Our Paths Might Cross: the Role of the Fungal Cell Wall Integrity Pathway in Stress Response and Cross Talk with Other Stress Response Pathways

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Fungi occupy diverse environments and are subjected to many extreme conditions. Among the stressful conditions faced by fungi are pH changes, osmotic changes, thermal changes, oxide radicals, nutrient deprivation, and exposure to chemicals. These adversities can be found either in the environment or in animal and human hosts. The cell wall integrity (CWI) pathway provides a means to fortify and repair damages to the cell wall in order to withstand stressful environments. The CWI pathway in comprised of cell wall stress sensors that lead to activation of a mitogen-activated protein kinase (MAPK) cascade. Signaling through the MAPK cascade leads to expression of transcription factors that facilitate biosynthesis of cell wall components and actin organization. Given the relatively limited number of components of the CWI pathway and the very diverse stimuli, there must be a means of expanding the pathway. To manage the diverse stress conditions, the CWI pathway cross talks with other pathways or proteins, and these cross talk events enhance the signaling capabilities of the CWI pathway. Lateral influences that facilitate maintaining the cell wall under stress conditions are TOR signaling, calcineurin signaling, the high-osmolarity glycerol pathway, the cyclic AMP-protein kinase A pathway, and additional proteins. In this article, we highlight several of the cross talk events that have been described for *Saccharomyces cerevisiae* and several other fungi.

Fungal environments range from soil to plants to animal and human hosts. Fungi can also inhabit extreme environments such as hydrothermal vents, bird excreta, the arctic, aquatic environments, salterns, mine drainages, or even bare rock surfaces in the case of lichens. Each environment presents challenges for fungi that must be overcome for them to survive and grow, including osmotic changes, oxidative stress, heat shock, pH changes, nutrient limitations, and chemical challenges. These stresses emanate either from natural environments or from the host immune system in response to pathogenesis. Exposure of fungal cells to any of these stress conditions results in altered gene expression to enable the cell to endure the adverse environment. Changes to gene expression require a coordinated effort from multiple pathways in order to allow a limited amount of proteins to achieve the complicated feat of surviving unfavorable conditions. The key defense to withstand environmental adversities is the fungal cell wall. Due to damage inflicted by the stressors, the cell wall is repaired and even fortified through cell wall biosynthesis and the integration of cell wall components into the cell wall when exposed to suboptimal or hostile environments. The cell wall is a cellular structure unique to fungi among eukaryotes (plant cells have a cell wall, but it is very different from the fungal cell wall). The cell wall varies between fungi, but the overall composition consists of α- and β-glucans (the principal polysaccharides of the bilayer-structured cell wall), N-acetylglucosamine, mannoproteins, and various other glycoproteins.

To elicit the necessary transcriptional changes due to extracellular stressors, fungal cells maintain transmembrane sensors that detect environmental changes and initiate signals that are conveyed from the cell surface to the nucleus. The model yeast *Saccharomyces cerevisiae*, for example, utilizes five separate mitogen-activated protein kinase (MAPK) cascades to respond appropriately to environmental changes (8). The cell wall integrity (CWI) pathway utilizes one of the MAPK cascades to facilitate the maintenance of the cell wall by mediating cell wall biosynthesis, actin organization, and other events necessary to maintain CWI (46). As *S. cerevisiae* is the model eukaryote, much of what is known about the CWI pathways as well as other signaling pathways has been defined in this organism. This review therefore highlights what is known about the pathways in *S. cerevisiae* and makes comparisons to other fungal species.

The CWI pathway utilizes GTPase-activating proteins and guanylnucleotide exchange factors (GEFs) to regulate the activation of the kinase cascade that leads to the activation of transcription factors. In *S. cerevisiae* this cascade is initiated by cell wall-associated stress sensors Mid2 and Wsc1 (38, 78) (Fig. 1). These proteins bind to Rom2, which is a GEF for Rhol (58, 62) (Table 1). Rhol affects compositional changes in the cell wall through activation of the glucan synthase Fks1 (19, 53, 63), which facilitates the production of the major cell wall component 1,3-β-D-glucan (18). Rhol also binds and activates Pck1 (36, 56), which in turn regulates the MAPK cascade. Pck1 phosphorylates Bck1, a MAPK kinase kinase (MAPKKK), which transmits the signal to MAPK kinases (MAPKKs) Mkk1.
and Mkk2. These two kinases finally activate the MAPK Slt2/Mpk1 (5). The stimulation of Slt2/Mpk1 leads to phosphorylation of the transcription factors Rlm1 and SBF (consisting of the two transcription factors Swi4 and Swi6), both of which initiate the expression of cell wall synthesis genes (17, 34, 50, 81).

Overall, the CWI pathway is conserved among fungi, including budding yeast, fission yeast, and filamentous fungi (61). However, components from some fungi are still being elucidated, and important differences in the pathway have been identified. An example is the redundant MAPKKs Mkk1 and Mkk2 of *S. cerevisiae*, for which there is only the MAPKK Mkk2 in *Cryptococcus neoformans* (27) (Table 1). Additionally, Slt2/Mpk1 is activated in *S. cerevisiae* in response to heat shock, cell wall stress compounds, or oxidative stress. However, in *Schizosaccharomyces pombe* the Slt2/Mpk1 homolog pmk1p is activated only in response to heat stress and sodium chloride (osmotic changes), indicating a more limited response by the CWI pathway of *S. pombe* (51).

Although there are slight differences, the CWI function of maintaining cell integrity is conserved between fungi, and CWI pathway activation is not restricted to an individual stimulus but can be elicited by a number of events. A genome-wide analysis of five cell wall mutants indicated transcriptional changes in 5% of the *S. cerevisiae* genome (or approximately 300 genes) (44), suggesting complex changes in gene expression not limited to a single signal cascade. Cross talk between the CWI pathway and other stress response pathways enable the CWI pathway to respond to numerous, diverse stress events with responses that appropriately alleviate cellular stress.

### RESPONSE TO VARIABLE OSMOTIC CONDITIONS

Osmoregulation is needed to maintain fungal cell turgor, volume, and the appropriate intracellular environment for biochemical reactions (32). For example, some fungi, such as *Aspergillus* spp., are found in the soil, an environment that can undergo osmotic change due to drought, rain, or other climate conditions. Furthermore, this fungus is capable of surviving both in the soil environment and within a mammalian host as a human pathogen. Fungi that make this transition must be able to compensate for the changes in environmental osmolarity.

In *S. cerevisiae*, external osmotic changes elicit responses from both the CWI pathway and the high-osmolarity glycerol (HOG) response pathway. In general, the CWI pathway can respond to hypoosmotic conditions but the HOG pathway of *S. cerevisiae* responds to hyperosmotic conditions. However, the HOG pathway of pathogenic fungi in which it has been studied is broader in scope than governing just the osmotic response, because it also regulates stress responses to stimuli such as oxidative stress, UV light, heavy metals, and high temperature (1, 4, 7).

A cellular response to hypoosmotic conditions is manifested through the CWI pathway as indicated by the activation of the MAPK Slt2/Mpk1. The phosphorylation of Slt2/Mpk1 occurs without specificity toward osmotic solutes and is observed with hypotonic solutions of sorbitol, NaCl, or glucose (14). The CWI pathway response to hypoosmotic conditions relies on

![Diagram of the CWI pathway](image-url)
components of the MAPK module. Deletion of the MAPK module components prevents Slt2/Mpk1 phosphorylation under hypoosmotic conditions (14). Further, the resulting pck1, bck1, and mkk1 mkk2 mutant strains lyse in the absence of osmotic stabilizers (33, 45, 47).

The role of the CWI pathway in hypoosmotic stabilization is retained in human-pathogenic fungi. In the case of Candida albicans, hypoosmotic conditions initiate the phosphorylation of Mkc1 (an Slt2/Mpk1 homolog) (Table 1) (54). Activation of Mkc1 is Pkc1 dependent, and a pck1/pkc1 homozygous mutant lyses in the absence of osmotic support (60). A pck1 deletion strain of C. neoformans also lyses in the absence of osmotic stabilization (26).

In contrast to the response elicited by hypoosmotic conditions, hyperosmotic conditions experienced by S. cerevisiae trigger a response from the HOG pathway (as noted above) in which the inactivation of Sln1 kinase leads to the dephosphorylation of Ssk1. The unphosphorylated Ssk1 activates the MAPKKs Ssk22 and Ssk2 (32). Ssk2 then activates the MAPKK Pbs2. Pbs2 phosphorylates the MAPK Hog1, which then is transported to the nucleus to activate transcription factors. An alternative pathway to initiate Hog1, known as the Sho1-dependent pathway, also exists for S. cerevisiae. In this pathway, Sho1 activates Ste20 in response to osmotic stress via the Cdc42 GTPase, which then stimulates the MAPKKS Ste11. Ste11 with Ste50 is then able to activate Pbs2 for the phosphorylation of Hog1 (3). The Sho1-dependent alternative pathway for Hog1 activation does not appear to be conserved in all other fungi (3).

Thus, opposing osmotic conditions activate the CWI and HOG pathways; therefore, it is perceivable that there is some type of cross talk to coordinate the regulation of the two pathways. A cross talk event indeed occurs between the CWI and HOG pathways, involving Slt2/Mpk1. While hypotonic solutions induce the phosphorylation of Slt2/Mpk1 in a protein kinase C (PKC) pathway-dependent manner, hypotonic solutions induce phosphorylation of SLT2/MPK1 that is Hog1 and Rlm1 dependent (28) (Fig. 2). The Hog1-dependent transcription of the CWI pathway gene SLT2/MPK1 under hyperosmotic conditions and the CWI-induced activation of Slt2/Mpk1 suggest regulation between the HOG and the CWI pathways for osmotic homeostasis.

Though S. cerevisiae demonstrates a model in which hypoosmotic conditions trigger the CWI pathway and hyperosmotic conditions trigger the HOG pathway, this division to regulate osmolarity is not conserved in all fungi. S. pombe poses an exception where the CWI pathway component Slt2/Mpk1 homolog Pmk1 (72, 83) (Table 1) is activated regardless of whether the osmotic stressor is hyperosmotic or hypoosmotic (51). Madrid and colleagues demonstrated that the activation of Pmk1 relies on the MAPK cascade elements Mkh1 and Pek1 (Table 1) (51).

A general perspective is that the CWI pathway responds to osmotic stress regardless of the type of osmotic stress in some fungi but that in other types of fungi the HOG pathway is needed in conjunction with the CWI pathway to respond to osmotic stress. A cross talk event occurs to coordinate the effort of stabilizing the cell in hypertonic solutions.

RESPONSE TO pH STRESS

Much like the osmotic changes that can occur as fungal cells are transferred from one environment to another, pH changes can be encountered upon environmental transitions. The CWI pathway also functions to facilitate tolerance to pH changes. Alkaline stress in the environment is sensed by the Wsc1 transmembrane sensor of S. cerevisiae (68). Other genes from the CWI pathway important to alkaline tolerance are BCK1 and SLT2/MPK1, indicating that the Wsc1 signal is transmitted to the CWI MAPK cascade. This is exemplified by alkaline tolerance for a wsc1 mutant strain that contains a bck1-20 allele to induce constitutive activation of Slt2/Mpk1 (68).

It is interesting to note that the wsc1 mutation does not completely abolish phosphorylation of Slt2/Mpk1 in alkaline stress, but rather Slt2/Mpk1 phosphorylation is only decreased, indicating the potential for additional means of Slt2/Mpk1 activation either through activation of the CWI pathway utilizing a different sensor or through proteins outside of the linear CWI pathway in response to alkaline conditions. The remnant phosphorylation of Slt2/Mpk1 could possibly be contributed through activation of the CWI pathway via the Mid2 sensor under alkaline conditions. However, a mid2 mutant strain exhibits only a marginal decrease in Slt2/Mpk1 phosphorylation under alkaline conditions and is not alkali sensitive (68). Thus, if Mid2 does contribute to CWI activation under alkaline conditions it is likely a minor contribution. Whatever the additional alkaline-induced signaling that results in Slt2/Mpk1 phosphorylation may be, it is likely independent of the HOG pathway, which shows interactions with the CWI pathway under other stress conditions, because Hog1 is not phosphorylated under alkaline conditions (68).

The phosphorylated form of Slt2/Mpk1 has the capacity to activate either the Rlm1 or SBF transcription factor. Mutation
of the transcription factor \textit{RLM1} results in resistance to alkaline conditions, but strains with SBF defects are sensitive to alkaline media (68), indicating that alkali tolerance is SBF dependent but Rlm1 independent.

Fungal cells adapt not only to alkaline conditions but also to low-pH conditions. This stress response is particularly important to withstand pH changes encountered within hosts. Phagocytic cells introduce low-pH conditions when foreign bodies are phagocytosed as a means of host defense. Further, acidic conditions are also produced in the environment occupied by growing fungal cells as a result of active proton extrusion into the medium by the plasma membrane ATPase (77). Low-pH conditions lead to Slt2/Mpk1 phosphorylation. Indeed, both \textit{SLT2}/\textit{MPK1} and \textit{BCK1} from the CWI pathway were found to be required for growth under low-pH medium conditions (9), indicating the requirement for the MAPK cascade.

Under low-pH stress conditions, a mid2 mutant strain exhibits a decrease in the transcription of \textit{PST1}, a gene regulated by the Slt2/Mpk1-activated transcription factor Rlm1 from the CWI pathway. Transcription of \textit{PST1} under acidic stress conditions is not particularly reliant on the stress sensor Wsc1 (9), but rather the stress sensor Mid2 mediates a response to acidic conditions that leads to activation of the Rlm1 transcription factor.

The CWI pathway receives lateral influence under acidic conditions from Rgd1 (Fig. 3). The \textit{RGD1} gene encodes a Rho GTPase-activating protein that acts on Rho3 and Rho4. It plays a role in tolerance to acidic conditions, which is exemplified by \textit{rgd1} mutant strain sensitivity to low-pH conditions. The defects observed for the \textit{rgd1} mutant strain are exacerbated in an \textit{rgd1 mid2} double mutant strain (9), suggesting that Rgd1 influences the CWI pathway. The \textit{rgd1} mutant strain viability phenotype is suppressed by an increase in PKC pathway activity, and defects in the \textit{RGD1} gene lead to defects in PKC pathway activity (15). It is likely that the PKC pathway functions downstream of Rgd1. Furthermore, the expression of \textit{RGD1} under acidic shock conditions was Hog1 dependent (25). Again, this is another stress condition indicating cross talk between the HOG and CWI pathways.

Taken together, the findings so far indicate that Wsc1 is the CWI pathway stress sensor necessary for alkaline conditions and that Mid2 is the stress sensor required for acidic conditions. Alkaline-activated Wsc1 leads to transcription of SBF-dependent genes, and Mid2 low-pH activation leads to transcription of Rlm1-dependent genes. Hence, the difference stress sensors for the CWI pathway lead to activation of two different transcription factors. Further, the CWI pathway manages alkaline stress in a Hog1-independent manner but acidic stress conditions in a Hog1-dependent manner.

**RESPONSE TO HEAT SHOCK**

For fungi that maintain a niche within soil or plant environments, heat shock can occur with changes in climate temperature. Moreover, the difference in temperate conditions between a soil environment and a mammalian host can constitute a “heat shock” (for example, when fungi are relocated from the soil to the lungs of a host). In \textit{S. cerevisiae}, heat shock causes the activation of Slt2/Mpk1 (28, 35, 78). However, phosphorylation of Slt2/Mpk1 is diminished under heat shock conditions in a \textit{wsc1} mutant strain (78). An \textit{S. cerevisiae wsc1} deletion mutant exhibits thermosensitive growth under high-temperature growth conditions at 37°C (78), indicating that Wsc1 plays a role as a sensor for heat shock.

There is evidence that Wsc1 mediates cross talk between the CWI pathway and the cyclic AMP (cAMP)-protein kinase A (PKA) pathway of \textit{S. cerevisiae} for the heat stress response. The cAMP-PKA pathway is a nutrient-sensing mechanism that controls the cell cycle. In this signaling cascade, Ras is activated by a GEF, Cdc25, and is inactivated by GAP-activating proteins Ira1 and Ira2. The activation of Ras yields an increase in adenyl cyclase activity, which produces the second messenger cAMP for the activation of PKA, which targets several downstream proteins involved in metabolism. Among the downstream proteins regulated by PKA are the transcription factors Msn2/Msn4, which regulate the stress-inducible MAPK phosphatase Sdp1 (71).

Verna and colleagues (78) used an \textsc{ira2} deletion strain, which is heat shock sensitive (70), to identify inhibitors of Ras activity. A \textit{wsc1} deletion mutation was found to suppress the heat shock phenotype of the \textsc{ira2} mutant strain and is speculated to negatively regulate targets of Ras. Indeed, a \textsc{ras2} deletion mutation rescues the heat shock sensitivity of a \textit{wsc1} mutant strain. Further, a \textit{wsc} strain that overexpresses \textsc{ira2} is not sensitive to heat shock (78). Examination of which \textit{WSC} gene is involved in this signaling cascade reveals that \textit{WSC1} but
not WSC2 suppresses the heat shock sensitivity phenotype of a ras1Δras2Δ pCYR1 strain in which the RAS genes are deleted and adenylyl cyclase (CYR1) is overexpressed. This suggests that Wsc1 but not Wsc2 functions in conjunction with RAS signaling and that RAS and Wsc1 have opposing affects on a downstream target (78).

Thus, the Wsc1 transmembrane stress sensor of the CWI pathway facilitates cross talk with the cAMP-PKA pathway through the downstream activation of Slt2/Mpk1 in the CWI pathway and negative regulation of a downstream target of RAS in the cAMP-PKA pathway in response to thermal stress (Fig. 4). Additional cross talk with the cAMP-PKA pathway occurs at the Slt2/Mpk1 position of the CWI signaling cascade. Slt2/Mpk1 is negatively regulated by Msn2/Msn4-dependent Sdp1 from the cAMP-PKA pathway by direct dephosphorylation of Slt2/Mpk1 under heat shock conditions (28).

The phosphorylation of Slt2/Mpk1 is dependent on the MAPK cascade of the CWI pathway during several stress responses, and thus it is presumably that the MAPK cascade is necessary for the heat shock response. Several genes in the PKC pathway are indeed required for thermal tolerance, in-necessary for the heat shock response. Several genes in the

**RESPONSE TO LIMITED NITROGEN SOURCES**

Nutrient stress occurs when fungal cells transition from a nutrient-rich to a nutrient-poor environment. The environment in this case could be the soil, a host, or even the in vitro conditions of yeast culture media. The availability of nutrients dictates cellular functions such as cell growth and cell cycle progression. TOR (target of rapamycin) is named for the ability of the agent rapamycin to inhibit this particular signaling cascade. The TOR signaling cascade senses the availability of nitrogen sources to regulate gene expression. To study the effects of nitrogen-limited conditions, rapamycin is used to inhibit the TOR signaling cascade. The agent rapamycin forms a complex with FKBP12, which is then able to complex with the serine/threonine kinase Tor, inhibit TOR signaling, and prevent association between Sit4 and Tap42. In contrast, Tor activated by nitrogen sources promotes the association of the type 2A phosphatase Sit4 with Tap42. Tor protein is conserved in many fungi (10), but two fungi from which much has been elucidated about TOR signaling are *S. cerevisiae* and *S. pombe*, each encoding two Tor proteins, Tor1 and Tor2 (30, 37).

There is evidence for cross talk between TOR-mediated signaling and the CWI pathway. *S. cerevisiae* cells with inhibited TOR signaling due to exposure to rapamycin exhibit activation of Slt2/Mpk1. Strains carrying a TOR1-1 or TOR2-1 allele are insensitive to rapamycin. The TOR1-1 strain exhibits no change in phosphorylation of Slt2/Mpk1 in the presence of rapamycin, and a TOR2-1 mutant experiences mild phosphorylation of Slt2/Mpk1 (73). Thus, inhibition of Tor1 leads to CWI activation. This is further illustrated by the lack of Slt2/Mpk1 phosphorylation in response to rapamycin in *TAP42* and *SIT4* mutant strains. Interestingly, the TOR inhibited-induced activation of Slt2/Mpk1 is dependent on components of the CWI pathway, including Pkc1 and Bck1, as indicated by a lack of Slt/Mpk1 activation in *pkc1* and *bck1* deletion strains. Ex-
amino-terminal for the requirement of the upper part of the CWI signal cascade reveals that deletion of ROM2 diminishes phosphorylation of Slt2/Mpk1 in the presence of rapamycin (73). The decrease in phosphorylation rather than the complete lack of phosphorylation may be due to the presence of Rom1, which has a role partially redundant with that of Rom2 (58). Thus, the activation of Slt2/Mpk1 by TOR inhibition is dependent on PKC pathway components as well as upstream elements of the CWI pathway.

TOR signaling and the CWI pathway share the common function of actin organization. Although there are two TOR genes in *S. cerevisiae*, only TOR2 plays a role in actin organization (67). ROM2 restores growth in a *tor2*Δ mutant strain when overexpressed (31, 66). However, TOR2 overexpression does not repair the growth defect of a *rom2*Δ mutant strain (66). Further evidence that TOR influences Rom2 is demonstrated by the reduction in Rom2 GEF activity in a *tor2*Δ mutant strain. Thus, Rom2 likely functions downstream of Tor2 (66) (Fig. 5). Although it is unclear from the reports by Schmidt et al. (66) and Torres et al. (73) whether Tor directly or indirectly influences the CWI pathway, it is agreed by both groups that the most likely candidate to receive Tor-directed signals is Rom2.

Cross talk events between the TOR signaling and CWI pathways in the pathogenic fungus *C. albicans* are also observed. In *C. albicans* there is a single copy of TOR (10). Cross talk occurs through a G protein of the Ras superfamily, Rhb1, involved in the activation of Tor. A *rhh1/rhb1* deletion mutant strain is sensitive to rapamycin (75), indicating a relationship to Tor. Of note, the *RHH1* ortholog in *Aspergillus fumigatus* is *rhbA*. A role for *rhbA* in TOR signaling is conserved in *A. fumigatus*, where an *rhhA* deletion mutant strain is sensitivity to rapamycin (59).

Further, wild-type *C. albicans* cells exposed to rapamycin exhibit enhanced phosphorylation of Mkc1 (encoded by a *SLT2/MPK1* homolog) (Table 1), suggesting that TOR influences the CWI pathway (75). The *rhb1/rhb1* strain is also sensitive to cell wall stress induced by Congo red and calcofluor white (75), which indicate defects in CWI. Congo red and calcofluor white exposure are known to stimulate MAPK induction of Mkc1 (16, 38). Exposure of cells to calcofluor white indeed induces Mkc1 phosphorylation, but in the *rhh1/rhb1* strain activation of Mkc1 is diminished. This suggests that Mkc1 phosphorylation is Rhb1 dependent during cell wall stress (75).

**CALCIUM SIGNALING-INDUCED STRESS RESPONSE**

It has been established that calcium functions as a second messenger, and changes in calcium homeostasis have a number of physiological effects in cells. It is not well known what type of calcium exposure fungal cells encounter in the environment, but indeed they are able to react to calcium. Calcium has been shown to induce Mkc1 in *C. albicans* (54), and an *mkc1/mkc1* mutant is sensitive to calcium (55). The calcium-induced phosphorylation of Mkc1 is a Hog1-dependent event (54). Calcium is also known to be an activator of calcineurin in cells. Calcineurin is a Ca2+/calmodulin-dependent serine/threonine protein phosphatase. It is the target of such drugs as cyclosporine and FK506. In the inactive state calcineurin is a heterodimer composed of A and B subunits. When activated, it is in a Ca2+/calmodulin-bound state, causing a change that frees the active site of an autoinhibitory domain (21). *S. cerevisiae* carries two redundant calcineurin A genes, *CNA1* and *CNA2* (12, 49), and a single copy of the calcineurin B gene, *CNB1* (13, 42). Only a single copy of the calcineurin A subunit is encoded by the fungi *S. pombe* (82), *C. albicans* (65), *C. neoformans* (57), and *Aspergillus nidulans* (64). Calcineurin participates in a number of physiological processes in fungi, including cell cycle progression, morphological changes, and cell wall biosynthesis. Among pathogenic fungi, calcineurin is also important for virulence (57, 65). Inhibition of *C. neoformans* calcineurin function induces the phosphorylation of Mpk1 and the induction of *FKS1*. *FKS1* is also induced by calcineurin in *C. albicans* (65). Both Mpk1 and Fks1 are under the control of the CWI pathway (40). Fks1 is activated by Rho1, which also activates Pkc1 to induce the MAPK cascade. Interestingly, the deletion of *FKS1* in *S. cerevisiae* leads to increased sensitivity to the calcineurin inhibitor cyclosporine (20), suggesting an interplay between the CWI pathway and calcineurin.

The potential interplay between calcineurin and the CWI pathway is further strengthened by evidence from *S. cerevisiae* in which constitutively active calcineurin partially suppresses the lysis phenotypes of *pck1* and *slt2/mpk1* mutant strains (24). Garrett-Engle and colleagues (24) further demonstrated that the *pck1* mutant strain grown with the osmotic stabilizer was sensitive to the calcineurin inhibitors cyclosporine and FK506. An *slt2/mpk1* mutant strain also exhibited sensitivity to these calcineurin inhibitors (24). Thus, calcineurin compensates for some of the defects in the CWI pathway. The calcineurin overexpression may have produced more osmotically stable
conditions for the \( \text{pkc}1 \) and \( \text{slt2/mpk1} \) mutant strains, or it could have activated a subset of CWI pathway transcription factors.

**RESPONSE TO OXIDATIVE STRESS**

Oxidative stress occurs in the natural environment with exposure to aerobic conditions and UV light. For pathogenic fungi, oxidative stress is manifested through the host’s production of reactive oxygen species, hydrogen peroxide, and hydroxyl radicals by phagocytic cells. Oxide radicals are also the consequence of normal cell metabolism. The first line of defense by fungi to prevent cell damage from exogenous oxide radicals is the cell wall. The cell senses oxide stress, and reactive measures are taken by signal transduction pathways to protect fungi from the adverse environment. A key defense against oxidative stress is the HOG pathway (1, 2, 69). The CWI pathway provides additional oxide defense through cross talk with the HOG pathway. Oxidative stress resistance is also dependent upon the transcription factors Skn7, Yap1, Msn2, and Msn4. These transcription factors control the expression of numerous genes in response to oxidative stress. Orthologous sequences for the transcription factors are shared between the budding yeast \( S. \text{cerevisiae} \), the fission yeast \( S. \text{pombe} \), and the pathogens \( C. \text{albicans}, \text{Candida glabrata} \), and \( U. \text{maydis} \) (11). A number of other enzymes are produced to combat oxide attack, including superoxide dismutases, catalases, peroxidases, and glutathione peroxidases.

In \( S. \text{cerevisiae} \), diamide and hydrogen peroxide exposure aid in the effort to deduce the CWI pathway response to oxide stress. Diamide depletes glutathione and oxide thiols. Hydrogen peroxide promotes lipid peroxidation, protein oxidation, and DNA damage. Diamide or hydrogen peroxide is capable of activating the dual phosphorylation of \( \text{Slt2/Mpk1} \), although with differing kinetic activity. Diamide-induced oxide stress is sensed in \( S. \text{cerevisiae} \) by the transmembrane proteins \( \text{Mid2} \) and \( \text{Wsc1} \) (79). In contrast, hydrogen peroxide does not require one of the known CWI stress sensors to provoke the phosphorylation of \( \text{Slt2/Mpk1} \) and is suspected to elicit an intracellular means of activation (79). Exposure of a \( \text{rom2} \) mutant strain to either diamide or hydrogen peroxide results in diminished phosphorylation of \( \text{Slt2/Mpk1} \). \( \text{Pck1} \) is also required, but the MAPK module downstream of \( \text{Pck1} \) is not vital to the cellular response in defense of oxidative stress (79). Thus, the activation of the CWI pathway by oxidative stress by either inducer requires the upper part of the CWI pathway for cell survival (79).

A similar oxidative stress response occurs in \( C. \text{neoformans} \) (26). In this case, phosphorylation of \( \text{Mpk1} \) indicates PKC pathway activity in response to oxidative stress elicited from diamide or hydrogen peroxide. \( \text{PKC1} \) is required to defend against oxidative stress. However, as seen in \( S. \text{cerevisiae} \), the \( C. \text{neoformans} \) MAPK module is dispensable for survival against oxidative stress (26). Those authors suggest that an alternative pathway extends from \( \text{Pck1} \) to provide for protection against oxidative stress (26). Thus, there is an alternative function of \( \text{Pck1} \) other than activation of the MAPK module that leads to an oxide-induced phosphorylation of \( \text{Mpk1} \). The greater severity of the \( \text{pck1} \) mutation compared to the \( \text{beck1} \) mutation evaluated for \( C. \text{neoformans} \) by Gerik and colleagues suggests further roles for \( \text{Pck1} \) that are not facilitated through the MAPK module (27), a hypothesis that has been postulated by others in regard to the role of \( \text{PKC} \) (39). Indeed, analysis of the synthetic genetic network around \( S. \text{cerevisiae PKC1} \) reveals several genes that influence the CWI pathways but are not themselves CWI pathway genes (41). Additional evidence for an extended role of \( \text{Pck1} \) beyond the activation of the MAPK module is the lethality of \( \text{pck} \) mutations in \( S. \text{cerevisiae} \), indicating that \( \text{PKC1} \) is essential (48), whereas mutations of the MAPK module are not lethal. Likewise, deletion of \( \text{PCK1} \) of \( C. \text{neoformans} \) is lethal unless the cells are protected by an osmotic stabilizer such as sorbitol, whereas the MAPK module mutants show no growth defects under regular medium conditions (26).

The CWI pathway is also activated by oxidative stress in \( C. \text{alicins} \) (54). \( \text{Mpk1} \) is phosphorylated in the presence of hydrogen peroxide, but interestingly, it is not required for cell viability in the presence of hydrogen peroxide (54). Phosphorylation of \( \text{Mpk1} \) with oxide stress is \( \text{Pck1} \) dependent (as seen with other fungi) except when diamide is used as the inducing agent. In the case of diamide, \( \text{Mpk1} \) phosphorylation is \( \text{Pck1} \) independent. The phosphorylation event is not simply a result of cell wall damage, because phosphorylation also occurs in the presence of the osmotic stabilizer sorbitol (54). The phosphorylation of \( \text{Mpk1} \) is also \( \text{Pbs2} \) and \( \text{Hog1} \) dependent, indicating a cross talk event between the CWI and HOG pathways in response to oxide stress (2).

A slightly different oxide response was observed with \( A. \text{fumigatus} \). The CWI pathway of \( A. \text{fumigatus} \) differentiates between the types of oxidative stressor. An \( \text{mpkA} \) (a homolog of \( \text{SLT2/MPK1} \)) deletion strain is sensitive to diamide and menadione, which generates superoxides, but resistant to hydrogen peroxide (76). Mechanistically there is similarity in that \( A. \text{fumigatus} \), like \( C. \text{albicans} \), does not require MAPK for cell survival when oxide stress is presented as hydrogen peroxide, indicating that various oxides elicit different cellular responses.

**RESPONSE TO NITROSATIVE STRESS**

The CWI pathway is not characterized to respond to nitrosative stress. However, \( C. \text{neoformans} \) presents a thus far unique situation by responding to nitrosative stress by activating the CWI pathway. In vitro exposure of \( C. \text{neoformans} \) to nitrosative stress in the form of sodium nitrite results in phosphorylation of \( \text{Mpk1} \). \( \text{PKC1} \) is vital to cell survival during nitrosative stress (26). Further examination of the cell response to nitrosative stress is needed to determine if the CWI pathway facilitates a response to nitrosative stress for other fungi. If the CWI pathway is not the primary means of responding to nitrosative stress in fungal cells, perhaps the CWI response to nitrosative stress observed in \( C. \text{neoformans} \) indicates a cross talk event with another pathway.

**RESPONSE TO ANTIFUNGAL AGENTS AND CHEMICAL COMPOUNDS**

There are several types of antifungal agents available for treating fungal diseases. Echinocandin antifungals such as caspofungin target (1,3)-\( \beta\)-D glucan synthase. When \( C. \text{albicans} \) cells are exposed to caspofungin, the chitin biosynthesis genes
CHS2 and CHS8 are upregulated (80), among other genes. In an slt2/mpk1 mutant strain, CHS2 and CHS8 exhibit only marginal increases in expression upon caspofungin exposure compared to a wild-type strain exposed to caspofungin, which exhibits significant increases in CHS2 and CHS8 expression (80). Thus, the CWI pathway is required for chitin biosynthesis in response to stress induced by caspofungin.

Additional compounds have been identified to elicit CWI stress; these compounds are not used to treat fungal infections in patients but can provide additional information about the pathway interactions that are induced to prevent or repair CWI damage. Zymolyase is an enzyme that affects the cell wall by hydrolyzing the β-(1,3)-glucan network. The cellular response to Zymolyase involves both the CWI and HOG pathways. Exposure of *S. cerevisiae* to Zymolyase results in the increased expression of *CRH1*, which encodes a glycosylphosphatidylinositol cell wall protein induced under cell wall damage growth conditions. Crh1 protein expression induced by Zymolyase is Hog1 and Slt2/Mpk1 dependent (6).

Zymolyase stimulation of the HOG pathway is through the Sho1-dependent branch. The genes *SHO1, STE11*, and *HOG1*, but not *SSK1* or *SSK2/SSK2*, are required in the response to Zymolyase treatment (6, 23). Zymolyase induction of Slt2/Mpk1 phosphorylation from the CWI pathway requires *PKC1, BCK1, MKK1/MKK2*, and *MPK1* but not the upstream element *ROM2* or the stress sensors *MID2* and *WSC1* (6, 23). This type of activation indicates that the MAPK module is required for Slt2/Mpk1 phosphorylation by Zymolyase but not the upstream portion of the CWI pathway or the stress sensors. Thus, it is not a linear activation of Slt2/Mpk1 through the CWI pathway but rather the influence of a lateral element. The evidence provided suggests that the response to Zymolyase is mediated through the Sho1-dependent branch of the HOG pathway and that a cross talk event which initiates the MAPK module of the CWI pathway occurs. Bermejo et al. (6) demonstrated that Hog1 phosphorylation was Slt2/Mpk1 independent. Further, they demonstrated that Slt2/Mpk1 was overphosphorylated in response to Zymolyase and that the overphosphorylation event was Hog1 dependent. Thus, the HOG pathway functions to activate the MAPK module of the CWI pathway in the presence of Zymolyase-induced cell stress (Fig. 6).

A different manifestation of the cellular CWI response is observed with exposure to the chemical caffeine. Although it is not likely that caffeine is a stress-inducing agent in the natural environment, it does provide insight into fungal cell signaling interactions. It has not yet been fully elucidated how caffeine affects cells, but it does indeed act as a cell wall stress agent. Kuranda and colleagues (43) identified that caffeine induces a CWI response in *S. cerevisiae* through phosphorylation of Slt2/Mpk1 in a Tor-dependent manner. Activation of the CWI pathway is independent of the stress sensors *WSC1* and *MID2*. It is, however, dependent on other members of the CWI pathway, including *ROM2, RHO1, BCK1*, and *MKK1* (43, 52). Thus, there is potentially a sensor other than Wsc1 and Mid2 to initiate the signal cascade or a lateral influence.

The chemical activation of the CWI pathway is not a uniform response but rather depends on the chemical that elicits the response. We have seen that the CWI pathway can interact with the HOG pathway or TOR signaling, and it is possible that other signaling mechanisms could be recruited by different chemical stresses.

**CONCLUSION**

Maintaining CWI is a difficult task given the numerous external influences that challenge fungal cells. Plant and animal hosts in particular present adverse conditions to fungi in multiple ways simultaneously through changes in pH, temperature, and nutrient availability and by actively combating fungi with oxidative radicals and phagocytosis of fungal cells. It would be prudent for fungi to maintain constitutively high activation of the CWI pathway at all times, but this is not the case. Perhaps it is too costly in energy and cellular resources to maintain the CWI pathway at defensive levels at all times. It is therefore strategically activated when stress conditions or growth warrants the activity.

It would be difficult for a single pathway to correctly attend to each of the very different adverse conditions to which a fungal cell is exposed in a linear manner. Therefore, the CWI pathway is a multifunctional pathway that receives lateral influences at many points along the signaling cascade. Lateral interactions encompass Tor signaling, calcineurin signaling, the HOG pathway, cAMP-PKA signaling, and input from proteins not yet elucidated. This complicated network of signaling enables the CWI pathway to be activated by numerous types of stressors and provides the necessary responses to fortify the cell and preserve fungal cell viability.

Although much of the information presented in this article is based on pathway interactions learned through research involving the model yeast *S. cerevisiae*, with additional information provided through research on several pathogenic fungi,
much more is likely to be revealed about the signal transduction interactions that lead to CWI maintenance from numerous other fungi that cope with a plethora of hostile environments. Further, there is still much that can be learned through fungal studies by creating double knockout and overexpression strains to elucidate how the pathways presented in this article interact to cope with stress conditions.

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