Secretory proteins are delivered to the septin-organized penetration interface during root infection by *Verticillium dahliae*

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Abstract

Successful infection of the host requires secretion of effector proteins to evade or suppress plant immunity. Secretion of effectors in root-infecting fungal pathogens, however, remains unexplored. We previously reported that *Verticillium dahliae*, a root-infecting phytopathogenic fungus, develops a penetration peg from a hyphopodium to infect cotton roots. In this study, we report that a septin ring, requiring VdSep5, partitions the hyphopodium and the invasive hypha and form the specialized fungus-host interface. The mutant strain, VdΔnoxb, in which NADPH oxidase B (VdNoxB) is deleted, impaired formation of the septin ring at the hyphal neck, indicating that NADPH oxidases regulate septin ring organization. Using GFP tagging and live cell imaging, we observed that several signal peptide containing secreted proteins showed ring signal accumulation/secretion at the penetration interface surrounding the hyphal neck. Targeted mutation for VdSep5 reduced the delivery rate of secretory proteins to the penetration interface. Blocking the secretory pathway by disrupting the vesicular trafficking factors, VdSec22 and VdSyn8, or the exocyst subunit, VdExo70, also arrested delivery of the secreted proteins inside the hyphopodium. Reduced virulence was observed when cotton roots were infected with VdΔsep5, VdΔsec22, VdΔsyn8 and VdΔexo70 mutants compared to infection with the isogenic wild-type V592. Taken together, our data demonstrate that the hyphal neck is an important site for protein secretion during plant root infection, and that the multiple secretory routes are involved in the secretion.

Author summary

Pathogens secrete effector proteins as molecular weapons to evade or suppress plant immunity. However, the mechanism(s) by which root-infecting fungal pathogens secrete secretory effector proteins remains unexplored. We previously reported that *Verticillium dahliae*, a root-infecting phytopathogenic fungus, forms a specialized infection structure known as a hyphopodium that develops a penetration peg to pierce plant roots. In this study, we observed that after penetration, the penetration peg-developed hyphal neck, partitioning the hyphopodium and invasive hypha, came into close contact with the host, forming the fungus-host penetration interface. NADPH oxidase B (VdNoxB) regulated
the cytoskeletal organization of the septin ring at the hyphal neck. Importantly, the penetration interface was a preferential site for secretion of signal peptide-containing proteins. Septin plays an important role in the efficient delivery of secretory proteins to the penetration interface. Moreover, the conventional fungal ER-to-Golgi secretion pathway, endosome-mediated transport and the exocyst complex are involved in the delivery of secretory proteins to the penetration interface. Together, our data demonstrate that the *V. dahliae* infection structure functions as a key signaling hub during plant infection and is the apparatus that not only breaches host cells but also provides a unique interface for the secretion of fungal effectors.

**Introduction**

Pathogens secrete effector proteins as molecular weapons to evade or suppress plant immunity. Most effectors are small secreted proteins [1,2,3], and in many cases, the expression of these genes is induced by infection, helping the microbe to successfully colonize on the surface or inside of the host [1]. Studies of the secretion system have revealed diverse manners for pathogen effector translocation into their host. Fungi secrete different effectors at different infection stages from stage-specific compartments at the host-pathogen interface [4]. Before penetrating host cells, some effector proteins are focally secreted from appressorial penetration pores and may function to suppress early plant defense responses, as in case of several *Colletotrichum* species [4,5]. After invasive hypha developed, effectors in *Colletotrichum orbiculare* accumulate at the pathogen-plant biotrophic interface, a ring-like region around the neck of the biotrophic primary hypha [5]. Effectors of some plant fungal pathogens are putatively translocated into the host cell, where they interact with cytoplasmic or nuclear R proteins [6]. For example, *Magnaporthe oryzae* has a highly localized structure to accumulate cytoplasmic effectors secreted by invasive hyphae, known as the biotrophic interfacial complex (BIC), which forms at the tip of the initially filamentous hypha in the host cell [7,8].

Secretion of effectors to the host is also important for soil-borne fungal and oomycete pathogens, such as *Verticillium dahliae* and *Phytophthora sojae*, for successful infection [9,10]. Transit of many oomycete or fungal effectors to host cell depends only on the RXLR motif or other host-entry motifs of the effectors and host molecules, but not pathogen-encoded machinery [11,12]. However, however, the mechanism(s) by which root-infecting fungal pathogens secrete secretory protein remains unknown. *V. dahliae* causes vascular wilt disease [13,14] and infects more than 200 host species worldwide, including many economically important crops, such as lettuce, cotton and tomato [15]. *V. dahliae* contains more than 100 small cysteine-rich potentially secreted proteins [3]. So far, only two effectors, Ave1 and VdIsc1, have been functionally studied. Ave1 contributes to fungal virulence in the absence of its corresponding R protein (Ve1) [9]; it is a small secreted protein containing 134 amino acids (aa) with four cysteines [9], however, its secretion process has not been studied. VdIsc1 suppresses salicylate-mediated innate immunity *in planta* [10]. VdIsc1 is an unconventionally secreted protein as it lacks an N-terminal signal peptide that can direct the protein to the conventional secretory pathway [10]. Therefore, whether the soil-borne *V. dahliae*, a root-infecting phytopathogenic fungus, have a highly localized structure to secret secretory effector proteins remains unexplored.

We recently identified and provided the molecular features of the infectious structure, the hyphopodium, in *V. dahliae* [16]. We demonstrate that *V. dahliae* NADPH oxidase B (VdNoxB) is required for local reactive oxygen species (ROS) production during infection,
and ROS-Ca\textsuperscript{2+} signaling in the hyphopodium plays key roles in regulating polarized penetration peg formation and pathogenicity in \textit{V. dahliae} [16]. In \textit{M. oryzae}, the Nox2 (NoxB)-NoxR complex spatially organizes a heteroligomeric septin ring at the appressorium pore [17,18]. Septins, small morphogenetic guanosine triphosphatases (GTPases), are thought to reorient and reorganize the cytoskeleton to determine cell shape [19]. Septin ring scaffolds a toroidal F-actin ring and recruits and organizes the exocyst to the appressorium pore where the penetration peg emerges [17,18,20].

In this study, to explore whether the secretion of effectors in \textit{V. dahliae} could be associated with the penetration peg, we first verified that VdNoxB was required for the cytoskeletal organization of a septin ring at the penetration peg and its derived hyphal neck in \textit{V. dahliae}. We observed that the septin-ring-organized hyphal neck acts as a functional fungus-host penetration interface for the delivery and secretion of signal peptide-containing secretory proteins. Using gene functional analyses, we further showed that VdSep5, the conventional fungal ER-to-Golgi secretion pathway, the endosome-mediated transport, and the exocyst complex are involved in the delivery of secretory proteins to the penetration interface.

**Results**

VdNoxB-dependent penetration peg and its derived hyphal neck are repeatedly developed during \textit{V. dahliae} infection of plant roots

\textit{Verticillium dahliae} infection requires the development of an infectious structure, hyphopodium, in which the NADPH oxidase catalytic subunit VdNoxB is specifically expressed to regulate formation of the penetration peg to pierce the cell wall [16]. To further understand the infection process, we used FITC-WGA (FITC-conjugated wheat germ agglutinin) to label the fungal hyphae of wild-type \textit{V. dahliae} strain V592 to assess cellophane penetration and root infection. We observed repeated development of hyphopodium for penetration inside the cellophane and roots (Fig 1A and 1B, S1 and S2 Movies). Transmission electron microscopy (TEM) images showed that after penetration, the invasive hypha grew, and the penetration peg acted as a hyphal neck that partitioned the hyphopodium and invasive hypha and came into close contact with the cellophane (Fig 1C) or the host (Fig 1D), forming the fungus-host penetration interface. WGA staining showed various uneven hyphal neck lengths after piercing the cellophane or root cell wall (Fig 1A and 1B). The length of the hyphal neck reflects the length of the penetration peg and might be dependent on the piercing sites of different cells, such as the root epidermis or cortical cells, and the piercing angle. Repeated development of hyphopodium inside the cellophane was observed using V592 expressing GFP-tagged VdNoxB under the native promoter (S2A Fig) [16]. Inside the cellophane, the GFP signal was observed in flattened irregular hyphopodia and at the tips of the penetration pegs (S2B Fig). The VdΔnoxb mutant, in which VdNoxB was deleted, was incapable of producing a penetration peg (S2C Fig). Together, our results demonstrate that \textit{V. dahliae} infection requires VdNoxB-dependent, repeated development of the hyphopodium and penetration peg for each cell wall penetration, and the penetration peg-derived hyphal neck connects the hyphopodium to the invasive hypha and marks a site of close fungus-host penetration interface contact.

Organization of the cytoskeletal septin ring at the penetration peg and its derived hyphal neck

To explore the specific features of the penetration peg and its derived hyphal neck, we first examined whether the cytoskeleton protein septin plays a role in determining penetration peg morphogenesis in \textit{V. dahliae}. The \textit{V. dahliae} homolog of Septin5 was identified (S1B Fig) and
named VdSep5. VdSep5-GFP was expressed in the V592 and VdΔnoxb mutant. In the wild-type V592 hyphopodium during penetration peg induction, confocal laser scanning microscopy (CLSM) observation revealed a continuous funnel-shaped VdSep5-GFP fluorescent signal from the base of the hyphopodium, outlining the curved contact area between the hyphopodium and cellophane membrane (Fig 2A, plane 1.5 μm), to the central protruded zone (refer to the hyphopodium pore) where the penetration peg was initially developed and spread throughout to its tip (Fig 2A, from plane 2.7 to 4.5 μm). In contrast, in the VdΔnoxb hyphopodium, the VdSep5-GFP signal outlined the curved contact area between the hyphopodium and the cellophane membrane but without the central septin ring at the base of the
hyphopodium (Fig 2B). In wild-type V592, compared with the widest part of the VdSep5-GFP signal area (Fig 2A, plane 1.5 μm), the diameter of the VdSep5-GFP ring at the hyphopodium pore (2.7 μm) was reduced approximately 58%, while there was no clear reduction of the diameter of the VdSep5-GFP signal area in the VdΔnoxb mutant, which is deficient in penetration peg formation (Fig 2C). After cellophane piercing and invasive hyphal growth, we observed the compact septin ring signal retained in the hyphal neck (Fig 2D). These results demonstrate that VdNoxB is required for VdSep5 organization of the cytoskeleton to determine the morphogenesis of the penetration peg and its derived hyphal neck.

After infecting Arabidopsis root with wild-type V. dahliae V592, a VdSep5-GFP ring was also observed at the hyphal neck partitioning the hyphopodium and invasive hyphae (Fig 2E). Two VdSep5-GFP rings were observed in the first and second hyphal necks in two CLSM planes within the same scanning view (Fig 2F, 3.6 μm for the first penetration and 13.2 μm for the second penetration), verifying the requirement of multiple penetrations for each new cell wall to reach the vascular bundle. Together, our results clearly demonstrate that septin-ring organization accompanies V. dahliae penetration of either cellophane or plant roots, and
VdNoxB plays a role in the organization of the septin ring at the penetration peg and its derived hyphal neck. Remarkably, once established, the VdSep5-GFP ring was retained at and framed the hyphal neck, forming the fungus-host penetration interface.

Consistent with previous findings that septin scaffolds a toroidal F-actin ring at the appressorium pore in *M. oryzae* [17,18], we also observed that F-actin was organized as a ring structure at the hyphal neck in either cellophane or root by live-cell imaging of V592 expressing LifeAct-GFP (S3 Fig).

**Secretory proteins are preferentially localized at penetration interfaces**

Next, we investigated whether the septin-ring-organized hyphal neck, in addition to its piercing role, could act as a functional fungus-host penetration interface for the delivery of secretory proteins. The arsenal of potentially secreted proteins in plant pathogens includes key pathogenicity molecules that are generally referred to as effectors (small cysteine-rich proteins, < 400 amino acids (aa) and ≥ 4 cysteine residues) [3]. Because the well-known Ave1 secreted effector has not been identified in the cotton isolate V592, we selected three small cysteine-rich proteins (SCP), VDAG_08085 (194 aa, 6 Cys, named VdSCP8), VDAG_00902 (375 aa, 16 Cys, VdSCP9) and VDAG_05717 (205aa, 4 Cys, VdSCP10), for analysis in this study. Each of these SCPS has an N-terminal signal peptide predicted by the SignalP 4.1 server (S4A Fig) [21]. Among them, VdSCP9 is a LysM domain-containing protein. The LysM effector family contains relatively conserved secretory proteins that are known to play significant roles in the pathogen-host interaction [3,22,23,24]. VdSCP8 was identified by liquid chromatography-mass spectrometry (LC-MS) of the V592 culture filtrate, and VdSCP10 was one of the potential pathogenicity genes in our previous screening of the T-DNA insertional mutant library [25] and confirmed by the targeted gene replacement mutant of VdSCP10 (S5 Fig). Transcript levels of these SCP genes were first examined using quantitative RT-PCR (qRT-PCR). The expression levels of *VdSCP9* and *VdSCP10*, but not *VdSCP8*, were significantly up-regulated at 4 days post-inoculation (dpi) of V592 on cellophane and at 2 dpi on cotton roots (S4B Fig). To observe the localization of these SCPs, VdSCP8-GFP, VdSCP9-GFP and VdSCP10-GFP were expressed under the native promoter. Only VdSCP8-GFP fluorescence was detectable as a ring signal at the penetration zone on cellophane (S4C Fig). Neither VdSCP9-GFP nor VdSCP10-GFP fluorescence was observed on cellophane. Therefore, these GFP fusion proteins were constructed under the oliC promoter. The *V. dahliae* small effector VdIs1 (190 aa, 1 Cys), which lacks a signal peptide and exhibits characteristics that lead to unconventional secretion [10], was also fused to GFP as a control. After *V. dahliae* invasion into cellophane, VdSCP8-GFP, VdSCP9-GFP and VdSCP10-GFP, but not the control VdIs1-GFP, showed ring signals surrounding the penetration zones (Fig 3A). From a picture of the penetration at an incline, the VdSCP10-GFP ring signal was clearly observed surrounding the hyphal neck linking the hyphopodium to the invasive hypha (Fig 3B). We also fused the signal peptides of SCPs to GFP and found that SPVdSCP8-GFP, SPVdSCP9-GFP and SPVdSCP10-GFP also showed ring signals outside the plasma membrane of the penetration zones (S4D Fig). To detect whether the SCP-GFP signal rings overlapping with the septin ring, VdSep5-RFP was co-expressed with VdSCP8-GFP in V592, and the results showed that the VdSep5-RFP ring was inside the VdSCP8-GFP ring (Fig 3C). Similar results were obtained for VdSCP9-GFP and VdSCP10-GFP, demonstrating that signal peptide-containing SCPS accumulate and/or are delivered to the hyphal neck for secretion. Together, these data suggest that the hyphal neck made up a fungus-host penetration interface for the delivery and/or exportation of secretory proteins.

Next, we inoculated strains of V592 expressing the GFP-tagged SCPS on *Arabidopsis* roots. VdSCP8-GFP accumulation was first observed at approximately 2 dpi. A strong VdSCP8-GFP
signal ring was observed at the hyphal neck, which partitioned the hyphopodium and the invasive hypha (Fig 4A and 4B). Some weak signals were also observed inside the invasive hyphae (Fig 4A and 4B). VdSCP9-GFP accumulation was first observed at 1 dpi. One weak and one strong VdSCP9-GFP signal ring was observed, respectively, at the first and second hyphal neck in two CLSM planes in the same scanning view (Fig 4C at planes of 1.8 µm and 4.8 µm).
observation is consistent with the requirement of the repeated development of the hyphopodium for each cell wall penetration during the colonization of V592 from the root surface to the vascular bundle (Fig 1B). The VdSCP10-GFP ring signal was observed at approximately 6 dpi. In the upper plane of the hyphopodium (0 μm), a weak VdSCP10-GFP speckle signal was observed at the periphery of the hyphal cell and on two sides of a septa (Fig 4D). A stronger signal was observed at the apex of the hyphopodium (Fig 4D). The clear and strongest VdSCP10-GFP ring signal was observed at and throughout the hyphal neck using a series of continuous scanning planes (Fig 4D). Taken together, our data demonstrate that successful invasion of plant roots and cellophane by V. dahliae has the common phenomenon of signal peptide-containing secretory protein recruitment at the hyphal neck for effective secretion through this fungus-host penetration interface. The detectable ring signals for each secretory protein at different time points suggest that their synthesis and/or rate of delivery were different, revealing a complex process for successful infection in plant roots by V. dahliae.

We then examined whether the directional ring-shaped accumulation of the small secretory proteins was derived from dynamic secretion towards the penetration interface. Fluorescence recovery after photobleaching (FRAP) was performed with the V592 strain expressing VdSCP10-GFP on cellophane at 6 dpi. We photobleached VdSCP10-GFP fluorescence at the...
penetration site and then monitored the fluorescence recovery over time. After near complete elimination, fluorescence recovered within 27 min (Fig 3D and 3E). The fluorescence recovery time for VdSCP8-GFP and VdSCP9-GFP was 22 min and 18 min, respectively (Fig 3F). These data suggest that secretory proteins were continuously synthesized and/or delivered to the penetration interface.

**VdSep5 plays a role in the delivery of protein secretion toward the penetration interface**

To detect whether septin also plays a role in the delivery of secretory proteins to penetration interfaces, the targeted gene knockout mutants VdΔsep5 and VdΔsep3 were generated in wild-type V592 (S6A and S6B Fig). The VdSep3 homologous sequence was searched from V592 based on a BLASTP search using MoSep3 and the VdLs.17 database (S1A Fig). The VdSep3 knockout mutant strain exhibited a reduced hyphal growth rate on PDA medium compared with V592 (S6D Fig), and developed an abnormal hyphopodium on cellophane without smooth swelling at the end of branching hypha (S6E Fig) that was incapable of forming a penetration peg to pierce the cellophane (S6E Fig). This result demonstrates that VdSep3 plays roles in hyphal growth and hyphopodium development. In contrast, the VdSep5 knockout mutant strain exhibited a normal growth rate on PDA medium (S6D Fig) but developed fewer hyphopodia on cellophane (S6F and S6G Fig) and displayed greatly delayed penetration of the cellophane compared with V592 (S6E Fig), demonstrating that VdSep5 plays a role in hyphopodium development. Consistently, both the VdΔsep3 and the VdΔsep5 mutant showed reduced virulence on cotton plants (S6H and S6I Fig). The reintroduction of Psep3::VdSep3::TIRPC and Ptef::VdSep5-GFP::TIRPC restored the hyphal morphologies and cellophane penetration abilities, as well as the pathogenicity (S6D, S6E, S6H and S6I Fig), confirming the targeted gene deletion. Our results suggest that VdSep5 plays an important role in the initiation of hyphopodium formation, whereas, VdSep3 is more important for development of the hyphopodium.

To observe the localization of secretory protein in the VdΔsep5 mutant, the targeted gene knockout mutant was generated in VdSCP10-GFP-expressing V592 strain. VdSCP10-GFP secretion in the VdSep5 deletion mutant was assessed. In contrast to the remarkable VdSCP10-GFP signal ring surrounding the hyphal neck in the wild-type V592 (Fig 5A), the VdSCP10-GFP signal was observed in both the hyphopodium and hyphal neck in the VdΔsep5 mutant, in which either the hyphopodium or the hyphal neck was stained with FM4-64 (Fig 5A and 5B). The VdSCP10-GFP signal at the hyphal neck in the VdΔsep5 mutant was clearly reduced compared with that in the wild-type V592 background (Fig 5B). The average signal intensity of VdSCP10-GFP in the hyphal neck of VdΔsep5 was approximately 78% of that in V592 (Fig 5C). The FRAP assay showed 86% recovery of VdSCP10-GFP fluorescence at the penetration interface within 97 min (Fig 5D and 5E), which was significantly longer than the recovery time of 27 min for the wild-type V592. Three FRAP tests on cellophane showed a similar delayed in secretion. These results demonstrate that VdSep5 plays a role in mediating the delivery of secretory proteins to the penetration interface, in addition to its functions in hyphopodium formation and cortical structure organization of the penetration peg and hyphal neck.

The vesicular trafficking factor SNAREs VdSec22 and VdSyn8 are involved in protein secretion toward penetration interfaces

We next investigated the role of vesicular traffic in the delivery of secretory proteins to penetration interfaces. SNAREs function as key elements in membrane fusion [26,27,28]. The R-SNARE Sec22 is important for modulating transport between the ER and the Golgi
The Qc-SNARE Syn8 in *S. cerevisiae* and *M. oryzae* (MoSyn8) localizes at endosomes and/or late endosome/prevacuolar compartments (PVCs) [26,28]. To identify functional proteins in the secretion of *V. dahliae*, homologous sequences were searched in V592 based on a BLASTP search using MoSec22 and MoSyn8 and the database for VdLs.17, designated VdSec22 and VdSyn8, respectively (S1C and S1D Fig). Targeted gene knockout mutants VdΔsec22 and VdΔsyn8 were generated (S7A and S7B Fig). Both mutants exhibited growth defects with a reduced vegetative hyphal growth rate; VdΔsyn8 also showed reduced melanin production (S7D Fig). The reintroduction of VdSec22 and VdSyn8 under the control of each native promoter recovered the growth ability and hyphal morphologies (S7C and S7D Fig), confirming the targeted gene deletion of VdSec22 and VdSyn8.

VdSCP10-GFP secretion in the VdΔsec22 and VdΔsyn8 mutants was assessed. VdSCP10-GFP expressed under the oliC promoter was transformed into VdΔsec22 and VdΔsyn8 mutants. The single copy insertion strains determined by Southern blot were used for further analysis (S8 Fig). In contrast to the remarkable VdSCP10-GFP signal ring in wild-type V592 (Fig 6A), the VdSCP10-GFP signal was observed in both the hyphopodia and hyphal necks in both deletion mutant strains (Fig 6B and 6C). The VdSCP10-GFP signal in the hyphal neck in
both mutants was also clearly reduced, and most of the VdSCP10-GFP signal rings were overlapping with or inside the plasma membrane compared with that in the wild-type V592 background (Fig 6B and 6C). The average signal intensity of VdSCP10-GFP in the hyphal neck of VdΔsec22 and VdΔSyn8 was approximately 54% and 70%, respectively, of that in V592 (Fig 6E), suggesting that ER-Golgi transport is a predominant route of transport of SCPs. The FRAP assay was also performed with VdSCP10-GFP-expressing VdΔsec22 and VdΔSyn8 mutants. Fluorescence was recovered after approximately 52 min and 45 min in VdΔsec22 and VdΔSyn8 mutants, respectively (Fig 6F), which was much longer than the recovery time of 27 min observed for wild-type V592. Our data demonstrate that VdSec22 and VdSyn8 play roles in mediating the delivery of secretory proteins to the penetration interface.

Fig 6. Deletion of VdSec22, VdSyn8 or VdExo70 decreased secretory protein delivery to the hyphal necks. (A) VdSCP10-GFP signal ring surrounding the hyphal neck in wild-type V592. (B–D) The mutant strains VdΔsec22 (B), VdΔSyn8 (C) and VdΔexo70 (D) retained most of the VdSCP10-GFP inside the hyphopodium and reduced the VdSCP10-GFP signal in the hyphal neck. The plasma membrane of HP and HN was stained with FM4-64 (red). (E) Quantitative analysis of the effect of VdSec22, VdSyn8, and VdExo70 on secretory protein delivery to penetration interfaces. More than 30 hyphal necks with a visible “ring” signal were investigated for each VdSCP10-GFP-expressing mutant strain to determine the intensity of the ring signal at the HN. The mean and SD for (E) were calculated from three independent fungal transformants for each mutant (*P<0.05; t-test). Bar = 2.5 μm. (F) FRAP assay for the dynamic accumulation/delivery of VdSCP10-GFP at the hyphal neck in VdΔsec22, VdΔSyn8 and VdΔexo70 on cellophane. Three FRAP tests were performed for each mutant strain.
To further determine whether the VdSCP10-GFP signal in the hyphopodium and hyphal neck was due to decreased transport from the ER to the Golgi apparatus in the mutant strains, a VdSCP10-GFP-expressing VdΔsec22 mutant on cellophane was stained with ER-Tracker Blue-White DPX. The VdSCP10-GFP signal was observed to overlap with the ER in the hyphopodium (S9 Fig), suggesting that the deletion of VdSec22 resulted in retention of VdSCP10-GFP in the ER.

Taken together, our data demonstrate that the transport route between the hyphal ER and Golgi apparatus and endosome-mediated transport are involved in protein secretion toward penetration interfaces.

Efficient secretion of secretory proteins at the penetration interface requires the exocyst complex

The exocyst was discovered as a tethering complex that mediates the initial encounter of arriving exocytic vesicles with the plasma membrane [30]. The exocyst complex is an evolutionarily conserved doctameric protein complex comprising Sec3, Sec5, Sec6, Sec8, Sec10, Sec15, Exo70, and Exo84 [31,32]. To test the role of the exocyst in the accumulation of small secretory proteins at the penetration interface in V. dahliae, two predicted exocyst components, VdSec8 and VdExo70, were identified (S1E and S1F Fig). VdSec8-GFP and VdExo70-GFP were expressed under either their native promoter or the oliC promoter and introduced into V592. Similar localization profiles were observed for both GFP-tagged proteins under either the native or the oliC promoter; however, the GFP signal derived from the native promoter was weak, and thus the fluorescence signals derived from the oliC promoter were photographed. Both VdSec8-GFP and VdExo70-GFP were observed as a crescent structure at the growing tip of vegetative hyphae (S10A Fig). VdSec8-GFP and VdExo70-GFP were organized as a ring at the base of the hyphopodium that was observed before penetration peg formation on either cellophane or Arabidopsis root (S10B and S10C Fig). After the development of invasive hyphae, VdSec8-GFP was organized at the hyphal neck on either cellophane or root (S10D Fig). Together, our data demonstrate that the exocyst is active at the base of the hyphopodium and the hyphal neck.

To characterize the localization relationship between the exocyst complex and VdSep5, VdSep5-RFP was transformed into the V592-expressing Polic:VdSec8-GFP:Ttrpc strain. Red septin rings were observed in all 20 observed hyphal necks, and VdSec8-GFP signal rings were observed in 14 of the detected septin rings. The corresponding linescan confirmed the co-localization of VdSec8-GFP and VdSep5-RFP (S11 Fig).

To further study the role of exocyst subunits on secretory protein accumulation at the penetration interface, we tried to knockout VdExo70 and VdSec8 in V592. VdΔexo70 mutants carrying the VdExo70 deletion were obtained (S7B Fig), but the deletion of VdSec8 was not successful, in agreement with a previous study in which M. oryzae exocyst-encoding gene knockouts generated only Δsec5 and Δexo70 mutants [20,33]. Thus, the failure to delete VdSec8 was possibly due to the lethality of the absence of Sec8 in filamentous fungi including V. dahliae and M. oryzae. The VdΔexo70 mutant exhibited growth defects with a low growth rate on PDA medium (S7D Fig). The reintroduction of Polic:VdExo70-GFP:Ttrpc into the VdΔexo70 mutant recovered the growth ability and hyphal morphologies (S7C and S7D Fig), confirming the targeted disruption of VdExo70. VdSCP10-GFP was then transformed into the VdΔexo70 mutant and incubated on cellophane for hyphopodium induction. The single copy insertion strains determined by Southern blot were used for further analysis (S8 Fig). The VdSCP10-GFP signal was observed inside of the hyphopodium in the VdΔexo70 mutant (Fig 6D). Weak signals were observed in the hyphal neck, but most of them overlapped with the FM4-64-stained
plasma membrane ring (Fig 6D). The average intensity of the green fluorescence ring of 
VdSCP10-GFP in the VdΔexo70 mutant was approximately 65% of that in V592 (Fig 6E). These 
results demonstrate that VdExo70 plays a role in secreting VdSCP10-GFP out of the hyphal 
neck. The FRAP assay on cellophane was also performed with VdSCP10-GFP in the VdΔexo70 
mutant. The fluorescence recovered after 63 min (Fig 6F), which was significantly longer than 
the recovery time of 27 min determined for wild-type V592. Taken together, our results demon-
strate that exocyst components also organize at the hyphal neck and take part in the delivery of 
secretory proteins to penetration interfaces.

Proteins involved in the secretion pathway play critical roles in the 
pathogenicity of V. dahliae

To explore the roles of proteins involved in the secretion pathway in the pathogenicity of V. 
dahliae, we inoculated VdΔexo70, VdΔsec22 and VdΔsyn8 mutants on cotton plants and found 
a significant (P< 0.05) reduction in the disease index for the three mutants (Fig 7A and 7B). 
The loss of virulence was restored when the VdΔexo70, VdΔsec22 and VdΔsyn8 mutants were 
complemented with Polic:VdExo70-GFP:Ttrpc, Psec22:VdSec22:Ttrpc or Psyn8:VdSyn8:Ttrpc, 
respectively (Fig 7A and 7B). The significant loss of pathogenicity for the VdΔexo70, VdΔsec22 
and VdΔsyn8 mutants was presumably consistent with their inefficient secretion of effector-
related secretory proteins, which are required for successful fungal pathogen infection by evad-
ing or suppressing host plant immunity. Therefore, attributed to critical roles in the efficient 
secretion of secretory proteins at fungus-host penetration interfaces, VdExo70, VdSec22 and 
VdSyn8 play important roles in the pathogenicity of V. dahliae.

Discussion

Septin assembly at the penetration peg-derived hyphal neck framing an 
interface between V. dahliae hypha and the host

Penetration of the intact cuticles of the host is a very important step for successful infection by 
phytopathogens, either for leaf- or root-infecting fungal pathogens, such as M. oryzae and the 
anthracnose disease-causing Colletotrichum species or V. dahliae [4,5,16,17,34]. The foliar fun-
gal pathogen M. oryzae forms conspicuous melanized appressoria with an average diameter of 
8.0 μm when it inflates to full turgor and develops penetration pegs with an average diameter 
of 780-nm to breach the hydrophobic, waxy leaf cuticle [18,35]. In contrast, we found herein 
that the root-infecting fungus V. dahliae developed hyphopodia with an average diameter of 
3.4 μm and formed a penetration peg with an average diameter of 1.3-μm, suggesting that less 
pressure is needed for V. dahliae to breach the root cuticle (Fig 1C and 1D).

We observed that the nature of the interface between V. dahliae hyphae and the host is the 
penetration peg-derived hyphal neck, in which a septin ring was organized. In M. oryzae, sep-
tsins are found to provide the cortical rigidity and membrane curvature necessary for protru-
sion of the rigid penetration peg to breach the leaf surface [18]. Similarly, we found that the 
septin ring framed a recognizable cytoskeletal region of the hyphal neck in which F-actin was 
also organized as a ring structure, partitioning the hyphopodium and invasive hypha on both 
cellophane and roots. On cellophane, we also observed a funnel-shaped septin structure prior 
to invasive hyphal growth, suggesting that V. dahliae septins also function in the membrane 
curvature necessary for protrusion of the penetration peg at the base of the hyphopodium. In 
the VdNoxB knock out mutant, the VdSep5-GFP signal at the base of the hyphopodium sug-
gests that septins provide membrane curvature, but the mutant strain failed to show protrusion 
of the penetration peg in the absence of VdNoxB. Previous studies in yeast and in fungal
pathogen *Aspergillus fumigatus* suggest the importance of septin phosphorylation/dephosphorylation in controlling septin assembly [36,37]. In yeast, Rts1, a protein phosphatase 2A (PP2A) subunit, regulates septin dephosphorylation during telophase, and this dephosphorylation contributes to cytokinesis [36]. Dephosphorylation of the core septin, AspB, in a PP2A-dependent manner also impacts hyphal septation in *A. fumigatus* [37]. In animals, PP2A is a well-known tumor suppressor. ROS accumulation in cancer cells causes nitration and inactivation of PP2A, which interferes with the interaction of Bcl-2 with the PP2A catalytic core, leading to increased phosphorylation and antiapoptotic activity of Bcl-2 [38]. We recently reported that
V. dahliae VdNoxB is required for local ROS production during infection and plays key roles in regulating polarized penetration peg formation [16]. Together with the regulated synthesis of ROS by M. oryzae Nox complexes directly control septin and F-actin dynamics [17], and the septin ring assembles in a kinase Chm1-dependent manner [18], we speculate that fungal Nox-dependent ROS might also play a role in inactivation of PP2A-like phosphatase, leading to increased Chm1-dependent septin phosphorylation, which is key for controlling septin assembly.

We speculate that septins also provide membrane curvature for polarity determination during penetration peg development on roots, although funnel-shaped septin signal was barely observed in the hyphopodium-penetration peg on the infected root, presumably due to a fast piercing process on the roots. The targeted gene deletions of VdSep5 or VdSep3 exhibited defects in hyphopodium and/or hyphal development, suggesting that core V. dahliae septins also act cooperatively to form heteroligomers during hyphal growth and infection. This result is consistent with previous observation in M. oryzae that septins formed rings at the neck of nascent appressoria and a wider range of structures in hyphae and during invasive growth, including bars, gauzes, collars and rings [18], in addition to an appressorium pore-located large septin ring [18]. Nevertheless, our data demonstrate the requirement for VdNoxB-dependent ROS in the regulation of cytoskeleton septin ring remodeling at the base of the hyphopodium, leading to rapid polarized growth of the penetration peg in V. dahliae. Each occurrence of penetration requires septin ring organization at the penetration peg and hyphal neck, supporting that successful colonization of extracellular hyphae to the vascular bundle requires repeated development of the hyphopodium and penetration peg, which repeatedly form penetration interfaces between V. dahliae hyphae and the host.

The hyphal neck-associated penetration interface as the site of delivery of secretory proteins

Plant infection by pathogens involves the deployment of effector proteins that suppress plant immune responses and facilitate proliferation of the pathogen within plant tissues [30,35]. The delivery of effectors has been shown by extra-invasive hyphal membrane (EIHM) and BIC in the first-differentiated bulbous invasive hyphae in M. grisea [7,33]. In C. higginsianum, sequential delivery of host-induced effectors by the appressorium pore and intracellular hyphae has been observed [4]. In C. orbiculare, the accumulation of effectors occurred in the ring-like region around the neck linking the penetration peg to the biotrophic primary hyphae [5]. In this study, we found that during the penetration of cellophane or plant roots, the tested SCP-GFP and SP-GFP accumulated on the penetration interfaces, indicating a general role of the penetration interface as an active secretory protein delivery zone in V. dahliae. The FRAP assay revealed the dynamic accumulation of SCPs at the penetration interfaces. Secretion of the three SCPs into the hyphal neck is likely not dependent on the biological host. However, the cellophane membrane was used to mimic the hydrophobic niche for induction of appressoria in M. grisea [39] and hyphopodia in V. dahliae [16]. Together with the identification of VdSCP8 by LC-MS of the V592 culture filtrate without any treatment, and transcripts of VdSCP9 and VdSCP10, but not VdSCP8, were induced upon incubation of V. dahliae on both cellophane and roots, we speculate that both VdSCP9 and VdSCP10 are probably in planta-expressed secretory proteins in V. dahliae. Moreover, the targeted gene deletion of VdSCP10 caused a significant decrease in virulence toward cotton plants, suggesting that VdSCP10 may function as an effector to suppress plant immune responses. Although the LysM effector family contains relatively conserved secretory proteins that are known to play significant roles in the pathogen-host interaction [3,22,23], it has been recently reported that deletion of the VdSCP9
homologous core LysM protein, Vd4LysM, in *V. dahliae* strain JR2, did not compromise virulence during infection in *Arabidopsis*, tomato or *Nicotiana benthamiana* [24]. Whether VdSCP9 and VdSCP8 function as effectors to suppress plant immune responses or facilitate proliferation of *V. dahliae* within plant tissues requires further investigation. Remarkably, the ring signals of the tested SCPs were outside and around the hyphal neck and septin ring (Fig 3), and they were reduced in the hyphal neck in the VdΔsep5 mutant. These observations indicate that septins are not only required to organize the hyphal neck to form a fungus-host interface but also participate in the delivery and exportation of secretory proteins.

**Delivery of secretory proteins to the penetration interface via vesicular trafficking coupled with exocytosis**

Phytopathogenic fungi express numerous small proteins that possess classical N-terminal signal peptides that direct them to the endoplasmic reticulum (ER) [3,4,40]. The three signal peptide-containing SCPs, but not the unconventional secretion protein VdIscI, accumulated around the hyphal neck, suggesting that secretion into the penetration interface depends on ER processing. The retention of VdSCP10-GFP in the ER of the VdΔsec22 mutant demonstrates the importance of transport between the ER and the Golgi apparatus in secretory protein delivery to penetration interfaces in *V. dahliae*. In *M. oryzae* during the invasion of rice cells, ER-to-Golgi trafficking is involved in the secretion of apoplastic effectors by EIHMM [33]. The Δsec22 mutants of *C. orbiculare* also show a decreased accumulation of effectors at biotrophic interfaces [5]. Similar to *M. oryzae* and *C. orbiculare* [5,41], the absence of Sec22 weakens the virulence of *V. dahliae* (Fig 7), suggesting that conventional ER-Golgi transport has a conserved function in the transport of some pathogen secretory proteins to interact with host molecules.

Endosomes participate in endocytosis and secretion during fungal infection in the host [28,42]. The long-distance retrograde motility of early endosomes is necessary to perceive plant cues and trigger the transcription of effector-coding genes during plant infection by the pathogenic fungus *Ustilago maydis*, which regulates effector production and secretion during host cell invasion [43]. Syn8 in *M. oryzae* is involved in the secretion of BIC-localized AVR proteins but not the apoplastic effector (Bas4) *in planta* [28]. In VdΔsyn8 mutants, the retention of VdSCP10-GFP in the hyphopodium and inside the hyphal neck, suggests that the induction of SCP delivery in *V. dahliae* requires cues from the fungus-contacting surface and that VdΔsyn8 mutants prevent the perception of information from the contact surface, and resulting in VdSCP10-GFP retention. Thus, effective delivery of secretory proteins during fungal infection in the host requires Syn8-mediated transport/cue-sensing via endosomes.

The final steps of the secretory pathway, which occur in the vicinity of the plasma membrane, are regulated by an array of small GTPases, the exocyst tethering complex, and SNARE proteins [20,30,44]. Co-localization of VdSec8-GFP and VdSep5-RFP and the absence of VdSep5 or VdExo70 to impair the delivery of secretory proteins to the penetration interface also support septin-dependent assembly of the exocyst in *V. dahliae*. Together with previous reports demonstrating that cytoplasmic effector accumulation in BICs of *M. oryzae* also required the exocyst components Exo70 and Sec5 [33], we assume that the effective delivery of secretory proteins during infection of plant hosts requires the exocyst coupled with SNARE proteins, such as Sec22 and Syn8, to tether vesicles loaded with secretory proteins to the plasma membrane.

In summary, we provide evidence that hyphopodium-specific VdNoxB-regulated penetration peg formation accompanied by cytoskeletal organization of the septin-ring, form a fungus-host interface that functions as a site for the dynamic delivery of secretory proteins. The
exocyst, VdSec22-mediated transport between the ER and Golgi apparatus and VdSyn8-mediated transport/cue-sensing via endosomes are involved in the secretion of secretory proteins, possibly including effectors, towards the interfaces (Fig 8). We assume that the fungal infectious structures function as key signaling hubs during plant infection and are the apparatus that not only breaches host cells but also generates unique interfaces for the secretion of fungal secretory proteins and associated regulatory components.

**Materials and methods**

**Fungal isolates, culture conditions, infection assays and DNA analysis**

The virulent defoliating *V. dahliae* isolate V592 from cotton that originated in Xinjiang, China, was used in this study. This isolate and its transformants were stored at –80˚C and cultures were reactivated on potato dextrose agar (PDA) medium at 25˚C in the dark. The conidia for the infection assays were cultured in liquid Czapek-Dox medium. Hyphae for microscopic observation were incubated on M0 medium with urea modified as NaNO$_3$ [45].

For plant infection, cotton plants ('Xinluzao No. 16') were used in infection assays to evaluate the effect of *V. dahliae* isolate V592 and transformants on virulence using our laboratory’s unimpaired root-dip inoculation method, as described in our previous research [25]. Disease progression was recorded after 3 weeks of incubation. The infection assay for transformants was repeated three times. The symptoms were evaluated, and the disease grade was classified as follows: 0 (no symptoms), 1 (0–25% wilted leaves), 2 (25–50%), 3 (50–75%) and 4 (75–100%) [10]. The data were analyzed using the Student’s t-test.

Nucleic acid extraction and fungal transformation have all been previously described [25]. Single copy insertion was confirmed in transformants which were used to analyze the fluorescence intensity.

**Preparation of deletion constructs**

To generate the knockout plasmids pKOVdSCP10, pKOVdSep3, pKOVdSep5, pKOVdSec22, pKOVdSyn8 and pKOVdExo70, upstream and downstream genomic sequences were amplified with the primers shown in S1 Table. The upstream and downstream genomic sequence pairs were inserted into a position flanking the hygromycin resistance cassette of vector.
pGKO-HPT with the USER enzyme to generate knock-out plasmids, and transformation was performed as previously described [46].

**Preparation of GFP fusion constructs**

All of the GFP fusion constructs or RFP fusion constructs (next part) were generated by the infusion cloning method based on homologous recombination using the ClonExpress II or ClonExpress MultiS kit (Vazyme, China). The primers are shown in S1 Table. In each case, the primers contain a 15-20-bp overlap with adjoining fragments to allow the assembly of fragments by homologous recombination.

To select transformants with G418, the pSUL-NEO binary vector was created by insertion of the G418 resistance cassette amplified with Neo-F/R primers from pKOV21 into XbaI-digested pSULPH-GFP [25].

For convenient expression of the GFP fusion protein under the constitutive Tef promoter and TrpC terminator, we generated a binary vector pSUL-NEO-Tef-EKGFPTrpC using the following steps: (1) a 0.7-kb 3GA-EGFP fragment, amplified with primers 3GAGFP-F (bearing three repeats of nucleotides encoding ‘GA’) and GFP-Nt-R from pNPP9 [47], was cloned into the EcoRI/NotI sites of pNPP94 [48], resulting in pNPP94-3GAGFP; (2) we amplified the Ptef:3GAGFP:Ttrpc fusion from pNPP94-3GAGFP using primers psul-ppn-HindIII-F and psul-ppn-EcoRI-R and recombined with the product HindIII/EcoRI-linearized pSUL-NEO to generate plasmid pSUL-NEO-Tef-3GAGFP-TrpC; (3) to separate GFP and the protein of interest, we introduced a linker with five repeats of nucleotides encoding the ‘EAAK’ motif [49] into EcoRI-digested pSUL-NEO-Tef-3GAGFP-TrpC, generating plasmid pSUL-NEO-Tef-EKGFP-TrpC.

To generate the Ptef:VdSep5-GFP:Ttrpc construct, VdSep5 was amplified from the cDNA of V592 and fused into pSUL-NEO-Tef-EKGFP-TrpC.

To generate a binary vector that included the Tef promoter and TrpC terminator but lacked GFP, we amplified the Tef promoter and TrpC terminator from pNPP94 using primer psul-ppn-HindIII-F and psul-ppn-EcoRI-R and recombined with the product HindIII/EcoRI-linearized pSUL-NEO to generate pSUL-NEO-Tef-TrpC.

To generate the Ptef:LifeAct-GFP:Ttrpc fusion construct, we amplified Ptef:LifeAct-GFP from pAB261 [50] and integrated into HindIII/EcoRI-linearized pSUL-NEO-Tef-EKGFP-TrpC.

To generate the C-terminal GFP fusion construct under the oliC promoter, the primer pair oliC-HindIII-F and oliC-BamHI-R were used to amplify the template pNAH-Grx1-roGFP2 plasmid [51], and the resulting PCR products were fused into HindIII/BamHI-digested pSUL-NEO-Tef-EKGFP-TrpC to generate plasmid pSUL-NEO-oliC-EKGFP-TrpC. To generate VdSCP8-GFP, VdSCP9-GFP, VdSCP10-GFP, VdSec8-GFP and VdExo70-GFP constructs with the oliC promoter, genomic sequences were amplified and fused into pSUL-NEO-oliC-EKGFP-TrpC.

To generate VdSCP8-GFP, VdSCP9-GFP, VdSCP10-GFP, VdSec8-GFP and VdExo70-GFP constructs with the native promoter, genomic sequences spanning 1.5–2 kb upstream of the start codon were amplified and fused into pSUL-NEO-Tef-3GAGFP-TrpC using homologous recombination.

**Preparation of the RFP fusion construct**

To co-express the RFP fusion construct with the GFP fusion construct, a nourseothricin resistance cassette was amplified using the primer pair nat-F/R with the pAL6-LifeAct plasmid as
template [52] and fused into XbaI/Xhol-digested pSULPH-GFP to generate pNat-GFP. We amplified the Tef promoter and TrpC terminator from pNPP94 using the primers psul-ppn-HindIII-F and trpC-xbaI-R, and we recombined them with HindIII/EcoRI-linearized pNat-GFP to generate pNat-Tef-TrpC.

To study the localization relationship between VdSep5 and VdSCP10-GFP, we constructed Ptef:VdSep5-RFP:TrpC which was selected by nourseothricin. First, we cloned the RFP sequence from plasmid pAL6-LifeAct [52] by EK-RFP-F and RFP-R into EcoRI-digested pNat-Tef-TrpC, resulting in pNat-Tef-EKRFP-TrpC. A VdSep5 cDNA fragment was fused into BamHI/EcoRI-digested pNat-Tef-EKRFP-TrpC, resulting in Ptef:VdSep5-RFP:TrpC fusion. Transformants were selected in the presence of nourseothricin (50 μg/mL).

Preparation of complementary constructs

For complementary VdΔsep3, VdΔsec22 and VdΔsyn8 mutants, the corresponding genomic sequences, including 1.5–2 kb upstream of the start codon, were amplified with the primers listed in S1 Table and fused into HindIII/EcoRI-linearized pNat-Tef-TrpC by homologous recombination. Transformants were selected in the presence of nourseothricin (50 μg/mL). The Ptef:VdSep5-GFP:TrpC construct and Polie:VdExo70-GFP:TrpC were used to complement VdΔsep5 and VdΔexo70 mutants, respectively.

RNA extraction and cDNA

Total RNA was isolated from frozen mycelium collected from M0 medium cultured for 3d. V. dahliae cDNA was reverse-transcribed using SuperScript® III (Invitrogen).

Quantitative real-time PCR

Before reverse transcription, residual DNA was removed from the total RNA using gDNA wiper (Vazyme). cDNA was reverse transcribed using HiScript II Q RT Supermix (Vazyme), and qRT-PCR was performed using ChamQ SYBR qPCR MasterMix (Vazyme) with the Bio-Rad CFX96 Real-Time system. The transcription levels of the target genes were quantified relative to the constitutively expressed elongation factor 1-α of Verticillium dahliae (VdElf). The gene-specific primers are listed in S1 Table. Biological replicates were performed three times.

Light microscopy

To observe the infection of V. dahliae, A. thaliana roots were immersed in a conidial suspension (~10^5 conidia/mL in water solution) for 10 min and then transferred onto a 0.75% agar plate at 25˚C in the dark. To observe the protein localization of V. dahliae on cellophane, conidia were placed on cellophane and incubated at 25˚C. The mycelium grown on cellophane for 3–9 days was used for protein localization assays. To compare the secretory difference between V592 and VdΔsec22/VdΔsyn8 mutants, the fungi were collected from the outer zone of the colony at the earliest time point for most V592 hyphal necks with ring signals. The fluorescence intensity data were analyzed using the Student’s t-test. Small pieces (~0.5 cm^2) of cellophane with mycelium at the margin of the fungal colonies were cut with a scalpel and mounted in water. Images were obtained under a confocal laser microscope (Leica TCS SP8; Leica Microsystems) with a 100x oil immersion objective lens. The excitation wavelengths and emission filters were as follows: 488 nm/band-pass 500 to 550 nm for GFP, 561 nm/ band-pass 570 to 670 nm for RFP and FM4-64, and 405 nm/band-pass 400 to 600 nm for ER-Tracker. Confocal images were captured with a Leica hybrid detector and analyzed with Leica LAS AF software.
For each microscopy-based experiment, at least 20 images with three biological independent samples were observed for each micrograph to make conclusions. Each experiment was repeated at least twice.

Transmission electron microscopy

For TEM observation, *V. dahliae*-infected cotton root and *V. dahliae* on cellophane were fixed immediately in 2.5% glutaraldehyde, buffered with PBS (pH 7.4) at 4˚C overnight, washed with the same buffer four times and post-fixed with 1% osmium tetroxide for 1 h. Dehydration was then performed in an acetone series (50%, 75%, 85%, 95%, 100%), and the slices were embedded in Spurr’s resin mixture. Ultrathin serial sections (70 nm thickness) were cut from resin blocks, followed by uranyl acetate staining, and observed with a JEM-1400 electron microscope.

Staining of fungi

For plasma membrane staining, FM4-64 (ThermoFisher) was used according to the manufacturer’s protocol. For ER-Tracker staining, cultures were incubated at 30˚C for 30 min with PBS containing 1 μM ER-Tracker™ Blue–White DPX (Molecular Probes) that had been prewarmed at 30˚C for 30 min, washed once with fresh PBS without the dye, and subjected to microscopic observation [53]. Next, 100 μg/mL FITC-conjugated wheat germ agglutinin (FITC-WGA, Sigma) was used to stain the fungal cell wall.

Fluorescence recovery after photobleaching (FRAP) analysis

FRAP analyses was carried out with fungi on cellophane under a spinning disk confocal microscope (UltraVIEW VoX, Perkin Elmer, Beaconsfield, Buckinghamshire, UK) equipped with a Yokogawa Nipkow CSU-X1 spinning disk scanner, Hamamatsu EMCCD 9100–13, and Nikon TiE inverted microscope with the Perfect Focus System. We used the UltraVIEW PK Device to photobleach GFP. For the FRAP analyses, the specific region of interest (ROI) covering the entire fluorescence in the ring was selected for bleaching. Twenty bleaching iterations were performed using a 488 laser power of 60%. Image scans were obtained with 15% 488 laser power before and after bleaching. For quantitative analyses, the GFP fluorescence recovery curves were measured as the mean intensity of the ROI pixels, normalized using the using Volocity software (Perkin Elmer), and graphed using Microsoft Excel.

Supporting information

S1 Fig. Alignment of the predicted fungal amino acid sequences used in this study. (A-F) The *V. dahliae* amino acid sequences were aligned with each sequence of the putative homologs in *M. oryzae* (Mo) and *E. festucae* (Ef)/*S. cerevisiae* (Sc). Sequences were aligned using ClustalX2 and shaded using GeneDoc. Amino acid residues within a black background were identical among all of the listed proteins, gray residues were identical in two out of three of the listed proteins, and those shown on a white background did not show any similarity. VdSep3 (VDAG_00736) is aligned with *M. oryzae* (EHA54688.1) and *S. cerevisiae* (DA09624.2) (A). VdSep5 (VDAG_04382) is aligned with *M. oryzae* (EHA45843.1) and *S. cerevisiae* (DA08862.1) (B). VdSec22 (VDAG_08386) is aligned with *M. oryzae* (EHA47424.1) and *S. cerevisiae* (DA09582.1) (C). VdSyn8 (VDAG_01236) is aligned with *M. oryzae* (EHA50711.1) and *S. cerevisiae* (DA06974.1) (D). VdExo70 (VDAG_09051) is aligned with *M. oryzae* (EHA54952.1) and *S. cerevisiae* (DA08714.1) (E). VdSec8 (VDAG_08435) is aligned with *M.
oryzae (EHA47501.1) and S. cerevisiae (DAA11477.1). (F).

(S2) Fig. Hyphopodium-specific VdNoxB-dependent development of the penetration peg in V. dahliae. (A) Hyphopodium-specific expression of GFP-tagged V. dahliae NADPH oxidase B (VdNoxB) under the native promoter (left) and localization of GFP-VdNoxB at the penetration peg (right). (B) Observation of the development of the VdNoxB-dependent hyphopodium and penetration peg in cellophane. Bar = 2.5 µm. (C) Observation of the penetration peg on cellophane at 3 dpi. The micrographs show two scanning layers of the upper side and base (below) of the hyphopodium. A thin penetration peg of wild-type V592 differentiated from the base of the hyphopodium and pierced the cellophane (below); VdΔnoxB developed a hyphopodium without the formation of a penetration peg.

(S3) Fig. Localization of the F-actin ring at the hyphal neck. (A–B) Micrographs of F-actin organization in the hyphal neck visualized by expression of LifeAct-GFP in V592. F-actin organized in the hyphal neck on cellophane (A) and on Arabidopsis thaliana root (B). Bar = 2.5 µm.

(S4) Fig. Secretion of signal peptide containing proteins on the penetration interface. (A) Signal peptide analysis of VdSCP8, VdSCP9, VdSCP10 and VdIscI in V. dahliae. The signal peptide of each protein was predicted using the SignalP 4.1 server. The predicted signal peptides are marked in red color, and the 30 amino acids from the initiation codon are displayed. (B) Detection of the expression levels of selected SCPs by qRT-PCR. RNA samples isolated from 2-day-old fungal culture harvested in liquid Czapek-Dox (CD) medium, 4-day-old fungi cultured on cellophane and 2-day-old fungi on cotton roots. The relative expression levels were estimated using the 2^ΔΔCt method. The expression level of each SCP gene in liquid culture was arbitrarily set to 1. The mean and standard errors were calculated from three independent replicates. The asterisks indicate significant differences (*P<0.05; Dunnett’s test). (C) VdSCP8-GFP expressed under the native promoter was detected at the penetration zone. The V592 transformant expressing VdSCP8-GFP under the native promoter was observed after growth on cellophane for 8 d. Bar = 2.5 µm. (D) The ring signals of SP_VdSCP8-GFP, SP_VdSCP9-GFP and SP_VdSCP10-GFP at the penetration zone. V. dahliae transformants expressing SP_VdSCP8-GFP, SP_VdSCP9-GFP and SP_VdSCP10-GFP under the control of the oliC promoter were used for the assay. The plasma membrane was stained with FM4-64 (red). Bar = 2.5 µm.

(S5) Fig. Targeted deletion of VdSCP10 decreased virulence of V. dahliae in cotton plants. (A) Physical maps of the VdSCP10 locus and the homologous recombination construct obtained by fusion of the VdSCP10 5’flank, hygromycin B resistance gene cassette and VdSCP10 3’flank. The probe and relative positions of primers used for PCR are indicated. (B) Southern blot analysis of targeted gene deletion mutants. EcoRI digested genomic DNA from the V592 wild type strain and two putative Vdscp10 transformants were blotted with the probe indicated in the schematic diagram. (C) PCR amplification of genomic DNA from the complemented transformants using the primer pair in-F and in-R produced a banding pattern consistent with the integration of an intact gene in V592. (D) The colony morphology of the wild-type V592 and Vdscp10 mutant strains and the corresponding complemented strains on PDA plates after a 2-week incubation. (E, F) Disease symptoms (E) and disease grades (F) of cotton plants infected with wild-type V592, VdΔscp10 mutant and the complementary strains at 21 dpi. The disease grade (DG) was calculated as previously described. Four is the highest DG, meaning that the entire plant died, while 0 is the lowest DG with no visible wilting. Three
replicates of 36 plants were used for each inoculum (*P<0.05; t-test).

(PDF)

S6 Fig. Targeted deletions of VdSep3 and VdSep5 genes in V. dahliae strain V592. (A) Physical maps of the VdSep3 locus and of the homologous recombination constructs obtained by fusion of the VdSep3 5’flack, hygromycin B resistance gene cassette and VdSep3 3’flack. Probes and relative positions of the primers used for PCR are indicated. hph, hygromycin resistance gene. (B) Southern blot analysis of targeted gene deletion mutants. BamHI digested genomic DNA from the V592 strain and two putative VdΔsep3 transformants were blotted with the probe indicated in the schematic diagram. KpnI and HindIII digested genomic DNA from V592 wild type strain and two putative VdΔsep5 transformants were analyzed as described above. (C) PCR amplification of genomic DNA from the complemented transformants using the primer pair in-F and in-R produced a banding pattern consistent with the integration of an intact VdSep3 and VdSep5. Lanes 1–4 were using for the verification of VdΔsep3 complementation and lanes 5–8 were for the verification of VdΔsep5 complementation. (D) Colony morphology of V592, VdΔsep3 and VdΔsep5 mutants and the complementary strains on PDA plates incubated for 2 weeks. (E) Penetration ability analysis of VdΔsep3 and VdΔsep5. Colonies of V592, VdΔsep3 and VdΔsep5 on M0 medium overlaid with cellophane (Before) and removal of the cellophane membrane (After). Images in the first row were obtained at 6 dpi, and the colonies below the cellophane were obtained at 9 dpi. (F) Hyphopodium morphology analysis of V592, VdΔsep3 and VdΔsep5. images were obtained at 6 dpi. (G) Deficient development of the hyphopodium in VdΔsep3 and VdΔsep5. Fungi incubated on cellophane at 5 dpi were used for the observation. The numbers of hyphopodia were counted in three fields of the culture under a light microscope at x1000 magnification with three replicates. The mean and SD for (G) were calculated from three clones for each mutant (*P<0.05; t-test). (H, I) Disease symptoms (H) and disease grades (I) of cotton plants infected with wild-type V592, VdΔsep3 and VdΔsep5 mutants and the complementary strains at 21 dpi. Three replicates of 36 plants were used for each inoculum. The asterisks indicate significant differences compared with V592 infection (*P<0.05; t-test).

(PDF)

S7 Fig. Gene disruptions of VdSec22, VdSyn8 and VdExo70 in V. dahliae. (A) Physical maps of the VdSec22 locus and the homologous recombination construct obtained by fusion of the VdSec22 5’flack, hygromycin B resistance gene cassette and VdSec22 3’flack. The probe and relative positions of primers used for PCR are indicated. The same approach was used for disruption of VdSyn8 and VdExo70. (B) Southern blot analysis of targeted gene deletion mutants. NcoI-digested genomic DNA from V592 and two putative VdΔsec22 transformants were blotted with the probe indicated in the schematic diagram. NcoI-digested genomic DNA from the V592 and two putative VdΔsyn8 transformants were analyzed as described above. SmaI and BamHI-digested genomic DNA from the V592 wild type strain and two putative VdΔexo70 transformants were analyzed as described above. (C) PCR amplification of genomic DNA from the complemented transformants using the primer pair in-F and in-R produced a banding pattern consistent with the integration of an intact gene in V592. Lanes 1–4, 5–8 and 9–12 were for the verification of VdΔsyn8, VdΔsec22 and VdΔexo70 complementation, respectively. (D) Colony morphology of wild-type V592 and VdΔexo70, VdΔsec22 and VdΔsyn8 mutant strains and the corresponding complemented strains on PDA plates 2 weeks post-incubation.

(PDF)

S8 Fig. Determination of the T-DNA insertional copy number of VdSCP10-GFP in V592, VdΔsyn8, VdΔsec22 and VdΔexo70. Genomic DNA isolated from V592, VdΔsyn8, VdΔsec22
and VdΔexo70 expressing VdSCP10-GFP were digested with EcoRI for Southern blot analysis. Red arrowheads indicate selected colonies with single copy insertions for further study. Hybridization was performed with the 32P-labeled oliC promoter-specific DNA probe as shown below.

(PDF)

S9 Fig. Deletion of VdSec22 caused VdSCP10-GFP retention in the ER. VdSCP10-GFP expressed under the oliC promoter was transformed into V592 and VdΔsec22. Staining the hyphal ER with ER-Tracker Blue-White DPX. Bar = 2.5μm.

(PDF)

S10 Fig. Localization of V. dahliae exocyst subunits at the hyphal tips, hyphopodium base and hyphal neck. (A) Localization of VdSec8-GFP and VdExo70-GFP in the hyphal tips of V. dahliae on cellophane. (B) Localization of VdSec8-GFP and VdExo70-GFP at the base of the hyphopodium on cellophane. (C) VdSec8-GFP organized at the base of the hyphopodium on the surface of the root. (D) After invasive hyphae developed, VdSec8-GFP was organized at the hyphal neck on cellophane and plant roots. Bar = 2.5μm.

(PDF)

S11 Fig. Colocalization of V. dahliae exocyst subunit VdSec8 and VdSep5 at the hyphal neck. Confocal laser scanning microscopy (CLSM) images and linescan graph showing co-localization of VdSec8-GFP and VdSep5-RFP at the hyphal neck. Bar = 2.5μm.

(PDF)

S1 Table. Primers used in this study.

(XLSX)

S1 Movie. Verticillium dahliae colonization in cellophane. Confocal laser scanning microscopy (CLSM) images of hyphopodium development and penetration of V. dahliae V592 in cellophane. Fungal cell wall was stained with FITC-WGA. Movie was taken at 7dpi.

(AVI)

S2 Movie. Verticillium dahliae colonization in Arabidopsis thaliana root. Confocal laser scanning microscopy (CLSM) images of hyphopodium development and penetration of V. dahliae V592 on Arabidopsis thaliana root. Fungal cell wall was stained with FITC-WGA. Movie was taken at 1 dpi.

(AVI)

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References


