Putative Stress Sensors WscA and WscB Are Involved in Hypo-Osmotic and Acidic pH Stress Tolerance in *Aspergillus nidulans*†‡

Taiki Futagami,1 Seiki Nakao,1 Yayoi Kido,1 Takuji Oka,2 Yasuhiro Kajiwara,3 Hideharu Takashita,3 Toshiro Omori,3 Kensuke Furukawa,3 and Masatoshi Goto1*

Department of Bioscience and Biotechnology, Faculty of Agriculture, Kyushu University, 6-10-1, Hakozaki, Fukuoka 812-8581, Japan1; Department of Applied Microbial Technology, Faculty of Biotechnology and Life Science, Sojo University, 4-22-1 Ikeda, Kamamoto 860-0082, Japan2; Research Laboratory, Sanwa Shurui Co., Ltd., 2231-1, Usa, Oita 879-0495, Japan3; and Department of Food and Bioscience, Faculty of Food Science and Nutrition, Beppu University, 82 Kitaiishigaki, Beppu 874-8501, Japan4

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Wsc proteins have been identified in fungi and are believed to be stress sensors in the cell wall integrity (CWI) signaling pathway. In this study, we characterized the sensor orthologs WscA and WscB in *Aspergillus nidulans*. Using hemagglutinin-tagged WscA and WscB, we showed both Wsc proteins to be N- and O-glycosylated and localized in the cell wall and membrane, implying that they are potential cell surface sensors. The *wscA* disruptant (*ΔwscA*) strain was characterized by reduced colony and conidia formation and a high frequency of swollen hyphae under hypo-osmotic conditions. The deficient phenotype of the *ΔwscA* strain was facilitated by acidification, but not by alkalization or antifungal agents. In contrast, osmotic stabilization restored the normal phenotype in the *wscA* disruptant strain, but to a lesser extent. In addition, a double *wscA* and *wscB* disruptant (*ΔwscA ΔwscB*) strain was viable, but its growth was inhibited to a greater degree, indicating that the functions of the products of these genes are redundant. Transcription of α-1,3-glucan synthase genes (*agsA* and *agsB*) was significantly altered in the *wscA* disruptant strain, resulting in an increase in the amount of alkali-soluble cell wall glucan compared to that in the wild-type (wt) strain. An increase in mitogen-activated protein kinase (MpKA) phosphorylation was observed as a result of *wsc* disruption. Moreover, the transient transcriptional upregulation of the *agsB* gene via MpKA signaling was observed in the *ΔwscA ΔwscB* strain to the same degree as in the wt strain. These results indicate that *A. nidulans* Wsc proteins have a different sensing spectrum and downstream signaling pathway than those in the yeast *Saccharomyces cerevisiae* and that they play an important role in CWI under hypo-osmotic and acidic pH conditions.

Signal transduction plays an important role in sensing environmental stimuli and the subsequent regulation of gene expression required for appropriate cell development and morphology. In the yeast *Saccharomyces cerevisiae*, the cell wall integrity (CWI) signaling pathway has been identified as a major regulator of cell wall biogenesis in adaptation to environmental stresses such as heat shock and hypo-osmolarity (23, 24). Recent genomics studies revealed that the core component of this pathway is highly conserved in yeast and other fungal genomes (31, 38).

The CWI pathway has been well studied in *S. cerevisiae* (23, 24). The five plasma membrane sensor proteins, Wsc1, Wsc2, Wsc3, Mid2, and Mid1, act as mechanosensors and detect cell wall perturbations caused by stressful events (35, 48). Wsc1 and Mid2 act as the main sensor proteins in this system. The activated sensors initiate the signaling cascades of downstream cytoplasmic transducers (23, 24). The sensor proteins interact with GDP/GTP exchange factors (Rom1 and Rom2) that activate a small G protein (Rho1). Rho1 GTPase activity in turn activates a mitogen-activated protein kinase (MAPK) module through sequential phosphorylation. The MAPK module consists of MAPKKK (Bck1), MAPKKs (Mkk1 and Mkk2), and MAPK (Slt2 [Mpk1]). Eventually, the phosphorylation relay activates the transcriptional factors Rlm1 and SBF (composed of Swi4 and Swi6), which regulate the transcriptional levels of cell wall-related genes.

A BLAST search using the sequences of these functionally characterized CWI components revealed that their orthologous genes, except for the stress sensor genes *MID2* and *MTL1*, are conserved in the filamentous fungi *Aspergillus fumigatus*, *A. nidulans*, and *A. niger* (8). Several downstream signal-transducing orthologous genes, including *rhoA*, *pkcA*, *mpkA*, and *rlmA*, that are orthologs of the *RHO1*, *PKC1*, *MPK1*, and *RLM1* genes, respectively, have been demonstrated to play essential roles in cell polarity, differentiation, and CWI in *Aspergillus* species. In *A. nidulans*, the Rho family GTPase RhoA was shown to be involved in establishing polarity, branching, and cell wall synthesis (13). It was suggested that the protein kinase C (PKC)-encoding gene *pkcA* plays an essential role in the maintenance of cell integrity and polarized growth in *A. nidulans* (18, 41, 46). The *A. nidulans* MpKA protein is involved in germination of conidial spores and polarized growth (4), and transcription of the *mpkA* gene seems to be autoregulated by the CWI pathway via MpKA (8). The transcription of most cell wall–related genes except for the
α-1,3-glucan synthase genes \textit{agsA} and \textit{agsB} is independent of RimA, unlike the case in the yeast model (8).

Although the downstream signal transducers and transcription factors involved in CWI signaling have been characterized in \textit{Aspergillus} species, the functional roles of the upstream cell wall sensor WSC1-3 orthologs have not been elucidated in filamentous fungi. The Wsc family of sensor proteins has been characterized in \textit{S. cerevisiae} and its close relative \textit{Kluyveromyces lactis} (24, 40). A structural feature of the Wsc proteins is the presence of a cysteine-rich domain (also referred to as a WSC domain), a serine/threonine-rich region, a transmembrane region, and a highly charged C-terminal cytoplasmic region. The WSC domain contains up to eight conserved cysteine residues that may form S-S bonds and is believed to mediate noncovalent binding with cell wall glucans. The 1,3-exoglucanase of \textit{Trichoderma harzianum} also contains two copies of the WSC domain that may bind glucan chains (5). A GFP fusion localization study revealed that Wsc1 resides in membrane patches within the plasma membrane in both \textit{S. cerevisiae} and \textit{K. lactis} (39, 44). Recently, single-molecule atomic force microscopy revealed that Wsc1 behaves like a linear nanospring that is capable of resisting a high level of mechanical force and of responding to cell surface stress (6). The WSC domain of Wsc1 is required for clustering stimulated by stress-activated force and of responding to cell surface stress (6). The WSC domain of Wsc1 is required for clustering stimulated by stress-activated force and of responding to cell surface stress (6).


during the course of our previous study on posttranslational modification by O-glycosylation in \textit{A. nidulans}, we identified the gene for the Wsc family sensor ortholog \textit{wscA} (11). The \textit{wscA} gene was used to investigate the substrate specificities of the protein O-mannosyltransferases PmtA, PmtB, and Pmc in \textit{A. nidulans}, since the conserved serine/threonine-rich region was shown to be highly O-mannosylated by the O-mannosyltransferases Pmt2 and Pmt4 in \textit{S. cerevisiae} (26). O-mannosylation is believed to confer the rod-like structure upon the sensors by extending and stiffening the polypeptide. Our results also suggested that \textit{A. nidulans} \textit{wscA} is O-mannosylated in \textit{A. nidulans} by PmtA and PmtC, but not by PmtB, and that the O-glycan attachment has a significant impact on the stability and degradation of WscA (11, 20). Protein O-glycosylation plays an essential role in fungal species, as demonstrated by the observation that the \textit{pmtA} disruptant exhibits abnormal cell morphology and altered cell wall composition (10, 32). Notably, disruption of \textit{pmtC} leads to a higher level of growth repression than disruption of the other \textit{pmt} genes. The hypha in the \textit{pmtC} disruptant is swollen and frequently branched, and cells lose the ability to form conidia under normal growth conditions. Accordingly, impairment of Pmt substrates, including WscA, may result in a crucial defect in the hyphal structure, germination, and CWI in \textit{A. nidulans}.

In this study, we sought further understanding of how the substrates of Pmt function in \textit{Aspergillus} species. We also identified and characterized the stress sensor ortholog \textit{wscB} and examined its role in influencing cell morphology and cell wall biogenesis in \textit{A. nidulans}. Using phenotypic and transcriptional analysis, we showed that WscA and WscB are involved in CWI in \textit{A. nidulans} subjected to hypo-osmotic and acidic pH conditions.

### TABLE 1. \textit{A. nidulans} WscA AND WscB

| Strain | Genotype | Source or reference |
|--------|----------|---------------------|
| AKU89  | bi1 \textit{argB}2 \textit{akb1} \textit{bza1}A | 11 This study |
| Δ\textit{wscA} | \textit{bi1 argB}2 \textit{akb1} \textit{bza1}A | This study |
| \textit{wscA} Δ\textit{wscB} | \textit{bi1 argB}2 \textit{akb1} \textit{bza1}A | This study |
| Δ\textit{wscA} | \textit{wscA} \textit{ptrA} pyG/wscC | This study |
| Δ\textit{wscB} | \textit{wscB} \textit{ptrA} pyG/wscC | This study |
| AKU89-\textit{wscA}-HA | \textit{alcF} (\textit{wscA}+3:\textit{H3A}-\textit{Bip1} \textit{ptrA}A) | This study |
| AKU89-\textit{wscB}-HA | \textit{alcF} (\textit{wscB}+3:\textit{H3A}-\textit{Bip1} \textit{ptrA}A) | This study |
| Δ\textit{mpkA} | \textit{mpkA}::\textit{argB} | This study |

### MATERIALS AND METHODS

\textit{A. nidulans} strains, media, and culture conditions. The \textit{Aspergillus} strains used in this study are listed in Table 1. Fungi were grown at 30°C in YM medium (0.5% [wt/vol] yeast extract, 1% [wt/vol] glucose) or MM medium (1% [wt/vol] glucose, 0.6% [wt/vol] NaNO3, 0.052% [wt/vol] KCl, 0.052% [wt/vol] MgSO4·7H2O, 0.152% [wt/vol] KH2PO4, 0.211% [wt/vol] arginine, 5 μg/ml of biotin, and Hunter's trace elements [pH 6.5]). Media were adjusted to the required pH with HCl and NaOH. To express genes under the control of the \textit{alcA} promoter, 100 mM threonine and 0.1% fructose were added as carbon sources instead of glucose in MM medium. For the cultivation of pGEM-negative strains, 0.056% (wt/vol) uracil and 0.122% (wt/vol) uridine were added to MM medium.

Construction of \textit{wscA} and \textit{wscB} disruptants. \textit{A. nidulans} AKU89 served as the wild-type (wt) strain in this study (11). The \textit{wscA} and \textit{wscB} genes were disrupted in \textit{wt A. nidulans} by insertion of the pyrithiamine resistance gene (\textit{ptrA}). The gene replacement cassette encompassing 1 kb at the 5' end of \textit{wscA}, 2 kb of \textit{ptrA}, and 1 kb at the 3' end of \textit{wscA} was constructed by recombinant PCR using the primer pairs F5660-1015/R5660-2042, F5660-ptrA/R5660-ptrA, and F5660-2939/R5660-3992, respectively (see Fig. S1A and Table S2 in the supplemental material). For amplification of the \textit{ptrA} gene, pGEM-T Easy \textit{ptrA} was used as the template DNA. The resultant DNA fragment amplified with primers F5660-1015 and R5660-3992 was used to transform \textit{wt A. nidulans}. The gene replacement cassette encompassing 1.5 kb at the 5' end of \textit{wscB}, 2 kb of \textit{ptrA}, and 1.5 kb at the 3' end of \textit{wscB} was constructed by recombinant PCR using the primer pairs F4674-599/R4674-1990, F4674-ptrA/R4674-ptrA, and F4674-3065/R4674-4505, respectively (see Fig. S1B and Table S2 in the supplemental material), and the resultant DNA fragment amplified with primers F4674-599 and R4674-4505 was used to transform \textit{wt A. nidulans}. Standard transformation procedures for \textit{A. nidulans} were used (53). For the selection of transformants, MM agar plates supplemented with 0.1 μg/ml of pyrithiamine were used. Disruption of \textit{wscA} and \textit{wscB} was confirmed by Southern blot analysis (see Fig. S1A and B in the supplemental material).

\textit{A. nidulans} genomic DNA was prepared as previously described (32). The digoxigenin (DIG)-labeled probes were constructed using primer sets F5660-2939/R5660-3992 for \textit{wscA} and F4674-3065/R4674-4505 for \textit{wscB} and a DIG labeling kit (Roche) according to the manufacturer’s protocols.

To construct a \textit{wscA} and \textit{wscB} double disruptant strain, the \textit{wscB} gene was disrupted in a \textit{wscA} disruptant strain using \textit{argB} insertion. The gene replacement cassette encompassing 1.4 kb of the 5' end of \textit{wscB}, 1.8 kb of \textit{argB}, and 1.5 kb of the 3' end of \textit{wscB} was constructed by recombinant PCR using the primer pairs F4674-599/RargB4674-1990, F4674-argB/R4674-argB, and FargB4674-3065/R4674-4505, respectively (see Table S2 in the supplemental material), and the resultant DNA fragment amplified with primers F4674-599 and R4674-4505 was used to transform a \textit{wscA} disruptant strain. The transformants were selected on MM agar plates without arginine.

Introduction of the \textit{argB} gene into the \textit{wscB} locus was confirmed by PCR using the primer pair F-AnwscB/R4674-4505 (see Fig. S1C in the supplemental material).

Complementation of \textit{wscA} and \textit{wscB} deletion strains. For analysis of complementation of the \textit{wscA} disruptant with \textit{wt wscA}, a gene replacement cassette encompassing 1.5 kb of the 5' end of \textit{pyrG}, 2.9 kb of \textit{wt wscA}, and 1.2 kb of the...
3' end of ppyG was constructed by recombinant PCR using the primer pairs F1-AnypyG/R1-AnypyG, F-Awnsac/R-Awnwsac, and F2-AnypyG/R2-AnypyG, respectively (see Table S2 in the supplemental material). The resultant DNA fragment amplified with primers F1-AnypyG and R2-AnypyG was used to transform the wscA disruptant. Transformants were selected on MM agar with 10 mM arginine, 5 mM 5-fluoroorotic acid, 5 mM uridine, and 5 mM uracil. Introduction of the wt wscA gene into the wscA disruptant at the ppyG locus was confirmed by PCR using the primer pair F3-AnypyG/R2-AnypyG (see Fig. S1D in the supplemental material).

For analysis of complementation of the A. nidulans wscB disruptant with wt wscB, the plasmid pGTasal-ppyG-wscB was constructed as follows. A 3.1-kb DNA fragment of wscB was amplified by PCR using F-Awnwsac and R-Awnwsac and ligated into the SalI site of pGTasal-ppyG (11) (see Table S2 in the supplemental material), yielding pGTasal-ppyG-wscB. Using pGTasal-ppyG-wscB as a template, we amplified the DNA fragment carrying 1.3 kb of the 5' end of ppyG, 3.1 kb of wscB, and 1.7 kb of the 3' end of ppyG with primers F1-AnypyG and R2-AnypyG and used it to transform the wscA disruptant. Introduction of the wt wscB gene into the wscB disruptant at the ppyG locus was confirmed by PCR using the primer pair F3-AnypyG/R2-AnypyG (see Fig. S1D in the supplemental material).

**Analysis of conidiation efficiency.** Approximately 10^5 conidia were spread onto an 84-mm agar plate containing YG medium or YG medium with 0.6 M KCl. After 5 days of incubation at 30°C, a 1-cm^2 agar block containing newly formed conidia was suspended in 0.01% (wt/vol) Tween 20 solution and the conidia were counted using a hemocytometer. The mean number of conidia formed was determined from the results of 10 agar blocks from two independent plates.

**Analysis of glucon and chitin content.** Strains were grown for 24 h at 30°C with shaking at 120 rpm in flasks containing YG liquid medium. Analysis of the glucon and chitin contents was carried out according to previously described methods (3, 37). Briefly, cells were dissolved in 250 mM phosphate buffer (pH 7.0) and placed on ice for 30 min. The cells were then disrupted using a French press (Ohtake Seikasaku, Tokyo, Japan) and centrifuged at 27,300 × g for 10 min at 4°C. The pellet was washed with 10 ml of phosphate buffer. The centrifugation and washing steps were repeated five times, and the resulting pellet was dissolved in 10 ml of ultrapure water. The centrifugation and washing steps using ultrapure water were repeated five times. The resulting pellet was dispersed and used as a cell wall. Ten milligrams of cell wall pellet was dissolved in 1 ml of 1 M KOH and incubated for 30 min at 100°C. After centrifugation at 20,000 × g for 5 min, the resulting supernatant was centrifuged in 1 ml of 1 M KOH and incubated for 30 min at 70°C. After centrifugation at 20,000 × g for 5 min, the supernatant was used as the alkali-soluble fraction, while the pellet was dissolved in 1 ml of formic acid and incubated for 20 min at 100°C. After centrifugation at 20,000 × g for 5 min, the supernatant was used as the alkali-insoluble fraction. For chitin quantification, 10 mg of cell wall was digested with 2 mg/ml of chitinase (Watalase [Takara] in 250 mM phosphate buffer (pH 7.0) for 16 h at 30°C. After centrifugation at 20,000 × g for 5 min, the supernatant was washed with 10 ml of phosphate buffer. The centrifugation and wash steps were repeated three times, and the resulting supernatant was subjected to high-performance liquid chromatography (HPLC) using a CARBOSep COREGEL-87P column (Transgenomic) and a refractive index detector. The ultrapure water was used as a mobile phase at a flow rate of 1 ml per min. The column temperature was maintained at 85°C. The standards for polysaccharides, including those for trehalose, glucose, mannitol, and glycerol, were used.

Microscopy. To observe aerial hyphae, we inoculated conidia on a slide culture of YG agar medium. After incubation at 30°C for 48 h, the aerial hyphae were observed, using an inverted IX70 light microscope (Olympus). To observe submerged A. nidulans hyphae, we inoculated 2 × 10^5 conidia into 100 ml of YG liquid medium or liquid medium with 0.6 M KCl and incubated them with shaking at 120 rpm at 30°C. After 24 h, the mycelia were transferred to a 12-well plate and incubated with 10 mg/ml of fluorescent brightener 28 (calcofluor white; Sigma) for 10 min. The mycelia were mounted on the slide glass and observed using a FluoView FV10i confocal laser scanning microscope (Olympus). Image acquisition was performed using the Z-stack mode.

**Construction of A. nidulans strains expressing HA-tagged WscA or WscB.** For the expression of hemagglutinin (HA)-tagged WscA and WscB in A. nidulans, plasmids pAGTB-wscA-HA-ptrA and pAGTB-wscB-HA-ptrA, which contained a wscA::HA or wscB::HA fusion between the alteA promoter and bip4 terminator, were constructed as follows. HA was amplified using the F1-3HA-R1-3HA primer set and ligated into the Sall and PstI sites of pSAST0-Thp carrying Puc and Thp (see Table S2 in the supplemental material), yielding pAGTB-HA. The wscA and wscB genes were amplified using primer sets F1-S660.2-exp/R1-S660.2-exp and F1-4674.2-exp/R1-4674.2-exp, respectively (see Table S2 in the supplemental material), and ligated into the Noel and Sall sites of pAGTB-HA, yielding pAGTB-wscA-HA and pAGTB-wscB-HA. The ptrA gene was obtained by digesting pGEM-T Easy (Promega) with HindII1 and ligating pAGTB-wscA-HA and pAGTB-wscB-HA into the HindIII site, yielding pAGTB-wscA-HA-ptrA and pAGTB-wscB-HA-ptrA. The integrative plasmids were then transformed into wt A. nidulans. Southern blot analysis was used to confirm the pyrithiamine-resistant transformants carrying wscA::HA (AKU89-wscA-HA) and wscB::HA (AKU89-wscB-HA). The DIG-labeled probes were constructed using primer set F1-S660.2-exp/R1-S660.2-exp for AKU89-wscA-HA and F1-4674.2-exp/R1-4674.2-exp for AKU89-wscB-HA using a DIG-labeling kit (Roche) according to the manufacturer's protocols.

**Preparation of A. nidulans extracts and immunoblot analysis.** To investigate the localization of HA-tagged WscA and WscB, total, membrane, microsomal, and cytosolic fractions were prepared. The A. nidulans membrane fraction was prepared according to the method of Yamazaki et al., with some modifications (52). The conidia (2 × 10^7) of AKU89-wscA-HA and AKU89-wscB-HA were inoculated into 100 ml of MM medium with 100 mM threonine and 0.1% fructose as carbon sources instead of glucose. After cultivation at 30°C with shaking at 120 rpm for 24 h, mycelia were harvested by filtration. Approximately 0.1 mg of cell pellets was mechanically broken using a beadshot shocker instrument (2,500 rpm, 10 sec), and proteins were extracted with 2 ml of LY buffer (50 mM Tris·HCl [pH 7.4], 150 mM NaCl, 5 mM EDTA) supplemented with 4% (vol/vol) Complete protease inhibitor cocktail (Roche) using a beadshot shocker (2,000 rpm, 60 sec). The homogeneous suspension was centrifuged at 2,430 × g for 5 min at 4°C, and the supernatant was centrifuged twice under the same conditions. The resulting supernatant represented the total fraction, and this supernatant was centrifuged at 21,880 × g for 15 min at 4°C. The pellet was washed with LY buffer. The centrifugation and wash steps were repeated three times, and the pellet was dissolved in 200 μl of LY buffer and used as the membrane fraction.

The supernatant remaining after the pellet was removed was centrifuged at 100,000 × g for 2 h, and the resulting supernatant was used as the cytoplasmic fraction, while the pellet was used as the microsomal fraction. Each fraction was precipitated with trichloroacetic acid (TCA) and centrifuged, and the pellet was washed three times with ice-cold acetone. The pellet was then dissolved in SDS-PAGE sample buffer. After separation of proteins on 10% SDS-polyacrylamide gels, WscA-HA and WscB-HA were detected using immunoblot analysis with anti-HA monoclonal antibody (Sigma). Proteins were visualized with nito-betacyclooctyl-3-bromochloroiodophosphate (Roche) according to the manufacturer's instructions.

To determine the levels of MpkA and actin expression and the level of MpkA phosphorylation, we inoculated 2 × 10^5 conidia of each strain into 100 ml of YG medium or YG medium with 0.6 M KCl. After cultivation at 30°C with shaking at 120 rpm for 16 h, mycelia were harvested by filtration. Approximately 10^7 conidia of each strain were suspended in 2 ml of PBS buffer (pH 7.4), and the resulting suspension was used as the cytoplasmic fraction. MpkA, actin, and phosphorylated MpkA were detected using anti-p44/42 mitogen-activated protein (MAPK) antibodies (Cell Signaling Technology), anti-actin monoclonal antibody clone C4 (MP Biomedical), and anti-phospho-p44/42 MAPK (Cell Signaling Technology), respectively.

**N-glycosidase treatment of WscA-HA and WscB-HA and immunoblot analysis.** Membrane fraction proteins isolated as described above were treated with N-glycosidase F (Roche) according to the manufacturer's protocol. Briefly, the membrane fraction was denatured at 95°C for 5 min with 1% SDS and treated with N-glycosidase in 200 mM sodium phosphate (pH 8.0), 10 mM EDTA, 1%
2-mercaptoethanol, 0.5% Nonidet P-40, and 0.1% SDS at 37°C for 12 h. The proteins were then precipitated with TCA, washed three times with acetone, and subjected to SDS-PAGE and immunoblot analysis using anti-HA monoclonal antibody (Sigma).

TFMS treatment of WscA-HA and WscB-HA and immunoblot analysis.

The freeze-dried mycelia (20 mg) were broken with metal corn by the multibead shocker at 2,000 rpm for 10 sec. The resultant cells were dissolved in TNE buffer (50 mM Tris-HCl [pH 7.5], 150 mM NaCl, 5 mM EDTA, 1% Triton X-100) as described previously (52) and further broken by a multibead shocker with 0.15 g of 0.5-mm beads at 2,000 rpm for 60 sec. To remove debris, we centrifuged the mixture at 19,060 \(\times g\) for 5 min. The supernatant containing 1 mg protein was transferred to a glass bottle and freeze dried. The glass bottle was sealed with a Teflon-coated butyl rubber stopper and purged with nitrogen gas. Five hundred \(\mu\)l of trifluoromethanesulfonic acid (TFMS; Sigma) was added, and the mixture was incubated on ice for 40 min. It was neutralized by the addition of 5 ml of 1 M Tris solution. After TCA precipitation, the proteins were collected by centrifugation at 2,430 \(\times g\) for 10 min, washed by ethanol, and subjected to SDS-PAGE and immunoblot analysis using anti-HA monoclonal antibody (Sigma).

Indirect immunofluorescence microscopy.

Fixed cells were prepared and processed as described by Harris et al. and Takeshita et al. except that Yatalase (30 mg/ml; Takara) and cellulase R10 (30 mg/ml; Yakult Pharmaceutical) were used for cell lysis (15, 45). A rabbit anti-HA antibody (Sigma) at 1:500 dilution and a...
mouse anti-actin monoclonal antibody C4 clone (MP Biomedical) at 1:200 dilution were used as primary antibodies. Cy3-conjugated anti-mouse IgG antibody (Sigma) at 1:1,000 dilution and Alexa Fluor 488-conjugated anti-rabbit antibody (Invitrogen) at 1:1,000 were used as secondary antibodies. The chitin was stained with calcofluor white. The pictures were taken by a FluoView FV10i confocal laser scanning microscope (Olympus).

Preparation of A. nidulans total RNA. For investigation of the transcription of cell wall-related genes, the conidia of wt and wscA and wscB disruptants (2 x 10⁸) were grown in 100 ml of YG liquid medium or YG liquid medium with 0.6 M KCl in flasks for 18 h at 30°C with shaking at 120 rpm. The mycelia were collected and frozen at -80°C. The frozen mycelia were disrupted using a multibead shocker instrument (2,000 rpm, 10 sec). One milliliter of RNAiso (Takara) was added, and the mycelia were again homogenized using a multibead shocker (2,000 rpm, 5 sec). Chloroform treatment was performed according to the manufacturer's protocol. The resulting RNA was precipitated with isopropanol and rinsed with 70% ethanol, and the pellet was air dried and then dissolved in diethylpyrocarbonate-treated water. Finally, DNase treatment was performed using RQ1 RNase-Free DNase (Promega) according to the manufacturer's protocol.

For analysis of agsB transcription after micafungin treatment, 2 x 10⁸ conidia from the wt and wsc disruptant strains were inoculated into 100 ml of YG medium in a flask and incubated with shaking at 120 rpm at 30°C. After 18 h, micafungin was added to a final concentration of 10 ng/ml. Mycelia were collected at 0, 30, 60, and 120 min and immediately frozen at -80°C. Total RNA was extracted according to the method described above.

Real-time RT-PCR analysis. cDNAs were synthesized using a PrimeScript Perfect real-time reagent kit (Takara) according to the manufacturer's protocol using 400 ng of total RNA as the template. Real-time reverse transcription (RT)-PCR analysis was performed using a LightCycler Quick 330 system (Roche) with SYBR Premix DimerEraser (perfect real time) reagent (Takara). The following primers were used: wscA-RT-F and wscA-RT-R for wscA, wscB-RT-F and wscB-RT-R for wscB, gpdA-RT-F and gpdA-RT-R for gpdA, agsA-RT-F and agsA-RT-F for agsA, gelA-RT-F and gelA-RT-R for gelA, csmA-RT-F and csmA-RT-R for csmA, chsB-RT-F and chsB-RT-R for chsB, fksA-RT-F and fksA-RT-R for fksA, rhoA-RT-F and rhoA-RT-R for rhoA, and histone-RT-F and histone-RT-R for the histone H2B gene (see Table S2 in the supplemental material). The histone H2B gene was used to standardize the mRNA levels of the target genes.

RESULTS

In silico identification of A. nidulans WscA and WscB. Previously, WscA (encoded by ANID_05660.1) was identified as an ortholog of Wsc1 in S. cerevisiae (11). In this study, BLASTP analysis using the amino acid sequences of S. cerevisiae Wsc sensor proteins as the search queries identified three Wsc family sensor candidates, including WscA, ANID_04674.1, and ANID_06927.1. The amino acid se-
sequences of these putative Wsc family proteins were analyzed using Pfam and TMHMM Server v. 2.0. Although WscA and ANID_04674.1 showed structural homology to Wsc1-4 from S. cerevisiae with regard to the domain constitution and N-terminal signal sequence, WSC motif, serine/threonine-rich region, transmembrane domain, and the highly charged C-terminal cytoplasmic region, in silico analysis indicated that ANID_06927.1 does not possess a region after the serine/threonine-rich domain, including a transmembrane domain that is important for membrane localization (Fig. 1). Thus, ANID_04674.1 was termed WscB as a putative stress sensor. Alignments of the WscA, WscB, and Wsc1 sequences from S. cerevisiae and K. lactis showed that the WSC domain (C_1-X-S-X_12–16–Φ-O-S-X_5–7-C_5–7-X_5–8-A-L(I)-X_6–7-C_4–8-X_2–5–11–C_6–9-X_1–2–C_8–9-G-X_6(30)-VY) found in Wsc proteins from S. cerevisiae is highly conserved in WscA and WscB, with the exception of several aromatic amino acids (designated by the symbol Φ) (48) (Fig. 1). Although the conserved KXYQ sequence in the C terminus of S. cerevisiae Wsc proteins was not found in A. nidulans WscA and WscB, the conserved DXXD sequence was found (48). ANID_06927.1 does not possess both KXYQ and DXXD sequences. The primary difference was in the length of the serine/threonine-rich region, which was shorter in WscA and WscB than in S. cerevisiae Wsc1 (81 amino acids [aa] in A. nidulans WscA and 73 aa in WscB). In addition, three putative N-glycosylation sites were found at N135, N176, and N258 in WscA and N41, N250, and N257 in WscB using the NetN Glyc 1.0 server.

The A. nidulans Wsc sequences had 19.4% to 35.8% amino acid identity with the functionally characterized S. cerevisiae Wsc1-4 proteins (see Table S1 in the supplemental material). Phylogenetic analysis showed that WscA and WscB can be classified into the Wsc1 and Wsc4 branches, respectively, with relatively high bootstrap values (data not shown). Wsc1 localizes in the plasma membrane and acts as a main sensor transducer for the CWI signaling pathway in S. cerevisiae (12, 25, 30). On the other hand, Wsc4 localizes in the plasma membrane and acts as a main sensor transducer for the CWI signaling pathway in A. nidulans (12, 25, 30).

Localization and glycosylation of HA-tagged WscA and WscB. WscA and WscB fused with a triple HA tag were expressed in wt A. nidulans under the control of the alcA promoter, and their localization was investigated by immunoblotting (Fig. 2A) and indirect immunostaining (Fig. 2B) using an anti-HA antibody. Both proteins were detected in the total, membrane, and microsomal fractions, but not in the cytoplasmic fraction (Fig. 2A). In addition, indirect immunostaining of WscA-HA and WscB-HA using an anti-HA antibody did not show any fluorescence intensity in intracellular structures, including the ER (Fig. 2B), indicating that Wsc proteins are localized in the cytoplasmic membrane. This result was in agreement with the prediction that WscA and WscB function as cytoplasmic membrane-spanning sensors. The WSC domain is believed to interact with cell wall carbohydrates. The molecular masses of WscA-HA and WscB-HA as determined by SDS-PAGE were 43 to 50 kDa and 55 to 63 kDa, respectively (Fig. 2A). Because the predicted molecular masses of WscA-HA and WscB-HA are 30 and 32 kDa, respectively (see Table S1 in the supplemental material), these two proteins appeared to be posttranslationally modified.

Since WscA and WscB possess putative N-glycosylation sites, we examined the proteins for the presence of N-glycans. Total and membrane fraction proteins were treated with N-glycosidase, and HA-Wsc proteins were subsequently analyzed by SDS-PAGE and immunoblotting (Fig. 3A). After the N-glycosidase treatment, the apparent molecular masses of WscA-HA and WscB-HA decreased to 43 to 48 kDa and 48 to 63 kDa, respectively. The size differences before and after N-glycosidase treatment were consistent with the existence of three N-glycosylation sites with the typical N-glycan structure [Man_3–12, GlcNAc_2] that has been described for Aspergillus glycoproteins (1, 19, 49, 50). Thus, our results suggest that both WscA and WscB are N-glycosylated in A. nidulans. Because WscA and WscB also have O-glycosylation sites, the remaining difference between the observed and native protein molecular weight is believed to be due to the presence of N-glycans.
masses of 30 kDa and 32 kDa, respectively, seems to be due to modification by O-linked glycosylation (11).

We demonstrated previously that WscA-HA is O-glycosylated by the O-mannosyltransferases PmtA and PmtC (11). To further confirm the O-glycosylation, we treated the WscA-HA and WscB-HA proteins expressed in A. nidulans (Fig. 3B) with TFMS, which removed both N- and O-glycans, resulting in a significant decrease in the molecular masses of Wsc proteins. The extents of reduction in molecular mass of Wsc proteins by treatment with TFMS are much larger than those by treatment with N-glycosidase. Thus, these two proteins were also shown to be O-glycosylated.

Deletion of wscA and wscB results in sensitivity to hypo-osmolarity and low pH. To explore the functional roles of wscA and wscB, we generated wsc deletion strains (ΔwscA, ΔwscB, and ΔwscA ΔwscB) (see Fig. S1 in the supplemental material). The ΔwscA strain colonies were significantly smaller in size on YG and MM media than the wt strain colonies (Fig. 4A). The ΔwscA strain conidiophores were of similar structure to those of the wt strain on YG medium (Fig. 3B) with and without 0.6 M KCl. The relative radial growth rates of both the wt strain and ΔwscA strain grown on YG medium with KCl than when grown on YG alone (Fig. 5). On the other hand, the ΔwscB strain formed slightly smaller colonies than did the wt strain. The relative radial growth rates of the ΔwscB strain in YG medium and YG medium with KCl were reduced to 0.91 and 0.93, respectively (Fig. 4B). A double disruption of the wscA and wscB genes (ΔwscA ΔwscB) inhibited growth to a greater degree than did the single disruptions (Fig. 4B). These results indicate that the functions of WscA and WscB are redundant.

The defect in colony formation was facilitated under low-pH conditions (pH 3.8) in the ΔwscA and ΔwscB strains (Fig. 6). The relative radial growth rates of both the ΔwscA and ΔwscB strains were reduced when cells were cultivated on acidified YG medium (pH 3.8) but not when they were cultivated on an alkalinated medium. Colony formation was also compared in the presence of several growth inhibitors (caffeine, calcofluor white, hygromycin B, Congo red, and micafungin). The growth of the ΔwscA and ΔwscB strains did not decrease relative to that of the wt strain when conidia were inoculated into YG medium with growth inhibitors at different concentrations (data not shown).

Effect of wsc disruptions on conidia formation. To examine the effects of wsc deletions on conidiation, mutants were cultivated on YG agar plates or YG agar plates with 0.6 M KCl for 5 days, at which time the numbers of conidia were counted. The number of conidia/cm² in the ΔwscA strain was significantly reduced, to 3% and 2% of the number produced by the wt strain cultured on both YG and YG with KCl, respectively (Fig. 7). Thus, the addition of KCl did not improve conidiation in the ΔwscA strain. On the other hand, the number of conidia produced by the ΔwscB strain was reduced to 71% and 59% of the number produced by the wt strain grown on YG and YG with 0.6 M KCl, respectively. Therefore, conidiation in the ΔwscB strain was inhibited by both hypo-osmotic and hyperosmotic conditions.

Effect of wsc disruptions on cell wall components. The CWI pathway is known to regulate cell wall-related genes during cell wall biosynthesis. We therefore compared the cell wall contents of the wt, ΔwscA, ΔwscB, and ΔwscA ΔwscB strains (Table 2). The alkali-soluble fraction containing α-glucan and soluble β-1,3-/1,6-glucan and the alkali-insoluble fraction containing β-1,3-/1,6-glucan covalently linked to chitin were quantified after 24 h of cultivation in YG liquid medium (3, 7, 22). In the ΔwscA and ΔwscA ΔwscB strains, the amount of α-glucan and soluble β-1,3-/1,6-glucan increased to 130% and 125%, respectively, of that in the wt strain. The alkali-insoluble fraction containing insoluble β-1,3-/1,6-glucan is believed to be responsible for fungal cell wall rigidity (7). However, we found
no significant difference in cell wall composition between the wt and wsc disruptants with regard to the alkali-insoluble fraction and chitin.

**Glycerol concentration in wt and wsc disruptants.** Because polyol concentrations in the cells were associated with stress tolerance (42), we determined them in the wt and wsc disruptants. HPLC analysis revealed that significant amounts of glycerol are present but only trace amounts of trehalose, glucose, and mannitol are present in the wt and wsc disruptants. The glycerol concentration was significantly decreased in the wscA disruptive and the wscA and wscB double disruptant compared to that in the wt strain in the YG medium (Fig. 8).

In addition, the reduced glycerol concentration was retrieved by the addition of 0.6 M KCl into the YG medium and a statistically significant difference in glycerol content among the wt and the wsc disruptant was not shown. This result was consistent with the results that the hyphal growth rates of the wscA disruptant and the wscA and wscB double disruptant were significantly reduced in the YG medium and that the reduced growth was retrieved by the addition of 0.6 M KCl.

**Transcriptional analysis of wscA, wscB, gpdA, and cell wall-related genes.** Since disruption of wscA altered the cell wall...
composition, leading to an increase in the amount of the alkali-soluble fraction containing α-glucan and soluble β-1,3/1,6-glucan in YG medium, we hypothesized that WscA may be involved mainly in the transcriptional regulation of cell wall biogenesis-related genes under hypo-osmotic conditions. To elucidate the regulation of these genes in the wsc disruptants, we compared the transcription of wscA, wscB, gpdA (glycerol-dehydro-3-phosphate dehydrogenase), and cell wall-related genes between the wt and wsc disruptant strains using real-time RT-PCR (Fig. 9). We used RNA templates extracted from log-phase cells cultured for 18 h in YG liquid medium or YG liquid medium supplemented with 0.6 M KCl. Since osmotic stabilization restored the phenotype of wscA disruptants upon the addition of NaCl, KCl, and sorbitol. The complete disappearance of gpdA in the wscA disruptant strain, and similar levels of mRNA for these two genes were detected in the wt and wsc disruptant strains compared to that in the wt strain only when the cells were cultured in YG medium (Fig. 9A). Thus, the response to high osmotic conditions seemed to be active in all four strains.

Next, we examined the transcription of several cell wall biogenesis-related genes, including α-1,3-glucan synthase genes (agsA and agsB), the β-1,3-glucan synthase gene (fksA), the β-1,3-glucanosyl transferase gene (gelA), and chitin synthase genes (csmA and chsB). Among the genes examined, agsA and agsB had significantly altered transcription in the wsc disruptant strains compared to that in the wt strain only when the cells were cultured in YG medium (Fig. 9B and C). The transcription of agsA decreased in all the wsc disruptant strains compared to that in the wt strain. The reduction in transcription was proportional to the degree of deficiency of the wsc disruptant strain phenotypes. On the other hand, the transcription of agsB increased in the ΔwscA strain compared to that in the wt strain. This result agrees with the observation that changes in the cell wall composition of the ΔwscA strain led to an increase in the amount of α-glucan extracted in the alkali-soluble fraction (Table 2). The differences observed among the wt, ΔwscA, ΔwscB, and ΔwscA ΔwscB strains cultured in YG medium were remedied when the cells were cultured in YG supplemented with 0.6 M KCl, as transcription of agsA and agsB was similar in the wt, ΔwscA, ΔwscB, and ΔwscA ΔwscB strains. This result was consistent with the determination that the reduced colony formation and abnormal hyphal structures found in ΔwscA could be reversed in the presence of KCl.

Effect of wsc disruptions on phosphorylation of MpkA. Because the mitogen-activated protein kinase MpkA is a response factor for the CWI signaling pathway, we examined the level of MpkA phosphorylation in the wt and wsc disruptant strains. Interestingly, in YG medium, phosphorylation of MpkA in the wsc disruptants increased even in the ΔwscA ΔwscB strain (Fig. 10). Thus, both Wsc proteins are not essential for the phosphorylation of MpkA. On the other hand, in the presence of 0.6 M KCl, the level of MpkA expression was similar among strains, but its phosphorylation was significantly repressed in the wt and wsc disruptant strains (Fig. 10), indicating that the remedial effect of KCl was due to a reduction in MpkA signaling under the hyperosmotic condition.

Effect of micafungin treatment on transcription of agsB and phosphorylation of MpkA. It has been shown that micafungin treatment transiently upregulates the transcription of agsB via MpkA-RlmA signaling in A. nidulans (8). Therefore, we compared this transcriptional response in the wt and wsc disruptant strains to obtain direct evidence that WscA and WscB are placed upstream of MpkA-RlmA signaling. However, we found that exposure to micafungin activated the transcription of agsB in both the wt strain and the wsc disruptants (Fig. 11A), indicating that the Wsc proteins are not involved in sensing stress associated with exposure to micafungin. A consistent increase in the level of MpkA phosphorylation after micafungin treatment was observed in the ΔwscA ΔwscB strain, to the same degree as that observed in the wt strain (Fig. 11B). The result was also consistent with the observation that the ΔwscA and ΔwscB strains are not hypersensitive to micafungin, in contrast to the wt strain.

**DISCUSSION**

In a previous study, we hypothesized that impairment of WscA is involved in producing some of the morphological deficiencies observed in pmtA and pmtC disruptants (11). In

**TABLE 2. Cell wall compositions of A. nidulans wild-type strain AKU89 and wsc disruptants**

| Strain   | Total sugar (%) | Alkali-soluble fraction (%) | Alkali-insoluble fraction (%) | Total GlcNAc (%) |
|----------|-----------------|-----------------------------|-------------------------------|-----------------|
| AKU89    | 475 ± 71 (100)  | 254 ± 9 (100)               | 176 ± 30 (100)                | 309 ± 8 (100)   |
| ΔwscA    | 537 ± 84 (114)  | 329 ± 15 (130)              | 176 ± 12 (100)                | 297 ± 24 (96)   |
| ΔwscB    | 441 ± 97 (93)   | 278 ± 40 (109)              | 179 ± 40 (102)                | 298 ± 22 (96)   |
| ΔwscA ΔwscB | 536 ± 95 (113) | 317 ± 40 (125)              | 160 ± 10 (91)                 | 291 ± 20 (94)   |

*Measurements are in μg per mg of cell wall.
the present study, we observed several overlapping deficiencies between the wsc and pmt disruptants. The ΔwscA strain had a phenotype similar to that of the pmtA and pmtC disruptants, such as swollen hyphal structures, reduced conidiation, and reduced colony formation when cells were cultured under hypo-osmotic conditions. Osmotic stabilization of the medium remedied these defects in all wscA, pmtA, and pmtC disruptants. In contrast, disruption of pmtB had no significant effect on colony formation. The phenotypes we observed are in close agreement with the determination that O-mannosylation of WscA is catalyzed by PmtA and PmtC, but not by PmtB.

In our previous study on protein O-mannosylation in A. nidulans (11), N-glycosidase and TFMS treatments suggested that WscA and WscB are modified by both O- and N-glycosylation. S. cerevisiae Wsc1 also possesses two potential N-glycosylation sites; however, it is posttranslationally modified only by O-glycosylation (35). Both O- and N-glycosylation of the S. cerevisiae Mid2 sensor protein have been reported (17). O-mannosylation determines the stability of Mid2. On the other hand, N-glycosylation near the N-terminal end (N35) is required for proper function of Mid2. It has been suggested that the N-glycan may be directly involved in Mid2 sensing.

The sensing spectrum in A. nidulans seems to be different from that in the well-studied S. cerevisiae Wsc proteins. In our study, both the ΔwscA and ΔwscB strains exhibited hypersensitivity to stress associated with acidic but not alkaline conditions, indicating that WscA and WscB sense the perturbations in the cell wall caused by exposure to low pH. In contrast, S. cerevisiae Wsc1 is involved in the response to stress associated with exposure to alkaline conditions (43). Interestingly, phenotypic analysis suggested that WscA and WscB are not involved in sensing stress associated with exposure to antifungal agents, including caffeine, calcofluor white, hygromycin B, Congo red, and micafungin. In S. cerevisiae, Wsc1, but not Wsc2, Wsc3, or Mid2, mediates caspofungin-induced PKC pathway activation (36). Caspofungin is an echinocandin class β-1,3-glucan synthase inhibitor similar to micafungin. In A. nidulans, the downstream components of the CWI signaling pathway are involved in tolerance to antifungal agents. For example, the dominant negative rhoA<sup>E40I</sup> allele confers hypersensitivity to calcofluor white and caspofungin (13), while the ΔmpkA strain exhibits sensitivity to calcofluor white and micafungin, and the ΔrlmA strain is sensitive to calcofluor white (8). Consistently, the transcriptional level of rhoA was unchanged among wt and wsc disruptants (data not shown). Moreover, micafungin treatment transiently upregulates the transcription of agsB via MpkA-RlmA signaling in A. nidulans (8). In this study, a double disruptant strain of wscA and wscB demonstrated a transient upregulation of agsB and increased phosphorylation of MpkA after treatment with micafungin, similar to the wt strain. This suggests that WscA and WscB are not required for MpkA signaling and either that the sensor responsible for responding to stress associated with micafungin exposure has not been identified or that micafungin independently activates the downstream pathway of CWI signaling.

Although the BLASTP search did not identify any orthologous MID2 sensor genes in the Aspergillus genomes, a S. cerevisiae Cwh43 homolog was found (8). Cwh43 is a putative sensor/transporter protein that acts upstream of the BCK2 branch of the PKC1-dependent cell wall integrity pathway and is involved in cell wall biogenesis (29). The principal structural feature of Cwh43 is the presence of 14 to 16 transmembrane segments and several putative glycosylation and phosphorylation sites.
The colony formation and abnormal hyphal structures observed with the wsc disruptant strains were remedied when cells were cultured under high osmotic conditions. Therefore, the osmoreponsive activation pathway is retained in the wt and both wsc disruptants, indicating that WscA and WscB are dispensable in osmoadaptation. The osmoinducible gpdA gene was consistently upregulated in the wsc disruptant strains, and the level of MpkA phosphorylation was equally depressed under high osmotic conditions in the wt strain and all of the wsc disruptants. Osmotic signals are known to activate the high-osmolarity glycerol (HOG) pathway. Genome sequencing analyses have revealed that A. nidulans has genes orthologous to all the genes of the HOG response MAPK pathway of S. cerevisiae (14). Activation of the A. nidulans HOG pathway depends solely on the two-component signaling system, and MAPKK activation mechanisms in the A. nidulans HOG pathway differ from those in the yeast model (9). Although abnormal colony size was remedied by the addition of KCl in our study, conidiation efficiency was not recovered in the wsc disruptant strains. Wsc proteins seem to be required for the conidiation event even under high-osmolarity conditions.

In A. nidulans, the putative stress sensors WscA and WscB are involved in CWI under hypo-osmotic and acidic pH conditions. However, direct evidence of transcriptional response to cell wall-related stress has not been obtained. Further study is therefore needed to fully elucidate the stress sensor functions of these Wsc family proteins in A. nidulans.

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