Transcriptomic, Protein-DNA Interaction, and Metabolomic Studies of VosA, VelB, and WetA in Aspergillus nidulans Asexual Spores

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ABSTRACT In filamentous fungi, asexual development involves cellular differentiation and metabolic remodeling leading to the formation of intact asexual spores. The development of asexual spores (conidia) in Aspergillus is precisely coordinated by multiple transcription factors (TFs), including VosA, VelB, and WetA. Notably, these three TFs are essential for the structural and metabolic integrity, i.e., proper maturation, of conidia in the model fungus Aspergillus nidulans. To gain mechanistic insight into the complex regulatory and interdependent roles of these TFs in asexual sporogenesis, we carried out multi-omics studies on the transcriptome, protein-DNA interactions, and primary and secondary metabolism employing A. nidulans conidia. RNA sequencing and chromatin immunoprecipitation sequencing analyses have revealed that the three TFs directly or indirectly regulate the expression of genes associated with heterotrimeric G-protein signal transduction, mitogen-activated protein (MAP) kinases, spore wall formation and structural integrity, asexual development, and primary/secondary metabolism. In addition, metabolomics analyses of wild-type and individual mutant conidia indicate that these three TFs regulate a diverse array of primary metabolites, including those in the tricarboxylic acid (TCA) cycle, certain amino acids, and trehalose, and secondary metabolites such as sterigmatocystin, emericellamide, austinol, and dehydroaustinol. In summary, WetA, VosA, and VelB play interdependent, overlapping, and distinct roles in governing morphological development and primary/secondary metabolic remodeling in Aspergillus conidia, leading to the production of vital conidia suitable for fungal proliferation and dissemination.

IMPORTANCE Filamentous fungi produce a vast number of asexual spores that act as efficient propagules. Due to their infectious and/or allergenic nature, fungal spores affect our daily life. Aspergillus species produce asexual spores called conidia; their formation involves morphological development and metabolic changes, and the associated regulatory systems are coordinated by multiple transcription factors (TFs). To understand the underlying global regulatory programs and cellular outcomes associated with conidium formation, genomic and metabolomic analyses were performed in the model fungus Aspergillus nidulans. Our results show that the fungus-specific WetA/VosA/
VelB TFs govern the coordination of morphological and chemical developments during sporogenesis. The results of this study provide insights into the interdependent, overlapping, or distinct genetic regulatory networks necessary to produce intact asexual spores. The findings are relevant for other *Aspergillus* species such as the major human pathogen *Aspergillus fumigatus* and the aflatoxin producer *Aspergillus flavus*.

**KEYWORDS** sporulation, asexual development, velvet, WetA, secondary metabolites, *Aspergillus*, transcription factor, genetic regulatory network

Fungal asexual spores are key reproductive cells that are essential for the long-term survival of filamentous fungi under a variety of environmental conditions (1). These spores can easily disperse into various environmental niches and act as infectious units for some pathogenic fungi (2–4). Asexual development in *Aspergillus* involves the formation of multicellular structures called conidiophores, each bearing hundreds of asexual spores called conidia. The production of intact conidia (conidiation) requires highly specialized cellular and structural differentiation and metabolic remodeling, which is governed by the coordinated activities of multiple positive and negative regulators (5, 6). Current knowledge about conidiogenesis is derived from numerous studies in model filamentous fungi such as *Aspergillus nidulans* (7–10).

The entire process of conidiogenesis is regulated by distinct gene sets, including central, upstream, and feedback regulators (6, 11). These components are highly conserved in *Aspergillus* species (12). In order to initiate conidiation, upstream developmental activators (FluG and FlbA, FlbB, FlbC, FlbD, and FlbE) induce the activation of *brlA*, an essential initiator of conidiation (13). This occurs when the fungal cells have acquired developmental competence that involves the removal of repressive effects imposed by the key negative regulators SfgA, NsdD, and VosA (14–16). Upon the activation of *BrlA*, it turns on AbaA and WetA, and together they sequentially control the conidiation-specific genetic regulatory networks, thereby governing the formation of conidiophores consisting of aerial stalks, vesicles, metulae, phialides, and conidia (9, 17). These three regulators are considered to form the central regulatory pathway (*BrlA* → *AbaA* → *WetA*) in *Aspergillus* species (18). *BrlA* is a key transcription factor (TF) that activates the expression of *abaA* and other genes in the early stage of conidiation (19, 20). *AbaA* is a TEF1 (transcriptional enhancer factor 1) family TF governing the expression of certain genes such as *wetA*, *vosA*, *velB*, and *rodA* in the metulae and phialides (21–23). *WetA* plays an important role in conidial wall integrity and conidial maturation during the late phase of conidiogenesis (24, 25). Our recent studies have shown that *WetA* functions as a DNA-binding protein that regulates spore-specific gene expression (25, 26). Along with *WetA*, two velvet regulators, *VosA* and *VelB*, which are fungus-specific TFs, coordinate morphological, structural, and chemical developments and exert feedback control of *BrlA* in conidia (27–30).

Previous studies have found that single-knockout mutants of *vosA*, *velB*, and *wetA* share multiple conidial phenotypes, including reduced spore viability, impaired trehalose biosynthesis, defective cell wall integrity, and reduced stress tolerance (25, 31, 32). The mRNA levels of these three regulators are high in wild-type (WT) conidia (25, 27, 28, 33). Results of chromatin immunoprecipitation (ChIP) analyses have demonstrated that *VosA* and *WetA* recognize certain DNA sequences in the promoter regions of target genes and regulate the mRNA expression of spore-specific genes in asexual spores (25, 29). In addition, the deletion of *vosA* or *wetA* affects the mRNA levels of multiple secondary metabolite cluster genes (25, 30, 34). Biochemical studies have determined that *VosA* interacts with *VelB* in conidia, and this complex controls trehalose and β-glucan biosynthesis (30, 35). Importantly, the roles of these three TFs are conserved in *Aspergillus* species (36–39). Considered jointly, these results suggest that *VosA*, *VelB*, and *WetA* are key TFs that orchestrate spore-specific gene expression in *A. nidulans*. Although the role of each regulator has been studied, the regulatory networks between these proteins have not, to date, been investigated in detail. In addition, the effects of these three proteins on primary and secondary metabolism are yet to be elucidated.
In this study, we aimed to determine the cross-regulatory mechanisms of VosA/ VelB/WetA in fungal conidiation using comparative transcriptomic and metabolomic analyses of WT and null mutants of wetA, velB, and vosA in A. nidulans conidia. In addition, the direct targets of these regulators were identified by combining the results from the VosA- and VelB-chromatin interactions using ChIP sequencing (ChIP-seq) analysis with WetA direct targets identified in a previous study (25). The results clarify the detailed molecular mechanisms by which VosA/VelB and WetA control defined common and distinct regulons and increase the overall understanding of the regulatory networks that govern fungal cell differentiation and metabolism.

RESULTS

VosA-, VelB-, and WetA-mediated gene regulation in A. nidulans conidia. To understand the conserved and divergent regulatory roles of VosA, VelB, and WetA in A. nidulans conidia, a comparative analysis of gene expression differences between the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1). Totals of 40.98% (4,503/10,988), 45.61% (5,012/10,988), and 51.96% (5,729/10,988) of genes of the A. nidulans genome are differentially regulated in the WT and null mutant conidia was carried out (Fig. 1).
experiments followed by high-throughput sequencing of the enriched DNA fragments were carried out. ChIPs from strains containing FLAG epitope-tagged versions of VosA and VelB were compared to ChIPs from WT conidia that did not contain the FLAG epitope. Totals of 1,734 and 655 genes that were VosA and VelB peak associated,

**FIG 1** Genome-wide analyses of the genes differentially affected by VosA, VelB, and WetA in A. nidulans conidia. (A) Venn diagram showing the genes whose mRNA levels are downregulated (left) or upregulated (right) by the absence of VosA, VelB, or WetA in conidia. (B) Gene Ontology (GO) term enrichment analysis of downregulated (left) or upregulated (right) genes in the ΔvosA, ΔvelB, and ΔwetA conidia.
respectively, were identified using the same analysis pipeline as the one described previously (25). To identify the VosA/VelB response elements, DNA sequences in the 100 bp surrounding each peak were subjected to Multiple Em for Motif Elicitation (MEME) analysis, which led to the predicted VosA response element (VoRE) and the predicted VelB response element (VbRE) (Fig. 2A). Interestingly, the predicted VbRE (5’-CCXTGG-3’) was quite similar to the predicted VoRE (5’-CCXXGG-3’). The VoRE was found in 278/1,404 peak sequences, had an E value of 1.6e-256, and was the only motif identified by MEME with an E value of < 1. The VbRE was found in 188/511 peak sequences, had an E value of 2.4e-285, and was one of only two motifs identified by MEME with an E value of < 1 (the other motif had an E value of 4.0e-5 and was found in only 72 peak sequences).

We then compared the results of the ChIP-seq and RNA sequencing (RNA-seq) analyses to identify potential direct target genes of the three TFs (Table S3). There were 66 genes associated with the peaks of all three TFs (Table S4). Among them, 22 genes, including \(fb\)A, \(xg\)E, \(at\)B, \(tps\)A, \(vad\)A, \(cet\)A, \(nop\)A, and \(pps\)A, were DEGs in all three null mutants (Fig. 2B). Importantly, 532 genes were considered to be potential direct target genes of VosA, VelB, and WetA.
genes for both VosA and VelB but not WetA. A total of 166 genes were upregulated in both $\Delta$vosA and $\Delta$velB mutant conidia. These genes, including brlA, fadA, rosA, steA, steC, and veA, were found to be involved primarily in asexual or sexual developmental processes. Taking these results together with the previously reported results (27, 35), we suggest that VosA works with VelB and that the VosA-VelB complex coordinates the processes involved in conidial production and maturation in $A.$ nidulans.

**Roles of VosA, VelB, and/or WetA in conidial wall integrity.** Previous studies have shown that the deletion of vosA, velB, or wetA leads to decreased amounts of trehalose and increased $\beta$-glucan levels in conidia (25, 30). The results of transmission electron microscopy analyses revealed that three TFs are needed for the proper formation of the conidial wall (25, 30, 37), suggesting that these genes play a conserved role in regulating the expression of genes associated with conidial structural integrity. High-performance liquid chromatography (HPLC) analysis demonstrated that the trehalose contents of the three null mutant conidia were dramatically decreased (Fig. 3A). The mRNA expression levels of most genes involved in trehalose biosynthesis were down-regulated (Fig. 3B and Table S5). In addition, tpsA, a putative trehalose synthase gene, is the direct target of three TFs (Fig. 2B). These results suggest that three TFs directly or indirectly control the mRNA levels of genes associated with trehalose biosynthesis, thereby regulating the trehalose contents in conidia. Most genes associated with chitin and $\beta$-(1,3)-glucan biosynthesis were upregulated in the $\Delta$vosA, $\Delta$velB, and $\Delta$wetA mutant conidia (Fig. 3C and D). These results suggest that VosA, VelB, and WetA govern the mRNA expression of genes associated with conidial wall integrity in $A.$ nidulans.
Alterations to primary metabolites in ΔvosA, ΔvelB, and ΔwetA conidia. As mentioned above, the deletion of vosA, velB, or wetA led to alterations in the mRNA expression of genes involved in metabolic processes (glycerol metabolic process, ketone metabolic process, and amino sugar metabolic process) and amino acid metabolism (Table S6), implying that the amounts of primary metabolites may be affected by the absence of vosA, velB, or wetA in conidia. To test this hypothesis, the abundances of several primary metabolites involved in the tricarboxylic acid (TCA) cycle and amino acid biosynthesis were examined in WT and mutant conidia (Fig. 4). The abundances of pyruvate, α-ketoglutarate, and malate were increased in the conidia of the three null mutants. The abundances of acetyl-CoA and succinate were decreased in both ΔvosA and ΔvelB, but not ΔwetA, mutant conidia. The amounts of lactate in both ΔvosA and ΔvelB mutant conidia were significantly large compared with those in WT conidia.

The abundances of 13 amino acids (alanine, isoleucine, methionine, leucine, phenylalanine, tryptophan, valine, threonine, serine, asparagine, glutamine, aspartate, and glutamate) were affected in at least one null mutant. Moreover, the levels of nine amino acids were high in all three mutant conidia. The effects of deleting vosA-velB or wetA on the abundances of glutamate, glutamine, aspartate, and asparagine differed. The deletion of wetA caused decreased levels of glutamate, glutamine, and asparagine in conidia, whereas the levels of these amino acids were increased or not affected by the absence of vosA or velB. The genes involved in the biosynthesis of these amino acids and primary metabolites were differentially regulated in the three null mutants. Overall, these results show that WetA, VosA, and VelB regulate the expression of genes involved in both the TCA cycle and amino acid biosynthesis; however, the three lists of primary metabolites that they affect contain both shared and unique molecules.

Abundances of secondary metabolites in ΔvosA, ΔvelB, and ΔwetA conidia. Previous studies found that these three TFs are important for the production of several
secondary metabolites in *Aspergillus* species (34, 36, 40). In addition, according to the GO analysis results, the deletion of vosA, velB, or wetA results in an alteration of the mRNA expression of biosynthetic gene clusters involved in the production of multiple secondary metabolites, including monodictyphenone, sterigmatocystin, and asperfurane (Fig. 1, Fig. S2, and Table S7). To elucidate the conserved and divergent regulatory effects of secondary metabolism in the three conidial mutants, the secondary metabolites were extracted and subjected to liquid chromatography-mass spectrometry (LC-MS) analysis. A principal-component analysis showed differences between the four different conidial samples (Fig. S3). The secondary metabolite content of the WT conidia was relatively similar to that of the ΔwetA conidia, indicating similar abundances and types of secondary metabolites. Conidia from the ΔvosA and ΔvelB mutants clustered far apart, which suggested that a unique set of secondary metabolites or different levels of metabolites were expressed and extracted. This is interesting considering that the two TFs can interact and that their binding motifs and regulated gene lists were so similar to one another (Fig. 1A and Fig. 2A).

Next, we applied analysis of variance to identify the most different molecular entities detected as mass/charge (m/z) values and retention time (RT) pairs in the LC-MS analysis-derived metabolomics data. As shown in Fig. 5, the abundances of several secondary metabolites were different in the positive and negative ionization modes. For example, the abundance of arugosin A was high in the ΔwetA conidia, compared with the WT conidia, but not in the ΔvosA and ΔvelB mutant conidia. However, the amounts of sterigmatocystin intermediates were different in ΔvosA and ΔwetA conidia. Levels of norsolorinic acid and nidurufin were low in the ΔvelB and ΔwetA conidia, while the level of versiconol was

![FIG 5](levels_of_secondary_metabolites_in_the_ΔvosA,_ΔvelB,_and_ΔwetA_conidia. Differentially regulated secondary metabolites in WT, ΔvosA, ΔvelB, and ΔwetA conidia are shown. The heat map is color-coded and represents high abundances (red) or low abundances (blue) of ions/retention time pairs detected by LC-MS analysis.)
Regulation of key secondary metabolites in ΔvosA, ΔvelB, and ΔwetA conidia of A. nidulans. (A, top) Chemical structures of the compounds. (Middle) Abundances of norsolorinic acid, nidurufin, versiconol, and sterigmatocystin in WT, ΔvosA, ΔvelB, and ΔwetA conidia. (Bottom) The sterigmatocystin gene cluster and differentially expressed genes involved in sterigmatocystin biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia. (B, top) Abundance of emericellamide in WT, ΔvosA, ΔvelB, and ΔwetA conidia with the emericellamide A structure. (Bottom) The emericellamide gene cluster and mRNA expression of genes associated with emericellamide biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia. (C, left) Abundances of austinol and dehydroaustinol in WT, ΔvosA, ΔvelB, and ΔwetA conidia with their structures. (Right) The austinol gene cluster and mRNA expression of genes associated with austinol biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia.

FIG 6 Regulation of key secondary metabolites in ΔvosA, ΔvelB, and ΔwetA conidia of A. nidulans. (A, top) Chemical structures of the compounds. (Middle) Abundances of norsolorinic acid, nidurufin, versiconol, and sterigmatocystin in WT, ΔvosA, ΔvelB, and ΔwetA conidia. (Bottom) The sterigmatocystin gene cluster and differentially expressed genes involved in sterigmatocystin biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia. (B, top) Abundance of emericellamide in WT, ΔvosA, ΔvelB, and ΔwetA conidia with the emericellamide A structure. (Bottom) The emericellamide gene cluster and mRNA expression of genes associated with emericellamide biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia. (C, left) Abundances of austinol and dehydroaustinol in WT, ΔvosA, ΔvelB, and ΔwetA conidia with their structures. (Right) The austinol gene cluster and mRNA expression of genes associated with austinol biosynthesis in ΔvosA, ΔvelB, and ΔwetA conidia.
high only in the ΔvelB conidia. The RNA-seq results indicated that the mRNA levels of almost all of the genes in the sterigmatocystin gene cluster were increased in both the ΔvosA and ΔwetA conidia, whereas the mRNA expression of these genes in the ΔvelB conidia was less consistent. In particular, the mRNA levels of stcL, stcN, stcQ, stcS, stcT, stcU, stcV, and stcW were decreased in the ΔvelB conidia compared with the WT conidia. These results suggest that VosA and VelB play diverse roles in the regulation of sterigmatocystin biosynthesis.

Emericellamide compounds are cyclopeptides that are produced by several Aspergillus species (43, 44). The abundances of these compounds, relative to WT production, were high in ΔvosA and ΔvelB conidia, and the mRNA levels of easA, easB, easC, and easD were also high in both mutant conidia, implying that VosA and VelB repress emericellamide biosynthesis in WT conidia (Fig. 6B). In the ΔwetA conidia, however, the mRNA expression of the emericellamide gene cluster was increased, but the quantity of emericellamide compounds did not increase, suggesting that the regulatory mechanism of emericellamide biosynthesis in the ΔwetA conidia is more complex than the influence of ΔvosA and ΔvelB on emericellamide production in conidia. In the three types of null mutant conidia, the abundances of two fungal meroterpenoids, austino1 and dehydroausti1nol (45), were decreased, compared with the WT conidia (Fig. 6C). Furthermore, the expression levels of several austinol cluster genes were decreased in the ΔvelB and ΔwetA conidia. Taken together, these results demonstrate that the ways in which VosA, VelB, and WetA govern the expression of secondary metabolite gene clusters, and the production of their associated metabolites, in A. nidulans conidia are divergent from one another.

**DISCUSSION**

Asexual developmental processes in filamentous fungi are regulated by a variety of TFs (6). These TFs orchestrate the spatial and temporal transcriptional expression of development-specific genes, leading to physiological and metabolic changes. During the processes of conidium formation from phialides and conidial maturation, conidium-specific TFs, including VosA, VelB, and WetA, regulate spore-specific gene expression patterns and metabolic changes (25, 30). In this study, we investigated the transcript and metabolite changes that are regulated by VosA, VelB, and WetA in A. nidulans conidia.

Transcriptomic analyses indicated that about 20% of the A. nidulans genome (2,143 genes) is differentially expressed in ΔvosA, ΔvelB, and ΔwetA mutant conidia. ChIP-seq results identified 66 direct target genes that are shared between VosA, VelB, and WetA in conidia. These results offered some explanation of how these TFs control phenotypic changes in conidia. First, the deletion of vosA, velB, or wetA caused increased mRNA expression of certain development-specific genes, including abaA (23), brlA (19), flbA (46), flbC (47), nsdC (48), nosA (49), and mpkB (50), which are involved in the formation of asexual and sexual structures during the early and middle stages of conidium formation, but decreased transcript accumulation of spore-specific genes such as vadA (51), catA (52), wa (53), conF (54), conJ (54), cetA (55), cetJ (56), and cetL (56), which are important for conidial germination, morphogenesis, and dormancy (see Table S1 in the supplemental material). Alteration of the mRNA expression levels of development-specific genes or spore-specific genes affect spore maturation, dormancy, and germination. For example, misscheduled expression of key asexual developmental regulators, especially BrlA and AbaA, can affect proper sporulation (9, 57). In the case of the spore-specific genes, the deletion of vadA or catA affects conidial germination and the conidial stress response (51, 52, 58). Based on these results, we propose that alteration of the mRNA expression levels of development-specific genes or spore-specific genes caused by the deletion of vosA, velB, or wetA affect conidial maturation, dormancy, morphology, and germination. However, the detailed molecular mechanism of how these TFs act as activators or repressors for the expression of development-specific genes and spore-specific genes will be elucidated in further studies.
Another important phenotype of the ΔvosA, ΔvelB, and ΔwetA mutant conidia was the differences in conidial wall integrity and the components of the conidial wall (25, 30). As shown in Fig. 3, most of the genes involved in chitin and β-glucan biosynthesis were upregulated in all three mutant conidia. The dynamic expression of these genes is required mainly for the remodeling of the cell wall during isotropic growth and mobilization of energy for differentiation (59) but is not required in dormant conidia. However, by altering the mRNA expression of these genes in the mutant conidia, the dormancy of conidia could be broken, affecting long-term viability as well as conidial germination.

Another feature of fungal spores is their ability to resist various environmental stresses (1). However, ΔvosA, ΔvelB, and ΔwetA mutant conidia are more sensitive to several environmental stresses (25, 35). It is speculated that this is regulated by alterations in the expression of genes involved in environmental stress tolerance. The data that we show here support this hypothesis. First, these regulators govern the mRNA expression of genes involved in the trehalose biosynthetic pathway, thereby affecting the amount of conidial trehalose, a key component in stress protection and fungal virulence (60). Second, VosA, VelB, and WetA directly or indirectly regulate genes previously associated with stress responses. CatA is a spore-specific catalase, and compared with WT spores, catA deletion mutant spores are sensitive to oxidative stress (52). AtfB is a bZIP TF (61), and the AtfB homolog is crucial for the stress response in Aspergillus oryzae conidia (62). These two genes are putative direct target genes of the three regulators reported in this study, and the mRNAs of catA and atfB can be positively regulated by VosA, VelB, and WetA in conidia (Fig. 2 and Table S3). Along with these genes, the mRNA level of hogaA, a key component of osmotic stress signaling (63), was downregulated in all mutant conidia. These results contribute to our understanding of the ways in which these three regulators influence the environmental stress response in conidia.

VosA, VelB, and WetA are key functional regulators in the formation of conidia and control spore-specific gene expression. However, our data have shown that their gene regulation networks are slightly different. RNA-seq results showed that VosA and VelB coregulate the expression of spore-specific genes. Importantly, the predicted VbRE is quite similar to the predicted VoRE (Fig. 2A). In addition, biochemical results from previous studies (27, 35) suggested that VosA and VelB form a heterocomplex in asexual spores. However, WetA is not directly related to VosA and VelB. WetA’s putative binding site is different from the VosA/VelB binding site. Moreover, the WetA peak-associated genes and the VosA/VelB peak-associated genes did not overlap much. These results imply that WetA-mediated gene regulation may be different from the VosA- or VelB-mediated gene regulatory network.

The velvet domain is a fungus-specific DNA-binding domain that recognizes specific DNA sequences. Previously, Ahmed et al. proposed that the VosA velvet domain recognizes a DNA sequence (5′-TGGCCGCGG-3′) based on ChIP-chip analysis and electrophoretic mobility shift assays (EMSAs) (29). Further EMSAs demonstrated that both TGG and CCGCGG sequences are necessary for DNA binding of the VosA velvet domain. In the present study, we conducted ChIP-seq analyses in conidia and proposed the predicted VbRE (5′-CCXTGG-3′) and VoRE (5′-CCXXGG-3′) (Fig. 2). In our experimental results, the TGG sequence does not appear for the VbRE or VoRE, but the 5′-CCXXGG-3′ sequence is conserved in the VbRE and VoRE. The reason why these DNA sequences are not the same is likely because the experimental methods and analyses are different from those used to obtain the previous results. Ahmed et al. used 15 DNA sequences based on chromatin immunoprecipitation with microarray technology (ChIP-chip) analysis and EMSAs, whereas the motif in Fig. 2A was built from running MEME with every peak sequence that we identified. Nevertheless, the 5′-CCXXGG-3′ sequence appears common in previous and current results. Based on these data, we propose that the 5′-CCXXGG-3′ sequence may be crucial for DNA binding of the velvet
domain, and further studies will be needed to fine-tune the precise velvet protein-binding sequence.

During the asexual development of A. nidulans, the abundance of amino acids other than phenylalanine changes, and the expression of genes related to amino acid biosynthesis is altered (64). Overall, our analyses confirmed that the amounts of most amino acids, and the expression of related genes, increased in all mutant spores. In addition, the abundances of metabolites involved in the TCA cycle increased in all mutant conidia. However, the abundances of some primary metabolites such as glutamate, glutamic acid, lactate, and acetyl-CoA were decreased in the ∆wetA conidia (Fig. 4). It is not yet clear how these metabolic changes affect spore production and maturation, and further studies will be needed to understand this.

Our multi-omics analyses found that VosA, VelB, and WetA regulate the expression of several secondary metabolite gene clusters (Table S7) and the production of secondary metabolites, especially sterigmatocystin, in conidia. The process of sterigmatocystin production and its regulation involves 25 genes, and this metabolite is produced via steps involving several intermediate products. In ∆vosA conidia, the mRNA expression of sterigmatocystin gene clusters was induced, and the amounts of sterigmatocystin produced were similar to those in the WT conidia. These results were similarly observed in sexual spores (34). While the ∆vosA conidia contained sterigmatocystin, the metabolite was not detected in ∆velB conidia. We reported that the VosA-VelB complex is a functional unit in conidia, but this particular result indicates that VosA and VelB play different roles in sterigmatocystin production. It is possible that VelB forms another complex, such as the VelB-VeA-LaeA complex (40), to participate in sterigmatocystin production in conidia. For the ∆velB conidia, we speculated that the mRNA expression levels of genes such as stcB, stcC, stcF, and stcI, which are associated with the early stages of sterigmatocystin biosynthesis, were increased, and that the amount of versiconol, a putative sterigmatocystin/aflatoxin intermediate, was also increased in comparison with the WT. However, the mRNA levels of genes associated with the late phase of sterigmatocystin biosynthesis, such as stcl, stcN, stcQ, and stcT, were decreased in ∆velB conidia. It might be possible that VelB (or VelB/VeA/LaeA) can regulate some expression of sterigmatocystin gene clusters by epigenetic means rather than through the canonical method of aflR expression or activity. Although changes in the expression of secondary metabolite gene clusters and secondary metabolites affected by three TFs were studies, detailed molecular mechanisms have not been studied yet. Therefore, it is necessary to study how these three TFs work together or separately through further research. In the ∆vosA and ∆wetA conidia, the mRNA levels of most of the genes in the sterigmatocystin gene cluster were increased compared to those in WT conidia, but the amounts of sterigmatocystin were similar to those in WT conidia. There are some speculations about this phenomenon. The expression of genes may not directly affect the biosynthesis of secondary metabolites. Alternatively, the translation of mRNA molecules to proteins and the posttranslational modification of those metabolite-producing proteins are two factors that can create discrepancies between RNA and metabolite abundances. To further explain this, further experiments should be conducted to determine how the three TFs regulate the biosynthesis of secondary metabolites.

In conclusion, this study provides a systematic dissection of the gene regulatory network and molecular mechanisms of VosA, VelB, and WetA (Fig. 7). In conidia, VosA, VelB, and WetA directly or indirectly control the expression of spore-specific or development-specific genes, thereby altering conidial wall integrity and conidial viability. In addition, these TFs regulate multiple secondary metabolite gene clusters, thus inducing secondary metabolic changes. These results provide an advance in the knowledge of conidial formation and will provide the basis for future insights into spore formation in other filamentous fungi.

MATERIALS AND METHODS

Strains, media, and culture conditions. The fungal strains used in this study are listed in Table 1. Fungal strains were grown on solid or liquid minimal medium with 1% glucose (MMG) and appropriate
supplements for general purposes as previously described (65). For conidium samples, WT and mutant conidia were inoculated onto solid MMG plates and incubated for 48 h. Next, conidia were collected from plates using Miracloth (Calbiochem, San Diego, CA, USA) and stored at -20°C.

RNA sequencing analysis. To isolate total RNA for RNA sequencing (RNA-seq) analysis, total RNA from WT and mutant conidia was extracted using TRIzol reagent (Invitrogen, USA), according to the manufacturer’s instructions, with modifications. To remove DNA contamination from the RNA samples, DNase I (Promega, USA) was added, and RNA was then purified using an RNasea minikit (Qiagen, USA). Three technical replicates of each sample were analyzed. RNA sequencing was performed as previously described (34). RNA samples were submitted to the University of Wisconsin Gene Expression Center (Madison, WI, USA) for library preparation and sequencing. A strand-specific library was prepared using an Illumina TruSeq strand-specific RNA sample preparation system. The libraries of all the replicates were sequenced using an Illumina HiSeq 2500 system.

Data analysis of the ΔvosA and ΔvelB RNA-seq experiments was performed using the same analysis pipeline as the one previously described for the ΔwetA RNA-seq analysis (25). Reads were mapped to the A. nidulans FGSC4 transcriptome using TopHat2 version 2.1.1 (66) and the parameter "-max-intron-length 4000." On average, 19.9 million reads per sample mapped to the genome, and the number of reads aligning to each gene was counted with HTseq-Count version 0.9.1 (67). DESeq version 1.14.1 (68)

TABLE 1 Aspergillus strains used in this study

| Strain | Relevant genotype | Source or reference |
|--------|-------------------|---------------------|
| FGSC4  | A. nidulans wild type; veA⁺ | FGSC⁺ |
| THS15  | pyrG89; pyroA4; ΔvosA::AfupyG⁺; veA⁺ | 27 |
| THS16  | pyrG89; pyroA4; ΔvelB::AfupyG⁺; veA⁺ | 27 |
| THS20.1| pyrG89; pyroA::velB[p];velB::FLAG; “pyroA`; ΔvelB::AfupyG⁺; veA⁺ | 27 |
| THS28.1| pyrG89; pyroA::vosA[p];vosA::FLAG; “pyroA`; ΔvosA::AfupyG⁺; veA⁺ | 27 |
| TMY4   | pyrG89; pyroA4; ΔwetA::AfupyG⁺; veA⁺ | 25 |

FGSC, Fungal Genetic Stock Center.

*The 3/4 pyroA marker causes targeted integration at the pyroA locus.
was used to determine significantly differentially expressed genes, and genes were considered regulated if they exhibited an adjusted \( P \) value of \(<0.05\) and a \( \log_2 \) fold change either greater than 1 or less than \(-1\).

**Chromatin immunoprecipitation sequencing analysis.** Samples for chromatin immunoprecipitation sequencing (ChIP-seq) analysis were prepared according to methods described previously (29, 30). DNA samples from each strain were extracted using a MAGnify chromatin immunoprecipitation system (Invitrogen, USA) according to the manufacturer's protocol, with modification. Two-day-old conidia from the WT strain or strains containing VosA-FLAG or VelB-FLAG were cross-linked, washed, homogenized with a Mini-Beadbeater 16 instrument (Biospec, USA), sonicated, and separated by centrifugation. The chromatin extracts were incubated with an anti-FLAG antibody–Dynabead complex. Next, samples were eluted from the beads at 55°C using proteinase K. Enriched DNA was purified using DNA purification magnetic beads. DNA samples from each strain were submitted to the University of Wisconsin Gene Expression Center (Madison, WI). Libraries were prepared using a TruSeq CHIP library preparation kit (Illumina, CA). The libraries of all the replicates were sequenced using an Illumina HiSeq 2500 system.

Raw reads were trimmed using Trimmomatic version 0.36 (69) and the parameters “ILLUMINA1CLIP:2:30:10 LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:36.” Trimmed reads were mapped to the *A. nidulans* A4 genome using version 0.7.15 of BWA-MEM (70), and shorter split hits were marked as secondary alignments. Mapped reads with mapping quality (MAPQ) values of \(<1\) as well as unmapped, secondarily aligned, supplemental, and duplicated reads were discarded with SAMtools version 1.6 (71). On average, 2.3 million and 7.2 million reads per sample were used for peak calling in the VosA and VelB experiments, respectively. Mapped reads with mapping quality (MAPQ) values of \(\geq 1\) were aligned, and duplicate peaks were combined with a tolerance of 0.1 and a correlation value of \(\geq 0.99\) using Deseq v2 as well as unmapped, secondarily aligned, supplemental, and duplicated reads were discarded with SAMtools version 1.6 (71). On average, 2.3 million and 7.2 million reads per sample were used for peak calling in the VosA and VelB experiments, respectively. Mapped reads that survived our filter were pooled, and extension sizes were estimated with version 1.15.2 of SPP (72, 73). Peaks were called with MACS2 (74) version 2.1.2 using the extension sizes estimated by SPP, a genome size of 3.9Mb, and the “-nomodel” parameter. Peaks with a fold change of \(\geq 2\) and a \( q \) value of \(<0.001\) were further analyzed. Peak lists were combined from both of the VosA biological replicates, as \(>99\%\) of the peaks from the first replicate were found in the second replicate. Motifs were identified in the 100 bps of sequences surrounding each peak summit using MEME-ChIP (75). Motifs that occurred zero times or once in the sequences around the peaks and that were 4 to 21 nucleotides (nt) long were further analyzed.

**Functional enrichment analysis.** Enriched terms from the GO Biological Process, KEGG, InterPro, and Pfam databases were identified using the tools available at AspGD (76), FungiDB (77), and ShinyGO v0.60 (78). Unless otherwise stated, default settings were used in ShinyGO v0.60. The settings were as follows: database, *Emericella nidulans* STRINGdb; \( P \) value cutoff (FDR), 0.05; number of most significant terms to show, 30.

**Primary metabolite analysis.** WT and \(\Delta wetA\), \(\Delta vosA\), and \(\Delta velB\) mutant conidia were inoculated onto solid MMG plates and incubated for 48 h, and fresh conidia were then harvested using Miracloth with HPLC-grade water. For each sample, \(2 \times 10^6\) conidia were mixed with 500 \(\mu\)l HPLC-grade acetonitrile-methanol-water (40:40:20, vol/vol/vol) and 300 \(\mu\)l beads, homogenized by using the Mini-Beadbeater, and centrifuged. The supernatant was filtered using a 0.45- \(\mu\)m polytetrafluoroethylene (PTFE) Mini-UniPrep filter vial (Agilent), collected, and immediately snap-frozen with liquid nitrogen. The samples were stored at \(-80°C\) until primary metabolite analysis.

The samples were then analyzed as described previously (79, 80). Samples were analyzed using an HPLC-MS system consisting of a Dionex ultrahigh-performance liquid chromatography (UHPLC) instrument coupled by electrospray ionization (ESI) (negative mode) to a hybrid quadrupole–high-resolution mass spectrometer (Q Exactive orbitrap; Thermo Scientific) operated in full-scan mode. Metabolite peaks were identified by their exact mass and matching retention times to those of pure standards (Sigma-Aldrich).

**Secondary metabolite analysis.** The conidia of WT and \(\Delta wetA\), \(\Delta vosA\), and \(\Delta velB\) mutant strains were extracted by adding 1.5 ml of a methanol-acetonitrile (2:1) mixture followed by sonication for 60 min. The suspension was then left overnight before centrifugation at 14,000 rpm for 15 min. The supernatant (1 ml) was removed, filtered, and evaporated to dryness in vacuo. Extracts for the metabolomics analysis were normalized to 10 mg/ml in methanol for LC-MS analysis.

Analytical HPLC was performed using an Agilent 1100 HPLC system equipped with a photodiode array detector. The mobile phase consisted of ultrapure water (mobile phase A) and acetonitrile (mobile phase B) with 0.05% formic acid in each solvent. A gradient method from 10% mobile phase B to 100% mobile phase B in 35 min at a flow rate of 0.8 ml/min was used. The column (Phenomenex Kinetex C18, 5 \(\mu\)m by 150 mm by 4.6 mm) was reequilibrated before each injection, and the column compartment was maintained at 30°C throughout each run. All samples were filtered through a 0.45- \(\mu\)m nylon filter before LC-MS analysis.

Extracts from the WT and mutant conidia were analyzed in duplicate on an Agilent 1100 series LC-MS platform (81, 82). The negative ionization mode was found to detect the most metabolites. The first 5 min of every run was removed due to a large amount of coeluting, low-molecular-weight, polar metabolites. Data sets were exported from Agilent’s Chemstation software as .netCDF files and imported into MZmine 2.38 (83). Peak picking was performed with established protocols (84), resulting in 123 marker ions. Briefly, mass detection was centroid with a Sez minimum height. Chromatogram building was limited to peaks greater than 0.1 min with a tolerance of 0.05 \( m/z \) and a minimum height of 1e3. Data smoothing was performed at a filter width of 5. Chromatogram deconvolution was performed by utilizing a local minimum search with a chromatographic threshold of 95%, a minimum relative height of 10%, a minimum absolute height of 3e3, a minimum ratio of peak to edge of 1, and a peak duration range of 0.1 to 5.0 min. The spectra were de-isotoped with a 1-ppm \( m/z \) tolerance before all treatments were aligned, and duplicate peaks were combined with a tolerance of 0.1 \( m/z \) and a 3.0-min RT. Peak finder gap filling was performed with 50% intensity tolerance and 0.1 \( m/z \) tolerance. Peak lists were
exported to Metaboanalyst (85), where missing values were replaced with half the minimum positive value, data were filtered by the interquartile range, and log transformation of the data was employed. **Data availability.** All RNA-seq and ChIP-seq data files are available from the NCBI Gene Expression Omnibus database (wetA RNA-seq, accession number GSE114143; vosA and velB RNA-seq, accession number GSE154639; WetA ChIP-seq, accession number GSE114141; VosA and VelB ChIP-seq, accession number GSE154630).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 0.5 MB.

**FIG S2**, TIF file, 0.5 MB.

**FIG S3**, TIF file, 0.3 MB.

**TABLE S1**, XLSX file, 0.02 MB.

**TABLE S2**, XLSX file, 0.02 MB.

**TABLE S3**, XLSX file, 0.1 MB.

**TABLE S4**, XLSX file, 0.01 MB.

**TABLE S5**, XLSX file, 0.02 MB.

**TABLE S6**, XLSX file, 0.02 MB.

**TABLE S7**, XLSX file, 0.04 MB.

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