Fungal plant pathogens secrete proteins that manipulate the host in order to facilitate colonization. Necrotrophs have evolved specialized proteins that actively induce plant cell death by co-opting the programmed cell death machinery of the host. Besides the broad host range pathogen *Botrytis cinerea*, most other species within the genus *Botrytis* are restricted to a single host species or a group of closely related hosts. Here, we focused on *Botrytis squamosa* and *B. elliptica*, host specific pathogens of onion (*Allium cepa*) and lily (*Lilium* spp.), respectively. Despite their occurrence on different hosts, the two fungal species are each other’s closest relatives. Therefore, we hypothesize that they share a considerable number of proteins to induce cell death on their respective hosts. In this study, we first confirmed the host-specificity of *B. squamosa* and *B. elliptica*. Then we sequenced and assembled high quality genomes. The alignment of these two genomes revealed a high level of synteny with few balanced structural chromosomal arrangements. To assess the cell death-inducing capacity of their secreted proteins, we produced culture filtrates of *B. squamosa* and *B. elliptica* that induced cell death on their respective hosts. In this study, we first confirmed the host-specificity of *B. squamosa* and *B. elliptica*. Then we sequenced and assembled high quality genomes. The alignment of these two genomes revealed a high level of synteny with few balanced structural chromosomal arrangements. To assess the cell death-inducing capacity of their secreted proteins, we produced culture filtrates of *B. squamosa* and *B. elliptica* that induced cell death responses upon infiltration in host leaves. Protein composition of the culture filtrate was analysed by mass spectrometry, and we identified orthologous proteins that were present in both samples. Subsequently, the expression of the corresponding genes during host infection was compared. RNAseq analysis showed that the majority of the orthogroups of the two sister species display similar expression patterns during infection of their respective host. The analysis of cell death-inducing proteins of *B. squamosa* and *B. elliptica* provides insights in the mechanisms used by these two *Botrytis* species to infect their respective hosts.

**KEYWORDS**
*Botrytis* spp., plant-necrotroph interaction, host specificity, cell death-inducing proteins, synteny
Introduction

In the battle for survival, plant pathogenic fungi have evolved different strategies to colonize their hosts. While biotrophic fungi obtain nutrients from living host cells via specialized feeding structures, necrotrophs kill the cells of their host to acquire nutrients. For a long time, fungi with a necrotrophic lifestyle have been considered to invade their hosts in a rather unsophisticated manner. Using lytic and degradative enzymes, the fungus first kills and subsequently grows into host cells and eventually colonizes host tissue. In the past decades however, evidence has been accumulating that the interaction between necrotrophs and their hosts is more subtle and sophisticated than previously appreciated. Necrotrophs actively induce plant cell death by co-opting the programmed cell death machinery of the host using specialized effector proteins, to a certain extent similar to biotrophs (Dickman et al., 2001; van Kan, 2006). The induction of plant cell death using proteinaceous effectors is vital for the success of infection (Tan et al., 2010; Stergiopoulos et al., 2013; Veloso and van Kan, 2018). For some necrotrophic effectors the corresponding plant receptor proteins required for effector recognition and triggering of the cell death response have been identified (Schürch et al., 2004; Abeysekara et al., 2009; Gao et al., 2015; Shi et al., 2016). The cell death response to effectors of necrotrophs is based on similar mechanisms as the recognition of effectors from biotrophic pathogens, but leads to disease susceptibility instead of disease resistance as is the case for biotrophs (Lorang et al., 2007; Tan et al., 2010).

The role of effectors of necrotrophs in plant-pathogen interactions is best studied in Septoria nodorum blotch in wheat, caused by the Ascomycete Parastagonospora nodorum. The first identified effector from P. nodorum was SnToxA, a small secreted protein that induced necrosis upon infiltration in wheat lines carrying the sensitivity gene Snn1 (Liu et al., 2004). SnToxA knockout mutants lost virulence on Snn1-carrying wheat lines, and introduction of SnToxA into an avirulent P. nodorum isolate conferred pathogenicity on wheat lines harbouring Snn1 (Liu et al., 2012). By genetic analysis, more P. nodorum effectors and corresponding wheat sensitivity genes were identified such as SnToxA-Tsn1, SnToxA-Snn4, and SnToxA-Snn6 (Abeysekara et al., 2009; Faris et al., 2010; Tan et al., 2012; Gao et al., 2015). To date, nine necrotrophic effectors and corresponding host sensitivity loci have been identified in the P. nodorum-wheat pathosystem (Duba et al., 2018; Cowger et al., 2020). The severity of infection depends in a quantitative manner on the number and identity of pathogen effectors and corresponding wheat sensitivity genes (Oliver et al., 2012; Phan et al., 2016; Rud et al., 2017; Haugrud et al., 2019).

The broad host range model necrotroph Botrytis cinerea is also known to secrete proteins that induce cell death upon recognition by the host. B. cinerea endopolygalacturonases (PGs) induce cell death upon infiltration in Arabidopsis thaliana, but only in lines harbouring leucine-rich-repeat (LRR) receptor-like protein RBPG1 (Zhang et al., 2014). Other proteinaceous virulence factors that function as cell death-inducing effectors in B. cinerea are BcXyl11a, BcXyl1 and BcXYG1, hemicellulases that induce necrosis in plant tissue independent from their catalytic activity (Noda et al., 2010; Zhu et al., 2017; Yang et al., 2018). Xylanases are recognized by the LRR receptor-like proteins LeEix1 and LeEix2 and the cell death response to BcXyl1 and BcXYG1 requires the LRR receptor-like kinases BAK1 and SOBIR1 (Ron and Avni, 2004; Zhu et al., 2017; Yang et al., 2018). In addition to the cell wall degrading enzymes with cell death-inducing activity, often considered as PAMPs or catalytic necrosis-inducing proteins (NIPs), B. cinerea also secretes cell death-inducing proteins that lack a known enzymatic domain and are referred to as effectors. Amongst others BcNep1, BcNep2, BcSpI and BcIEB1 have been heterologously produced and induced cell death upon infiltration in plant leaf tissue (Arenas et al., 2010; Frias et al., 2011; Frias et al., 2016).

The genus Botrytis comprises ~40 described species of which a few, including B. cinerea, are generalist plant pathogens with very broad host ranges (Hyde et al., 2014; Garfinkel et al., 2017). Most Botrytis species, however, are considered host-specific since they are pathogenic on a single host or a few taxonomically related hosts. An inventory by Mercier et al., 2019 of polyphagy indices of Botrytis species indicated that B. elliptica (commonly considered to be a pathogen specifically to lily and known as “lily fire blight”) was reported to infect as many as 21 host species from 10 distinct genera, and has a polyphagy index of 7.1. By contrast, its close relatives B. deweyae, B. sinoalli and B. squamosa, that cluster with B. elliptica in a subclade of the phylogenetic tree of the genus Botrytis (Valero-Jimenez et al., 2020), have polyphagy indices below 1.5 (Mercier et al., 2019). In this study we focussed on B. elliptica and B. squamosa, two closely related sister species that are reported to cause disease on lily and onion, respectively. For both species, induction of programmed cell death is a key step in the infection process and it is achieved by means of secreted proteinaceous effectors (Van Baarlen et al., 2004; Malvestiti et al., 2021; Steentjes et al., 2022). We aimed to analyse whether the shared secreted proteins between these two closely related species can contribute to cell death induction in their respective host and non-host plant. We verified the host specificity and sequenced and assembled high quality genomes of B. squamosa and B. elliptica and analysed their synteny. We compared the cell death inducing capacity of the culture filtrates of B. squamosa and B. elliptica, and analysed the protein composition of these culture filtrates and the expression of the corresponding genes during host infection.
Materials and methods

Plant material and growth conditions

Bulbs of Lilium spp. cultivar “Asiatic” were planted in plastic crates containing potting soil and grown in a greenhouse under natural day light at a minimum night temperature between 12 and 15°C and maximum day temperature between 24 and 26°C. Mature leaves as described by Bar and Ori (2014) were harvested before the flowering stage and used for disease and infiltration assays.

Nicotiana benthamiana plants were grown in potting soil in a greenhouse with 16 h light at 24°C and 8 hours darkness at 22°C and 75% relative humidity. Both disease and infiltration assays were carried out on leaves that were still attached to the plant. Allium cepa (cv Ceresco F1) plants were grown from seeds in potting soil in a climate chamber with 12h light, 70% relative humidity, 18°C day temperature and 16°C night temperature. Fully grown leaves of 8 to 10 weeks-old onion plants were used for inoculation and infiltration assays. Disease assays were conducted on cut onion leaves, while for infiltration assays the leaves remained attached to the plant.

Fungal material and growth conditions

Botrytis squamosa isolate MUCL31421 (Steentjes et al., 2021), Botrytis elliptica isolate 9401 (Malvestiti et al., 2021) and Botrytis cinerea isolate B05.10 (Van Kan et al., 2017) used in this research were stored as conidia suspensions in 20% glycerol at -80°C. To obtain conidia for inoculation, in this research were stored as conidia suspensions in 20% glycerol at -80°C. To obtain conidia for inoculation, Botrytis squamosa and Botrytis elliptica were obtained by growing each fungus in a 250 mL culture was inoculated with a spore concentration of 10⁵ conidia/mL and stored until use in darkness at 4°C.

DNA extraction, genome assembly and annotation

High molecular weight genomic DNA for Nanopore sequencing was extracted from in vitro grown mycelium as described in Valero-Jiménez et al. (2020). Both genomes of B. elliptica and B. squamosa were assembled with Oxford Nanopore reads. The adapters of the reads were firstly trimmed with pochepoch, version 0.2.4 (https://github.com/rwick/Porechop). NECAT (Chen et al., 2021) was then launched for genome assembly, with genome sizes of 48 Mb and 55 Mb for B. elliptica and B. squamosa respectively and other parameters were set as default. After the assembly, we polished the B. elliptica genome with Illumina short reads using Pilon (Walker et al., 2014) and the B. squamosa genome by aligning with the previously published genome assembly (Valero-Jiménez et al., 2020). Subsequently, the two genomes were aligned with MUMer 3.07 (Kurtz et al., 2004) using PROmer (-l 150, -c 500, -g 100, -b 200). The genome of B. elliptica was annotated using Funannotate (https://funannotate.readthedocs.io). The previously predicted secretome of B. elliptica strain 9612 was subsequently mapped to the new genome assembly and the missing secretome genes in the new assembly were manually added. The previously published annotation of B. squamosa of the same strain was mapped to the new assembly.

Production of culture filtrate, ammonium sulfate precipitation and leaf infiltration

B. squamosa and B. elliptica Culture Filtrate (CF) samples were obtained by growing each fungus in a 250 mL flask containing 50 mL of liquid medium with 3 g/L Gamborg B5 salts (Duchefa, Haarlem, Netherlands), 10 mM potassium phosphate pH=6.0, 0.1% D-Glucose and 5mL of lily or onion leaf extract in demi water, respectively. The leaf extract was obtained by grinding 30g of fresh harvested leaf material in 250 mL demineralized water. The homogenate was centrifuged (3500 rpm, 20 min) and the supernatant was concentrated by freeze drying. The concentrate was redissolved in 10mL water, filter-sterilized and added to the liquid medium. The liquid culture was inoculated with a spore concentration of 10⁵
conidia/mL medium. The cultures were incubated at 140 rpm, at 20°C in the dark. After 5 days of growth, the CF was passed through a layer of Miracloth (Calbiochem, San Diego, CA, United States), filter-sterilized (0.45 μm pore size, Millipore, Amsterdam, Netherlands) and kept on ice. A mock liquid medium was prepared without fungal conidia. Of the ca 40 mL harvested CF samples, 2mL were stored at 0°C for one night. To the remaining volume ammonium sulfate (LabChem, Tiel, Netherlands) was gradually added and mixed on a rolling bench at 4°C until the CFs were saturated with salt. Samples were centrifuged 30min at 4°C (4000rpm). After centrifugation a solid pellet of precipitated compounds was collected. The pellet was redissolved in 10mM potassium phosphate pH=6.0 and transferred to a prewetted dialysis membrane (Spectra/Por 3 Dialysis Membrane 3.5 kD MWCO, Repligen), and dialyzed overnight at 4°C in 5 L 10mM potassium phosphate pH=6.0. The dialyzed samples were collected and stored at 0°C until infiltration. The same plant genotypes used in the disease assays were tested for the infiltration assays. For lily, ca 100 μL of each sample (crude CFs, dialyzed pellets and mock liquid media) was infiltrated with a 1 mL syringe in two spots of two different detached leaves on their abaxial side. The leaves were placed in moist plastic boxes and incubated at 20°C under ambient light. The same volume of sample was infiltrated in two spots on two onion and N. benthamiana leaves whereby onion and N. benthamiana leaves were not detached from the plant. After 3 days incubation all leaves were photographed and the response was evaluated using red light imaging (Villanueva et al., 2021).

Proteomic analysis

B. elliptica and B. squamosa proteins that were analyzed with mass spectrometry were sampled from the liquid culture as described in the above section. Proteins were prepared for mass spectrometry as described in Supplementary Information 1. The samples were subjected to LC-MS/MS according to the parameters described in (Xiong et al., 2020). For protein identification and quantification, each run with all MS/MS spectra obtained was analyzed with Maxquant 2.0.3.0 with the Andromeda search engine. The B. squamosa and B. elliptica annotated genomes were used for protein mapping. A maximum of two missed cleavages and a mass deviation of 20 ppm for the fragment MS/MS peaks were allowed. The false discovery rate (FDR) was set to 1% on both peptide and protein level. The length of peptides was set to at least seven amino acids. Protein identification required minimally two distinct peptides of which at least one unique and at least one unmodified. Intensity based absolute quantification (iBAQ) algorithm was calculated as the sum of all peptide intensities divided by the number of theoretically observable tryptic peptides.

RNA extraction and gene expression analysis

For B. elliptica, RNAs used to create RNAseq library and to analyze gene expression profiles were isolated from 5 days old mycelial tissue of B. elliptica 9401 grown in vitro on MAE and from lily leaf tissue of cultivar “Asiatic” inoculated with conidia suspension of B. elliptica 9401 harvested at 16, 24 and 40 hours post inoculation (hpi), with three replicate samples per timepoint. RNA extraction was carried out from 100 mg freeze-dried, graded material using the RNA extraction protocol of Maxwell® RSC Plant RNA Kit (Product AS 1500, Promega). For B. squamosa RNAs were extracted from 5 days old mycelial tissue grown on MAE supplemented with onion extract and from onion leaf tissue inoculated with B. squamosa isolate MUCL31421 at 16, 24 and 48 hpi with three replicate samples per timepoint. 100 mg of freeze-dried graded material was incubated with Trizol (Ambion, Life Technology) and treated with chloroform. Technical EtOH (Sigma Aldrich) was added to the aqueous phase and the mixture was used as input for RNeasy Plant Mini Kit (Qiagen) for RNA extraction. RNAs of both Botrytis species were sequenced at Beijing Genomic Institute (BGI), Shenzhen, China. Mapping and quantifying gene transcripts from sequenced RNA-seq reads of both B. elliptica and B. squamosa were performed using Kallisto (Bray et al., 2016). The +1 log10 TPM value was calculated for plotting the expression heatmap.

Results

Host specificity of Botrytis spp.

B. squamosa and B. elliptica are reported to be host-specific pathogens of onion and lily, respectively. To assess this host specificity under laboratory conditions, we tested the virulence of B. squamosa and B. elliptica on their hosts and non-host plants and compared them to the broad host range pathogen B. cinerea (Figure 1). All three species were inoculated on onion, lily and N. benthamiana leaves and lesion development was monitored. The generalist B. cinerea was able to develop expanding lesions exclusively when inoculated on N. benthamiana leaves (Figure 1A) whereas the lesions on lily and onion leaves remained limited to the size of the inoculation droplet. Lesion expansion and disease progression were observed when B. squamosa was inoculated on onion leaves (Figure 1B) and when B. elliptica was inoculated on lily leaves (Figure 1C). In contrast, when B. squamosa and B. elliptica were cross-inoculated on lily and onion, respectively, the primary necrotic lesions did not expand over time and except for a few cases, they remained limited to the size of the inoculation droplet. In addition, inoculation of B. squamosa and B.
elliptica on N. benthamiana leaves did not cause formation of expanding lesions. In conclusion, B. squamosa and B. elliptica caused expanding lesions on their reported natural host but not on the other tested plants.

Novel genome sequence assembly and annotation

High molecular weight genomic DNA of B. squamosa and B. elliptica was isolated from mycelial cultures and was used for whole genome sequencing using Nanopore technology. Assembly of the reads resulted in genomes with a size of 55 and 50 Mb for B. squamosa and B. elliptica, respectively (Table 1). The assembly sizes are in accordance to the available genomes that were sequenced using PacBio technology as described in Valero-Jiménez et al. (2020) (55 and 48 Mb), but the quality of the assemblies was significantly improved. The new assemblies consisted of 19 and 26 contigs for B. squamosa and B. elliptica, respectively (Table 1), as compared to 29 and 137 contigs in the previously published genomes (Valero-Jiménez et al., 2020). Many contigs resembled complete chromosomes. For B. squamosa we assembled 10 out of the 19 contigs with telomeric repeats at both ends, while the remaining 9 contigs contained telomeric sequences at one end. For the genome of B. elliptica, we obtained 8 complete chromosomes containing telomeres at both ends.

A whole genome synteny analysis of B. elliptica and B. squamosa was performed by pairwise alignment of the assemblies. The genomes displayed a high level of synteny, with perfect alignment along 6 of the 16 contigs (Figure 2). There was evidence for 3 structural rearrangements between the contigs of B. elliptica and B. squamosa. For instance contig 1 of B. elliptica aligned to contig 3, 4, and 16 of B. squamosa, suggesting chromosomal rearrangements between the species.

![Figure 1](image_url)

**TABLE 1** Assembly information of genomes of B. elliptica and B. squamosa sequenced in this study.

| Species   | #Contigs | Assembly size (Mb) | Largest contig (bp) | N50 (bp) | #Proteins | #Secreted proteins |
|-----------|----------|-------------------|---------------------|----------|-----------|-------------------|
| B. squamosa | 19       | 54.68             | 6112245             | 3203072  | 11933     | 897               |
| B. elliptica | 26       | 49.77             | 4879067             | 2914681  | 13585     | 925               |

Information regarding proteins and secretome was obtained using gene prediction tools.
Similarly, contig 10 from *B. elliptica* aligned with contig 4 and 16 from *B. squamosa* (Figure 2).

The manually curated annotation of Valero-Jimeñez et al. (2020) was used as input for annotation of the *B. squamosa* and *B. elliptica* genomes, together with RNAseq data from mycelial cultures and infected plant material. The genomes of *B. squamosa* and *B. elliptica* were predicted to contain 11,933 and 13,585 genes respectively, and we specifically examined and manually curated the gene models of the secretome (Table 1).

The predicted secretome of *B. squamosa* consists of 897 genes (7.52%) while the predicted secretome of *B. elliptica* consists of 925 genes (6.81%) (Table 1). Based on the orthogroups described in Valero-Jimeñez et al. (2020), we analyzed the predicted secretome for shared proteins between *B. squamosa* and *B. elliptica* and the more distantly related species *B. cinerea* which functions as the model species for the genus *Botrytis* (Pedro et al., 2019). The vast majority of the predicted secreted proteins could be assigned to 734 orthogroups that were shared between all three species (Supplementary Figure 1). A total of 91 orthogroups is shared only between *B. squamosa* and *B. elliptica* which reflects the recent divergence of these sister species as compared to *B. cinerea*. Only a small proportion of the predicted secretome is shared between *B. cinerea* and either *B. elliptica* or *B. squamosa* (31 and 20 orthogroups, respectively). Only 17, 26 and 55 predicted secreted proteins are unique for *B. squamosa*, *B. elliptica* and *B. cinerea*, respectively (Supplementary Figure 1; Supplementary Data 1).

The genome of *B. elliptica* isolate 9401 contains two mini-chromosomes (comprising both telomeres), of which the largest one is 199 Kbp. These mini-chromosomes have no detectable synteny to the genome of *B. squamosa* (Figure 2).

The largest mini-chromosome (contig 23, 199 kb) contains 18 predicted genes, of which two appeared interesting. The gene BELL_9401_23g00040 encodes a predicted effector protein and has homologs in only *B. tulipae* (BTUL_0255g00010) and *B. hyacinthi* (BHYA_0337g00090) (Valero-Jimeñez et al., 2019). The gene BELL_9401_23g00180 encodes a cysteine protease with potential ubiquitin-deconjugating activity, and it is orthologous to the gene Bcin18g00060 that is located on *B. cinerea* isolate B05.10 mini-chromosome Chr18. The second *B. elliptica* mini-chromosome (contig 25, 175 kb) contains nine predicted genes, of which three have interesting features. BELL_9401_25g00050 has an ortholog in *B. cinerea* B05.10 on mini-chromosome Chr18 (Bcin18g00040, of unknown function), while BELL_9401_25g00060 encodes a putative Forkhead-domain protein kinase with a homolog in *B. cinerea* on mini-chromosome Chr17 (Bcin17g00060). The orthologous *B. cinerea* genes on Chr9 encode a CAZyme GH43 protein and a secreted effector-like protein, whereas one of the genes on Chr17 (Bcin17g00060) encodes a Forkhead-domain protein kinase and is also orthologous to the gene in the *B. elliptica* mini-chromosome contig 25, as mentioned above.
Liquid cultures of *Botrytis* spp. contain cell-death inducing proteins

In order to analyze the cell death-inducing activity of proteins secreted by the different *Botrytis* spp., we collected culture filtrate (CF) samples from *B. squamosa* and *B. elliptica* grown in liquid cultures. The CFs were infiltrated into *N. benthamiana*, onion and lily to assess plant cell death responses. Plant responses were recorded at 3 dpi (Figure 3) and red light imaging (Villanueva et al., 2021) was used to visualize plant cell death responses (Supplementary Figure 2). *B. elliptica* grown in presence of lily leaf extract, caused the most severe response in all tested plant species. Upon infiltration of this sample (CF number 1 in Figure 3, left panel), we observed collapse of the tissue, drying and intense yellowing/browning at the infiltrated area. By contrast, leaf infiltration with compounds collected from *B. elliptica* grown in presence of onion leaf extract (CF number 2 in Figure 3, left panel) caused a milder response and only in lily leaves, where the infiltrated tissue showed a slight translucence. On the other hand, compounds collected from *B. squamosa* grown in presence of lily leaf extract (CF number 3 in Figure 3, left panel) caused a more severe response when compared to the compounds collected from *B. squamosa* grown in presence of onion leaf extract (CF number 4 in Figure 3, left panel). Both *B. squamosa* CF samples did not cause any clear response upon infiltration in lily and *N. benthamiana* leaves, although in the highly sensitive red light imaging system some cell death response could still be observed (Supplementary Figure 2). When CF samples from *B. squamosa* grown in the presence of lily leaf extract were infiltrated in onion leaves, tissue collapse, drying and intense yellowing was observed. Infiltration of the medium used for the fungal liquid culture caused no detectable response in any of the tested plants (Supplementary Figure 2). CF samples contain proteins and secondary metabolites secreted by the fungus, both of which potentially possess cell death-inducing activities. To distinguish the proteinaceous activity from the activity of other compounds, ammonium sulfate precipitation was performed to concentrate the proteins secreted by the fungi in the liquid medium. After dialysis of the protein pellet, the resuspended precipitates were infiltrated in *N. benthamiana*, onion and lily leaves. The ammonium sulfate precipitation of the proteins did not notably affect the cell death inducing activity of *B. elliptica* proteins, since the precipitated proteins derived from *B. elliptica* liquid culture grown in presence of lily leaf extract showed the strongest cell death inducing activity in all tested plants (sample A in Figure 3, right panel; Supplementary Figure 3). The symptoms appeared similar to those observed upon infiltration of the crude CF. In contrast, ammonium sulfate precipitation of the proteins contained in *B. squamosa* CF sample grown in presence of lily leaf extract substantially lost their cell death inducing activity when infiltrated in onion leaves (Figure 3, right panel sample C; Supplementary Figure 3). This result suggests that the cell death-inducing activity of the culture filtrate does not originate from proteins but from phytotoxic secondary metabolites such as botrydial, for which a functional biosynthetic gene cluster is present in *B. squamosa* (Colmenares et al., 2002; Valero-Jiménez et al., 2020). Infiltration of the dialysis buffer caused no visible response in any of the plants tested (Supplementary Figure 3).

Protein composition of the cell death-inducing culture filtrates

The identity of proteins present in the CF samples that showed cell death-inducing activity was analyzed using mass spectrometry. The *B. elliptica* CF sample contained significant hits for a total of 165 proteins identified in the predicted

![Figure 3](image)

**Figure 3**

Leaf responses observed at 3dpi caused by infiltration of crude CF samples (left panel) and AS precipitated compounds (right panel) obtained from liquid cultures of *B. squamosa* and *B. elliptica* on leaves of *N. benthamiana*, onion and lily. 1 = *B. elliptica* grown in GB5_lily; 2 = *B. elliptica* grown in GB5_onion; 3 = *B. squamosa* grown in GB5_lily; 4 = *B. squamosa* grown in GB5_onion. (A) = AS precipitation of *B. elliptica* CF grown in GB5_lily; (B) = AS precipitation of *B. elliptica* CF grown in GB5_onion; (C) = AS precipitation of *B. squamosa* CF grown in GB5_lily; (D) = AS precipitation of *B. squamosa* CF grown in GB5_onion.
To investigate the potential contribution to virulence of the 85 shared proteins detected in the CF samples, we analyzed the transcript levels of their corresponding genes at different timepoints during host infection (Supplementary Data 2). Figure 4 shows a heatmap of the gene expression profile of orthologous genes of *B. squamosa* and *B. elliptica* during infection. When examining the transcript levels in the two *Botrytis* spp., there are three orthogroups with relatively high expression at 40-48 hpi (G17, 49 and 50). For example, G17 which is orthologous to the *B. cinerea* cell death-inducing protein Sp1 (Frias et al., 2011) and G49 which encodes an aspartic protease, both peaked in expression in both species at 40-48 hpi. In contrast, some orthogroups had the highest relative expression at 16 hpi (G24, G29, G46, 58, 62 and 84) in both *B. squamosa* and *B. elliptica*. G24 encodes a putative endoglucanase (GH12) and G58 is predicted as a putative effector. Some orthologs showed different expression patterns between the two fungi, such as G09, G11, G18, G21, G23, G35, G38, G58. For instance, the orthogroup G21, which encodes a polygalacturonase also found in *B. cinerea*, was relatively highly expressed throughout infection in *B. elliptica*, but not in *B. squamosa*. spp.

**Discussion**

Previously the genomes of 16 *Botrytis* spp. with distinct host specificity were analyzed in order to identify unique genes that might encode virulence factors associated to host specificity (Valero-Jiménez et al., 2020). However, no specific elements were found, suggesting that host specificity may be determined by a combination of different (quantitative) traits rather than by presence or absence of unique virulence factors. The aim of this study was to analyse shared features of the secretome in *B. elliptica* and *B. squamosa* that contribute to cell death induction and virulence on their host and non-host plants. This was done by comparing their genomes and by testing the cell death-inducing activity of their secreted proteins.

Genome sequences were available for both species (Valero-Jiménez et al., 2020). However, we considered it useful to further improve the assembly of the *B. squamosa* reference genome and to newly sequence a different *B. elliptica* isolate 9401 (Malvestiti et al., 2021) that was much more virulent than the isolate *B. elliptica* 9612 previously sequenced by Valero-Jiménez et al. (2020). The new genome assemblies of *B. elliptica* 9401 and *B. squamosa* MUCL31421 indeed provided significant improvement over the previous assemblies and enabled us to perform a whole genome alignment and synteny analysis between *B. squamosa* and *B. elliptica*. They display a high degree of synteny, with evidence for only three balanced chromosomal rearrangements. A reconstruction of the ancestral genome of the genus *Botrytis* by Valero-Jiménez et al. (2020) had provided indications for the occurrence of very low numbers of balanced chromosomal rearrangements between the common ancestor of the genus *Botrytis* and the

secretome. More specifically, we detected 70 CAZymes (42% of total number), 15 putative effectors, 8 oxidoreductases and 24 proteases (14.5%). The remaining proteins represented enzymes related to lipid metabolism, different phosphatases, sugar and ion transporters, or proteins without a described domain. In the *B. squamosa* CF sample, 114 proteins with a predicted signal peptide could be identified. Compared to the CF of *B. elliptica*, the *B. squamosa* CF sample contains a considerably higher proportion of CAZymes (69 correspondig to 60% of the total number) but a lower proportion of proteases (7%) and only three putative effectors. In total 85 proteins are present both in *B. squamosa* and *B. elliptica*. More specifically, we detected 70 CAZymes (42% of total number), 15 putative effectors, 8 oxidoreductases and 24 proteases (14.5%). The remaining proteins represented enzymes related to lipid metabolism, different phosphatases, sugar and ion transporters, or proteins without a described domain. In the *B. squamosa* CF sample, 114 proteins with a predicted signal peptide could be identified. Compared to the CF of *B. elliptica*, the *B. squamosa* CF sample contains a considerably higher proportion of CAZymes (69 correspondig to 60% of the total number) but a lower proportion of proteases (7%) and only three putative effectors. In total 85 proteins are present both in *B. squamosa* and *B. elliptica*. When examining the transcript levels in the two *Botrytis* spp., there are three orthogroups with relatively high expression at 40-48 hpi (G17, 49 and 50). For example, G17 which is orthologous to the *B. cinerea* cell death-inducing protein Sp1 (Frias et al., 2011) and G49 which encodes an aspartic protease, both peaked in expression in both species at 40-48 hpi. In contrast, some orthogroups had the highest relative expression at 16 hpi (G24, G29, G46, 58, 62 and 84) in both *B. squamosa* and *B. elliptica*. G24 encodes a putative endoglucanase (GH12) and G58 is predicted as a putative effector. Some orthologs showed different expression patterns between the two fungi, such as G09, G11, G18, G21, G23, G35, G38, G58. For instance, the orthogroup G21, which encodes a polygalacturonase also found in *B. cinerea*, was relatively highly expressed throughout infection in *B. elliptica*, but not in *B. squamosa*. spp.

Discussion

Previously the genomes of 16 *Botrytis* spp. with distinct host specificity were analyzed in order to identify unique genes that might encode virulence factors associated to host specificity (Valero-Jiménez et al., 2020). However, no specific elements were found, suggesting that host specificity may be determined by a combination of different (quantitative) traits rather than by presence or absence of unique virulence factors. The aim of this study was to analyse shared features of the secretome in *B. elliptica* and *B. squamosa* that contribute to cell death induction and virulence on their host and non-host plants. This was done by comparing their genomes and by testing the cell death-inducing activity of their secreted proteins.

Genome sequences were available for both species (Valero-Jiménez et al., 2020). However, we considered it useful to further improve the assembly of the *B. squamosa* reference genome and to newly sequence a different *B. elliptica* isolate 9401 (Malvestiti et al., 2021) that was much more virulent than the isolate *B. elliptica* 9612 previously sequenced by Valero-Jiménez et al. (2020). The new genome assemblies of *B. elliptica* 9401 and *B. squamosa* MUCL31421 indeed provided significant improvement over the previous assemblies and enabled us to perform a whole genome alignment and synteny analysis between *B. squamosa* and *B. elliptica*. They display a high degree of synteny, with evidence for only three balanced chromosomal rearrangements. A reconstruction of the ancestral genome of the genus *Botrytis* by Valero-Jiménez et al. (2020) had provided indications for the occurrence of very low numbers of balanced chromosomal rearrangements between the common ancestor of the genus *Botrytis* and the...
extant species, despite the estimate that they diverged ~5 Million years ago (Shen et al., 2020). The chromosome configurations of *B. squamosa* and *B. elliptica* provide further indications for the remarkable stability of genomes in the genus *Botrytis*, as proposed by Valero-Jiménez et al. (2020). Moreover, the manually curated annotations by Valero-Jiménez et al. (2020) could largely be transferred to the new assemblies of *B. squamosa* and *B. elliptica*, and provided a reliable insight into their proteome and especially secreted protein repertoires. The predicted secretomes of *B. squamosa* and *B. elliptica* were more similar to each other than either of the two species resembled the *B. cinerea* secretome, in concordance with their recent divergence and larger evolutionary distance to *B. cinerea*.

Interestingly, the genome assemblies yielded evidence for the occurrence in both species of mini-chromosomes with lengths of 175-253 kb. One mini-chromosome was identified in *B. squamosa* and two in *B. elliptica*. These mini-chromosomes contain few (predicted) genes, however, remarkably they contain orthologs of genes that are also present on the mini-chromosomes of *B. cinerea*, which is believed to have diverged from *Botrytis* species in clade 2 around 5 Million years ago (Shen et al., 2020). Intriguingly, homologs of several genes on *B. elliptica* and *B. squamosa* mini-chromosomes were detected in distant species (*B. tulipae* and *B. hyacinthi*) but none were detected in the close relatives *B. deweyae* and *B. sinoalli*, suggesting that certain more distant *Botrytis* species may possess homologous mini-chromosomes, while related species might not possess them. One gene that seems to be shared among mini-chromosomes in *B. cinerea*, *B. elliptica* and *B. squamosa* encodes a putative Forkhead-domain protein kinase, while other genes that are shared between two of the three species include genes encoding a putative effector, a glycosyl hydrolase (GH43) and a cysteine peptidase with potential ubiquitin-deconjugating activity. The presence of such genes on mini-chromosomes of three distinct species might explain their retention in the species and the population. The function of these genes in the biology of these pathogens. Based on transcriptome data of *B. elliptica*-infected lily and *B. squamosa*-infected onion, there were 15-20 genes in each of the species that appeared to display significant expression, including a subset of genes that displayed a transient expression peak in early phases of infection. Especially proteins from G58, encoding a putative effector, and G17 which encodes orthologs of the *B. cinerea* cell death-inducing protein SpH1 (Frias et al., 2011) are interesting candidates for analysis.

Both CF samples from *B. squamosa* and *B. elliptica* contained proteins that were uniquely present in that sample. Since the focus of this study is on the shared cell death inducing proteins between the two *Botrytis* spp., these unique proteins were not analysed. However, such unique proteins showing high gene expression during infection can be further investigated to explore their contribution to virulence in the respective host plants. This list still comprises a few dozen genes, and experiments are ongoing to purify CF samples through ion exchange chromatography and to test the activity of fractions for cell death-inducing activity. These efforts should result in reducing the number of candidate genes to a manageable number for functional analysis through CRISPR/Cas9-mediated deletion (Leisen et al., 2020; Leisen et al., 2022).

**Data availability statement**

The Whole Genome Shotgun project of *B. elliptica* 9401 has been deposited at DDBJ/ENA/GenBank under the accession JANCTG000000000. The version described in this paper is version JANCTG010000000. The Whole Genome Shotgun project of *Botrytis squamosa* MUCL31421 has been deposited at DDBJ/ENA/GenBank under the accession RCTC000000000. The version described in this paper is version RCTC02000000.

**Author contributions**

MM and MS contributed equally to this work. MM, MS, XS-K and JK designed the study. MM, MS, HB and SB
performed the lab experiments and analysed the data. XS-K performed the genome assemblies and computational analyses. MM, MS, XS-K and JK wrote the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

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

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.993325/full#supplementary-material

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