The NADPH Oxidase Complexes in *Botrytis cinerea*: Evidence for a Close Association with the ER and the Tetraspanin Pls1

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Abstract

NADPH oxidases (Nox) are major enzymatic systems that generate reactive-oxygen species (ROS) in multicellular eukaryotes. In several fungi they have been shown to be involved in sexual differentiation and pathogenicity. However, in contrast to the well characterized mammalian systems, basic information on the composition, recruitment, and localization of fungal Nox complexes and on the molecular mechanisms of their cellular effects are still lacking. Here we give a detailed analysis of components of the Nox complexes in the gray mold fungus *Botrytis cinerea*. It had previously been shown that the two catalytic transmembrane subunits BcNoxA and B are important for development of sclerotia and for full virulence, with BcNoxA being involved in spreading of lesions and BcNoxB in penetration; BcNoxR functions as a regulator of both subunits. Here we present evidence (using for the first time a functional GFP fusion able to complement the ΔbcnoxA mutant) that BcNoxA localizes mainly to the ER and at the plasma membrane; BcNoxB shows a similar localization pattern, while the regulator BcNoxR is found in vesicles throughout the hyphae and at the hyphal tip. To identify possible interaction partners, which could be involved in the localization or recruitment of the Nox complexes, we functionally characterized the tetraspanin Pls1, a transmembrane protein, which had been suggested to be a NoxB-interacting partner in the saprophyte *Podospora anserina*. Knock-out experiments and GFP fusions substantiate a link between BcNoxB and BcPls1 because both deletion mutants have overlapping phenotypes (especially a defect in penetration), and the proteins show a similar localization pattern (ER). However, in contrast to the corresponding protein in *P. anserina* BcPls1 is important for female fertility, but not for ascospore germination.

Introduction

Reactive oxygen species (ROS) play a dual role in eukaryotic cells: as unwanted by-products of many metabolic reactions they have to be scavenged to avoid deleterious effects on macromolecules, but their diffusible nature enables them to play a role as messenger molecules for signaling within the cell and in cell-cell communication [1]. Multiple roles of ROS are also evident in host-pathogen interactions. Plants react to pathogen attack with so-called oxidative burst, which generates ROS serving primarily as signaling molecules for induction of effective plant defense reactions like the hypersensitive response (HR). In addition ROS can have direct toxic effects on the invading pathogen. On the other hand, ROS are involved in differentiation processes of fungal pathogens, which are important for virulence [2]. The necrotrophic gray mold fungus *Botrytis cinerea* initiates a massive oxidative burst upon infection of its host plants and even requires the host plant’s response to achieve full pathogenicity: in HR-deficient *Arabidopsis thaliana* strains the virulence of *B. cinerea* is strongly reduced [3]. *B. cinerea* - like many other fungi - was shown to produce ROS within and around hyphal tips also in planta [4]. Therefore, it was postulated that the fungus uses ROS to actively trigger the host response [5,6]. Searching for the source of this extracellular ROS we initiated a functional characterization of the major enzymatic ROS-generating system in eukaryotic cells, the NADPH oxidase complex, in *B. cinerea*.

NADPH oxidases are transmembrane proteins which reduce oxygen to superoxide using NADPH as an electron donor. The best studied member of this group is the mammalian phagocytic gp91phox (named also Nox2), which is responsible for the defense response of neutrophils [7]. To activate its function a multi-subunit complex has to be formed: the integral membrane protein flavocytochrome b558, consisting of the catalytic subunit gp91phox (named also Nox2), and the adaptor protein p22phox [8], is assembled with the cytosolic components p40phox, p47phox, p67phox and the small GTPase Rac upon phosphorylation of p47phox [9]. NADPH oxidases have been found in a wide range of eukaryotic organisms, and have originally been linked with multicellularity [10]. However, recently a genuine NADPH oxidase was identified in *Saccharomyces cerevisiae* [11] showing that this correlation is no longer valid. In filamentous fungi three distinct subfamilies of Nox enzymes have been discovered: NoxA, NoxB and NoxC [12,13]. NoxA and NoxB are homologs of the mammalian catalytic transmembrane subunits gp91phox (named also Nox2) and p22phox, respectively.
subunit Nox2 and are widely distributed within filamentous fungi. The third isoform NoxA, present in some species, carries a calcium binding EF-hand, like the human Nox5 and the plant Nox homologs (Rbohs) [14]. Furthermore, in filamentous fungi homologs of the mammalian regulatory subunit p67phox (NoxR) and the small GTPase Rac are present [15,16]. Other components of the fungal Nox complexes are yet to be discovered, as neither homologs of the mammalian cytosolic subunits p47phox and p40phox nor the flavocytochrome component p22phox have been identified in fungi [16]. Possible components might be the scaffold protein Bem1 and the guanine exchange factor (GEF) Cdc24, as a direct interaction of these proteins with the putative regulator NoxR via their PB1 domains was shown in Epichloë festucae [17]. Functional analyses of Nox genes in filamentous fungi have revealed that Nox are involved in various differentiation processes. This was first demonstrated in Aspergillus nidulans, where NoxA is required for fruiting body formation [12]. Subsequently, it was shown that also in the ascomycetes Podospora anserina and Neurospora crassa Nox1 is necessary for development of fruiting bodies. The second Nox enzyme present in P. anserina and N. crassa (Nox2) is involved in germination of ascospores [18,19]. In B. cinerea both BcNoxA and BcNoxB are involved in sclerotial formation [20]. Additionally, it was shown recently that in P. anserina both Nox complexes are required for cellulose degradation via specialized needle-like hyphal structures [21]. Besides their involvement in sexual differentiation processes, Nox are also needed during pathogenicity. In B. cinerea deletion mutants of bcnoxA penetrate the host tissue, but colonization of the host is slower compared to the wild-type (WT), and bcnoxB mutants are slower in forming primary lesions. The deletion of the regulator bcnoXR shows an additive effect [20]. In Claviceps purpurea the deletion mutant of cpnox1 is impaired in colonization of host tissue: it produces conidia-containing honeydew and immature sclerotia only in rare cases [22]. In contrast cpnox2 deletion results in significantly increased production of honeydew. However, the mutants are unable to form mature sclerotia and are therefore restricted to the sporulating stage (D. Buttermann, pers. communication). In the endophyte Epichloë festucae deletion of ephox1 leads to a switch from a mutualistic to an antagonistic interaction, while deletion of both ephox1 and ephox2 resulted in non-sporulating stage (D. Buttermann, pers. communication). In the production of honeydew. However, the mutants are unable to form mature sclerotia: it produces conidia-associated honeydew and immature sclerotia only in rare cases [22]. In contrast ephox2 deletion results in significantly increased production of honeydew. However, the mutants are unable to form mature sclerotia and are therefore restricted to the sporulating stage (D. Buttermann, pers. communication). In the endophyte Epichloë festucae deletion of ephox1 leads to a switch from a mutualistic to an antagonistic interaction, while deletion of both ephox1 and ephox2 caused apathogenicity of the fungus [24]. Interestingly, the effect of nos deletion on ROS production varies between the different species. Some show unchanged cellular ROS levels (B. cinerea, M. oryzae single mutants), in some cases they are enhanced (M. oryzae vegetative hyphae of ANox1Δnox2, P. anserina Δnox1, Δnox1/2 and ΔnoxR) [20,21,24], and in some cases they are reduced (A. nidulans ΔnoxΔox during sexual development; E. festucae ΔnoxA; M. oryzae Δnox1/2 during appressorium formation) [12,24,25].

Given that the catalytic subunits NoxA and NoxB are transmembrane proteins, it is still not known to which cellular membranes they localize. So far the only localization study was accomplished by Egan et al., who showed Nox1 localization in M. oryzae at the appressorium periphery starting from 4 h and in the central appressorium vacuole after 24 h [24]. This gfp-construction was under control of the native promoter, but was not shown by complementation studies to be functional.

As the constitution of the fungal Nox complex is still incompletely elucidated, possible partners for direct or indirect interaction are of high interest. In P. anserina the Nox complex was recently linked to another transmembrane protein, the tetrasmus Pls1, because the deletion of pls1 and nos2 led to a similar germination defect in ascospores [26]. Tetrasmus are small eukaryotic integral membrane proteins [27] known to have varying functions. In the animal kingdom they function as organizers of multimolecular membrane complexes and regulate, amongst others, cell migration, fusion and signaling events [28]. In filamentous fungi three families of tetrasmus have been identified (Pis1, Tsp2, Tsp3) with different distribution amongst phyla [29]. All these tetrasmus proteins display low sequence similarities, but they share highly conserved secondary structures.

These include four transmembrane domains and a cytosine-based pattern in the large extracellular loop EC2 [30]. The most important tetrasmus of ascomycetes seems to be Pis1, which was first identified as a virulence factor in M. oryzae. Expression of pis1 was shown in appressoria, where the protein localizes to the plasma membrane and in vacuoles [31]. As shown for M. oryzae, Pis1 is also necessary for appressoria-mediated penetration in Colletotrichum lindemuthianum and the B. cinerea strain T4 [32,33]. Interestingly, pis1 and nos2 seem to have a similar distribution within fungal genomes, as they are either both present or both absent, suggesting a connection between these genes or their products, respectively [26]. Here we present evidence that the two NADPH oxidase catalytic subunits BcNoxA and BcNoxB localize in the ER and at the plasma membrane, while their regulator NoxR localizes in vesicles throughout the hyphae and at the hyphal tip. Furthermore, the tetrasmus Pis1 is functionally characterized in the more aggressive B. cinerea strain B05.10 to substantiate a possible connection between NoxB and Pis1. Indeed, both deletion mutants have overlapping phenotypes and the proteins show a similar localization pattern.

**Results**

In a previous study NADPH oxidase catalytic subunits bcnxA (BC1G_10823.1) and bcnxB (BC1G_14597.1), encoding the catalytic subunits, and bcnXR (BC1G_06200.1), encoding the regulatory subunit, were identified in B. cinerea B05.10. Functional characterization of the respective deletion mutants revealed an involvement of all components in pathogenicity and sclerotia production [20]. Even though the composition and possible up- and downstream partners of the Nox complex are intensely investigated in several fungi, so far many components remain to be elucidated. As Lambou et al. [26] recently proposed a connection between the tetrasmus Pis1 and the catalytic NADPH oxidase subunit NoxB, we were interested in the characterization of the tetrasmus Pis1 in B. cinerea B05.10 to investigate a possible connection to the BcNox complex.

**Identification of the Gene bcpls1 in B. cinerea B05.10**

A blastx search in the B. cinerea B05.10 database (Broad Institute, http://www.broadinstitute.org/annotation/genome/botrytis_cinerea/Home.html) using the Magnaporthe oryzae Pis1 sequence as a query identified an MgPls1 homolog protein annotated as hypothetical protein similar to tetrasmus (BC1G_09439.1). According to the annotation, the open reading frame (ORF) BC1G_09439.1 consists of 612 bp and encodes a protein of 176 aa. However, 88 bp upstream of the annotated start codon, there was no sequence information in the B05.10 sequence. Comparison of the genomic B05.10 DNA sequence with expressed sequence tags (ESTs) of the strain B05.10 and the genomic sequence of B. cinerea T4 (http://urgi.versailles.inra.fr/Species/Botrytis) indicated that the actual start codon of BC1G_09439.1 is 147 bp upstream of the annotated start codon (data not shown).

Therefore the ORF BC1G_09439.1, named bcpls1, consists of 759 bp. Comparison of ESTs with bcpls1 confirmed an intron of 81 bp. Therefore, bcpls1 encodes a protein of 226 aa.
predicted protein contains no conserved domains. Still, a prediction of transmembrane helices in proteins using the TMHMM server version 2.0 (http://www.cbs.dtu.dk/services/TMHMM) indicated the structural hallmarks of tetraspanins with four transmembrane domains and a large extracellular loop between TM3 and TM4 (data not shown).

**Generation of Δbcpls1 Mutants**

The gene *pls1* had already been characterized in the wild-type strain T4 of *B. cinerea* [33]. However, this strain shows some phenotypical differences to the strain B05.10 (which is widely used as reference strain in the *Botrytis* community): it sporulates in darkness, does not produce any sclerotia or oxalic acid and is less aggressive during infection. In order to analyze the function of BcPls1 in *B. cinerea* B05.10 and compare it to the function of other proteins in this strain, a *bcpls1* deletion strain was created using a replacement approach. The ORF of *bcpls1* in the wild-type strain B05.10 was replaced by a hygromycin resistance cassette by homologous recombination (Fig. S1A). Replacement of *bcpls1* was detected in several transformants by diagnostic PCR. Single-spore isolations resulted in homokaryotic deletion mutants, which were confirmed by diagnostic PCR (Fig. S1B). Southern blot analysis confirmed homologous integration into the *bcpls1* locus without further ectopic integration events of the replacement cassette in three independent transformants, termed Δbcpls1 (T20), Δbcpls1 (T21) and Δbcpls1 (T23) (Fig. S1C). The mutants Δbcpls1 (T20) and Δbcpls1 (T21) showed the same phenotype in all experiments, while the mutant Δbcpls1 (T23) showed a stronger phenotype when grown on minimal medium. Therefore Δbcpls1 (T20) was taken for all further analyses.

**BcPls1 has no Influence on Light-dependent Differentiation but Affects Sexual Development**

Differentiation and growth rates of the Δbcpls1 deletion mutant were compared to the wild type (WT) in plate assays. These analyses revealed that light-dependent differentiation of Δbcpls1 was not disturbed: the mutant produced conidia when grown in light and sclerotia of normal appearance when grown in darkness (Fig. S2A). Furthermore, Δbcpls1 was neither affected in growth on complete and minimal medium nor in growth on media containing different oxidative or osmotic stressors (Fig. S2B). In addition, the Δbcpls1 mutant did not show significantly altered extracellular H$_2$O$_2$ production, as shown by DAB staining and an Amplex Red peroxide assay (although the latter showed a slight induction of H$_2$O$_2$ production by the mutant (P<0.05)) or acidification of the surrounding medium as compared to the WT (Fig. S3B). The same analyses were also done for ΔbcnoxA, ΔbcnoxB, ΔbcnoxR and ΔbcnoxAB. All of these *bcnox* mutants showed WT-like production of oxalic acid and H$_2$O$_2$ production was not reduced in any of them. Indeed the ΔbcnoxA mutant even showed an increased production of H$_2$O$_2$ (P<0.01). In contrast, treatment of the WT with the flavoenzyme inhibitor diphenyl-
neiodonium chloride (DPI) in the Amplex Red assay showed a reduction of $H_2O_2$ production of about 50% compared to the untreated WT ($P<0.01$), indicating that besides the Nox enzymes other flavoenzymes are involved in extracellular $H_2O_2$ production (Fig. S3).

However, results of sexual crosses with $B$. cinerea SAS405 (the MAT1-2 reference strain commonly used as opposing mating partner of B05.10) showed an unusual behavior of the $\Delta$bcpls1 mutant. Although the $\Delta$bcpls1 sclerotia exhibited no morphological differences during their development, they failed to develop apothecia after fertilization with SAS405 microconidia. In contrast, the sclerotia of SAS405 when fertilized with microconidia of the $\Delta$bcpls1 mutant produced apothecia of normal appearance (Fig. 1). In conclusion, sclerotia of the $\Delta$bcpls1 mutant are non-functional in sexual development, i.e. the mutant is female sterile, while the microconidia of the mutant are functional. Ascospores were collected from apothecia obtained from the cross between SAS405 sclerotia and $\Delta$bcpls1 microconidia and germinated at low density (~50–100 spores/plate) on non-selective medium. The ascospores showed normal germination rates. Upon transfer to plates containing hygromycin, at least half of the germlings were hygromycin-resistant, indicating that these germlings carried the $\Delta$bcpls1 mutation. Therefore, in contrast to $P$. anserina [26], $\text{Pls}1$ is not required for ascospore germination in $B$. cinerea. We previously showed that $\text{BcNoxA, BcNoxB}$ and $\text{BcNoxR}$ are involved in production of sclerotia. Deletion of $\text{bcnoxA}$ and $\text{bcnoXR}$ led to complete loss of sclerotia, whilst deletion of $\text{bcnoxB}$ resulted in the formation of fewer and morphologically aberrant sclerotia. However, ascospores derived from sexual crosses between microconidia of these deletion mutants and SAS405 sclerotia resulted in normal apothecia and the ascospores developing from these showed normal germination rates [20].

**BcPls1 is not Involved in Germination of Asexual Conidia or Vegetative Hyphal Fusion Events**

Recently it was shown that $B$. cinerea, as many other fungi, forms conidial anastomosis tubes (CATs) resulting in vegetative hyphal fusions. In this process the BcNox complex is involved: while the deletion of $\text{bcnoxA}$ leads to reduced hyphal fusion rates, deletion of $\text{bcnoxB}$ and $\text{bcnoXR}$ leads to complete loss of germling fusions [34]. To compare $\text{BcPls1}$ and the subunits of the BcNox complexes the ability of $\Delta$bcpls1 to form CATs was analyzed.

Interestingly, the fusion frequency of $\Delta$bcpls1 germlings was slightly enhanced when compared to the WT (Fig. 2A). However, 3 h after inoculation also the sugar-induced germination of asexual conidia was significantly increased in the deletion mutant. Hence, the increased germination rate might account for the more frequent fusion events observed in the $\Delta$bcpls1 mutant (Fig. 2B). Therefore, in summary, regarding the CAT formation the $\Delta$bcpls1 phenotype is clearly distinct from the $\text{ΔbcnoxA}$ phenotype, and closer to the $\text{ΔbcnoxB}$ phenotype, which is still able to form CATs [34].

**BcPls1 has an Impact on Pathogenicity**

The main defect of the $\text{ΔbcnoxB}$ mutant is its performance in the early stages of pathogenicity [20]. According to these findings $\text{BcPls1}$ might also be required for infection in the strain B05.10. Therefore, the virulence of $\Delta$bcpls1 was compared to the WT in a pathogenicity test (Fig. 3A). This test showed that indeed the virulence of the $\text{bcpls1}$ deletion mutant was reduced. However, the B05.10-derived mutant was not completely apathogenic, as reported for the T4-derived mutant [33]. In fact, the lesion diameter of the B05.10-derived mutant at 3 dpi was only moderately reduced compared to controls, when considering only the diameters of lesions in successful infections (Fig. 3C). However, by that assessment time only ca. 30% of $\text{Δbcpls1}$ inoculated bean leaves showed any symptoms of infection, while the inoculations of the WT were 100% successful, as demonstrated in a separate set of experiments in Fig. 3B: the infection efficiencies of the mutant gradually increased until at $6 \text{dpi}$ the infection was successful in 100%. Whenever there was an infection the mutant was able to complete the whole life cycle nearly as rapidly as in the WT. In conclusion, $\text{Δbcpls1}$ showed reduced penetration efficiency, while the course of infection was normal once the mutant had entered the host. In comparison the infection pattern of $\text{ΔbcnoxB}$ showed delayed formation of primary lesions in all cases (1 day slower than the WT), but once it entered the host the infection was also normal [20].

**BcPls1, BcNoxB and BcNoxR Are Required for Penetration**

As pathogenicity assays revealed that $\text{BcNoxB, BcNoxR}$ and $\text{BcPls1}$ seem to be necessary for penetration in $B$. cinerea B05.10, the ability of the $\Delta$bcpls1 deletion mutant to penetrate onion epidermal strips was investigated. The deletion mutants of the $\text{bcnoX}$ genes as well as the double mutant of the catalytic subunits $\text{BcNoxA, BcNoxB}$, $\text{BcNoxAB}$ and $\text{BcNoxR}$ were included for comparison. To exclude the possible impact of host defense reactions affecting the assay, onion epidermises were inactivated prior to inoculation with conidial suspensions and the penetration ability of the strains was studied microscopically 24 hpi (Fig. 4). These analyses showed that $\text{BcPls1}$ still produced swollen hyphal tips (appressoria-like structures) resembling those produced by the WT, but whereas the WT successfully penetrated the epidermis via these structures, the mutant was incapable of doing so. A similar phenotype has been observed for the $\text{ΔbcnoxB}$ mutant in prior studies and was confirmed again here. Comparable phenotypes were also observed for $\text{ΔbcnoxAB, ΔbcnoxR}$, while the $\text{ΔbcnoxA}$ mutant was able to penetrate the onion epidermis 24 hpi [20]. On bean leaves a similar effect was expected. To analyze penetration ability in this system leaves of bean plants were inoculated with conidial suspensions of the WT, the $\Delta$bcpls1 mutant and the $\text{ΔbcnoX}$ mutants; 18 hpi the samples were prepared for analysis via scanning electron microscopy (SEM) (Fig. 5). This analysis revealed that in most cases the WT as well as $\text{ΔbcnoxA}$ produced short germ tubes before penetration of the host tissue. In contrast, the majority of germ tubes observed for $\text{ΔbcnoxB, ΔbcnoXR}$ and $\Delta$bcpls1 elongated on the surface of the epidermis of the host leaf and instead of penetrating the host by appressoria-like structures the mutants continually produced new hyphal outgrowths and cycles of appressoria-like structures. This phenotype has previously been described for the $\text{ΔbcnoxR}$ mutant [2]. The observation that $\text{Δbcpls1, ΔbcnoxB and ΔbcnoxR}$ mutants all show a similar defect in the penetration via appressoria-like structures suggests a connection between $\text{BcPls1}$ and the Nox complex.

**BcNoxA, BcNoxB and BcPls1, but not BcNoxR, are Localized in the ER**

In mammalian cells Nox have been shown to fulfill various functions in a range of tissues, therefore localization of the Nox isoforms varies considerably in mammals. In fungi only three isoforms have been identified and in $B$. cinerea only two catalytic subunits (BcNoxA and BcNoxB) and their putative regulator (BcNoxR) were found. Differing functions have been ascribed to BcNoxA and BcNoxB. Therefore, the question arises as to whether their distinct functions are based on different localizations. In order to localize the proteins N-terminal (BcNoxA) or C-
Figure 2. Germination assay and vegetative hyphal fusions of Δbcpls1 and the WT. 

A: Hyphal germling fusions of the WT and the Δbcpls1 mutant. 3 × 10⁶ conidia were plated on Vogel’s minimal medium and incubated for 14 h in darkness. CAT formation was analyzed using light microscopy. The diagram represents the mean values of three replicates evaluating 300 spores each. The fusion frequency (CAT fusion %) was slightly enhanced in the Δbcpls1 mutant; statistical analysis (T-test) revealed no significant differences. 

B: Comparison of germination efficiencies of B. cinerea B05.10 WT and Δbcpls1 conidia after 3, 6 and 24 hours. Germination was monitored in liquid Gamborg-B5 medium amended with 10 mM glucose. The diagram represents one experiment with triplicate counting. Germination rates of Δbcpls1 conidia were increased at 3 hpi. Asterisks above the bars denote significant differences in the measurements of the indicated strains to the WT. ** = P < 0.01.

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Figure 3. Virulence of the Botrytis cinerea strain Δbcpls1 in comparison to the wild-type B05.10 (WT) and the complemented strain Δbcpls1:GFP-Pls1. 

A: Time course of infection of French bean (Phaseolus vulgaris). Primary leaves of 10 day old plants were inoculated with 7 μl droplets of conidial suspensions (2 × 10⁵ spores/ml in Gamborgs B5+2% glucose). Progression of infections was documented 3, 5 and 7 days after the infection (dpi). Results of the mutant were not consistent as in some cases infection could be detected between 3 and 5 dpi (see middle row and B). However, at 7 dpi all the leaves infected with the mutant strain showed strong infection symptoms (see right row). 

B: Statistical evaluation of infection efficiencies at different time points. At 3 dpi in only about 50% of the Δbcpls1-infected leaves symptoms could be detected while infection efficiencies of the WT and the complemented strain amounted to 100%. At 6 dpi also in the Δbcpls1 strain infection symptoms were detected in 100% of infected leaves (separate set of experiments from A: repeated 3 times with a sample size of n = 16, 16, 12 per strain; standard deviations are indicated by the error bars).

C: Statistical evaluation of infection spreading. Only the lesion diameters of successful infection events were measured at 3 dpi. Lesion diameters of Δbcpls1 infections were only slightly reduced when compared to WT and to the complemented strain at this early time point (The indicated values are means of 16 different infection events; standard deviations are indicated by the error bars; a biological replicate yielded similar results). Asterisks above the bars denote significant differences in the measurements of the indicated strains to the WT. ** = P < 0.01; *** = P < 0.001.

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terminal (BcNoxB) gfp-fusions were generated and introduced into B. cinerea B05.10 (B05.10:GFP-NoxA, B05.10:NoxB-GFP). The constructs were controlled by the constitutive ohiC promoter. Conidia of strains containing NoxB:GFP and GFP:NoxA, respectively, were allowed to germinate on a glass slide and the germlings were then examined by epifluorescence microscopy (Fig. 6A and B). Surprisingly, despite differing functions, BcNoxA and BcNoxB both localize to similar structures. The strongest fluorescence was visible for dynamic, and to some extent spherical, structures within the hyphae. To identify these structures germlings were stained with ER-Tracker™ Blue-White DPX. Co-localization of the ER-tracker and GFP-NoxA showed partial overlap, with accentuated localization of NoxA around the nuclei. For NoxB-GFP colocalization with the ER-tracker showed largely consistent signals (Fig. 6A and B). Additionally germlings were stained with Hoechst 33342. The fluorochromic property of this dye is attributed to its ability to bind with DNA structures, but it can also stain cell walls and septae, depending on the pH [35]. Hoechst staining showed that the structures visible seemed to enclose the nuclei (Fig. S5). A similar localization pattern has been described for the endoplasmatic reticulum (ER) membrane system in connection with the outer membrane of the nuclear envelope [36]. In addition, faint fluorescence was at times visible in both strains at the plasma membrane surrounding the hyphae (Fig. S5).

Due to its homology to the mammalian Nox regulator subunit p67phox NoxR is thought to be necessary for regulation of NoxA and NoxB [16]. Furthermore, it was previously shown that NoxR is connected to NoxA and NoxB activation [15, 18, 20]. Therefore we studied the localization of the regulator BcNoxR by introducing a GFP-BcNoxR fusion construct in B. cinerea B05.10. Germinated conidia were examined as described above for this construct and it strikingly showed a quite distinct localization from the previously observed localization of BcNoxA and BcNoxB (Fig. 6C). BcNoxR most prominently localized to cellular granules distributed irregularly throughout the hyphae. These granules seemed to move randomly within the hyphae (Brownian motion), however, time lapse showed that the granules had the tendency to stay within the young parts of the hyphae, close to the hyphal tips (see movie S1(472,778),(554,857)). In addition, in growing hyphae BcNoxR localized also to hyphal tips, where it seemed to be associated with the origin of outgrowth and the growth direction (Fig. 6C). In some of these

Figure 4. Penetration assay on onion epidermis. The hydrophobic side of onion epidermis was inoculated with 10 μl droplets of conidial suspensions (5x10^6 spores/ml). After 24 h incubation at 18°C lactophenole blue staining was used to visualize the fungus on top of the onion epidermis. Fungal cells within the onion cells are not stained. Conidia of the WT and ΔbcnoxA form a short germ tube with a terminal thickening, which are able to directly penetrate the plant surface. Conidia of the deletion strains ΔbcnoxB, ΔbcnoxA, ΔbcnoxB and Δbcpls1 germinate on the plant epidermis and differentiate appressoria-like structures, but fail to penetrate the plant via these structures. Black arrows indicate appressoria-like structures. Scale bars = 10 μm.

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hyphae additional material coming from the granules was transported to the growing hyphal tip (see movie S2).

In order to make sure that the used gfp-fusion constructs are functional, complementation studies were performed. The constructs GFP-NoxA and GFP-NoxR showed similar localization patterns in the deletion mutants as the respective construct in the WT (Fig. S4) and complemented their phenotypes (Fig. 7). No transformants were obtained with the BcNoxB-GFP construct in D bcnoxB background.

Since in Botrytis the deletion mutants D bcpls1, D bcnoxA, D bcnoxB and D bcnoxR show similar penetration defects on onion epidermis and bean leaves, respectively, we also performed localization studies with BcPls1. Therefore an N-terminal fusion of bcpls1 and gfp was transformed into the B05.10 WT and the D bcpls1 deletion mutant (B05.10:GFP-BcPls1, D bcpls1:GFP-BcPls1). The construct was controlled by its native promoter and hence showed its native expression pattern. Accordingly, the phenotype of the deletion mutant could be restored (Figs. 1 and 3). To observe localization of the protein, the conidia were treated as described above and analyzed microscopically. These analyses revealed that BcPls1 localized to similar intracellular membrane structures (presumably ER and plasma membrane) as BcNoxB and BcNoxA (Fig. 8). However, while BcNoxB/BcNoxA localized to those membranes within the whole germling, BcPls1 was only detected in the last thickened compartments of some hyphae, which might be appressoria-like structures (Fig. 8A, see also movie S3). Nevertheless, localization to similar membrane structures might hint at a spatial connection between the Nox complex and BcPls1 during formation of specific differentiation structures (in this case appressoria-like structures).

In conclusion BcNoxA, BcNoxB and BcPls1 localize to the ER and at times also to the plasma membrane, while strikingly BcNoxR is, under the tested conditions, found in different structures: it localizes to growing hyphal tips and in cellular granules within the hyphae.

**Discussion**

Following our first report on the Nox complexes in B. cinerea [20], we present here a deeper analysis of the functions of the complex components focusing on the localization of the catalytic NADPH oxidase subunits BcNoxA and BcNoxB and their regulatory subunit BcNoxR. Furthermore, we initiated a detailed analysis and localization studies of the tetraspanin Pls1 of B. cinerea in strain B05.10, which had been suggested to be connected to NoxB, due to their associated distribution in fungal genomes and similar phenotypes of the respective deletion mutants in P. anserina [26].

Little is known about the localization of Nox complexes in filamentous fungi: only in M. owygae has Nox1 been shown directly to localize at the appressorium during pathogenic development. We showed here that the catalytic subunits BcNoxA and BcNoxB in B. cinerea both localize to the nuclear envelope, the ER and at times to the plasma membrane, whereat BcNoxA showed more
concentrated localization around the nuclei, while BcNoxB localization was more dominant within the ER. Since Nox are transmembrane proteins postulated to be located in the outer membrane and to produce extracellular ROS, an ER localization of the complex was not assumed. However, until now there is no clear evidence for extracellular ROS derived from Nox. During pathogenic development extracellular ROS could either be used directly to attack the host, or they could serve as messenger molecules that are perceived to activate internal signaling processes. Interestingly, in the bcnox deletion mutants (and in Δbcplsl) decreased ROS levels were neither detected in axenic culture by DAB staining or use of Amplex Red nor in planta using NBT staining [20]. In fact for ΔbcnoxA and Δbcplsl even a moderate enhancement was detected, which was also previously reported for other fungi [20,21,24]. The observed reduction of H$_2$O$_2$ production after DPI treatment substantiates the unspecific effect of this inhibitor and supports the now generally accepted idea that secreted ROS in fungi is not produced by Nox but by alternative flavoenzymes. Hence, in B. cinerea a contribution of the Nox to extracellular ROS production is doubtful, and accordingly, the Nox might localize to a compartment other than the outer membrane. Even though we show a localization of BcNoxA and possibly BcNoxB in the ER, both proteins do not possess any ER retention signals. However, retention of a protein in the ER can

**Figure 6. Cellular localization of BcNoxA, BcNoxB and BcNoxR.** Protein localization was determined by epifluorescence microscopy in germlings of the strains WT:GFP-NoxA, WT:NoxB-GFP and WT:GFP-NoxR expressing gfp-bcnoxA, bcnoxB-gfp and gfp-noxR gene fusions, respectively. 10-μl droplets of a conidial suspension in GB5 medium (10$^5$ conida/ml) were placed on a glass slide and allowed to germinate over night before microscopic analyses were performed. A: BcNoxA localized to intracellular membrane structures and at times also to the plasma membrane. Staining with the ER-Tracker™ Blue-White DPX and a respective overlay show that the intracellular structures are partly consistent with the ER (from left to right: GFP-NoxA, ER-tracker, overlay, white light). B: BcNoxB localized to similar intracellular membrane structures and to the plasma membrane as visible for BcNoxA. Staining with the ER-Tracker™ Blue-White DPX and a respective overlay show that the intracellular structures visual are similar to ER structures (from left to right: NoxB-GFP, ER-tracker, overlay, white light). C: BcNoxR accumulated in cellular granules, which were distributed irregularly throughout the hyphae (left) and at the hyphal tip, where it seemed to determine the point of outgrowth (right, see also movie S2). Scale bars = 10 μm.

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result from various factors like the lack of a sorting signal, the presence of a signal for retrieval from pre-Golgi compartments or properties of the transmembrane domain (TMD) [37]. The exact mechanism of ER retention remains to be elucidated [38–40]. Integral predictions of protein locations (ProtComp v. 9.0) were done resulting in a prediction of all analyzed Nox proteins (S. sclerotiorum, M. oryzae, E. festucae, N. crassa, A. nidulans, P. anserina, B. cinerea, Trichoderma reesei, Homo sapiens) at the plasma membrane (see Table S1), except Nox2 from T. reesei which is predicted at the ER. The fact that the H. sapiens Nox1, 2 and 4 have repeatedly been shown to localize and function at the ER [41–42] shows that software based localization predictions may differ from real localization. Recently, the yeast Nox Yno1 was also shown to be present in the ER [11] like mammalian Nox1, Nox2 and Nox4 [41,42]. Besides a primary localization in the ER [43], Nox4 was reported to translocate to the plasma membrane when complexed with the stabilizing component p22phox [44], revealing that localization and function of a protein might not be restricted to a specific compartment or that proteins can be located in the ER for a period of time until they are needed and released. For example the chitin synthase 2 (Chs2) is expressed in the metaphase and retained in the ER through phosphorylation until further mechanisms stop this phosphorylation and release Chs2 from the ER [45–48]. According to these findings it is difficult to decide whether the ER is the final location of the catalytic BcNox subunits, or whether they are stored and modified there in order to be translocated. The fact that the GFP-BcNoxA fusion construct complemented the phenotype of the deletion mutant proves the functionality of the construct and supports a correct localization. However, the function of BcNoxA and BcNoxB within the ER is still unclear. A link between the ER and Nox might be the protein disulfide isomerase (PDI), which is a dithiol–disulfide oxidoreductase chaperone. In mammalian systems Nox1, Nox2, Nox1 and p22phox were linked to PDI [49,50]. PDI carries out different

Figure 7. Complementation analyses of ΔbcnoxA and ΔbcnoXR with gfp-constructs. Strains ΔbcnoxA:GFP-NoxA and ΔbcnoXR:GFP-NoxR expressing gfp-noxA and gfp-noxR gene fusions were tested for their phenotypical characteristics. The strains WT, ΔbcnoxA and ΔbcnoXR served as a control. The GFP-fusion constructs did complement the deletion phenotypes of ΔbcnoxA and ΔbcnoXR A: The ability to form sclerotia after 2 weeks incubation in darkness on CM media at 18°C is restored. B: The pathogenicity defect of the mutants is reinstated. Primary leaves of 10 day old French bean plants were inoculated with 7 μl droplets of conidial suspensions (2×10⁷ spores/ml in Gamborgs B5+2% glucose). Progression of infections was documented 3, 4 and 6 days after the infection (dpi).

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activities, amongst others it serves as a protein-folding catalyst for intramolecular relocation of disulfide bonds [51,52]. The main interactor of PDI is the ER flavoprotein Ero1 (ER oxidoreductin 1) [53]. Together, PDI and Ero1 play an essential role in oxidative protein folding [54] and they integrate ER function to cellular redox homeostasis [51]. Therefore the superoxide producing Nox might be involved in maintaining the redox level within the ER in connection with PDI (see also Fig. 9).

In contrast to the catalytic subunits, their regulator BcNoxR localized to cellular granules and the hyphal tip. An interaction, and therefore a similar localization, of NoxA and NoxA/B seems probable as they are homologs to the mammalian Nox2 and p67phox subunits, which come together to form the active complex [16]. Nevertheless, attempts to show a direct interaction between NoxR and NoxA or B in fungi have so far been unsuccessful. BiFC and yeast two-hybrid analysis (based on the split-ubiquitin system) couldn’t proof this interaction in B. cinerea so far [U. Siegmund and P. Tuzdynski, unpublished data]. It has to be taken into account that the formation of the active NADPH oxidase complex is a strictly regulated mechanism, which – in fungi – has not been completely elucidated. The interaction, and therefore similar localization, may only take place under specific conditions in the cell, when the active complex is needed. Thus BcNoxR might be stored in cellular compartments (the visible granules) and upon Nox activation translocates towards the ER-membrane where it interacts with the catalytic subunits. Alternatively, the catalytic subunits BcNoxA and B could have BcNoxR-independent roles, as shown for NoxA in mammals [43] (see also Fig. 9).

Another possible role for Nox worth investigating in the future could be their involvement in cytoskeleton organization. This was previously postulated for the mammalian system [55] and the recently identified yeast Nox (Yno1) was shown to have an impact on the cell cycle and possibly influences actin assembly [11].

Clearly not all the components of fungal Nox complexes have been identified yet. The previously postulated link between NoxB and Pls1 (similar phenotypes and linked presence in fungal genomes [26]) encouraged us to investigate this connection. Here, we show that as in B. cinerea T4 BcPls1 is also necessary for appressoria-mediated penetration in the more aggressive strain B05.10. However, in contrast to Pls1 in T4, lack of BcPls1 in B05.10 does not lead to apathogenicity. The infection rate is reduced, but not completely blocked, and once the fungus has entered the host, infection is comparable to the WT. Preliminary expression data supports an involvement of BcPls1 in formation of appressoria (6–24 hpi), as bepl1 is primarily expressed at 12 and 24 hpi (J. Heller and J. van Kan, unpublished data). This fits to the localization studies presented here indicating the presence of BcPls1 in the ER and plasma membrane of hyphae only in the last

Figure 8. Cellular localization of the protein BcPls1. Protein localization was determined by confocal laser scanning microscopy (CLSM) in germlings of the strain Δbcpls1:gfp-Pls1 expressing a gfp-bcpls1 gene fusion. 10-μl droplets of a conidial suspension in GB5 medium (10^5 conida/ml) were given on a glass slide and were allowed to germinate over night before microscopic analyses were performed. A: Overview of BcPls1 localization in a Botrytis cinerea germling. BcPls1 mainly localizes to the border of circular structures in germinating hyphae that seem to be coherent (middle row). Higher magnification showed that the protein also localizes to structures at the edge of the hyphal tip (top row). Fluorescence is clearly limited to the last segment of the germling. The septal pore shows no fluorescence (white arrow, bottom row). B: Germlings were treated with the endocytosis marker FM4-64 (red, top right panel) to stain intracellular membranes. BcPls1 localization is shown in green (top left panel). White arrows indicate co-localization of BcPls1 fluorescence and the FM4-64 stained plasma membrane (see overlay, bottom left panel). Yellow arrows indicate localization of BcPls1 in membrane structures that are not stained by FM4-64 (indicative for the nuclear envelope/endoplasmic reticulum). C: Germlings were treated with the dye Hoechst (blue, top right panel) to stain nuclei. BcPls1 localization is shown in green (top left panel). White arrows indicate localization of BcPls1 to the interior space of Germlings were treated with the dyes FM4-64 (red, second row) and Hoechst (blue, third row), simultaneously. BcPls1 localization is shown in green (first row). In addition to the membrane structures seen before (nuclear envelope), white arrows indicate localization of BcPls1 to structures at the edge of the hyphal tip (top row). Fluorescence is clearly limited to the last segment of the germling. The septal pore shows no fluorescence (white arrow, bottom row).

The NADPH Oxidase Complexes in Botrytis cinerea
a separated physiological unit, sealed by the first septum. The precise role of Pls1 remains to be elucidated. Besides the impediment during appressoria formation, differentiation and growth of the bcpls1 mutant appears normal. Interestingly, Δbcpls1 forms sclerotia in WT-like manner, but crossing experiments showed that these sclerotia are sterile. This is an uncommon phenotype, because most female sterile mutants in *B. cinerea* thus far described, with the exception of the Δbcfrp1 mutant, fail to form any sclerotia [57].

There are several overlapping phenotypes of ΔbcnoxB and Δbcpls1 supporting the assumed functional link of both proteins, the most obvious one being the penetration defect. However, there are slight differences: while in the *bcnoxB* mutant primary lesion formation is generally retarded [20], Δbcpls1 is able to penetrate in 50% of the cases 3 dpi like the WT, in the other cases primary lesion formation is also delayed. Nevertheless, both mutants once they have entered the host are able to colonize tissue like the WT. More detailed analyses of the penetration 18–24 hpi on onion epidermis and French bean leaves revealed a similar defect of Δbcplsl and ΔbcnoxB resulting in appressoria formation and repeated outgrowth instead of penetration as observed for the WT. In contrast, the ΔbcnoxA mutant penetrates the host like the WT. Another observation supporting the functional link of BcNox and BcPls1 is the ability of the mutants to form hyphal fusions via CATs. Deletion mutants of *bcnoxA*, *bcnoxB* and the double mutant *bcnoxA/B* do not form any hyphal fusions, while ΔbcnoxB is able to form fusions (though with reduced frequency) [34]. Nonetheless, the fact that Δbcplsl forms WT-like hyphal fusions again relates BcPls1 to BcNoxB rather than to BcNoxA. BcNox and BcPls1 have also corresponding characteristics regarding their control of female fertility. Δbcplsl produces normal looking sclerotia, but crossing experiments revealed that these sclerotia are sterile. As the *bcnoxA* mutants are all impaired in sclerotial development (although ΔbcnoxB still produces smaller sclerotia) they are also regarded as female sterile [20]. A final link between BcNox and BcPls1 is their similar localization in the ER and the plasma membrane, although for BcPls1 this localization is only found in the last (apical) segment of the hyphae. This could suggest that the BcNox complexes are involved in different mechanisms controlled by differential regulation systems and that BcPls1 is involved in the activation of BcNoxB e.g. during early stages of infection.

Taken together we have shown that the catalytical Nox subunits BcNoxA and BcNoxB as well as the tetraspanin Pls1 are localized in the ER and the plasma membrane. A distinct localization of the regulator BcNoxR at the hyphal tip and in cellular granules suggests translocation of the cytosolic subunits to the catalytical subunit (NoxA/B) only when the active Nox complex is needed. A better understanding of the activation and assembly mechanisms of the fungal Nox complexes in more detail will be in the focus of further investigations.

**Materials and Methods**

**Fungal Strains**

*B. cinerea* ([teleomorph] *Botryotinia fuckeliana* (de Bary) Whetzel) strain B05.10 is a putative haploid derivate of SAS56, an isolate obtained from *Vitis vinifera* after benomyl treatment [58]. This strain is used as a host strain for gene replacement experiments and localization studies, and as a wild-type control in all experiments. All other strains used in this study are listed in Table 1.

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**Figure 9. Hypothetical schematic overview of the localization and putative functions of Nox in *B. cinerea*.** Nox in *B. cinerea* might function at two different locations. On the one hand (top) they are located in the ER (green) membrane, where they possibly contribute to the ER redox status. Besides GSH (glutathion), Nox might be a second supplier of electrons to PDI. PDI is oxidized by Ero1 by transfer of electrons to O₂, which in turn becomes H₂O₂ (scheme of Ero1-PDI supplier of electrons to PDI). Ero1 is oxidized by Nox oxygen to superoxide, using NADPH as an electron donor. The BcNoxA complex then transfers electrons through the membrane, reducing other so far unknown components (possibly Bem1 and Cdc24 [17]). This suggests translocation of the cytosolic subunits to the catalytical complex is necessary for CAT fusions, colonization of plant tissue and other strictly localized physiological reactions [56]. The results presented here indicate that the whole final hyphal compartment constitutes (apical) segment, especially in hyphae forming appressoria-like structures (see also movie S5). The infection structures in *B. cinerea* had been termed “appressoria-like” because they lack the typical closed septum allowing the generation of turgor pressure and other strictly localized physiological reactions [56].

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**Table 1.**

| Fungal Strains | Materials and Methods |
|---------------|-----------------------|
| *Botrytis cinerea* (teleomorph) *Botryotinia fuckeliana* (de Bary) Whetzel | Strain B05.10 is a putative haploid derivate of SAS56, an isolate obtained from *Vitis vinifera* after benomyl treatment [58]. This strain is used as a host strain for gene replacement experiments and localization studies, and as a wild-type control in all experiments. All other strains used in this study are listed in Table 1. |
Table 1. Strains used in this study.

| Strain | Genotype | Reference |
|--------|----------|-----------|
| B. cinerea SAS405 | MAT1-2 | [67] |
| AbcnoxA | B05.10ΔbcnoxA::nat1 | [29] |
| B05.10:gfp-noxA | B05.10::gfp-noxA hph | this study |
| AbcnoxA::gfp-noxA | AbcnoxA::gfp-noxA hph | this study |
| AbcnoxB | B05.10ΔbcnoxB::hph | [29] |
| B05.10::gfp-noxB | B05.10::gfp-noxB hph | this study |
| AbcnoxR | B05.10ΔbcnoxR::nat1 | [29] |
| B05.10::gfp-noxR | B05.10::gfp-noxR hph | this study |
| AbcnoxR::gfp-noxR | AbcnoxR::gfp-noxR hph | this study |
| AbcnoxB::gfp-noxR | AbcnoxB::gfp-noxR hph | this study |
| AbcplA | B05.10ΔabcplA::nat1 | [29] |
| AbcplA::gfp-pls1 | AbcplA::gfp-pls1 nat1 | this study |

Media and Culture Conditions

Yeast cells were either cultivated in complex media (YPD) (0.5% (w/v) yeast extract, 2% (w/v) glucose, 2% (w/v) peptone, pH 5.8) or for selection in selective drop-out (SD) medium (20 g/l glucose, 6.7 g/l Difco Yeast Nitrogen Base w/o amino acids [BD, Franklin Lakes, NJ USA], 0.77 g/l DO-supplement [Clontech, Mountain View, CA, USA], pH 5.8) at 30°C.

E. coli cells were grown in lysogeny broth (LB) medium (10 g/l peptone, 5 g/l yeast extract, 10 g/l NaCl) [59] at 37°C.

WT and mutant strains were grown on different complex media. Potato dextrose agar (PDA) (Sigma-Aldrich Chemie, Steinheim, Germany) was supplemented with 100 g/l homogenized leaves of French beans (PDA-B) (Phaseolus vulgaris). Synthetic complete medium (CM) was set according to Pontecorvo et al. [60]. Growth behavior was also tested on Czapek Dox minimal medium (CD) (20 g/l sucrose, 3 g/l NaNO₃, 1 g/l K₂HPO₄, 0.5 g/l KCl, 0.01 g/l FeSO₄·7H₂O, 0.5 g/l MgSO₄·7H₂O, pH 5.2). In order to obtain conidia, strains were incubated for 6–8 days at 18°C under light conditions (12 h light, 12 h darkness); for sclerotial production, strains were grown 3 weeks at 18°C in darkness. For DNA preparations, mycelium was grown 3–4 days at 18°C on CM agar overlaid with Cellophane. Stress sensitivity tests were performed by inoculating strains on CM plates supplemented with H₂O₂, menadione, sorbitol and NaCl.

Transformation

A knock-out construct for hpl61 was obtained using the homologous recombination system in yeast, as described previously [61]. The 5’- and 3’-regions of hpl61 (1178 bp and 724 bp) were amplified from genomic DNA using the primer pairs 7/8 and 9/10 (Table S2). These primers not only contain sequences to amplify the 5’- and 3’-regions of the corresponding genes, but also the sequences homologous to the resistance cassette and the yeast plasmid pRS426 [62]. The hygromycin resistance cassette containing the hph gene of E. coli under control of the trpC promoter of A. nidulans was generated with primers 11/12 using pCSN44 as a template. The three PCR fragments, together with the EcoRI/BglII linearized vector pRS426, were co-transformed into the Saccharomyces cerevisiae strain FY834 [63], where homologous recombination took place. Transformants were selected on SD plates lacking uracil. Total DNA from uracil-prototrophic yeast colonies was isolated with the GeneJET™ Plasmid Miniprep Kit (Fermentas), including an initial step to disrupt yeast cells by addition of glass beads. The DNA was then used as template to amplify the hpl61 replacement construct by use of primers 7/10 (Table S2). The complete replacement fragment was used to transform B. cinerea strain B05.10.

For construction of the gfp-hpl61 complementation construct, a nourseothricin resistance cassette had to be integrated into the existing vector pBHT2-GFP-BePls1 (Mathieu Gourgues and Marc-Henri Lebrun, unpublished data). The nourseothricin resistance cassette from the vector pEH1nat1 was amplified using the primers 13/14 and cloned into the vector pSTBlue1 for sequence confirmation. From the vector pSTBlue1 the resistance cassette was excised using the restriction enzyme EcoRI and cloned into the EcoRI-digested pBHT2-GFP-BePls1. The resulting vector was linearized using the restriction enzymes PstI, SalI and SpeI and transformed into strain Beplsl1.

For construction of the gfp-bcnR, gfp-noxA and bcnoxB-gfp fusion constructs the homologous recombination system in yeast was also used [61]. The genes bcnR, bcnoxA and bcnoxB were amplified using the primers 15/16, 19/20 and 17/18, respectively. These primers contain sequences homologous to the glucanase terminator and the gfp of the vector pNAH-OGG (bcnoxB, bcnoxA) and to the oblC promoter and gfp of the vector pNaN-OGG (bcnxR) [64]. These vectors include the bcnR-Banks to replace the nitrite reductase-encoding gene of B. cinerea, ensuring a defined integration site. Deletion of this gene has no effect on B. cinerea, provided nitrogen sources other than nitrate are available [64]. The PCR products were co-transformed into the uracil-prototrophic S. cerevisiae strain FY834 with pNaN-OGG linearized with NodI or pNaN-OGG linearized with NosI, respectively, as described above. Plasmid DNA from uracil-prototrophic yeast colonies was isolated using the SpeedPrep Yeast plasmid isolation kit (DualsystemsBiotech, Switzerland) and transformed into E. coli. Single plasmids were isolated, and after sequencing the expression cassette was excised using the enzymes PstI/SalI (bcnoxA, bcnoxB plasmids) and PstI/SalI (bcnxR plasmid) and transformed into strain B05.10.

For transformation of B. cinerea, protoplasts were generated using a mixture of glucanase (Novozymes, Denmark), lysing enzyme (Sigma-Aldrich, St Louis, MO, USA) and yatalase (Takara Bio Inc, Shiga, Japan). Protoplasts were transformed according to Schulze-Gronover et al. [65] using 300 μl of PCR-product (knock-out) or 25 μg of the linearized vector (GFP-fusions), respectively. Resistant colonies were transferred to agar plates containing GB5 agar, supplemented with 70 μg/ml of hygromycin B (Invitrogen, San Diego, USA) or 70 μg/ml of nourseothricin [Werner-Bioagents, Jena, Germany]. Single conidial isolates were obtained by spreading conidial suspensions on GB5 plates containing 70 μg/ml of hygromycin B or 70 μg/ml of nourseothricin. The conidia were germinated and single colonies transferred individually to new plates containing the selection marker. Homologous integration of the different transformation constructs was proven by diagnostic PCR (Primers see Table S2) and Southern blots.

Standard Molecular Methods

Fungal genomic DNA was isolated as described by Cenis [66]. Southern blot analysis was performed according to Sambrook et al. [67]. Hybridisation was carried out in 6× SSPE, 5× Denhardt’s solution, 0.1% SDS, and 50 mM phosphate buffer, pH 6.6, at 65°C for 16 to 20 h in the presence of a random-primed [α-32P]dCTP labeled probe. Filters were washed for 10 min in 2× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate), 0.1%
sodium dodecyl sulfate (SDS) and for 10 min in 1 x SSC, 0.1% SDS. Hybridization and washing of the filters were carried out at 65 °C.

**Sexual Crosses**

Crosses were performed as described by van der Vlugt-Bergmans et al. [68]. Cap structures of mature apothecia were collected and placed on a glass microscope slide in a droplet of sterile water. The cap was crushed with a second microscope slide in order to release asci and ascospores. The sample was suspended in 2–3 ml of sterile water and filtered over glass wool to remove large debris. The resulting flow-through contained a suspension of ascospores which was dispersed on appropriate media for further analysis.

**Conidial Germination Tests**

For analyses of conidiospore germination on glass surfaces in nutrient dependency [69], conidia were harvested from CM agar plates, filtered over Nyrex and suspended in 10 ml of sterile water and washed three times. A conidial suspension [25 μl] with a concentration of 5 x 10⁶ spores/ml was placed in the center of a cover slip and 475 μl of B5 nutrient solution were added to the conidia supplemented with glucose at a final concentration of 10.5 mM. The cover slips were incubated in the dark at 20 °C as indicated. Germination progress was monitored by light microscopy after 3, 6 and 24 h.

**Vegetative Hyphal Fusion Assay**

Vegetative hyphal fusions were tested as described by Roca and Weichert et al. [34]. Conidial suspensions were plated on Vogel’s minimal medium [70] and incubated for 14 h. Samples were analyzed using differential interference contrast (DIC) microscopy. For quantification three independent replicates were performed evaluating at least 300 spores each time.

**Pathogenicity Assays**

Infection assays were performed with conidia from 10-day-old PDAB-agar cultures. Primary leaves of *Phaseolus vulgaris* L. genotype N90598 (originating from J. D. Kelly, Michigan State University, East Lansing, MI) were inoculated with 7.5 x 10⁶ conidia/ml in GB5 medium supplemented with (NH₄)₂HPO₄ (1 mM). Germinated conidia were analyzed microscopically 24 h after inoculation with an inverted microscope (Leica DMIRE2) equipped with a Leica TCS SP2 laser scanning device (Leica Microsystems) using a 63 x water-immersion lens. GFP fluorescence was excited using a 488 nm laser line. Images were collected with a resolution of 8 bit using an emission range between 705 nm and 750 nm with a frame average and a line average of 4. For FM4-64 staining a 1:1000 dilution of FM4-64 (Invitrogen, USA) in GB5 medium was added to the germinating conidia. After a short incubation time FM4-64 fluorescence was excited using a 488 nm laser line. Images were collected with a resolution of 8 bit using an emission range between 420 nm and 470 nm.

**Epifluorescence Microscopy**

A glass slide was inoculated with 10 μl of a conidial suspension (1 x 10⁷ conidia/ml) in GB5 medium supplemented with glucose. Germinated conidia were analyzed 12–24 h after inoculation microscopically with Axio Imager 2 (Zeiss, Jena, Germany) using a 63 x objective lens or Observer Z.1 (Zeiss, Jena, Germany) using a 20 x objective lens.

Hoechst staining was conducted as described for CLSM. For staining of the ER a 1:5 dilution of ER-Tracker Blue-White DPX (Life Technologies, Germany) in Mc Ilvaine standard buffer was added to the germinating conidia. Hoechst staining and the ER-Tracker were examined using the filter set 49 DAPI shift free (excitation G 365, beam splitter FT 395, emission BP 445/50), GFP fluorescence was detected with filter set 38 (excitation BP 470/40, beam splitter FT 495, emission BP 525/50). Images were captured with a Zeiss AxioCam MRm camera and analyzed using the Axiovision Rel 4.8 software package.

**ROS Staining**

In order to stain exogenous H₂O₂ the strains to be tested were grown on CM plus Cellophane for 3 days. Afterwards, 25 mg of fresh mycelium were over-layered with 1 ml DAB solution (0.5 mg/ml DAB in 100 mM citric acid pH 3.7). The incubation took place for 1.5–2 h in the dark at RT, the evaluation was done visually. Only DAB solution, and DAB solution supplemented with 1 μl H₂O₂ (30%) served as a negative controls. As a positive
oxidative stress was induced using 5 mM H₂O₂, and the WT. Production of H₂O₂ was monitored using DAB solution for 1.5 h in the dark. As negative controls, one well was added to the media and turns yellow with decreasing pH value. Asterisks above the bars denote significant differences in the measurements of the indicated strains to the WT. * = P<0.05; ** = P<0.01.

**Figure S4** Cellular localization of BcNoxA and BcNoxR in the respective deletion mutants. Protein localization was determined by epifluorescence microscopy in germlings (in Gamborgs B5+2% glucose) of the strains ΔbcnoxA-GFP-NoxA and ΔbcnoxB-GFP-NoxB expressing gfp-bcnoxA and gfp-bcnoxB gene fusions, respectively. A: BcNoxA localized to intracellular membrane structures and at times also to the plasma membrane. B: BcNoxB accumulated in cellular granules, which were distributed irregularly all over the hyphae. Scale bars = 10 μm.

**Supporting Information**

**Figure S1** Deletion of the gene bcpls1. A: Schematic overview of the deletion strategy. For the detection of homologous integration of the replacement fragments primers were used that bind upstream the 5′-region and within the terminator of the resistance cassette (P3/P4; 5′), or in the promoter of the resistance cassette and downstream the 3′-prime region (P5/P6; 3′), respectively. For detection of purified mutants, primers binding within the wild-type bcpls1 were used (P1/P2; WT). For Southern blot analyses genomic DNA was digested using the restriction enzyme NcoI that cuts within the resistance cassette but not within bcpls1. Using the 5′ flank as a probe, in the wild-type a fragment larger than 5 kb was expected and in the mutant a fragment of 3.7 kb. B: Diagnostic PCR showing homologous integration of the knock-out fragment at bcpls1. C: Southern blot showing no further integrations of the knock-out fragment.

**Table S1** Predicted localization of Nox proteins from various organisms. Protein sequences of NoxA/1 and NoxB/2 from B. cinerea, A. nidulans, M. oryzae, S. sclerotiorum, E. festucae, P. anserina, N. crassa and T. reesi as well as Nox1, Nox2 and Nox4 from Homo sapiens were used to predict their cellular localization using ProtComp v. 9.0 (http://linux1.softberry.com/berry.phtml?topic = protcompan&group = programs&subgroup = protloc).

**Figure S2** Comparison of growth and differentiation of B. cinerea B05.10 WT and Δbcpls1. A: For analysis of light-dependent differentiation, strains were cultivated on CM for 7 days with a rhythm of 12 h light and 12 h darkness (light) or for 3 weeks in complete darkness (dark). Δbcpls1 shows no differences to the WT. B: Plate assays showing growth rates of B. cinerea B05.10 and Δbcpls1 on stress causing media. As a control CM was used, oxidative stress was induced using 5 mM H₂O₂, 10 mM H₂O₂ and 500 mM manodine, osmotic stress was induced using 1 M NaCl and 1 M sorbitol. Colony diameters were measured 3 days after the inoculation. The indicated values are means of five different plates; standard deviations are indicated by the error bars. Asterisks above the bars denote significant differences in the measurements of the indicated strains to the WT. * = P<0.05; ** = P<0.01; *** = P<0.001.

**Figure S3** Production of H₂O₂ and oxalic acid in Δbcpls1, ΔbcnoxA, ΔbcnoxB, ΔbcnoxAB, ΔbcnoxR and the WT. A: Production of H₂O₂ was monitored using DAB staining. 25 mg of fresh mycelia were placed in each well to exclude staining differences evolving from differing growth rates of the strains. Strains were previously grown on CM overlayed with cellophane for 3 days. The mycelium was inoculated with DAB solution for 1.5 h in the dark. As negative controls, one well was only filled with DAB solution (-) and one with DAB solution and 1 μl H₂O₂ (-). DAB solution, 1 μl H₂O₂ and 1 μl horseradish peroxidase served as a positive control (+). B: Quantitative analysis of H₂O₂ production using Amplex Red Peroxidase Assay. 20 mg of fresh mycelia were incubated in the Amplex Red working solution. Strains were previously grown on complete medium (CM) overlayed with cellophane for 3 days, for inhibition of flavonenzymes 100 μM DPI were added to the medium. Fluorescence emission was detected after 30 min at 590 nm (excitation 560 nm).

**Table S2** Oligonucleotide primers used in this study.
the hyphal tip. Additional material is added into cellular granules coming from the back of the hyphae.

(AVI)

Movie S3 Cellular localization of the protein BcPsl1 in growing hyphae. Protein localization was determined by epifluorescence microscopy in germinating conidia (0–14 hours after inoculation in Gamborgs B5+2% glucose) of the strain Abcpl1::gfp-psl1 expressing a gfp-bcpl1 gene fusion. GFP fluorescence can be detected from 12 hours in the apical segment of hyphae, showing thickening and growth arrest, which develop into appressoria-like structures.

(AVI)

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Author Contributions

Conceived and designed the experiments: US JH JvK PT. Performed the experiments: US JH JvK. Analyzed the data: US JH JvK PT. Contributed reagents/materials/analysis tools: US JH JvK PT. Wrote the paper: US JH PT.
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