The dark septate endophyte *Phialocephala sphaeroides* suppresses conifer pathogen transcripts and promotes root growth of Norway spruce

Kai Wang1,2,3, Zilan Wen2 and Fred O. Asiegbu2,3

1College of Forestry, Fujian Agriculture and Forestry University, Fuzhou 350002, China; 2Department of Forest Sciences, University of Helsinki, PO Box 27, Helsinki FIN-00014, Finland; 3Corresponding authors: K.Wang (wangkai1072@163.com; kai.wang@helsinki.fi) and F.Asiegbu (fred.asiegbu@helsinki.fi)

Received December 28, 2021; accepted July 14, 2022; handling Editor Malin Elfstrand

Plant-associated microbes including dark septate endophytes (DSEs) of forest trees play diverse functional roles in host fitness including growth promotion and increased defence. However, little is known about the impact on the fungal transcriptome and metabolites during tripartite interaction involving plant host, endophyte and pathogen. To understand the transcriptional regulation of endophyte and pathogen during co-infection, Norway spruce (*Picea abies*) seedlings were infected with DSE *Phialocephala sphaeroides*, or conifer root-rot pathogen *Heterobasidion parviporum*, or both. *Phialocephala sphaeroides* showed low but stable transcripts abundance (a decrease of 40%) during interaction with Norway spruce and conifer pathogen. By contrast, *H. parviporum* transcripts were significantly reduced (92%) during co-infection. With RNA sequencing analysis, *P. sphaeroides* experienced a shift from cell growth to anti-stress and antagonistic responses, while it repressed the ability of *H. parviporum* to access carbohydrate nutrients by suppressing its carbohydrate/polysaccharide-degrading enzyme machinery. The pathogen on the other hand secreted cysteine peptidase to restrict free growth of *P. sphaeroides*. The expression of both DSE *P. sphaeroides* and pathogen *H. parviporum* genes encoding plant growth promotion products were equally detected in both dual and tripartite interaction systems. This was further supported by the presence of tryptophan-dependent indolic compound in liquid culture of *P. sphaeroides*. Norway spruce and Arabidopsis seedlings treated with *P. sphaeroides* culture filtrate exhibited auxin-like phenotypes, such as enhanced root hairs, and primary root elongation at low concentration but shortened primary root at high concentration. The results suggested that the presence of the endophyte had strong repressive or suppressive effect on *H. parviporum* transcripts encoding genes involved in nutrient acquisition.

**Keywords:** dark septate endophyte, fungi–fungi–plant interaction, growth promotion, transcriptome.

Introduction

All macro-organisms including plants and forest trees are associated with huge and diverse number of microbiota in natural condition. These microbes play various functional roles either positively or negatively on the growth, development and health of their host plants. Microbe–microbe interaction might also positively affect the health of host plants. For instance, many antagonistic microbes are regarded as potential biocontrol agents to suppress plant diseases (Chan et al. 2007, Macarisin et al. 2010, Liu et al. 2013, Tang et al. 2018, Naik et al. 2019, Eitzen et al. 2021). In natural forest ecosystem, tree-associated fungal endophytes have health beneficial and tree growth promoting effects (Terhonen et al. 2019). Biocontrol microbial agents with plant growth-promotion ability are considered as a safe alternative to chemical control in agriculture and forest environments. Several antagonistic microbes have been applied in practice, such as the application of biocontrol fungus *Phlebiopsis gigantea* (Rotstop®) in coniferous forest to control...
root and butt rot disease caused by *Heterobasidion* species (Pratt et al. 2000, Berglund and Rönnberg 2004, Nicolotti and Gonthier 2005, Oliva et al. 2017).

Antagonistic microbes usually have the capability of competing for nutrients and space, and/or secreting enzymes, antibiotics to suppress plant pathogens (Liu et al. 2013, Naik et al. 2019). For instance, a protein member of glycoside hydrolase family 25 (GH25) of yeast *Moesziomyces bullatus ex Albugo* has been demonstrated for its antagonistic effect against pathogenic oomycete *Albugo laibachii*, contributing to reduced infection in Arabidopsis (Eitzen et al. 2021). A subtilisin-like protease, secreted by biocontrol fungus *Trichoderma harzianum*, displays antifungal activity against multiple phytopathogenic fungi (Yan and Qian 2009, Fan et al. 2014). Moreover, conifer endophyte *Phialocephala scopiformis* is reported to produce anti-insect secondary metabolites (Sumarah and Miller 2009).

A number of plant growth-promoting (PGP) endophytes have been previously described, including bacteria *Bacillus* spp., *Pseudomonas* spp., *Streptomyces* spp., *Burkholderia* spp. (Sun et al. 2009, Kurepin et al. 2015, Afzal et al. 2017, Vurukonda et al. 2018, Wang et al. 2021), filamentous fungi (Alberton et al. 2010, Khan et al. 2012) and some yeasts (Nassar et al. 2005, Nutaratat et al. 2014, Firisinči et al. 2015). The mechanisms of PGP by microbes include nitrogen fixation, alteration of plant hormone network, production of plant hormones, siderophore, phenazine (Sessitsch et al. 2012, Nutaratat et al. 2014, Naik et al. 2019). Plant root elongation, height and mass increase, yield improvement are also common phenotypic effects of PGP. However, many puzzles related to the mechanism of PGP by beneficial microbe remain to be unravelled.

*Phialocephala* species, the dark septate endophytes (DSEs) that belong to the group of ascomycetous fungi, are widely distributed and associated with forest trees (Kowalski and Kehr 1995, Wilson et al. 2004, Schlegel et al. 2016, Walker et al. 2016). PGP fungi and pathogen/insect antagonistic fungi in this genus have previously been reported (Terhonen et al. 2014, Walker et al. 2016, Narisawa 2017, 2018). The DSE *Phialocephala sphaeroides*, used in this study, was originally isolated from Norway spruce root (Terhonen et al. 2014). PGP and antagonistic effect of *Phialocephala* species have previously been demonstrated (Mandyam and Jumpponen 2005, Alberton et al. 2010, Tellenbach et al. 2013, Terhonen et al. 2014, 2016). Although many PGP microbes and antagonistic microbes have been documented, our knowledge on the interaction mechanisms, especially on how they interact on transcriptomic and proteomic levels, is still limited. The physiological and metabolic changes of PGP microbes and antagonistic microbes are largely unknown. Wen et al. (2022) recently reported that Norway spruce seedlings inoculated with DSE *P. sphaeroides* exhibited PGP effect in promoting primary root elongation. In addition, there was no negative effect on seedlings co-infected with both the endophyte and pathogen *Heterobasidion parviporum*, compared with those infected with pathogen alone. In addition, multiple host transcriptional responses in Ps-inoculated seedlings could have contributed to the root growth promotion (Wen et al. 2022). However, the transcriptomic and metabolic insights from the perspective of the interacting fungi (Ps and Hp) were not evaluated. The aim of this study was to identify transcriptomic and metabolic features associated with spruce growth promotion and pathogen inhibition by the endophyte. Norway spruce seedlings inoculated with Ps-alone (PaPs), Hp-alone (PaHp), and co-infected with Ps and Hp (PaPsHp) were selected for total RNA isolation. Thus, transcripts of both spruce (Wen et al. 2022) and fungi were sequenced. The results supplement the earlier published findings on the host transcriptome (Wen et al. 2022). In the present study, we further analysed the deep sequenced RNA sequencing (RNAseq) data. The fungal transcripts or sequences were mapped to the reference genomes of *P. scopiformis* or *H. parviporum*. Transcriptomic evidences of pathogen suppression and PGP effects by the endophyte Ps were revealed. Moreover, we detected the presence of indolic compounds with auxin activity and PGP effect in the liquid culture of the endophyte.

**Materials and methods**

**Seedling inoculation and RNAseq**

Norway spruce seedlings that were inoculated with either *P. sphaeroides* (Ps) and/or the pathogen *H. parviporum* (Hp) in co-cultivation were used for total RNA isolation. The total RNAs include the RNAs from Norway spruce, *P. sphaeroides* and/or *H. parviporum*. The co-cultivation, RNA isolation and RNAseq were conducted as previously described (Wen et al. 2022). Briefly, Norway spruce seeds were surface sterilized and germinated on 1% water agar plate for 2 weeks. The DSE *P. sphaeroides* was cultured on malt extract agar (MEA) plate for 1 month. The cellophane membrane containing the DSE mycelium was moved to petri dish. Two-week-old seedlings were transferred to the medium and laid on the hyphae of the DSE for 1 month, before transferring onto rectangular plates (230 × 82 × 18 mm, Radia Industry Co. Ltd, Japan) with sterile peat-based substrate for another month for adaptation. Agar plugs of 17-day-old *H. parviporum* culture were picked to inoculate seedling roots with/without *P. sphaeroides* inoculation. Thus, Ps seedlings (2.5 months old) were divided into two subgroups: one subgroup was inoculated with *H. parviporum* (PaPsHp), another without Hp (PaPs). Four systems including seedlings with *H. parviporum* inoculated (PaP), seedlings with *P. sphaeroides* inoculated (Ps), seedlings with *H. parviporum* inoculated (PaP), and seedlings with both *P. sphaeroides* and *H. parviporum* inoculated (PaPsHp) were set up with three biological replicates (three to four seedlings per replicate). The whole seedlings (needles, hypocotyl, roots) were collected 1 month post *H. parviporum* inoculation for RNA isolation with cetyl trimethyl ammonium bromide (CTAB) method (Chang et al. 1993). Paired-end (150 bp) transcriptome sequencing
RNAseq mapping to fungi

As to read mapping to fungal genomes, after checking RNAseq raw data quality with FastQC v0.11.8 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and MultiQC v1.9 (Ewels et al. 2016), SortMeRNA V4.2.0 (Kopylova et al. 2012) and trimmomatic (Bolger et al. 2014) were conducted to remove ribosomal RNA, adaptor and low-quality sequences (SLIDINGWINDOW: 5: 20 MINLEN: 50). The trimmed sequences were mapped into fungal genome *P. scopiformis* DAOMC 229536 (LKN1000000000) (Walker et al. 2016) or *H. parviporum* (assembly ASM299478v1, PDUQ0000000000) with alignment tool STAR (Dobin et al. 2013). Aligned sam files were converted into bam files with SAMTools (Li et al. 2009). Mapped transcripts in bam files were counted with HTSeq (Anders et al. 2015).

RNAseq and statistical analysis

Principal component analysis (PCA) and hierarchical clustering of samples were carried out with DESeq2 (Love et al. 2014) using combined count tables from different samples, which were transformed with variance stabilizing transformation (vst) and were normalized for library size and RNA composition effect. DESeq2 (Love et al. 2014) was applied with adjusted P-value 0.05 to produce differentially expressed gene (DEG). DESeq2 normalized values were used for heatmap (scaled by row) and dendrogram with package gplots heatmap.2 (Warnes et al. 2016) in R. For a comparison, DEG was also produced with edgeR (Robinson et al. 2010) with trimmed mean of M-values (TMM) normalization and P-value cutoff 0.05. DEG annotations were manually checked to avoid false computational annotation. Gene ontology (GO) enrichment of DEG (DESeq2 method) was conducted using package clusterProfiler 4.0 (Wu et al. 2021), with P-value cutoff 0.01 and P adjust method Benjamini-Hochberg (BH). Gene-list enrichment tool in KOBAS 3.0 (Bu et al. 2021) was utilized for Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment of DEGs, with Fisher’s exact test and false discovery rate (FDR) (Benjamini–Hochberg adjusted) <0.05. For DEG analysis of pathogen Hp transcripts, default adjusted P-value 0.1 was used as cutoff in DESeq2 analysis.

*Phialocephala scopiformis* mRNAs that mapped the transcripts from PaPs and PaPsHp samples were searched for PGP-related proteins, such as phytohormone biosynthesis enzymes, 1-aminocyclopropane-1-carboxylate (ACC) deaminase, nitrogen fixation, phenazine biosynthesis, non-ribosomal peptide synthase, polyketide synthase, siderophore biosynthesis (Sessitsch et al. 2012, Wang et al. 2021). Blastx was performed to select hits with cutoff e-value <1e-70 and bitscores > 200. Transcriptional abundance of those PGP-related genes from PaPs and PaPsHp samples are listed in Tables 2 and 3.

Candidate of effector genes and annotations of *H. parviporum* were applied as previously described (Zeng et al. 2018, Wen, Zeng, et al. 2019). The expression of predicted *H. parviporum* effector genes and total read counts were used for heatmap production with DESeq2 package. Raw count of transcripts with less than (≤) 5 in more than (≥) 80% samples were removed. Read counts were normalized and transformed with vst method. The details of DEGs were further investigated with the gene annotation, which were extracted from genome annotation of reference genomes.

Statistical analysis of relative transcript abundance and indolic compound production were conducted by t-test in R (version 4.1.2). Data of indolic compound production was plotted as boxplot with package ggplot2 (Wickham 2011) in R.

Fungal culture, indolic compound production and culture filtrate treatment on root

*Phialocephala sphaeroides* was cultured in MEA plate at 20 °C. For indolic compound production assay, *P. sphaeroides* was cultivated in malt extract (ME) or ME + 0.1% tryptophan liquid culture. Culture supernatant was centrifuged at 12,000 rpm for 2 min, and upper layer was chosen to detect indolic compounds with ME or ME + 0.1% tryptophan as blank reference. Salkowski reagent was chosen to quantify indolic compound production by Ps (Glickmann and Dessaux 1995, Wang et al. 2016), with indole-3-acetic acid (IAA) applied as standard. Spruce seeds were sterilized in hydrogen peroxide solution (Merck, USA) for 10 min and then were washed with clean water. Sterile seeds were germinated in 2% water agar plate at 16 h/8 h light/dark at 20 °C. Two-week-old spruce seedlings with similar sizes were transferred into new water agar plates. Filtered culture supernatant was added alongside the root, with culture medium as control. The root regions were covered with autoclaved filter papers. Growth of primary roots was monitored and recorded 4 weeks post treatment. Sterile Arabidopsis seeds germinated in half Murashige and Skoog (MS) agar plates at 16 h/8 h light/dark at 20 °C. Root tips of 1-week-old Arabidopsis seedlings were washed with clean water. Sterile seeds were treated with filtered culture supernatant (∼5 μl per root), with the same volume of 2 μM IAA as positive control and ME + 0.1% tryptophan medium as negative control. Photos of root hair phenotype were taken 24 h post treatment with ZEISS SteREO Discovery.V20 microscope.

Results

Heterobasidion parviporum transcripts decreased dramatically in co-infection

Variable effects of fungi inoculation on the growth of spruce seedlings were observed. The endophyte *P. sphaeroides* (Ps) significantly promoted the growth of primary root of spruce, whereas the pathogen *H. parviporum* (Hp) had the opposite effect. Equally, no negative effect on plant growth was observed in seedlings co-infected with endophyte and pathogen (Figure 1A). In order to unravel the activity of Ps and Hp in

---

with Illumina NovaSeq 6000 platform was conducted by Novogene, UK.
Figure 1. Phenotyping of co-cultivation systems and fungal relative transcription abundance. (A) Norway spruce seedlings were cultured in autoclaved soil with inoculation of pathogen Hp, endophyte Ps and Ps + Hp. Seedlings were kept at growth chamber with 16 h/8 h light/dark at 20 °C. Photos were taken 2 months post inoculation. (B) Relative transcription abundance of Ps from PaPs and PaPsHp samples, as well as of Hp from PaHp and PaPsHp samples. Transcripts were counted as the total transcripts that mapped to the reference genome of Phialocephala scopiformis DAOMC 229536 (LKNI00000000) or Heterobasidion parviporum (assembly ASM299478v1, PDUQ00000000). Pa, Picea abies; Ps, Phialocephala sphaeroides; Hp, Heterobasidion parviporum. *P < 0.05; **P < 0.01.

Table 1. Number and percentage of uniquely mapped reads of total RNAseq reads that aligned to reference fungal genomes of Phialocephala scopiformis or Heterobasidion parviporum from our co-cultivation system, we mapped and counted the total transcripts of Ps and Hp from spruce seedlings inoculated with Ps-alone (PaPs), Hp-alone (PaHp) and Ps + Hp (PaPsHp). Total mapped transcripts of both Ps and Hp decreased in PaPsHp samples, compared with PaPs or PaHp samples (Figure 1B). The decrease indicated that the presence of Ps had significant repressive or suppressive effect on Hp transcripts. The Ps total mapped reads in PaPsHp samples were 60% of those in PaPs samples. However, the Hp total mapped reads in PaPsHp samples were only 8% of those from PaHp samples (Figure 1B). Total mapping counts and mapping rates are listed in Table 1.

Enrichment of GO, KEGG pathway and transcriptomic changes of Ps. sphaeroides in response to co-infection
We compared the Ps transcriptomic patterns from PaPs and PaPsHp samples. Uniquely mapped reads to Ps reference genome...
were utilized for further analysis. PCA of PaPs and PaPsHp samples revealed that the two groups were well separated by principal component 1, which accounted for 67% variance (Figure 2A). Hierarchical clustering and heatmap of samples also confirmed the grouping of PaPs and PaPsHp samples (Figure 2B).

Two DEG analysis methods, edgeR and DESeq2, were performed. In summary, there were 74 upregulated genes and 52 downregulated genes in PaPsHp samples compared with those in PaPs samples by edgeR method. With DESeq2, 166 upregulated genes and 146 downregulated genes were detected. The two analysis methods offered similar results and conclusion with DEGs. We observed transcriptomic evidence for the shift from Ps growth to Ps stress response when Hp was added into the cultivation system. GO terms related to glycolytic process, proton-transporting ATPase activity, gluconeogenesis and serine kinase activity in Ps transcriptome were enriched in PaPsHp samples. On the other hand, GO terms about ribosome, RNA binding and translation were enriched in PaPs samples (Figure 2C). Similar to GO enrichment, KEGG pathway enrichment indicated that Ps cell growth-related pathways were enriched in PaPs sample, whereas anti-stress related pathways were enhanced in PaPsHp sample (Figure 2D). This comparison depicted that the Ps translation process was hindered in co-inoculation with Hp. More specifically, genes encoding for 40S-, 60S-ribosomal proteins, elongation factors, cell division control proteins were downregulated, representing a lower speed of Ps growth. By contrast, multiple genes encoding for superoxide dismutase (SOD), vacuolar ATP synthase (V-ATPase), kinase-like proteins and calcium-dependent kinases were upregulated, suggesting an amplified stress response. Transcripts mapped to two SOD genes, copper/zinc SOD and manganese and iron SOD, were upregulated to 5–6-fold in PaPsHp system. Three homologues of V-ATPase genes from Ps were upregulated to 5–8-fold in the co-cultivation system. More interestingly, homologues of genes encoding for inhibitory enzymes, such as glycoside hydrolases, subtilisin-like serine protease and polyketide synthase, were upregulated significantly in PaPsHp samples (Figure 2E).

Transcriptomic changes of *H. parviporum* in response to co-infection

Hp transcriptomic patterns from PaHp samples were also compared against PaPsHp samples. Sample PaPsHp3 was excluded in the analysis because of extremely low read counts. PCA plot (PC2 with 28% variance) and hierarchical clustering indicated the separation of the two sample groups (Figure 3A and B). With DESeq2 analysis, 25 genes were considered as DEGs, of which 16 were annotated with known functions (Figure 3C). The expression of genes encoding for carbohydrate/polysaccharide-degrading enzymes (cellobiose dehydrogenase, glycogen debranching enzyme and glycoside hydrolase) and cytochrome P450 monoxygenase were significantly decreased. By contrast, phenolic and epoxide degradation enzyme (Aryl-—alcohol dehydrogenase, epoxide hydrolase) genes were upregulated in PaPsHp samples (Figure 3C).

In addition, we monitored the expression of Hp effector candidate genes (*Wen, Zeng, et al. 2019*) in PaHp and PaPsHp samples. Although none of the potential effector gene was detected as DEG with DESeq2 analysis, there was a trend that some clusters of pathogenicity-related genes were expressed at a lower level in PaPsHp samples (Figure 4). Some secreted protein genes with high expression in both PaHp and PaPsHp samples were also identified, which include genes encoding for hydrophobin, cytochrome P450, cerato-platanin and other proteins that might play important roles during the interaction with their host Norway spruce (Figure 4).

**Fungal expression of PGP genes**

To understand growth promotion effect by Ps, we investigated the Ps mapped transcripts from both PaPs and PaPsHp samples. Interestingly, transcripts mapped to auxin and cytokinin biosynthesis, polyketide synthase and molecular chaperone HscA genes were discovered (Table 2). Among them, one indole-3-acetaldehyde dehydrogenase (Ald) gene, one polyketide synthase gene and four molecular chaperone HscA genes were highly expressed, which might reflect PGP traits. Notably, the expression of those Ps genes were not negatively affected in PaPsHp samples, which was consistent with the same level of PGP phenotype in PaPs and PaPsHp samples (Figure 1A). Other microbial genes related to PGP, such as ACC deaminase gene, nitrogen fixation nifH, nifD, nifK genes and phenazine biosynthesis genes were not found in transcripts of PaPs or PaPsHp samples. Interestingly, PGP-related genes were detected, with an overall higher diversity and abundance, in the transcripts aligned to pathogen Hp genome. Transcripts mapped to auxin and cytokinin biosynthesis gene were evenly expressed in both PaHp and PaPsHp samples. Similarly, the expression of genes for polyketides and iron—sulphur proteins were also observed from the pathogen Hp (Table 3).

**Indolic compound production and plant growth promotion by *P. sphaeroides* filtrate**

Plant hormone auxin production was mimicked by detection of indolic compounds in Ps liquid culture. Ps produced indolic compounds at 1.3 μg/ml (IAA equivalent) at 9th day in liquid ME medium. The indolic compounds production significantly increased to 7.2 μg/ml at 9th day in liquid culture supplemented with tryptophan (Figure 5A), indicating the presence of tryptophan-dependent auxin synthesis pathways. The accumulation of indolic compounds increased with time, reaching 6.8 and 17.6 μg/ml at 18th day without or with additional supplementation with tryptophan (Figure 5A). To explore endophyte PGP mechanism at metabolic level, we cultured spruce and
Figure 2. RNAseq profiling of *Phialocephala sphaeroides* in PaPs and PaPs Hp samples. (A) PCA plot of total uniquely mapped reads of endophyte *P. sphaeroides* from PaPs and PaPs Hp samples. (B) Heatmap of hierarchical clustering of uniquely mapped reads from PaPs and PaPs Hp samples. (C) GO enrichment of endophyte *P. sphaeroides* (Ps) from samples PaPsHp and PaPs. List of differential expressed gene (DESeq2 method) was used as input for clusterProfiler 4.0 in R with P-value cutoff 0.01 and P adjust method BH. (D) KEGG pathway enrichment of *P. sphaeroides* upregulated genes (DESeq2 method) in PaPs and PaPsHp samples (adjusted P-value < 0.05, gene count > 3). (E) Heatmap of selected differential expressed genes (DESeq2 method) related to cell growth, stress response and antifungal activity. ∗: It was annotated as Ca-dependent kinases, which is a mistake from computational annotation. Pa, *Picea abies*; Ps, *Phialocephala sphaeroides*; Hp, *Heterobasidion parviporum*.
Figure 3. Differential expressed genes of *Heterobasidion parviporum*. (A) PCA plot of total uniquely mapped reads of pathogen *H. parviporum* from PaHp and PaPsHp samples. (B) Heatmap of hierarchical clustering of uniquely mapped reads from PaHp and PaPsHp samples. Sample PaPsHp3 was not included in PCA and hierarchical clustering because of extremely low mapping rate. (C) Heatmap of annotated differential expressed genes from PaHp and PaPsHp samples. Default adjusted P-value 0.1 was used as cutoff in DESeq2 analysis. Pa, *Picea abies*; Ps, *Phialocephala sphaeroides*; Hp, *Heterobasidion parviporum*.

Table 2. The transcripts of endophyte *Phialocephala sphaeroides* related to plant growth promotion enzymes discovered from PaPs and PaPsHp samples.

| Annotation                        | Ps gene     | PaPs 1 | PaPs 2 | PaPs 3 | PaPsHp 1 | PaPsHp 2 | PaPsHp 3 | %    | e-value | Bit-score |
|-----------------------------------|-------------|--------|--------|--------|----------|----------|----------|------|---------|-----------|
| Hormone                           |             |        |        |        |          |          |          |      |         |           |
| Indole-3-acetaldehyde dehydrogenase | XM_018205505.1 | 236    | 217    | 180    | 206      | 470      | 292      | 64   | 0        | 657       |
| Cytokinin riboside 5′-monophosphate phosphoribohydrolase | XM_018206012.1 | 1      | 1      | 4      | 4        | 12       | 53       | 9e-73 | 224      |           |
| Polyketide synthase               | XM_018218844.1 | 12     | 6      | 11     | 23       | 14       | 37       | 33   | 7e-110   | 387       |
| Iron–sulphur protein HscA         | XM_018205861.1 | 208    | 326    | 185    | 91       | 97       | 74       | 39   | 2e-114   | 358       |
| Molecular chaperone HscA          | XM_018209936.1 | 68     | 63     | 42     | 16       | 20       | 27       | 42   | 3e-130   | 395       |
| Molecular chaperone HscA          | XM_018210372.1 | 390    | 449    | 404    | 135      | 130      | 128      | 39   | 2e-95    | 306       |
| Molecular chaperone HscA          | XM_018214238.1 | 370    | 373    | 181    | 736      | 903      | 253      | 41   | 5e-118   | 365       |

DESeq2 normalized transcript counts of Ps from different samples were shown. E-value <1e-70 and bitscores > 200 were applied in blastx hit selection. Pa, *Picea abies*; Ps, *Phialocephala sphaeroides*; Hp, *Heterobasidion parviporum*; %, amino acid sequence identity.

Arabidopsis seedlings with exposure of the roots to Ps culture supernatant (ME + tryptophan medium, 18 days). Arabidopsis root tips treated with 1/10× filtered Ps culture supernatant displayed root hair enrichment, similar to the phenotype of those with 2 μM IAA treatment, whereas those treated with ME + tryptophan medium exhibited normal root hair phenotype (Figure 5B). With the application of 1/10× filtered Ps culture supernatant on spruce seedlings, the significant elongation of...
Table 3. The transcripts of pathogen *Heterobasidion parviporum* related to plant growth promotion enzymes discovered from PaHp and PaPsHp samples.

| Annotation                  | Hp gene     | PaHp 1 | PaHp 2 | PaHp 3 | PaPsHp 1 | PaPsHp 2 | %   | e-value | Bit-score |
|-----------------------------|-------------|--------|--------|--------|----------|----------|------|---------|-----------|
| **Hormone**                 |             |        |        |        |          |          |      |         |           |
| Amine oxidase               | scaffold32.110 | 46     | 36     | 57     | 78       | 54       | 40   | 4e-160  | 478       |
| Indole-3-acetamide hydrolase| scaffold1.668 | 36     | 47     | 31     | 4        | 57       | 36   | 3e-79   | 256       |
| Indole-3-acetamide hydrolase| scaffold16.14 | 31     | 42     | 19     | 48       | 51       | 40   | 2e-96   | 298       |
| IAAld dehydrogenase         | scaffold2.297 | 80     | 132    | 93     | 69       | 137      | 34   | 8e-86   | 281       |
| IAAld dehydrogenase         | scaffold20.70 | 35     | 20     | 33     | 26       | 31       | 36   | 3e-95   | 297       |
| IAAld dehydrogenase         | scaffold3.523 | 2,166  | 2,945  | 2,240  | 2,295    | 2,478    | 60   | 0       | 623       |
| IAAld dehydrogenase         | scaffold2.626 | 80     | 92     | 80     | 117      | 66       | 53   | 0       | 529       |
| IPyA decarboxylase          | scaffold11.216 | 62     | 68     | 50     | 61       | 83       | 51   | 0       | 562       |
| Tryptophan aminotransferase | scaffold49.11 | 55     | 41     | 43     | 39       | 23       | 34   | 3e-85   | 272       |
| Tryptophan decarboxylase    | scaffold3.307 | 24     | 15     | 28     | 17       | 17       | 35   | 1e-94   | 295       |
| YUCCA flavin monoxygenase   | scaffold16.22 | 33     | 12     | 29     | 9        | 26       | 33   | 5e-98   | 311       |
| YUCCA flavin monoxygenase   | scaffold2.1024 | 10     | 6      | 10     | 4        | 9        | 40   | 5e-155  | 458       |
| YUCCA flavin monoxygenase   | scaffold27.43 | 206    | 183    | 213    | 234      | 174      | 43   | 2e-168  | 493       |
| YUCCA flavin monoxygenase   | scaffold8.64  | 4      | 11     | 2      | 4        | 6        | 34   | 1e-94   | 304       |
| Cytokinin dehydrogenase     | scaffold1.306 | 41     | 27     | 62     | 35       | 54       | 38   | 5e-109  | 336       |
| Cytokinin dehydrogenase     | scaffold12.90 | 37     | 27     | 18     | 35       | 71       | 51   | 2e-175  | 528       |
| Cytokinin dehydrogenase     | scaffold7.26  | 42     | 11     | 48     | 74       | 9        | 50   | 2e-159  | 466       |
| **Polyketide**              |             |        |        |        |          |          |      |         |           |
| Polyketide synthase         | scaffold12.21 | 27     | 34     | 30     | 52       | 29       | 35   | 4e-152  | 517       |
| Polyketide synthase         | scaffold47.18 | 63     | 55     | 65     | 48       | 63       | 57   | 4e-90   | 267       |
| **Iron–sulphur protein**    |             |        |        |        |          |          |      |         |           |
| Molecular chaperone HscA    | scaffold145.1 | 22     | 32     | 17     | 35       | 31       | 37   | 5e-102  | 322       |
| Molecular chaperone HscA    | scaffold19.1  | 80     | 117    | 62     | 135      | 71       | 37   | 5e-104  | 327       |
| Molecular chaperone HscA    | scaffold2.914 | 135    | 180    | 137    | 139      | 151      | 38   | 1e-102  | 323       |
| Molecular chaperone HscA    | scaffold3.76  | 237    | 151    | 235    | 187      | 117      | 38   | 3e-112  | 350       |
| Molecular chaperone HscA    | scaffold58.28 | 109    | 367    | 119    | 108      | 123      | 42   | 6e-127  | 388       |
| Cysteine desulphurase IscS  | scaffold4.267 | 61     | 134    | 56     | 48       | 66       | 56   | 7e-169  | 481       |

DESeq2 normalized transcript counts of Hp from different samples were shown. *E*-value <1e-70 and bitscores > 200 were applied in blastx hit selection. Pa, *Picea abies*; Ps, *Phialocephala sphaeroides*; Hp, *Heterobasidion parviporum*; IAAld, indole-3-acetaldehyde; IPyA, indole-3-pyruvate; %, amino acid sequence identity.
Dark septate endophyte suppresses conifer pathogen

Figure 4. Heatmap of the expression of secreted protein genes of *Heterobasidion parviporum*. The expression of predicted *Hp* effector genes and total read counts were taken into heatmap production with DESeq2 package. Raw count of transcripts with less than (≤) 5 in more than (≥) 80% samples were removed. Read counts were normalized and transformed with variance stabilizing transformation (vst) method. Genes with lower expression in PaPs*Hp* sample were indicated in red rectangle, and genes with high expression in both Pa*Hp* and PaPs*Hp* samples were indicated in blue rectangle. Pa, *Picea abies*; Ps, *Phialocephala sphaeroides*; *Hp*, *Heterobasidion parviporum*.

primary root was observed on water agar plate at 4 weeks post treatment (Figure 5C).

**Discussion**

There was large variation of *Hp* survival rates (8, 17 and 1%) in PaPs*Hp* samples. The possible reason could be that interaction system involving three partners (host–endophyte–pathogen) are much more complicated than host–endophyte or host–pathogen dual-interaction systems. The significant reduction of *Hp* transcripts in co-infection suggests that metabolic activity of *Hp* was largely restricted or inhibited by the presence of Ps. Many of the affected *Hp* transcripts were found to be those encoding carbohydrate-degrading enzymes. Ps probably repressed *Hp* ability to access carbohydrate nutrients by suppressing its carbohydrate/polysaccharide-degrading enzymes machinery as well as its ability to tolerate toxic environment through negative impact on its cytochrome P450. Conversely, the pathogen secreted cysteine peptidase, most probably as an attempt to restrict the free growth of endophyte, which might contribute to 40% reduction in Ps transcript. Interestingly, increased expression of genes encoding phenolic oxidizing enzymes by pathogen might represent a response to enhanced host defences and/or allelopathic phenolic acid analogues secreted by the endophyte. Equally, some downregulated genes in Ps were related to cell division, elongation factor and ribosomal protein synthesis. This lends credence to the idea that growth of Ps was partly negatively affected. Ribosomal proteins are essential for assembly and function of ribosome, which is required for protein synthesis and cell growth (Warner 1999, Wilson and Nierhaus 2005). All the detected homologues of ribosomal protein genes in DEGs were downregulated, indicating deceleration of protein synthesis and cell growth rate. Interestingly, despite of the negative effect on growth of Ps in PaPs*Hp* system at transcriptomic level, Ps still conferred growth fitness benefits on its host.

Several anti-stress genes were upregulated in Ps transcriptome during co-infection with *Hp*. SODs are key enzymes in
reactive oxygen species (ROS) scavenging systems. Fungal SOD plays a role in detoxification and stress tolerance to induced ROS (González-Guerrero et al. 2010, Xie et al. 2010). Thus, upregulation of two homologues of SOD genes in Ps might function as ROS detoxification in co-infection. V-ATPase functions in the acidification of intracellular compartments in both prokaryotic and eukaryotic cells. Previous studies demonstrated a protection role of V-ATPase when yeast cells are under endogenous oxidative stress and biotic stress (Milgrom et al. 2007, Zhang et al. 2008). Many kinases and kinase-like proteins function in defence/stress signalling in plants and microbes (Lee et al. 2016, Yip Delormel and Boudsocq 2019). The upregulation of stress responsive genes further highlights the ability of Ps to deal with the hostile environment.

The changes of Ps transcriptome also provided evidence of suppressive effect of Hp by Ps. GH25 is an antagonistic effector from phyllosphere yeast to inhibit pathogenic oomycete and reduce infection (Eitzen et al. 2021). Uregulation of glycoside hydrolase genes of Ps in PaPsHp system indicated possible antagonistic mechanism. Subtilisin-like serine protease from biocontrol fungi has been demonstrated to possess antifungal activity (Yan and Qian 2009, Fan et al. 2014). Ps might secrete subtilisin-like serine protease as a strategy to suppress the growth of other competing fungi. Although no obvious inhibition zone of Ps against the growth of Hp was observed on MEA medium (Wen et al. 2022), it is most probable that Ps had a repressive or suppressive impact on Hp metabolism and pathogenicity transcripts in PaPsHp system. This also underlies the complex nature of interaction outcome in artificial media versus in natural forest ecosystem.

Downregulation of some carbohydrate usage enzymes of Hp in response to co-infection indicated the reduced capability to use carbohydrates as energy source. Genes of cytochrome P450 enzymes, which are vital for metabolism and adaptation (Chen et al. 2014) and for potential pathogenicity (Mbgeheuru et al. 2017, Fessner et al. 2021), were downregulated in co-inoculation system. This evidence, together with the lower expression of some candidate effector genes, suggested that the normal growth and metabolism of Hp was disrupted by Ps during co-infection. Surprisingly, Hp genes encoding phenolic oxidizing enzymes were upregulated, indicating activation of an alternative pathway to source for nutrients. It is also likely that Hp might have secreted some inhibitory compounds to counter the effects of Ps. These defences included upregulation of cysteine peptidase of Hp to restrict the growth of other fungi, and lower expression of oxalate decarboxylase effector gene that could lead to accumulation of oxalic acid for antifungal activity (Mäkelä et al. 2010). In addition, we detected highly expressed candidate effector genes that are necessary for Hp growth, including hydrophobin and cytochrome P450 genes, and that for Hp toxicity such as cerato-platanin gene (Chen et al. 2015, Luti et al. 2020) in both groups of samples. The expression of effector HpSSP35.8 gene, which induces cell death and tree defence responses (Wen, Raffaello, et al. 2019), was kept at comparable levels in both groups of samples. These indicated that Hp growth was largely restricted by Ps in this system. Moreover, host priming by Ps might also contribute to Hp inhibition. Host genes related to jasmonic acid (JA) biosynthesis, plant hormone transduction, mitogen-activated protein kinase (MAPK) signalling and Ca2+ signalling were upregulated in PaPs samples (Wen et al. 2022). It suggests immunity responses of Norway spruce primed by Ps. As a conclusion, inhibition to Hp in co-infection system was probably due to a combination of direct antagonistic effect by Ps and indirect effect from the host responses primed by Ps.

Transcripts related to PGP were documented in Ps and Hp transcriptomes. Several transcripts encoding for hormone IAA synthesis, cytokinin synthesis, secondary metabolite polyketide synthesis and iron–sulphur protein maturation (Sessitsch et al. 2012, Wang et al. 2021) were detected in all fungi inoculated samples. Indole-3-acetaldehyde (Ald) functions in microbial IAA synthesis to convert indole-3-acetaldehyde (IAAld) to IAA (McClerklin et al. 2018). Ps Ald genes were highly expressed, indicating the active occurrence of IAA biosynthesis by Ps. Moreover, the presence of necrotrophic pathogen Hp did not affect the expression of Ps Ald gene. This transcriptomic pattern was consistent with the phenotype that PGP effect in PaPs and PaPsHp samples were at the same level. One polyketide synthase and four molecular chaperone HscA (S–Fe protein maturation) genes also exhibited similar expression level. The PGP transcripts were also detected from Hp, which might account for the minor growth promotion observed in PaHp samples compared with uninoculated control samples.

PGP was also recorded with treatment of diluted Ps culture filtrate on spruce seedlings in plates. However, with higher concentration of filtrate, the length of primary root growth was decreased, which is similar to the plant phenotype noted with high auxin concentration. Taken together, we concluded that the Ps secreted compounds had similar active auxin role for Norway spruce. Microbial auxin production plays multiple functional roles in plant growth and development (Vorholt 2012, Sukumar et al. 2013, Chanclud and Morel 2016, Lawren et al. 2016). We revealed that Ps indolic compounds production increased with addition of tryptophan, which represented a tryptophan-dependent IAA synthesis. Indolic compounds in Ps culture filtrate had active auxin-like roles, as seen by enhanced root hairs and changes of primary root. However, Ps culture filtrate contains multiple secreted compounds. We could not exclude the possibility of impacts of other metabolites, such as polyketides.

We characterized the mechanisms of PGP and antagonism of Ps at transcriptomic and metabolic level. Other mechanisms, such as competition for space and nutrients, activation of plant defence might also exist simultaneously for pathogen inhibition effect. In addition, pathogen inhibition could also be mediated by host immune responses primed by endophyte Ps. This is
the first exploratory study providing insight on the regulatory suppressive effect at transcriptomic level of this endophyte on the pathogen in a tripartite interaction. Future studies may wish to address the functional relevance of the identified genes in the beneficial interaction between the endophyte and its conifer host.

Acknowledgments

The authors acknowledge the support from Ministry of Agriculture, Finland (grant no. 4400T-2002). Z.W. was supported by Chinese Scholarship Council (CSC) and Alfred Kordelin Foundation (grant no. 200382). Computing platform in Finnish IT Center for Science (CSC) is gratefully acknowledged.

Conflict of interest

None declared.

Data and materials availability

RNAseq raw reads are available in NCBI database (https://www.ncbi.nlm.nih.gov/, Bioproject PRJNA734471) under accession number SRX11066041-SRX11066042, SRX11066044-SRX11066050.

Authors’ contributions

F.O.A., K.W. and Z.W. conceived the study. K.W. and Z.W. performed the experiments. K.W. analysed the data and wrote the manuscript. All authors edited and proofread the manuscript.

References

Afzal I, Igkar I, Shinwari ZK, Yasmin A (2017) Plant growth-promoting potential of endophytic bacteria isolated from roots of wild Dodonaea viscosa L. Plant Growth Regul 81:399–408.

Alberton O, Kuyper TW, Summerbell RC (2010) Dark septate endophytic fungi increase growth of Scots pine seedlings under elevated CO2 through enhanced nitrogen use efficiency. Plant Soil 328:459–470.

Anders S, Pyl PT, Huber W (2015) HTSeq—a Python framework to work with high-throughput sequencing data. Bioinformatics 31:166–169.

Berglund M, Rönning J (2004) Effectiveness of treatment of Norway spruce stumps with Phlebiopsis gigantea at different rates of coverage for the control of Heterobasidion. For Pathol 34:233–243.

Bolger AM, Lohse M, Usadel B (2014) Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30:2114–2120.

Bu D, Luo H, Huo P et al. (2021) KOBASE-i: intelligent prioritization and exploratory visualization of biological functions for gene enrichment analysis. Nucleic Acids Res 49:W317–W325.

Chan Z, Qin G, Xu X, Li B, Tian S (2007) Proteome approach to characterize proteins induced by antagonist yeast and salicylic acid in peach fruit. J Proteome Res 6:1677–1688.

Chanclud E, Morel JB (2016) Plant hormones: a fungal point of view. Mol Plant Pathol 17:1289–1297.

Chang S, Puryear J, Cairney J (1993) A simple and efficient method for isolating RNA from pine trees. Plant Mol Biol Rep 11:113–116.

Chen H, Quintana J, Kovalchuk A, Ubhayasekera W, Asiegbu FO (2015) A cerato-platanin-like protein HaCPL2 from Heterobasidion annosum sensu stricto induces cell death in Nicotiana tabacum and Pinus sylvestris. Fungal Genet Biol 84:41–51.

Chen W, Lee M-K, Jefcoate C, Kim S-C, Chen F, Yu J-H (2014) Fungal cytochrome P450 monoxygenases: their distribution, structure, functions, family expansion, and evolutionary origin. Genome Biol Evol 6:1620–1634.

Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M, Gingeras TR (2013) STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29:15–21.

Eltzen K, Sengupta P, Kroll S, Kemen E, Doehlemann G (2021) A fungal member of the Arabidopsis thaliana phyllosphere antagonizes Albugo laibachii via a GH25 lysozyme. Elife 10:e65306.

Ewels P, Magnusson M, Lundin S, Käller M (2016) MultiQC: summarize analysis results for multiple tools and samples in a single report. Bioinformatics 32:3047–3048.

Fan H, Liu Z, Zhang R, Wang N, Dou K, Mijiti G, Diao G, Wang Z (2014) Functional analysis of a subtilisin-like serine protease gene from biocontrol fungus Trichoderma harzianum. J Microbiol 52:129–138.

Fessner ND, Nelson DR, Glieder A (2021) Evolution and enrichment of CYP5035 in Polyoparas: functionality of an understudied P450 family. Appl Microbiol Biotechnol 105:6779–6792.

Firrinicelli A, Otlilar R, Salamov A, Schmutz J, Khan Z, Redman RS, Fleck ND, Lindquist E, Grigoriev IV, Doty SL (2015) Genome sequence of the plant growth promoting endophytic yeast Rhodotorula graminis WP1. Front Microbiol 6:978.

Glickmann E, Dessaux Y (1995) A critical examination of the specificity of the Salkowski reagent for indolic compounds produced by phytobacterial bacteria. Appl Environ Microbiol 61:793–796.

González-Guerrero M, Öger E, Benabdelhak K, Azcón-Aguilar C, Lanfranco L, Ferrol N (2010) Characterization of a CuZn superoxide dismutase gene in the arbuscular mycorrhizal fungus Glomus intraradices. Curr Genet 56:265–274.

Khan AL, Hamayun M, Kang S-M, Kim Y-H, Jung H-Y, Lee J-H, Lee I-J (2012) Endophytic fungal association via gibberellins and indole acetic acid can improve plant growth under abiotic stress: an example of Paecilomyces fumosoroseus LHL10. BMC Microbiol 12:1–14.

Kopylova E, Noé L, Touzet H (2012) SortMeRNA: fast and accurate filtering of ribosomal RNAs in metatranscriptomic data. Bioinformatics 28:3211–3217.

Kowalski T, Kehr RD (1995) Two new species of Phialocephala occurring on Picea and Alnus. Can J Plant Sci 73:26–32.

Kurepin LV, Park JM, Lazarovits G, Bernardas MA (2015) Burkholderia phytofirmans-induced shoot and root growth promotion is associated with endogenous changes in plant growth hormone levels. Plant Growth Regul 75:199–207.

Lareen A, Burton F, Schäfer P (2016) Plant root-microbe communication in shaping root microbiomes. Plant Mol Biol 90:575–587.

Lee K-T, So Y-S, Yang D-H, Jung K-W, Choi J, Lee D-G, Kwon H, Jang J, Wang LL, Cha S (2016) Systematic functional analysis of kinases in the fungal pathogen Cryptococcus neoformans. Nat Commun 7:1–16.

Li H, Handsaker B, Wyssoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R (2009) The sequence alignment/map format and SAMtools. Bioinformatics 25:2078–2079.

Liu J, Sui Y, Wisniewski M, Droby S, Liu Y (2013) Utilization of antagonistic yeasts to manage postharvest fungal diseases of fruit. Int J Food Microbiol 167:153–160.

Love MI, Huber W, Anders S (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol 15:1–21.
Luti S, Sella L, Quarantin A, Pazzagli L, Baccelli I (2020) Twenty years of research on cerato-platanin family proteins: clues, conclusions, and unsolved issues. Fungal Biol Rev 34:13–24.

Macarini D, Droby S, Bauchan G, Wisniewski M (2010) Superoxide anion and hydrogen peroxide in the yeast antioxidant–fruit interaction: a new role for reactive oxygen species in postharvest biocontrol? Postharvest Biol Technol 58:194–202.

Mäkelä MR, Hildén K, Lundell TK (2010) Oxalate decarboxylase: biotechnological update and prevalence of the enzyme in filamentous fungi. Appl Microbiol Biotechnol 87:801–814.

Mandayam K, Jumpenson A (2005) Seeking the elusive function of the root-colonising dark septate endophytic fungi. Stud Mycol 53:173–189.

McClerklin SA, Lee SG, Harper CP, Nwumeh R, Jez JM, Kunkel BN (2018) Indole-3-acetaldehyde dehydrogenase-dependent auxin synthesis contributes to virulence of *Pseudomonas syringae* strain DC3000. PLoS Pathog 14:e1006811.

Mgbaburuikke AC, Kovalchuk A, Ubbayasekera W, Nelson DR, Yadav JS (2017) CYPome of the conifer pathogen *Heterobasidion irregulare*: inventory, phylogeny, and transcriptional analysis of the response to biocontrol. Fungal Biol 121:158–171.

Milgrom E, Diab H, Middleton F, Kane PM (2007) Loss of vacuolar proton-translocating ATPase activity in yeast results in chronic oxidative stress. J Biol Chem 282:7125–7136.

Nalik K, Mishra S, Srichandhan H, Singh PK, Sarangi PK (2019) Plant growth promoting microbes: potential link to sustainable agriculture. J Biol Chem 293:1504–1514.

Narisawa K (2017) The dark septate endophytic fungus *Phialocephala fortinii* is a potential decomposer of soil organic compounds and a promoter of *Asparagus officinalis* growth. Fungal Ecol 28:1–10.

Narisawa K (2018) The inhibitory role of dark septate endophytic fungus *Phialocephala fortinii* against Fusarium disease on the *Asparagus officinalis* growth in organic source conditions. Biol Control 121:159–167.

Nassar AH, El-Tarabily KA, Sivasithamparam K (2005) Promotion of plant growth by an auxin-producing isolate of the yeast *Williopsis saturnus* endophytic in maize (*Zea mays* L.) roots. Biol Fertility Soils 42:97–108.

Nicolotti AW, Robinson JNA, Pratt JR, Nutaratat N, Zehnter MH, Piacente E (2017) Quantitative interections between the biocontrol fungus *Phlebiopsis gigantea* and the forest pathogen *Heterobasidion annosum* and the fungal community inhabiting Norway spruce stumps. For Ecol Manage 402:253–264.

Patt J, Niemi M, Sierota Z (2000) Comparison of three products based on *Phlebiopsis gigantea* for the control of *Heterobasidion annosum* in Europe. Biocontrol Sci Technol 10:467–477.

Robinson MD, McCarthy DJ, Smyth GK (2010) edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26:139–140.

Schlegel M, Münsterkötter M, Güldener U, Bruggmann R, Duö A, Hainaut M, Henrisatt B, Sieber CM, Hofmeister D, Grünig CR (2016) Globally distributed root endophyte *Phialocephala subalpina* links pathogenic and saprophytic lifestyles. BMC Genomics 17:1–22.

Sessitsch A, Hardoim P, Döring J, Weilharter A, Krause A, Woyke T, Mitter B, Hauberg-Lotte L, Friedrich F, Rahalkar M (2012) Functional characteristics of an endophyte community colonizing rice roots as revealed by metagenomic analysis. Mol Plant Microbe Interact 25:28–36.

Sukumar P, Legue V, Vaysierres A, Martin F, Tuskan GA, Kalluri UC (2013) Involvement of auxin pathways in modulating root architecture during beneficial plant–microorganism interactions. Plant Cell Environ 36:909–919.

Sumarah MW, Miller JD (2009) Anti-insect secondary metabolites from fungal endophytes of conifer trees. Nat Prod Commun 4:1934578X0900401112.

Sun Y, Cheng Z, Glick BR (2009) The presence of a 1-aminoacyclopropane-1-carboxylate (ACC) deaminase deletion mutation alters the physiology of the endophytic plant growth-promoting bacterium *Burkholderia phytofirmans* PSN. FEMS Microbiol Lett 296:131–136.

Tang JY, Bullen NP, Ahmad S, Whitney JC (2018) Diverse NADase effector families mediate interbacterial antagonism via the type VI secretion system. J Biol Chem 293:1504–1514.

Tellenbach C, Sumarah MW, Grüning CR, Miller JD (2013) Inhibition of *Phytophthora* species by secondary metabolites produced by the dark septate endophyte *Phialocephala europaea*. Fungal Ecol 6:12–18.

Terhonen E, Blumenstein K, Kovalchuk A, Asiegbu FO (2019) Forest tree microbicides and associated fungal endophytes: functional roles and impact on forest health. Forests 10:42.

Terhonen E, Kerio S, Sun H, Asiegbu FO (2014) Endophytic fungi of Norway spruce roots in boreal pristine mire, drained peatland and mineral soil and their inhibitory effect on *Heterobasidion parviporum* in vitro. Fungal Ecol 9:17–26.

Terhonen E, Sipari N, Asiegbu FO (2016) Inhibition of phytopathogens by fungal root endophytes of Norway spruce. Bio Control 99:53–63.

Vorholt JA (2012) Microbial life in the phyllosphere. Nat Rev Microbiol 10:828–840.

Vurukonda SSKP, Giovannardi D, Stefani E (2018) Plant growth promoting and biocontrol activity of *Streptomycetes* spp. as endophytes. Int J Mol Sci 19:952.

Walker AK, Frasz SL, Seifert KA, Miller JD, Mondo SJ, LaButti K, Lipzen A, Dockter RB, Kennedy MC, Grigoriev IV (2016) Full genome of *Phialocephala scopiformis* DAOMC 229536, a fungal endophyte of spruce producing the potent anti-insectan compound rugulosin. Genome Announc 4:e01768–e01715.

Wang K, Sipilä TP, Overmyer K (2016) The isolation and characterization of resident yeasts from the phylloplane of *Arabidopsis thaliana*. Sci Rep 6:39403.

Wang K, Wu Y, Ye M, Yang Y, Asiegbu FO, Overmyer K, Liu S, Cui F (2021) Comparative genomics reveals potential mechanisms of plant beneficial effects of a novel bamboo-endophytic bacterial isolate *Paraburkholderia sacchari* Suichang626. Front Microbiol 12:1597.

Warner JR (1999) The economics of ribosome biosynthesis in yeast. Trends Biochem Sci 24:437–440.

Warness MGR, Bolker B, Bonebakker L, Gentleman R, Huber W (2016) Package ‘ggplot2’: various R programming tools for plotting data. https://cran.r-project.org/web/packages/ggplot2/index.html

Wen Z, Raffaello T, Zeng Z, Pavicic M, Asiegbu FO (2019) Chlorophyll fluorescence imaging for monitoring effects of *Heterobasidion parviporum* small secreted protein induced cell death and in planta defense gene expression. Fungal Genet Biol 126:37–49.

Wen Z, Terhonen E, Asiegbu FO (2022) The dark septate endophyte *Phialocephala sparaehoides* confers growth fitness benefits and mitigates pathogenic effects of *Heterobasidion* on Norway spruce. Tree Physiol 42:891–906.

Wen Z, Zeng Z, Ren F, Asiegbu FO (2019) The conifer root and stem rot pathogen (*Heterobasidion parviporum*): effectome analysis and roles in interspecific fungal interactions. Microorganisms 7:658.

Wickham H (2011) ggplot2. Wiley Interdiscip Rev Comput Stat 3:180–185.
Wilson B, Addy H, Tsuneda A, Hambleton S, Currah R (2004) *Phialocephala sphaeroides* sp. nov., a new species among the dark septate endophytes from a boreal wetland in Canada. Can J Plant Sci 82:607–617.

Wilson DN, Nierhaus KH (2005) Ribosomal proteins in the spotlight. Crit Rev Biochem Mol Biol 40:243–267.

Wu T, Hu E, Xu S, Chen M, Guo P, Dai Z, Feng T, Zhou L, Tang W, Zhan L (2021) clusterProfiler 4.0: a universal enrichment tool for interpreting omics data. Innovation 2:100141.

Xie X-Q, Wang J, Huang B-F, Ying S-H, Feng M-G (2010) A new manganese superoxide dismutase identified from *Beauveria bassiana* enhances virulence and stress tolerance when overexpressed in the fungal pathogen. Appl Microbiol Biotechnol 86:1543–1553.

Yan L, Qian Y (2009) Cloning and heterologous expression of SS10, a subtilisin-like protease displaying antifungal activity from *Trichoderma harzianum*. FEMS Microbiol Lett 290:54–61.

Yip Delormel T, Boudsocq M (2019) Properties and functions of calcium-dependent protein kinases and their relatives in *Arabidopsis thaliana*. New Phytol 224:585–604.

Zeng Z, Sun H, Vainio EJ, Raffaello T, Kovalchuk A, Morin E, Duplessis S, Asiegbu FO (2018) Intraspecific comparative genomics of isolates of the Norway spruce pathogen (*Heterobasidion parviporum*) and identification of its potential virulence factors. BMC Genomics 19:220.

Zhang X, Liu S, Takano T (2008) Overexpression of a mitochondrial ATP synthase small subunit gene (*AtMtATP6*) confers tolerance to several abiotic stresses in *Saccharomyces cerevisiae* and *Arabidopsis thaliana*. Biotechnol Lett 30:1289–1294.