A Homeobox Transcription Factor UvHOX2 Regulates Chlamydospore Formation, Conidiogenesis, and Pathogenicity in Ustilaginoidea virens

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Rice false smut fungus (teleomorph: Villosiclava virens; anamorph: Ustilaginoidea virens) can generate chlamydospores and survive winter under field conditions. The chlamydospore is considered as an important infection source of the disease. However, little is known about the regulatory mechanism of the chlamydospore production. In this study, we identified a defective homeobox transcription factor (designated as UvHOX2) gene in a U. virens random insertional mutant B-766 that could not form chlamydospores. To confirm the regulatory function of UvHOX2, an Agrobacterium tumefaciens mediated transformation- and CRISPR/Cas9- based targeted gene replacement method was developed. The UvHox2 deletion mutants completely failed to produce chlamydospores, showed reduced conidia production and decreased virulence, and was hyper-sensitive to oxidative, osmotic, and cell wall stresses. We confirmed that UvHOX2 is located in the nuclei of U. virens, and the expression of UvHox2 was the strongest during the early stage of chlamydospore and conidium formation. Global transcription pattern of UvHOX2 was also determined by RNA-seq in this study, and several genes that might be down-stream of UvHOX2 regulation were identified. The results will better our understanding of the molecular mechanism of chlamydospore formation in U. virens as a model fungus.

Keywords: Villosiclava virens, chlamydospore, conidiation, pathogenicity, homeobox transcription factor

INTRODUCTION

The heterothallic ascomycetous fungus Ustilaginoidea virens (Cooke) Takah (Teleomorph: Villosiclava virens) is the causal agent of rice false smut (RFS), which has become an emerging disease in China and most rice-growing areas in the world (Guo et al., 2012; Nessa et al., 2015; Yu J.J. et al., 2015). The fungus infects stamen filaments of rice at the booting stage, hijacks rice nutrients, and transforms kernels into false smut balls containing a large number of yellow or dark green-pigmented chlamydospores (Hu et al., 2014; Meng et al., 2015; Song et al., 2016; Zheng et al., 2016). Occasionally, sclerotia could form on the surface of false smut balls in late autumn when the temperature falls (Yu et al., 2016; Yong et al., 2018). Previous studies have revealed that chlamydospores could survive in nature and play an important role in the epidemiology of RFS disease between seasons (Fan et al., 2016).
In the human pathogenic yeast *Candida albicans*, several genes were found to be involved in chlamydospore formation, including homeobox transcription factor (TF) gene *grf10* (Ghosh et al., 2015), phosphate mannose synthase encoding genes (Juchimiuk et al., 2015), mitogen-activated protein kinase gene *hgl* (Eisman et al., 2006), gene encoding dolichol phosphate mannose synthase (Juchimiuk et al., 2015), chromatin remodeling complex gene *isw2* (Nobile et al., 2003), MDS3, RIM101, RIM13, SCH9, and SUV3 (Nobile et al., 2003). Meanwhile, very few genes in filamentous fungi were found relative to chlamydospore formation, the limited examples include VELVET gene *vell* in biocontrol fungus *Trichoderma virens* (Mukherjee and Kenerley, 2010); and a group of genes in *Clonostachys rosea* identified from a study using RNA-seq (Sun et al., 2018). So far, the molecular mechanisms of chlamydospore formation remain elusive in filamentous fungi.

Homeobox genes encode a group of TFs, which contain a conserved homeobox domain and bind to specific DNA sequences (Gehring, 1987). In eukaryotic cells, these homeobox TFs play an important role in regulation of cell differential and development (Liu et al., 2010; Antal et al., 2012). The first reported homeobox gene in filamentous ascomycetes is *pah1* in *Podospora anserine* (Arnaise et al., 2001). *Pah1* deletion mutant showed increased production of microconidia and reduced growth rate of mycelia. In model fungus *Neurospora crassa*, three homebox genes were characterized (Colot et al., 2006). Specifically, deletion of *kal1*(*pah1*) homologled to defects in mycelia growth and conidiation; *bek-1* was found to be essential for perithelial development whereas the third homebox gene (Genbank accession number: NCU03070) was not described.

In recent years, several homebox genes were systematically studied in filamentous fungi *Porthe oryzea* and *Podospora anserine*, and the results confirmed that these homebox genes play a regulatory role in conidium and fruiting body development, as well as host infection (Kim et al., 2009; Coppin et al., 2012).

In this study, we identified a chlamydospore formation defect *U. virens* mutant B-766 from a random insertional mutant library that was constructed previously (Yu M.N. et al., 2015). A homebox gene (annotated as *UvHox2*) was confirmed to be involved in the regulation of chlamydospore formation and pathogenicity in *U. virens*. A CRISPR/Cas9 system based on *Agrobacterium tumefaciens* mediated transformation (ATMT) was developed for targeted gene deletion. Moreover, comparative transcriptional analysis of *UvHox2* deletion mutant and a wild-type strain was performed in this study. Taken together, the findings from this work will help us understand the regulatory mechanism of chlamydospore formation better.

**MATERIALS AND METHODS**

**Strains, Rice Variety, Plasmids, and Nucleotide Acids Manipulation**

A virulent wild-type *U. virens* strain P-1 was used as starting strain in this study. A rice variety susceptible to *U. virens*, Liangyoupeijiu, was used in the inoculation experiments. The plasmid pCas9-tRp-gRNA was kindly provided by Dr. Jingrong Xu at Northwest A&F University (Liang et al., 2018). *A. tumefaciens* strain AGL-1, plasmid pmCherry-hph, pCambia-1300, pBH12, pKHT, and pCN3EXPS were from our lab. Southern blot and thermal asymmetric interlaced PCR (TAIL-PCR) were performed as described previously (Yu M.N. et al., 2015).

**Phenotypic Analysis of *U. virens* Strains/Mutants**

The *U. virens* wild-type strain P-1 was routinely cultured on a potato sucrose agar medium (PSA) at 28°C for 10–15 days (Zheng et al., 2017). The transformants of P-1 were cultured on the PSA amended with 100 µg/ml hygromycin and/or 600 µg/ml geneticin 418 (G418). We used YT medium and broth to test mycelial growth rate and conidiation ability of *U. virens*, respectively (Tanaka et al., 2011). To determine the chlamydospore formation and the pathogenicity of *U. virens* strains, we inoculated rice following the method described previously (Zheng et al., 2017). Fifteen spikes were inoculated for each strain, and the number of false smut balls was counted 25 days after the inoculation. The chlamydospore formation structures on the surface of false smut balls were observed by scanning electron microscope (SEM). To stimulate chlamydospore formation in *U. virens*, mycelia dishes cut from the edge of fresh colonies were put on PSA medium. The cultures were incubated at 28°C under diffuse light for 2–3 months.

**Ustilaginoidea virens** strains were cultured on PSA medium to determine the growth rate. YT medium amended with 0.05% H₂O₂, 0.4 mol/l NaCl, 0.03% SDS, and 100 mg/l congo red were used to test sensitivity of stains to abiotic stresses. The cultures were incubated at 28°C for 15 days in darkness, and then the growing diameter was measured and the morphology of the colonies were characterized. Four duplicates were performed for each treatment. The sporulation capacity of the strains was determined as follows. Ten pieces of fresh mycelial dishes from each treatment were cultured in a 250 ml conical flask containing 100 ml YT liquid medium. The conical flasks were incubated at 28°C, 150 r/min for 6 days, and then the thin-wall conidia were counted with a blood cell counting chamber. To observe conidium generation structures, strains were cultured on minimal media (MM) (Gupta and Chattoo, 2008) for 10 days.

**Generation of ATMT Binary Vector for Gene Deletion With CRISPR/Cas9**

**Generation of Gene Deletion Vector With CRISPR/Cas9**

We constructed CRISPR-guideRNA(gRNA) cassettes from pCRISPR-UvrR and gene replacement cassettes [upstream flank (UF)-hygromycin resistant gene(Hyg⁺)-downstream flank (DF)] of *UvHox2* into two T-DNA regions of binary vector pCccd-dTN3, respectively. The details of vectors construction were described in Supplementary Figure S1.

**Generation of Gene Deletion Mutants**

*Agrobacterium tumefaciens* mediated transformation was performed as described previously (Yu M.N. et al., 2015). The
A. tumefaciens strains AGL-1 containing pdTN3-HX2-Cas9I or pdTN3-HX2-Cas9II was employed in transformation of U. virens wild-type strain P-1. The U. virens P-1 and A. tumefaciens AGL-1 were co-cultured on nitrocellulose membrane for 3 days and then transferred onto 2% TB3 [0.3% yeast extract, 0.3% casamino acid, 1% glucose, 2% sucrose (w/v)]. To make a selective medium, 400 µg/ml cefotaxime and 150 µg/ml timentin were added into 2% TB3 medium to inhibit the growth of A. tumefaciens, and 100 µg/ml hygromycin and 600 µg/ml G418 were added into 2% TB3 medium to select transformants containing both cassettes of UF-HYG⁺-DF and CRISPR/Cas9-gRNA, respectively. The UvHox2 deletion mutants were screened and verified by PCR with primers P19∼P26 listed in Supplementary Table S1.

**Generation of UvHox2-eGFP Constructs**

The open reading frame (ORF) of UvHox2 was amplified from cDNA that was generated through reverse transcription of total RNA using the primer pair P27–P28 (Supplementary Table S1). The enhanced green fluorescent protein (eGFP) fragment was amplified with primer pair P29–P30 (Supplementary Table S1). UvHox2-eGFP fusion cassette was generated via double-joint PCR and ligated to BamH I-EcoR I digested pCN3EXPS to construct UvHox2-eGFP fusion vector pCN3EXPS-HX2-eGFP, in which the UvHox2-eGFP cassette was under the control of glyceraldehyde-3-phosphate dehydrogenase promoter of Cochliobolus heterostrophus. Subsequently, the vector was used to transform U. virens via ATMT protocol to generate UvHox2-eGFP over-expression mutants.

**qRT-PCR Assays**

Vegetative mycelia were collected from 2-day-old YT cultures that started with 1 × 10⁶ conidia/ml. To stimulate sporulation in U. virens, mycelial dishes were cultured in YT broth by shaking for 3 days (initial stage of sporulation) or 7 days (later stage of sporulation). To collect samples undergoing chlamydospores formation, 20 rice spikes were inoculated for each strain/mutant. Rice smut balls at the initial stage [yellowish with intact membrane] and the later-stage [yellowish without membrane] of chlamydospore development were collected as described by Fan et al. (2016). PrimeScript™ RT reagent Kit with gDNA Eraser (Takara) and SYBR® Premix Ex Taq™ II (Takara) were used to synthesize cDNA and quantitative RT-PCR (qRT-PCR). Relative expression levels of genes were calculated with the 2^-ΔΔCt method. The α-tubulin gene was employed as the endogenous reference. Three biological replicates were performed to calculate the mean and the standard deviation.

**Comparative Transcriptional Analysis of U. virens**

Total RNA of U. virens was extracted employing TRIZOL (Invitrogen). RNA integrity was determined using Bioanalyzer 2100 RNA-6000 Nano Kit (Agilent Technologies). The
construction and sequencing of mRNA-seq libraries and preprocessing and mapping of Illumina reads were performed as described previously (Yu et al., 2016). The DESeq software (Anders and Huber, 2012) was used to generate base mean based on FPKM, and to evaluate significant differences in base mean between different samples. Three biological replicates were performed for each strain/mutant.

RESULTS

Characterization of Genes Relative to Chlamydospore Formation in Mutant B-766

In a preliminary study, we identified a T-DNA insertional mutant B-766 of *U. virens*, which failed to form chlamydospores on false smut balls (Figure 1). To determine the copy number of T-DNA inserted in B-766, 1.4 kb hygromycin resistant cassette was employed as a probe in southern blot. The result showed that three copies of T-DNA were detected in mutant B766 (Figure 2A). T-DNA flanking regions were amplified by TAIL-PCR (Yu M.N. et al., 2015). Three copies of T-DNA were inserted into the upstream of ORFs that encode proteins KDB15727 (Genbank accession number), KDB15728, KDB14847, KDB14848, and KDB18871 (Figure 2B). We then performed qRT-PCR to screen genes relative to chlamydospore formation in mutant B-766. The expression of KDB14847 in B-766 comparing to P-1 was reduced in a higher level than other genes that might be infected by T-DNA insertion in mutant B-766 (Figure 2C). Because KDB14847 is homologous to homeobox TF MoHOX2 in *Magnaporthe oryzae*, we designated KDB14847 as UvHOX2.

Homeobox TFs in *U. virens*

In eukaryotic cells, homeobox TFs contain a ∼60 aa long conserved homeodomain that binds to specific DNA sequences and regulates transcription (Coppin et al., 2012). We identified seven homeobox TFs in *U. virens* using the InterPro term
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**Figure 3** Phylogenetic analysis of homeobox transcription factors (TFs) in *U. virens*. (A) Positions of homeobox domain in homeobox TFs in *U. virens*. (B) Phylogenetic analysis of homeobox TFs that homologous to UvHOX2 in filamentous fungi. (C) Positions of homeobox domain (red) in UvHOX2 homologs. (D) Sequences of homeobox domain in filamentous fungi.

(IPR001356) and the NCBI assembly data of *U. virens* genome (accession number: GCA_000687475.1). The sequences of the homeobox TFs were deposited in the genbank with the accession number KDB17966, KDB17963.1, KDB17264, KDB14847, KDB13469, KDB13074, and KDB13172 (Figure 3A). The position of the homeodomain varied in the homeobox TFs in *U. virens*. The homeobox motif located at the N-terminus of KDB14847 (UvHOX2), KDB13469, and KDB13172, in the middle of KDB17963 and KDB13074, and at the C-terminus of KDB17966 (Figures 3A–D). Besides the homeobox domain, KDB17963 possessed a rhodanese-like domain at the C-terminus, KDB13172 harbored three zinc finger C2H2 DNA binding domains in the middle and an HTH CenpB-type DNA binding domain at the C-terminus (Figure 3A).

**Generation of UvHOX2 Deletion Mutants**

To delete *UvHox2* in the *U. virens* wild-type strain P-1, we constructed two binary vectors pdTN3-HX2-Cas9I or pdTN3-HX2-Cas9II containing two T-DNA regions, respectively. The first T-DNA inserted with UF-HYG+ RF cassette, and the other one harbors CRIPSR/Cas9-gRNA (gRNA1 or gRNA2) and NEO+ cassettes. The ATMT method was used to co-transfer the two T-DNAs into the *U. virens* wild-type strain P-1. The transformants resistant to hygromycin B and G418 were picked from the selective medium after about 7–10 days culture at 28°C in darkness. For pdTN3-HX2-Cas9I, 9 out of 33 transformants were confirmed as *UvHox2* deletion mutants. For pdTN3- HX2-Cas9II, 10 out of 45 transformants were confirmed as *UvHox2* deletion mutants (Figure 4). The homologous gene replacement frequency was 27.3 and 22.2% for pdTN3-HX2-Cas9I and pdTN3-HX2-Cas9II, respectively. Potential off-target sites were checked using protocol described by Liang et al. (2018), and no off-target event was detected in all transformants.

**UvHox2 Is Essential for Chlamydospore Formation and Contributes to Pathogenicity**

*UvHox2* deletion mutants DHOX17 and DHOX21 generated by pdTN3-HX2-Cas9I and DHOX51 and DHOX61 generated by pdTN3-HX2-Cas9II were then used to inoculate rice at booting stage on 15 spikes. Disease incidence of RFS was detected 25 days after inoculation. The rice smut balls were observed once again 60 days after inoculation to
check the chlamydospore formation. The results showed that the virulence of the *UvHox2* deletion mutants was significantly reduced compared to the wild-type strain P-1. The number of RFS balls on the spikes inoculated with the mutants DHOX-17, DHOX-21, DHOX-51, and DHOX-61 were 3.8 ± 1.6, 4.7 ± 2.6, 4.3 ± 1.6, and 4.3 ± 1.8, respectively; meanwhile, the number of smut balls on spikes inoculated with the wild-type strain P-1 was 21 ± 3.5 (Figure 5A and Table 1). The false smut ball samples were fixed with 2.5% glutaraldehyde in phosphate buffer pH 7.4 and subsequently observed under a SEM. The pictures showed that the chlamydospore formation structure was normal on the surface of false smut balls on the wild-type strain inoculated spikes. Short peg-like branches (sterigmata) were observed on the parallel arranged sporangiophore-like mycelia, and chlamydospores formed on these sterigmata in the wild-type strain P-1. In contrast, no special structure of hypha was observed on the false smut balls of *UvHox2* deletion mutants, and the hyphae were abnormally curved and twined (Figure 5B). To test whether conidia of *U. virens* wild-type strain (WT)/mutants could occasionally be converted into chlamydospores in the absence of water, conidia of *U. virens* WT/mutants were placed on glass slides and kept in the 9 mm-diameter petri dishes. After 10-days of incubation at 28°C in darkness, we found a small portion of conidia wild-type strain P-1 and *UvHox2* deletion mutants could both convert into chlamydospores (Figure 5C). The finding implied that *UvHox2* plays a key regulatory role in formation of chlamydospores rather than conversion from specific cells into chlamydospores.

**UvHOX2 Regulates Conidiogenesis**

The conidial production of *UvHox2* deletion mutants in YT broth was significantly reduced in comparison to that of the wild-type strain P-1 (Table 2). To observe the intact conidial sporulating structures, we cultured these strains (WT/mutants) on MM media. Wild-type strains produced typical conidial sporulating structures, but 86% of the *UvHox2* deletion mutants produced abnormal sporulating structures (Figure 5D). Whereas, the conidia produced by the
wild-type strain and the UvHox2 deletion mutant were the same in appearance.

**UvHOX2 Is Not Critical for Mycelia Growth and Conidium Germination**

In addition, mycelium growth rate and conidium germination rate of the UvHox2 deletion mutants was detected on YT media. The mycelium growth of UvHox2 deletion mutants were slightly reduced comparing to wild-type strains. And no significant differences in the germination of conidia were observed between the UvHox2 deletion mutant DHOX-61 and the wild-type strain P-1 (Table 2). This indicates that UvHOX2 is not critical in the regulation of mycelium growth and conidium germination in *U. virens*.

**UvHox2 Deletion Mutants Exhibited Increased Sensitivity to Oxidative, Osmotic, and Cell Wall Stresses**

When cultured on YT media amended with 0.05% H₂O₂ and 0.4 mol/l NaCl, the colony diameter of the UvHox2 deletion mutant DHOX-61 was significantly smaller than that of the wild-type strain P-1. When cultured on YT amended with
showed that the *UvHox2* deletion mutants were more sensitive to oxidative, osmotic and cell wall stresses than the wild-type strains. It suggests that UvHOX2 is also involved in responses to oxidative, osmotic, and cell wall stresses.

### Subcellular Location and Expression Patterns of UvHOX2

The UvHOX2-eGFP construct was transformed into wild-type strain P-1. The mutant HOX-GFP-2 was picked from a batch of transformants for its strong signal of the over-expressed eGFP fusion protein. The eGFP signal in the HOX-GFP-2 mycelia located in the nuclei (Figure 7A). 4′,6-diamidino-2-phenylindole (DAPI) was used to dye the mycelia and show the location of the nuclei in the cells.

The expression levels of *UvHox2* at the stages of mycelium growth, conidium generation, and chlamydospore formation were determined by qRT-PCR. The results showed that the expression of *UvHox2* was the highest during the early stage of chlamydospore formation and decreased at the later stage of chlamydospore formation. Meanwhile, the expression of *UvHox2* was also high at the sporulating stage and the early stage of infection on rice. In contrast, the expression of *UvHox2* was low during vegetative growth (Figure 7B).

### The Global Transcription Pattern of UvHox2 Deletion Mutant Differs From That of the Wild Type at the Early Stage of Chlamydospore Development

To identify putative regulated targets of the homeobox TF UvHOX2 during the formation of the chlamydospore generation structure, we compared the global gene transcription patterns in the *UvHox2* deletion mutant DHOX-61 with that in the wild-type strain P-1 by RNA-seq analysis. We inoculate P-1 (WTC) and DHOX-61 (DH) on rice as described above and collected rice false balls after 21 days. For false smut ball generated by P-1 (WTC), the membrane-like structure covering the false smut balls were intact then, and chlamydospore formation was at the early stage (Figure 8A).

The raw data of the RNA-seq was deposited in GenBank (accession number: SUB438508). There was high Pearson correlation among duplicates. We found that the global transcription patterns of DH and WTC at the early stage of

0.03% SDS and 100 mg/l congo red, the colony diameter of DHOX-61 was equivalent to that of P-1. However, the colony of DHOX-61 cultured on 100 mg/l congo red had shrinks and less aerial mycelia than P-1 (Table 3 and Figure 6). These findings

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### Table 1: Strains and vectors used in the study.

| Strain | Description in brief | References |
|--------|----------------------|------------|
| P-1    | Wild-type strain of *U. virens* | Zheng et al., 2017 |
| DHOX-17 | UvHOX2 deletion mutant | This study |
| DHOX-21 | UvHOX2 deletion mutant | This study |
| DHOX-51 | UvHOX2 deletion mutant | This study |
| DHOX-61 | UvHOX2 deletion mutant | This study |
| Trans-3 | UF-HYG′-RF ectopic insertion mutant | This study |
| HOX-GFP-2 | HOX-GFP fusion expression mutant | This study |

### Table 2: Pathogenicity, conidiation, and conidial germination in UvHOX2 deletion mutants.

| Strain | Mycelium growth (mm) | Pathogenicity | Concentration of conidia in YT broth (Log10 of spores/ml) | Percentage of abnormal sporulating structures (%) | Percentage of conidial germination (%) |
|--------|----------------------|--------------|----------------------------------------------------------|-----------------------------------------------|----------------------------------------|
| P-1    | 53.88 ± 2.61 A      | 21 ± 3.5 A   | 6.88 ± 0.02 A                                           | 19.25 ± 4.66 B                               | 90.56 ± 1.45 A                         |
| DHOX-17 | 44.75 ± 1.92 B     | 3.8 ± 1.6 B  | 6.78 ± 0.02 B                                           | 79.5 ± 4.72 A                                | 90.71 ± 1.19 A                         |
| DHOX-21 | 47.00 ± 1.37 B     | 4.7 ± 2.6 B  | 6.74 ± 0.02 B                                           | 73.75 ± 5.31 A                               | 89.33 ± 1.27 A                         |
| DHOX-51 | 45.13 ± 2.70 B     | 4.3 ± 1.6 B  | 6.65 ± 0.07 C                                           | 74 ± 6.67 A                                  | 92.65 ± 1.84 A                         |
| DHOX-61 | 43.00 ± 1.62 B     | 4.3 ± 1.8 B  | 6.75 ± 0.04 BC                                           | 74.5 ± 5.41 A                                | 90.05 ± 2.60 A                         |

*Number of rice false smut balls on the inoculated spike. †Data from at least four replicates were analyzed with the protected Fisher’s least significant difference (LSD) test. Different letters mark statistically significant differences (*P* ≤ 0.01).*
TABLE 3 | Responses of mycelium growth to abiotic stresses of UvHOX2 deletion mutants.

| Strain   | H2O2 Inhibition rate of mycelium growth (%) | NaCl Inhibition rate of mycelium growth (%) | SDS Inhibition rate of mycelium growth (%) | Congo red Inhibition rate of mycelium growth (%) |
|----------|-------------------------------------------|-------------------------------------------|-------------------------------------------|-----------------------------------------------|
| P-1      | 67.52 ± 3.55 B9                           | 48.78 ± 2.14 C                           | 35.37 ± 1.43 A                           | 36.43 ± 3.76 A                                |
| DHOX-17  | 79.33 ± 1.26 A                            | 55.87 ± 2.60 AB                         | 24.80 ± 2.91 AB                         | 27.04 ± 3.16 AB                              |
| DHOX-21  | 79.26 ± 1.93 A                            | 60.64 ± 2.63 AB                         | 32.98 ± 5.97 A                          | 33.78 ± 4.91 AB                              |
| DHOX-51  | 76.18 ± 3.02 A                            | 62.66 ± 2.14 A                         | 28.82 ± 5.93 AB                         | 28.26 ± 5.45 AB                              |
| DHOX-61  | 77.91 ± 1.87 A                            | 54.94 ± 3.93 BC                         | 23.26 ± 1.83 B                          | 23.72 ± 3.79 B                               |

aProtected Fisher’s least significant difference (LSD) test was used for statistical analysis. Different letters mark statistically significant differences (P ≤ 0.01).

FIGURE 6 | Growth of the UvHOX2 deletion mutant in the presence of different biotic stresses. The wild-type strain P-1, UvHOX2 deletion mutant DHOX-61 were cultured on YT medium or amended with 0.05% H2O2, 0.4 mol/l NaCl, 0.03% SDS and 100 mg/l congo red. Photographs were taken after incubation at 28°C for 15 days.

FIGURE 7 | Location and expression pattern of UvHOX2 in U. virens. (A) UvHOX2-eGFP was activated and located in nuclei of mycelia. 4',6-diamidino-2-phenylindole (DAPI) was used to stain nuclei in cells. (B) Expression pattern of UvHOX2 was determined by qRT-PCR. House-keeping gene elongation factor 1-α (EF1-α) was employed as a reference gene.

chlamydospore formation differed considerably. In a total of 8429 genes identified in U. virens previously (Zhang et al., 2014), 75 genes excluding UvHox2 were only expressed in WTC, 34 genes were barely expressed in DH, and 7008 overlapping genes were expressed in both WTC and DH (Figure 8B and Supplementary Table S2). Furthermore, we identified 1877 genes that have differential expression by at least twofold between WTC and DH, including 1185 genes being up-regulated and 692 genes being down-regulated in DH vs. WTC. These genes constituted approximately one-half of the differentially expressed genes, and they could be assigned to three major functional groups: biological process, cellular component, and molecular function (Figure 8C and Supplementary Table S2). In the biological process group, the top three subgroups of differentially
regulated genes in DH vs. WTC were “metabolic process,” “cellular process,” and “single-organism process.” In the cellular component group, the top three subgroups of differentially expressed genes in DH vs. WTC were “membrane,” “cell,” and “cell part.” In the molecular function group, the top three subgroups of differentially expressed genes in DH vs. WTC were “catalytic activity,” “binding,” and “transporter activity.” To validate the RNA-seq data, quantitative real-time polymerase chain reaction (qRT-PCR) was performed to confirm the differential expression of six selected genes. The qRT-PCR data for these genes were consistent with those obtained from RNA-seq (Supplementary Figure S2).

**Genes Involved in Signal-Transduction Pathway**
Several differentially expressed genes were detected in WTC, which were regarded as components in signal recognition and transduction system (Table 4). We also found that 43 genes encoding TFs were up-regulated during chlamydospore formation (Supplementary Table S2). And 19 of these genes were up-regulated beyond four folds (Table 4).

**Genes Involved in Cell Wall Synthesis**
Several differentially expressed genes were found to be closely linked to cell wall integrity. A gene encoding chitin deacetylase (KDB11455) were specially expressed in WTC but not in DH. Meanwhile, a chitin synthase (KDB11224) gene was up-regulated in WTC compared to DH. Chlamydospores of *U. virens* have thick cell walls. Chitin is an important component in cell wall, and fungi might mask chitin by deacetylating it into chitosan (Cord-Landwehr et al., 2016). These chitin synthases and deacetylase may play a key role in the thin cell wall synthesis in chlamydospores.

**Genes Involved in Ubiquitination and Autophagy**
Autophagy is a kind of intracellular recycling system that degrade cytoplasmic materials in lysosome/vacuole during development and in response to cell stresses in eukaryotic cells (Liu et al., 2017).
During chlamydospore formation, a lot of cytoplasmic materials may be degraded and reutilized. Here we identified three genes involved in autophagy that were differentially expressed in DH vs. WTC (Table 4).

**Genes Involved in Osmotic Response and Cell Membrane Integrity**

We found ten osmotic stress responsive genes were differentially expressed in DH vs. WTC. Nine of these genes encoding components for osmolarity two-component response system were up-regulated in DH vs. WTC. Meanwhile, several genes encoding cell membrane components were found to be up-regulated in WTC (Table 4). Most of them are transporter genes, which suggested that *U. virens* cells need to exchange substances with the environment more frequently during chlamydospore formation.

**Generation of Chlamydospores and Conidia Might Share a BrlA-AbaA-WetA Regulatory Pathway**

In *Aspergillus nidulans*, Myb-like DNA-binding protein FlbD is required for early conidiophore development (Wieser and Adams, 1995; Dong et al., 2015; Matheis et al., 2017). FluG regulates FlbD via repressing SfgA, a negative regulator of FlbD. FlbD delivers signals to the down-stream regulatory component FlbB to activate conidiogenesis regulatory cascade BrlA-AbaA-WetA (Wu et al., 2018). In *U. virens*, we found that homologs of FluG (KDB12888), FlbD (KDB18803), BrlA (KDB11753), AbaA (KDB11305), and WetA (KDB15008) were expressed at a higher level at the initial stage of chlamydospore and conidium formation than that at the vegetative growth stage. Homologs of FluG and FlbD were up-regulated, while homologs of *BrlA*, *AbaA*, and *WetA* were down-regulated in both initial sporulation mycelia and false smut ball at the initial stage of chlamydospore formation in DH compared to WTC (Figure 9). This suggested that the generation of chlamydospores and conidia may share the *BrlA*-AbaA-WetA regulatory pathway, and *BrlA*-AbaA-WetA signal cascade was downstream the UvHOX2 regulation.

**DISCUSSION**

Chlamydospores are a type of asexual spores that allow fungi to survive unfavorable conditions. These thick-wall spores play important roles in epidemic of pathogenic fungi as asexual resting spores and/or infection resources. Famous chlamydospore producing plant pathogenic fungi include *U. virens* (Zhang et al., 2014), *Fusarium oxysporum* (Klein et al., 2011), *F. sporotrichioides*, and *F. graminearum* (Goh et al., 2009). Some well-studied fungal biocontrol agents in livestocks, for example, *Trichoderma* spp. (Li et al., 2005), *Metarhizium anisopliae* (Ment et al., 2010), *Pochonia chlamydosporia* (Wang et al., 2005), and *Clonostachys rosea* (Ahmed et al., 2014), also produce chlamydospores. The development of chlamydospore is generally controlled by regulatory networks stimulated by the environment. Here, we report a homebox TF UvHOX2 that is essential for chlamydospore formation and also contributes to pathogenicity in *U. virens*. Additionally, we identified a group of genes that may participate in the downstream regulatory network of UvHOX2 during chlamydospore formation.

Homeobox domain-containing proteins play a critical role in the regulatory network of fungal development and pathogenicity as downstream elements in plant pathogenic fungi, but the
number of homeobox genes varies (Kim et al., 2009). Since the homolog of the transcription motif of STE12 in fungi is distinct from the typical homeodomain, we did not take account of the homolog of STE12 (KDB11415) as a homeobox gene. As a result, we identified seven genes encoding homeobox containing proteins in U. virens. Two of them contain multiple DNA-binding motifs besides homeobox motif. Only one homeobox DNA-binding motif was identified in the N-terminus of UvHOX2. In ascomycetous fungi, orthologs of UvHOX2 may play conserved roles in the development of specific structures during sporulation. In Verticillium dahliae, deletion of vhl1 (homolog of UvHox2) reduced sporulation rates in liquid medium (Sarmiento-Villamil et al., 2018). In Fusarium species, the deletion mutants failed to form conventional phialides and had obstructions in generating microconidia, but it could still produce macroconidia which was formed from hyphae through a budding-like mechanism (Zheng et al., 2012). In plant pathogen Magnaporthe oryzae, the Mohtf1 gene (homolog of UvHox2) is essential for conidiation but not for hypha growth and pathogenicity. The Mohtf1 deletion mutants generated more conidiophores, which failed to develop into sterigmata-like structures (Kim et al., 2009; Liu et al., 2010). Accordingly, deletion of the homolog of UvHox2 caused obstructions in conidiophoregenesis and completely abolished the generation structure of chlamydospores.

Conidia and chlamydospores are asexual spores produced by U. virens. In A. nidulans, regulatory factors, BrlA, AbaA, and WetA, were considered as cell developmental regulators that were critical for the development of conidiophore and phialide, as well as spore maturatio (Cary et al., 2017; Wu et al., 2018). Here, we provided a clue that BrlA-AbaA-WetA cascade may also participate in the regulation of chlamydospore formation in U. virens. Deletion of UvHox2 reduced the expression of BrlA, AbaA, and WetA, but the deletion did not affect the upstream regulatory factors FluG and FlbD. This suggested that UvHox2 and FlbD regulatory pathways could coordinate to regulate the downstream BrlA-AbaA-WetA cascade during sporulation and chlamydospore formation in U. virens. Although deletion of UvHox2 did not completely block the BrlA-AbaA-WetA signal cascade, the UvHox2 deletion mutant lost the ability to

| Genes (accession numbers) | Biological function | Up-/down-regulated |
|--------------------------|---------------------|-------------------|
| KDB11673                 | Transcription regulatory protein TUP1 | Up |
| KDB12278                 | MFS transporter       | Up |
| KDB17072                 | MFS multidrug transporter | Up |
| KDB12738                 | Calcium ion binding protein | Up |
| KDB18607                 | ABC transporter       | Up |
| KDB16091                 | Mitochondrial calcium uniporter protein | Up |
| KDB11501                 | Phosphatase beta protein | Up |
| KDB16963                 | Plasma membrane channel protein Azy1 | Up |
| KDB17556                 | Glycoside hydrolase family 3 protein | Up |

4Up- or down-regulated genes at the early stage of chlamydospore formation of wild-type strain P-1 comparing to UvHOX2 deletion mutant DHOX-61.

### TABLE 4 | A portion of genes that may under the control UvHOX2.

| Genes (accession numbers) | Biological function | Up-/down-regulated |
|--------------------------|---------------------|-------------------|
| KDB16312                 | G-protein coupled receptors | Up |
| KDB11861                 | G-protein coupled receptors | Up |
| KDB16287                 | G-protein coupled receptors | Up |
| KDB11050                 | G-protein coupled receptors | Down |
| KDB19029                 | Gtr1/RagA G protein | Up |
| KDB19030                 | Gtr1/RagA G protein | Up |
| KDB14836                 | Rho GTPase protein | Up |
| KDB11145                 | Serine/Threonine-protein phosphatase 6 subunit | Up |
| KDB12061                 | Serine/Threonine-protein kinase IO1 | Up |
| KDB17370                 | Protein kinase       | Up |
| KDB17112                 | Ethanolamine kinase | Up |
| KDB18374                 | TBC domain containing protein | Up |
| KDB15779                 | C2H2-type transcription factors | Up |
| KDB12683                 | C2H2-type transcription factors | Up |
| KDB12684                 | C2H2-type transcription factors | Up |
| KDB17197                 | C2H2-type transcription factors | Up |
| KDB17551                 | C2H2-type transcription factors | Up |
| KDB11104                 | bZIP transcription factors | Up |
| KDB14749                 | bZIP transcription factors | Up |
| KDB17948                 | C6 transcription factors | Up |
| KDB18664                 | C6 transcription factors | Up |
| KDB11753                 | Zinc finger protein odd-paired-like protein | Up |
| KDB13074                 | Homeobox transcription factor | Up |
| KDB14479                 | ACE1 | Up |
| KDB17109                 | bHLH family transcription factor | Up |
| KDB18696                 | Transcription factor | Up |
| KDB12822                 | Transcription factor | Up |
| KDB15421                 | Transcription factor | Up |
| KDB12143                 | Transcription factor | Up |
| KDB17685                 | Autophagy relative protein 3 | Up |
| KDB14954                 | Autophagy relative protein 5 | Up |
| KDB12188                 | Autophagy relative protein 7 | Up |
| KDB15378                 | Autophagy relative protein 12 | Up |
| KDB13656                 | Autophagy relative protein 26 | Up |
| KDB16605                 | Autophagy relative protein 22 | Up |
| KDB14455                 | Sm2 family ubiquitin-protein ligase | Up |
| KDB14890                 | Cullin 3 (McEwan and Dikic, 2014; Casanova et al., 2015) | Up |
| KDB17984                 | Target of rapamycin complex 2 (TORC2) subunit (Inoue and Klionsky, 2010) | Up |
| KDB18369                 | Sensor histidine kinase TcsA | Up |
| KDB18143                 | Sensor histidine kinase TcsA | Up |
| KDB18379                 | NIK1 | Up |
| KDB11585                 | SLN1 | Up |
| KDB12998                 | Phosphorelay intermediate protein YPD1 | Up |

(Continued)
form special structures for chlamydospore generation. UvHOX2 must control other regulatory pathways, which are critical for generating special structures during chlamydospore formation. Moreover, because conidia generated by UvHox2 deletion mutant could convert into chlamydospore as wild-type strains do, UvHOX2 may not be critical for maturation of chlamydospores in U. virens.

The two-component signaling proteins responsive to osmotic stress play an important role in cell tolerance under adverse environmental conditions. Our results revealed that Sln1 and Skn7, which were differentially expressed in DH vs. WTC, may function downstream of the UvHOX2 signaling pathway. Although the histidine kinase protein Sln1 and TF Skn7 play important roles in responses to osmotic, oxidative, and cell wall stresses (Zhang et al., 2010; Tang et al., 2013; Fan et al., 2014; Zhang et al., 2014; Yu J.J. et al., 2015; Song et al., 2016). The exact roles of these responsive proteins in cell development and pathogenesis need to be uncovered in the future.

The low frequency of homologous gene replacement in U. virens had limited its genetic study at the molecular level in the past years. Recently, a successful gene-deletion system based on CRISPR-Cas9 has been developed by Liang et al. (2018). It makes it convenient and efficient to perform gene deletion in U. virens. Using the optimized CRISPR-Cas9 cassettes, we developed a gene deletion system based on ATMT transformation and CRISPR-Cas9. This gene-deletion system shortens culture-selection-period by 7–10 days and provides a valuable tool for molecular genetic study in U. virens.

AUTHOR CONTRIBUTIONS

YL and JY conceived and designed the experiments. JY, MYu, ZQ, MYo, RZ, and XY performed the experiments. JY, TS, HC, XP, and YD analyzed the data. YL contributed to reagents, materials, and analysis tools. JY and YL wrote the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.01071/full#supplementary-material

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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