Roles of the pro-apoptotic factors CaNma111 and CaYbh3 in apoptosis and virulence of Candida albicans

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Candida albicans, a commensal and opportunistic pathogen, undergoes apoptosis in response to various stimuli, including hydrogen peroxide, acetic acid, and antifungal agents. Apoptotic processes are highly conserved among mammals, plants, and fungi, but little is known about the apoptosis-regulating factors in C. albicans. In this study, C. albicans homologs of the putative apoptosis factors were identified by database screening followed by overexpression analysis. CaNma111, a homolog of the pro-apoptotic mammalian HtrA2/Omi, and CaYbh3, a homolog of BH3-only protein, yielded increased apoptotic phenotypes upon overexpression. We showed that CaNma111 and CaYbh3 functions as pro-apoptotic regulators by examining intracellular ROS accumulation, DNA end breaks (TUNEL assay), and cell survival in Canma111/Canma111 and Caybh3/Caybh3 deletion strains. We found that the protein level of CaBir1, an inhibitor-of-apoptosis (IAP) protein, was down-regulated by CaNma111. Interestingly, the Canma111/Canma111 and Caybh3/Caybh3 deletion strains showed hyperfilamentation phenotypes and increased virulence in a mouse infection model. Together, our results suggest that CaNma111 and CaYbh3 play key regulatory roles in the apoptosis and virulence of C. albicans.

Apoptosis is a form of programmed cell death, that is highly conserved in mammals, plants, and fungi, including unicellular yeasts. The pathogenic yeast Candida albicans exhibits typical apoptotic markers when treated with various stimuli, including hydrogen peroxide (H$_2$O$_2$), acetic acid, and UV irradiation$^1$-$^3$. In addition, antifungal agents, such as amphotericin B and the quorum-sensing molecule, farnesol, can induce apoptosis$^4$-$^5$. The metacaspase CaMca1 is a caspase-related protease in C. albicans, that shows homology to the yeast Saccharomyces cerevisiae metacaspase, Yc1$^6$-$^9$. Metacaspases are known to be involved in the stress-induced cell death of the yeasts, S. cerevisiae and C. albicans, the plant, Arabidopsis thaliana, the fungal species, Aspergillus nidulans, the protozoa, Leishmania major$^2$,$^6$-$^8$,$^{10}$. Metacaspases are distinguished from mammalian caspases by various biochemical features, including their proteolytic processing ability and/or Arg/Lys substrate specificity.

In mammals, the activation or regulation of caspases requires various pro- and anti-apoptotic proteins, including the Bcl-2 (B-cell lymphoma) family members (Bax, Bak, Bcl-2, and Bcl-xL) and the inhibitor-of-apoptosis proteins (IAPs)$^{11}$-$^{17}$. S. cerevisiae appears to lack homologs of the Bcl-2 proteins with the exception of the yeast BH3-only protein, Ybh3$^{18}$-$^{14}$. A few other apoptotic regulators have been identified in S. cerevisiae, including the single IAP (inhibitor-of-apoptosis), Bir1, the pro-apoptotic protease, Omi/HtrA2 (Nma111) and an apoptosis-inducing factor (Aif1)$^{15}$-$^{18}$. In C. albicans, little is known about the regulation of apoptosis or metacaspase activation. Recently, a single IAP, CaBir1, was identified to inhibit apoptosis by lowering intracellular caspase-like activity in C. albicans$^{17}$-$^{19}$. The Cabir1/Cabir1 deletion mutant showed increased apoptotic features, including ROS accumulation and nuclear segmentation.

To investigate the regulatory mechanisms underlying apoptotic processes in C. albicans, we searched for putative pro-apoptotic or anti-apoptotic regulators by employing an overexpression strategy. Among five candidates screened, CaNma111 and CaYbh3 exhibited pro-apoptotic activity and were further characterized by constructing the deletion mutant strains. We also showed hyperfilamentation phenotypes and increased virulence of the Canma111/Canma111 and Caybh3/Caybh3 deletion strains.

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Results

Overexpression of putative apoptosis factors in C. albicans. To investigate putative apoptosis-regulating factors in C. albicans, we constructed overexpression strains of five genes: CaBIR1, CaNMA111, CaYBH3, CaDHH1 and CaPAT1 (Fig. 1). These genes were identified from the Candida Genome Database based on their amino acid sequence similarities with homologous proteins in S. cerevisiae and mammals. CaBir1, a single IAP in C. albicans, was shown to inhibit apoptosis by reducing caspase-like activity under an oxidative stress condition. CaBir1 and its homolog in S. cerevisiae, CaBir1 (nuclear mediator of apoptosis), is a homolog of the pro-apoptotic serine protease, HtrA2/Omi. In mammals and S. cerevisiae, HtrA2/Omi regulates apoptosis by binding and degrading cellular IAPs. Ybh3 is the yeast homolog of the BH3-only protein, which contains a BCL-2 homology domain. Ybh3 was identified in a previous study as a member of the BH3-only family. Ybh3, which is known as the components of P-bodies (processing bodies, mRNA granules) in S. cerevisiae, function as mRNA-decapping activators. CaDhh1, which is a homolog of CaDhh1, was identified in a previous work, shown to be localized to P-bodies in C. albicans. CaPat1 was identified as a CaPat1 homolog in this study.

For ectopic overexpression, each target gene was cloned downstream of the ACT1 promoter in plasmid pPR671, and the constructed plasmid was chromosome-integrated into the wild-type C. albicans strain.

Overexpression of CaNma111 or CaYbh3 yields increased apoptotic phenotypes. Apoptosis is characterized by several morphological and biochemical features including chromatin condensation, accumulation of reactive oxygen species (ROS), and increased caspase activity. In each overexpression strain, we determined the intracellular amount of ROS by staining cells with the fluorescent dye, H2DCFDA (Fig. 2). Compared to wild-type BWP17 cells, overexpression strains OEcaNma111 and OEcaYbh3 showed increased frequencies of H2DCFDA-positive cells before and after apoptosis-inducing H2O2 treatment. OEcaBIR1 and OEcaPat1 showed decreased ROS accumulation compared to the wild-type. OEcaDhh1 showed a ROS level higher than that of wild-type cells prior to H2O2 treatment but similar to that of wild-type cells after H2O2 treatment.

Next, we measured the caspase-like activity in the overexpression strains using the caspase substrate, D2R. In the wild-type strain, few cells were fluorescent in H2O2-untreated cells, but numerous fluorescent cells were observed after 30 min of H2O2 stress. OEcaNma111 and OEcaYbh3 showed increased frequencies of fluorescent cells compared to the wild-type strain with or without oxidative stress. In contrast, OEcaBIR1 and OEcaPat1 showed little fluorescence, regardless of H2O2 treatment. OEcaDhh1 showed an increase in the number of fluorescent cells before H2O2 treatment but only a slight increase after H2O2 treatment. These patterns of caspase-like activity in each overexpression strain were closely associated with the ROS accumulation level.

Deletion of CaNma111 or CaYbh3 decreased apoptotic phenotypes. To further investigate the roles of CaNma111 and CaYbh3 in apoptosis, we sequentially deleted the two copies of CaNma111 or CaYbh3 to construct the Canma111/Canma111 and Caybh3/Caybh3 deletion strains, respectively. The wild-type and deletion strains were compared for apoptotic hallmarks, including ROS accumulation, nuclear...
segmentation (TUNEL assay), and cell survival under oxidative stress (Fig. 4). Cell survival after H\textsubscript{2}O\textsubscript{2} treatment was much higher in Canma111/Canma111 and Caybh3/Caybh3 mutant cells than wild-type cells (Fig. 4A, B). The amount of ROS, which was determined using the fluorescent dye, H\textsubscript{2}DCFDA, was lower in Canma111/Canma111 and Caybh3/Caybh3 cells than wild-type cells upon H\textsubscript{2}O\textsubscript{2} treatment. The TUNEL assay, which measures DNA breaks, revealed that Canma111/Canma111 and Caybh3/Caybh3 mutant cells showed lower frequencies of TUNEL-positive nuclei than wild-type cells upon H\textsubscript{2}O\textsubscript{2} treatment. The mutant strains showed decreased caspase-like activity compared to wild-type cells following treatment with H\textsubscript{2}O\textsubscript{2} for 30 min (Fig. 4C, F). These results collectively suggest that CaNma111 and CaYbh3 are required for apoptotic cell death in C. albicans.

CaNma111 downregulates the apoptosis inhibitor, CaBir1. The mammalian serine protease, Omi/HtrA2, promotes apoptosis by binding and degrading IAP family proteins\textsuperscript{25,26}. Consistently, the yeast IAP, Bir1, was shown to be a substrate for Nma111 in S. cerevisiae\textsuperscript{15}. We repeatedly observed a very faint protein band when we assessed chromosome-tagged CaBir1-GFP or CaBir1-myc in a wild-type background (data not shown). To ask whether CaNma111 is one of the proteases responsible for the degradation of CaBir1, we compared CaBir1-myc levels in wild-type and Canma111/Canma111 cells. Here, CaBir1-myc was expressed under the control of the ACT1 promoter of the pPR671 vector. We observed an increased level of CaBir1-myc in Canma111/
Canma111 cells, compared to wild-type cells (Fig. 4G). This result suggests that CaNma111 downregulates CaBir1 in *C. albicans*.

**Deletion of CaNMA111 or significantly increases filamentous growth and virulence.** *C. albicans* is an opportunistic pathogen and switches rapidly among the budding yeast, pseudohyphal, and hyphal forms in response to environmental changes. This morphogenetic switching is particularly associated with virulence. In addition, it has been suggested that the morphological state affects apoptotic cell death.

We therefore examined whether the pro-apoptotic regulators, CaNma111 and CaYbh3, are involved in the filamentous growth or virulence of *C. albicans*. Colony morphologies of the wild-type, Canma111/Canma111, and Caybh3/Caybh3 strains were examined on hyphae-inducing solid medium. As shown in Fig. 5A, Canma111/Canma111 and Caybh3/Caybh3 mutant cells exhibited hyperfilamentation phenotypes on solid Spider medium, compared with wild-type cells. Interestingly, the mutant strains showed filamentous growth on YEPD complete
Figure 4. The apoptotic phenotypes of Canma111/Canma111 and Caybh3/Caybh3 mutant strains. (A) Survival of the wild-type, Canma111/Canma111, and Caybh3/Caybh3 strains was examined by spot assays. The cells were grown to early log phase and treated with 7.5 mM H2O2 for 2 h. Serial dilutions of cells were spotted onto YEPD plates, which were incubated at 30 °C and photographed after 2 days. (B) Relative viabilities of the wild-type, Canma111/Canma111, and Caybh3/Caybh3 strains were determined upon H2O2 treatment. Cells in early log phase were treated with 5 mM H2O2 for the indicated time. Culture samples were diluted and plated in duplicate. Viabilities were scored as a percentage of the number of colonies formed at time zero. (C) Fluorescence microscopy of ROS, TUNEL, or D2R staining of the wild-type, Canma111/Canma111, and Caybh3/Caybh3 strains following treatment with H2O2. Cells were observed under an Olympus BX51 microscope with a 60 × objective. Scale bar, 5 μm. (D) Graphs represent the quantification of ROS-stained cells (%) (n = 3 replicates, >200 cells). Values are presented as the mean ± SD; ** p < 0.01. (E) Quantification of TUNEL-positive cells are graphed (n = 3 replicates, >200 cells). TUNEL assays were carried out after cells were exposed to 7.5 mM H2O2 for 2.5 h. The percentages of values are presented as mean ± SD; ** p < 0.01, *** p < 0.005. (F) Graphs represent the quantification of D2R-stained cells (%) (n = 3 replicates, >200 cells). Cells were treated with 10 mM H2O2 for 30 min. Values are mean ± SD. * p < 0.05. (G) Detection of the CaBir1-myc protein band in the Canma111/Canma111 mutant strain. The PR671-derived ACT1-CaBIR1-MYC construct was chromosomally integrated in the wild-type and Canma111/Canma111 mutant strains. Western blotting was conducted using anti-myc antibody. Tubulin was detected as a loading control.
medium (data not shown). The hyperfilamentation phenotypes of Canma111/Canma111 and Caybh3/Caybh3 mutant strains were also evident in liquid medium supplemented with 10% serum (Fig. 5B).

The virulence of the Canma111/Canma111 and Caybh3/Caybh3 strains was tested in a tail vein-infection model with BALB/c mice. Two groups of mice (n = 10) were challenged with wild-type and mutant cells and survival was monitored for up to 30 days. Mice infected with Canma111/Canma111 or Caybh3/Caybh3 mutant strain showed more rapid weight loss and decreased survival than those infected with wild-type BWP17 (Fig. 5C). These results demonstrate that CaNMA111 and CaYBH3 play important roles in the virulence of C. albicans.

Discussion

We analyzed five candidate regulators for apoptosis in C. albicans and found that overexpression of CaNma111 or CaYbh3 yielded pro-apoptotic features, while that of CaBir1 or CaPat1 yielded anti-apoptotic feature. CaNma111 and CaYbh3 were further characterized by constructing the deletion mutant strains. Gene overexpression mimics gain-of-function mutations, and thus offers a useful approach for revealing pathways or pathway components in the diploid pathogen, C. albicans30–32. The overexpression phenotypes of the apoptosis inhibitor, CaBir1, were consistent with our recent report that the Cabir1/Cabir1 mutant strain showed increased apoptotic phenotypes, such as ROS accumulation and DNA fragmentation, under apoptosis-inducing conditions19. We analyzed CaDhh1 and CaPat1 because our previous work showed that CaEdc3, another component of P-bodies, is involved in apoptosis33. CaEdc3 contributes to the expression of CaMca1 expression and thereby functions as a pro-apoptotic factor. In our overexpression analysis, CaPat1 was suggested to be anti-apoptotic factor. We suggest that CaDhh1, CaEdc3, and CaPat1 could all participate in apoptosis, with each playing a distinct role. CaDhh1 and CaPat1 show protein interactions with each other but differ in their functional domains, intracellular locations, and mRNA targets30,34.

Here, we report our results from the deletion mutant analysis of CaNMA111 and CaYBH3. The decreased apoptotic phenotypes of Canma111/Canma111 and Caybh3/Caybh3 mutant cells suggest that CaNma111 and CaYbh3 function as pro-apoptotic regulators in C. albicans. HtrA2/Omi, which is a mammalian counterpart of CaNma111, has been identified as a direct IAP-binding protein16,26. It exerts pro-apoptotic character effects, possibly by disruption of the IAP-caspase interaction. Studies have shown that the serine protease, HtrA2/Omi,
can degrade mammalian IAP and XIAP\textsuperscript{25,26}. We repeatedly observed very faint protein band corresponding to CaBir1-GFP or CaBir1-myc in a wild-type background (data not shown). We speculated that the full-length CaBir1 protein could be a target of proteolytic degradation. Our observation that the CaBir1-myc protein level was increased in \textit{Canma111/Canma111} cells may support this notion. However, future work is needed to assess whether CaNma111 could be among the proteases responsible for CaBir1 degradation.

The hyperfilamentation phenotypes and increased virulence of the \textit{Canma111/Canma111} and \textit{Caybh3/Caybh3} mutant strains were particularly interesting, as these findings suggest that the pro-apoptotic regulators, CaNma111 and CaYbh3, exert repressive actions on filamentation and pathogenicity in \textit{C. albicans}. It remains unknown whether the pro-apoptotic roles of CaNma111 and CaYbh3 overlap with their functions during morphogenesis. One possible explanation is that CaNma111, which is a serine protease responsible for degrading the apoptosis inhibitor, CaBir1, could be involved in the processing or breakdown of regulatory factors crucial for filamentous growth. Further studies will be needed to uncover the downstream targets of CaNma111 or CaYbh3 protease activity during morphogenesis. Regarding CaYbh3, we speculate that a putative BH3 domain within this \textit{C. albicans} protein could be responsible for mitochondria-driven ROS accumulation and/or the release of apoptotic factors\textsuperscript{43}. During hyphal morphogenesis, \textit{C. albicans} produces a burst of ROS that is mainly located at the hyphal tip\textsuperscript{35,36}. Further investigation is needed to examine whether the repressive function of CaYbh3 during filamentous growth could also be associated with changes in the ROS level.

It is noteworthy that the pro-apoptotic regulator, CaMca1 metacaspase, was shown to be required for filamentation and pathogenicity\textsuperscript{37}. Cells harboring the apoptosis-defective deletion of \textit{CaMCA1} or the catalytic-site mutation \textit{CaMCA1}\textsuperscript{292}, showed defects in filamentation and virulence. It has been suggested that \textit{S. cerevisiae} metacaspases, which are responsible for apoptosis, are also involved in nonapoptotic characteristics and processes, such as longevity, the fitness of growing cells, and protein clearance\textsuperscript{38,39}. We speculate that the downstream targets of CaNma111 protease and CaMca1 metacaspase could act to either promote or repress filamentous growth and other nonapoptotic processes.

Various regulatory elements involved in the yeast-to-hyphal transition has been identified in \textit{C. albicans}\textsuperscript{27,40,41}. The Ras-CAMP-PKA and the MAPK pathway operate to promote the yeast-to-hyphal transition and the transcription factors, such as Cph1 and Efg1, are targets of these pathway responses in \textit{C. albicans}. Activation of Ras-signaling was shown to accelerate apoptotic responses under treatment with acetic acid or H\textsubscript{2}O\textsubscript{2}\textsuperscript{42}. However, little is known about the interrelationship between cell death and morphogenesis. The quorum-sensing molecule, farnesol, inhibits the yeast-to-hyphal switch, but this triggers apoptosis\textsuperscript{43}. Going forward, additional work is needed to improve our understanding of the detailed regulatory points and components involved in the apoptotic responses and pathogenicity of \textit{C. albicans}.

Materials and methods

\textbf{Strains, plasmids, and culture conditions.} The \textit{C. albicans} strains and plasmids used in this study are listed in Table S1. Constructions of the \textit{Canma111/Canma111} and \textit{Caybh3/Caybh3} deletion strains were essentially as described previously\textsuperscript{25,40}. We used plasmids pJl434 and pJl435 for \textit{Canma111} deletion, and plasmids pJl436 and pJl437 for \textit{CaYbh3} deletion. These plasmids carried the deletion cassettes, \textit{hph-URA3-hph} and \textit{hisG-URA3-hisG}, respectively. Each disruption was verified by PCR. Overexpression strains were constructed using the pPR671-derived plasmids, pJl426-pJl432. Each target gene was amplified using a primer set (Table S2), and the PCR fragment was digested with \textit{Mlu}\textsubscript{I} and ligated into the \textit{Mlu}\textsubscript{I} and \textit{Xma}\textsubscript{I} sites of pPR671\textsuperscript{22}. Each pPR671-derived plasmid was linearized by \textit{Stu}\textsubscript{I} and transformed into the wild-type BPW17 strain. Chromosome integrations were verified by PCR and protein expressions were analyzed by Western blot.

\textit{C. albicans} strains were cultured in YEPD (1% yeast extract, 2% peptone, 2% dextrose) or SC (synthetic complete; 0.67% yeast nitrogen base w/o amino acid, 2% glucose, all required amino acids) medium. The filamentation phenotype of \textit{C. albicans} cells was tested in serum-containing medium (YEPD with 10% new born calf serum) and Spider medium (1% mannitol, 1% nutrient broth, 0.2% K\textsubscript{2}HPO\textsubscript{4}, pH7.2) as described previously\textsuperscript{44}. Yeast cells were fixed with 3.7% formaldehyde, digested with 12 μg/ml zymolyase 100 T (106 units/g; US Biological, USA) at 30 °C for 45 min, and applied to a poly-lysine-coated slide. Each slide was rinsed with PBS and incubated in permeabilization solution (0.1% Triton X-100, 0.1% sodium citrate) for 2 min on ice. The slides were incubated with a TUNEL reaction mixture containing terminal deoxynucleotidyl transferase and DNA strand breaks were demonstrated by TUNEL (TdT-mediated DUTP nick end labeling) assay using an In Situ Cell Death Detection kit (Roche Molecular Biochemicals, Germany), as described previously\textsuperscript{44}. Yeast cells were fixed with 3.7% formaldehyde, digested with 12 μg/ml zymolyase 100 T (10 units/g; US Biological, USA) at 30 °C for 45 min, and applied to a poly-lysine-coated slide. Each slide was rinsed with PBS and incubated in permeabilization solution (0.1% Triton X-100, 0.1% sodium citrate) for 2 min on ice. The slides were incubated with a TUNEL reaction mixture containing terminal deoxynucleotidyl transferase and
FITC-labeled dUTP, and mounted with a drop of VECTASHIELD antifading agent (Vector Laboratories Inc., USA). Observations were made with an Olympus BX51 microscope equipped with a 60× objective.

**Western blot analysis.** Total protein preparation and Western blotting were performed as previously described. Myc-tagged proteins were detected with anti-myc antibody (Roche, USA); HRP-conjugated anti-mouse IgG antibody (Santa Cruz Biotechnology, USA) was used as the secondary antibody. Tubulin protein was used as a loading control, and was detected with a monoclonal anti-a-tubulin antibody (Sigma-Aldrich, USA). Protein bands were visualized using an Enhanced Peroxidase Detection (EPD) Western reagent kit (Eliptic-Biotech, KR).

**Assessment of virulence in a murine infection model.** Cells were grown overnight in SC-Ura medium and washed twice with sterile physiological saline. Seven-week-old female BALB/c mice were infected via lateral tail vein injection with 6 x 10^6 CFU (colony forming unit) in a 100-μl volume. Ten mice were inoculated per test strain, and host survival was monitored over 30 days. All animal experiments were approved by the Animal Experiment Ethics Committee of Chungnam National University (approval No. 202006A-CNU-120, July 2020) and performed in accordance with the guidelines of the Ethics Training Guidelines for Experiments on Animals of CNU Animal Research Center. This study additionally adheres to standards articulated in the ARRIVE guidelines.

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Author contributions

M.N., S.H.K.: Conceptualization, Methodology, Experimental work, Data collection and analysis, Writing-original draft. J. K.: Project planning, Funding acquisition, Data analysis and interpretation, Writing- Reviewing and Editing. J.-H. J., S.-Y. K.: Experimental work, Data collection and analysis. All authors reviewed and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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