### NAOSITE: Nagasaki University's Academic Output SITE



Title	Dissection of Ire1 Functions Reveals Stress Response Mechanisms Uniquely Evolved in Candida glabrata
Author(s)	Miyazaki, Taiga; Nakayama, Hironobu; Nagayoshi, Yohsuke; Kakeya, Hiroshi; Kohno, Shigeru
Citation	PLoS Pathogens, 9(1), e1003160; 2013
Issue Date	2013-01-31
URL	http://hdl.handle.net/10069/31913
Right	© 2013 Miyazaki et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

This document is downloaded at: 2020-09-18T13:48:36Z



# Dissection of Ire1 Functions Reveals Stress Response Mechanisms Uniquely Evolved in *Candida glabrata*

Taiga Miyazaki<sup>1</sup>\*, Hironobu Nakayama<sup>2</sup>, Yohsuke Nagayoshi<sup>1</sup>, Hiroshi Kakeya<sup>1</sup>, Shigeru Kohno<sup>1</sup>

1 Department of Molecular Microbiology and Immunology, Nagasaki University School of Medicine, Nagasaki, Japan, 2 Faculty of Pharmaceutical Sciences, Suzuka University of Medical Sciences, Mie, Japan

#### **Abstract**

Proper protein folding in the endoplasmic reticulum (ER) is vital in all eukaryotes. When misfolded proteins accumulate in the ER lumen, the transmembrane kinase/endoribonuclease Ire1 initiates splicing of *HAC1* mRNA to generate the bZIP transcription factor Hac1, which subsequently activates its target genes to increase the protein-folding capacity of the ER. This cellular machinery, called the unfolded protein response (UPR), is believed to be an evolutionarily conserved mechanism in eukaryotes. In this study, we comprehensively characterized mutant phenotypes of *IRE1* and other related genes in the human fungal pathogen *Candida glabrata*. Unexpectedly, Ire1 was required for the ER stress response independently of Hac1 in this fungus. *C. glabrata* Ire1 did not cleave mRNAs encoding Hac1 and other bZIP transcription factors identified in the *C. glabrata* genome. Microarray analysis revealed that the transcriptional response to ER stress is not mediated by Ire1, but instead is dependent largely on calcineurin signaling and partially on the SIt2 MAPK pathway. The loss of Ire1 alone did not confer increased antifungal susceptibility in *C. glabrata* contrary to UPR-defective mutants in other fungi. Taken together, our results suggest that the canonical Ire1-Hac1 UPR is not conserved in *C. glabrata*. It is known in metazoans that active Ire1 nonspecifically cleaves and degrades a subset of ER-localized mRNAs to reduce the ER load. Intriguingly, this cellular response could occur in an Ire1 nuclease-dependent fashion in *C. glabrata*. We also uncovered the attenuated virulence of the *C. glabrata* Δ*ire1* mutant in a mouse model of disseminated candidiasis. This study has unveiled the unique evolution of ER stress response mechanisms in *C. glabrata*.

Citation: Miyazaki T, Nakayama H, Nagayoshi Y, Kakeya H, Kohno S (2013) Dissection of Ire1 Functions Reveals Stress Response Mechanisms Uniquely Evolved in Candida glabrata. PLoS Pathog 9(1): e1003160. doi:10.1371/journal.ppat.1003160

Editor: Scott G. Filler, David Geffen School of Medicine at University of California Los Angeles, United States of America

Received September 14, 2012; Accepted December 12, 2012; Published January 31, 2013

**Copyright:** © 2013 Miyazaki et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This research was supported by a grant from the Global Centers of Excellence Program, Nagasaki University, Grant-in-Aid for Scientific Research (JSPS KAKENHI no. 24791027 to TM and no. 21390305 to SK) from the Japanese Ministry of Education, Culture, Sports, Science and Technology, and a grant of the Ueda award from the Japanese Society of Chemotherapy to TM. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript

1

Competing Interests: The authors have declared that no competing interests exist.

\* E-mail: taiga-m@nagasaki-u.ac.jp

#### Introduction

In eukaryotic cells, the majority of secretory and transmembrane proteins are folded and modified in the lumen of the endoplasmic reticulum (ER). Contingent on proper folding, they are either transported to the Golgi apparatus or degraded [1]. Impairment of these vital cellular machines can be caused by various factors, such as chemical compounds and mutations in genes involved in ER quality control, resulting in the accumulation of unfolded or misfolded proteins in the ER, collectively called ER stress [2,3]. Unsolved ER stress has detrimental consequences for eukaryotic cells. For example, in humans, ER stress is implicated in the pathology of various diseases including metabolic disease, inflammation, neurodegenerative disorders, and cancer [4,5]. It has also been revealed in several pathogenic fungi that ER quality control is important for antifungal resistance and virulence [6,7,8,9,10,11].

In response to ER stress, eukaryotic cells activate a signaling pathway, termed the unfolded protein response (UPR), to induce a comprehensive gene expression program that adjusts the proteinfolding capacity of the ER (reviewed in [2,12]). In Saccharomyces cerevisiae, the ER-resident transmembrane stress transducer Ire1, which contains an ER luminal sensing domain and cytosolic protein kinase/endoribonuclease domains, is responsible for

detecting unfolded proteins and triggering the UPR [13,14]. Ire1 is activated by direct interaction with the unfolded proteins, resulting in the oligomerization of its luminal domain. Signaling is transduced to the cytosolic domain through the sequential steps of autophosphorylation and oligomerization of the endoribonuclease domain, leading to the activation of the Irel ribonucrease [15,16]. Active Ire1 cleaves the HAC1 mRNA to excise the intron, allowing translation of the basic-leucine zipper (bZIP) transcription factor Hac1 that subsequently induces transcription of the UPR target genes [17,18]. ER-stressed cells attempt to reduce the load of abnormally folded proteins in the ER by facilitating protein folding (e.g. upregulation of genes encoding ER-resident chaperones and protein-modifying enzymes) and by translocating misfolded proteins from the ER to the cytosol where they are degraded by the proteasome. The latter mechanism is called ERassociated protein degradation (ERAD) (reviewed in [3]). An alternative mechanism of degradative response is autophagy, which degrades organelles including damaged ER. In addition to ER-resident chaperones and protein-modifying enzymes, many of the components that mediate ERAD and autophagy have also been identified as UPR targets [19,20,21].

In pathogenic fungi, the molecular basis of ER quality control has been poorly understood, but several recent studies in *Aspergillus* 

#### **Author Summary**

The majority of secretory and transmembrane proteins are structurally matured in the endoplasmic reticulum (ER). The accumulation of misfolded proteins in the ER (ER stress) activates the ER-resident stress transducer Ire1, which has two distinct outputs: the unfolded protein response (UPR) and regulated Ire1-dependent decay (RIDD). The UPR is a transcriptional response to increase the protein folding capacity of the ER. RIDD induces degradation of ER-localized mRNAs to reduce the ER load. To date, the UPR has been believed to be an evolutionarily conserved pathway in almost all eukaryotic species, while RIDD has been found only in metazoans. Recent studies in several pathogenic fungi revealed that the UPR is implicated in antifungal resistance and virulence, and thus it has attracted attention as a therapeutic target. Here, we demonstrate that the important fungal pathogen Candida alabrata has lost the canonical UPR, but instead possesses the RIDD pathway and is relatively tolerant to ER stress. The transcriptional response to ER stress was dependent mainly on calcium signaling mediated by the protein phosphatase calcineurin in C. glabrata. Our results provide novel insights into ER quality control mechanisms and are useful for understanding evolutionary biology and the development of antifungal agents targeting the UPR.

fumigatus and Cryptococcus neoformans have found that Ire1 and Hac1 homologs are key components of the UPR and are indeed required for the ER stress response [7,8,10]. Interestingly, these studies have also discovered that the UPR is implicated in fungal pathogenicity and antifungal resistance. In A. fumigatus, the loss of HacA/Hac1 results in increased susceptibility to caspofungin, amphotericin B, and azole antifungals, and decreased virulence in mouse models of invasive aspergillosis [10]. Recently, the necessity of IreA/Ire1 for A. fumigatus virulence has also been reported [8]. C. neoformans mutant strains lacking Irel or its downstream transcription factor Hxl1 display increased azole susceptibility, failure to grow at 37°C, and avirulence in a mouse model of systemic cryptococcosis [7]. It is also known in Candida albicans that Hac1 plays a role in hyphal development [11], and a  $\Delta ire1$  mutant is hypersusceptible to caspofungin [6]. These observations shed light on the UPR as an attractive target for the development of novel antifungal therapies.

Candida glabrata has emerged as an important fungal pathogen due in part to its intrinsic or rapidly acquired resistance to azole antifungals such as fluconazole [22,23]. In addition, recent surveillance data have revealed an increase of *C. glabrata* clinical isolates that display resistance to not only azoles, but also echinocandin-class antifungals [24,25]. Considering the limitations of the currently available antifungals in clinical settings, there is an urgent need to develop an effective antifungal strategy for a broad range of fungal pathogens, including *C. glabrata*.

Since how C. glabrata cells deal with ER stress has not been explored, we first functionally characterized the C. glabrata Irel and Hacl orthologs. It has been believed that the UPR mediated by the Irel-Hacl linear pathway is evolutionarily conserved in most eukaryotic species, but surprisingly, we found that C. glabrata Irel plays a role in the ER stress response in a Hacl-independent manner, despite the presence of an apparent HAC1 ortholog. The present study revealed that C. glabrata has lost the canonical Irel-Hacl UPR, but has developed alternative mechanisms for ER quality control. In addition, our comprehensive analyses of  $\Delta irel$  mutant phenotypes revealed significant diversities of Irel-mediated stress response mechanisms between C. glabrata and other fungi.

Here, we describe the unique evolution of ER quality control systems in *C. glabrata*.

#### Results

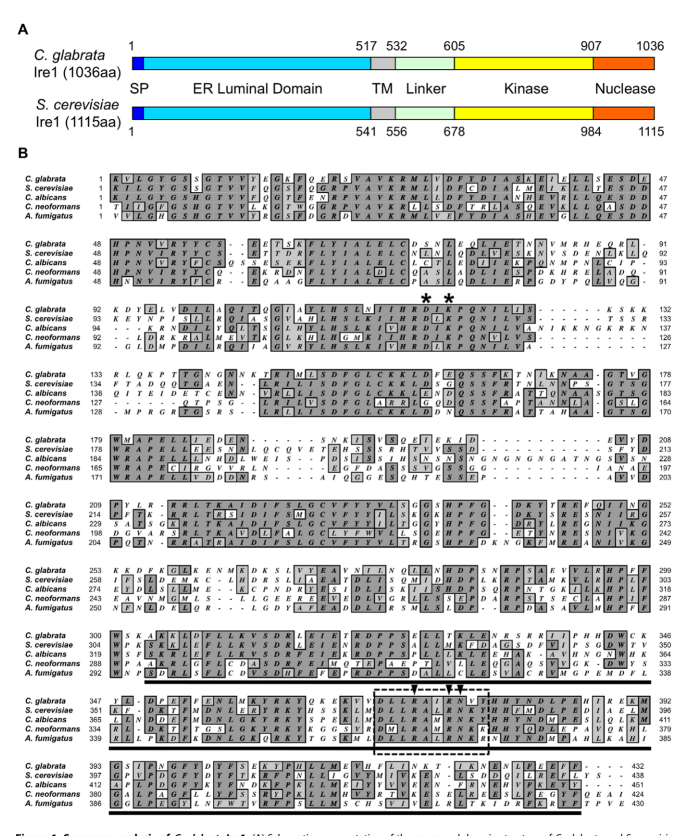
### Comparison of growth ability in the presence of ER stress between *S. cerevisiae*, *C. neoformans*, and *Candida* spp

The ability of fungal cells to cope with ER stress was assessed by monitoring cell growth in the presence and absence of two wellknown ER stress inducers that interfere with protein folding in the ER by different mechanisms: tunicamycin (TM), an N-linked glycosylation inhibitor, and dithiothreitol (DTT), an inhibitor of disulfide bond formation. Compared to S. cerevisiae and C. neoformans wild-type strains, C. glabrata was relatively tolerant to both TM and DTT independently of its strain backgrounds and culture media, although S. cerevisiae displayed strain dependent susceptibilities (Figure S1). Other representative Candida species also exhibited higher tolerance to these agents than S. cerevisiae and C. neoformans with intriguing exceptions: Candida krusei was highly susceptible to TM, but not to DTT, while Candida tropicalis was hypersusceptible to DTT, but not to TM. These results imply that diverse aspects of ER stress response mechanisms may exist even in closely related yeast species. In this report, the following studies focused on C. glabrata, which is phylogenetically close to S. cerevisiae, but has gained increased tolerance to ER stress and pathogenic potential to humans.

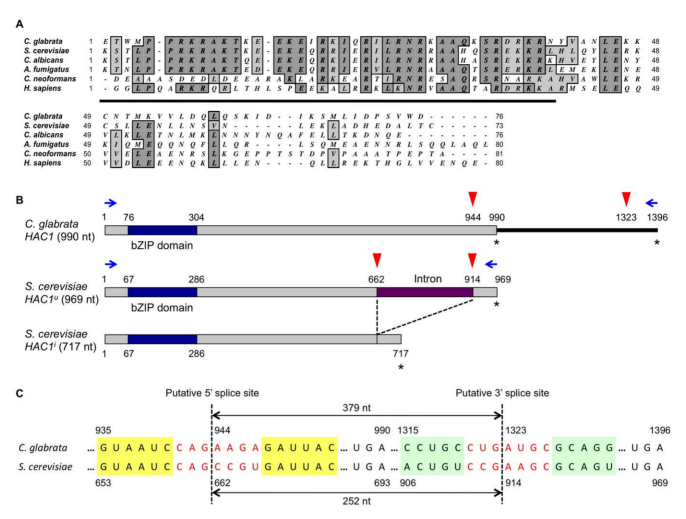
### Identification and structural characterization of the *IRE1* and *HAC1* orthologs in *C. glabrata*

The C. glabrata IRE1 and HAC1 orthologs were identified by BLASTp searches using the NCBI (http://www.ncbi.nlm.nih. gov/BLAST/) and Genolevures (http://cbi.labri.fr/Genolevures/ elt/CAGL) databases. The respective amino acid sequences of S. cerevisiae IRE1 (YHR079c) and HAC1 (YFL031w) were used as queries. The deduced amino acid sequence of C. glabrata IRE1 No.: XP 446111, accession Genolevures CAGL0F03245g) displayed 53.6% similarity and 35.8% identity with that of S. cerevisiae IRE1. As presented schematically in Figure 1A, C. glabrata Irel consists of typical Irel-domain structures [16,26], including an N-terminal hydrophobic signal sequence, an ER luminal domain, a transmembrane domain, and a serine-threonine protein kinase followed by a nuclease domain. The signal sequence and the ER luminal domain are important for ER localization and unfolded protein sensing, respectively [14,27,28,29]. The most conserved segment is the cytoplasmic C-terminal region containing the protein kinase linked to the nuclease domain (Figure 1B). The protein kinase domain is required for autophosphorylation and concomitant activation of the nuclease domain, which provides the endoribonuclease activity [30,31,32]. C. glabrata IRE1 is syntenic with S. cerevisiae IRE1 situated on chromosome VIII, but the 3' region of C. glabrata IRE1 is orthologous to S. cerevisiae chromosome IV (Figure S2). This suggests that a genomic arrangement between S. cerevisiae chromosomes IV and VIII has occurred to evolve the IRE1surrounding region on chromosome F in C. glabrata.

A BLASTp search using the NCBI and Genolevures databases also identified a single *HAC1* ortholog (NCBI accession No.: XP\_448761, Genolevures ID: CAGL0K12540g) in the *C. glabrata* genome, exhibiting 36.9% similarity and 24.2% identity with the deduced amino acid sequence of the unspliced form of *S. cerevisiae HAC1*. It has been known that functional orthologs of transcription factors often display limited sequence similarity outside their DNA-binding domains [11]. Indeed, *C. glabrata HAC1* is not highly conserved at the overall sequence level, but contains a bZIP



**Figure 1. Sequence analysis of** *C. glabrata* **Ire1.** (A) Schematic representation of the conserved domain structure of *C. glabrata* and *S. cerevisiae* Ire1. Abbreviations: SP, signal peptide; and TM, transmembrane. (B) Sequence alignment of the kinase and nuclease domains of fungal Ire1 orthologs. The asterisk indicates the conserved two catalytic residues in the nucleotide-binding pocket of Ire1 kinase (D797 and K799 in *S. cerevisiae* and D723 and K725 in *C. glabrata*). The predicted nuclease domain is underlined. Ten residues (boxed with dotted line) including the highly conserved three-nuclease active sites (arrowheads) are deleted in *C. glabrata IRE1*-ND. GenBank accession number: *Candida glabrata* Ire1, XP\_446111; *Saccharomyces cerevisiae* Ire1, NP\_011946; *Candida albicans* Ire1, XP\_717532; *Cryptococcus neoformans* Ire1, XP\_568837; and *Aspergillus fumigatus* IreA, AEQ59230. doi:10.1371/journal.ppat.1003160.g001



**Figure 2. Sequence analysis of** *C. glabrata* **Hac1.** (A) Sequence alignment of bZIP domains in Hac1 homologs from fungi and humans. The DNA binding domain is underlined. GenBank accession number: *Candida glabrata* Hac1, XP\_448761; *Saccharomyces cerevisiae* Hac1, NP\_116622; *Candida albicans* Hac1, XP\_718538; *Aspergillus fumigatus* HacA, XP\_748727; *Cryptococcus neoformans* Hxl1, XP\_568439; and *Homo sapiens* Xbp1, NP\_005071. (B) A schematic representation of the putative splice sites and ORF lengths in *C. glabrata* and *S. cerevisiae HAC1*. The *HAC1* ORFs and bZIP domains were shown in grey and blue boxes, respectively. In *S. cerevisiae*, the intron (purple box) in the uninduced form of *HAC1* (*HAC1*<sup>1</sup>). Red arrowheads and asterisks indicate putative splice sites and stop codons, respectively. Blue arrows indicate primers that were used for RT-PCR assays to examine *HAC1* splicing. (C) Alignment of the RNA sequence surrounding the predicted intron in *C. glabrata* and *S. cerevisiae HAC1*. In *S. cerevisiae*, *HAC1* mRNA is known to form stem-loop structures that enclose the intron between the two loops of seven residues (shown in red) held in place by short stems (yellow and green boxes). The predicted 5' and 3' splice sites and intron lengths in the *S. cerevisiae* and *C. glabrata HAC1* mRNAs are indicated. doi:10.1371/journal.ppat.1003160.g002

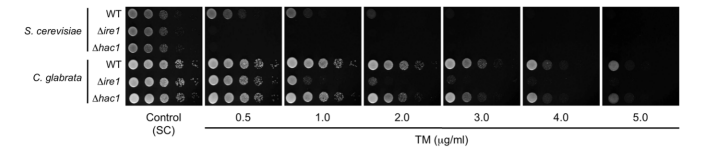
domain at the N-terminal region. The putative DNA binding domain of *C. glabrata* Hac1 displays strong sequence similarity to those of other Hac1 homologs (Figure 2A). In addition, there is a clear syntenic relationship between *S. cerevisiae* and *C. glabrata HAC1* loci (Figure S2). Based on these sequence similarities and synteny conservation, we considered the genes CAGL0F03245g and CAGL0K12540g to be *C. glabrata IRE1* and *HAC1*, respectively.

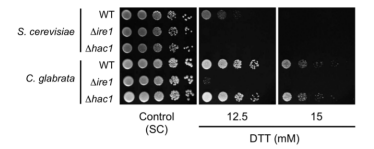
It has been known in *S. cerevisiae* [33,34,35] that the uninduced form of *HAC1* (*HAC1*") contains the 252 nucleotides intron, which is excised by Ire1 to generate the induced form of *HAC1* (*HAC1*") (Figure 2B). The well-conserved Ire1 recognition motifs, CNG'CNGN and CNG'AAGC at the 5' and 3' boundaries of the intron, respectively [36], locate in the loop regions of the *HAC1* mRNA stem-loop structures [34,37]. Based on the conservation of common features around the predicted intron (Figure 2C), *C. glabrata HAC1* mRNA is also likely to form stem-loop structures

[36]. If excision of the predicted intron in *C. glabrata HAC1* mRNA occurs, it changes the last 15 amino acids of the putative ORF and extends it by only 9 amino acids (Figures 2B and 2C). However, *C. glabrata HAC1* has mutations in the consensus sequences of the Irel recognition motifs at both splice sites: a C to A mutation at the 4th position of the 5' splice site and an A to U mutation at the 5th position of the 3' splice site (Figure 2C).

### Ire1, but not Hac1, is required for ER stress response in *C. glabrata*

To examine involvement of Ire1 and Hac1 in the ER stress response in C. glabrata, we constructed a mutant of each in which the entire open reading frame (ORF) of IRE1 or HAC1 was deleted, and examined growth on plates containing TM and DTT (Figure 3). Both S. cerevisiae  $\Delta ire1$  and  $\Delta hac1$  mutants were used as controls and they exhibited severe growth defects in the presence of TM or DTT. In C. glabrata, the  $\Delta ire1$  mutant displayed





**Figure 3. Growth assay in the presence of ER stress-inducing agents.** Logarithmic-phase cells of *S. cerevisiae* and *C. glabrata* strains were adjusted to  $2\times10^7$  cells/ml, and then 5 μl of serial 10-fold dilutions were spotted onto synthetic complete (SC) plates containing either tunicamycin (TM) or dithiothreitol (DTT) at the indicated concentrations. Plates containing TM and DTT were incubated at 30°C for 24 and 48 h, respectively. *S. cerevisiae* strains: WT, BY4742; Δ*ire1*, BY4742Δire1; and Δ*hac1*, BY4742Δhac1. *C. glabrata* strains: WT, CBS138; Δ*ire1*, TG121; and Δ*hac1*, TG141. doi:10.1371/journal.ppat.1003160.g003

decreased tolerance to both TM and DTT compared with the wild-type strain, whereas surprisingly, the  $\Delta hac1$  mutant exhibited wild-type growth in the presence of these ER stress-inducing agents (Figure 3). Southern blot analysis demonstrated that HAC1 was truly absent and there was no ectopic integration of the deletion construct in the  $\Delta hac1$  mutant (Figure S3). The results indicate that Ire1, but not Hac1, is required for the ER stress response in C. glabrata. This is quite a contrast to S. cerevisiae, which is dependent equally on Ire1 and Hac1 to survive ER stress [38], implying that C. glabrata may possess a unique mechanism, which is mediated by Ire1 independently of Hac1, to cope with ER stress.

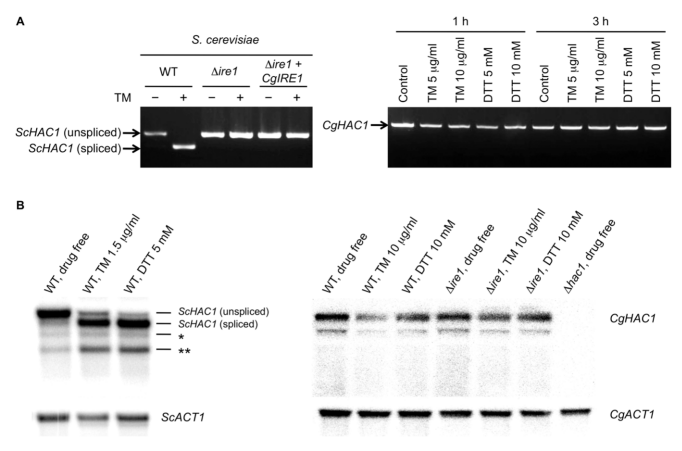
#### Ire1 does not cleave HAC1 mRNA in C. glabrata

Ire1-mediated splicing of HAC1/XBP1 mRNA was first uncovered in S. cerevisiae [32], and similar mechanisms in HAC1/XBP1 homologs were later described in some fungi, including Trichoderma reesei, Aspergillus nidulans [39], C. albicans [11], Pichia pastoris [40], Yarrowia lipolytica [41], A. fumigatus [8,10], and C. neoformans [7], as well as in metazoans including Caenorhabditis elegans, mice [37,42], humans [43], and Drosophila melanogaster [44]. We examined HAC1 mRNA splicing in ER-stressed S. cerevisiae and C. glabrata cells by RT-PCR using the *HAC1* specific primer pairs denoted in Figure 2B. As expected, S. cerevisiae HAC1 mRNA was spliced in response to treatment with 1.5 µg/ml TM in wild-type control but not in the  $\Delta \textit{ire1}$ strain (Figure 4Å, left panel). As C. glabrata wild-type cells were relatively tolerant to both TM and DTT (Figure S1), they were treated with higher concentrations of TM and DTT for 1 and 3 h. However, contrary to S. cerevisiae, splicing of HAC1 mRNA was not induced in C. glabrata under any of the conditions tested in this study (Figure 4A, right panel). The absence of a HAC1 splicing event was also verified by sequencing the RT-PCR products of the entire HAC1 mRNA (data not shown), suggesting that Hac1 is not a downstream target of Ire1 in C. glabrata.

To further confirm this aspect, we also performed Northern blot analysis using the probes directed against the 5' region of the HAC1 ORF in S. cerevisiae and C. glabrata. In S. cerevisiae, HAC1 mRNA was efficiently spliced upon ER stress induced by 1.5 µg/ ml TM or 5 mM DTT (Figure 4B, left panel). In addition to the unspliced and spliced HAC1 mRNAs, detection of two smaller bands (indicated with asterisks in Figure 4B) were consistent with previous reports [17,45,46], although it is unclear whether these bands correspond to splicing intermediates or dead-end products [17,46]. The C. glabrata wild-type and  $\Delta ire1$  cells were incubated under the normal conditions and in the presence of 10 µg/ml TM or 10 µM DTT for 3 h. Under all conditions tested, the C. glabrata wild-type strain somehow exhibited 2 bands but these bands were also detected similarly in the  $\Delta ire1$  mutant (Figure 4B, right panel), indicating that it was not due to Irel-mediated splicing. These results and the phenotypic differences between the  $\Delta ire1$  and  $\Delta hac1$ mutants strongly suggest that C. glabrata Irel plays a role in the ER stress response in a Hacl-independent manner.

### Exploration of other bZIP transcription factors as potential downstream targets of Ire1 in *C. glabrata*

To address the possibility that a HAC1-like gene encoding a bZIP transcription factor may exist as a downstream target of Irel in C. glabrata, we performed a low-stringent BLASTp search (e-val $ue = 10^{-2}$ ) using only the bZIP domain sequences of S. cerevisiae Hacl, C. neoformans Hxll, and Homo sapiens Xbpl as described previously [7]. In addition to HAC1, the following eight putative genes were identified in the C. glabrata genome (Table S1): CAGL0H04631g (YAPI), CAGL0J06182g (SKOI), CAGL0K02585g (YAP3), CAGL0M10087g (YAP3), CAGL0L02475g (GCN4), CAGL0F03069g (CAD1/YAP2), CAGL0F01265g (YAP7), and CAGL0M08800g (YAP6). We sequenced the RT-PCR products of these mRNAs, which were obtained from C. glabrata wild-type cells treated with TM at concentrations of 1.5 and 5  $\mu g/ml$  for 3 h, but none of them displayed evidence of a splicing event (data not shown). The results suggest that another Hac1 homolog is unlikely to exist in C. glabrata.



**Figure 4. Assays for** *HAC1* **mRNA splicing.** (A) RT-PCR analysis. The *S. cerevisiae* strains containing either empty vector or pRS415-ADH-CgIRE1, in which *C. glabrata IRE1* was expressed under the control of the *S. cerevisiae ADH1* promoter, were incubated in SC-leu broth in the presence and absence of 1.5 μg/ml tunicamycin (TM) for 3 h. *S. cerevisiae* strains: WT, BY42-1;  $\Delta$ ire1, BY42l-1; and  $\Delta$ ire1+CgIRE1, BY42l-2. The *C. glabrata* wild-type strain CBS138 was incubated in SC broth in the presence and absence of TM and dithiothreitol (DTT) at the indicated concentrations for 1 and 3 h. RT-PCR products of the entire *HAC1* mRNA in *S. cerevisiae* (left panel) and *C. glabrata* (right panel) were electrophoresed on a 1% agarose gel. (B) Northern blot analysis. *S. cerevisiae* WT (BY4742) cells were treated with 1.5 μg/ml TM or 5 mM DTT for 1 h (left panel). *C. glabrata* cells were treated with 10 μg/ml TM or 10 mM DTT for 3 h (right panel). Both *ScHAC1* and *CgHAC1* probes were generated from the 5' regions of the *HAC1* ORFs. The same blot was probed for *HAC1* mRNA, stripped, and then probed for *ACT1* mRNA. The asterisks indicate potential splicing intermediates (\*: 5' exon plus the intron; \*\*: 5' exon alone). *C. glabrata* strains: WT, CBS138;  $\Delta$ ire1, TG121; and  $\Delta$ hac1, TG141. doi:10.1371/journal.ppat.1003160.g004

### Complementation assays of the IRE1 and HAC1 orthologs between S. cerevisiae and C. glabrata

To investigate whether *C. glabrata* Ire1 could complement *S. cerevisiae* Ire1 function, *C. glabrata* IRE1 was expressed under the *S. cerevisiae* ADH1 promoter in the *S. cerevisiae* Δire1 mutant. After treatment with 1.5 μg/ml TM for 3 h, a spliced form of *S. cerevisiae* HAC1 was generated in the wild-type strain, but not in the Δire1 strain regardless of the presence of *C. glabrata* IRE1 (Figure 4A). The absence of a HAC1 splicing event in the *S. cerevisiae* Δire1 mutant containing *C. glabrata* IRE1 was also verified by sequencing analyses of the RT-PCR products (data not shown). As expected, *C. glabrata* Ire1 did not rescue the growth of the *S. cerevisiae* Δire1 mutant in the presence of TM (Figure 5A). These results indicate that *C. glabrata* Ire1 is unable to initiate the UPR due to its inability to cleave the HAC1 mRNA in *S. cerevisiae*, accounting for the growth defects of the *S. cerevisiae* Δire1 mutant containing *C. glabrata* IRE1 in the presence of TM.

Although *HAC1* was dispensable for *C. glabrata* cells to cope with ER stress (Figure 3), the fact that the bZIP domain is conserved in this gene prompted us to investigate whether *C. glabrata* Hac1 behaves as a transcription factor similarly to *S. cerevisiae* Hac1. To address this question, *C. glabrata HAC1* was constitutively expressed

under the ADH1 promoter in the S. cerevisiae  $\Delta hac1$  mutants, and the transcription of representative UPR target genes, including KAR2 (ER-resident chaperone), PDI1 (protein disulfide isomerase), DER1 (ER-associated degradation), and FPR2 (ER protein trafficking), was examined. Compared to the wild-type strain, the expression levels of these genes were decreased in the  $\Delta hac1$  mutant, but reversed by the heterologous expression of C. glabrata HAC1 in the mutant (Figure 5B). Although a direct interaction between C. glabrata Hacl and UPR targets in S. cerevisiae was not examined, the overexpression of C. glabrata HAC1 conferred the increased expression levels of these genes (e.g. 3.5-fold induction of KAR2 expression relative to the wild-type control). Since sustained activation of the UPR is toxic to cells [47], the constitutive overexpression of C. glabrata HAC1 and the induced form of S. cerevisiae HAC1 (ScHAC1<sup>i</sup>) similarly impaired cell growth even under normal growth conditions (Figures 5C and 5D). Furthermore, growth of the S. cerevisiae  $\Delta hac1$  mutant in the presence of TM was rescued by C. glabrata Hac1 to the wild-type level (Figure 5C). The sequences of C. glabrata HAC1 obtained by RT-PCR from TMtreated and -untreated cells were identical (data not shown). These results indicate that C. glabrata Hac1 was matured without splicing in S. cerevisiae. In agreement with this, C. glabrata Hac1 also rescued growth of the S. cerevisiae  $\Delta ire1$  mutant in the presence of TM

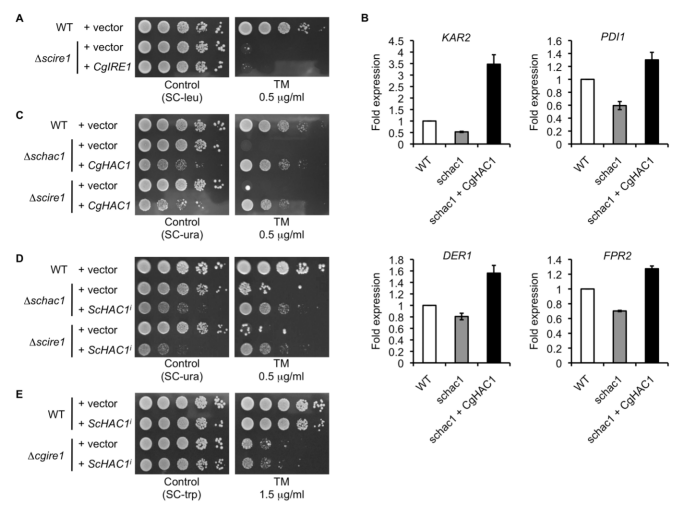


Figure 5. Functional complementation assays of the IRE1 and HAC1 orthologs in S. cerevisiae and C. glabrata. (A) C. glabrata IRE1 was expressed under the control of the S. cerevisiae ADH1 promoter in the S. cerevisiae  $\Delta ire1$  mutant. Logarithmic-phase cells were adjusted to  $2\times10^7$ cells/ml, and then 5 µl of serial 10-fold dilutions were spotted onto agar plates in the presence and absence of 0.5 µg/ml tunicamycin (TM). Plates were incubated at 30°C for 48 h. S. cerevisiae strains: WT+vector, BY42-1; Δscire1+vector, BY42l-1; and Δscire1+CqIRE1, BY42l-2. (B) C. qlabrata HAC1 was expressed under the control of the S. cerevisiae ADH1 promoter in the S. cerevisiae Δhac1 mutant. Expression levels of representative UPR target genes in S. cerevisiae were analyzed by gRT-PCR as described in Materials and Methods. Results are presented as fold expression relative to the levels in the wild-type control. The means and standard deviations for three independent experiments are shown. S. cerevisiae strains: WT. BY42-2; schac1, BY42H-1; and schac1+CgHAC1, BY42H-2. (C) C. glabrata HAC1 was expressed under the control of the S. cerevisiae ADH1 promoter in the S. cerevisiae Δhac1 and Δire1 mutants. The assay was performed as in part A. S. cerevisiae strains: WT+vector, BY42-2; Δschac1+vector, BY42H-1; and Δschac1+CqHAC1, BY42H-2; Δscire1+vector, BY42I-3; and Δscire1+CqHAC1, BY42I-4. (D) The induced form of S. cerevisiae HAC1, denoted as ScHAC1<sup>1</sup>, was expressed under the control of the S. cerevisiae PGK1 promoter in the S. cerevisiae Δhac1 and Δire1 mutants. The assay was performed as in part A. S. cerevisiae strains: WT+vector, BY42-2; Δschac1+vector, BY42H-1; and Δschac1+ScHAC1<sup>i</sup>, BY42H-3; Δscire1+vector, BY42l-3; and Δscire1+ScHAC1<sup>i</sup>, BY42I-5. (E) ScHAC1' was expressed under the control of the S. cerevisiae PGK1 promoter in the C. glabrata wild-type and  $\Delta ire1$  mutant strains. The assay was performed as in part A except TM concentration (1.5 μg/ml). C. glabrata strains: WT+vector, TG11; WT+ScHAC1<sup>†</sup>, TG13; Δcgire1+vector, TG122; and  $\triangle cgire1 + ScHAC1^i$ , TG126. doi:10.1371/journal.ppat.1003160.g005

(Figure 5C). These results suggest that *C. glabrata* Hac1 retains a function as a transcription factor to activate the UPR targets; however, unlike *S. cerevisiae* HAC1, the *C. glabrata* HAC1 mRNA does not need to be spliced by Ire1 before translation.

We also examined the effects of heterologous expression of *S. cerevisiae HAC1* in *C. glabrata*. Since *C. glabrata* Ire1 was unable to splice *S. cerevisiae HAC1* (Figure 4A), *ScHAC1<sup>i</sup>* was expressed under the control of the *PGK1* promoter in *C. glabrata*. Although *C. glabrata* Hac1 complemented *S. cerevisiae* Hac1 functions (Figures 5B and 5C), *ScHAC1<sup>i</sup>* did not rescue growth of the *C. glabrata*  $\Delta ire1$  mutant in the presence of TM (Figure 5E). *ScHAC1<sup>i</sup>* partially rescued growth of the *S. cerevisiae*  $\Delta hac1$  and  $\Delta ire1$  mutants in the presence of TM, confirming its functionality (Figure 5D). Since *C.* 

glabrata Hac1 induced transcription of the UPR targets in S. cerevisiae (Figure 5B), the Hac1 orthologs in C. glabrata and S. cerevisiae may share a common binding motif in the promoter regions of the target genes. These results support the idea that the impaired ER stress response of the C. glabrata  $\Delta ire1$  mutant is not associated with Hac1 functions.

## The Ire1, calcineurin, and Slt2 signaling pathways are coordinately required for ER stress response in *C. alabrata*

We next investigated the possibility that the cellular response to ER stress might involve potential crosstalk between Irel and other signaling pathways in C. glabrata. In addition to the UPR, the serine-threonine-specific protein phosphatase calcineurin and the Slt2 mitogen-activated protein kinase (MAPK) pathway are also required for the ER stress response in S. cerevisiae [48,49,50,51]. We examined whether this is the case in C. glabrata using four C. glabrata mutant strains lacking a key component of these signaling pathways, including Ire1, the regulatory B subunit of calcineurin Cnb1, the calcineurin-regulated transcription factor Crz1, and the last member of the PKC1-MAPK cascade Slt2. While all mutants displayed wild-type growth on a drug-free control plate, the  $\Delta ire1$ ,  $\Delta cnb1$ , and  $\Delta slt2$  mutants exhibited decreased tolerance to both TM and DTT (Figure 6A). Growth of the  $\Delta crz1$  mutant was at nearly wild-type levels in the presence of TM, but drastically impaired on the plate containing DTT. These results were in agreement with a previous proposal that C. glabrata calcineurin is involved in various stress responses via Crz1-dependent and independent mechanisms depending on the type of stress [52]. All mutant phenotypes were recovered to wild-type levels by reintroducing the corresponding wild-type genes into the mutants (Figure 6A). Taken together, the results indicate that Irel, calcineurin, and Slt2 are required for the ER stress response in C. glabrata, consistent with previous findings in S. cerevisiae.

Next, we deleted IRE1 in the  $\Delta cnb1$ ,  $\Delta crz1$ , and  $\Delta slt2$  backgrounds and monitored the viabilities of the single and double deletants after exposure to 1.5 µg/ml TM in liquid medium for up to 5 h. The double deletants displayed more rapid and greater reduction of viability than each single deletant in the presence of TM (Figure 6B). The results suggest that Ire1, calcineurin, and Slt2 function in parallel to cope with ER stress in C. glabrata.

### Gene expression profiling of *C. glabrata* mutants in the presence of ER stress

To examine the transcriptional response to ER stress in C. glabrata, the wild-type,  $\Delta ire1$ ,  $\Delta cnb1$ ,  $\Delta crz1$ , and  $\Delta slt2$  mutants were exposed to 1.5 µg/ml TM for 0.5 and 3 h, and genome-wide analyses were conducted using DNA microarrays. The complete dataset can be found at the NCBI Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo/) with accession number GSE29855. It has been reported that at least 381 genes, corresponding to more than 5% of the ORFs in the genome, were induced in response to TM and DTT under the control of the UPR in S. cerevisiae [20]. In our assay, a total of 325 genes were differentially regulated (greater than 2.0-fold change) and 75 genes of them were upregulated in the C. glabrata wild-type strain after treatment with TM for 3 h (Figure 7 and Table S2). In contrast to previous findings in S. cerevisiae [20], the induced dataset in C. glabrata was not enriched for genes involved in folding capacity of the ER (Table S2). Therefore, to rule out the possibility that unfolded protein stress might not be induced sufficiently by the conditions used in our study, C. glabrata wild-type cells were treated with higher concentrations of TM and DTT, and then expression levels of five C. glabrata genes (KAR2, PDI1, DER1, FPR2 and HACI), which are well-known UPR targets in S. cerevisiae [20,53], were examined by qRT-PCR. The induction levels of KAR2 in the presence of high concentrations of TM and DTT were similar to those observed in the presence of 1.5 µg/ml TM (Figure S4A). In addition, expression levels of the other 4 genes were not increased even in cells treated with high concentrations of TM and DTT (Figure S4B). Collectively, these results suggest that ER quality may be controlled differently in C. glabrata than S. cerevisiae, which relies primarily on transcriptional induction of genes that increase the protein folding capacity of the ER.

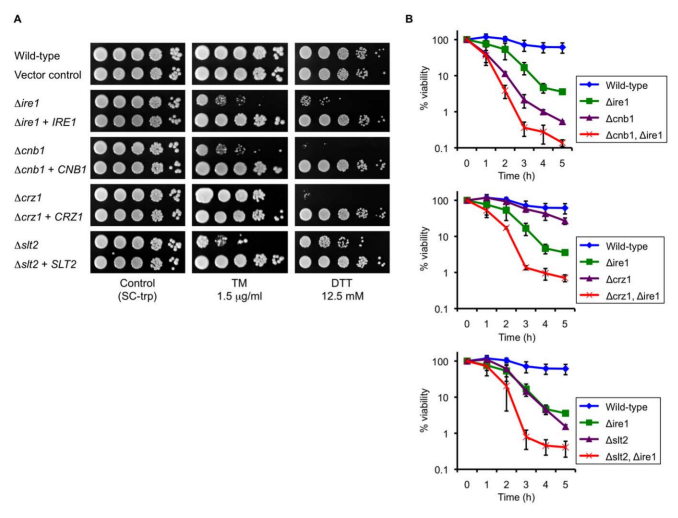
Furthermore, in contrast to S. cerevisiae, the vast majority of genes induced in the C. glabrata wild-type strain were also similarly upregulated in the  $\Delta ire1$  mutant, but at less than two-fold change, or were downregulated in the  $\Delta cnb1$  mutant (Figure 7 and Table S2). The loss of Crz1 and Slt2 had partial effects on the upregulation of those genes. Many of the genes induced by TM overlapped with calcineurin-regulated genes reported in a recent study of calcineurin signaling in C. glabrata [54], indicating the activation of calcineurin signaling in response to TM. A subset of genes that displayed calcineurin-dependent upregulation in the presence of 1.5 µg/ml TM in our microarray experiments were further evaluated by qRT-PCR after treatment with a higher concentration of TM (10 µg/ml) for 3 h (Figure 8). All the genes examined in this assay were upregulated upon TM treatment in a calcineurin- and Crz1-dependent manner but independently of Ire1 and Hac1. In addition to calcineurin, Slt2 was also required for the induction of CAGL0I07249g, consistent with the microarray data. Although these assays cannot differentiate a direct from an indirect effect, the results suggest that calcineurin, but not Irel, plays important roles in transcriptional response to ER stress in C. glabrata.

Transcriptional induction of the ER-resident chaperone KAR2 is a well-known marker for UPR activation in yeast [55,56]. In response to TM treatment, expression levels of C. glabrata KAR2 were increased in the wild-type,  $\Delta slt2$ , and  $\Delta ire1$  strains, but not in the  $\Delta cnb1$  and  $\Delta crz1$  mutants (Figure 8). These results are in contrast to previous observations in S. cerevisiae, where Irel-Hacl signaling is required for upregulation of KAR2 in response to ER stress [13,17]. A Hac1-binding site, termed the unfolded protein response element (UPRE), was originally defined as a 22-bp sequence element of the S. cerevisiae KAR2 promoter and subsequently refined to a seven-nucleotide consensus, 5'-CAGNGTG-3' [57,58]. Mutations in any of the six conserved nucleotides, or deletion of the central nucleotide, cause it to lose its function as an autonomous upstream activating sequence [57]. Consistent with the KAR2 expression data, the consensus UPRE sequence was absent in the 1-kb upstream region of C. glabrata KAR2. On the other hand, it is known in S. cerevisiae and C. glabrata that the promoters of most calcineurin-dependent genes contain one to six copies of the Crz1-binding sequence, 5'-GNGGC(G/ T)-3' [59,60]. Five copies of the Crz1-binding sequence, including one copy of the full consensus sequence, 5'-GNGGCTCA-3' [60], were found within the 1-kb upstream region of C. glabrata KAR2. These results support our KAR2 expression data, collectively suggesting that transcriptional induction of KAR2 in response to ER stress is mediated by the calcineurin-Crz1 pathway, but not by Ire1 signaling, in C. glabrata.

In our microarray analyses, there was a subset of  $\sim 33$  genes whose mRNA abundance was decreased in the wild-type strain, but not in the  $\Delta ireI$  mutant at a relatively early phase (0.5 h) of ER stress (Figures S5 and Table S4). This cluster predominantly contained genes encoding membrane proteins involved in transferase activity (Table S5). Most of these gene products are known to be ER-resident or pass through the ER before translocating to their final destinations. Further analyses were made to interpret this phenomenon, which will be discussed below.

### Functional dissection of the protein kinase and nuclease domains in *C. glabrata* Ire1

To further investigate how Irel is involved in the ER stress response in *C. glabrata*, we examined the contribution of the protein kinase and nuclease functions of *C. glabrata* Irel to cell growth in the presence of TM and DTT. In *S. cerevisiae*, mutations of two



**Figure 6. Multiple signaling pathways are coordinately involved in the ER stress response in** *C. glabrata.* (A) The *C. glabrata*  $\Delta ire1$ ,  $\Delta cnb1$ ,  $\Delta crz1$ , and  $\Delta slt2$  deletion mutants were transformed with either an empty vector or a plasmid containing the corresponding wild-type gene. Logarithmic-phase cells were adjusted to  $2\times10^7$  cells/ml, and then 5 μl of serial 10-fold dilutions were spotted onto synthetic complete medium without tryptophan (SC-trp) plates in the presence and absence of tunicamycin (TM) and dithiothreitol (DTT) at the indicated concentrations. Plates were incubated at 30°C for 48 h. (B) Time-kill analysis of the *C. glabrata* deletion mutants in the presence of TM. Logarithmic-phase cells  $(5\times10^5$  CFU/ml) were incubated in SC medium containing 1.5 μg/ml TM. The number of viable cells was determined by plating the appropriate dilutions on YPD plates at the indicated time points. Data are expressed as the percentages of viability relative to the untreated (time point 0) control in each strain. The means and standard deviations for three independent experiments are shown. *C. glabrata* strains: Wild-type, CBS138; Vector control, TG11; Δ*ire1*, TG122; Δ*ire1*+*IRE1*, TG123; Δ*cnb1*, TG162; Δ*cnb1*+*CNB1*, TG163; Δ*crz1*, TG172; Δ*crz1*+*CRZ1*, TG173; Δ*slt2*, TG152; Δ*slt2*+*SLT2*, TG153; Δ*cnb1* Δ*ire1*, TG1612; Δ*crz1* Δ*ire1*, TG1712; and Δ*slt2* Δ*ire1*, TG1512. doi:10.1371/journal.ppat.1003160.g006

catalytic residues (D797 and K799) in the nucleotide-binding pocket of Ire1 kinase to asparagines abolish phosphorylation, but preserve RNase activity [61]. Therefore, these mutations (D797N, K799N) allow uncoupling of kinase and nuclease activities of Ire1. Since these two residues are highly conserved in the Ire1 orthologs of fungi, including *C. glabrata* (Figure 1B, asterisks), we created *C. glabrata* kinase-dead *IRE1* (*IRE1*-KD) harboring the corresponding mutations (D723N, K725N) in the Ire1 kinase domain. The growth defects of the Δ*ire1* null mutant in the presence of TM and DTT were complemented by wild-type *IRE1*, but not by *IRE1*-KD (Figure 9), indicating that the protein kinase function of Ire1 is required for the ER stress response in *C. glabrata*.

Unlike in other eukaryotes, *C. glabrata* Ire1 did not cleave *HAC1* mRNA in response to ER stress; we therefore hypothesized that the nuclease function of Ire1 would be dispensable for the ER stress response in *C. glabrata*. To test this hypothesis, we engineered *C. glabrata* nuclease-dead *IRE1* (*IRE1*-ND) containing a 10 residue

(D973-Y982) internal deletion within the nuclease domain (Figure 1B). This deleted region contains three highly conserved residues (indicated by arrowheads in Figure 1B) that are the nuclease active sites of *S. cerevisiae* Ire1 [16]. A recent study has shown that deletion of the corresponding region (D1076-R1085) in *A. fumigatus* IreA abrogates its nuclease activity without an apparent effect on any nuclease-independent functions [8]. Contrary to our expectations, the  $\Delta ire1$  mutant carrying *IRE1*-ND displayed growth defects in the presence of TM and DTT, similar to the  $\Delta ire1$  null mutant in *C. glabrata* (Figure 9). These results suggest that both protein kinase and nuclease functions of Ire1 are required for the ER stress response in *C. glabrata*.

### The nuclease activity of Ire1 is required for degradation of ER-associated mRNAs

It has been found in higher eukaryotes that Irel induces degradation of mRNAs encoding proteins that are translocated

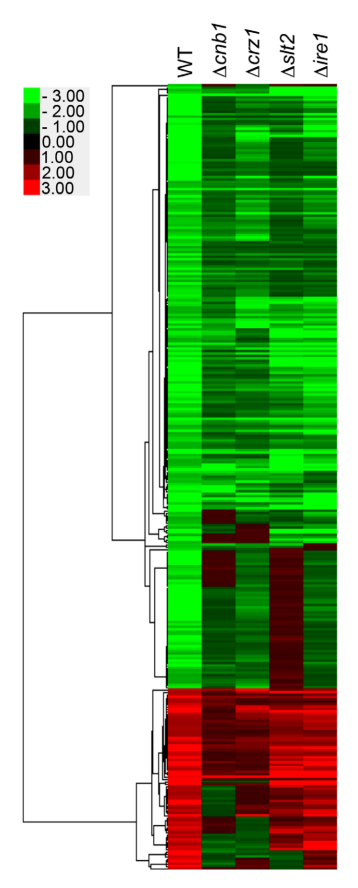


Figure 7. Genome-wide gene expression profiles in response to tunicamycin (TM) exposure for 3 h. Hierarchical clustering of genes whose expression levels were changed more than 2-fold after treatment with 1.5  $\mu$ g/ml TM for 3 h. Genes were clustered with centroid linkage. *C. glabrata* strains: WT, 2001T;  $\Delta cnb1$ , TG161;  $\Delta crz1$ , TG171;  $\Delta slt2$ , TG151; and  $\Delta ire1$ , TG121. doi:10.1371/journal.ppat.1003160.g007

into the ER lumen [62,63,64]. This cellular response is called regulated Ire1-dependent decay (RIDD) and represents an Xbp1independent posttranscriptional mechanism to selectively relieve the burden of incoming proteins on the ER [63]. In our microarray analysis, a subset of genes displayed Irel-dependent "downregulation" in response to TM (Figure S5 and Table S4). There was a strong enrichment for genes encoding GPI-anchored cell wall and membrane proteins in this cluster (Table S5). Thus, we further analyzed the expression levels of 3 representative genes, GAS2 encoding a GPI-anchored cell wall protein that functions as β-1,3-glucanosyltransferase, GAS4 encoding a member of the GAS family similar to GAS2, and ECM33 encoding a GPI-anchored membrane protein involved in apical bud growth, by qRT-PCR using C. glabrata cells treated with DTT and TM. In response to the treatment with 10 mM DTT for 2 h, RNA abundance of these genes was significantly decreased in the wild-type strain compared to the results with the  $\Delta ire1$  mutant (Figure 10A). Similar results were obtained for cells treated with 1.5 µg/ml TM in the microarray analysis (Table S4) and by a confirmatory qRT-PCR assay (data not shown). This  $\Delta ire1$  phenotype was reversed by intact IRE1, but not by IRE1-ND (Figure 10A), indicating that the degradation of these mRNAs was dependent on the nuclease activity of Ire1. C. glabrata IRE1-KD harboring the simultaneous mutations D723N and K725N had moderate effects on the mRNAs decay, implying that Ire1's nuclease activity may be partially impaired by these mutations in C. glabrata.

Blocking transcription with 1,10-phenanthroline (PHEN) inhibited the transcriptional upregulation of CAGL0K11946g and CAGL0L06776g by DTT but had no or little effect on the mRNA dacay of GAS2, GAS4 and ECM33 (Figure 10A), indicating that the decrease in the mRNA abundance was not due to transcriptional repression. This was further supported by the observation that  $\beta$ -galactosidase activities of the C. glabrata GAS2 promoter- $lac\mathcal{Z}$  fusion gene were increased, not decreased, in both the wild-type and  $\Delta inel$  strains in response to DTT exposure (Figure 10B). Although it is unclear why the activity of the GAS2 promoter is increased in response to DTT, this paradoxical behavior suggest that transcriptional regulation and Ire1-dependent decay may not coordinately function in ER-stressed C. glabrata cells.

## Phenotypic differences between the $\Delta ire1$ mutant in *C. glabrata* and UPR defective mutants in other fungal species

To further investigate the roles of Irel in stress response in *C. glabrata*, we examined the growth of the  $\Delta irel$  mutants under various stress conditions. Pathogenic fungi commonly encounter a variety of adverse environmental conditions, such as low oxygen availability and nutrient depletion, in the infected host. It has been reported in *A. fumigatus* that IreA is required for growth in hypoxia and under iron-depleted conditions, which is induced by adding the Fe<sup>2+</sup> chelator bathophenantroline disulphonate (BPS) to the culture medium [8]. In *C. glabrata*, the loss of Irel had no apparent effect on cell growth under low oxygen tension (Figure S6A) and on plates containing BPS or the bacterial siderophore desferrioxamine (DFO) (Figure S6B). *C. glabrata* cells are unable to utilize DFO, which therefore induces iron-depletion for this fungus [65]. Growth of both *C. glabrata* wild-type and  $\Delta irel$  strains in the presence of BPS or DFO was rescued by supplementation of the media with 1 mM FeCl<sub>3</sub>.

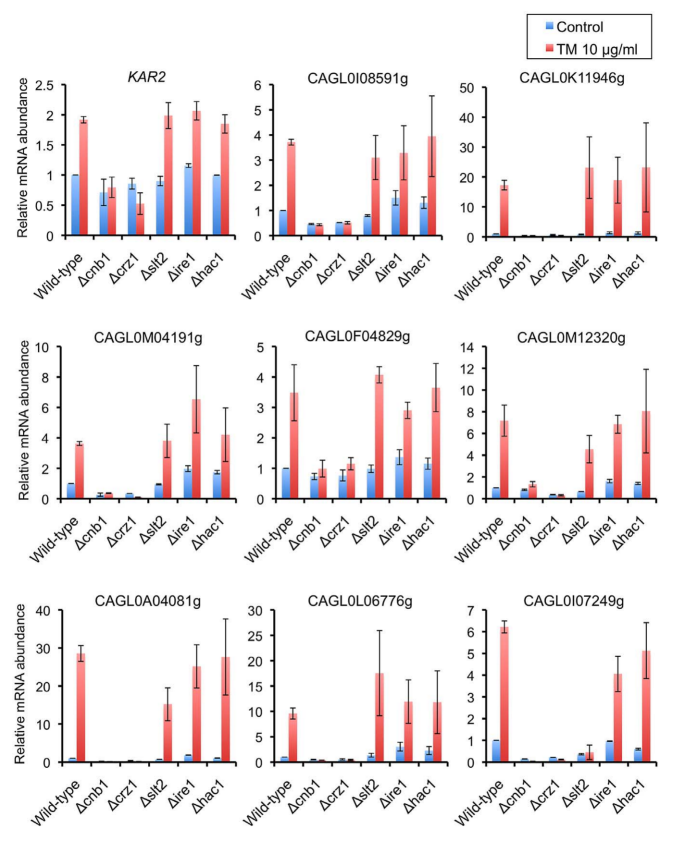
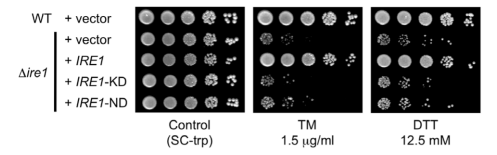
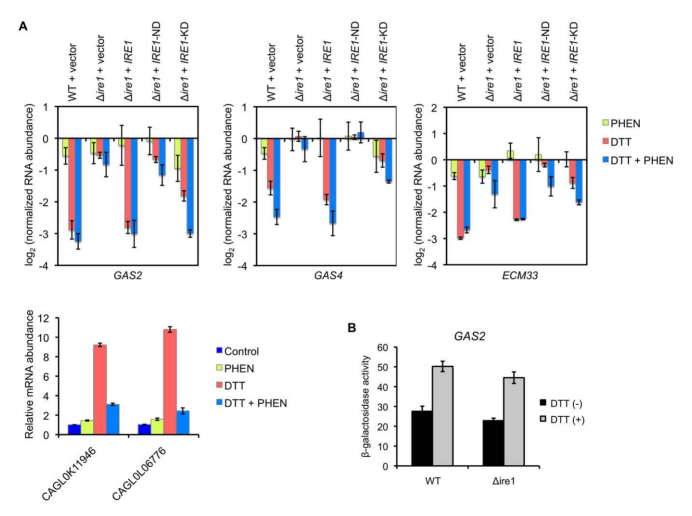


Figure 8. qRT-PCR validation of transcriptional profiles in the presence of tunicamycin (TM). Logarithmic-phase *C. glabrata* cells were incubated in the presence and absence of 10  $\mu$ g/ml TM. qRT-PCR was performed as described in Materials and Methods. The means and standard deviations for three independent experiments are shown. *C. glabrata* strains: WT, 2001T;  $\Delta cnb1$ , TG161;  $\Delta crz1$ , TG171;  $\Delta slt2$ , TG151;  $\Delta ire1$ , TG121; and  $\Delta hac1$ , TG141.

doi:10.1371/journal.ppat.1003160.g008



**Figure 9. Dissection of Ire1 functions required for the ER stress response in** *C. glabrata*. (A) Both the kinase and nuclease activity of Ire1 are required for cell growth in the presence of ER stress. The *C. glabrata*  $\Delta ire1$  mutant was transformed with an empty vector or a plasmid containing wild-type *IRE1*, kinase-dead *IRE1* (*IRE1*-KD), or nuclease-dead *IRE1* (*IRE1*-ND). Logarithmic-phase cells were adjusted to  $2 \times 10^7$  cells/ml, and then 5 μl of serial 10-fold dilutions were spotted onto synthetic complete medium without tryptophan (SC-trp) plates in the presence and absence of tunicamycin (TM) and dithiothreitol (DTT) at the indicated concentrations. Plates were incubated at 30°C for 48 h. *C. glabrata* strains: WT+vector, TG11;  $\Delta ire1$ +vector, TG122;  $\Delta ire1$ +*IRE1*, TG123;  $\Delta ire1$ +*IRE1*-ND, TG125; and  $\Delta ire1$ +*IRE1*-KD, TG124. doi:10.1371/journal.ppat.1003160.g009



**Figure 10. Analysis of Ire1-dependent mRNA decay in** *C. glabrata.* (A) Logarithmic-phase *C. glabrata* cells were incubated in the presence and absence of 10 mM dithiothreitol (DTT) for 2 h. To examine the effects of transcription, cells were treated with the transcription inhibitor 1,10-phenanthroline (PHEN) (50 μg/ml) 5 min before DTT addition. qRT-PCR was performed as described in Materials and Methods. Expression levels of *GAS2* (CAGL0M13849g), *GAS4* (CAGL0F03883g), and *ECM33* (CAGL0M01826g) are expressed as mRNA abundance relative to the untreated control in  $log_2$  scale. The means and standard deviations for three independent experiments are shown. (B) β-galactosidase assay. Logarithmic-phase cells of the *C. glabrata* wild-type and Δ*ire1* strains containing pEM14-GAS2 were exposed to 3 mM DTT for 2 h. β-galactosidase activities of the *GAS2* promoter-*lacZ* fusion gene are expressed in Miller units. The means and standard deviations for three independent experiments are shown. doi:10.1371/journal.ppat.1003160.g010

It is also reported in several pathogenic fungi, including *C. albicans*, *C. neoformans*, and *A. fumigatus*, that mutant strains lacking either Ire1 or Hac1 display growth defects in the presence of cell wall-damaging agents such as caspofungin, Congo red, and calcofluor white [6,7,8,10,11,66]. However, in contrast to those fungi, *C. glabrata* strains lacking *IRE1*, *HAC1*, or both did not exhibit decreased tolerance to these agents at various concentrations tested (data not shown), suggesting that neither Ire1 nor Hac1 is primarily involved in cell wall stress response in *C. glabrata*.

Importantly, disruption of Ire1/IreA confers a drastic increase in azole susceptibility in C. neoformans and A. fumigatus [7,8]. In contrast to these findings, deletion of IRE1 alone did not affect azole susceptibility in C. glabrata (Figure 11A). However, we found that the  $\Delta cnb1 \Delta ire1$  double mutant, but not the  $\Delta crz1 \Delta ire1$  double mutant (data not shown), was more susceptible to azole antifungals than either single mutant (Figure 11A). The results are in agreement with our previous report that calcineurin is required for azole tolerance through a Crz1-independent mechanism in C. glabrata [52]. Furthermore, deletion of either CNB1 or IRE1 alone did not affect cell growth in the presence of caffeine and NaCl, whereas the simultaneous loss of these genes resulted in severe growth defects (Figure 11B). None of these effects was observed when *IRE1* was deleted in the  $\Delta stt2$  background (data not shown). These results suggest that calcineurin and Irel serve redundant roles in cell growth under certain stress conditions in C. glabrata. Collectively, these phenotypic analyses revealed that loss of Irel alone does not induce diverse phenotypes in C. glabrata unlike in other fungi.

#### Ire1 is required for virulence in C. glabrata

We have previously demonstrated in C. glabrata that calcineurin and Slt2 are required for virulence in murine models of disseminated candidiasis [52,67], but the involvement of Ire1 in C. glabrata virulence has not been examined. The C. neoformans  $\Delta ire1$  and A. fumigatus  $\Delta ireA$  mutants are avirulent in murine models of systemic cryptococcosis and invasive aspergillosis, respectively, which can be explained primarily by their growth defects at 37°C [7,8]. In C. glabrata, the loss of Irel did not affect cell growth at  $37^{\circ}$ C (generation time of the wild-type and  $\Delta ire1$  strains in Yeastpeptone-dextrose (YPD) broth was 75 and 78 min, respectively). Therefore, we compared the virulence of the *C. glabrata* wild-type,  $\Delta ire1$ , and IRE1-complemented strains in a mouse model of disseminated candidiasis. Immunocompetent mice infected with the  $\Delta ire1$  mutant displayed significantly reduced fungal burden in both kidney and spleen relative to those infected with the wild-type and Ire1-complemented strains (Figure 12A). To evaluate mortality of mice with disseminated candidiasis, mice were immunosuppressed by cyclophosphamide prior to C. glabrata infection. Consistent with the results of organ fungal burden, mice infected with the wild-type and IRE1-complemented strains exhibited higher mortality rates than mice infected with the  $\Delta ire1$  mutant (Figure 12B). These results suggest that Ire1 is important for virulence in C. glabrata.

#### Discussion

According to the current evidence, the UPR regulated by Ire1-Hac1 signaling is highly conserved and is a key pathway to cope with ER stress in eukaryotes [2,12]. However, the present study demonstrates the lack of this pathway and the development of alternative mechanisms for the ER stress response in *C. glabrata*. The gene CAGL0K12540g, named *C. glabrata HAC1*, seems to be the sole *HAC1* ortholog in *C. glabrata*, since it is the only gene showing sequence similarity and conserved synteny to *S. cerevisiae* 

HAC1. In addition, C. glabrata Hac1 was able to complement S. cerevisiae Hac1 functions. However, surprisingly, C. glabrata Hac1 did not necessitate Ire1-mediated splicing of mRNA before translation, and thus could recover the growth of the S. cerevisiae  $\Delta irel$  mutant in the presence of ER stress. A recent study has comprehensively characterized the RNA structures of the Hac1/ Xbp1 orthologs in various eukaryotic species [36]. In C. glabrata HAC1, although the overall RNA structure is maintained, the predicted intron (379 nucleotides) is much longer than those in other species and has mutations in the Ire1 recognition motifs at both splice sites, suggesting that the Irel-dependent unconventional splicing mechanism may not be present in C. glabrata [36]. This has been experimentally confirmed in our studies: the heterologous expression of C. glabrata HAC1 complemented S. cerevisiae Hac1 functions without splicing, and C. glabrata HAC1 mRNA was not spliced by Ire1 even in ER-stressed C. glabrata cells. Furthermore, Hooks and Griffiths-Jones [36] revealed that the non-canonical Hac1/Xbp1 intron structure is not conserved in 28 out of 156 eukaryotic species searched, including several Candida species such as Candida parapsilosis, Candida lusitaniae, and Candida guilliermondii, despite the fact that the HAC1 ORFs are intact in these Candida species. These findings suggest that these species may have lost the unconventional splicing mechanism, but instead acquired an alternative mechanism to regulate the UPR. Future studies are warranted to determine whether Hac1 is involved in the ER stress response in these species.

Although some Hacl-independent functions of Irel have been reported in some eukaryotic species [2,7,8,63], mutants lacking Hac1 or its homolog always phenocopy  $\Delta ire1$  strains with respect to growth defects in the presence of ER stress-inducing agents (e.g. TM and DTT) in all eukaryotic species tested so far, and the UPR mediated by Ire1-Hac1 signaling has been believed to be a phylogenetically conserved pathway in eukaryotes [2,68,69]. However, our study demonstrated that C. glabrata Irel plays a role in the ER stress response in a Hacl-independent manner. Exploration of potential Irel targets in this fungus by a genomewide analysis will be of interest in future investigations (e.g. RNAseq analysis of ER-stressed and unstressed cells of wild-type and  $\Delta ire1$  strains to detect Ire1-dependent spliced RNAs). The results of our gene expression assays suggest that Ire1 does not mediate transcriptional response to ER stress in C. glabrata, raising the question of how Ire1 is involved in the ER stress response in this fungus.

In metazoans, ER stress triggers two distinct outputs of Ire1's nuclease activity, XBP1 splicing and RIDD [69,70]. The latter is an Xbp1-independent pathway that selectively degrades a small subset of ER-associated mRNAs and remodels the repertoire of proteins translated in ER-stressed cells [63,64]. This cellular response is predicted to reduce the ER load by limiting protein influx and unfolded protein load into the ER lumen. The RIDD pathway was first uncovered in D. melanogaster [64], and later, its conservation was confirmed in mammalian cells [62,63], but has yet to be fully determined in fungi. Interestingly, in S. cerevisiae, RIDD does not occur [63,70], but the classic Ire1-Hac1 UPR mediates downregulation of some mRNAs encoding membrane proteins [64,71]. Here, we have demonstrated that mRNA abundance of a subset of genes encoding GPI-anchored cell wall and membrane proteins was diminished during the response to ER stress in the wild-type strain dependent on the nuclease activity of Ire1. There was no overlap between these repressed C. glabrata genes and the previously identified S. cerevisiae gene that were transcriptionally downregulated by the UPR in response to TM [71]. It is thought that RIDD targets are nicked by the Irel endoribonuclease at sites that do not display an identifiable

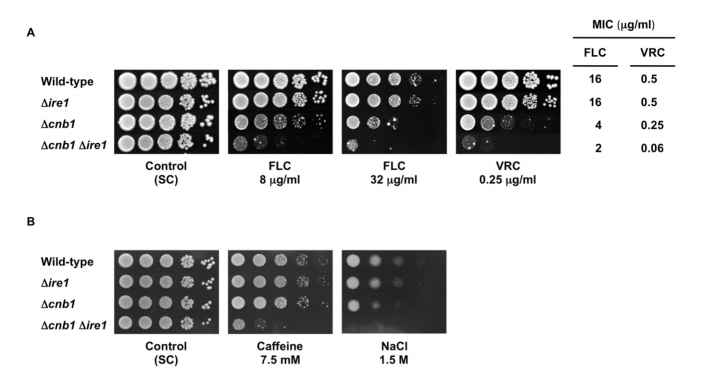


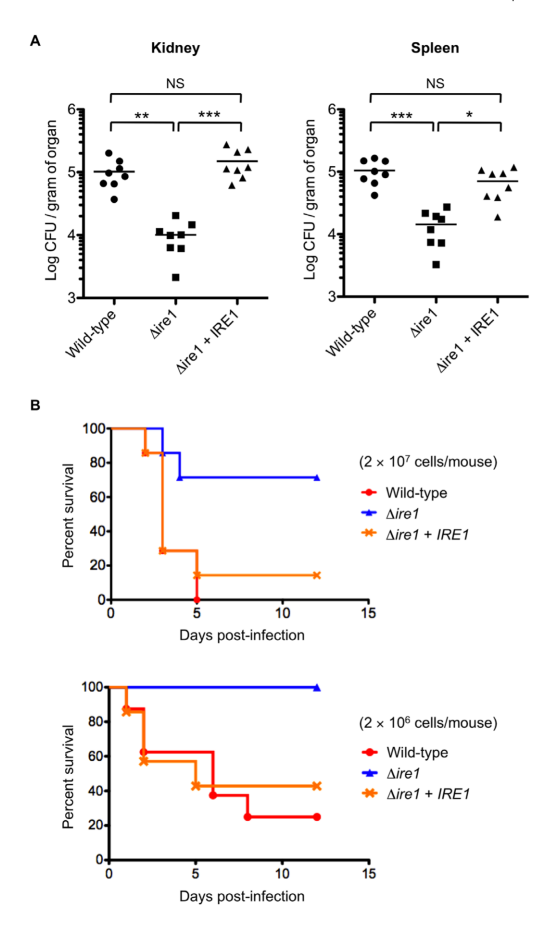
Figure 11. Effects of *IRE1* deletion on stress response in wild-type and  $\Delta cnb1$  backgrounds in *C. glabrata*. (A) Ire1 plays a role in azole tolerance in *C. glabrata* when calcineurin is absent. Logarithmic-phase cells of each *C. glabrata* strain were adjusted to  $2 \times 10^7$  cells/ml, and then 5  $\mu$ l of serial 10-fold dilutions were spotted onto synthetic complete (SC) plates containing either fluconazole (FLC) or voriconazole (VRC) at the indicated concentrations. Plates were incubated at  $30^{\circ}$ C for 48 h. MICs were determined by a broth microdilution method. (B) Ire1 and calcineurin serve redundant roles in cell growth under certain stress conditions in *C. glabrata*. The assay was performed as in part A. *C. glabrata* strains: Wild-type, CBS138;  $\Delta ire1$ , TG121;  $\Delta cnb1$ , TG161; and  $\Delta cnb1$   $\Delta ire1$ , TG1612. doi:10.1371/journal.ppat.1003160.g011

consensus sequence, in contrast to Ire1-mediated splicing of *HAC1/XBP1* mRNA at the highly conserved splice junctions [69]. As excessive activation of RIDD seems to be detrimental to cell integrity [62], reducing the load of proteins entering the ER must be balanced with the need to sustain synthesis of essential proteins. In contrast to Ire1 in metazoans, *C. glabrata* Ire1 may not need to switch between specific and nonspecific modes of cleavage. We are currently investigating how the RIDD activity is regulated in this fungus.

Although the only known substrate of the Irel kinase is Irel itself in S. cerevisiae [30,31], we also addressed the possibility that Ire1 may integrate with other signaling pathways in C. glabrata. In S. cerevisiae, calcineurin and Slt2 are also involved in the ER stress response through different mechanisms [48,49,50,51]. Slt2 plays a role in the ER stress surveillance (ERSU) pathway that ensures transmission of only functional ER to daughter cells during cell division [72]. Upon ER stress, the ERSU pathway delays ER inheritance and cytokinesis to prevent death of both mother and daughter cells. Calcineurin mediates the calcium cell survival pathway by regulating intracellular Ca<sup>2+</sup> homeostasis. TM increases Ca<sup>2+</sup> uptake by stimulating the Cch1-Mid1 high affinity Ca<sup>2+</sup> channel, while calcineurin dephosphorylates the Cch1 subunit of the channel to inhibit Ca<sup>2+</sup> influx and prevents nonapoptotic cell death in S. cerevisiae [48,51]. In C. glabrata, loss of either calcineurin or Slt2 resulted in decreased tolerance to ER stress, and the additional deletion of IRE1 in the  $\Delta cnb1$  and the  $\Delta slt2$  backgrounds had synthetic effects on viability loss in the presence of TM. The results suggest that Ire1 functions in parallel with the calcineurin and Slt2 MAPK pathways to cope with ER stress in C. glabrata, consistent with previous findings in S. cerevisiae [48,49,50]. However, the possibility that Ire1 might interact with other signaling pathways has not been completely ruled out. Whether the Ire1 kinase domain has an effector other than Ire1 itself remains one of the major unsolved questions in this field [69].

The term UPR is derived from a cellular response to chemicals that interfere with proper protein folding (e.g. TM and DTT), but it is now known that the signaling pathway is involved in various stress responses from yeast to humans [73,74]. Indeed, downstream targets of the UPR include not only ER chaperones, but also genes involved in diverse functions, including protein trafficking and quality control, lipid and sterol metabolism, heme biosynthesis, and cell wall biogenesis in S. cerevisiae [20]. To extend our understanding of Irel functions in C. glabrata, phenotypes of the  $\Delta ire1$  mutant were investigated in various stress conditions in view of clinical significance. The UPR is critical for azole tolerance in C. neoformans and A. fumigatus, and thus loss of either Ire1/IreA or Hxl1/HacA leads to increased susceptibility to azole antifungals in these species [7,8,10]. In contrast, loss of Ire1, Hac1, or both did not affect azole tolerance in C. glabrata (Figure 11A and data not shown), further supporting the idea that the classic UPR mechanism regulated by Ire1-Hac1 signaling is not conserved in this fungus. However, the additional deletion of *IRE1* in the  $\Delta cnb1$ background increased azole susceptibility in C. glabrata. One possible explanation is that azole antifungals induce severe ER stress in calcineurin-defective C. glabrata cells, which therefore require Irel to survive this stress condition.

Previous studies in several yeasts and filamentous fungi have demonstrated that the UPR plays a role in maintaining cell wall integrity, and that mutants lacking Irel or Hacl exhibit decreased tolerance to a variety of cell wall perturbing agents [6,7,8,10,11,66].



**Figure 12.** *C. glabrata* **Ire1 is required for virulence in a murine model of disseminated candidiasis.** (A) Groups of 8 immunocompetent mice were intravenously inoculated with  $8 \times 10^7$  cells for each *C. glabrata* strain. Bilateral kidneys and spleen were excised 7 days after injection. Appropriate dilutions of organ homogenates were plated, and the numbers of CFU were counted after 2 days of incubation at 30°C. Numbers of recovered CFU from each organ are indicated for individual mice in the scatter plots. The geometric mean is shown as a bar. Statistical analyses were performed using the Kruskal-Wallis test with Dunn's multiple comparison post-test. Asterisks indicate statistically significant differences (\*\*: P < 0.05; \*\*\*: P < 0.01; \*\*\*: P < 0.001). NS indicates no significance (P > 0.05). Representative data of two independent experiments are shown. *C. glabrata* strains: Wild-type, TG11; Δ*ire1*, TG122; and Δ*ire1+lRE1*, TG123. (B) Groups of 7 mice were immunosuppressed by intraperitoneal administration of cyclophosphamide (200 mg/kg/day) 72, 48, and 24 h before challenge with *C. glabrata* cells. The mice were infected intravenously with *C. glabrata* cells (2×10<sup>7</sup> and 2×10<sup>6</sup> cells/mouse) no Day 0 of the experiment, and survival was monitored for 12 days post-infection. Kaplan-Meier curves were created and compared with the log rank (Mantel-Cox) test. Upper panel (2×10<sup>7</sup> cells/mouse): P = 0.0078 for wild-type vs. Δ*ire1*, P = 0.0284 for Δ*ire1* vs. Δ*ire1+lRE1*, and P = 0.6768 for wild-type vs. Δ*ire1+lRE1*; lower panel (2×10<sup>6</sup> cells/mouse): P = 0.0043 for wild-type vs. Δ*ire1*, P = 0.0222 for Δ*ire1* vs. Δ*ire1+lRE1*, and P = 0.7440 for wild-type vs. Δ*ire1+lRE1*. doi:10.1371/journal.ppat.1003160.g012

TM also induces cell wall stress, thereby explaining why certain genes implicated in cell wall biogenesis are upregulated during the UPR in TM-treated S. cerevisiae and C. albicans cells [11,20,75]. In contrast to other fungi investigated to date, the loss of Ire1 alone did not confer a cell wall-defective phenotype, and upregulation of some cell wall-related genes in response to TM exposure was mainly dependent on calcineurin, but not on Ire1, in C. glabrata (Table S2). These results provide additional evidence that there are diversities in Irel-dependent stress response mechanisms between C. glabrata and other fungal species. In addition,  $\Delta ire1/ireA$  mutants of C. neoformans and/or A. fumigatus exhibit growth defects under various environmental conditions, such as growth temperature at 37°C, hypoxia, and iron limitation, which are commonly encountered by pathogenic fungi in the infected host [7,8]. However, none of these phenotypes were observed in the C. glabrata  $\Delta ire1$  mutant. In agreement with our proposal that the canonical Ire1-Hac1 UPR has been lost in C. glabrata, phenotypes of the C. glabrata  $\Delta ire1$  mutant were confined relative to those of UPR-defective mutants in other fungi where the UPR is involved in various stress responses.

Diversity in the transcriptional induction system in response to ER stress has developed during evolution [68]. For instance, the Ire1-Hac1 pathway is required for upregulation of KAR2 in yeast while the pathway is dispensable for induction of the Kar2 homolog GRP78/BiP in mammalian cells [17,76]. In C. glabrata, the majority of genes, including KAR2, induced in response to ER stress were dependent on the calcineurin-Crz1 pathway, but not on Ire1 signaling. In addition, several lines of evidence described above demonstrate that C. glabrata has lost the classic Irel-Hacl UPR, but instead possesses an alternative mechanism, RIDD, like metazoans. To our knowledge, these unusual characteristics have been demonstrated for the first time in any eukaryote, and perhaps they were developed in C. glabrata after its divergence from S. cerevisiae due to its ecological niche. Our results provide novel insights into ER stress response mechanisms as well as fundamental information for evolutionary biology to further understand how eukaryotic cells have developed ER quality control systems.

#### **Materials and Methods**

#### Ethics statement

All animal experiments were performed in full compliance with the Guide for the Care and Use of Laboratory Animals (National Research Council, National Academy Press, Washington DC, 2011) and all of the institutional regulations and guidelines for animal experimentation after pertinent review and approval by the Institutional Animal Care and Use Committee of Nagasaki University under protocol number 0904130747.

#### Strains and culture conditions

The *C. glabrata* and *S. cerevisiae* strains used in this study are listed in Table S6. Cells were routinely propagated at 30°C in YPD,

synthetic complete (SC) medium lacking appropriate amino acids, or synthetic defined (SD) medium [77], unless otherwise indicated. The nitrogen starvation medium, SD-N, consists of 0.17% yeast nitrogen base without amino acids and ammonium sulfate (BD Biosciences, Franklin Lakes, NJ) plus 2% glucose as described previously [78]. The Anaeropack system (Mitsubishi Gas Chemical Company Inc, Tokyo, Japan) was used for cultures under conditions of low oxygen tension (hypoxic). To induce iron-depleted conditions, the Fe<sup>2+</sup> chelator BPS (MP Biomedicals, Solon, OH) or the bacterial siderophore DFO (EMD Chemicals, San Diego, CA) were added to SC media at final concentrations of 20 and 50  $\mu$ M, respectively.

#### Plasmid and strain construction

The primers and plasmids used in this study are listed in Tables S7 and S8, respectively. Transformation of *C. glabrata* and *S. cerevisiae* was performed using a lithium acetate protocol as described previously [79]. *C. glabrata* deletion mutants were constructed using a one-step PCR-based technique as described previously [67]. Briefly, a deletion construct was amplified from pBSK-TRP or pBSK-HIS using primers tagged with the 100 bp sequences homologous to the flanking regions of the target ORF. *C. glabrata* parent strains were transformed with the deletion construct, and the resulting transformants were selected by tryptophan or histidine prototrophy. PCR and Southern blotting were performed to verify that the desired homologous recombination occurred at the target locus.

To generate complementation plasmids, *C. glabrata* and *S. cerevisiae* genes were amplified from the genomic DNA of CBS138 [80] and BY4742 [81], respectively. Procedures for each plasmid construction are summarized in Table S8. The spliced form of *S. cerevisiae HAC1* mRNA was obtained from BY4742 cells treated with 1.5 μg/ml TM for 3 h. The extracted RNA was then reverse-transcribed using a QuantiTect Reverse Transcription kit (Qiagen, Valencia, CA), and the resulting cDNA was used as a template to amplify the mature *S. cerevisiae HAC1*, *ScHAC1*<sup>i</sup> ("i" for induced).

The plasmid pCgACT-PIRE was used as the template to generate pCgACT-PIRE-KD and pCgACT-PIRE-ND. pCgACT-PIRE-KD was generated by mutating two residues, D723 and K725, in the kinase domain of *C. glabrata* Ire1 to asparagines using the KOD-Plus-Mutagenesis Kit (Toyobo, Osaka, Japan), and mutagenic primers CgIRE1-mut-F2171 and CgIRE1-mut-R2170. Similarly, pCgACT-PIRE-ND was created by introducing a 10 residue (D973-Y982) internal deletion within the nuclease domain of *C. glabrata* Ire1 using mutagenic primers, CgIRE1-F2947 and CgIRE1-R2916. All of the plasmids constructed using PCR products were verified by sequencing before use.

#### Drug susceptibility assay

Spot dilution tests and time-kill assays were performed as described previously [52]. Voriconazole was kindly provided by Pfizer (New York, NY). Other drugs were purchased from Sigma (St Louis, MO). TM, voriconazole, and caffeine were dissolved in dimethyl sulfoxide (DMSO), and others were dissolved in distilled water. DMSO alone did not interfere with cell growth at the final concentrations used in this study. MICs of azole antifungals were determined by a broth microdilution test using a commercially prepared colorimetric microdilution panel (ASTY; Kyokuto Pharmaceutical Industrial Co., Ltd.) [82]. All sensitivity tests were performed on at least two separate occasions to ensure reproducibility.

### Analysis of *HAC1* mRNA splicing by RT-PCR and Northern blotting

HAC1 mRNA splicing was analyzed using a modification of a reported protocol [66]. Logarithmic-phase S. cerevisiae cells grown in SC medium lacking leucine were treated with 1.5 µg/ml TM or 5 mM DTT for 3 h. Logarithmic-phase C. glabrata cells grown in SC medium were treated with either TM (5 and 10 µg/ml) or DTT (5 and 10 mM) for 1 and 3 h. These cells were harvested, and total RNA was extracted using a FastRNA Red Kit (Qbiogene, Carlsbad, CA) according to the manufacturer's instructions. cDNA was synthesized from 1 µg of total RNA using a QuantiTect Reverse Transcription kit (Qiagen) in a final volume of 20 µl, and 5 µl of resulting cDNA were then used as the template for individual PCR. The amplified PCR products were analyzed by both electrophoresis on a 1% agarose gel and DNA sequencing. For Northern blot analysis, twenty micrograms total RNA was separated on a 2% agarose gel and transferred to a Hybond-N<sup>+</sup> membrane (GE Healthcare, Buckinghamshire, UK), which was incubated in PerfectHyb Hybridization Solution (Toyobo) with probes directed against the 5' region of the HAC1 ORF. All probes were generated by PCR with primers listed in Table S7 and were labeled with [α-32P]dCTP using a Random Primer DNA Labeling Kit Ver. 2.0 (Takara Bio Inc., Shiga, Japan). Images were obtained using Image Reader FLA-5000 (FUJIFILM Corporation, Tokyo, Japan).

#### gRT-PCR

Total RNA extraction and cDNA synthesis were performed as described above, and 3  $\mu$ l of resulting cDNA were used as the template for individual PCR with a QuantiTect SYBR Green PCR kit (Qiagen). qRT-PCR was carried out in triplicate in a 96-well plate format, using a 7500 Real-Time PCR System (Applied Biosystems, Foster City, CA). The mRNA abundance of the target genes was normalized to 18S rRNA (CAGL0L13398r). The qRT-PCR assays were repeated at least twice on independent occasions.

#### Microarray analysis

C. glabrata cells were grown in SC medium to exponential phase (OD600 = 0.8) and exposed to 1.5  $\mu$ g/ml TM at 30°C. Total RNAs were extracted using the FastRNA Red Kit (Qbiogene). The quality of RNA was checked with a RNA 6000 Nano Kit and Agilent 2100 Bioanalyzer. Double-stranded cDNA was synthesized using the Invitrogen SuperScript Double-Stranded cDNA Synthesis Kit and oligo (dT) primers. The resulting cDNA samples were labeled with Cy3 using the NimbleGen One-Color DNA Labeling Kit and subsequently hybridized to a custom-made  $4\times72$  K C. glabrata array (Roche NimbleGen, Tokyo, Japan) wherein each chip measures the expression levels of 5,217 genes from C. glabrata CBS138 with six 60-mer-probe pairs per gene, with two-fold technical redundancy. The arrays were washed using the NimbleGen Wash Buffer Kit and scanned with a NimbleGen MS 200 Microarray Scanner. Data were quantified

using the NimbleScan v2.6 software and normalized as described previously [83,84]. Hierarchical clustering was performed using Gene Cluster 3.0 and visualized using Java TreeView ver. 1.1.5r2. Gene Ontology (GO) enrichment was searched using the Candida Genome Database GO term finder (http://www.candidagenome.org/cgi-bin/GO/goTermFinder) with default parameters [85].

#### β-galactosidase assay

A 699-bp DNA fragment containing the 5' untranslated region (UTR) and the first 17 codons of the C. glabrata GAS2 gene was amplified from the genomic DNA of CBS138 and fused to the *lac2* reporter in pEM14 [86] to construct pEM14-GAS2 as described in Table S8. Logarithmic-phase cells of the C. glabrata wild-type and  $\Delta ire1$  strains containing pEM14-GAS2 were grown in SC-ura broth, adjusted to  $1 \times 10^7$  cells/ml, and then incubated in the presence and absence of 3 mM DTT for 2 h. β-galactosidase assay was performed as described previously [59]. The cell cultures were harvested and washed twice with ice-cold phosphate-buffered saline. Cells (100 ul) were re-suspended in 300 ul Reporter Lysis Buffer (Promega, Madison, WI) containing 5 ul Protease Inhibitor Cocktail (Sigma). Cell extracts were prepared using acid-washed glass beads (Sigma) and cleared by centrifugation at  $14,000 \times g$  for 30 min at 4°C. Protein concentrations were determined by the Bio-Rad Protein Assay (Bio-Rad, Richmond, CA) using bovine serum albumin as a standard. β-galactosidase activities were measured using the β-Galactosidase Enzyme Assay System (Promega) according to the manufacturer's instructions and calculated in Miller units (nmoles/min/mg of protein) at 37°C [87]. All assays were performed in triplicate on separate days.

#### Virulence assay

Animal experiments were performed as described previously [52,67]. Briefly, to prepare cells for injection, logarithmic-phase C. glabrata cells were harvested, washed, resuspended in sterile saline, and adjusted to  $4 \times 10^8$  cells/ml after counting the number of cells using a hemocytometer. The actual CFU in the inocula were confirmed by plating serial dilutions of cell suspension on YPD plates. Groups of 8 female, 8-week-old, BALB/c mice (Charles River Laboratories Japan Inc., Japan) were injected with 0.2 ml of the C. glabrata cell suspension via the lateral tail vein. The mice were euthanized 7 days after injection and the spleen and bilateral kidneys were then excised. No mice died before euthanasia in this experiment. Appropriate dilutions of organ homogenates were plated on YPD plates. Colonies were counted after 2 days of incubation at 30°C and the CFUs per organ were calculated. Statistical analyses were performed using the Kruskal-Wallis test with Dunn's multiple comparison post-test (GraphPad Prism 5, La Jolla, CA).

To examine mortality of mice with disseminated candidiasis due to C. glabrata, groups of 7 female BALB/c mice (20–23 g body weight, mean 20.6–20.9 g for each group) were immunosuppressed by intraperitoneal administration of cyclophosphamide at a concentration of 200 mg/kg/day on days -3, -2, and -1 of infection, and housed under sterile conditions. The mice were inoculated intravenously with 0.2 ml of C. glabrata cell suspensions (1×10<sup>8</sup> and 1×10<sup>7</sup> cells/ml) on day 0. Checks were made for dead or moribund mice in a blinded fashion twice daily over a period of 16 days. A group of 4 immunosuppressed mice was injected with sterile saline instead of C. glabrata cell suspension, and no mice died due to immunosuppression only. Survival was plotted on a Kaplan-Meier curve for each C. glabrata strain, and the log rank (Mantel-Cox) test was used for pairwise comparison of percent survival (GraphPad Prism 5). A P value of <0.05 was considered

statistically significant. These animal experiments were conducted on two separate occasions to ensure reproducibility.

#### Accession numbers for genes and proteins mentioned in the text (NCBI Entrez Gene ID or GenBank accession number)

C. glabrata: 18S rRNA (9488051); ACT1 (2890423); CAD1/YAP2 (2887750); CAGL0A04081g (2886450); CAGL0F04829g (2887774); CAGL0I07249g (2889193); CAGL0I08591g (2888926); CAGL0K11946g (2889962); CAGL0L06776g (2890816); CAGL0M04191g (2891538); CAGL0M12320g (2891428); CNB1 (2890566); CRZI (2891693); DERI (2891627); ECM33 (2891194); FPR2 (2888506); GAS2 (2891237); GAS4 (2887679); GCN4 (2890908); HACI (2890416); HIS3 (2890838); IREI (2887705); KAR2 (2887030); PDII (2886594); SKOI (2889579); SLT2 (2889880); TRPI (2886909); YAP1 (2886604); YAP3/CAGL0K02585g (2890264); YAP3/CAGL0M10087g (2891349); YAP6 (2891289); YAP7 (2887954).

S. eerevisiae: ACT1 (850504); ADH1 (854068); CCH1 (853131); DER1 (852500); FPR2 (852131); HAC1 (850513); IRE1 (856478); KAR2 (853418); MID1 (855425); PDI1 (850314); PGK1 (850370); SLT2 (856425).

Others: A. fumigatus HacA/Hac1 (3506096); A. fumigatus IreA/Ire1 (AEQ59230); A. nidurans HacA (CBF87535); C. albicans Hac1 (3639758); C. albicans Ire1 (3640823); C. elegans Xbp1 (175541); C. neoformans Hxl1 (3255200); C. neoformans Ire1 (3255994); D. melanogaster Xbp1 (44226); H. sapiens Xbp1 (7494); P. pastoris HACI (8199414); T. reesei HACI (AJ413272); Y. lipolytica HACI (2906724).

#### **Supporting Information**

**Figure S1** Comparison of cell growth in the presence of ER stress between various fungal species. Logarithmic-phase cells of each fungal strain were harvested after appropriate incubation in YPD broth. Cell concentration was adjusted with optical density at 600 nm, and then 5 µl of serial 10-fold dilutions were spotted onto YPD or synthetic complete (SC) plates containing either tunicamycin (TM) or dithiothreitol (DTT) at the indicated concentrations. Plates were incubated at 30°C for 48 h. (TIF)

**Figure S2** Synteny relationships of the *IRE1* and *HAC1* loci between *S. cerevisiae* and *C. glabrata*. (A) Schematic representation of the *C. glabrata IRE1* (CAGL0F03245g) locus compared to syntenic regions of the *S. cerevisiae* genome. Colored arrows indicate orthologs in the two species. White arrows indicate *S. cerevisiae* genes of which orthologs are not located around *IRE1* in the *C. glabrata* genome. (B) Synteny analysis of the *C. glabrata HAC1* (CAGL0K12540g) and *S. cerevisiae HAC1* loci. (TIF)

Figure S3 Disruption of HAC1 in C. glabrata. The entire open reading frame (ORF) of HAC1 was replaced with the deletion construct containing a TRP1 marker. Southern blot analysis of ClaI-digested genomic DNA using a HAC1 probe, which was designed within the HAC1 ORF, identified the predicted 4.3 kb band in the parent strain 2001HT ( $\Delta his3 \Delta trp1$ ) but no band in the  $\Delta hac1$  mutant TG141 ( $\Delta hac1::TRP1 \Delta his3$ ). A second Southern blot analysis using a TRP1 probe confirmed that the desired homologous recombination had occurred at the HAC1 locus without ectopic integration of the deletion construct in the mutant. (TIF)

**Figure S4** qRT-PCR analysis to validate expression profiles of representative UPR target genes in the presence of tunicamycin

(TM) and dithiothreitol (DTT). *C. glabrata* wild-type (CBS138) cells were treated with TM or DTT at the indicated concentrations for 1 and 3 h. Expression levels of *KAR2* (A) and other known UPR targets (B) were examined by qRT-PCR as described in Materials and Methods. The means and standard deviations for three independent experiments are shown.

**Figure S5** Genome-wide gene expression profiles in response to tunicamycin (TM) exposure for 0.5 h. Hierarchical clustering of genes whose expression levels were changed more than 1.5-fold after treatment with 1.5  $\mu$ g/ml TM for 0.5 h. Genes were clustered with average linkage. *C. glabrata* strains: WT, 2001T;  $\Delta cnb1$ , TG161;  $\Delta crz1$ , TG171;  $\Delta slt2$ , TG151; and  $\Delta ire1$ , TG121. (TIF)

**Figure S6** Irel is dispensable for growth under conditions of hypoxia and iron depletion in *C. glabrata*. (A) Logarithmic-phase cells of each *C. glabrata* strain were adjusted to 2×10<sup>7</sup> cells/ml, and then 5 μl of serial 10-fold dilutions were spotted onto YPD plates. One plate was incubated under normal (aerobic) condition for 44 h, and the other plate was incubated under conditions of low oxygen tension (hypoxic) for 68 h. (B) Serial dilutions of *C. glabrata* cells were prepared as described above and spotted onto synthetic complete (SC) plates containing either 20 μM bathophenantroline disulphonate (BPS) or 50 μM desferrioxamine (DFO) in the presence and absence of 1 mM ferric chloride (FeCl<sub>3</sub>). Plates were incubated at 30°C for 24 h. (TIF)

**Table S1** Putative *C. glabrata* bZIP transcription factors identified by a BLASTp search using the bZIP domain sequences of *S. cerevisiae* Hacl, *H. sapiens* Xbp1, and *C. neoformans* Hxl1 as queries. (DOC)

**Table S2** List of genes whose expression levels were altered by tunicamycin treatment for 3 h. (XLSX)

**Table S3** Classification of the putative Slt2-repressed genes. (XLSX)

**Table S4** List of genes whose expression levels were altered by tunicamycin treatment for 0.5 h.

(XLSX)

**Table S5** Classification of the putative Ire1-repressed genes. (XLSX)

**Table S6** Strains used in this study. (DOC)

**Table S7** Primers used in this study. (DOC)

**Table S8** Plasmids used in this study. (DOC)

#### **Acknowledgments**

We thank Dr. Brendan Cormack for providing BG2 and pGRB2.2 and Dr. Fritz Muhlschlegel for pEM14. We also thank Shunsuke Yamauchi, Asuka Minematsu, and Naoki Hosogaya for technical assistance and other members of the Kohno lab for their support.

#### **Author Contributions**

Conceived and designed the experiments: TM HN SK. Performed the experiments: TM HN YN HK. Analyzed the data: TM HN YN HK SK. Wrote the paper: TM HN SK.

#### References

- Ellgaard L, Helenius A (2003) Quality control in the endoplasmic reticulum. Nat Rev Mol Cell Biol 4: 181–191.
- Ron D, Walter P (2007) Signal integration in the endoplasmic reticulum unfolded protein response. Nat Rev Mol Cell Biol 8: 519–529.
- Vembar SS, Brodsky JL (2008) One step at a time: endoplasmic reticulumassociated degradation. Nat Rev Mol Cell Biol 9: 944–957.
- Kaufman RJ (2002) Orchestrating the unfolded protein response in health and disease. J Clin Invest 110: 1389–1398.
- Wang S, Kaufman RJ (2012) The impact of the unfolded protein response on human disease. J Cell Biol 197: 857–867.
- Blankenship JR, Fanning S, Hamaker JJ, Mitchell AP (2010) An extensive circuitry for cell wall regulation in Candida albicans. PLoS Pathog 6: e1000752.
- Cheon SA, Jung KW, Chen YL, Heitman J, Bahn YS, et al. (2011) Unique evolution of the UPR pathway with a novel bZIP transcription factor, Hxl1, for controlling pathogenicity of Cryptococcus neoformans. PLoS Pathog 7: e1002177
- Feng X, Krishnan K, Richie DL, Aimanianda V, Hartl L, et al. (2011) HacA-independent functions of the ER stress sensor IreA synergize with the canonical UPR to influence virulence traits in Aspergillus fumigatus. PLoS Pathog 7: e1002330.
- Richie DL, Feng X, Hartl L, Aimanianda V, Krishnan K, et al. (2011) The virulence of the opportunistic fungal pathogen Aspergillus fumigatus requires cooperation between the endoplasmic reticulum-associated degradation pathway (ERAD) and the unfolded protein response (UPR). Virulence 2: 12–21.
- Richie DL, Hartl L, Aimanianda V, Winters MS, Fuller KK, et al. (2009) A role for the unfolded protein response (UPR) in virulence and antifungal susceptibility in Aspergillus fumigatus. PLoS Pathog 5: e1000258.
- Wimalasena TT, Enjalbert B, Guillemette T, Plumridge A, Budge S, et al. (2008) Impact of the unfolded protein response upon genome-wide expression patterns, and the role of Hac1 in the polarized growth, of Candida albicans. Fungal Genet Biol 45: 1235–1247.
- Bernales S, Papa FR, Walter P (2006) Intracellular signaling by the unfolded protein response. Annu Rev Cell Dev Biol 22: 487–508.
- Cox JS, Shamu CE, Walter P (1993) Transcriptional induction of genes encoding endoplasmic reticulum resident proteins requires a transmembrane protein kinase. Cell 73: 1197–1206.
- Mori K, Ma W, Gething MJ, Sambrook J (1993) A transmembrane protein with a cdc2+/CDC28-related kinase activity is required for signaling from the ER to the nucleus. Cell 74: 743–756.
- Korennykh AV, Egea PF, Korostelev AA, Finer-Moore J, Zhang C, et al. (2009) The unfolded protein response signals through high-order assembly of Ire1. Nature 457: 687–693.
- Lee KP, Dey M, Neculai D, Cao C, Dever TE, et al. (2008) Structure of the dual enzyme Irel reveals the basis for catalysis and regulation in nonconventional RNA splicing. Cell 132: 89–100.
- Cox JS, Walter P (1996) A novel mechanism for regulating activity of a transcription factor that controls the unfolded protein response. Cell 87: 391– 404.
- Mori K, Kawahara T, Yoshida H, Yanagi H, Yura T (1996) Signalling from endoplasmic reticulum to nucleus: transcription factor with a basic-leucine zipper motif is required for the unfolded protein-response pathway. Genes Cells 1: 803–817.
- Bernales S, McDonald KL, Walter P (2006) Autophagy counterbalances endoplasmic reticulum expansion during the unfolded protein response. PLoS Biol 4: e423.
- Travers KJ, Patil CK, Wodicka L, Lockhart DJ, Weissman JS, et al. (2000) Functional and genomic analyses reveal an essential coordination between the unfolded protein response and ER-associated degradation. Cell 101: 249–258.
- Yorimitsu T, Nair U, Yang Z, Klionsky DJ (2006) Endoplasmic reticulum stress triggers autophagy. J Biol Chem 281: 30299–30304.
- Pfaller MA, Diekema DJ (2007) Epidemiology of invasive candidiasis: a persistent public health problem. Clin Microbiol Rev 20: 133–163.
- Li L, Redding S, Dongari-Bagtzoglou A (2007) Candida glabrata: an emerging oral opportunistic pathogen. J Dent Res 86: 204

  –215.
- Pfaller MA (2012) Antifungal drug resistance: mechanisms, epidemiology, and consequences for treatment. Am J Med 125: S3–13.
- Pfaller MA, Castanheira M, Messer SA, Moet GJ, Jones RN (2011) Echinocandin and triazole antifungal susceptibility profiles for Candida spp., Cryptococcus neoformans, and Aspergillus fumigatus: application of new CLSI clinical breakpoints and epidemiologic cutoff values to characterize resistance in the SENTRY Antimicrobial Surveillance Program (2009). Diagn Microbiol Infect Dis 69: 45–50.
- Urano F, Bertolotti A, Ron D (2000) IRE1 and efferent signaling from the endoplasmic reticulum. J Cell Sci 113 Pt 21: 3697–3702.
- Credle JJ, Finer-Moore JS, Papa FR, Stroud RM, Walter P (2005) On the mechanism of sensing unfolded protein in the endoplasmic reticulum. Proc Natl Acad Sci U S A 102: 18773–18784.
- 28. Gardner BM, Walter P (2011) Unfolded proteins are Ire1-activating ligands that directly induce the unfolded protein response. Science 333: 1891-1894.
- 29. Zhou J, Liu CY, Back SH, Clark RL, Peisach D, et al. (2006) The crystal structure of human IRE1 luminal domain reveals a conserved dimerization

- interface required for activation of the unfolded protein response. Proc Natl Acad Sci U S A 103: 14343–14348.
- 30. Papa FR, Zhang C, Shokat K, Walter P (2003) Bypassing a kinase activity with an ATP-competitive drug. Science 302: 1533–1537.
- Shamu CE, Walter P (1996) Oligomerization and phosphorylation of the Ire1p kinase during intracellular signaling from the endoplasmic reticulum to the nucleus. EMBO I 15: 3028–3039.
- Sidrauski C, Walter P (1997) The transmembrane kinase Irelp is a site-specific endonuclease that initiates mRNA splicing in the unfolded protein response. Cell 90: 1031–1039.
- 33. Chapman RE, Walter P (1997) Translational attenuation mediated by an mRNA intron. Curr Biol 7: 850–859.
- Gonzalez TN, Sidrauski C, Dorfler S, Walter P (1999) Mechanism of nonspliceosomal mRNA splicing in the unfolded protein response pathway. EMBO J 18: 3119–3132.
- Kawahara T, Yanagi H, Yura T, Mori K (1997) Endoplasmic reticulum stressinduced mRNA splicing permits synthesis of transcription factor Hac1p/Ern4p that activates the unfolded protein response. Mol Biol Cell 8: 1845–1862.
- Hooks KB, Griffiths-Jones S (2011) Conserved RNA structures in the noncanonical Hac1/Xbp1 intron. RNA Biol 8: 552–556.
- Calfon M, Zeng H, Urano F, Till JH, Hubbard SR, et al. (2002) IRE1 couples endoplasmic reticulum load to secretory capacity by processing the XBP-1 mRNA. Nature 415: 92–96.
- Frost A, Elgort MG, Brandman O, Ives C, Collins SR, et al. (2012) Functional Repurposing Revealed by Comparing S. pombe and S. cerevisiae Genetic Interactions. Cell 149: 1339–1352.
- Saloheimo M, Valkonen M, Penttila M (2003) Activation mechanisms of the HAC1-mediated unfolded protein response in filamentous fungi. Mol Microbiol 47: 1149–1161.
- 40. Guerfal M, Ryckaert S, Jacobs PP, Ameloot P, Van Craenenbroeck K, et al. (2010) The HAC1 gene from Pichia pastoris: characterization and effect of its overexpression on the production of secreted, surface displayed and membrane proteins. Microb Cell Fact 9: 49.
- Oh MH, Cheon SA, Kang HA, Kim JY (2010) Functional characterization of the unconventional splicing of Yarrowia lipolytica HAC1 mRNA induced by unfolded protein response. Yeast 27: 443

  –452.
- Shen X, Ellis RE, Lee K, Liu CY, Yang K, et al. (2001) Complementary signaling pathways regulate the unfolded protein response and are required for C. elegans development. Cell 107: 893–903.
- Yoshida H, Matsui T, Yamamoto A, Okada T, Mori K (2001) XBP1 mRNA is induced by ATF6 and spliced by IRE1 in response to ER stress to produce a highly active transcription factor. Cell 107: 881–891.
- Ryoo HD, Domingos PM, Kang MJ, Steller H (2007) Unfolded protein response in a Drosophila model for retinal degeneration. EMBO J 26: 242–252.
- Ruegsegger U, Leber JH, Walter P (2001) Block of HAC1 mRNA translation by long-range base pairing is released by cytoplasmic splicing upon induction of the unfolded protein response. Cell 107: 103–114.
- Sidrauski C, Cox JS, Walter P (1996) tRNA ligase is required for regulated mRNA splicing in the unfolded protein response. Cell 87: 405–413.
- Chawla A, Chakrabarti S, Ghosh G, Niwa M (2011) Attenuation of yeast UPR is essential for survival and is mediated by IRE1 kinase. J Cell Biol 193: 41–50.
- Bonilla M, Cunningham KW (2003) Mitogen-activated protein kinase stimulation of Ca(2+) signaling is required for survival of endoplasmic reticulum stress in yeast. Mol Biol Cell 14: 4296–4305.
- Bonilla M, Nastase KK, Cunningham KW (2002) Essential role of calcineurin in response to endoplasmic reticulum stress. EMBO J 21: 2343–2353.
- Chen Y, Feldman DE, Deng C, Brown JA, De Giacomo AF, et al. (2005) Identification of mitogen-activated protein kinase signaling pathways that confer resistance to endoplasmic reticulum stress in Saccharomyces cerevisiae. Mol Cancer Res 3: 669–677.
- Dudgeon DD, Zhang N, Ositelu OO, Kim H, Cunningham KW (2008) Nonapoptotic death of Saccharomyces cerevisiae cells that is stimulated by Hsp90 and inhibited by calcineurin and Cmk2 in response to endoplasmic reticulum stresses. Eukaryot Cell 7: 2037–2051.
- Miyazaki T, Yamauchi S, Inamine T, Nagayoshi Y, Saijo T, et al. (2010) Roles
  of calcineurin and Crz1 in antifungal susceptibility and virulence of Candida
  glabrata. Antimicrob Agents Chemother 54: 1639–1643.
- Ogawa N, Mori K (2004) Autoregulation of the HAC1 gene is required for sustained activation of the yeast unfolded protein response. Genes Cells 9: 95– 104
- Chen YL, Konieczka JH, Springer DJ, Bowen SE, Zhang J, et al. (2012)
   Convergent Evolution of Calcineurin Pathway Roles in Thermotolerance and Virulence in Candida glabrata. G3 (Bethesda) 2: 675–691.
- Kaufman RJ (1999) Stress signaling from the lumen of the endoplasmic reticulum: coordination of gene transcriptional and translational controls. Genes Dev 13: 1211–1233.
- Kohno K, Normington K, Sambrook J, Gething MJ, Mori K (1993) The promoter region of the yeast KAR2 (BiP) gene contains a regulatory domain that responds to the presence of unfolded proteins in the endoplasmic reticulum. Mol Cell Biol 13: 877–890.

- Mori K, Ogawa N, Kawahara T, Yanagi H, Yura T (1998) Palindrome with spacer of one nucleotide is characteristic of the cis-acting unfolded protein response element in Saccharomyces cerevisiae. J Biol Chem 273: 9912–9920.
- Mori K, Sant A, Kohno K, Normington K, Gething MJ, et al. (1992) A 22 bp cis-acting element is necessary and sufficient for the induction of the yeast KAR2 (BiP) gene by unfolded proteins. EMBO J 11: 2583–2593.
- Miyazaki T, Izumikawa K, Yamauchi S, Inamine T, Nagayoshi Y, et al. (2011) The glycosylphosphatidylinositol-linked aspartyl protease Yps1 is transcriptionally regulated by the calcineurin-Crz1 and Slt2 MAPK pathways in Candida glabrata. FEMS Yeast Res 11: 449–456.
- Yoshimoto H, Saltsman K, Gasch AP, Li HX, Ogawa N, et al. (2002) Genomewide analysis of gene expression regulated by the calcineurin/Crz1p signaling pathway in Saccharomyces cerevisiae. J Biol Chem 277: 31079–31088.
- Rubio C, Pincus D, Korennykh A, Schuck S, El-Samad H, et al. (2011) Homeostatic adaptation to endoplasmic reticulum stress depends on Irel kinase activity. J Cell Biol 193: 171–184.
- Han D, Lerner AG, Vande Walle L, Upton JP, Xu W, et al. (2009) IRE1alpha kinase activation modes control alternate endoribonuclease outputs to determine divergent cell fates. Cell 138: 562–575.
- Hollien J, Lin JH, Li H, Stevens N, Walter P, et al. (2009) Regulated Ireldependent decay of messenger RNAs in mammalian cells. J Cell Biol 186: 323– 331
- Hollien J, Weissman JS (2006) Decay of endoplasmic reticulum-localized mRNAs during the unfolded protein response. Science 313: 104–107.
- Nevitt T, Thiele DJ (2011) Host iron withholding demands siderophore utilization for Candida glabrata to survive macrophage killing. PLoS Pathog 7: e1001322.
- 66. Scrimale T, Didone L, de Mesy Bentley KL, Krysan DJ (2009) The unfolded protein response is induced by the cell wall integrity mitogen-activated protein kinase signaling cascade and is required for cell wall integrity in Saccharomyces cerevisiae. Mol Biol Cell 20: 164–175.
- Miyazaki T, Inamine T, Yamauchi S, Nagayoshi Y, Saijo T, et al. (2010) Role of the Slt2 mitogen-activated protein kinase pathway in cell wall integrity and virulence in Candida glabrata. FEMS Yeast Res 10: 343–352.
- Mori K (2009) Signalling pathways in the unfolded protein response: development from yeast to mammals. J Biochem 146: 743–750.
- Walter P, Ron D (2011) The unfolded protein response: from stress pathway to homeostatic regulation. Science 334: 1081–1086.
- Pavitt GD, Ron D (2012) New insights into translational regulation in the endoplasmic reticulum unfolded protein response. Cold Spring Harb Perspect Biol 4: pii: a012278.
- Kimata Y, Ishiwata-Kimata Y, Yamada S, Kohno K (2006) Yeast unfolded protein response pathway regulates expression of genes for anti-oxidative stress and for cell surface proteins. Genes Cells 11: 59–69.

- Babour A, Bicknell AA, Tourtellotte J, Niwa M (2010) A surveillance pathway monitors the fitness of the endoplasmic reticulum to control its inheritance. Cell 142: 256–269.
- Rutkowski DT, Hegde RS (2010) Regulation of basal cellular physiology by the homeostatic unfolded protein response. J Cell Biol 189: 783–794.
- Thibault G, Ismail N, Ng DT (2011) The unfolded protein response supports cellular robustness as a broad-spectrum compensatory pathway. Proc Natl Acad Sci U S A 108: 20597–20602.
- Torres-Quiroz F, Garcia-Marques S, Coria R, Randez-Gil F, Prieto JA (2010)
   The activity of yeast Hog1 MAPK is required during endoplasmic reticulum stress induced by tunicamycin exposure. J Biol Chem 285: 20088–20096.
- Oda Y, Okada T, Yoshida H, Kaufman RJ, Nagata K, et al. (2006) Derlin-2 and Derlin-3 are regulated by the mammalian unfolded protein response and are required for ER-associated degradation. J Cell Biol 172: 383–393.
- Kaiser C, Michaelis S, Mitchell A (1994) Laboratory Course Manual for Methods in Yeast Genetics. Cold Spring Harbor, New York: Cold Spring Harbor Laboratory Press.
- Budovskaya YV, Stephan JS, Reggiori F, Klionsky DJ, Herman PK (2004) The Ras/cAMP-dependent protein kinase signaling pathway regulates an early step of the autophagy process in Saccharomyces cerevisiae. J Biol Chem 279: 20663– 20671.
- Cormack BP, Falkow S (1999) Efficient homologous and illegitimate recombination in the opportunistic yeast pathogen Candida glabrata. Genetics 151: 979

  987
- 80. Dujon B, Sherman D, Fischer G, Durrens P, Casaregola S, et al. (2004) Genome evolution in yeasts. Nature 430: 35–44.
- Winzeler EA, Shoemaker DD, Astromoff A, Liang H, Anderson K, et al. (1999)
   Functional characterization of the S. cerevisiae genome by gene deletion and parallel analysis. Science 285: 901–906.
- Pfaller MA, Arikan S, Lozano-Chiu M, Chen Y, Coffman S, et al. (1998) Clinical evaluation of the ASTY colorimetric microdilution panel for antifungal susceptibility testing. J Clin Microbiol 36: 2609–2612.
- 83. Irizarry RA, Bolstad BM, Collin F, Cope LM, Hobbs B, et al. (2003) Summaries of Affymetrix GeneChip probe level data. Nucleic Acids Res 31: e15.
- Irizarry RA, Hobbs B, Collin F, Beazer-Barclay YD, Antonellis KJ, et al. (2003) Exploration, normalization, and summaries of high density oligonucleotide array probe level data. Biostatistics 4: 249–264.
- Boyle EI, Weng S, Gollub J, Jin H, Botstein D, et al. (2004) TermFinder-open source software for accessing Gene Ontology information and finding significantly enriched Gene Ontology terms associated with a list of genes. Bioinformatics 20: 3710–3715.
- El Barkani A, Haynes K, Mosch H, Frosch M, Muhlschlegel FA (2000) Candida glabrata shuttle vectors suitable for translational fusions to lacZ and use of betagalactosidase as a reporter of gene expression. Gene 246: 151–155.
- Miller JH (1972) Experiments in Molecular Genetics. Cold Spring Harbor, New York: Cold Spring Harbor Laboratory Press.