenotype. Taken together, these results support a model that rapidly destroying cyclin C by the UPS restricts it from translocating to the cytoplasm following nitrogen starvation, thereby preventing mitochondrial fragmentation and RCD (see Discussion). Therefore, manipulating the location of a single protein allows the cell to quickly adapt to changing environmental cues such as oxidative stress or nitrogen starvation to deliver the correct molecular response.

**DISCUSSION**

Correctly adapting to environmental signals requires the cell to translate signals into a molecular response. Previous studies have shown that this response triggers a change in mitochondria morphology, with fragmented mitochondria preceding RCD (see Discussion). Therefore, manipulating the location of a single protein allows the cell to quickly adapt to changing environmental cues such as oxidative stress or nitrogen starvation to deliver the correct molecular response.

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a role in directing cell fates. Combined with our previous studies (Cooper et al., 1997, 2014), these results demonstrate that cyclin C’s role as a repressor of SRGs is identical following both survival and cell death cues, whereas it only mediates mitochondrial fragmentation in response to cell death cues.

Although the exact location of cyclin C destruction following nitrogen starvation was not determined, these studies imply that it is different from oxidative stress. Exactly where active proteasomes are located is still controversial and was recently reviewed (Enenkel, 2014). Solid genetic experiments with nuclear and cytoplasmic substrates suggest that active proteasomes are located on the outside of the NPC (Chen and Madura, 2014). Similar genetic approaches by others with different substrates suggest that proteasomes destroy substrates in the nucleus (Boban et al., 2014). Likewise, Ump1-GFP, and other proteasomal subunits, have been reported to primarily localize to the nucleus in dividing cells (Lehmann et al., 2002; Fehlker et al., 2003). Last, it has been suggested that active proteasomes are located in clusters surrounding the NPC (Chen and Madura, 2014; Albert et al., 2017), which may help resolve the conflicting data obtained by different groups. Further studies with cyclin C may not only be useful in understanding how the CKM controls the response to various stresses but also could be used as a test substrate to investigate what cellular compartment maintain active proteasomes. It was interesting that the cyclin C mutation (A110V) that stabilizes cyclin C in the nucleus in response to oxidative stress (Jin et al., 2014a) did not have the same effect following nitrogen starvation. This same mutation also partially stabilizes cyclin C following heat shock where cytoplasmic foci are seen and during meiosis (Cooper and Strich, 1999; Cooper et al., 1997, 2012). This suggests that different upstream signal-

**FIGURE 9:** Model for regulator network controlling the final subcellular address for cyclin C and subsequent cell fate decisions. In unstressed cells growing in replete media (left-hand panel), cyclin C-Cdk8 repress the expression of a subset of stress-responsive genes; 80–90% of the mitochondria show a reticular fused phenotype. After treatment with H$_2$O$_2$ (middle panel), cyclin repression on cyclin C–Cdk8-controlled genes is relieved by cyclin C translocation to the cytoplasm (1). There it interacts with the fission machinery (and Bax in mammalian cells) to induce stress-induced mitochondrial fission and promote cell death (2). Thereafter, cyclin C is destroyed by the UPS dependent on the Not4 E3 ligase. Following nitrogen depletion (right panel), the cyclin C-Cdk8 repression of the autophagy gene ATG8 is relieved by destroying cyclin C before it can affect mitochondrial morphology (possibly in the nucleus or at the NPC) by the UPS system. The 26S proteasome and an unknown E3 ligase(s) are required. As a result, the mitochondria remain reticular or may even become hyper-elongated to promote ATP production under starvation conditions. $N =$ nucleus, $C =$ cytoplasm.

**Cyclin C destruction following nitrogen depletion promotes cell survival by up-regulation of ATG8 transcription**

Results presented in this paper argue that the second reason for cyclin C destruction following nitrogen starvation is to prevent mitochondrial fission. As cyclin C-mediated mitochondrial fission promotes H$_2$O$_2$-induced RCD, the simplest interpretation would be that preventing cyclin C–mitochondrial association promotes the autophagic response over the death program (see Figure 9). This model is strongly supported by our finding that misdirecting cyclin C to the mitochondria changes the starvation signal from survival to death. The relationship between mitochondrial morphology and cell survival has been studied in mammalian cells (Rambold et al., 2011a; Gomes and Scorrano, 2013). In these studies, autophagy induction by starvation or mTORC1 inhibition stimulates mitochondria hyperfusion. As a result, mitochondria possess more cristae which in turn optimizes ATP production (Buck et al., 2016; Gandin et al., 2016; Mishra and Chan, 2016), thereby linking mitochondrial dynamics to the metabolic state of the cell (Morita et al., 2017). This protective effect of mTORC1 inhibition against oxidative stress is also evident in reactive oxygen species-associated pathologies such as aging and neurodegenerative diseases (Rubinsztein et al., 2011), playing a role in

Cyclin C destruction following nitrogen depletion promotes cell survival by inhibiting extensive mitochondrial fragmentation

Why does cyclin C destruction promote cell survival following nitrogen starvation? We propose that there are two main explanations for this observation. First, destroying cyclin C inactivates its cognate kinase Cdk8, promoting induction of the essential autophagy gene ATG8. The studies presented here, combined with those of others, show that ATG8 derepression upon nitrogen starvation requires multiple steps. First, Ume6 function is negated on phosphorylation by the great wall kinase Rim15 (Backues et al., 2012). Epistasis experiments place cyclin C-Cdk8 into this pathway in an unknown role. Consistent with this model, both cyclin C-Cdk8 (Figure 5) and Ume6 do not regulate ATG7 or ATG14 mRNA expression (Jin et al., 2014b). One possibility is that this kinase phosphorylates a member of the Ume6–HDAC complex that in turn maintains ATG8 repression. An alternative, but not mutually exclusive possibility, is that cyclin C-Cdk8 represses Rim15 activity. Two targets of this pathway are the transcriptional activators Msn2–Msn4 which up-regulate ATG8 expression by an unknown mechanism upon starvation (Vlahakis et al., 2017). Although we currently do not understand the precise molecular mechanism, the results presented here show that that precocious ATG8 induction in the absence of cyclin C-Cdk8 provides a growth advantage. Importantly, this growth advantage was dependent on autophagy induction, as it was not observed in atg8Δ cells (Figure 6). This suggests a model in which the increased GFP-Atg8 foci seen in cnclΔ or cdkΔ cells primes them to deal with starvation conditions, but an external starvation signal is required to trigger autophagic activity. Last, although these results show that cyclin C-Cdk8 regulate ATG8, it is possible that this kinase could negatively regulate other autophagy genes. Consistent with this, Cdk8 was identified in a recent proteomics screen as a negative regulator of general autophagy (Müller et al., 2015). Different to our experiments, here autophagic activity was measured after 24 h rapamycin stress using the alkaline phosphatase-based assay which utilizes a cytosol-targeted proenzyme of ALP (cytALP) (Mendel et al., 2011).
maintaining long-term fitness of cells, predominantly by up-regulating transcriptional responses that support adaptation (Aramburu et al., 2014). However, the role mitochondria play is just starting to emerge (Sun et al., 2016; Srivastava, 2017) with recent studies showing that inhibition of mitochondrial fission attenuates disease progression in models of neurodegenerative and cardiovascular diseases (Cui et al., 2010; Givimani et al., 2012; Ong et al., 2010). Mitophagy is also attenuated in cells with reduced mitochondrial fission, suggesting that fission is a prerequisite for mitophagy to occur (Arnoult et al., 2005; Lee et al., 2011; Liu et al., 2020). Likewise, it is not completely understood how mitochondrial dynamics affect cell death mechanisms (Liu et al., 2020). However, our results in mammalian cells strongly suggest that cyclin C plays a role, as it binds to Drp1 following stress (Ganesan et al., 2019). Moreover, we also found that cyclin C also associates directly with Bax (Jezeck et al., 2019a), affirming its previously characterized role in regulating MOMP and intrinsic cell death mechanisms (Wang et al., 2015). Given the identification of cyclin C as a tumor suppressor (Li et al., 2014; Jezeck et al., 2019c) and the contradictory role of autophagy as a mechanism that both promotes and inhibits tumorgenesis (Lorente et al., 2020), this work emphasizes the need for further studies to investigate cyclin C’s role in promoting survival in response to nutrient depletion in mammals.

MATERIALS AND METHODS

Yeast strains and plasmids

Experiments performed primarily in the S. cerevisiae W303 background (Ronne and Rothstein, 1988) are listed in Supplemental Table S2 and have an RSY prefix. Experiments performed in different strain backgrounds are noted in the figure legend. In accordance with the Mediator nomenclature unification efforts (Bourbon et al., 2004), the cyclin C (SSN8/UME3/SRB11) and Cdk8 (SSN3/UMES/SRB10) will use CNC1 and CDK8 gene designations, respectively. Plasmids used in this study are listed in Supplemental Table S3. Plasmids construction details are available on request. All constructs were verified by sequencing. The previously described transmembrane domain of Fis1 (amino acids 121-155; Horie et al., 2003) was used to construct the cyclin C-YFP-Fis1 chimeric protein.

Yeast cell growth

Yeast were grown in either rich, nonselective medium (YPD: 2% [wt/vol] glucose, 2% [w/v) Bacto peptone, 1% (w/v) yeast extract, 0.001% [w/v) adenine sulfate or synthetic minimal dextrose (SD: 0.17% [wt/vol) yeast nitrogen base without amino acids and ammonium sulfate, 0.5% [wt/vol) ammonium sulfate, 1x supplement mixture of amino acids, 2% [wt/vol) glucose) allowing plasmid selection as previously described (Cooper et al., 1997). For all experiments, the cells were grown to mid-log phase (6×10^6 cells/ml) in 2% glucose media selecting for plasmids when appropriate before directly adding rapamycin (Biovision, dissolved in 90% Tween, 10% ethanol) or 0.8 mM H2O2 (Millipore Sigma HX0635-3) at the concentrations indicated into the SD media. For the nitrogen starvation experiments, mid-log phase cells were washed in H2O then resuspended in synthetic dextrose medium lacking ammonium sulfate or supplemental amino acids (SD-N). Protein extracts were prepared from 25-ml culture samples per timepoint and washed in H2O, and the pellet was flash frozen in liquid nitrogen (see below for more details). E. coli cells for isolating plasmids were grown in LB medium with selective antibiotics as previously described (Cooper et al., 1997). For the rapamycin survival assays (Figures 6C and 8C), mid-log cells were grown in YPDA medium, serially diluted 10-fold, and then plated onto SD complete plates containing either 200 or 2.5 ng/ml rapamycin. For growth rate studies (Figure 6F), strains were grown in YPDA to 2×10^6 cells/ml and then followed for 6 h after the addition of 2.5 ng/ml rapamycin.

For the cycloheximide chase assay (Supplemental Figure S1E), mid-log phase cells were treated with 50 µg/ml cycloheximide dissolved in dimethyl sulfoxide for 4 h.

Yeast cellular assays

RT-qPCR analysis was executed as previously described (Cooper et al., 2012). Oligonucleotides used are available on request. All RT-qPCR studies were conducted with three biological samples in technical duplicates. For all other assays, the experiments were performed in biological replicates and the number of replicates for each experiment is given in the figure legends. P values were determined using the Student’s t test. GFP-Atg8 cleavage assays were executed as described (Nair et al., 2011) using the single-copy GFP-Atg8 expression plasmid (Abeliovich et al., 2003) except that protein extracts were prepared using the NaOH lysis method described below. For the three (Figure 8D), the cells were grown in SD, washed, and then resuspended in SD-N. After 3 d, the cells were stained with phloxine B (Millipore Sigma P2759) at 5 µg/ml essentially as described (Noda, 2008). The percentage of phloxine positive (dead) cells was measured using FACs as previously described (Khakhina et al., 2014). To determine cell viability in Figure 7G, the cells were stained, grown in SD medium to mid-log, split into four, and treated as shown. H2O2 was added directly to rapamycin-treated cells at the time indicated. After the treatment, the cells were spun down and resuspended in phosphate-buffered saline and stained with FDA (Millipore Sigma P7378) at 10 µg/ml essentially as described (Kwolek-Mirek and Zadrag-Tecza, 2014). The percentage of FDA-positive (live) cells was analyzed by FACs and plotted as the amount of unstained (dead) cells. For both FDA and phloxine stains, 30,000 cells were counted per timepoint and samples were analyzed in biological triplicate. The plating assay shown in Figure 7F was essentially executed in a similar manner. The cells were grown in SD to mid-log, split, and treated as shown in the figure with H2O2 being added to rapamycin-treated cells. After the time shown, 10-fold dilutions of the cells were made and plated on YPDA media.

Western blot analysis

Protein extracts for Western blot studies were prepared using a NaOH lysis procedure exactly as described (Stieg et al., 2018; Willis et al., 2018). To detect epitope tagged proteins 1:5000 dilutions of all primary antibodies were used (anti-myc [UpState New York], anti-HA and anti-GFP [Abcam]) except that 1:2500 dilutions of anti-Pgk1 (Invitrogen) antibodies were used. Western blot signals were detected using 1:5000 dilutions of either goat anti-mouse or goat anti-rabbit secondary antibodies conjugated to alkaline phosphatase (Abcam) and the CDP-Star chemiluminescence kit (Thermo). Signals were quantified (relative to Pgk1 controls) by CCD camera imaging (Kodak). All degradation assays were performed more than once on biological replicates (see figure legends for the exact number). The SD was calculated from the mean (error bars) using GraphPad Prism 7. The half-life of cyclin C degradation was calculated from these graphs.

E3 ligase screen

The putative E3 ligases screened are listed in Supplemental Table S1 and were previously identified by others (Finley et al., 2012;
Zhu et al., 2016). Viable null mutants from the yeast knockout collection (Chu and Davis, 2008) were transformed with cyclin C-myc (pLR101) and were starved of nitrogen for 4 h. Cyclin C was detected by Western blot analysis as described above and images were quantified as described above. The blots were stripped and probed for Pgk1. A representative Western blot image is shown in Supplemental Figure S2E. To create the rps5 end-N rule doxycycline inducible degron (Supplemental Figure S2A), pMK632 was integrated into cells to create RSY2301 (Ubi-leucine::3HA-Rsp5). Next, RSY2301, harboring cyclin C-myc, was grown to mid-log, a sample was removed to control for Rsp5-HA expression, and then treated with 2 μg/ml doxycycline (Sigma D9891) for 5 h. Thereafter, the cells were washed in water and resuspended in SD-N plus doxycycline and samples were taken for NaOH lysis and Western analysis as described above. The same method was used to create the skp1 and tfb3 end-N rule degron strains, except that pMK-Arg was integrated into cells to create RSY2366 and RSY2367, respectively.

Fluorescence microscopy

YFP-cyclin C subcellular localization and mitochondrial morphology was monitored as described previously (Cooper et al., 2012, 2014). For all experiments, the cells were grown to mid-log (6 × 10^6 cells/ml), treated with either H_2O_2, rapamycin, or resuspended in SD-N (after being washed in water) for the timepoints indicated, then analyzed by fluorescence microscopy. Images were obtained using a Nikon 90i microscope with a 100x objective with 1.2x camera magnification (Plan Fluor Oil, NA 1.3) and a CCD camera (Hamamatsu model C4742). Data were collected using NIS software and processed using Image Pro software. All images of individual cells were optically sectioned (0.2-μm slices at 0.3-μm spacing) and deconvolved, and the slices were collapsed to visualize the entire fluorescent signal within the cell. Cyclin C-YFP foci were scored as being cytoplasmic when three or more foci were observed outside of the nucleus (Jin et al., 2014a). Mitochondrial fission assays were performed on live cells as described (Cooper et al., 2014). In brief, mitochondrial fission was scored positive if no reticular mitochondria were observed that transversal half the cell diameter. Fission was scored when cells exhibited one or more reticular mitochondria the diameter of the cell. Fission and fusion was scored for 200 cells from three independent isolates. GFP-Atg8 foci were scored in unstressed cells for at least 200 cells from three independent isolates as described (Torggler et al., 2017). Statistical analysis was performed using the Student’s t test.

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REFERENCES

Abeliovich H, Zhang C, Dunn WA Jr, Shokat KM, Klionsky DJ (2003). Chemical genetic analysis of Apg1 reveals a non-kinase role in the induction of autophagy. Mol Biol Cell 14, 477–490.

Albert S, Schaffer M, Beck F, Mosalaganti S, Asano S, Thomas HF, Plitzko JM, Beck M, Baumeister W, Engel BD (2017). Proteasomes tether to two distinct sites at the nuclear pore complex. Proc Natl Acad Sci USA 114, 13726–13731.

Allen BL, Taates DJ (2015). The Mediator complex: a central integrator of transcription. Nat Rev Mol Cell Biol 16, 155–166.

Aramburu J, Ortell MC, Tejedor S, Buxade M, Lopez-Rodriguez C (2014). Transcriptional regulation of the stress response by mTOR. Sci Signal 7, re2.
BH3 domains activate Bax and nucleate its oligomerization to induce apoptosis. Cell 152, 519–531.

Dalny M, Geng F, Muratani M, Geisinger JM, Salghetti SE, Tansey WP (2008). Modulation of RNA polymerase II subunit composition by ubiquitylation. Proc Natl Acad Sci USA 105, 19649–19654.

Delorme-Ax#for(0,460)lf E, Guimaraes RS, Reggiori F, Klionsky DJ (2015). The yeast Saccharomyces cerevisiae: an overview of methods to study autophagy progression. Methods 75, 3–18.

Enenkel C (2014). Nuclear transport of yeast proteasomes. Biomolecules 4, 940–955.

Fehlker M, Wendler P, Lehmann A, Enenkel C (2003). Bim3 is part of nascent proteasomes and is involved in a late stage of nuclear proteasome assembly. EMBO Rep 4, 957–963.

Finley D, Ulrich HD, Sommer T, Kaiser P (2012). The ubiquitin-proteasome system of Saccharomyces cerevisiae. Genetics 192, 319–360.

Galluzzi L, Baehrelle EH, Ballabio A, Boya P, Bravo-San Pedro JM, Cecconi F, Choi AM, Chu CT, Codogno P, Colombo MI, et al. (2017). Molecular definitions of autophagy and related processes. EMBO J 36, 1811–1836.

Gandin Y, Mavridis L, Hulea L, Gravel SP, Cargnello M, McLaughlin S, Cai Y, Balanathan M, Morita M, Rajakumar A, et al. (2016). nanoCAGE reveals S’ UTR features that define specific modes of translation of functionally related MTOR-sensitive mRNAs. Genome Res 26, 636–648.

Ganesan V, Willis SD, Chang KT, Beluch S, Cooper KF, Strich R (2019). Cyclin C directly stimulates Drp1 GTP affinity to mediate stress-induced mitochondrial fission. Mol Biol Cell 30, 302–311.

Givvimani S, Munjal C, Tyagi N, Sen U, Metreveli N, Tyagi SC (2012). Fast protein-depletion system utilizing tetracycline repressible promoter and N-end rule in yeast. Mol Biol Cell 23, 762–768.

Gomes LC, Di Benedetto G, Scorrano L (2011). During autophagy mitochondria elongate, are spared from degradation and sustain cell viability. Nat Cell Biol 13, 589–598.

Gomes LC, Scorrano L (2013). Mitochondrial morphology in mitophagy and macroautophagy. Biochim Biophys Acta 1833, 205–212.

Gray JV, Petsko GA, Johnston GC, Ringo D, Singer RA, Werner-Washburne M (2004). “Sleeping beauty”: quiescence in Saccharomyces cerevisiae. Microbiol Mol Biol Rev 68, 187–206.

Gross A, Katz SG (2017). Non-apoptotic functions of BCL-2 family proteins. Mol Biol Cell 28, 1987–2014.

Gnanasundram SV, Kos M (2015). Uib4 and Ulb4: two ubiquitin conjugating enzymes Ubc4 and poly-ubiquitination in ethanol-induced down-regulation of targeted proteins. Biosci Biotechnol Biochem 73, 1836–1840.

Hartmann-Petersen R, Seeger M, Gordon C (2003). Transferring substrates at the yeast proteasome. Trends Biochem Sci 28, 26–31.

Hirai H, Okada M, Ohtsu I, Takagi H (2009). A functional analysis of yeast Dss1, regulates exocytosis and pseudohyphal differentiation in Schizosaccharomyces pombe. Mol Biol Cell 20, 8268–2273.

Holstege FC, Jennings EG, Wyrick JJ, Lee TI, Hengartner CJ, Green MR, et al. (1998). A circuitry of a eukaryotic genome. Cell 95, 717–728.

Jin C, Strich R, CooperKF (2014a). Slt2p phosphorylation induces cyclin C nuclear-to-cyttoplasmic translocation in response to oxidative stress. Mol Biol Cell 25, 1396–1407.

Jin M, He D, Backues SK, Freeberg MA, Liu X, Kim JK, Klionsky DJ (2014b). Transcriptional regulation by Pho23 modulates the frequency of autophagosome formation. Curr Biol 24, 1314–1322.

Jin M, Kim SK, Willis SD, CooperKF (2015). The MAPKks Ste11 and Bck1 jointly transduce the high oxidative stress signal through the cell wall integrity MAP kinase pathway. Microb Cell 2, 329–342.

Kadosh D, Struhl K (1997). Repression by Um6 involves recruitment of a complex containing Sin3 corepressor and Rpd3 histone deacetylase to target promoters. Cell 89, 365–371.

Karren MA, Coonrod EM, Anderson TK, Shaw JM (2005). The role of Fis1p-Mdr1p interactions in mitochondrial fission complex assembly. J Cell Biol 171, 291–301.

Kasahara A, Scorrano L (2014). Mitochondria: from cell death executioners to regulators of cell differentiation. Trends Cell Biol 24, 761–770.

Khakhina S, CooperKF, Strich R (2014). Med13p prevents mitochondrial fission and programmed cell death in yeast through nuclear retention of cyclin C. Mol Biol Cell 25, 2807–2816.

Kirisako T, Baba M, Ishihara N, Ohsumi Y, Ohsumi T, Noda T, Ohsumi Y (1999). Formation process of autophagosome is traced with Apg8/Aut7p in yeast. J Cell Biol 147, 435–446.

Klosinska MM, Crutichfield CA, Bradley PH, Rabonwitz JD, Broach JR (2011). Yeast cells can access distinct quiescent states. Genes Dev 25, 336–349.

Krasley E, CooperKF, Mallory MJ, Dunbrack R, Strich R (2006). Regulation of the oxidative stress response through Sltp2p-dependent destruction of cyclin C in Saccharomyces cerevisiae. Genetics 172, 1477–1486.

Krause SA, Gray JV (2002). The protein kinase C pathway is required for viability in quiescence in Saccharomyces cerevisiae. Curr Biol 12, 588–593.

Kwolek-Mirek M, Zadrag-Tecza R (2014). Comparison of methods used for assessing the viability and vitality of yeast cells. FEBS Lett 580, 1068–1079.

Lander GC, Estrin E, Matsyekla ME, Bashore C, Nogales E, Martin A (2012). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Mitochondrial autophagy by Bnip3 involves Drp1-mediated mitochondrial fission and recruitment of Parkin in cardiac myocytes. Am J Physiol Heart Circ Physiol 301, H1924–H1931.

Lehmann A, Janek K, Braun B, Kloetzl PM, Enenkel C (2002). 20 S proteasomes are imported as precursor complexes into the nucleus of yeast. J Mol Biol 317, 401–413.

Lee Y, Lee HY (2005). Cell wall integrity signaling in Saccharomyces cerevisiae. Microbiol Mol Biol Rev 69, 262–291.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.

Lee Y, Lee HY, Hanna RA, Gustafsson AB (2011). Complete subunit architecture of the proteasome regulatory particle. Nature 482, 186–191.
Vlahakis A, Lopez Muniozguren N, Powers T (2017). Stress-response transcription factors Msn2 and Msn4 couple TORC2-Ypk1 signaling and mitochondrial respiration to ATG8 gene expression and autophagy. Autophagy 13, 1804–1812.

Waite KA, De-La Mota-Peynado A, Vontz G, Roelofs J (2016). Starvation induces proteasome autophagy with different pathways for core and regulatory particles. J Biol Chem 291, 3239–3253.

Wang K, Yan R, Cooper KF, Strich R (2015). Cyclin C mediates stress-induced mitochondrial fission and apoptosis. Mol Biol Cell 26, 1030–1043.

Wang G, Yang J, Huibregtse JM (1999). Functional domains of the Rsp5 ubiquitin-protein ligase. Mol Cell Biol 19, 342–352.

Wang X, Yen J, Kaiser P, Huang L (2010). Regulation of the 26S proteasome complex during oxidative stress. Sci Signal 3, ra88.

Willis SD, Stieg DC, Ong KL, Shah R, Strich R, Grose JH, Cooper KF (2018). Snf1 cooperates with the CWI MAPK pathway to mediate the degradation of Med13 following oxidative stress. Microbial Cell 5, 357–370.

Wissing S, Ludovico P, Herker E, Buttner S, Engelhardt SM, Decker T, Link A, Proksch A, Rodrigues F, Corte-Real M, et al. (2004). An AIF orthologue regulates apoptosis in yeast. J Cell Biol 166, 969–974.

Xie Z, Nair U, Klionsky DJ (2008). Atg8 controls phagophore expansion during autophagosome formation. Mol Biol Cell 19, 3290–3298.

Yukawa M, Yo K, Hasegawa H, Ueno M, Tsuchiya E (2009). The Rpd3/HDAC complex is present at the URS1 cis-element with hyperacetylated histone H3. Biosci Biotechnol Biochem 73, 378–384.

Zhang D, Chen T, Ziv I, Rosenzweig R, Matsuhih Y, Bronner V, Glickman MH, Fushman D (2009). Together, Rpn10 and Dsk2 can serve as a polyubiquitin chain-length sensor. Mol Cell 36, 1018–1033.

Zhu J, Deng S, Lu P, Bu W, Li T, Yu L, Xie Z (2016). The Ccl1-Kin28 kinase complex regulates autophagy under nitrogen starvation. J Cell Sci 129, 135–144.