The false smut pathogen *Ustilaginoidea virens* requires rice stamens for false smut ball formation

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Summary

Rice false smut has emerged as a serious grain disease in rice production worldwide. The disease is characterized by the transformation of individual rice florets into false smut balls, which is caused by the fungal pathogen *Ustilaginoidea virens*. To date, little is known about the host factors required for false smut ball formation by *U. virens*. In this study, we identified histological determinants for the formation of false smut balls by inoculating *U. virens* into rice floral mutants defective with respect to individual floral parts. The results showed that *U. virens* could form mature false smut balls in rice floral mutants with defective pistils, but failed to develop false smut balls in the *superwoman* mutant lacking stamens, identifying that *U. virens* requires rice stamens to complete its infection cycle. Comparative transcriptome analysis indicated a list of candidate host genes that may facilitate nutrient acquisition by *U. virens* from the rice stamens, such as *SWEET11*, *SWEET14* and *SUT5*, and genes involved in the biosynthesis of trehalose and raffinose family sugars. These data pinpoint rice stamens as the key target organ of *U. virens* infection and provide a valuable starting point for dissecting the molecular mechanism of false smut ball formation.

Introduction

Rice false smut disease is caused by the ascomycete fungal phytopathogen *Ustilaginoidea virens* (Cooke) Takahashi (teleomorph: *Villosciava virens*) (Fan et al., 2016a). This disease has been reported with increasing occurrence in most rice production areas worldwide (Brooks et al., 2009; Ladhalakshmi et al., 2012; Jecmen and TeBeest, 2015). It not only causes considerable losses in terms of grain yield and quality but also acts as a health hazard to humans and animals due to poisonous mycotoxins contaminating the rice grains and straws of infected plants. The mycotoxins, including ustiloxins and ustilaginoidins, are produced from the fungal colonies in rice florets, called false smut balls (Koiso et al., 1994; Nakamura et al., 1994; Zhou et al., 2012; Lu et al., 2015; Wang et al., 2016a; Sun et al., 2017; Wang et al., 2017; Wang et al., 2019).

The false smut ball is the only visible symptom of rice false smut disease (Fan et al., 2016a). The process of false smut ball formation has been extensively investigated and can be summarized as follows. At the late booting stage of rice, spores of *U. virens* germinate on the surface of a rice spikelet (Ashizawa et al., 2012). Then, hyphae of *U. virens* extend into the inner space of the spikelet though the gap between the lemma and the palea and infect inner floral parts, including stamen filaments, lodicules and pistil (Ashizawa et al., 2012; Tang et al., 2013; Li et al., 2013b; Hu et al., 2014; Song et al., 2016). During infection, *U. virens* interrupts the process of rice flowering and fertilization and hijacks host nutrients for mycelial growth and false smut ball formation (Fan et al., 2015; Song et al., 2016). False smut balls are covered with chlamydospores that become orange or green (Wang et al., 2019) and generate sclerotia when
exposed to large day/night temperature differences (Fan et al., 2016b). As chlamydospores and sclerotia are produced from false smut balls and are the primary inocula of the rice false smut disease (Ikegami, 1960; Fan et al., 2010; Yong et al., 2018), understanding the factors involved in the formation of false smut ball is significant for controlling the disease.

To dissect the mechanism of false smut ball formation, the first step is to identify the infection sites of *U. virens* in rice flowers. It has been reported that *U. virens* primarily infects stamen filaments and, to a lesser extent, the lodicules (Tang et al., 2013). *U. virens* has also been reported to infect stigmas and styles (Song et al., 2016). Occasionally, *U. virens* hyphae can infect the ovary (Li et al., 2013b; Song et al., 2016). Several studies have consistently found that at all the observed infection sites *U. virens* extends only intercellularly and cannot form infection structures such as appressoria and haustoria (Tang et al., 2013; Hu et al., 2014; Song et al., 2016). So far, *U. virens* infection sites have not been detected in other floral parts such as anthers, lemma, palea or rac- chilla (Tang et al., 2013; Song et al., 2016). Therefore, we speculated that successful colonization of *U. virens* may require the presence of stamen filaments, lodicules or pistil. Nevertheless, it is still an open question as to which floral parts are essential to support false smut ball formation by *U. virens*.

In the current study, we investigated the events in detail at early infection stages by utilizing an efficient artificial inoculation system with a Green Fluorescent Protein (GFP)-tagged *U. virens* isolate and a highly susceptible rice accession Pujiang6 (Pu6) for artificial inoculation. We examined the outer surface and inner space of more than 500 rice spikelets collected from multiple inoculated panicles every 2 days post-inoculation (dpi) and recorded the infection status of *U. virens* in each floral part of each spikelet. Our data demonstrated that P4 hyphae were restricted to the outer surface of all the examined spikelets at 1 and 3 dpi, extending to the inner floral parts at 5 dpi (Figs 1 and 2A-C).

Taking the data from year 2017 as an example (Fig. 1A), P4 hyphae reached the stamen filaments of 2.78% of the examined spikelets and the lodicules in that *U. virens* preferentially attacked rice stamens. Moreover, we performed artificial inoculation on several floral rice mutants deficient with respect to either stamens or pistils and demonstrated that rice stamens were essential for the formation of false smut ball. To identify potential molecular determinants involved in false smut ball formation, we further conducted RNA-Seq analysis on rice stamens infected with *U. virens*, which provided novel insights into the underlying mechanisms of nutrient acquisition from rice florets by *U. virens*.

**Results**

*Ustilaginoidea virens* preferentially attacks stamen filaments of rice

To identify the key target site of *U. virens* in rice spikelets, we first examined whether *U. virens* showed any preference for specific floral parts during infection. We monitored the infection process at early infection stages in detail by utilizing a GFP-tagged *U. virens* strain P4 and a highly susceptible rice accession Pujiang6 (Pu6) for artificial inoculation. We examined the outer surface and inner space of more than 500 rice spikelets collected from multiple inoculated panicles every 2 days post-inoculation (dpi) and recorded the infection status of *U. virens* in each floral part of each spikelet. Our data demonstrated that P4 hyphae were restricted to the outer surface of all the examined spikelets at 1 and 3 dpi, extending to the inner floral parts at 5 dpi (Figs 1A and 2A–C).

Taking the data from year 2017 as an example (Fig. 1A), P4 hyphae reached the stamen filaments of 2.78% of the examined spikelets and the lodicules in...
2.19% of the examined spikelets at 5 dpi. The value fell to 0.2% for both anthers and stigmas at 5 dpi. At 7 dpi, the percentage infection increased rapidly, with P4 hyphae reaching all the inner floral parts (Fig. 2D), at percentages of 19.85, 15.24, 15.43, 4.52 and 2.54 for stamen filaments, lodicules, anthers, stigmas and ovaries respectively (Fig. 1A). The percentage infection was the least for ovaries at both 5 and 7 dpi (Fig. 1A). At 9 dpi, P4 hyphae had reached all inner floral parts of most (over 86%) of the infected spikelets (Fig. 2E). Subsequently, fungal growth increased rapidly to embrace all the inner floral parts (Fig. 2F and G). Although the values of infection proportions varied between years 2017 and 2018, similar trends were observed, i.e. the proportion was the highest for P4 hyphae attacking stamen filaments, followed by lodicules, with ovaries showing the least pathogen colonization (Fig. 1B). These data indicate that *U. virens* preferentially attacked stamen filaments.

We also monitored the infection status of P4 in rice florets of Pu6 at a very late infection stage (25 dpi), by which time mature false smut balls had already formed (Fig. 2H). We found that approximately 25% of inoculated spikelets had been converted into false smut balls in both 2017 and 2018 (Fig. 1C). Interestingly, 12.5%–18.9% of the examined spikelets had P4 mycelia embracing all the inner floral parts but were not converted into false smut balls (Fig. 1C). These observations indicate that not all infected spikelets could develop into false smut balls, but
colonization of all the inner floral parts is the prerequisite for the conversion of a spikelet into a false smut ball. Therefore, it is important to determine which floral part needs to be colonized in order to achieve the formation of false smut balls.

**Ustilaginoidea virens fails to develop false smut balls in stamen-/pistil-deficient rice mutant**

We screened floral organ deficient mutants from our previous ethyl methanesulfonate (EMS)-mutagenesis library derived from an indica rice X3107 (Liao et al., 2018), for their ability to form false smut balls following inoculation with *U. virens* PJ52-2-5 (Fig. 3A). The mean number of false smut ball per diseased panicle reached 22 and the disease incidence was up to 85% (Table 1), following inoculation of PJ52-2-5 onto wild-type X3107B, indicating compatible interaction.

We isolated a floral mutant without any stamens or pistil in its floret but possessing two lodicules (Fig. 3B). We inoculated 30 panicles of this stamen-/pistil-deficient mutant with PJ52-2-5, but did not observe false smut balls formed from the mutant spikelets, although there was limited pathogen growth on the lodicules of 16 infected spikelets from seven diseased panicles (Fig. 3B; Table 1). Using an environmental scanning electron microscope, we observed hyphae of the pathogen on anthers, stamen filaments, lodicules and ovaries of X3107B (Fig. 3C–E). We also observed hyphae of the pathogen covering the lodicules of the mutant florets (Fig. 3G and I), whereas no hyphae were seen on the mock-inoculated florets (Fig. 3F and H). These data demonstrate that *U. virens* hyphae could still invade into the inner space of the stamen-/pistil-deficient mutant spikelets but could not form false smut balls. These results indicate that lodicules could support the limited growth of pathogen hyphae but were not sufficient to support false smut ball development.

**Ustilaginoidea virens could successfully colonize pistil-defective rice mutants**

To identify the role of rice pistil in *U. virens* infection, we inoculated a pistil-defective mutant, which was a spontaneous mutant derived from rice accession W2555 in our germplasm collection. This floral mutant also exhibited abnormal lemma and palea, resulting in imperfect closure of the spikelet (Fig. 4A). In the innermost whorl of the mutant flower, the pistil was replaced by unknown translucent structures (Fig. 4B). Environmental scanning electron microscopy (ESEM) analysis showed that lamellar structures were generated in the carpel whorl (Fig. 4E–H), in contrast to the normal appearance of W2555 pistil (Fig. 4C and D). In addition, trichome-like tissues were observed on the lamellar structures of the mutant floret (Fig. 4G and H). As expected, abnormal development of the mutant pistil resulted in sterility of the mutant floret. When this pistil-defective mutant was artificially inoculated with *U. virens* PJ52-2-5, mature false smut balls were normally formed in the inoculated panicles, with the ball size being larger than that in the wild-type W2555 (Fig. 4I; Table 1). These results indicate that *U. virens* does not require the presence of a rice pistil for the development of false smut ball.

We also inoculated a pistil-sterile rice mutant *ptb1* in the rice false smut disease assay. *ptb1* was a spontaneous mutant derived from an indica rice X3107 and caused by loss-of-function of the POLLEN TUBE BLOCKED 1 gene (Li et al., 2013a). *ptb1* did not allow pollen tube growth towards ovary due to intense callose deposition on the ovule surface, thus resulting in female-specific sterility (Li et al., 2013a). Following inoculation with *U. virens* PJ52-2-5, *ptb1* supported the formation of false smut balls (Fig. 4J). The average number of false smut ball per diseased panicle and the mean false smut ball size in *ptb1* were not significantly different from those in the wild-type X3107 (Fig. 4J; Table 1). Taken together, the dysfunction of rice pistil fertilization did not stop *U. virens* from developing false smut balls.

**Ustilaginoidea virens fails to generate false smut balls in stamen-deficient rice mutant**

In the EMS-mutagenesis library of rice accession Yixiang1B (Liao et al., 2018), we found another floral mutant without any stamens but with multiple pistils in its floret (Fig. 5), resembling the superwoman mutant phenotype (Nagasawa et al., 2003). Cytological analysis displayed that the mutant floret had five pistils with partially fused ovaries and two palea-like organs instead of lodicules (Fig. 5C, D, and G). No stamens or other floral parts were observed. Artificial inoculation analysis showed that this superwoman mutant could not support false smut ball formation in spikelets, while the wild-type Yixiang1B produced mature false smut balls after *U. virens* infection (Fig. 5A; Table 1). Interestingly, considerable growth of mycelium was observed on the superwoman ovaries, which were enlarged in 19 of 30 infected spikelets at 36 dpi (Fig. 5B, E, and F).

To identify the infection sites of *U. virens* in the superwoman mutant, inoculated spikelets were collected at 16 dpi (ovaries were not enlarged; inset in Fig. 5G) and 36 dpi (ovaries were enlarged; inset in Fig. 5M), and subjected to anatomical analysis. At 16 dpi, large quantities of *U. virens* hyphae were observed surrounding the ovaries (Fig. 5G) and stigmas (Fig. 5J) under a light microscope. Further transmission electron microscopy (TEM) analysis showed that *U. virens* hyphae were unable to
Infection of *Ustilaginoidea virens* in a stamen-pistil-deficient rice mutant.

A. *U. virens* isolate PJ52-2-5 infected an *indica* rice accession Yixiang1B (wild-type, WT) and formed false smut balls (black arrows). B. A stamen-pistil-deficient rice mutant (MT) was derived from an EMS mutagenesis mutant library of Yixiang1B. In the inner space of MT spikelet, only lodicules were present. PJ52-2-5 failed to form false smut balls in MT but did grow on the lodicules with visible fungal mass (white arrows). C–E. Scanning electron microscopy (SEM) analysis of Yixiang1B floret infected with *U. virens* at 11 dpi (day post-inoculation). F, H. SEM image of MT floret without *U. virens* infection. G, I. SEM image of MT floret infected with *U. virens* at 11 dpi. an, anther; lo, lodicule; ov, ovary; pa, palea; sf, stamen filament. White arrows indicate *U. virens* hyphae. Bar size = 1 cm (right panel in A), 1 mm (A, B), 200 μm (C–E) and 500 μm (F–I). [Color figure can be viewed at wileyonlinelibrary.com]
infected stamens, compared with control stamens revealed 4431 differentially expressed genes (DEGs) in clean reads for each replicated sample. Data analysis investigating tissues (Fig. 2E), representing a suitable time point for infected stamens could be clearly separated from other hyphae mostly invaded stamen tissues and Pu6 at 9 dpi, because at this infection stage the pathogen inoculated and P4-inoculated panicles of rice accession criptome analysis. We collected spikelets from mock-Seq. In overview, RNA-Seq generated 54.7 million reads, with 22.2-fold (Table S5) respectively. The expressions of Synthase 1b (Os01g0633100), PPase LARGE SUBUNIT 2 (Os08g0345800), PYROPHOSPHORYLASE 1 (Os07g0687900) and putative mRNA family sugars and trehalose tended to be reduced by more than twofold (Fig. 6, Fig. S1; Table S4). Particularly, a number of genes responsible for starch synthesis were significantly downregulated, such as ADP GLUCOSE PYROPHOSPHORYLASE 1 (Os08g0345800), ADPGLC-PASE LARGE SUBUNIT 2 (Os01g0633100), Granule-bound starch synthase 1b (Os07g0412100) and Soluble starch synthase 3 (Os05g0533600), with expression levels reduced by more than twofold (Fig. 6, Fig. S1; Table S4). On the contrary, genes responsible for the synthesis of rafinose family sugars and trehalose tended to be upregulated. For instance, the expressions of galactinol synthase 2 (Os07g0687900) and putative raffinose synthase (Os06g0172800) were increased by 22.2-fold and 4.6-fold (Table S5) respectively. The expressions of

Table 1. Disease assay data of rice floral mutants inoculated with U. virens.

| Material | Background | Number of inoculated panicles | Number of diseased panicles | Disease incidence (%) | Average number of false smut balls per diseased panicle | Diameter of false smut ball (cm) |
|----------|------------|-------------------------------|------------------------------|-----------------------|---------------------------------------------------------|---------------------------------|
| Yixiang1B | N/A        | 27                            | 23                           | 85.19                 | 22.00 ± 0.65                                            | 0.84 ± 0.16                     |
| Stamen-pistil-deficient mutant | Yixiang1B | 30                            | 0                            | 0                     | N/A                                                     | N/A                             |
| Stamen-deficient/multi-ovary mutant | Yixiang1B | 50                            | 0                            | 0                     | N/A                                                     | N/A                             |
| W2555    | N/A        | 20                            | 13                           | 65.00                 | 16.38 ± 6.90                                            | 0.64 ± 0.09                     |
| Pistil-defective mutant | W2555 | 20                            | 12                           | 60.00                 | 13.58 ± 5.81                                            | 0.92 ± 0.23***                 |
| X3107    | N/A        | 17                            | 10                           | 58.82                 | 31.40 ± 18.05                                           | 0.53 ± 0.09                     |
| pbt1     | X3107      | 18                            | 11                           | 61.11                 | 47.36 ± 14.08                                           | 0.50 ± 0.03                     |

N/A, not applicable.

a. Diseased panicles indicate panicles with false smut balls.

b. Data points refer to the mean ± standard deviation. Student’s t-test was performed to examine the significance of differences between wild-type and mutant. **P-value < 0.01.

infect ovary (Fig. 5H and I) but invaded into stigma and extended intercellularly among stigma cells (Fig. 5K and L). Note that the epidermal cells of the ovary were tightly organized, while the stigma cells were irregularly and loosely arranged. At 36 dpi, some ovaries were enlarged in the infected floret, and the remaining ones became degenerated (Fig. 5M). In the middle of the enlarged ovaries, a batch of cells had collapsed. Surprisingly, the outer layer of the enlarged ovaries was found to be infected intercellularly by U. virens hyphae (Fig. 5N). The intercellular infection of U. virens in the ovary was further confirmed by TEM analysis (Fig. 5O).

Taken together, these data indicate that U. virens was unable to form false smut balls in rice mutant florets lacking stamens, despite being able to initiate infection in stigmas or in enlarged ovaries.

Ustilaginoidea virens infection results in global transcriptional changes in rice stamens

As U. virens needs rice stamens for successful colonization and false smut ball formation (Fig. 1; Fig. 5), we decided to screen for molecular determinants in rice stamens required for U. virens infection through transcriptome analysis. We collected spikelets from mock-inoculated and P4-inoculated panicles of rice accession Pu6 at 9 dpi, because at this infection stage the pathogen hyphae mostly invaded stamen tissues and U. virens-infected stamens could be clearly separated from other tissues (Fig. 2E), representing a suitable time point for investigating U. virens–rice stamen interaction. Stamens were then carefully separated and subjected to RNA-Seq. In overview, RNA-Seq generated 41.5–54.7 million clean reads for each replicated sample. Data analysis revealed 4431 differentially expressed genes (DEGs) in infected stamens, compared with control stamens (criteria: P < 0.05 and absolute Log2 Fold change ≥1). Among the DEGs, 2562 genes were upregulated and 1869 genes were downregulated (Table S1). The data from the DEGs were further subjected to MapMan software for pathway analysis. The results showed that a number of pathways were significantly modulated by U. virens infection, such as biotic stress, regulation of transcription, receptor kinases, hormone metabolism, secondary metabolism, major carbohydrate metabolism, lipid metabolism, and so forth. (Table S2). For example, DEGs were mostly enriched in pathways of biotic stress (P = 1.53E-11) and receptor kinase signalling (P = 2.99E-11). In particular, 78 out of 93 (83.87%) biotic stress-related DEGs and 97 out of 117 (82.91%) DEGs encoding receptor kinases were coordinately upregulated in U. virens-infected stamens (Table S3). These data indicate that defence responses are transcriptionally activated in rice stamens in response to U. virens infection; nevertheless, these defence responses could not restrict the pathogen growth in the host tissue.

To facilitate visualization of the expression changes of DEGs involved in different metabolisms, we mapped the data of DEGs to metabolism pathways in MapMan. As displayed in Fig. 6, genes involved in amino acid synthesis, phenylpropanoids synthesis and minor carbohydrate metabolism were coordinately upregulated in rice stamens infected with U. virens, whereas genes involved in starch-sucrose metabolism and phospholipid synthesis were coordinately downregulated (Fig. 6; Table S4). Particularly, a number of genes responsible for starch synthesis were significantly downregulated, such as ADP GLUCOSE PYROPHOSPHORYLASE 1 (Os08g0345800), ADPGLC-PASE LARGE SUBUNIT 2 (Os01g0633100), Granule-bound starch synthase 1b (Os07g0412100) and Soluble starch synthase 3 (Os05g0533600), with expression levels reduced by more than twofold (Fig. 6, Fig. S1; Table S4). On the contrary, genes responsible for the synthesis of rafinose family sugars and trehalose tended to be upregulated. For instance, the expressions of galactinol synthase 2 (Os07g0687900) and putative raffinose synthase (Os06g0172800) were increased by 22.2-fold and 4.6-fold (Table S5) respectively. The expressions of
Nine trehalose synthesis-related genes were increased by 2.2-fold–477.7-fold (Table S4). Moreover, a sucrose export gene $SU75$ (Os02g0576600) was highly upregulated by more than fivefold (Fig. S1; Table S4). Bidirectional sugar transporter genes $SWEET11$ (Os08g0535200) and $SWEET14$ (Os11g0508600) were also upregulated by 3.4-fold and

Fig. 4. Infection of *Ustilaginoidea virens* in pistil-defective rice mutants. A, B. Spikelet phenotype of a pistil-defective mutant derived from accession W2555. Compared with the wild-type (WT), the mutant (MT) spikelet has normal stamens but no pistils. In addition, the MT spikelet has abnormal lemma and palea, resulting in imperfect closure. Image areas marked with dotted line rectangles were enlarged and presented in (C) and (D). White arrowheads indicate unknown lamellar structures replacing pistil.

E–J. Scanning electron microscopy analysis of WT (E, F) and MT (G–J) floral organ. Note that stigmas and ovary were not observed in MT; instead, unknown lamellar structures were present (white arrowheads).

K, L. *U. virens* isolate PJ52-2-5 was inoculated into panicles of WT (K) and MT (L). Disease phenotype was recorded at 4–5 weeks post-inoculation (wpi). Mature false smut balls were observed in both WT and MT panicles.

M, N. PJ52-2-5 was inoculated into panicles of an indica rice accession X3107 (M) and a female-sterile mutant $ptb1$ derived from X3107 (N) (Li et al., 2013a). Disease phenotype was recorded at 4–5 wpi. Mature false smut balls were observed in both X3107 and $ptb1$ panicles. sf, stamen filament; lo, lodicule; an, anther; st, stigma; ov, ovary. White arrows indicate rice false smut balls. Bar size = 5 mm (A, B), 1 mm (C, D, E, G), 0.4 mm (F), 0.5 mm (H–J) and 1 cm (K–N). [Color figure can be viewed at wileyonlinelibrary.com]
5.8-fold respectively (Table S1). These data indicate that *U. virens* infection may cause inhibition of polysaccharide synthesis but activation of disaccharide/oligosaccharide synthesis in rice stamens, with the sugars that accumulated in the stamen cells being exported into apoplast space to facilitate *U. virens* growth.
To validate the RNA-Seq data, we selected 11 DEGs involved in different pathways for quantitative polymerase chain reaction (qPCR) analysis. The results showed that all the tested genes demonstrated the same trend of expression changes in RNA-Seq and qPCR experiments (Fig. S2), indicating that the RNA-Seq data were reliable in this study.

Fig. 5. Infection of Ustilaginoidea virens in superwoman rice mutant.
A. Disease phenotype of Yixiang1B (wild-type, WT) and a superwoman mutant (MT) derived from an EMS mutagenesis mutant library of Yixiang1B, at 1 month post-inoculation of U. virens isolate PJ52-2-5. Mature false smut balls (white arrows) were observed in WT panicles but never observed in MT panicles.
B. Observation of inner floral parts of MT spikelets under infected or uninfected conditions. White mycelium was seen (white arrows) and ovaries were enlarged in infected MT spikelets at 36-day post-inoculation (dpi).
C-F. Scanning electron microscopy analysis of MT floral organs under uninfected (C, D) or infected (E, F) conditions. Image areas marked with dotted line rectangles in (C) were enlarged and presented in (D), (E) and (F) are views of different parts from the same infected mutant spikelet. Considerable fungal growth (white arrows) was detected on enlarged ovaries in infected spikelets.
G-L. Anatomical analysis of infected MT spikelets at 16 dpi, when the ovaries were not enlarged (inset in G). Under light microscopy (G, I), intact ovary cells were observed and were not infected by pathogen hyphae (red arrows). Infection sites on stigmas were examined under both light microscopy and transmission electron microscopy. Large-scale fungal hyphal accumulation (white or red arrows) was detected around stigmas (J) and in the extracellular space among cells (N, O), st. stigma; ov. ovary. Arrowheads in Fig. 5C–E indicate palea-like organs. Arrows indicate U. virens pathogen. *, collapsed cells. Bar size = 2 cm (A), 5 mm (inset in A and B), 1 mm (C, E), 300 μm (D), 500 μm (F), 5 μm (H, I, L, O), 10 μm (K), 20 μm (N), 200 μm (G, J) and 300 μm (M). [Color figure can be viewed at wileyonlinelibrary.com]

Fig. 6. Expression changes of metabolism genes in rice stamens in response to Ustilaginoidea virens infection. Rice stamens were collected from mock- and U. virens P4-inoculated spikelets of rice accession Pu6 and subjected to comparative RNA-Seq analysis. Differentially expressed genes (DEGs) (absolute Log2 Fold change ≥1 and P value < 0.05) were mapped to the Metabolism pathway with MapMan software. Red points represent upregulated genes, while blue points represent downregulated ones. Dotted line rectangles represent pathways of particular interest. Expression data are included in Table S4. [Color figure can be viewed at wileyonlinelibrary.com]
Discussion

False smut ball is critical for the disease cycle as it generates chlamydospores and sclerotia, representing the primary inocula for this pathogen, but it is detrimental to humans and animals as it produces mycotoxins (Yong et al., 2018; Wang et al., 2019). Failure of false smut ball formation is believed to interrupt the disease cycle as a result of the depletion of the primary inoculum. Therefore, understanding the mechanism of false smut ball formation is of great potential for controlling the disease. Here, we demonstrated that *U. virens* preferentially attacked rice stamen (Fig. 1) and was unable to form false smut balls in rice mutant florets lacking stamens (Fig. 5). In addition, by conducting comparative transcriptome analysis on rice stamens infected with *U. virens*, we found that *U. virens* modulated host sugar- and amino acid-related metabolic pathways to promote nutrient acquisition.

*U. virens* is a biotrophic fungal pathogen. In an intercellular manner, it infects a subset of rice tissues, primarily stamen filaments, stigmas, the outer layer of lodicules, and epidermal cells of seedling roots and coleoptiles (Ikegami, 1963; Tang et al., 2013; Song et al., 2016; Prakobsub and Ashizawa, 2017). However, *U. virens* could not infect leaves, stems and pedicels of rice (Tang et al., 2013; Fan et al., 2014). The organ-specificity of *U. virens* infection is suggested to be associated with the ultrastructure and components of cell walls in different rice organs (Yong et al., 2016; Rong et al., 2017). For example, the packing of cells in stamen filaments is loose and the cell walls are flexible, whereas, in stem and ovary, the cell walls are compact (Tang et al., 2013; Yong et al., 2016; Rong et al., 2017). Therefore, *U. virens* may find it easier to colonize stamen filaments, an observation that is supported by our data in this study (Fig. 1). *U. virens* can also easily attack lodicules and stigmas, as demonstrated by the observation that *U. virens* hyphae could invade the inner space of stamen-/pistil-deficient mutant florets and proliferate on lodicules (Fig. 3), and that a large amount of *U. virens* mycelia could be accommodated inside *superwoman* mutant flowers, when only stigmas were infected (inset in Fig. 5G). On the contrary, *U. virens* could not infect ovaries with compact cell walls in the *superwoman* mutant (Fig. 5G–I), although *U. virens* infection occurred in some ovaries that were enlarged by unknown mechanisms (Fig. 5M–O). These data indicate that *U. virens* can infect and proliferate in a few rice organs or tissues, whereas false smut ball formation is restricted to florets.

Which floral part is required for *U. virens* to form false smut balls? Previously, Song et al. (2016) had proposed that *U. virens* infects rice stigmas and styles to mimic ovary fertilization and to continually acquire nutrients for the development of false smut ball, indicating that pistil may be required for false smut ball formation. However, here we found that pistil-defective rice flowers could still support the false smut ball development (Fig. 4), while stamen-deficient mutant flowers failed to develop false smut balls after *U. virens* infection (Fig. 5). These observations suggest that rice stamen is essential for the formation of false smut balls. How does rice stamen facilitate *U. virens* to develop false smut balls? It is unlikely that local nutrients from stamens provide sufficient nutrients for *U. virens* to form false smut balls. As stamen filaments were infected and ultimately replaced by *U. virens* hyphae, and stamen filaments are connected to the base of ovary, and tap into rachilla via vascular bundles that transport nutrients from source organs (Wang et al., 1994), we speculated that *U. virens* hyphae may acquire abundant nutrients by establishing an interface at rachilla through stamen filaments. In turn, lacking of rice stamens could lead to the failure of establishing the proposed nutrient interface, preventing nutrient transport and blocking the formation of false smut balls.

To understand how *U. virens* manipulates rice stamens for successful infection, we further investigated *U. virens* modulation of nutrient-related pathways in rice stamens via transcriptome analysis. Our data demonstrated that *U. virens* appears to manipulate rice stamens for nutrient acquisition in the following ways. First, *U. virens* upregulated the expression of sugar transporter genes, such as *SWEET11, SWEET14* and *SUT5* (Fig. S1; Table S1). *SWEET11* and *SWEET14* mediate efflux of glucose and can be hijacked by the bacterial phytopathogen *Xanthomonas oryzae* to pump sugars from host cells into apoplast space, which the pathogen inhabits (Chen et al., 2010). *SUT5* has been demonstrated to transport sucrose (Sun et al., 2010). Upregulation of these genes in *U. virens*-infected rice stamens likely facilitates sugar supply to support the intercellular growth of *U. virens*. Second, *U. virens* infection induced the expression of genes encoding galactinol synthase 2 and putative raffinose synthase, which are involved in biosynthesis of galactinol and raffinose family sugars (Smith et al., 1991) respectively (Fig. 6; Table S5). Raffinose could either be utilized directly by *U. virens* as a carbon source (Wang et al., 2016b) or in concert with galactinol, to biosynthesize stachyose. Stachyose has been shown to be a preferential carbon source for the growth of *U. virens* (Wang et al., 2016b), although rice genes responsible for stachyose biosynthesis have not been identified. Therefore, raffinose family sugars may be favourably manipulated by *U. virens*. Third, *U. virens* infection led to the upregulation of a subset of genes related to trehalose biosynthesis and hormone metabolisms of cytokinin and ethylene (Fig. 5; Table S4, Table S6). Trehalose could be utilized as a carbon source by *U. virens* (Fig. S3); additionally, trehalose may promote acquisition of nitrogen-
containing nutrients in *U. virens*-inhabited intercellular space, as has been reported for *Pseudomonas aeruginosa* in plant leaf (Djonovic et al., 2013). Interestingly, increasing concentrations of cytokinin and ethylene, promoted by the rice endophyte *Phomopsis liquidambaris*, can also enhance the uptake of nitrogen-containing nutrients in rice (Li et al., 2017). Thus, *U. virens* may stimulate rice to take up and supply with more nutrient-containing nutrients to appropriate locations to achieve pathogen proliferation, via modulating trehalose and hormone metabolisms. Finally, *U. virens* upregulated the expression of a subset of genes involved in the biosynthesis of amino acids (Fig. 6; Table S4), which may contribute to meeting the nitrogen requirements of *U. virens* growth.

Compared with previous transcriptome analyses on compatible interactions between different *U. virens* isolates and rice accessions, expressions of *SWEET11* and/or *SWEET14* were consistently upregulated by *U. virens* infection (Table S1) (Chao et al., 2014; Fan et al., 2015), suggesting that *SWEET* gene family in rice may also be targeted by the flower-infecting fungal pathogen *U. virens*, in addition to the leaf-infesting bacterial pathogen *Xanthomonas oryzae* pv. *oryzae* (Xu et al., 2019). Pathways associated with biotic stress, secondary metabolism and hormone metabolism were also consistently modulated in rice flowers infected with *U. virens*, according to the previous transcriptomic analyses and this study (Table S1) (Chao et al., 2014; Yang et al., 2014; Fan et al., 2015; Han et al., 2015). In a previous transcriptomic analysis (Fan et al., 2015), up to 35 rice grain-filling-related genes were activated by *U. virens* infection; whereas, in the present work, only three genes related to grain filling (*Os07g0182000*, *Os03g0188500* and *Os01g0644600*) were induced in rice stamens infected with *U. virens* (Table S1). Moreover, the present study uniquely identified, in rice stamens infected with *U. virens*, a subset of upregulated genes that are involved in the biosynthesis of trehalose and raffinose family sugars (Table S5), which may point to a novel nutrient acquisition strategy deployed by *U. virens*. These differences in expression patterns between different studies are probably attributable to the use of different rice accessions and/or different time points after *U. virens* inoculation for transcriptome analysis in these studies.

Overall, we demonstrated that *U. virens* requires rice stamens for successful colonization and development of false smut ball, and identified a set of candidate genes in rice stamens, which may be employed by *U. virens* to facilitate nutrient acquisition from the host. In the future, these candidate genes need to be further investigated to determine whether they are required for initial infection and/or false smut ball formation of *U. virens*.

**Experimental procedures**

**Plant materials and fungal isolates**

Rice accessions Pu6, W2555, X3107, Yixiang1B, and the corresponding floral mutants were grown in an experimental field of Sichuan Agricultural University, China. *Ustilaginoidea virens* isolate PJ52-2-5 was obtained from a rice false smut ball in Pu6 by amerosporous purification (Fan et al., 2019). The GFP-tagged *U. virens* P4 was a gift from Prof. Yongfeng Liu from Jiangsu Academy of Agricultural Sciences, China.

**Artificial inoculation**

Artificial inoculation of *U. virens* was conducted as described in a previous report (Fan et al., 2015). Briefly, *U. virens* mycelium was cultured on potato-sucrose-agar medium and transferred into potato-sucrose broth (PSB) incubated at 28°C and 120 rpm for seven days. The blended mixture of mycelia and conidia (concentration adjusted to 1 × 10⁶ conidia/ml) was injected into rice panicles at late booting stages (5–7 days before heading) until the inoculum dripped out. PSB alone was inoculated into a separate set of panicles as mock-inoculation controls.

**Fluorescence microscopy and ESEM**

For fluorescence microscopy, rice spikelets were collected from P4-inoculated panicles of Pu6 at multiple time points. Half of the lemma and palea was removed from spikelets for better observation of the inner floral organs under a fluorescence microscope (Zeiss Axio Imager A2, Carl Zeiss). For ESEM, spikelets were sampled from pistil- and/or stamen-deficient floral mutants and their corresponding wild-type plants, and directly examined with an environmental scanning electron microscope (FEI Quanta 450, FEI) at low-vacuum (70 Pa), after removing the lemma and the palea.

**Light microscopy and TEM**

Sample preparation for anatomical analysis was performed as previously described with minor modifications (Tang et al., 2013). In brief, *U. virens*-infected spikelets were collected from superwoman mutant derived from EMS mutagenesis of Yixiang1B, and pre-fixed in 2.5% glutaraldehyde in 0.1 M PBS (pH 7.0) overnight at 4°C. The pre-fixed samples were washed three times with 0.1 M PBS (pH 7.0), 15–20 min each time, and post-fixed with 1% osmium tetroxide. After a series of washing and dehydration steps, the samples were embedded in Spurr resin and polymerized overnight at 70°C.
For light microscopy, a series of 1 μm sections were prepared from the embedded samples, stained with 0.05% methylene blue, and observed under a Nikon Eclipse 80i microscope. For TEM, 6–7 nm ultrathin sections were prepared and examined under a TEM (Hitachi H-7650) as previously described (Tang et al., 2013).

**RNA-Seq analysis**

Total RNAs were isolated from Pu6 rice stamens at 9 day post-inoculation (dpi) of either P4 or PSB. Quantity and integrity of RNAs were analysed by using a Nanodrop 2000 spectrophotometer (Thermo Scientific, Waltham) and Bioanalyzer 2100 (Agilent Technologies, Santa Clara). Library construction and Illumina HiSeq 2500 sequencing were carried out at Novogene (Beijing, China). Sequencing data were filtered to remove adaptor and low-quality reads. The generated clean data were then mapped to Oryza_sativa.IGRSP-1.0.26 genome (Ensembl) by using TopHat2 with default parameters (Kim et al., 2013). Subsequently, gene expression was determined by HTSeq (Anders et al., 2015). Differential gene expression analysis was conducted with DESeq2 (Love et al., 2014) under the criteria of the absolute Log2 Fold change ≥1 and P < 0.05.

**Quantitative PCR analysis**

Total RNAs were reverse transcribed with Primerscript RT reagent kit with gDNA Eraser (Takara) according to the manufacturer's instructions and then subjected to quantitative PCR (qPCR) using SYBR Green Mix (Takara) and gene-specific primers (Table S7). Rice gene GAPDH was used as the reference to calculate the relative expression of investigated genes using comparative CT method 2^{-\Delta\Delta CT} (Livak and Schmittgen, 2001).

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658  J. Fan et al.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Fig. S1.** Expression changes of sucrose-starch and raffinose metabolism genes in rice stamens upon *Ustilaginoidea virens* infection. Differentially expressed genes (DEGs) (absolute Log2 Fold change ≥1 and $P$ value<0.05) were mapped to Sucrose-starch metabolism (A) and Raffinose metabolism (B) pathways with MapMan software. Red points represent up-regulated genes, while blue points represent down-regulated ones. Expression data are included in Table S5.

**Fig. S2.** QPCR validation of differentially expressed genes randomly selected from RNA-Seq analysis. Samples of rice stamens were collected from mock-inoculated (CK) and *U. virens* P4-infected (P4) spikelets, and subjected to qPCR analysis using rice GAPDH as the reference gene (Fan et al., 2015). Log2 Fold change of gene expression was presented. ‘-Inf.’ indicates that expression Os06g0229800 was detected in CK, but not in P4-infected stamens.

**Fig. S3.** Trehalose can be utilized by *Ustilaginoidea virens*. *U. virens* PJ52-2-5 was cultured at 28°C for 30 d in Czapek-agar medium supplemented with a single carbon source of sucrose, or trehalose. Bar size = 1.5 cm.

**Table S1.** Differentially expressed genes in rice stamens upon *Ustilaginoidea virens* infection (absolute Log2 Fold change ≥1 and $P$ value<0.05).

**Table S2.** Pathway enrichment analysis of differentially expressed genes in rice stamens upon *Ustilaginoidea virens* infection.

**Table S3.** Differentially expressed genes involved in biotic stress or encoding receptor kinases.

**Table S4.** Differentially expressed genes involved in metabolism pathways.

**Table S5.** Differentially expressed genes involved in sucrose-starch and raffinose metabolisms.

**Table S6.** Differentially expressed genes involved in hormone-related pathways.

**Table S7.** Primers used in this study.

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