RESEARCH ARTICLE

Functional Characterization of NAC and MYB Transcription Factors Involved in Regulation of Biomass Production in Switchgrass (Panicum virgatum)

Ruiqin Zhong, Youxi Yuan, John J. Spiekerman, Joshua T. Guley, Jane Frances C. Egboisiuba, Zheng-Hua Ye*

Department of Plant Biology, University of Georgia, Athens, Georgia, 30602, United States of America

* yh@uga.edu

Abstract

Switchgrass is a promising biofuel feedstock due to its high biomass production and low agronomic input requirements. Because the bulk of switchgrass biomass used for biofuel production is lignocellulosic secondary walls, studies on secondary wall biosynthesis and its transcriptional regulation are imperative for designing strategies for genetic improvement of biomass production in switchgrass. Here, we report the identification and functional characterization of a group of switchgrass transcription factors, including several NACs (PvSWNs) and a MYB (PvMYB46A), for their involvement in regulating secondary wall biosynthesis. PvSWNs and PvMYB46A were found to be highly expressed in stems and their expression was closely associated with sclerenchyma cells. Overexpression of PvSWNs and PvMYB46A in Arabidopsis was shown to result in activation of the biosynthetic genes for cellulose, xylan and lignin and ectopic deposition of secondary walls in normally parenchymatous cells. Transactivation and complementation studies demonstrated that PvSWNs were able to activate the SNBE-driven GUS reporter gene and effectively rescue the secondary wall defects in the Arabidopsis snd1 nst1 double mutant, indicating that they are functional orthologs of Arabidopsis SWNs. Furthermore, we showed that PvMYB46A could activate the SMRE-driven GUS reporter gene and complement the Arabidopsis myb46 myb83 double mutant, suggesting that it is a functional ortholog of Arabidopsis MYB46/ MYB83. Together, these results indicate that PvSWNs and PvMYB46A are transcriptional switches involved in regulating secondary wall biosynthesis, which provides molecular tools for genetic manipulation of biomass production in switchgrass.

Introduction

Switchgrass (Panicum virgatum L.) is considered to be one of the leading renewable and sustainable feedstock crops for the production of biofuels. It is a perennial grass species with a
high yield of aboveground biomass and high productivity when grown on marginal land, which is in part due to its beneficial physiological traits such as the production of deep roots and the utilization of C₄ photosynthetic metabolism [1,2]. Furthermore, when compared with first-generation biofuel crops, switchgrass has been shown to be extremely efficient in its consumption of water and nutrients with low fertilizer inputs during development [1,2]. Because the switchgrass biomass used for biofuel production is mainly the lignocellulosic cell wall residues from stems and leaves, much attention has been paid toward altering cell wall composition in order to reduce biomass recalcitrance for conversion into biofuels.

Several lignin biosynthetic genes have been targeted for reduction in lignin content in switchgrass as lignin is one of the main factors contributing to biomass recalcitrance. Downregulation of Caffeic Acid O-Methyltransferase (COMT) in switchgrass by RNA interference (RNAi) resulted in a reduction in lignin content and consequently, a higher yield of biomass conversion into ethanol compared with the wild type under the same saccharification and fermentation conditions [3]. Similarly, downregulation of Cinnamyl Alcohol Dehydrogenase (CAD) or 4-Coumarate:Coenzyme A Ligase (4CL) in switchgrass by RNAi caused a reduction in lignin content and an increase in biomass saccharification efficiency [4,5]. A reduction in lignin content and an improvement in saccharification efficiency in switchgrass were also achieved by overexpression of a MYB transcription factor, PvMYB4, which represses the expression of lignin biosynthetic genes [6]. These studies provide convincing evidence that a reduction in lignin content in switchgrass is a feasible approach to overcome biomass recalcitrance.

To develop switchgrass as a bioenergy feedstock, it is important to unravel the mechanisms underlying the biomass biosynthesis and the transcriptional regulation of biomass production. Uncovering genes involved in biomass biosynthesis and its regulation may provide additional tools to further manipulate switchgrass biomass yield and composition tailored for biofuel production. Genes involved in lignin biosynthesis in switchgrass have been identified using an inducible switchgrass cell suspension system and several of them have been confirmed for their involvement in lignin biosynthesis by RNAi downregulation of their expression in switchgrass [7]. Four Sucrose Synthase (SUS) genes, which are potentially involved in the supply of UDP-glucose for cellulose biosynthesis, have been functionally characterized in switchgrass. Overexpression of one of them, PvSUS1, in switchgrass led to increased plant height and biomass, suggesting the potential usefulness of SUS genes as targets to increase biomass production in switchgrass [8]. In addition, as mentioned above, PvMYB4 has been shown to be a transcriptional repressor regulating lignin biosynthesis [6]. Because the bulk of switchgrass biomass is from sclerenchyma cells with heavily thickened secondary walls, further studies of genes involved in secondary wall biosynthesis and its regulation will be important for better utilization of switchgrass biomass.

In this report, we present the identification and functional characterization of several NAC and MYB genes involved in regulation of secondary wall biosynthesis. In Arabidopsis, a group of secondary wall NACs (SWNs) function as master transcriptional switches activating the entire secondary wall biosynthesis program in vessels and fibers [9]. Arabidopsis SWNs are vessel-specific VND1 to VND7 and fiber-specific SND1/NST1/2, and when overexpressed, they are sufficient to activate the biosynthetic pathways for cellulose, xylan and lignin as well as the genes for programmed cell death [10–19]. They regulate their downstream target genes, including a number of transcription factors and secondary wall biosynthetic genes, by binding to the secondary wall NAC binding element (SNBE) consisting of a 19-bp imperfect palindromic consensus sequence, (T/A)NN(C/T)(T/C/G)TNNNNNNNA(A/C)GN(A/C/T)(A/T) [20]. The SWN-regulated downstream transcription factors include secondary wall-associated SND2, SND3, MYB20, MYB42, MYB43, MYB46, MYB52, MYB58, MYB63, MYB69, MYB83, MYB85, MYB103 and KNAT7 [21–24], and among them, MYB46 and MYB83 act as second-
level master switches and bind to the secondary wall MYB responsive element [SMRE; ACC (A/T)A(A/C)(T/C)] to activate their target genes [25,26]. Orthologs of SWNs and MYB46/83 from rice, maize and Brachypodium have also been demonstrated to regulate secondary wall biosynthesis, indicating the evolutionary conservation of the SWN-mediated transcriptional network controlling secondary wall biosynthesis [27–29]. By searching the switchgrass genome, we have identified 14 SWN homologs (PvSWNs) and two MYB46/MYB83 homologs (PvMYB46A/B). We show that PvSWNs and PvMYB46A are capable of activating the expression of the biosynthetic genes for cellulose, xylan and lignin, leading to ectopic deposition of secondary walls when overexpressed in Arabidopsis. We further demonstrate that PvSWNs can effectively activate the expression of the SNBE-driven GUS reporter gene and complement the Arabidopsis snd1 nst1 double mutant. Likewise, PvMYB46A is able to activate the SMRE-driven GUS reporter gene and rescue the Arabidopsis myb46 myb83 double mutant phenotypes. Our findings indicate that PvSWNs and PvMYB46A are functional orthologs of Arabidopsis SWNs and MYB46/MYB83, respectively, and are involved in regulating secondary wall biosynthesis in switchgrass.

Results
Expression analysis of PvSWNs
In the switchgrass (Panicum virgatum) genome, there exist 14 NAC genes that are close homologs of Arabidopsis secondary wall NAC master switches (SWNs). They appear to be divided into three subgroups based on the phylogenetic analysis (Fig 1A). PvSWN1/2A/2B are grouped closely with SND1/NST1/2, PvSWN3A/3B are grouped closely with VND7, and PvSWN4-8 are grouped closely with VND1-6. For each PvSWN, there exist close homologs in rice, maize and/or Brachypodium (Fig 1A). Except PvSWN1 and PvSWN5, the rest of PvSWNs form pairs, of which the two members share at least 92.4% similarity at the amino acid level. Since switchgrass is a tetraploid species, the gene pairing of PvSWNs is likely due to the increased ploidy level. In this study, eight PvSWNs, including PvSWN1, PvSWN5 and one member from each pair of other PvSWNs, were functionally characterized. Quantitative PCR analysis showed that PvSWN genes were all expressed in leaf blades, leaf sheaths and stems, and among them, PvSWN2A, PvSWN7A and PvSWN8A exhibited a high level expression in stems (Fig 1B). Examination of lignin deposition in mature switchgrass stems showed that in addition to its heavy deposition in the secondary walls in xylem and cortical sclerenchyma cells, lignin was present in the walls of ground cells (Fig 1C). A low level of lignin staining was also observed in the walls of epidermal cells and some phloem parenchyma cells. In situ hybridization revealed that PvSWN2A, PvSWN7A and PvSWN8A were expressed in xylem bundle cells, cortical sclerenchyma cells, and ground cells in switchgrass stems (Fig 1D–1F), which were matched with the lignin-staining pattern in these tissues (Fig 1C). However, hybridization signals were also seen in those patches of cortical cells below epidermis that did not deposit lignified secondary walls, which could be resulted from non-specific hybridization. The control stem section hybridized with the sense probe of PvSWN2A exhibited no hybridization signal (Fig 1G). These findings indicate that the expression of PvSWN2A, PvSWN7A and PvSWN8A is associated with secondary wall deposition and lignification in switchgrass stems.

Overexpression of PvSWNs in Arabidopsis results in ectopic deposition of cellulose, xylan and lignin
To find out whether PvSWNs are involved in regulating secondary wall biosynthesis, we investigated the effects of PvSWN overexpression on secondary wall deposition in transgenic
Arabidopsis. The full-length cDNAs of \textit{PvSWNs} driven by the \textit{Cauliflower mosaic virus} (CaMV) 35S promoter were introduced into wild-type Arabidopsis and at least 30 transgenic plants for each \textit{PvSWN}-OE construct were generated. At least one-third of the transgenic plants for each of the eight constructs had small rosettes with leaves curled upward (Fig 1A), a phenotype...
similar to that observed in the SND1 overexpressors [13]. Examination of PvSWN-OE leaves revealed that in contrast to the wild-type leaves, which only had lignified helical secondary wall thickening in veins (Fig 2B and 2C), the PvSWN overexpressors had lignified secondary walls in many mesophyll cells (Fig 2D–2L), which were deposited in a helical or reticulated pattern. The
ectopically deposited secondary walls in the leaf mesophyll cells most likely restricted the normal expansion of leaves, leading to the small, curly leaf phenotype.

The ectopic deposition of secondary walls also occurred in the epidermal and cortical cells of PvSWN-OE stems (Fig 3). The most heavily thickened secondary walls were observed in the epidermal and cortical cells of stems of PvSWN4A-OE (Fig 3E), PvSWN5-OE (Fig 3F) and
PvSWN7A-OE (Fig 3H). It was noted that the ectopic secondary walls in some cortical cells of the PvSWN7A-OE stems (Fig 3K) were 2 to 3 times thicker than those of the interfascicular fibers of wild-type stems (Fig 3J). To investigate whether the ectopically deposited cell walls contained all three secondary wall components, stem sections of PvSWN overexpressors were detected for cellulose and lignin by histological staining and for xylan by immunostaining. It was found that although the epidermal and cortical cells of wild-type stems showed no staining for lignin (Fig 4A) and xylan (Fig 5A) and little staining for cellulose (Fig 5J), the ectopically deposited secondary walls in the epidermal and cortical cells of PvSWN-OE stems were heavily stained for lignin (Fig 4B–4I), xylan (Fig 5B–5I) and cellulose (Fig 5K–5R). These results indicate that PvSWN overexpression activates the biosynthetic pathways of cellulose, xylan and lignin, leading to ectopic deposition of secondary walls in normally parenchymatous cells.

Overexpression of PvSWNs in Arabidopsis induces the expression of secondary wall-associated transcription factors and secondary wall biosynthetic genes

To further substantiate the finding that PvSWNs were capable of activating the secondary wall biosynthetic program, we investigated whether PvSWN overexpression induced the expression...
Fig 5. Ectopic deposition of xylan and cellulose in the epidermal and cortical cells in the stems of PvSWN overexpressors. Cross sections of stems were stained for xylan with the xylan-specific monoclonal antibody LM10 (A to I) and for cellulose with Calcofluor White. (A) Section of a wild-type stem showing xylan staining in the walls of interfascicular fibers and its absence in the epidermal and cortical cells. (B to I) Stem sections of PvSWN overexpressors showing ectopic xylan deposition in the epidermal cells (arrowheads) and cortical cells (arrows) in addition to its normal deposition in the interfascicular fibers and xylem cells. (J) Section of a wild-type stem showing the predominant cellulose staining in the walls of interfascicular fibers. (K to R) Stem sections of PvSWN overexpressors showing intense cellulose staining in the epidermal cells (arrowheads) and cortical cells (arrows) in addition to the interfascicular fibers and xylem cells. co, cortex; ep, epidermis; if, interfascicular fiber; xy, xylem. Bar in (A) = 256 μm for (A to R).
of secondary wall-associated transcription factors and secondary wall biosynthetic genes. 
SND1 has been shown to regulate the expression of a number of downstream transcription factors, including SND2, SND3, MYB46, MYB83, MYB58, MYB63, MYB85, MYB103 and KNAT7, all of which have been demonstrated to regulate secondary wall biosynthesis [9]. Quantitative PCR analysis revealed that the expression of all these SND1-regulated transcription factors was highly induced in the PvSWN-OE lines (Fig 6). The induction level of each transcription factor appeared to vary significantly among different PvSWN overexpressors, which may be due to the differential activation strength by PvSWNs.

Examination of the expression of secondary wall biosynthetic genes showed that PvSWN overexpression induced the expression of biosynthetic genes for secondary wall cellulose [represented by Cellulose SynthaseA4 (CesA4), CesA7 and CesA8], xylan [represented by Fragile Fiber8 (FRA8) and Irregular Xylem9 (IRX9)], and lignin [represented by Hydroxycinnamoyl CoA-Shikimate Hydroxycinnamoyl Transferase (HCT) and Caffeoyl CoA O-Methyltransferase (CCoAOMT)] [30] (Fig 7A). These results demonstrated that congruent with the observed ectopic secondary wall deposition, PvSWN overexpression activated the expression of both secondary wall-associated transcription factors and secondary wall biosynthetic genes.

Arabidopsis vessel-specific SWNs, including VND1 to VND7, have been previously shown to regulate the expression of not only secondary wall biosynthetic genes but also genes involved in programmed cell death [15,17,20]. Quantitative PCR analysis revealed that three representative genes involved in programmed cell death, Bifunctional Nuclease1 (BFN1), Ribonuclease3 (RNS3) and Xylem Cysteine Peptidase1 (XCP1), were highly induced by overexpression of PvSWN3A, PvSWN4A, PvSWN5, PvSWN6A, PvSWN7A and PvSWN8A (Fig 7B), which are close homologs of VND1 to VND7 (Fig 1A). In contrast, overexpression of PvSWN1 and PvSWN2A, which are close homologs of SND1 and NST1/2, did not significantly affect the expression levels of BFN1, RNS3 and XCP1 (Fig 7B).

Complementation of the Arabidopsis snd1 nst1 mutant by PvSWNs

To test whether PvSWNs were capable of complementing the secondary wall defects conferred by the snd1 and nst1 mutations, we expressed PvSWNs under the SND1 promoter in the Arabidopsis snd1 nst1 double mutant. The snd1 nst1 double mutant had a pendent stem phenotype due to weakened stem strength caused by lack of lignified secondary walls in interfascicular and xylary fibers [11,13,14] (Fig 8A–8D). Expression of PvSWNs in the snd1 nst1 double mutant rescued the pendent stem phenotype (Fig 8A) and restored the mechanical strength of stems (Fig 8B). Examination of cross sections of stems showed that PvSWNs completely restored the lignified secondary walls in interfascicular fibers and xylary fibers (Fig 8E–8L). These results provide additional lines of evidence demonstrating that PvSWNs are functional orthologs of Arabidopsis SWNs and are involved in regulating secondary wall biosynthesis.

It has been shown that Arabidopsis SWNs bind to and activate the 19-bp semi-palindromic Secondary Wall NAC Binding Element (SNBE) sequence in the promoters of their target genes [20]. Using transactivation analysis, we next tested whether PvSWNs were capable of activating the SNBE-driven GUS reporter gene expression. Coexpression of the CaMV 35S promoter-driven PvSWNs and the SNBE-driven GUS reporter gene in Arabidopsis protoplasts demonstrated that PvSWNs effectively activated the GUS reporter gene driven by SNBE sequences from the promoters of MYB46, MYB83, SND3 and KNAT7 (Fig 9), indicating that like Arabidopsis SWNs, PvSWNs bind to the SNBE sites to activate their target genes.
Fig 6. Induction in the expression of secondary wall-associated transcription factors in PvSWN overexpressors. Leaves of 4-week-old transgenic Arabidopsis plants overexpressing PvSWNs were extracted for RNA, which was subsequently used for quantitative PCR analysis of the expression of various transcription factors. The expression level of each gene in the wild type was set to 1. Error bars represent the SE of three biological replicates.

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Fig 7. Induction in the expression of secondary wall biosynthetic genes and programmed cell death genes in PvSWN overexpressors. Leaves of 4-week-old transgenic Arabidopsis plants overexpressing PvSWNs were extracted for RNA, which was subsequently used for quantitative PCR analysis of the expression of secondary wall biosynthetic genes (A) and programmed cell death genes (B). The expression level of each gene in the wild type was set to 1. Error bars represent the SE of three biological replicates.

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Fig 8. PvSWNs functionally complement the secondary wall defects in the Arabidopsis snd1 nst1 double mutant. Expression of PvSWNs in the Arabidopsis snd1 nst1 double mutant was driven by the SND1 promoter. (A) PvSWNs rescued the pendent inflorescence stems of snd1 nst1 to the upright form. (B) Measurement of stem breaking strength showing that PvSWNs restored the stem strength of snd1 nst1 to the wild-type level. Each bar denotes the breaking force of inflorescence stems of individual plants. (C) Cross section of a wild-type stem showing lignified secondary walls in the xylem and interfascicular fibers. (D) Cross section of an snd1 nst1 stem showing lignified secondary walls only in xylem vessels but not in interfascicular fibers and xylary fibers. (E to L) Cross sections of stems of PvSWN-complemented snd1 nst1 showing that PvSWNs restored the lignified secondary walls in interfascicular fibers and xylary fibers in snd1 nst1. The bottom parts of 7-week-old stems were sectioned and stained for lignin with phloroglucinol-HCl. if, interfascicular fiber; xy, xylem. Bar in (C) = 180 μm for (C to L).

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Fig 9. PvSWNs activate the expression of SNBE-driven GUS reporter gene. (A) Diagrams of the effector constructs and the GUS reporter constructs. Three copies of SNBE sequences were ligated upstream of the CaMV 35S minimal promoter linked with the GUS reporter gene. (B) SNBE sequences used for transactivation analysis are from the promoters of MYB46, MYB83, SND3 and KNAT7. The consensus SNBE sequence is shown at the bottom. (C) Transactivation analysis showing the activation of SNBE-driven GUS reporter gene by PvSWNs. Arabidopsis leaf protoplasts were co-transfected with the reporter and effector constructs as shown in (A) and assayed for GUS activity. The GUS activity in the control protoplasts transfected with the GUS reporter constructs and an empty effector construct without the PvSWN sequences is set to 1. Error bars denote the SE of three biological replicates.

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PvMYB46A is a functional ortholog of Arabidopsis MYB46/MYB83

To further expand our understanding of the transcription regulation of secondary wall biosynthesis in switchgrass, we blast-searched the switchgrass genome for MYB sequences that were close homologs of Arabidopsis MYB46/MYB83, which have been shown to be second-level master switches regulating secondary wall biosynthesis [21,23]. It was found that the switchgrass genome harbors two close homologs of MYB46/MYB83, which was named PvMYB46A and PvMYB46B (Fig 10A). They share 89% identity and 91% similarity and like paired...
PvSWNs, they are likely originated from the increased ploidy level. We thus chose one of them, PvMYB46A, for functional characterization. PvMYB46A/B are more closely grouped together with three other MYB46 homologs from grass species (OsMYB46, BdMYB46 and ZmMYB46) [27, 28]; Arabidopsis MYB46/83 are more closely grouped together with four wood-associated MYB46 homologs (PtrMYB3/20/2/21) from poplar [31, 32] (Fig 10A). Quantitative PCR analysis showed that PvMYB46A was highly expressed in leaf sheaths and stems (Fig 10B), and in situ mRNA localization revealed that in switchgrass stems, PvMYB46A was expressed in xylem bundles, cortical sclerenchyma cells and ground cells (Fig 10D), the tissues that underwent lignification (Fig 1C). However, hybridization signals were also seen in those patches of cortical cells below epidermis that did not deposit lignified secondary walls, which could be resulted from non-specific hybridization. In addition, expression of PvMYB46A in the Arabidopsis myb46 myb83 double mutant completely rescued the mutant phenotypes (Fig 10C), including the arrested seedling growth and the loss of helical secondary wall thickening in leaf veins [23]. These results indicate that PvMYB46A is a functional ortholog of Arabidopsis MYB46/MYB83 involved in regulating secondary wall biosynthesis.

PvMYB46A overexpression leads to ectopic deposition of secondary walls

We next investigated whether PvMYB46A could activate the secondary wall biosynthetic program when overexpressed in Arabidopsis. Transgenic seedlings overexpressing PvMYB46A had small, upward-curving leaves compared with the wild type (Fig 11A), a phenotype similar to that of PvSWN overexpressors (Fig 2A). Quantitative PCR analysis revealed that PvMYB46A overexpression induced the biosynthetic genes for secondary wall cellulose (CesA4, CesA7 and CesA8), xylan (IRX9 and FRA8), and lignin (HCT and CCoAOMT), but not the programmed cell death genes (BFN1, RNS3 and XCP1) (Fig 11B). The expression of a number of secondary wall-associated transcription factors, including SND3, MYB58, MYB63, MYB85 and KNAT7 was also induced by PvMYB46A (Fig 11C).

Consistent with the induction in the expression of secondary wall biosynthetic genes, PvMYB46A overexpressors had ectopic deposition of lignified secondary walls in the epidermal cells of leaves (Fig 11F and 11G) compared with the epidermis of wild-type leaves, which only displayed weak lignin autofluorescence in guard cells (Fig 11D and 11E). The epidermal cells of stems of PvMYB46A overexpressors also exhibited ectopic deposition of secondary walls with intense staining for lignin, xylan and cellulose (Fig 11H and 11J) compared with the wild type (Figs 4A, 5A and 5J). The control wild-type stem sections omitted with Calcofluor White (Fig 11H) or LM10 xylan antibody (Fig 11I) did not show any fluorescence signals under the same detection conditions. These results indicate that similar to Arabidopsis MYB46/MYB83, PvMYB46A is also able to activate the secondary wall biosynthetic program.

PvMYB46A activates the SMRE-driven GUS reporter gene

It has been shown that Arabidopsis MYB46/MYB83 activate their target genes by binding to the SMRE sequences in their promoters [25]. Transactivation analysis of the SMRE-driven GUS reporter gene revealed that PvMYB46A was able to activate the GUS reporter gene driven by all 8 variants of SMRE sequences (Fig 12A and 12B). The activation levels of the GUS reporter gene varied among the SMRE sequences, indicating that PvMYB46A exhibits differential activation strength toward the 8 variants of SMRE sequences. In contrast, the GUS reporter gene driven by a mutant form of SMRE1 (mSMRE1) showed no activation by PvMYB46A (Fig 12B). Together, these results indicate that like Arabidopsis MYB46/MYB83, PvMYB46A regulates its target gene expression by activating the SMRE sites.
Fig 11. PvMYB46A overexpression induces ectopic deposition of secondary walls in the epidermal cells of leaves and stems of Arabidopsis. (A) Three-week-old seedlings showing the smaller rosette size and curly leaves of the PvMYB46A overexpressor (PvMYB46A-OE) compared with the wild type. (B) Quantitative PCR analysis showing the induction in the expression of secondary wall biosynthetic genes (CesA4, CesA7, CesA8, FRA8, IRX9, PARVUS, PAL1, HCT and CCoAOMT1) in the PvMYB46A overexpressors. No induction of the programmed cell death genes (XCP1, BFN1 and RNS3) was detected. The expression level of each gene in the wild type is set to 1 and the error bars represent the SE of three biological replicates. (C) Quantitative PCR analysis showing the induction in the expression of several secondary wall-associated transcription factors in the PvMYB46A overexpressors. (D, E) Differential interference contrast (DIC) image (D) and lignin autofluorescence image (E) of a wild-type leaf showing the absence of lignin signals in the epidermis. (F, G) DIC (F) and lignin autofluorescence image (G) of a PvMYB46A-OE leaf showing lignified walls in the epidermal cells. (H) Control wild-type stem section that was omitted with Calcofluor White showed no fluorescence signals. (I) Control wild-type stem section that was incubated with fluorescein isothiocyanate-conjugated goat-anti-rat secondary antibodies but without LM10 xylan antibody showed no fluorescence signals. (J to L) Ectopic deposition of lignin (J), cellulose (K) and xylan (L) in the epidermal cells of PvMYB46A-OE stems. Cross sections of the stems were stained for lignin with phloroglucinol-HCl, cellulose with Calcofluor White, and xylan with the xylan-specific monoclonal antibody LM10. ep, epidermis; if, interfascicular fiber. Bars = 63 μm in (D) to (G) and 114 μm in (H) to (J).

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Fig 12. PvMYB46A activates the expression of SMRE-driven GUS reporter gene. (A) Diagrams of the effector construct and the GUS reporter constructs. Three copies of SMRE sequences were ligated upstream of the CaMV 35S minimal promoter linked with the GUS reporter gene. (B) Transactivation analysis showing the activation of SMRE-driven GUS reporter gene by PvMYB46A. The SMRE sequences used in the GUS reporter constructs are shown at the left of the corresponding bars. The effector construct and the GUS reporter constructs as shown in (A) were co-transfected into Arabidopsis leaf protoplasts, which were subsequently assayed for GUS activity. The GUS activity in the control protoplasts transfected with the GUS reporter constructs and an empty effector construct without the PvMYB46A sequence is set to 1. Error bars denote the SE of three biological replicates.

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Discussion

The main above-ground biomass of switchgrass targeted for biofuel production is from stems and leaves. Large amount of lignified secondary walls are deposited in the stems of switchgrass at maturity. In particular, the bundle sheath fibers and cortical sclerenchyma cells are heavily thickened with secondary walls, leaving extremely small intracellular spaces (Fig 1C). To accommodate the heavy deposition of secondary walls in these cells, it is conceivable that switchgrass possesses some mechanisms to sustain the robust secondary wall biosynthetic activity in these cells. One way to achieve this is through a strong, sustained activation of the expression of secondary wall biosynthetic genes, which can be fulfilled by transcription factors activating their expression. Our functional studies on switchgrass NAC and MYB transcription factors have revealed that among the secondary-wall NAC and MYB master switches, three of them, namely PvSWN4A, PvSWN5 and PvSWN7A, exhibit a strong activation of the secondary-wall biosynthetic program (Figs 3–5), and when overexpressed in Arabidopsis, they could lead to ectopic deposition of extremely thick secondary walls that are up to three times thicker than the walls of wild-type fiber cells (Fig 3). This finding indicates that PvSWN4A/5/7A are strong master switches activating secondary wall biosynthesis leading to massive deposition of secondary walls. If so, these PvSWNs could potentially be used as molecular tools to induce massive deposition of secondary walls in switchgrass and other biofuel crops so that the lignocellulosic biomass yield per land acreage could be increased.

The Arabidopsis SWNs can be divided into two groups based on their expression patterns, i.e., vessel-specific SWNs (VNDs) and fiber-specific SWNs (SND1/NSTs). Orthologs of both vessel-specific VNDs and fiber-specific SND1/NSTs are present in switchgrass and other grass species including rice, maize and Brachypodium (Fig 1A) [27–29]. However, SWNs from switchgrass, rice and maize appear to be expressed in all secondary wall-forming cells including both vessels and fibers in stems [27,29], indicating that unlike those in Arabidopsis, SWNs in grass species did not diverge into vessel- or fiber-specific groups.

The Arabidopsis vessel-specific SWNs (VNDs) have been shown to induce the expression of not only secondary wall biosynthetic genes, but also programmed cell death genes [15,17,20]. When overexpressed in Arabidopsis, PvSWN3A to PvSWN8A, which are orthologs of VNDs, were also capable of activating programmed cell death genes. In contrast, PvSWN1 and PvSWN2A, which are orthologs of Arabidopsis fiber-specific SND1 and NST1/2, had little effect on the expression of programmed cell death genes. Activation of programmed cell death genes by Brachypodium orthologs of VNDs but not those of SND1/NST1/2 was also observed [28]. These findings indicate that although switchgrass orthologs of the Arabidopsis VNDs do not exhibit vessel-specific expression [27,29], they retained similar functionality as VNDs in activating not only secondary wall biosynthetic genes but also programmed cell death genes.

Our finding that PvMYB64A is sufficient to activate secondary wall biosynthetic pathways for cellulose, xylan and lignin provides further molecular evidence demonstrating the evolutionary conservation of the SWNs- and MYB46/83-mediated transcriptional network. So far, several MYB46 homologs from dicots (poplar, Eucalyptus and pine) [16,31–35] and grass species (rice and maize) [27] have been shown to be functional orthologs of MYB46/83 regulating secondary wall biosynthesis. All these MYB46 homologs activate their target genes by binding to the SMRE sites, indicating that the functional conservation is achieved through their binding to the common cis-elements for transcriptional activation. It is interesting to note that PtMYB46A activates the expression of secondary wall biosynthetic genes but not the genes involved in programmed cell death. This finding implies that although PvSWNs and PtMYB46 co-regulate the secondary wall biosynthetic program, PtMYB46 is unlikely involved in activating the programmed cell death program during vessel development, which is consistent with
the previous finding showing that in Arabidopsis, the vessel-specific VNDs directly regulate the expression of programmed cell death genes through binding and activating the SNBE sites [15,17,18,20].

Considerable interest has recently been focused on developing perennial grasses, such as switchgrass and *Miscanthus*, as a source of renewable energy; therefore, a better understanding of genes involved in the biosynthesis of grass cell walls is deemed necessary [36]. Although secondary cell walls from both grasses and dicots are made of cellulose, hemicelluloses and lignin, they differ in their composition [37]. For example, xylan from grasses is substituted with arabinosyl residues, which are often cross-linked with lignin via ferulic acid, whereas xylan from dicots has few arabinosyl substitutions. In addition, lignin from grasses contain a much higher amount of p-hydroxyphenyl units than that from dicots [37]. The heterogeneity of cell walls can be controlled by cell wall biosynthetic genes as well as transcriptional regulators of the biosynthetic genes. So far, only a few studies have been devoted to the transcriptional regulation of grass cell wall biosynthesis [27–29, 38,39]. Available evidence indicates that the NAC and MYB master transcriptional switches activating the secondary wall biosynthetic program are conserved between grasses and dicots [27–29]. Furthermore, it appears that the grass orthologs of the Arabidopsis MYB transcriptional repressors for lignin biosynthesis are also functionally conserved [6,40,41]. However, since grass cell walls differ from those of dicots in their composition, it is conceivable that the transcription regulation of secondary wall biosynthesis in grass species might have some divergence from that in dicots. Further identification and functional characterization of switchgrass regulatory genes controlling secondary wall biosynthesis will not only help decipher how grass cell walls are constructed but also offer additional molecular tools to genetically modify grass cell wall composition better suited for biofuel production.

**Methods**

**Gene expression analysis**

Total RNA was isolated from switchgrass and Arabidopsis tissues with a Qiagen RNA isolation kit (Qiagen). After treatment with DNase I, the total RNA was converted into first strand cDNAs, which were then used as templates for PCR analysis. The real-time quantitative PCR was performed with the QuantiTect SYBR Green PCR kit (Clontech) using first strand cDNAs as templates. The PCR primers for *PvSWN1* are 5′-ggccgcgtcgtcagtgagca-3′ and 5′-ctacacgttgtcatcagctccgc-3′, those for *PvSWN2A* are 5′-gtatcaga gacttgcccatgattgaa-3′ and 5′-ctacacgacaagcttgctacag-3′, those for *PvSWN3A* are 5′-tccagctgcagtcgtcatctc-3′ and 5′-ctacatgactaggt cattctgttg-3′, those for *PvSWN4A* are 5′-gcagcagtcatatggcgccgaa-3′ and 5′-tgctgccacatctgttcaag-3′, those for *PvSWN5* are 5′-acaagttctgcgtgcagtgctccga-3′ and 5′-tcattttctcaacagtcagctcc-3′, those for *PvSWN6A* are 5′-ccacactccacccatcagccta-3′ and 5′-ctgtctattttccaccaagttgca-3′, those for *PvSWN7A* are 5′-tcagctcatg gatgactcgagta-3′ and 5′-ctgacactctccaccaagttgca-3′, those for *PvSWN8A* are 5′-ggccgttgagctgcgtacatga-3′ and 5′-ctatattctccaccaagttgca-3′, and those for *PvMYB46A* are 5′-aacaagttcgtcctgagaattgca-3′. The PCR primers for Arabidopsis secondary wall-associated transcription factors and secondary wall biosynthetic genes were according to Zhong et al. (2006). The relative expression level of each gene was calculated by normalizing the PCR threshold cycle number of each gene with that of an actin reference gene from switchgrass or the *EF1α* reference gene from Arabidopsis. The data were the average of three biological replicates.
In situ hybridization

Elongating switchgrass stems were fixed in 2.5% formaldehyde and 0.5% glutaraldehyde, embedded in paraffin, and sectioned (12 μm thick) according to McAbee et al. [42] and Zhou et al. [43]. The 250-bp 3' untranslated sequences of PvSWN cDNAs were used for synthesis of digoxigenin-labeled antisense and sense RNA probes with the DIG RNA labeling mix (Roche). The primers used for PCR-amplification of the 250-bp 3' untranslated sequences are 5’ - cacctcaacggcagggcggcga-3' and 5’ - ggccggcaggtcatacagcgacact-3’ for PvSWN2A, 5’ - tgtgactaatataaaccaaatatcct-3’ and 5’ - tgtgactaagag catctgagttct-3’ for PvSWN7A, 5’ - cttgctgaagaacctacaggaacca-3’ and 5’ - tacctgtttatgtgacataagt-3’ for PvSWN8A, and 5’ - tgtggagggc caccactcagtctgct-3’ and 5’ - agccagattttatgatgattatt-3’ for PvMYB46A. Stem sections were hybridized with the antisense and sense probes and the hybridization signals were detected by incubation with alkaline phosphatase-conjugated antibodies against digoxigenin and subsequent color development with alkaline phosphatase substrates.

Overexpression of PvSWNs and PvMYB46A

The full-length cDNAs of PvSWNs and PvMYB46A were ligated downstream of the CaMV 35S promoter in pBI121 to create overexpression constructs. The constructs were introduced into wild-type Arabidopsis thaliana (ecotype Columbia) by agrobacterium-mediated transformation. For each construct, at least 30 transgenic Arabidopsis plants were generated and used for morphological and histological analyses.

Complementation of Arabidopsis mutants

The full-length cDNAs of PvSWNs driven by the 3-kb SND1 promoter were cloned into the pGPTV-HPT vector and introduced into the Arabidopsis snd1 nst1 double mutant [14]. For each construct, at least 30 transgenic plants were generated and 8 of them were examined for stem strength and restoration of secondary walls in fibers. For stem strength analysis, the basal parts of the main inflorescence of 7-week-old plants were measured for breaking force using a digital force/length tester [44]. The breaking force was calculated as the force needed to break apart a stem segment.

The full-length cDNA of PvMYB46A driven by the 3-kb MYB46 promoter was cloned into the pGPTV-HPT vector and introduced into the myb46 (-/-; homozygous) myb83 (+/-; heterozygous) double mutant [23]. Transgenic plants were screened for double homozygous myb46 (-/-) myb83 (-/-) mutants by PCR-based genotyping as described previously [23]. At least 10 transgenic plants with the myb46 (-/-) myb83 (-/-) background were examined for plant growth and vessel morphology phenotypes.

Histology

Tissues were fixed in 2% formaldehyde and embedded in Low Viscosity (Spurr’s) resin (Electron Microscopy Sciences) and cut into thin sections with a microtome as described [45]. One-μm-thick sections were stained with toluidine blue for examination of cell wall morphology using light microscopy. Visualization of lignin in the stems were done by staining 50-μm-thick sections with phloroglucinol-HCl, which stains lignin bright red. For lignin autofluorescence visualization, leaves were cleared in methanol and examined using a UV fluorescence microscope [13]. Visualization of secondary wall cellulose was achieved by incubating 1-μm-thick sections with 0.01% Calcofluor White [46]. The fluorescence signals for cellulose staining were
detected with excitation at 360 nm and emission at 480 nm under a confocal microscope. No bright white signals were observed in sections without Calcofluor White staining under the same detection conditions. For immunostaining of xylan, 1-μm-thick sections were first incubated with the monoclonal LM10 antibody and then with fluorescein isothiocyanate-conjugated goat-anti-rat secondary antibodies according to McCartney et al. [47]. The fluorescence signals for xylan immunostaining were detected with excitation at 488 nm and emission at 600 nm under a confocal microscope. The control sections were omitted with the LM10 antibody and no fluorescence signals were observed under the same detection conditions.

Transactivation analysis

The reporter and the effector constructs were co-transfected into Arabidopsis leaf protoplasts [48]. A construct containing the firefly luciferase gene driven by the CaMV 35S promoter was included in each transfection for determination of the transfection efficiency. After 20-hr incubation, protoplasts were lysed and the supernatants were assayed for GUS and luciferase activities. The GUS activity was normalized against the luciferase activity in each transfection, and the data are the average of three biological replicates.

Statistical analysis

The experimental data of the quantitative PCR analysis and GUS activity assay were subjected to statistical analysis using the Student’s t test program (http://www.graphpad.com/quickcalcs/ttest1.cfm), and the quantitative difference between the two groups of data for comparison in each experiment was found to be statistically significant (p < 0.001).

Accession numbers

The gene identification numbers (Panicum virgatum v1.1) and the GenBank accession numbers for the genes used in this study are PvSWN1 (Pavir.J07835; KT075080), PvSWN2A (Pavir.J20698; KT075081), PvSWN2B (Pavir.J21162; KT075082), PvSWN3A (Pavir.J20890; KT075083), PvSWN3B (Pavir.J31179; KT075084), PvSWN4A (Pavir.Ib02477; KT075085), PvSWN4B (Pavir.Ia02924; KT075086), PvSWN5 (Pavir.Ib00161; KT075087), PvSWN6A (Pavir.J07126; KT075088), PvSWN6B (Pavir.Gb00744; KT075089), PvSWN7A (Pavir.J26987; KT075090), PvSWN7B (Pavir.Da02426; KT075091), PvSWN8A (Pavir.J09314; KT075092), PvSWN8B (Pavir.J39804; KT075093), PvMYB46A (Pavir.J11191; KT075094), and PvMYB46B (Pavir.Ca02370; KT075095). PvSWN1, PvSWN2A, PvSWN2B and PvSWN5 were previously described as PvNAC055, PvNAC101, PvNAC61/62 and PvNAC068, respectively [49]. PvMYB46A and PvMYB46B were previously described as AP13ISTG55477 and AP13ISTG55479, respectively [50].

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Author Contributions

Conceived and designed the experiments: RZ YY JJS JTG JCE ZHY. Performed the experiments: RZ YY JJS JTG JCE ZHY. Analyzed the data: RZ YY JJS JTG JCE ZHY. Contributed reagents/materials/analysis tools: RZ YY JJS JTG JCE ZHY. Wrote the paper: RZ YY JJS JTG JCE ZHY.
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