Infection of the Western Flower Thrips, *Frankliniella occidentalis*, by the Insect Pathogenic Fungus *Beauveria bassiana*

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1. Introduction

The broad-host-range insect pathogen, *Beauveria bassiana*, has long been recognized as a potential biological insecticide that could be incorporated into integrated pest management (IPM) practices for the control of thrips [1]. Interest in the use of *B. bassiana* as a biopesticide has led to its emergence as a model system for examining fungal development, potential biological insecticide that could be incorporated into integrated pest management (>72 h post-infection). Real-time RT-PCR analyses of the expression of select genes implicated in virulence including the *Pr1* protease, beauvericin synthase, involved in the production of the secondary metabolite beauvericin, two cytochrome P450 monoxygenases implicated in cuticular hydrocarbon degradation, two multidrug efflux proteins, a periplin involved in lipid storage, and the Hog1 MAP kinase and protein kinase A signaling factors revealed discrete patterns of infection-time dependent expression. These data provide basic insights into the process of *B. bassiana* infection of *F. occidentalis*.

Keywords: *Beauveria bassiana*; *Frankliniella occidentalis*; biopesticides; pathogenesis; histology; gene expression; real-time RT-PCR
cuticle penetration are essential to the pathogenic nature of entomogenous fungi [8–10]. *B. bassiana* displays a wide host range allowing it to be used against vectors of human-disease-causing agents [11–14] and a wide range of agricultural pests [15].

The western flower thrips, *Frankliniella occidentalis*, is a highly destructive insect pest that can cause significant damage in greenhouse crops, a niche in which this insect is particularly adept at proliferating, especially as it displays a marked ability to develop resistance to a wide range of different chemical insecticides [1]. Because of its mixed diet, wide plant host range, and ability to transmit lethal plant viral diseases [16], *F. occidentalis* is regarded as an important economic pest [17–19]. In China, *F. occidentalis* was first discovered in greenhouses in Beijing in 2003 and is considered an invasive pest along with a number of other thrips species [20,21]. A significant amount of literature exists on attempts at using entomopathogenic fungi for control of *F. occidentalis*. Various formulations and the effects of environmental conditions and application methods (e.g., foliar versus soil) have been used to evaluate and develop programs for thrips control using insect pathogenic fungi [22–27]. These data suggest the high potential for effective control of thrips especially in greenhouse conditions. Although the histopathology of *B. bassiana* infection has been examined in a number of insects, e.g., *Peregrinus maidis*, *Plutella xylostella*, and *Carposina sasakii* larvae, much more limited information exists concerning the *B. bassiana-F. occidentalis* interaction and/or the contributions of specific fungal (virulence) genes and pathways to the infection process.

Here, we examined the infection of *F. occidentalis* by *B. bassiana* (strain BbYT12) that displayed high virulence towards the thrips. Scanning electron microscopy was used to examine a time course of fungal infection on the insect surface, including initial attachment, germination, and penetration of the host cuticle. Internal fungal growth including hyphal growth, hyphal body formation and proliferation, and the invasion of various host tissues by the fungus was followed via histological sectioning. The expression of a number of fungal genes including those involved in cuticle penetration, secondary metabolite synthesis, hydrocarbon and xenobiotic degradation, and cellular signaling was followed during discrete stages of the infection using Real-Time RT-PCR. These data form a basis for understanding the cellular and molecular factors involved in *B. bassiana* targeting of *F. occidentalis*.

2. Materials and Methods

2.1. Entomopathogenic Fungi and Insects

*B. bassiana* strains BbYT12, 13, and 14 were obtained from infected giant scale insects (*Drosicha corpulenta*, Kuwana). Fungal strains were cultured on SDAY (4% glucose, 1% yeast extract, 1% peptone, and 2% agar, pH 6.0 ± 0.1) at 25 °C for 7 d. Typically, conidia were harvested from the surface of the agar plates into a sterile 0.1% Tween-80 solution. Conidial suspension concentrations were determined using a hemocytometer and adjusted to 1.0 × 10⁸ conidia/mL with a 0.1% Tween-80 solution as needed. Female adults of *F. occidentalis* were reared on clover (*Trifolium repens* L.) at Qingdao Agricultural University. *F. occidentalis* were fed on kidney bean (*Phaseolus vulgaris* L.) and purple cabbage (*Brassica oleracea* L.) at 25 ± 0.5 °C and 50–60% relative humidity (RH) with a photoperiod of 16:8 (light:dark) in the laboratory.

2.2. Insect Bioassays Using *F. occidentalis*

Adult *F. occidentalis* were randomly placed into inside insect-rearing boxes containing purple cabbage as food. Adult *F. occidentalis* were inoculated with fungal spores by direct spray application of 1.5 mL of the conidial suspension using an Aldrich chromatography sprayer (deposits of sprayed conidia: 510–525 conidia/mm²). Control insects were treated with sterile 0.1% Tween-80. At least fifty *F. occidentalis* adults were selected and put into the insect-rearing box. Following inoculation, the insects were transferred to rearing chambers at 25 ± 0.5 °C and 85 ± 5% RH with a photoperiod of 16:8 (L:D). Every treatment contained three replicates.
2.3. Histopathological Observations

2.3.1. Light Microscopy

Frozen sections of *F. occidentalis* adults were collected 4, 8, 12, and 18 h after inoculation by the fungal conidia. For each observation period, approximately 30 insects were fixed in 10% formaldehyde buffer for 24 h. After fixation, the samples were dehydrated in a graded series (20 and 30%) of sucrose solutions and embedded in embedding medium. Embedded specimens were serially sectioned to 10 µm thicknesses by a manual freezing microtome, mounted on glass slides, and stained with hematoxylin and eosin. Samples were covered with neutral balata and observed under an Olympus microscope. Paraffin sections: At 24, 48, 72, 96, and 120 h post-inoculation, live and dead insects were selected from the samples and rinsed using phosphate-buffered saline (PBS). The rinsed samples were gradually dehydrated in a series of ethanol solutions (35, 55, 75, 85, 95, and 100% (v/v)) and xylene infiltrated (35, 55, 75, 85, 95, and 100%) for 20 min at each level. The samples were embedded in xylene:paraffin (1:1, v/v) for 24 h at 56 °C and embedded in paraffin for 48 h at 56 °C. The embedded specimens were serially sectioned into 8 µm slices using a Leica microtome. The sections were mounted on glass slides and stained using Grocott’s methenamine silver stain (counterstained with light green) [28]. Slides were enclosed in neutral balsam, and the slide preparations were observed under an Olympus microscope.

2.3.2. Scanning Electron Microscopy

At 1, 4, 8, 12, 18, and 24 h post-inoculation, approximately 15 insects for each observation period were collected and immersed in 4% (v/v) glutaraldehyde (pH 7.2, 0.2 M phosphate buffer) for 48 h at 4 °C. After rinsing three times with 0.2 M phosphate buffer, the rinsed samples were dehydrated in a gradient of ethanol solutions (30% to 100%) for 15 min at each level. Then, tert-butyl alcohol was used to displace the ethanol. Samples were dried by critical point drying, coated with a gold palladium film, and examined and photographed using a scanning electron microscope (7500F, JEOL, Tokyo, Japan).

2.4. Genes Expression Analyses by Real-Time Reverse Transcription RT-PCR

*F. occidentalis* were infected with *B. bassiana* conidial suspensions (1 × 10⁸ conidia/mL) as above. Samples of treated and control untreated adults were collected at 4, 8, 12, 18, 24, 48, 72, and 96 h post-infection (at least 50 adults in each sample). Three samples were used for each treatment at a given time point. Total RNA from a sample was isolated using Trizol reagent (Invitrogen, Carlsbad, CA, USA). Briefly, 1 mL of the Trizol reagent was added to 50–100 mg of tissues and homogenized using a mortar and pestle. After homogenization 0.2 mL of chloroform was added to the suspension and mixed by shaking. After 2–3 min, the sample was centrifuged (12,000 × g, 15 min, 4 °C) and the resultant RNA precipitated from the upper aqueous phase using isopropanol. Synthesis of cDNA and real-time RT-PCR were performed according to the manufacturer’s instructions (PrimeScriptHRT reagent Kit with gDNA Eraser, TaKaRa). A total of 12 genes related to adhesion, appressorium formation, and penetration were selected for expression profiling by RT-PCR. Actin was used as an internal control. Primer sequences for amplification of indicated genes are given in Table 1. All real-time RT-PCR experiments included three replicates and the experiments were performed using three biological replicates.

2.5. Statistical Analysis

All experiments were repeated at least three times. For real-time RT-PCR experiments, expression levels between the groups were analyzed by the 2^{-\Delta\Delta C_{T}} method [29]. Briefly, CT values provided from real-time RT-PCR experiments (triplicate samples for each data point) were analyzed for the change in expression using the equation, 
\[\Delta\Delta C_{T} = (C_{T,\text{Target}} - C_{T,\text{Actin}})_{\text{Time}x} - (C_{T,\text{Target}} - C_{T,\text{Actin}})_{\text{Time}0}.\] All data were analyzed by one-way analysis of variance (ANOVA) using SPSS19.0 and the LSD test to compare differences between treatments.
Table 1. Primers used in this research.

| Gene                                | Gene Accession | Sequence (5'-3')                | Sequence (3'-5')                |
|-------------------------------------|----------------|---------------------------------|---------------------------------|
| ATP-binding cassette transporter (Pdr1) | BBA_07660      | TCTGCCCCCTTCTTCTCTGTCATG         | AGAGCAGCCGCCGACCATAG            |
| ATP-binding cassette transporter (Pdr2) | BBA_08779      | CGACGAGAGCCAGCTTCATTTTCCCAG     | GACAGCCCCAGGGGACCATG            |
| Protein kinase A                     | BBA_05916      | CATCAGGCAAGTCCGTCCAG            | TGCTGCGTTCTATTTGTTG             |
| cAMP-dependent subunit              | BBA_00443      | AGACAGTGGCTCGGITTCCG            | TCTGGCCGACATCTCCATT             |
| Cuticle-degrading protease (Pr1)     | BBA_09727      | TAAAGGGACTCGACATGCTCA           | GGGGCATCGCTATCCATCTTTGAC        |
| Beauvericin biosynthetic protein     | BBA_10207      | CCGCTGTATCCCACGAAATG            | AAGAACGGTGACAGGGAGCA            |
| β-1,3 Glucan synthase               | BBA_08759      | CAACGTGGCTGCTCTGC              | CGGCGAAGTCTTGTGAG               |
| Perilipin homolog 1                 | BBA_01244      | TGAACGAGACGCAGGTCTC             | GAGCTGCTGGGCGAACGATG            |
| Mitogen-activated protein kinases    | GU566074       | ACCGCAACCCGAAACCACATT          | CCGCATATCTAACATCTCCTGCTGTC      |
| hog1                                | BBA_04705      | GCTTGCACGGCGCTGCG              | CAACCTCTGACGTGTCAGCTGACTG       |
| Cytochrome P450                       | BBA_05718      | CGAAGGGTAGCTGAGGTGC            | CGACATTTCTGGCAGCTCTGAAAG        |
| Monooxygenase CYP52 × 1              | BBA_04860      | ACCCTCCGCTACCCCATG            | CGGCGGCGTGACGTGTC               |
| Cytochrome c synthase                |                |                                 |                                 |
| Actin                               |                |                                 |                                 |

3. Results
3.1. Insect Mortality and Fungal Attachment to the Insect Body

Three different B. bassiana strains, BbYT12, 13, and 14, isolated from scale insects, were used in laboratory bioassays to determine the rate of mortality against F. occidentalis (Figure 1). For BbYT12, the calculated mean lethal time to kill 50% of the insect hosts (LT50) using 1 × 10^8 conidia/mL was determined to be 90 ± 15.12 h. Accurate LT50 values could not be determined for the other two strains due to lower mortality levels. As strain BbYT12 showed the highest virulence, it was used in all subsequent experiments.

![Figure 1. Insect bioassays. Time course of corrected mortality of F. occidentalis infected with B. bassiana. Data were subjected to one-way ANOVA for each isolate at a given time point. Means in each column within each isolate followed by the different letter indicate significant difference at p < 0.05. Error bars = standard deviation.](image-url)
regions of the body. With respect to the thrips’ head region, conidia could be seen attached to antennae, around the mouthparts, and surrounding the compound eye (Figure 2A–C). In a magnified view, several grooves, as well as surface roughness, could be observed around the conidia at the basal area of the head (Figure 2D). Fungal conidia were found attached throughout the body surface, although relatively higher concentrations could be seen at the thorax (Figure 2E), where they were deposited in areas near the setae and at the base of the legs (Figure 2F). Masses of conidia were also observed on the intersegmental folds and at the end of the abdomen (Figures 2G,H and 3A), with fewer conidia observed on the alinotum. High conidial densities were seen on the wings, particularly at the base of the wings (Figure 3B,C). Furthermore, conidia were distributed over the cuticle, on the surfaces of the feet and the propodium, and around the apotelus (Figure 3D–F). In addition, *B. bassiana* conidia were found to be able to adhere to the smooth surfaces around the setae (Figure 3G), and large numbers of conidia were observed between the pereiopoda and metapodes (Figure 3H).

**Figure 2.** Representative scanning electron micrographs (SEMs) of the fungal infection process. *B. bassiana* fungal conidia visualized: (A) between the ommatidia of the compound eye, (B) on the antenna, (C) around the mouthparts, (D) on the basal area of the insect head, (E) attached to the thorax, (F) around the base of the legs, (G) at the end of the host abdomen, and (H) on the intersegmental folds in the abdomen.
3.2. Conidial Germination and Penetration of the Insect Cuticle

Within 4 h of inoculation on the insects, B. bassiana germ tubes could be seen, often oriented such that they appeared to be in the process of penetrating the host cuticle. Within this time frame (4 h p.i.), ~83% of the conidia had germinated (Figure 4A and Table 2), with clear indications of some of the germinated conidia beginning to penetrate the integument (Figure 4B). In several instances “excavation” of the cuticle was apparent (Figure 4C). Within 8 h p.i., almost 100% of the conidia had germinated (Figure 4D). Most conidia produced a single, short germ tube, although bipolar germination could be seen in a sub-population of the germinated conidia (Figure 4E). At this time point (8 h p.i.), the proportion of cuticle-penetrating germ tubes/hyphae appeared to be >80% (Figure 4F). In some instances shriveled conidia could be seen, although whether this may be due to fixation is unknown (Figure 4I).
Figure 4. Representative SEM photographs of *B. bassiana* attachment, germination, and penetration of the *F. occidentalis* cuticle. (A) Fungal germ tube (gt) formation 4 h post-inoculation (p.i.), (B) fungal penetration of the integument, 4 h p.i., (C) fungal penetration of the cuticle, 4 h p.i., (D) conidial germination 8 h p.i., (E) bipolar conidial germination, 8 h p.i., (F) fungal cuticle penetration/penetrating conidia (pc), 8 h p.i., (G) conidial germination, 8 h p.i., (H) cuticle penetration, 8 h p.i., and (I) shriveled penetrating conidia.

Table 2. Quantification of discrete morphological infection stages of *B. bassiana* on the cuticle of *F. occidentalis*.

| Rate (%) | Time (Post-Inoculation, h) |
|----------|---------------------------|
|          | 4    | 8     | 12    | 18    |
| Germination | 82.69 ± 1.33 | 100 + 0 | 100 + 0 | 100 + 0 |
| Penetration  | 46.15 ± 1.26 | 82.26 ± 0.61 | 100 + 0 | 100 + 0 |
| Shriveling   | 0 + 0  | 51.61 ± 1.22 | 57.41 ± 0.71 | 67.38 ± 0.82 |

Note: approximately 15 insects for each observation period.

At the 8 h p.i. time point, few ungerminated conidia were detected on the cuticle surface (Figure 5A), whereas mucilage and penetrating structure began to be evident (Figure 5B). After 12 h, many conidia appeared blended into the cuticle due to the production of an apparent mucilaginous coat around the conidia and germ tube, and penetrating conidia were increasingly found (Figure 5C and Table 2). Intriguingly, at this time point ~55% of the conidia appeared shriveled (Figure 5D). A similar pattern was seen at the 18 h post-inoculation time points, with mucilage around penetrating conidia (Figure 5E,G). At 18 h p.i., most conidia appeared to be in the process of penetrating the cuticle, and the proportion of shriveled conidia increased to ~67% (Table 2).
Figure 5. Representative SEM photographs of *B. bassiana* germ tubes, penetration, and shriveled conidia on the surface of *F. occidentalis*. (A) *B. bassiana* long germ tubes (lgt), 8 h p.i., (B) fungal mucilage production, 12 h p.i., (C) penetrating conidia (pc), 12 h p.i., (D) shriveled conidia (sc), 18 h p.i., (E) fungal mucilage production, 18 h p.i., (F) penetrating and shriveled conidia, 18 h p.i., (G) shriveled conidia, 18 h p.i., and (H) fungal germ tubes, penetrant structures, and shriveled conidia, 18 h p.i.

3.3. Histopathology: Fungal Hyphal Body Proliferation Inside the Insect Host

Sectioning of *B. bassiana*-infected *F. occidentalis* revealed extensive invasion of the insect internal tissues. Transition from hyphal growth to the production of fungal hyphal bodies (yeast-like cells), was apparent inside the insect body and hemocoel at 12 h p.i., with an increased number of hyphal bodies seen at 18 and 24 h p.i. (Figure 6A). At 24 h post-inoculation, hyphal bodies could be seen surrounding and invading the insect fat body (Figure 6B). At 48 h p.i., a few dead *F. occidentalis* could be seen (~10%). Hyphal growth increased in dead insects as compared to those remaining alive, which had greater proportions of hyphal bodies (Figure 6C,D). Post-mortality, the entire bodies of insect cadavers appeared overrun with fungal hyphae (~72 h p.i., ~50% mortality). Within the 24–72 h time points, the insect midgut and the Malpighian tubules were filled with hyphal bodies (Figure 6E,F), and invasion of the alimentary canal (Figure 6G) and the reproductive system (Figure 6H) was evident. At 96 h p.i., in those insects still alive, hyphal bodies could be seen overgrowing the insect body (Figure 7A), and covering the midgut and the lateral...
oviducts (Figure 7B). In addition, hyphal invasion of the ovaries in both live insects and dead insects could be seen. Extensive hyphal growth covering the bodies of both cadavers, and in some instances living insects, could also be seen (Figure 7C).

**Figure 6.** Representative histopathological images of *B. bassiana* infection of *F. occidentalis*. (A) *B. bassiana* hyphal bodies (hb) inside *F. occidentalis*, 12 h p.i., (B) fungal hyphal bodies surrounding the insect fat body (fb), 24 h p.i., (C) fungal hyphae (hy) in the live insects, 48 h p.i., (D) hyphal network in dead *F. occidentalis*, 48 h p.i., (E) insect midgut (mg) filled with the hyphal bodies, 72 h p.i., (F) insect Malpighian tubules (Mt) with fungal hyphal bodies, 72 h p.i., (G) hyphal bodies in the insect alimentary canal (ac), 72 h p.i., and (H) fungal invasion of insect ovariole (ova) 72 h p.i.

**Figure 7.** Representative histopathological images of sagittal sections of *F. occidentalis* infected with *B. bassiana*, (96 h p.i.). (A) Extensive hyphal bodies proliferation in the insect body, (B) hyphal bodies in the lateral oviducts (lov), and (C) insect cadavers covered with hyphae.
3.4. Expression of Fungal Virulence-Related Genes during the Infection Time Course

In order to examine the expression of virulence-related genes in *B. bassiana* during infection of *F. occidentalis*, the expression of 12 genes was examined during the infection process (Table 1). These genes included the Pr1 protease, beauvericin synthase, cytochrome P450s CYP52X1 and CYP5293A, perilipin, cytochrome c synthase, the Hog1 MAPK, protein kinase A cAMP-dependent subunit, the multidrug efflux proteins, Pdr2 and Pdr5, and β-1,3 glucan synthase. After normalization to actin expression levels, candidate genes were separated into three groups based upon expression levels for scaling and visualization purposes. Beauvericin synthase and CYP52X1 showed the highest levels of expression at early infection time points, with beauvericin synthase expression induced ~35–40 fold 12 h p.i., and CYP52X1 ~20–25 fold 18 h p.i. (Figure 8A). Pdr5, the Pr1 protease, and β-1,3 glucanase expression levels similarly peaked at ~18 h p.i. (5–7-fold above the initial 4 h p.i. time point), with expression of the latter gene typically 2–3-fold induced (as compared to the 4 h p.i. time point) over most of the time course examined (Figure 8B). In contrast expression of CYP5293A was highest at the 12 and 24 h (peak) p.i. time points (3–6-fold), with lower levels seen at the other time points. Expression of the protein kinase A cAMP-dependent subunit peaked at ~72 h p.i., whereas the perilipin showed high expression from the 24 to 72 h p.i. time points (Figure 8C). Maximal expression of the Hog1 MAP kinase was seen at the 48 h p.i. time point, with cytochrome c synthase levels low throughout the time course.

![Figure 8. Time course of *B. bassiana* gene expression during infection of *F. occidentalis*. The expression of select fungal genes implicated in the infection process was quantified by real-time RT-PCR as detailed in the Materials and Methods section. Means in each column within each gene followed by a different letter indicate significant difference at *p* < 0.05. Relative gene expression represents the log fold change, 2^−ΔΔCT.*](image)
4. Discussion

Insect management using living biological control agents differs significantly from the use of chemical pesticides. Chemical pesticides tend to directly target specific physiological processes, resulting in death of the host in a relatively short period of time (min to h) after contact. Chemical pesticides tend to have broad non-target and potential detrimental human health and environmental effects. In contrast, microbial biological control agents, and insect pathogenic fungi in particular, have significantly lower non-target effects and are not pathogenic towards vertebrates, with overall lower environmental and human health risks [30]. However, most microbial agents can only optimally function under appropriate environmental and handling conditions, and the process of infection and mortality is on a longer time scale (days) as compared to chemical insecticides. Thrips are a major agricultural pest, and the use of chemical pesticides has led to the emergence of resistant populations [1]. Thus, the use of the entomopathogenic fungus B. bassiana has been investigated either alone or in combination with chemical pesticides for thrips control [31,32]. In order to gain a better understanding of the process of B. bassiana infection of thrips, we have performed a histological examination of the infection process and examined the expression of select fungal genes implicated in virulence during a time course of infection of F. occidentalis.

The initial attachment of fungal conidia to the surface of the insect body depends on a combination of hydrophobic interactions and consolidation of attachment [33,34]. Our data show that conidia of B. bassiana strain BbYT12 were capable of binding to any site on the cuticle of adult F. occidentalis, although some preference were observed particularly to the surfaces of the setae, intersegmental folds, and grooves, a finding similar to what has been reported by others [35]. Conidia were frequently trapped by the setae, possibly because the environment of the setae is more humid as a result of their association with respiratory transpiration processes [36,37] These conditions would be beneficial for conidial germination, thus increasing the activity of the fungus. The pathogenic activity of B. bassiana has been shown to be enhanced in high humidity when infecting Phaedon brassicae Baly (Coleoptera: Chrysomelidae) and Bemisia tabaci nymphs [38] and humidity has been identified as an important factor for targeting F. occidentalis [23,24]. A large number of conidia were also seen gathered near the compound eye and around the mouthparts, similar to what has been observed during B. bassiana infection of C. sasakii larvae. Consolidation of adhesion via production of mucilage is known to play an important role in the adhesion of fungal spores to a wide range of hosts including plants, animals, and insects [34,39,40]. For insect pathogenic fungi, mucilage was found around spores that were penetrating the integument [41]. Here, we also observed mucilage around adhering and penetrating conidia/germ tubes. Examination of the adhesion of Lecanicillium lecanii adhered to the epicuticle of thrips, aphids, and mealybugs, revealed the secretion of dense mucilage during attachment [42,43]. In addition to promoting tighter attachment, mucilage may also serve as a vehicle for delivery of cuticle-degrading enzymes [44].

Our data show that within 8 h of infection, the B. bassiana conidia had germinated, and germ tubes and penetrating hyphae could be seen as early as 4 h post-infection (p.i.). Over time, an increase in penetrating hyphae was observed as well as a “shriveling” of the conidia and penetrating germings. This shriveling was notable as a reduction in 20–50% of the size of the cells was seen as well as distortions in the overall cellular morphology. While it cannot be excluded that these changes were due to sample fixation effects, these observations appear to indicate important changes in either the overall stability of the infecting/penetrating cells and/or in membrane components. Whether this is a consequence of the infection process or due to stress/host defense responses remains to be determined. Overall, the progression of germination and penetration agree with reports of others [45], as well as instances of bipolar germination and some specialized routes of entry, e.g., germinating conidia on F. occidentalis wings and germ tube penetration of the
cuticle through sockets at the base of bristles/hair situated on the forewing veins, the latter similar to what has been seen for *B. bassiana* penetration of the cuticle of elaterid larvae [46].

Fungal hyphal body production and proliferation inside the host was apparent within 24 h p.i., rapidly spreading to other tissues after 24–36 h p.i., where extensive fungal growth could be seen. An examination of *Leptolegnia chapmanii* infection of *Aedes aegypti* indicated invasion of host fat body tissues within 1–2 d of infection [42,46], and in our study, hyphal bodies surrounding the *F. occidentalis* fat body and host melanization responses (blackening of tissues) could be seen as early as 24 h p.i. Fungal cells were capable of rapidly infecting additional tissue including the alimentary canal, Malpighian tubules, and muscle tissues. Over time (>36 h) hyphae and hyphal bodies could be seen colonizing ovaries and other tissues, similar to what has been reported for *Isaria fumosorosea* infection of *Bemisia tabaci* [47].

Here, we also examined the expression patterns of a number of fungal genes implicated in virulence over the time course of infection. Secondary metabolites are known to impact infection and the expression of beauvericin synthase was induced over 35-fold during the time course examined, with peak expression seen at 12 h p.i. These data indicate that production of at least some (insect) toxic secondary metabolites is transient and occurs early in the infection process. Similarly, a cytochrome P450 (*CYP52X1*), was also highly induced early (18 h p.i.), consistent with its projected role in assimilating host cuticular hydrocarbons and participating in penetration events [7,48]. Expression of the *Pdr5* multidrug efflux protein as well as the *Pr1* protease, and *β*-1,3 glucanase gradually increased 8–12 h p.i., with a peak of expression seen at 18 h p.i. Expression of the critical protease gene, *Pr1*, implicated in targeting the host cuticle, was consistent with its function as a cuticle-degrading enzyme, and the *Pdr5* membrane efflux protein may act as a means for detoxifying host toxic molecules present on the surface of the insect host. The *β*-1,3 glucanase could be functioning in the degradation of host compounds during penetration, although it is not clear what substrates it may be acting on. Alternatively, the enzyme may be important for cell wall remodeling of the fungus, during its dimorphic transition to the production of hyphal bodies once the cuticle has been breached [5,49]. The importance of a *B. bassiana* GPI anchored-glucanosyltransfaerase has been examined [50], however whether it acts in concert with the *β*-glucanase remains to be determined. Intriguingly, the expression of a second cytochrome P450, *CYP5293A*, was found to peak 24 h p.i., and although the functions and substrates of this enzyme remain unknown, it is interesting to speculate that distinct hydrocarbon assimilation pathways may be (sequentially) induced depending upon changes and/or access to different host lipids during the infection process. In terms of signaling, the expression of the protein kinase A (cAMP-dependent subunit) and the Hog1 MAP kinase were examined. Our data show a slight elevation of protein kinase A expression early (12 h p.i.), with a more robust responses at 72 h p.i. In contrast Hog1 MAP kinase expression levels were highest from 24–72 h p.i. These data indicate that expression levels of the two different fungal signaling pathway components may differentially change during the time course of infection, implying that distinct pathways are triggered at discrete stages. Overall, our data provide a systematic overview of the histopathology of *B. bassiana* infection of *F. occidentalis*, and provide new information concerning the expression of key molecular determinants involved in the infection process.

**Author Contributions:** J.W. and Z.Z. designed the study. Z.Z. carried out most of the work. J.W. and Z.Z. drafted the manuscript. J.W., Z.Z. and C.Z. analysed the data. N.O.K. helped address intellectual content, made critical suggestions for experimental improvement and helped draft the manuscript. Y.G. helped interpretation of data and gave critical suggestions for revising the manuscript. All authors have read and agreed to the published version of the manuscript.

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