The Transient Receptor Potential Channel Yvc1 Deletion Recovers the Growth Defect of Calcineurin Mutant Under Endoplasmic Reticulum Stress in Candida albicans

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INTRODUCTION

Candida albicans, as the most well-known pathogenic fungi, may cause lethal systemic infection of humans (Dantas Ada et al., 2015). Calcium ion is an important signal messenger, which serves various activities and structural functions in all eukaryote cells. Normally, the cytosolic calcium ions are maintained at low concentrations in C. albicans, in which some calcium pumps and calcium exchangers play an important role (Cagnac et al., 2010; Ghanegolmohammadi et al., 2017). However, in response to specific stress, the extracellular calcium ions enter into the cytoplasm through the high-affinity calcium uptake system (HACS) (Cyert and Philpott, 2013). HACS consists of Cch1, Mid1, and Ecm7, which interact with each other (Iida et al., 2017). Besides, the organelle...
vacuole is the natural calcium ion pool in yeast cells, containing over 90% of the total cellular calcium ions (Bianchi et al., 2019). Under environmental stress, the calcium ions stored in vacuolar cavity are released through transient receptor potential (TRP) channel Yvc1 into the cytosol (Palmer et al., 2001).

In *C. albicans*, calcineurin is the conserved Ca\(^{2+}\)/calmodulin-dependent phosphatase, composed of catalytic subunit Cna1 and regulatory subunit Cnb1 (Connolly et al., 2018). By dephosphorylation of transcription factor Crz1, calcineurin triggers downstream signaling events and regulates the cytosolic ion concentration (Chow et al., 2017; Xu et al., 2020). In *Saccharomyces cerevisiae*, calcineurin is activated when exposed to severe stimulation, such as high pH, cell membrane damage, or antifungal drugs (LaFayette et al., 2010; Li Y. et al., 2018). The most important is that calcineurin regulated CCH1 negatively, which inhibits the expression of Cch1 once the cellular calcium ions are overloading (Karababa et al., 2006; Teng et al., 2013; Xu et al., 2019). Yvc1 is the unique TRP-type calcium ion channel in yeast cells, a homolog with the TRP family of mammalian cells (de Castro et al., 2014). CaYvc1, similar with ScYvc1, located on the vacuolar membrane, is important for the calcium transport under environmental stress. In the previous study, we found that the deletion of YVC1 caused hypersensitivity to oxidative stress (Yu et al., 2014b). Meanwhile, Yvc1 is important in the process of hyphal growth, and its specific localization on the vacuolar membrane is necessary for the normal function of V-ATPase in *C. albicans* (Peng et al., 2019, 2020).

Generally, the proteins are synthesized, folded, and secreted in the endoplasmatic reticulum (ER) (Zhang et al., 2019). Tunicamycin (TN) inhibits N-glycosylation and blocks the formation of glycoprotein, thereby leading to ER stress (Cherepanova et al., 2019). The unfolded protein response (UPR) is the typical strategy in yeast cells for relieving the stress (Bravo et al., 2013). bZIP transcription factor Hac1, a homolog with Xbp1 in mammalian cells, is spliced at the C terminus and transported from cytosol into nucleus, triggering the immediate expression of Pmt4 or Prb1 to alleviate the ER stress (Cheon et al., 2011; Cherry et al., 2019). In mammalian cells, the nuclear factor E2-related factor 2 (Nrf2) was activated under oxidative stress, thereby modulating ER calcium levels by the regulation of glutathione peroxidase (Granatiero et al., 2019). Besides, since ER stress contributes to intracellular calcium and stress response, overloaded calcium induced mitochondrial dysfunction in cardiac, especially complex I (Mohsin et al., 2020).

In this study, we found that cnb1Δ/Δ was sensitive to ER stress, which was related with the overexpression of cytoplasm membrane channel CCH1 and irrelevant to the oxidative stress reaction. Besides, we found that the growth of the cnb1Δ/ΔΔvc1Δ/Δ strain was faster than that of the cnb1Δ/Δ strain, and the cell death rate, vacuolar membrane permeability, and mitochondrial activity were relieved in the double mutant. Interestingly, the classical UPR pathway was activated normally in all of the strains, indicating that the mechanism of relieving the growth in cnb1Δ/ΔΔvc1Δ/Δ was unrelated with the UPR pathway. However, the calcium flux was enhanced and its concentration was increased in the cnb1Δ/Δ strain under ER stress, and their level decreased obviously in the cnb1Δ/ΔΔvc1Δ/Δ strain. CaCl\(_2\) or its chelating reagent EGTA was added to verify the regulatory role of Yvc1 in the calcium ion concentration, and we found that the addition of CaCl\(_2\) led to poor viability and weakened the functions of vacuole and mitochondria under ER stress of the cnb1Δ/Δ strain. However, the growth state or organelle activity was relieved under the treatment of EGTA, indicating that Yvc1 alters the cellular calcium concentration in response to ER stress to improve the cnb1Δ/Δ growth. Overall, our work shed a novel light on the interaction between Yvc1-mediated calcium homeostasis and ER stress response in *C. albicans*.

**MATERIALS AND METHODS**

**Strains and Culture Conditions**

The strains and primers used in our study are listed in Tables 1, 2. Wild-type (WT) strain BWP17 was used as the background strain to construct the cnb1Δ/Δ, yvc1Δ/Δ, and cnb1Δ/ΔΔvc1Δ/Δ mutant strains by the PCR-mediated homologous recombination method. The ARG4 cassettes were amplified and transformed into the WT. The heterozygous mutants (cnb1Δ/ARG4/CNB1) were identified by PCR. After that, the URA3 fragment was transformed into the heterozygous mutant above to construct the cnb1Δ/Δ mutant strains (cnb1Δ/ARG4/cnb1Δ/URA3). The yvc1Δ/Δ and double mutant cnb1Δ/ΔΔvc1Δ/Δ were constructed with a similar method.

Besides, the pPCCH1-GFP plasmid was digested with Stul for 1 h and transferred into the WT and other mutant strains to measure the CCH1 expression level. In general, the strains were cultured in YPD (10 g/l yeast extract, 20 g/l peptone, 20 g/l glucose) medium. The SC (2% glucose, 0.67% yeast nitrogen base, 0.2% amino acid mixture) medium without uracil was used to separate and select the URA3-tagged strains.

**Spot Assay**

The YPD plates containing different concentrations of TN, β-mercaptoethanol, dithiothreitol (DTT), or calcofluor white (CFW) were used to measure the sensitivity to ER stress or cell wall stress (Su et al., 2021). Besides, 5 mM reductive agent ascorbic acid (VG) was added into these stress-related plates to investigate the relationship between stress susceptibility and oxidative stress response (OSR).

**Cellular Calcium Levels**

The content of cellular calcium and the calcium flux were measured with a Fluo-4 (C\(_{50}\)H\(_{90}\)F\(_{2}\)N\(_{20}\)O\(_{23}\), Beyotime, Shanghai, China, dissolved in DMSO) probe (Bartoli et al., 2019). Log-phase cells were treated with 2 μg/ml TN for 2 h. The cells were collected, washed with phosphate-buffered saline (PBS, 8 g/l NaCl, 0.2 g/l KCl, 1.42 g/l Na\(_{2}\)HPO\(_{4}\), 0.27 g/l KH\(_{2}\)PO\(_{4}\), pH 7.4), and resuspended with PBS buffer. The 2-mM Fluo-4 probe was added and incubated with 70 r/min at 30° C for 1 h. The fluorescence intensity (excitation wavelength at 488 nm, emission wavelength at 525 nm) was detected using a fluorescent microplate reader (Bode et al., 2020). The scanning time was sustained for 7 min with a scan gap for 1 s.
added into the suspension. The cells were incubated for 1 h, washed, and resuspended in PBS buffer. Two micrograms per milliliter of JC-1 (2 mg/ml, dissolved in DMSO, Sigma) was added into the suspension. The cells were incubated for 1 h.

**TABLE 1 | Strains and plasmids used in this study.**

| Strains | Genotype and description | References |
|---------|--------------------------|------------|
| C. albicans strains | | |
| BWP17 (WT) | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG | Dana Davis |
| cnb1ΔΔ | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG cnb1::ARG4/cnb1::URA3-dp200 | This study |
| yvc1ΔΔ | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG yvc1::ARG4/yvc1::URA3-dp200 cnb1::URA3-dp200/cnb1::URA3-dp200 | Qilin Yu |
| cnb1ΔΔyvc1ΔΔ | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG yvc1::ARG4/yvc1::URA3-dp200 cnb1::URA3-dp200/cnb1::URA3-dp200 | This study |
| WT- pPCCH1-GFP | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG pPCCH1-GFP | This study |
| cnb1ΔΔ- pPCCH1-GFP | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG cnb1::ARG4/cnb1::URA3-dp200 pPCCH1-GFP | This study |
| yvc1ΔΔ- pPCCH1-GFP | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG yvc1::ARG4/yvc1::URA3-dp200 pPCCH1-GFP | This study |
| cnb1ΔΔyvc1ΔΔ- pPCCH1-GFP | ura3Δ::imm434/ura3Δ::imm434 his1::hisG/his1::hisG arg4::hisG/arg4::hisG yvc1::ARG4/yvc1::URA3-dp200 cnb1::URA3-dp200/cnb1::URA3-dp200 pPCCH1-GFP | This study |
| Plasmid | pPCCH1-GFP | Containing URA3 marker, Amp<sup>r</sup> |

**TABLE 2 | Primers used in this study.**

| Primers | Sequence (5′ → 3′) |
|---------|-------------------|
| ACT1-5RT | TGGAGTGGTGGTCCAGAAAGAC |
| ACT1-3RT | GTCAGCATTGAATGATGCG |
| CCH1-5RT | TTTTCAAGGACACACATAG |
| CCH1-3RT | GGAGTACATCGTTGAGT |
| PRB1-5RT | CATGGGCTTATCTTTACA |
| PRB1-3RT | GTGGTTGGGGTCTTTGAAC |
| PMT4-5RT | TCACTATGAGGCCATCACAT |
| PMT4-3RT | TGCAGATGAAAACGACACAGGAGAGA |
| HAC1-5RT | TC AGGATGAAACACACAAAGAAA |
| HAC1-3RT | CAAAGTCCAACCTGAAATGGAT |

**Vacular Membrane Permeability Assay**

5-(6)-Carboxy-2′,7′-dichlorofluorescein diacetate (C-DCFDA) or 7-amino-4 chloromethyl coumarin (CMAC) was used to measure the vacular membrane permeability (VMP) (Andrei-Selmer et al., 2001). In the normal cells, C-DCFDA with green fluorescence or CMAC with blue fluorescence was detected in the cells with decreased MMP. The green fluorescence (excitation wavelength at 490 nm, emission wavelength at 530 nm) with J-monomer could be detected in the cells with normal MMP. The red fluorescence (excitation wavelength at 525 nm, emission wavelength at 590 nm) with J-aggregates could be detected in the cells with decreased MMP (Yu et al., 2021).

**HAC1 Splicing Assay**

The strains treated with TN were cultured to log-phase. Total RNA was extracted with Eastep<sup>TM</sup> Total RNA Extraction Kit (Promega, Madison, WI, United States) and transcribed reversely to cDNA. The HAC1 splicing assay was detected by the PCR method with primers HAC1-5RT and HAC1-3RT. The PCR product was separated in agarose gel electrophoresis for 3 h (Li J. et al., 2018).

**MTT Assay**

The cellular viability was detected by MTT reagent [3-(4,5)-dimethylthiazol-(z-y1)-3,5-di-phenyltetrazoliumromide, 4 mg/ml, Beyotime, China] (Rong et al., 2020). The log-phase cells were collected and resuspended in PBS buffer. The 0.5-μg/ml MTT reagent was added and incubated with 70 r/min at 37°C for 1 h. The cells were centrifuged, and the supernatant was removed. Dimethyl sulfoxide (DMSO, Beyotime, China) was added into the precipitated cells, and the cells were centrifuged. The dissolved supernatant was collected to measure the absorption wavelength at 570 nm.

**Propidium Iodide Assay**

Propidium iodide (PI) was used to measure the cell death rate for the PI-positive cells which were stained with red fluorescence completely (Priante et al., 2018). The strains treated with TN were cultured to log-phase. Five micrograms per milliliter of PI (1 mg/ml, Sigma) was added into cells and incubated foranden the mitochondrial membrane potential (MMP) was recorded with the flow cytometer (FACS Calibur, BD, San Jose, CA, United States). The red fluorescence (excitation wavelength at 525 nm, emission wavelength at 590 nm) with J-monomer was recorded in the cells with decreased MMP. The green fluorescence (excitation wavelength at 490 nm, emission wavelength at 530 nm) with J-monomer was detected in the cells with normal MMP.
5 min. Mortality rate was represented by the PI-positive cells with a flow cytometer.

Real-Time PCR Assay
Real-time PCR assay was used to measure the expression level of the calcium channel-related gene CCH1 and UPR response-related genes PMT4 and PRB1. Cells were collected, and the total RNA was extracted with Eastep™ Total RNA Extraction Kit (Promega, United States) and transcribed reversely to cDNA (Meng et al., 2018). The RealMasterMix (SYBR Green) kit (TransGen, China) was used for real-time PCR analysis (Gomes-Neto et al., 2017). The following primers used are listed in Table 2: ACT1-5RT, ACT1-3RT, CCH1-5RT, CCH1-3RT, PRB1-5RT, PRB1-3RT, PMT4-5RT, and PMT4-3RT. The 2\(^{-\Delta\Delta CT}\) method was used to calculate the expression level of different genes, and ACT1 was used as the internal control.

Statistical Analysis
Each experiment mentioned above was repeated at least three times under the tested conditions. The standard deviations and means were calculated by the separate three replicates. The one-tailed Student’s t test was used to calculate p values. The p values less than 0.05 were considered as statistically significant difference.

RESULTS

The Deletion of CNB1 Caused Sensitivity to Endoplasmic Reticulum Stress and Led to the CCH1 Overexpression
Firstly, we constructed the cnb1\(\Delta/\Delta\) mutant and measured its sensitivity to ER stress reagent TN. The results showed that on the YPD plate, cnb1\(\Delta/\Delta\) grew normally as a wild-type (WT) strain, while cnb1\(\Delta/\Delta\) could hardly grow on the 2-\(\mu\)g/ml TN plate (Figure 1A). Besides, the growth of liquid medium indicated that the WT strain grew rapidly during the 24-h culture period, whose value increased from 0.8 to 24. However, the OD\(_{600}\) of cnb1\(\Delta/\Delta\) was always maintained at 0.8 and the maximum value at 24 h was just 3 (Figure 1B). Since calcineurin regulated the cellular calcium concentration through inhibition of plasma membrane channel CCH1 (Xu et al., 2019), we assumed that the deletion of CNB1 might have an impact on the CCH1 expression. It was interesting that under TN treatment, the expression level of the CCH1 promoter in the cnb1\(\Delta/\Delta\) strain was near as 1.5 times as WT (Figure 1C). Moreover, the qPCR analysis indicated that CCH1 was upregulated in the transcription level of cnb1\(\Delta/\Delta\) in response to ER stress (Figure 1D). In general, the deletion of CNB1 leads to the sensitivity to ER stress and overexpression of CCH1.

The Inhibition of the Plasma Membrane Calcium Channel Could Recover the Growth Defect of cnb1\(\Delta/\Delta\) Under Endoplasmic Reticulum Stress
Now that the sensitivity to ER stress of the cnb1\(\Delta/\Delta\) strain was related with the overexpression of CCH1, we speculated that the inhibition of the plasma membrane calcium channel might improve its growth status. Verapamil or nifedipine, as the universal calcium channel blocker, was usually applied to cure the hypertension or angina. A different concentration of verapamil or nifedipine was added into the YPD plates containing 2-\(\mu\)g/ml TN. The spot assay result showed that the colony of cnb1\(\Delta/\Delta\) could grow in both the plates treated with verapamil and nifedipine (Figure 2A). Meanwhile, the liquid
the cytosolic calcium concentration could recover the inhibition of the plasma membrane calcium channel to decrease growth under ER stress. We found that similar with the WT strain, \(cnb1\Delta/\Delta\) and \(yvc1\Delta/\Delta\) mutants grew slowly, \(cnb1\Delta/\Delta\) grew marginally faster than the \(cnb1\Delta/\Delta\) strain did (Figure 3B). Besides, the cell death rates under ER stress of these were calculated by flow cytometry with PI dye. All of the strains grew well in the control group, and their death rates were low (Figures 3C,D, control). Under the TN treatment, WT and \(yvc1\Delta/\Delta\) strains were still with the small dead cells, and the death rate was 26.8% in the \(cnb1\Delta/\Delta\) strains, which was increased to 39.72% in the \(cnb1\Delta/\Delta\) strain (Figures 3C,D). These results indicated that even if \(cnb1\Delta/\Delta\) and \(yvc1\Delta/\Delta\) showed sensitivity to TN, the cell death rate was lower than that of the \(cnb1\Delta/\Delta\) strain, indicating that the deletion of \(YVC1\) recovered the cell vitality of \(cnb1\Delta/\Delta\).

The Deletion of \(YVC1\) Recovered Vacuolar Membrane Permeability and Mitochondrial Membrane Potential of \(cnb1\Delta/\Delta\) Under Endoplasmic Reticulum Stress

Normally, the vacuole cavity could be dyed by CMAC, while the damaged cells were stained in the whole cell or failed to be stained (Andrei-Selmer et al., 2001). The vacuolar membrane permeability of the mutants showed that deletion of \(CNB1\) caused the damaged vacuolar membrane under TN treatment, with the whole cells stained by CMAC or failing to be stained (Figure 4A). Besides, the calculation of the VMP-positive rate showed that cultured in the TN-treated medium, the WT and \(yvc1\Delta/\Delta\) strains maintained the integrity of vacuoles, with a low percentage of the VMP-positive rate. However, the positive rate was up to 70% in the vacuolar severely impaired \(cnb1\Delta/\Delta\) strain, although for the \(cnb1\Delta/\Delta\) strain with impaired vacuolar membrane, the VMP-positive rate was lower than \(cnb1\Delta/\Delta\), just about 45% (Figure 4B).

The mitochondrial activities under ER stress were measured as well. In the control group, the \(cnb1\Delta/\Delta\) and \(yvc1\Delta/\Delta\) strains, as the WT strain, maintained the normal mitochondrial function, with the low percentage of damaged rate (Figures 4C,D, control). However, compared with the other strains, the mitochondrial function was interfered by TN in the \(cnb1\Delta/\Delta\) strain, and the rate of decreased MMP was up to 41.72%. Nevertheless, the rate of impaired mitochondria growth measurement indicated that during the 24-h culture, 40 or 80 \(\mu\)g/ml verapamil could improve the growth of \(cnb1\Delta/\Delta\) under ER stress (Figure 2B). These results implied that the inhibition of the plasma membrane calcium channel to decrease the cytosolic calcium concentration could recover the \(cnb1\Delta/\Delta\) growth under ER stress.

Deletion of \(YVC1\) Decreased the Cell Death Rate of \(cnb1\Delta/\Delta\) Under Endoplasmic Reticulum Stress

Since the inhibition of the cytosolic calcium concentration recovered the growth of \(cnb1\Delta/\Delta\) and \(YVC1\) was the unique vacuolar membrane calcium channel in yeast, we doubted whether the deletion of \(YVC1\) could increase the growth status of \(cnb1\Delta/\Delta\) under ER stress. We constructed \(yvc1\Delta/\Delta\) and \(cnb1\Delta/\Delta\) strains and measured their susceptibility to environmental stress. CFW and caspofungin could cause the cell wall stress. Besides, both \(\beta\)-mercaptoethanol and DTT could interfere the formation of disulfide bonds, and TN could inhibit glycosylation. These three reagents could lead to ER stress. We found that similar with the WT strain, \(cnb1\Delta/\Delta\), \(yvc1\Delta/\Delta\), and \(cnb1\Delta/\Delta\) strains were resistant to CFW, \(\beta\)- mercaptoethanol, and DTT (Figure 3A, panels 2–4). However, the \(cnb1\Delta/\Delta\) and \(cnb1\Delta/\Delta\) strains were sensitive to caspofungin and TN. Since \(YVC1\) regulated the OSR in yeast (Yu et al., 2014b), the antioxidant vitamin C (VC) was added into the plates to verify whether the sensitivity was related with abnormal OSR. It was interesting that the addition of VC recovered the strains on the caspofungin-treated plate, indicating that the sensitivity to cell wall stress was related with impaired OSR reaction (Figure 3A, panels 5–6). Nevertheless, the addition of VC could not change the strains’ susceptibility to TN (Figure 3A, panels 7–8), which revealed the novel interaction between \(YVC1\) and \(Cnb1\) with ER stress independently of OSR.
FIGURE 3 | Deletion of CNB1 and YVC1 recovered the growth defect of the cnb1Δ/Δ strain. (A) ER stress or cell wall stress-related reagents were added into YPD plates. WT, cnb1Δ/Δ, yvc1Δ/Δ, or cnb1Δ/Δyvc1Δ/Δ strains were cultured and spotted on the plates. (B) The growth curve of each strain was drawn. The initial OD600 value was rectified at 0.2 and strains were cultured in liquid YPD medium, and 2 μg/ml TN was added into these medium and cultured for 25 h. The OD600 value in each strain was measured at specific times. (C) The death rate of each strain under ER stress. Log-phase strains were collected and washed twice with PBS buffer. Five micrograms per milliliter of PI was added into the strains and incubated for 5 min. The cell death rate was measured by flow cytometry. The cells without TN treatment were as the control group. (D) The statistical analysis of PI death rate. * means significant difference between the cnb1Δ/Δ and cnb1Δ/Δyvc1Δ/Δ strains under TN-treated conditions (p < 0.05). The experiments were repeated three times separately.

The Unfolded Protein Response Pathway Was Activated Effectively in the cnb1Δ/Δ and cnb1Δ/Δyvc1Δ/Δ Strains Under Endoplasmic Reticulum Stress

The UPR pathway is the classical response process in the ER stress of yeast strains (Zhang et al., 2019). Since the deletion of YVC1 recovered the cell vitality of cnb1Δ/Δ under the treatment of TN, we speculated whether the UPR pathway was overactivated in this mutant. To verify the possibility, the total RNA of these three mutants and WT strains was extracted and transcribed reversely to cDNA (Gomes-Neto et al., 2017). The Hac1 splicing level and the expression level of PRB1 and PMT4 were measured. However, much unexpectedly, similar with the WT or yvc1Δ/Δ mutants, the UPR pathway was activated in cnb1Δ/Δ and cnb1Δ/Δyvc1Δ/Δ effectively. The unspliced HAC1 was 581 bp in the WT and mutant strains of the control group (Figure 5A). Moreover, under TN treatment for 2 h, HAC1 was spliced...
The Deletion of YVC1 Decreased the Calcium Flux and Cellular Calcium Concentration of cnb1Δ/Δ Under Endoplasmic Reticulum Stress

Yvc1 and Cnb1 were related with the cellular calcium regulation (Cyert and Philpott, 2013), and the inhibition of CCH1 to decrease the cytosolic calcium content could improve the growth status in the cnb1Δ/Δ strain. Therefore, we speculated whether the disruption of YVC1 reduces the vacuolar calcium release to improve the cell vitality. Moreover, the results showed that calcium fluctuation was enhanced in the cnb1Δ/Δ strain under ER stress. In the detected period, the calcium fluctuation was increased within 3 min, and the maximum concentration was higher than other strains. Nevertheless, the calcium flux of the cnb1Δ/Δ yvc1Δ/Δ strain was steady with a low peak value like the WT strain (Figure 6A).

Next, we measured the specific concentration of cellular calcium, which showed a similar tendency with the calcium flux. We found that the calcium concentration was increased significantly with TN treatment for 2 h of cnb1Δ/Δ, in which the concentration was much higher than those of other strains. The calcium content of the cnb1Δ/Δ yvc1Δ/Δ strain was close to that of yvc1Δ/Δ, lower than that of cnb1Δ/Δ (Figure 6B). In conclusion, the calcium concentration was decreased significantly and its normally in the cnb1Δ/Δ and cnb1Δ/Δ yvc1Δ/Δ strains and other strains with the size at 562 bp (Figure 5A).

Besides, qPCR analysis indicated that the expression levels of PMT4 and PRB1 in the cnb1Δ/Δ and cnb1Δ/Δ yvc1Δ/Δ strains were similar with WT, which were all upregulated in the transcriptional level under ER stress (Figures 5B,C). In summary, these figures showed that the URP pathway in all of the mutants did not fail to evoke, implying a novel regulation mechanism within susceptibility.
fluctuation was reduced obviously under ER stress in the cnb1ΔΔyvc1ΔΔ strain, indicating that the disruption of YVC1 could reduce the cellular calcium concentration, thereby improving the vitality.

**DISCUSSION**

In this study, we found that the deletion of YVC1 recovered the growth defect of cnb1ΔΔ in the regulation of calcium ions in response to ER stress (Figure 8). cnb1ΔΔ was hypersensitive to TN, which was relevant with the overexpression of the cytoplasm membrane channel CCH1 (Figures 1, 2). This revealed the interaction between the sensitivity to ER stress and the regulation of cytosolic calcium content of the cnb1ΔΔ strain. Although cnb1ΔΔyvc1ΔΔ was sensitive to TN, the growth was recovered and the death rate was obviously decreased (Figure 3). In the spot assay, the strains were sensitive to TN instead of DTT, the former inhibited the N-glycosylation, and the latter influenced the disulfide bond. The results indicated that the disruption of EGTA could recover the growth of cnb1ΔΔ. The VMP assay showed that the permeability was improved in the addition of EGTA treatment, the positive rate was down to 48% (Figure 7D). In conclusion, we determine that it is the overloaded calcium ions that cause the susceptibility to ER stress in the cnb1ΔΔ strain, and the disruption of YVC1 reduces the cytosolic calcium and improves the cell vitality (Figure 8).
strain might have no impact on the disulfide bond, which could be verified subsequently. Moreover, the vacuolar membrane permeability and mitochondrial activity were improved in the double mutant (Figure 4).

The UPR pathway was a classical regulation method in response to ER stress; much unexpectedly, the UPR pathway was evoked in all of the tested strains, indicating their normal activation (Figure 5). However, we measured the calcium flux and the cellular calcium content. The calcium fluctuation was enhanced with the highest ion concentration in the cnb1∆/∆ strain. Also, the calcium flux was steady with a lower content in the cnb1∆/∆yvc1Δ/Δ strain, indicating the effect of the deletion of YVC1 on the decrease in cellular calcium (Figure 6). The further results show that addition of EGTA recovered the cell growth, cell vitality, and vacuolar membrane permeability, which corresponded with our conjectures (Figure 7).

Endoplasmic reticulum stress leads to the activation of the Ire1/Xbp1 signal pathway in mammalian cells. Regulated by Hac1, Pmt4 is expressed in the nucleus to alleviate the misfolded or unfolded proteins. In S. cerevisiae, ER stress upregulated the expression of Ptp2 tyrosine phosphatase and Cmp2 calcineurin phosphatase; the former was mediated by Mpk1 MAP kinase, and the latter could downregulate the activity of Hog1 MAP kinase (Mizuno et al., 2018). The Cmp2 homolog with Cna1 in C. albicans is the catalytic subunit which maintains the principal role for the stress response. Besides, calcineurin is essential for cells in many biological processes and for the long-term survival of cells undergoing ER stress (Liu et al., 2015). Connected with our findings, we doubt whether the Hog1-MAPK pathway is related with the ER stress response in the mutants.

It was reported that for humans, the nephrotic syndrome was associated with the activation of the ER calcium release
FIGURE 8 | Schematic model of Yvc1 regulation cellular calcium concentration to improve growth in the cnb1Δ/Δyvc1Δ/Δ strain under ER stress. The calcineurin mutant strain cnb1Δ/Δyvc1Δ/Δ was sensitive to ER stress while the double mutant cnb1Δ/Δyvc1Δ/Δ recovered the growth. The URP pathway was effectively activated in these mutants. However, the deletion of TRP channel Yvc1 could inhibit the release of Ca2+ from vacuole and reduce the cellular calcium concentration in response to ER stress. Thereby, the vacuolar membrane permeability (VMP) was inhibited and the cell vitality was maintained, leading to the improvement of cell growth.

channel, which led to podocyte injury (Park et al., 2019). Moreover, the connection of ER and mitochondria was complex. Through shuttling of calcium ions, the connection was involved not only in ion homeostasis but also in many structural and apoptotic proteins (Kumar and Maity, 2021). A stimulator of IFN genes, called STING, regulated not only calcium homeostasis but also ER stress and T cell survival, which were associated with lung disease (Wu et al., 2019). In yeast, cadmium exposure led to the interrupted calcium homeostasis, induced the lipid dysregulation, and finally caused ER stress (Rajakumar et al., 2016).

Verapamil and nifedipine are the cytoplasm membrane channel blockers in mammalian cells, used as the clinical drugs in curing heart diseases like angina pectoris and supraventricular arrhythmias (Xing et al., 2020). In our previous work, verapamil inhibited the Hwp1 expression, indicating its regulatory role in both morphogenesis-associated proteins and the secretory pathway (Yu et al., 2014a). Moreover, the combined use of verapamil and antifungal drug fluconazole had a synergetic inhibitory effect on hyphal development (unpublished data). Thereby, the cumulative effect of verapamil and TN could be evaluated in-depth. Besides, the effect of CNB1 or YVC1 deletion was interesting since the sensitivity to TN was irrelevant with the UPR pathway; indeed, it was the ROS-independent and UPR-independent calcium overloading. Connected with our previous cognition, it was subversive and enlightening.

Moreover, the relationship between the calcium signaling pathway members and the environmental stress response needs further investigation, for instance, the relationship between Cch1 and Yvc1 for their regulation role in calcium ion content. Since Cnb1 regulates the ER stress response, does the transcription factor Crz1 or the other subunit of calcineurin Cna1 regulate CCH1 in C. albicans. Besides, the specific regulation site of CCH1, the mechanism by which deletion of Cnb1 and Yvc1 caused cellular calcium ions to be increased under TN treatment, and the sensitivity to other environmental stimulus of the mutant are still unknown. Furthermore, we will work in-depth to figure out more mechanisms about the calcium signal pathway regulation under environmental stress.

In summary, our study revealed a novel interaction between the Yvc1-regulated cellular calcium ions and ER stress response, which was independent on the antioxidative reaction or UPR pathway (Figure 8). This work will extend our knowledge of the cellular sensor role of the TRP channel and calcineurin under environmental stress and uncover the new targets against fungal infections.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

LP and ML conceived and designed the experiments and wrote the manuscript. LP, JD, and RZ performed the experiments. HZ, NZ, and QZ analyzed the data. ML and QY did supervision. All authors have read and agreed to the published version of the manuscript.

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