Evaluation of fungal endophytes to biological control of Dothistroma needle blight on Pinus nigra subsp. pallasiana (Crimean pine)

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Dothistroma needle blight (DNB), caused by Dothistroma septosporum and Dothistroma pini, is the most important forest disease of pine in many countries. This disease has recently emerged in Ukraine as a major threat to mostly Pinus nigra subsp. pallasiana and less to Scots pine. There is increasing evidence that some fungal and bacterial isolates can reduce the growth and pathogenicity of fungal plant pathogens.

In this research, infected needles were collected from 30-year-old Crimean pine (P. nigra subsp. pallasiana) in four locations in Southern Ukraine. In total, 244 of endophytic fungi were recovered from needles of Crimean pine during summer sampling of the host’s microbiome in Ukraine in 2012-2014.

Dothistroma spp. were detected using fungal isolation and species-specific priming PCR techniques.

Among all endophytes, eight fungal species were selected based on the commonness of their occurrence in the foliage of the host and their antagonistic activity. All selected species were tested for their antifungal activity against Dothistroma needle blight. Good antifungal activity against Dothistroma pini was achieved with the Trichoderma sp. and Gliocladium rosea, indicating their good potential possibility in preventing the Dothistroma needle blight on young pines. Moreover, the significant reduction in numbers of conidia and spore germination was found on needles treated with Trichoderma sp. and Gliocladium rosea, compared to conidia numbers following treatment with other fungi. Thus, the use of an effective biological control agent against Dothistroma could be of value in forest nurseries, where it is essential to reduce losses to D. pini infection prior to transferring pines to field sites for planting out.

Key words: DNB; Dothistroma septosporum; Dothistroma pini; pathogens; biological control; forest health.

Introduction. Dothistroma needle blight (DNB) is one of the most serious needle diseases of pine plantations and is responsible for many economic losses in landscape and forest settings through negative impacts on yields, quality of stands and visual amenity. The main symptoms of the disease are premature needle fall, reduce in photosynthetic capacity, followed by yield losses and, in some cases, tree mortality (Bradshaw 2004, Woods et al. 2005, Millberg et al., 2016). Two closely related ascomycetous fungi are...
known to cause DNB: *Dothistroma septosporum* (Dorogin) M. Morelet and *D. pini* Hulbary (Barnes et al., 2004). *Dothistroma septosporum* and *D. pini* do not show consistent differences in the length of their conidia, so it is necessary the use of specific molecular markers for their discrimination and identification of the two pathogens (Barnes et al. 2004, Ioos et al., 2010).

*Dothistroma septosporum* has a worldwide distribution and found in many parts of the world where pines are grown (Barnes et al., 2008, Bulman et al., 2013). *D. pini* was found from some countries in Europe including Ukraine and from the USA (Drenkhan et al., 2016).

*Dothistroma septosporum* was first recorded in the Kiev region (Smela town) of Ukraine on *Pinus sylvestris* L. by L. Kaznowski (Barnes et al., 2008). Since 2004, DNB outbreak has been reported several times in the Tsjurupinsk (Kherson region) and Mykolay region on 30-year-old Crimean pine (*P. nigra* subsp. *pallasiana* or *P. pallasiana*, or *P. nigra* var. *pallasiana* (D. Don in Lamb) resulted in massive pine dieback in the southern Ukraine (Usichenko & Akulov 2005, Drenkhan et al., 2016). The natural range of *P. nigra* subsp. *pallasiana* covers the southern Carpathians and the Crimean Peninsula in Ukraine, as well as the Balkan Peninsula, Cyprus, the Black sea coast of Caucasus and Turkey (Barnes et al., 2008, Lazarević et al., 2017). Outbreak of DNB has spread in southern Ukraine and south-western Russia devastating more than 8000 ha of pine forests (Usichenko & Akulov, 2005).

Presently, DNB occurs throughout southern Ukraine and its severity appears to be increasing (Drenkhan et al., 2016) alongside climate change. It may act together with native or invasive pathogens and reduce the populations of Crimean pine (Tubby and Webber, 2010, Adamson et al., 2018).

As DNB affects over 80 species of *Pinus*, as well as other conifers (Watt et al., 2005, Drenkhan et al., 2016), there is an urgent need to find effective methods for disease control and management, because DNB resulted in the rejection of planting susceptible *Pinus* species in some countries of Africa, Asia, Australasia, Europe and North America (Bulman et al., 2016). The main practices used to control of DNB in Ukraine are early detection of symptoms and signs of DNB, pathogen population monitoring. No chemicals applied in the Ukrainian forest due to the lack of permissible fungicides. However, copper and other fungicides were applied in the forest and landscape nurseries for pathogen spread prevention to uninfected areas, as well as an elimination of infected seedlings. According to Bulman et al. (2016), DNB may be effectively controlled using copper fungicides as it routinely applied in New Zealand and Australia, or by planting non-susceptible species, as is the most common form of management in Europe. Only a few studies demonstrated the possibility of using biological control agents to reduce the impact of this highly damaging pathogen in forest tree nurseries (Allenzi et al., 2015).

The aim of the study was to find potential biological agents against *Dothistroma* needle blight. This study reports the following results: i) screening needle endophytes to search potential agents of biological control against pine needle pathogen *Dothistroma pini*; ii) artificial inoculation experiments to test the potential of two fungal strains to provide control of Crimean pine infection by *D. pini*.

**Objects and methods.** *Fungal isolates and plant material.* Several endophytic fungi were recovered from needles of Crimean pine during summer sampling of the host’s microbiome in Ukraine in 2012-2014. Asymptomatic, healthy needles were collected from trees growing in the Kherson region (46°31’36.2”N 32°32’01.3”E). Fifty individual needles were sampled in the canopy from each of ten trees of Crimean pine for a total of 500 needles. Needles were surface-sterilized by serial sterilization (1 min in 95% ethanol, 5 min in 6% sodium hypochlorite (NaOCl), and 30 s in 95% ethanol) before plating onto 3% potato dextrose agar (PDA). Smaller sample groups were also taken from diseased pines that appeared to have lower disease severity. The hyphal tip of each morphologically different mycelium that emerged from a needle was subcultured and transferred to for later identification. Following incubation, fungal isolates recovered from each plant fragment were selected at random, purified and grouped based on phenotypic characteristics, e.g. colony morphology, colony colour, and growth rate. Isolates representing each fungal group of interest were selected for further identification by morphological traits (classic taxonomy) and/or rDNA sequencing.

*Dothistroma pini* was isolated from infected needles collected in 2012-2014 in southern Ukraine (Kherson region, 46°31’36.2”N 32°32’01.3”E), and was used in artificial inoculation experiments previously. Following recovery from the plant tissues, representative cultures of dominant genera were made according to morphotypes and stored at 4°C on PDA.

**Selection of endophytic fungi antagonists to *Dothistroma pini*.** Fungi were selected based on the commonness of their occurrence in the foliage of the host and their antagonistic activity. The in vitro selection of antagonists against *D. pini* was carried out on 8% malt extract agar (MEA) medium a paired-growth assay. For this, mycelial discs (5 mm) of *D. pini* were inoculated on Petri dishes (100 mm) containing MEA medium and incubated at 28°C (photoperiod of 12 hours). Due to the slow growth of *D. pini*, after 15 days, the endophytic microorganisms were inoculated 50 mm from *D. pini* colony. The ability of a root endophyte to antagonize the pathogen was determined based on the inhibition level over a given period of time. This was achieved by assessing and measuring the concurrent growth of both the endophyte and the pathogen simultaneously on a shared MEA nutrient media surface (Fig. 1).

The inhibitory effect of each fungal endophyte on the pathogen is reflected in the spherical index (α/β) of the respective organisms (Rigerte et al., 2019). Solitary
cultures of the respective endophyte and the pathogen were also plated—and observed—as controls in this experiment. The antagonism was detected also by the formation of an inhibition halo (Fig. 2). For dual-culture assay the null hypothesis was formulated as follows: the difference between the means of the spherical indices of pathogens under antagonism and the means of the spherical indices of pathogen controls zero (Rigerte et al., 2019). The alternative hypothesis was formulated as follows: the difference between the means of the spherical indices of pathogens under antagonism and the means of the spherical indices of pathogen controls is less than zero (Rigerte et al., 2019). The rationale for assuming this alternative hypothesis was that the spherical index of the pathogen under antagonism would be less than one (<1) while the controls, having grown in the absence of any biotic/abiotic pressure(s), should have a spherical index ≈1 (Rigerte et al., 2019).

Thus, if the effect of specific endophyte is real, then the means would also record the same behaviour, and the subtraction of the means of controls from the pathogen replicates involved in the test would then be a negative number (Rigerte et al., 2019).

**Plant material and fungal inoculation in planta.**

Two-year-old Crimean pine (*P. nigra* subsp. *pallasiana*) seedlings were grown from seeds in the State Forest Enterprise “Holoprystsanske LG” Kherson region. Five hundred seedlings without any symptoms of DNB were replanted in the nurseries at the Forest Protection Service enterprise “Kharkivlysozahist” (Kharkiv region, Ukraine) in March, 2017 for inoculation experiment.

A single plate of a mature culture of each selected fungus was used to obtain inoculum. To generate inoculum, approximately 20 ml of sterile distilled water was added to each Petri dish with mature culture and loosening the spores into suspension by passing sterile glass beads over the surface of the culture, yielding a spore density of from $3 \times 10^4$ to $7.3 \times 10^9$ cells (CFU) ml$^{-1}$ (Tab. 1). Conidial suspensions were adjusted to c. $1 \times 10^4 – 1 \times 10^9$ spores ml$^{-1}$ following replicate haemocytometer counts.

All suspensions were made to a volume of 200 ml with sterile distilled water and placed to the different spray bottles.

All pure cultures of eight selected fungi were made on MEA and PDA.

A solution of 200 ml of sterile distilled water was also included as a control. Concentrations of spores were not standardized across treatments since the fungi
were so diverse and their interactions with the host and the pathogen could not be expected to be comparable.

In July 2017, *P. nigra* subsp. *pallasiana* seedlings were assigned to inoculation treatments with eight endophytic fungi, with 20 replicate plants per treatment. For *in planta* testing for *D. pini* antagonists, 10 days after inoculation of endophytic fungi, spores of *D. pini* were introduced into the seedlings. For each treatment, the foliage was sprayed with a hand-held atomiser until large droplets formed. Control plants were inoculated just with sterile distilled water.

To stimulate DNB development, free water was maintained on needle surfaces. For this, plants were sprayed twice a day with water for 7 days. These incubation conditions were modified from Fraser et al. (2016) and were designed to be optimum for the development of DNB.

The symptoms were evaluated from 60 to 120 days and the data were statistically analyzed by the one-way variance ANOVA method (test compared the means). After 60 days, five needles were collected randomly from each of five plants within each treatment group to determine the fungal inocula loads and percentage conidial germination. Four months after inoculation with *D. pini*, all needles of the previous year were collected and inspected under a microscope. DNB severity was assessed by calculating the percentage of needles with *D. pini* conidiomata. DNB severity was evaluated using the assessment system of Schwelm et al. (2009).

### Table 1

Approximate propagule concentrations per ml of selected fungi suspension for testing their antagonistic activity

| Species             | Medium for cultivation | Spore concentration in suspension |
|---------------------|------------------------|----------------------------------|
| Cladosporium sp     | MEA                    | 1.8×10^6                         |
| Gliocladium roseum  | MEA, PDA               | 2.9×10^4                         |
| Ilyonectria sp.     | MEA                    | 2.7×10^8                         |
| Sydowia polyspora   | MEA                    | 1.9×10^6                         |
| *Trichoderma* sp.1* | MEA, PDA               | 7.3×10^6                         |
| *Trichoderma* sp.2* | MEA                    | 6.2×10^8                         |
| Unidentified Ascomycetes 23 | PDA | 2.3×10^5                         |
| Unidentified Ascomycetes 37 | PDA | 3.7×10^6                         |

* – One *Trichoderma* sp. strain was obtained from the Environmental Sciences culture collection, Natural Research Centre (Vilnius, Lithuania) showing string activity against pine pathogens in artificial inoculation experiments previously.

Molecular detection and identification. DNA was extracted from the selected symptomatic needles representing groups of different treatments. To avoid contamination on the needle surface, needle samples were washed in 96% ethanol for 60 seconds, 2% sodium hypochlorite for 5 minutes and rinsed in 96% ethanol for 30 seconds. Needles were transferred to a screw cap tube together with a screw and two nuts, freeze-dried and homogenized using a fast prep shaker (Precellys 24 Bertin Technologies). DNA was extracted following a CTAB protocol. Briefly, 1 ml of CTAB was added to each sample and incubated for one hour at 65°C. Samples were centrifuged and the supernatant was transferred to new tubes and cleaned with chloroform. DNA was precipitated with isopropanol, washed with 70% ethanol and eluted in 50 ml of milliQ water. After DNA extraction, samples were cleaned using JetQuick DNA purification kit (Genomed GmbH) and the concentration was measured with a NanoDropTM (Thermo scientific).

Conventional PCR with species specific primers was used to detect *D. septosporum* and *D. pini* in the needles, using species-specific primers DStub2-Forward (CGAACATGGACTGAGCAAAATGAC), and DStub2-Reverse (GCACGGCTCTTTTCAAAATGAC), and DPEtF-Forward (ATTTTTGGCTGTCTGTCACT) and DPEtF-Reverse (CAATGTGAGATGTTCGTCGT), respectively (Ioos et al. 2010). The PCR reaction contained 200 mM deoxyribonucleotide triphosphates, 0.2 mM of each of the two primers, 0.0265 u/ml DreamTaq polymerase with 10X DreamTaq Green Buffer (DreamTaq Green, Thermo Scientific, Waltham, MA, USA) and MgCl2 at a final concentration of 3.25 mM. The PCR conditions included an initial denaturation step at 95°C for 10 min followed by 35 amplification cycles of denaturation at 95°C for 30 s, annealing at 60°C for 30 s and extension at 72°C for 1 min, and thermal cycling was ended by a final extension step at 72°C for 10 min. The PCR products were purified with Qiagen DNA extraction PCR M kit (Qiagen, Hilden, Germany). PCR products were size separated on 1% agarose gels and visualized under UV light to confirm the presence of the 231 bp *D. septosporum* specific bands and 191 bp *D. pini* specific bands (Ioos et al. 2010).

Statistical analyses. Statistical analysis was carried out using the software JMP®, Version 11.0.0. SAS Institute Inc., Cary, NC, 1989-2007. Data were
statistically analyzed by the one-way variance method and the Tukey-Kramer multiple comparison. Test compared the means. One-way ANOVA tests were used to assess the impact of treatment on DNB severity. If ANOVA tests were significant, post-hoc Tukey tests were used to identify which treatments differed significantly. DNB severity was log-transformed before analysis. The impact of treatment on conidial germination was assessed with Kruskal–Wallis U-tests and post hoc Mann–Whitney U-tests with Holm corrections.

Results and discussion. The diversity of endophytic fungi was assessed in healthy (Site 1 and 2) and symptomatic (Site 3 and 4) trees of Crimean pines. To avoid contamination and to isolate endophytic fungi only from inner needle tissues, surface disinfection was applied. The endophytic fungal community which was isolated from needles included *Alternaria* sp., *Cladosporium* sp., *Colletotrichum* sp., *Diaporthe* sp., *Dothistroma* sp., *Fusarium* sp. (including *Gibberella avenacea*), *Gliocladium* sp., *Lophodermium* sp., *Mariannaea elegans*, *Sordariomycetes* sp., *Trichoderma* sp. mostly (Tab. 2). The number of most common and fast-growing endophytic fungi that were recovered using MEA and PDA medium was not significantly different within both categories of plants (healthy and symptomatic) evaluated.

**Table 2**

Pooled relative abundance of selected most common fungal taxa obtained from asymptomatic and symptomatic needles collected on Crimean pine grown in south Ukraine (Kherson region)*

| Fungal taxa                             | Site 1 healthy trees | Site 2 healthy trees | Site 3 symptomatic trees | Site 4 symptomatic trees |
|-----------------------------------------|----------------------|----------------------|--------------------------|--------------------------|
| **Ascomycota and other species**        | 1                    | 2                    | 3                        | 4                        |
| *Alternaria* sp.                        | 2.17                 | 2.1                  | 0                        | 6.94                     |
| *Aspergillus versicolor*                | 0.5                  | 2.52                 | 2.1                      | 2.3                      |
| *Bionectria* sp.                        | 5.33                 | 5.2                  | 4.17                     | 3.47                     |
| *Botryotinia fuckeliana*                | 5.14                 | 8.9                  | 1.2                      | 3.5                      |
| *Cadophora* sp.                        | 0.1                  | 0.5                  | 0.42                     | 0.25                     |
| *Chaetomium globosum*                  | 1.3                  | 1.2                  | 1.6                      | 2.51                     |
| *Chaetomium* sp.                       | 2.1                  | 1.05                 | 2.52                     | 3.13                     |
| *Cladosporium* sp.                     | 1.2                  | 2.1                  | 5.25                     | 2.52                     |
| *Cladosporium herbarum*                | 2.2                  | 2.52                 | 6.33                     | 5.25                     |
| *Cladosporium* sp.                     | 10.42                | 5.25                 | 19.6                     | 9.74                     |
| *Cordyceps* sp.                        | 4.17                 | 0                    | 5.14                     | 6.60                     |
| *Colletotrichum* sp.                   | 5.14                 | 1.05                 | 2.52                     | 0.1                      |
| *Cyclaneusma niveum*                   | 0.9                  | 1.3                  | 5.5                      | 2.15                     |
| *Dothistroma* pini*                    | 0.1                  | 0.3                  | 24.5                     | 52.3                     |
| *Dothistroma* sp.                      | 1.05                 | 2.52                 | 5.25                     | 1.05                     |
| *Diaporthe* sp.                        | 0.01                 | 12.5                 | 6.8                      | 0.01                     |
| *Gibberella* avenacea*                 | 18.75                | 15.6                 | 10.42                    | 17.3                     |
| *Gliocladium* roseum*                  | 7.25                 | 5.25                 | 8.15                     | 2.3                      |
| *Eupenicillium* sp.                    | 7.6                  | 1.3                  | 8.33                     | 5.21                     |
| *Fusarium* oxyysporum*                 | 2.08                 | 0.2                  | 0.1                      | 3.13                     |
| *Fusarium* sp.                         | 0.05                 | 4.17                 | 0.1                      | 2.43                     |
| *Ilyonectria* sp.                      | 8.8                  | 2.08                 | 4.25                     | 8.33                     |
| *Lophodermium* seditiosum*             | 4.75                 | 4.17                 | 6.3                      | 9.03                     |
| *Mariannaea* elegans*                  | 9.5                  | 6.8                  | 2.2                      | 2.52                     |
| *Penicillium* sp.                      | 0.01                 | 4.17                 | 0.9                      | 4.86                     |
| *Penicillium* roqueforti*              | 0.5                  | 0.1                  | 0.1                      | 1.74                     |
| *Phoma* sp.                            | 0.1                  | 0.1                  | 4.17                     | 5.21                     |
| *Phomopsis* sp.                        | 0.3                  | 0.1                  | 0.1                      | 0.1                      |
A total of 244 endophytic fungi isolated from needles were randomly picked up and this population was partially characterized by rDNA (partial 18S, ITS-1, 5.8S, ITS-2 and partial 23S) sequencing. The results (see Tab. 2) showed that the most common and fast-growing endophytic fungi associated with the Crimean pine belong mainly to Ascomycetes group (Fig. 3) being the Botryosphaeriaceae, Diaportheaceae, Dothideaceae, and Capnodiaceae families the most frequent. The fungi Fusarium spp. and Cladosporium spp. were the dominant genera and showed the highest diversity (see Tab. 2). No correlation between fungal groups and plant categories was observed. The frequency of fungi isolation was 0.54 and 0.75 for fungal groups and plant categories was observed. The highest diversity (see Tab. 2). No correlation between ITs-1, 5.8s, ITs-2 and partial 23s) sequencing. The was partially characterized by rDNA (partial 18s, needes were randomly picked up and this population was evaluated by a neighbour-joining algorithm (see Fig. 3). Using this strategy, some isolates could not be identified. Cladosporium sp. was the most common genus recovered from needles. Both Sydowia and Trichoderma spp. as well as two unidentified species were also recovered at high frequency in Dothistroma-infected foliage sample and visually healthy needles. Given these findings, seven commonly isolated fungi from the microbiome of Crimean pine, including Gliocladium roseum (anamorph, Clonostachys rosea) (see Tab. 1, 3) and Bionectria sp. were selected and employed as putative disease modifiers.

Clonostachys rosea is a known parasite and antagonist of other fungi (Moraga-Suazo et al., 2011) and the genus Bionectria includes destructive mycoparasites, some of which are used as biocontrol agents of fungal plant pathogens (Schroets, 2001). One Trichoderma sp. strain was obtained from the Environmental Sciences Culture collection, Natural Research Centre (Vilnius, Lithuania) showing high activity against pine pathogens in artificial inoculation experiments previously. So, a total of eight endophytic fungi were evaluated in vitro and in vivo against D. pini. Only two species were able to inhibit the growth of the causal agent of Dothistroma needle blight of Crimean pine in vitro and two in planta (Tab. 3).

Dothistroma needle blight severity expressed as the percentage of needles with conidiomata was significantly lower on plants treated only with Trichoderma sp. 1 and Gliocladium roseum than trees treated with either species. In contrast, DNB severity on plants treated with other fungi was not significantly different from that on plants treated with D. pini. No conidiomata were observed on control plants treated with purified water.

At 120 days after D. pini inoculation, conidial density on needles from plants treated with both Trichoderma sp. 1 and Gliocladium roseum was significantly lower than on plants treated with either antagonistic fungi (Tab. 4). There was no significant difference between the other fungal species and positive treatments in conidial density or germination at either time point. Although conidial density appeared to increase between days 60 and 120, this increase was not significant (paired t-test, p > 0.05).

Overall, disease severity varied significantly by a tree (see Tab. 3), where the tree represents the combined effects of tree resistance and fungi impact. Severity varied significantly from tree to tree from as little as 1.7% to as much as 18.1%. Modifying effects were strongly significant (P < 0.0001), as were their interactions with trees (P < 0.0001) (Tab. 5).

It has previously been shown that endophytic communities vary spatially in the plant or may be dependent on the interaction with other endophytic or pathogenic microorganisms (Allenzi et al., 2015). Moreover, plant susceptibility to DNB is often related to the stress level of the individual and stress can arise from mismatching of the planting stock’s ecological traits to the planting site, root deformities, damage, and desiccation, planting at improper depths in unsuitable soils, poor nutrient and water availability, and increased

|            | 1   | 2   | 3   | 4   | 5   |
|------------|-----|-----|-----|-----|-----|
| Pleosporales sp. | 2.1 | 0.1 | 2.2 | 5.6 |
| Sordariomycetes sp. | 7.9 | 0.1 | 2.3 | 5.6 |
| Sydowia polyspora | 13.00 | 19.5 | 15.9 | 15.4 |
| Trichoderma like asperellum | 12.3 | 15.8 | 6.35 | 15.6 |

Unidentified fungi

|                |       |       |       |       |
|----------------|-------|-------|-------|-------|
| Uncultured Ascomycetes clone H23 | 15.6 | 14.3 | 9.2 | 10.5 |
| Uncultured Ascomycetes clone H37 | 7.9 | 5.1 | 8.6 | 9.7 |
| Unidentified Basidiomycota FG14 | 0.1 | 0.1 | 0.1 | 0.1 |
| Unidentified Basidiomycota FG39 | 0.1 | 0.1 | 0.1 | 0.1 |
| Fungal sp HN18 | 0.1 | 0.1 | 0.1 | 0.1 |
| Fungal sp HN19 | 0.1 | 0.1 | 0.1 | 0.1 |
| Unidentified culture M12 | 0.1 | 0.1 | 0.1 | 0.1 |
| Unidentified culture M74 | 0.1 | 0.1 | 0.1 | 0.1 |

* – fungi for antagonistic tests were selected based on the commonness, so only species common for all four sites were included in the Tab. 2.
exposure to pollutants, xenobiotics and contaminants (Bulman et al., 2013, 2016). So, these endophytic fungi are ubiquitous and may increase the plant resistance by improving tolerance to drought, reducing the phytopathogen settling and promoting plant growth (Allenzi et al., 2015).

**Table 3**

| Impact of potential antagonistic fungal species treatments on *Dothistroma pini* isolates growth and on DNB disease severity on Crimean pine needles |
|---|---|---|
| Inoculation treatments | Inhibition halo, mm | Disease severity, % ± SE)* |
| | | Antagonistic fungi | *Dothistroma pini* |
| *Bionectria sp* | 32.5±2.9 | + | 17.95±0.1* |
| *Cladosporium sp* | 14.2±1.4 | + | 18.11±0.5* |
| *Gliocladium roseum* | 24.2±2.8 | + | 1.71±0.2* |
| *Sydowia polyspora* | 15.3±1.9 | + | 17.22±0.3* |
| *Trichoderma sp.1* | 29.8±1.8 | + | 2.31±0.4* |
| *Trichoderma sp.2* | 19.2±1.4 | + | 17.41±0.2* |
| Unidentified Ascomycetes 23 | 14.8±1.3 | + | 17.34±0.4* |
| Unidentified Ascomycetes 37 | 13.2±3.4 | + | 17.58±0.5* |
| Positive control (*D. pini*) | – | – | 8.34±0.3* |
| Negative control (sterile water) | – | – | 0±0 |

* – treatments followed by the same letter do not differ significantly (Tukey, p > 0.05). Negative control plants were excluded from statistical analysis.

**Table 4**

| Impact of potential antagonistic fungal species treatments on *Dothistroma pini* conidial density and germination on Crimean pine needles |
|---|---|---|---|---|
| Inoculation treatments | After 60 days | After 120 days |
| | *D. pini* conidia density (spores mm−2; mean ± SE) | *D. pini* conidia germination (%; mean ± SE) | *D. pini* conidia density (spores mm−2; mean ± SE) | *D. pini* conidia germination (%; mean ± SE) |
| *Bionectria ochroleuca* | 24.4±1.3* | 20.5±10.0a | 26.2±1.3* | 21.5±9.0a |
| *Cladosporium sp* | 19.3±1.5* | 19.8±9.2c | 19.8±1.4* | 20.0±9.1c |
| *Gliocladium roseum* | 2.4±1.2c | 12.8±6.0b | 2.6±1.1b | 12.3±6.5b |
| *Sydowia polyspora* | 22.1±0.9a | 24.5±5.2c | 22.1±1.1c | 24.9±5.8c |
| *Trichoderma sp.1* | 3.4±1.2c | 9.4±6.5b | 3.5±1.1c | 10.4±6.9c |
| *Trichoderma sp.2* | 14.8±3.1d | 19.5±6.8c | 15.2±3.1d | 20.2±6.8c |
| Unidentified Ascomycetes 23 | 20.5±1.8c | 22.3±7.3c | 21.0±1.7c | 23.1±6.3c |
| Unidentified Ascomycetes 37 | 19.4±1.8c | 24.5±6.8c | 19.9±1.8c | 26.5±4.8c |
| Positive control (*D. pini*) | 30.6±3.2c | 85.9±6.4c | 32.8±4.2c | 87.8±6.9c |
| Negative control (sterile water) | 0±0 | – | 0±0 | – |

**Table 5**

| Statistical data of modifying effects of antagonistic fungi including tree effect |
|---|---|---|---|---|
| Source | DF | F-value | P-value |
| Treatment | 7 | 50.07 | <.0001 |
| Tree | 19 | 957.98 | <.0001 |
| Treatment * tree | 54 | 104.44 | <.0001 |
| N* | Mean DS** | Standard deviation | Standard error | $R^2$ | CV*** | N* |
| 12.006 | 0.527 | 0.2286 | 0.0013 | 0.4337 | 33.3274 | 12.006 |
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Figure 3. Phylogenetic tree showing the relationship between Crimean pine endophytic fungi and other fungal species. The tree was constructed based on the rDNA (ITS1, 5.8S, and ITS2) fragment sequence by using the neighbour-joining method. The bootstrap analysis was performed with 1000 repetitions.
Current methods for control and management include pruning and thinning of stands to reduce humidity, planting of less susceptible or completely resistant host species and the application of copper-based and other modern fungicides (Bulman et al. 2013). With the exception of a few studies with bacteria, the possibility of using biological control agents to reduce the impact of DNB in forest tree nurseries has received little attention (Allenzi et al., 2015) and this study has increased in importance in Europe for application of biological methods in forest protection.

Plant ecosystems rely heavily on their microbial communities to optimize forest health, although this association might be a good possibility to find a balance between mutualism and disease (Rabiey et al., 2019). Endophytes might cover the capacity to directly inhibit pathogens by producing antifungal compounds (Allenzi et al., 2015, Bulman et al., 2016, Rabiey et al., 2019). Most tests and experiments have carried out in laboratory conditions, but it is unknown how the endophyte-pathogen interaction will alter in the presence of changing environmental conditions and competition with other organisms in the tree system (Rabiey et al., 2019).

Thus, much more field experiment should be taken place to recognize and confirm the optimal time and conditions for usage of biocontrol agents, as climate conditions and tree physiology could alter efficacy and efficiency of biocontrol agents may vary greatly depending on climate and tree traits.

**Conclusion.** The use of endophytes as biocontrol agents resulted in that Dothistroma needle blight was reduced on Crimean pine seedlings treated with Trichoderma sp. and Gliocladium rosea. The significant reductions in numbers of conidia and spore germination were found on needles treated with Trichoderma sp. and Gliocladium rosea, compared to numbers following treatment with other fungi. Our result suggested that both these species may possess potential in preventing the Dothistroma needle blight at least on young pines.

Although *D. pini* is present almost everywhere Crimean pine is grown, both severity of disease and area of outbreak vary significantly over time and efficiency of the biocontrol agent application may vary greatly depending on climate and tree traits. That’s why it is important to continue the search of endophytic biological control agents that may alter the microbial community of the host tree and could decrease DNB virulence or enable host resistance. Further work is required on the impact of the fungal species on Dothistroma infections under nursery conditions.

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### References

Adamson, K., Mullett, M.S., Solheim, H., Barnes, I., Müller, M.M., Hantula, J., ... Drenkhan R. (2018). Looking for relationship between the populations of *Dothistroma septosporum* in northern Europe and Asia. *Fungal genetics and biology*, 110, 15-25. https://doi.org/10.1016/j.fgb.2017.12.001

Alenzi, F.N., Fraser, S., Belka, M., Doğuð T., Heckova, Z., Oskay, ... Woodward, S. (2016). Biological control of Dothistroma needle blight of pine with *Aneurinibacillus migulanus*. *Forest Pathology*, 46, 555-558. https://doi.org/10.1111/efp.12237

Barnes, I., Crous, P.W., Wingfield, M.J., & Wingfield, B.D. (2004). Multigene phylogenies reveal that red band needle blight of *Pinus* is caused by two distinct species of *Dothistroma, D. septosporum* and *D. pini*. *Studies in Mycology*, 50, 551–565. Available at: http://www.wi.knaw.nl/images/ResearchGroups/Phytopathology/pdf/PDF%20OP%20NUMMER/201.pdf

Barnes, I., Cortinas, M.N., Wingfield, M.J., & Wingfield, B.D. (2008). Microsatellite markers for the red band needle blight pathogen, *Dothistroma septosporum*. *Molecular Ecology Resources*, 8 (5), 1026-1029. https://doi.org/10.1111/j.1439-0329.2008.00356.x

Bradshaw, R.E. (2004). Dothistroma (red-band) needle blight of pines and the dothistromin toxin: a review. *Forest Pathology*, 34 (3), 163-185. https://doi.org/10.1111/j.1439-0329.2004.00356.x

Bulman, L.S., Dick, M.A., Ganley, R.J., McDougal, R.L., Schwelm, A., & Bradshaw, R.E. (2013). 22 Dothistroma Needle Blight. In book: Infectious forest diseases. Croydon: CABI & CPI Group (UK) Ltd.

Bulman, L.S., Bradshaw, R.E., Fraser, S., Martin-Garcia, J., Barnes, I., Musolin, D.L., ... Tubby, K. (2016). A worldwide perspective on the management and control of Dothistroma needle blight. *Forest Pathology*, 46 (5), 472-488. https://doi.org/10.1111/efp.12305

Eschen, R., Roques, A., Santini, A., (2014). Taxonomic dissimilarity in patterns of interception and establishment of alien arthropods, nematodes and pathogens affecting woody plants in Europe. *Diversity and Distributions*, 21(1), 36-45. https://doi.org/10.1111/ddi.12267

Drenkhan, R, Tomešová-Haataja, V., Fraser, S., Vahalik, P., Mullett, M., Martin-Garcia, J., ... Barnes, I.,(2016). Global geographic distribution and host range of Dothistroma species: a comprehensive review. *Forest Pathology*, 46 (5), 408-442. https://doi.org/10.1111/efp.12290

Fraser, S., Martin-Garcia, J., Perry, A., Kabir, M.S., Owen, T., Solla, A., ... & Vasconcelos, M.W. (2016). A review of Pinaceae resistance mechanisms against needle and shoot pathogens with a focus on the Dothistroma-Pinus interaction. *Forest pathology*, 46 (5), 453-471. https://doi.org/10.1111/efp.12201
Ioos, R., Fabre, B., Saurat, C., Fourrier, C., Frey, P., & Marçais, B. (2010). Development, comparison, and validation of real-time and conventional PCR tools for the detection of the fungal pathogens causing brown spot and red band blights of pine. *Phytopathology, 100* (1), 105-14. https://doi.org/10.1094/PHYTO-100-1-0105.

Lazarević, J., Davydenko, K., & Millberg, H. (2017). Dothistroma needle blight on high altitude pine forests in Montenegro. *Baltic Forestry, 23* (1), 294-302. Available at: https://www.balticforestry.mi.lt/bf/PDF_Articles/2017-23%5B1%5D/Baltic%20Forestry%202017.1_294-302.pdf

Millberg, H., Hopkins, A. J. M., Boberg, J., Davydenko, K., & Stenlid, J. (2016). Disease development of Dothistroma needle blight in seedlings of *Pinus sylvestris* and *Pinus contorta* under Nordic conditions. *Forest pathology, 46* (5), 515-521. https://doi.org/10.1111/efp.12242

Moraga-suazo, P., Opazo, A., González, G., & Sanfuentes, E. (2011). Evaluation of *Trichoderma* spp. and *Clonostachys* spp. strains to control *Fusarium circinatum* in *Pinus radiata* seedlings. *Chilean journal of agricultural research, 71* (3), 412. http://biblioteca.inia.cl/medios/biblioteca/agritec/NR38113.pdf

Rabiey, M., Hailey, L. E., Roy, S. R., Grenz, K., & Jackson, R. W. (2019). Endophytes vs tree pathogens and pests: can they be used as biological control agents to improve tree health? *European Journal of Plant Pathology, 155*, 711-729. Available at: https://link.springer.com/article/10.1007/s10658-019-01814-y

Rigerte, L., Blumenstein, K., & Terhonen, E. (2019). New R-Based Methodology to Optimize the Identification of Root Endophytes against *Heterobasidion parviporum*. *Microorganisms, 7* (4), 102. https://doi.org/10.3390/microorganisms7040102.

Schoers, H. J. (2001). A monograph of *Bionectria* (Ascomycota, Hypocreales, Bionectriaceae) and its *Clonostachys* anamorphs. Utrecht: Centraalbureau voor Schimmelcultures.

Schwelm A., Barron N. J., Baker J., Dick M., Long P. G., Zhang S., & Bradshaw R. E. (2009). Dothistromin toxin is not required for Dothistroma needle blight in *Pinus radiata*. *Plant Pathology, 58*, 293-304. https://doi.org/10.1111/j.1365-3059.2008.01948.x

Usichenko, A. S. & Akulov A. Y. (2005). *Dothistroma septosporum* (teleomorph *Mycosphaerella pini*) – quarantine plant pathogenic fungus revealed in Ukraine. In: *Proceedings of the ‘Fungi in Natural and Anthropogenous Ecosystems*, 1 (pp. 248-253). Saint-Petersburg: V. L. Komarova Botanical Institute of Russian Academy of Sciences (In Russian).

Woods, A., Coates, K. D., & Hamann, A. (2005). Is an unprecedented Dothistroma needle blight epidemic related to climate change? *BioScience, 55* (9), 761-769. https://doi.org/10.1641/0006-3568(2005)055[0761:IAUDNB]2.0.CO;2

Davydenko, K. (2012). Oцінювання грибів-ендофітів для біологічного контролю дотистромозу сосни кримської (*Pinus nigra subsp. pallasiana*)

K. V. Davydenko

Дотистромоз зазвичай викликають два види патогенних грибів, *Dothistroma septosporum* та *Dothistroma pini*. Дотистромоз хвої є одним із найбільш важливих інвазійних хвороб сосни у багатьох країнах. Ця хворoba нещодавно виявлена в Україні та була оцінена як основна загроза для сосни кримської (*Pinus nigra subsp. pallasiana*); для *Pinus sylvestris* вона є менш загрозливою.

На сьогодні існують відомості, що грибні та бактеріальні ізоляти можуть зменшити ріст і патогеність грибів – збудників хвороб рослин. У цьому дослідженні інфікований грибні ізоляти були вибрані з 30-річних насаджень сосни кримської (*P. nigra subsp. pallasiana*) на півдні України. За допомогою фітопатологічних методів упродовж 2012-2014 рр. з хвоїм сосни отримано 244 ізоляти ендофітних грибів, які використовували для аналізу їхньої антагоністичної активності.

Збудники дотистромозу (*Dothistroma* spp.) та потенційно перспективні для подальшого експерименту грибні кулікури виявлені за допомогою ізоляції та застосування специфічних для виду методів полімеразної ланцюгової реакції.

Серед усіх ендотофідів відібрано вісім видів грибів, які були спільними для всіх зразків і мали ознаки антагоністичної активності або раніше були визначені як антагонисти грибних захворювань рослин. Усі відібрані види протестовані на антагоністичну активність стосовно дотистромозу (*Dothistroma* spp.) у чашках Петрі та на саджанцях сосни кримської у розсаднику. Найкращі показники антагоністичної активності стосовно *Dothistroma pini* досягнути під час використання грибів *Trichoderma* sp. та *Gliocladium rosea*. Це дає змогу припустити, що обидва види мають великий потенцій запобігання ураженню і поширенню дотистромозу, прийнятні для молодих деревцях сосни. Значне зменшення кількості конідій збудника та пригнічення росту спор виявлено на хвоїнках, оброблених грибами *Trichoderma* spp. та *Gliocladium rosea* у порівнянні з іншими вида- дями грибів.

Хоча патоген *D. pini* присутній в Україні майже скрізь, де росте кримська сосна, важливо викорис

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Автори використовують біологічні засоби контролю, які могли б зменшити вірусність збудника дотистромозу або забезпечити стійкість рослин-живителів. Та-ким чином, використання ефективного біологічно-го контролю дотистромозу може бути корисним у лісових розсадниках, де важливо зменшити втрати від інфекції D. pini до висаджування крымської со-син в польові умови.

**Ключові слова:** Dothistroma septosporum; Dothistroma pini; патоген; біологічний контроль.

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**Оценка использования грибов-эндофитов для биологического контроля дотистромоза сосны крымской (Pinus nigra subsp. pallasiana)**

Е. В. Давиденко

Дотистромоз обычно вызывают два вида патогенных грибов, Dothistroma septosporum и Dothistroma pini. Дотистромоз хвои является одной из важнейших инвазивных болезней сосны во многих странах. Эта болезнь недавно распростра-нилась в Украине и представляет угрозу преимущественно для сосны крымской (Pinus nigra subsp. pallasiana) и менее серьезную – для Pinus sylvestris.

Сегодня много исследований свидетельствуют о том, что некоторые грибные и бактериальные изоляты могут снизить интенсивность роста и патогенность гриб – возбудителей болезней растений. Для тестирования потенциальных анта-гонистов дотистромоза, в 2012-2014 гг. собраны инфицированные и ненфицированные хвоинки с 30-летней крымской сосны (P. nigra subsp. pallasiana) на юге Украины, и 244 вида эндофитных грибов проверены на возможную антигрибную ак-тивность. Возбудители дотистромоза (Dothistroma spp.) и потенциальные грибные антагонисты определены путем изоляции в чистую культуру и использования специфических видовых при-меров методом полимеразной цепной реакции.

Среди всех эндофитов отобрано восемь видов грибов, которые были общими для всех образцов и имели явные признаки антагонистической актив-ности. Все отобраные виды протестированы на наличие противогрибной активности относительно дотистромоза (Dothistroma spp.) на 2-летних культурах сосны крымской. Лучшие показатели такой активности против Dothistroma pini достигнуты при использовании грибов Trichoderma sp. и Gliocladium rosea. Это позволяет предположить, что оба вида потенциально способны предотвра-тить распространение дотистромоза, по крайней мере, на молодых деревцах сосны. Значительное уменьшение количества конидий патогена и за-медление прорастания спор выявлено на хвоинках, обработанных Trichoderma sp. и Gliocladium rosea, по сравнению с другими грибами. Хотя патоген D. pini присутствует в Украине везде, где растет крымская сосна, важно прилагать больше усилий, чтобы использовать биологические средства кон-троля, которые снижают вирулентность возбуди-теля дотистромоза или обеспечить устойчивость растения-хозяина. Таким образом, использование эффективного биологического контроля против дотистромоза может быть полезным в лесных пи-томниках, где важно уменьшить потери от инфек-ции D. pini до высадки крымской сосны в полевые условия.

**Ключевые слова:** Dothistroma septosporum; Dothistroma pini; патоген; біологічний контроль.