Potential Link between Ectomycorrhizal Community Composition and Host Tree Phenology

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Abstract: The timing of leaf phenology tends to be crucial in controlling ecosystem processes such as the acquisition of carbon and water loss as well as in controlling tree nutrient cycling. To date, tree phenology has mostly been associated with environmental control (e.g., temperature and photoperiod) in a relationship with inheritance, while it has rarely been linked with ectomycorrhizal community compositional changes through the host tree’s phenological stages. Seasonal variations of fungal communities have been widely studied, but little is known about mycorrhiza community composition changes along phenological stages. Therefore, we analyzed ectomycorrhizal communities associated with silver fir and their compositional changes during the transition between phenological stages. The phenological stages of each individual tree and time of bud break were monitored weekly for two years and, at the same time, ectomycorrhiza was harvested from selected silver fir trees. In total, 60 soil cores were analyzed for differences in the ectomycorrhizal community between phenological stages using Sanger sequencing of individual ectomycorrhizal morphotypes. A significant difference in beta diversity for an overall ectomycorrhizal community was confirmed between analyzed time periods for both sampled years. Species-specific reactions to transitions between phenological stages were confirmed for 18 different ectomycorrhizal taxa, where a positive correlation of Russula ochroleuca, Russula illota, Tomentella sublilacina, and Tylospora fibrillosa was observed with the phenological stage of bud burst.

Keywords: Abies alba (Mill.); ectomycorrhizal community; phenological stages

1. Introduction

Silver fir (Abies alba Mill.) is one of the most valuable relic conifer tree species in Europe [1]. It is considered a significant ecological and functional balancer tree species of European forests. Among other functions, it stabilizes soils and retains water [2]. However, little is known about its ectomycorrhizal symbionts and their ecological and physiological role in providing vigor to silver fir [1]. Furthermore, to date, ectomycorrhizal community shifts have rarely been linked to tree leaf phenology, with some of these studies only focusing on deciduous trees. Leaf phenology is an important trait affecting tree survival and growth through its impact on carbon balance and overall tree fitness [3,4]. Due to environmental conditions and endogenous rhythms, tree phenology changes seasonally, from bud bursting to full leaf/needle expansion over a growing season followed by a period of winter dormancy [5]. To date, the timing of belowground organs’ phenology, such as fine roots, paralleled to leaf phenology has received less attention even though it likely influences overall plant functioning and fitness [6].

Among others, mycorrhizal symbiosis is an important interaction and should not be ignored in phenological studies. For instance, root colonization by ectomycorrhiza has been shown to correlate with earlier bud burst in oak seedlings [7] and in our previous study we confirmed a significant difference in ectomycorrhizal community composition between early bud bursting and late bud bursting silver fir seedlings [8].
Although tree phenology and seasonal variations of fungal communities have been widely studied, their interdependence—the above–below ground link between tree phenology and ectomycorrhizal fungal community composition shifts—is largely unknown. Only scarce, early studies have correlated ectomycorrhizal root colonization and tree phenology [7,9,10]. Ectomycorrhizal root colonization most likely affects host phenology through improved mineral absorption capacity of mycorrhizal roots; Courty et al. [11] and Waldrop et al. [12] strongly suggested a mycorrhizal effect on host phenology by providing the carbon to the host tree at a time when demand is high and photoassimilates are not yet available. At the time of bud break, when carbon supply is provided exclusively by the metabolic activity of stored reserves, Dhuli et al. [13] and Courty et al. [11] recorded an increase in organic matter degrading enzymes produced by *Lactarius quietus* just before bud break, which mobilize carbon as glucose at the time of formation of new leaves. Changes in mycorrhizal communities throughout the season also affect nutrient and water availability, and thus directly impact the host tree [14].

Short-term shifts in the ectomycorrhizal community in relation to the ectomycorrhizal tree partner phenology stage of development had not yet been analyzed on a detailed time scale, thus we analyzed ectomycorrhizal fungal community composition changes along the transitions between phenological stages and species-specific reactions to those transitions. We described the ectomycorrhizal fungal community per individual phenological stage of silver fir in native silver fir-dominated forest and hypothesized that the whole ectomycorrhizal community structure will differ between phenological stages. At least for some in our study-identified ectomycorrhizal taxa, species-specific reactions to transitions between silver fir phenological stages are expected.

2. Materials and Methods

2.1. Study Site and Phenological Observations

This study was performed in a silver fir (*Abies alba* Mill.)-dominated forest site measuring 16.47 hectares and located in the ‘Lehen na Pohorju’ area (Pohorje, Slovenia, 46°33′ N 15°20′ E, 469–611 m a. s. l.). The phenological stages of silver fir trees in natural regenerating centers were assessed based on a simplified protocol [15] by visual observations for two years (2016 and 2017). Phenological observations of bud break were carried out once per week. Phenological stages were simplified and determined as (1) dormancy (buds enclosed by needles and not visible), (2) bud bursting (bud elongation, and elongating brush of soft needles emerge), and (3) full needle expansion (shoots with developed needles). Soil/root cores for ectomycorrhizal identification were collected at three different time periods, which were linked to host tree phenology (Table 1).

| Phenophase                      | Sampling Time Year 2016 | Sampling Time Year 2017 |
|--------------------------------|-------------------------|-------------------------|
| Dormancy                       | 16 March                | 20 March                |
| Bud bursting                   | 16 May                  | 28 April                |
| Full needle expansion          | 15 June                 | 25 May                  |

The observations of phenological stages ended when the final phenological stage was confirmed for all analyzed silver fir trees.

2.2. Mycorrhizal Sampling

Ectomycorrhiza sampling collection times were determined by the phenological stage of individual silver fir trees. Ectomycorrhizal samples were harvested by sequential soil coring with a soil corer from ten preselected mature silver fir trees. Each soil core was 3.2 cm in diameter and 20 cm deep. The sampling methodology followed that of Kraigher [16]. Altogether, we collected 60 soil samples in three different time periods and in 2 consecutive
years (Table 1). To avoid roots sampling from other adult tree species, soil samples were collected approximately 1 m from the trunk, whereas individual silver fir trees were at least 5–7 m apart to minimize sampling from the same common mycorrhizal network [17–20].

Individual soil core samples were placed in plastic bags and stored at 4 °C before being processed. All roots from individual soil core samples were soaked in tap water for approximately 15–20 min before being gently washed to remove all soil particles attached to the roots. For further processing, the roots from individual soil core samples were divided into coarse or fine roots. Fine roots were defined as roots with a diameter of less than 2 mm [19,21]. Fine roots were observed in tap water under an Olympus SZX12 stereomicroscope (Olympus, Hamburg, Germany). Altogether, sampling resulted in 105,085 ectomycorrhizal root tips characterized and included in further analyses. The pieces of fine roots were placed above a transparent plexi-glass tray with a grid. The roots were equally distributed throughout the grid, from which one further square was randomly chosen. Within the selected square, all root tips were characterized, counted, and separated into vital and non-vital (dead) ectomycorrhizal root tips. The number of root tips counted and characterized in each sample was 500. Vital ectomycorrhizal root tips were further separated based on morphological and anatomical features and characterized according to the method of Agerer [22]. Although non-vital ectomycorrhizal root tips could be due to specific morpho-anatomical characteristics categorized into morphotype, we ranked them among non-vital/dead ectomycorrhizal root tips. Altogether, we characterized 30,000 ectomycorrhizal root tips, including vital and non-vital (dead) mycorrhizas. The relative abundance of individual ectomycorrhizal morphotypes was thereby calculated as the number of individual ectomycorrhizal morphotype root tips/500 analyzed root tips. Further, three to five ectomycorrhizal root tips of each morphotype from individual soil core samples were kept at −20 °C and later freeze-dried at −92 °C for 48 h. After freeze-drying, roots were homogenized into a fine powder using a Retsch mixer mill (Retsch, Hahn, Germany) and analyzed using molecular methods.

2.3. Molecular Analysis

Total DNA from each morphotype was extracted with a DNeasy Plant Mini kit (Qiagen, Hilden, Germany), following the manufacturer’s instructions. The complete ITS region of nuclear ribosomal DNA was amplified from isolated DNA using the fungus specific primer pair ITS1F and ITS4 [23,24], following the modified procedure described in Sulzbacher et al. [25]. PCR products were run on 1.5% agarose gels in 0.5× TBE buffer and visualized with Gel Doc EQ System, PC (BioRad, Hercules, CA, USA). For morphotypes for which amplification failed, the extracted DNA was diluted 10:1 and re-amplified. The amplification success of morphotypes varies between 78% and 100% per individual sample. Amplified DNA fragments were cut from an agarose gel and purified with innuPREP DOUBLEpurify Kit (Analytik Jena AG, Jena, Germany), following the manufacturer’s instructions. After amplified DNA fragment purification, Sangar sequencing was performed at a commercial sequencing laboratory (Macrogen Inc., Seoul, Korea). All morphotypes were sequenced in both directions with ITS1F and ITS4 primers [23,24]. The obtained sequences were processed in Geneious version 11.1.4 (https://www.geneious.com, accessed on 3 September 2019) [26]. Nucleotide base calls with an error probability greater than 5% were trimmed from read ends to improve read quality, while reads were assembled into contigs at 90% base pair similarity. The BLASTN algorithm from the NCBI website (National Center for Biotechnology Information; https://blast.ncbi.nlm.nih.gov/Blast.cgi, accessed on 3 September 2019) and from the UNITE website (https://unite.ut.ee/ [27], accessed on 3 September 2019) was used to assess the similarity of obtained ITS sequences compared to the sequences in GenBank and UNITE public sequence databases. Sequences that remained unclassified at a kingdom or family level were discarded. The final criteria for a database match were as follows: query cover ≥80% and sequence similarity >92% (cut-off value at genus level) or sequence similarity ≥97–100% (cut-off at species level) [28,29].
The ectomycorrhizal status of identified OTUs (operational taxonomic units) was determined according to Rinaldi et al. [30] and Tedersoo et al. [31]. Sequences for ectomycorrhizal morphotypes were deposited in the GenBank database under accession numbers MK820070-MK820175.

2.4. Statistical Analyses

All statistical analyses were completed in R version 3.5.1 [32] using the community analysis specific package ‘vegan’ [33] and statistical packages ‘mvabund’ [34], ‘ggplot2’ [35], ‘multcomp’ [36], and ‘reshape2’ [37].

The correlation of phenological stages and sampling year with the composition of ectomycorrhizal community was tested with the ADONIS function (analogous for permutational multivariate analysis of variance (PERMANOVA)) in the vegan package [28] in R, which ran for 999 iterations using a Bray–Curtis distance matrix.

The differences in ectomycorrhizal community composition between phenological stages were visualized with a non-metric multidimensional scaling analysis (NMDS) ordination using the Bray–Curtis distance matrix.

Generalized linear models (GLMs) with Poisson regression were used to examine the effect of phenological stages on abundances of individual ectomycorrhizal taxa. For multiple comparisons between the time periods of each GLM, general linear hypotheses (GLHT) with Tukey contrast were used [33]. For taxa abundance comparison between time periods, when separated by sampling year, taxa that had an abundance higher than 2% were included for analysis.

A two-way ANOVA test was used to examine differences in the ratio of explorational types between phenological stages and sampled years. The homogeneity of variance assumptions was checked with a Levene test.

3. Results

3.1. Ectomycorrhizal Community Composition Changes along Transition between Phenological Stages

The environmental conditions in the previous month (PERMANOVA, \( p = 0.002 \)) and year (PERMANOVA, \( p = 0.003 \)) significantly affected ectomycorrhizal community composition in the month and year of sampling.

In sampling year 2016, the time between the phenological stages of dormancy and bud bursting was 9 weeks, and between bud bursting and full needle expansion this was 4 weeks. In sampling year 2017, the time between the phenological stages of dormancy and bud bursting was 5 weeks, and between bud bursting and full needle expansion this was 4 weeks.

For sampling year 2016, NMDS showed a significantly different ectomycorrhizal community of bud bursting silver fir trees compared to dormant trees (PERMANOVA, \( p = 0.005 \)) and trees with fully expanded needles (PERMANOVA, \( p = 0.006 \)), meanwhile, there was an overlap in ectomycorrhizal community between dormant trees and trees with fully expanded needles (PERMANOVA, \( p = 0.080 \)) (Figure 1A). A similar pattern was observed for sampling year 2017, where the ectomycorrhizal community of bud-bursting trees significantly differed from dormant trees (PERMANOVA, \( p = 0.012 \)) but not from trees with fully expanded needles (PERMANOVA, \( p = 0.305 \)). In contrast to sampling year 2016, in 2017 significant differences in ectomycorrhizal community between dormant trees and trees with fully expanded needles were observed (PERMANOVA, \( p = 0.022 \)) (Figure 1B).

3.2. Individual Responses of Ectomycorrhizal OTUs along Transition between Phenological Stages

Ectomycorrhizal morphotyping of vital root tips combined with ITS sequencing retrieved 35 distinct ectomycorrhizal fungal OTUs belonging to 12 different genera, among which the most abundant were *Tomentella*, *Russula*, *Lactarius*, and *Cenococcum*; together, these represented 60.8% of all sequences. For a general view, root tips numbers of individually identified fungi per each phenological stage are represented in Table S2.
Figure 1. Non-metric, multidimensional scaling ordination (MDS) of ectomycorrhizal community based on phenological stages, calculated by Bray–Curtis distance, separated by sampled year. Presented are centroids of each group of samples. (A) In sampling year 2016, bud-bursting trees differ significantly from dormant trees (PERMANOVA, \( p = 0.005 \)) and from trees with fully expanded needles (PERMANOVA, \( p = 0.006 \)). (B) In sampling year 2017, significant differences in dormant trees compared to bud-bursting trees and trees with fully expanded needles were recorded (PERMANOVA, \( p = 0.012 \); \( p = 0.022 \)).

We observed 18 distinct ectomycorrhizal OTUs differing between three different time periods in the two years of sampling. Changes in ectomycorrhizal OTUs abundances along the transition between time periods were analyzed separately per sampling year. In sampling year 2016, dormant silver fir trees were associated with a significantly higher abundance of *Amanita rubescens* Pers. and *Russula badia* Beeli. In contrast, bud-bursting trees were significantly associated with *Neoboletus erythopus* (Pers.) C. Hahn and *Tylospora fibrillosa* (Burt) Donk, while we observed a higher abundance of *Sebacina epigaea* (Berk. and Broome) Neuhoff and *Lactarius vellereus* (Fr.) Fr. in association with a fully expanded needle phenological stage. *Lactarius salmonicolor* R. Heim. and Leclair, *Tomentella stuposa* (Link) Stalpers, and *Tricholoma virgatum* (Fr.) Kumm. were associated equally with dormant and bud-bursting trees and with a significantly lower abundance in the phenological stage of full needle expansion. *Cenococcum geophilum* Fr. had the lowest abundance during dormancy in 2016 while *Russula ochroleuca* Pers. had comparable abundance in all three phenological stages (Figure 2).

Significant differences in ectomycorrhizal OTUs abundances along the transition between time periods were further confirmed for sampling year 2017. Dormant trees were associated significantly with *Elaphomyces granulatus* Fr., *Lactarius lignyotus* Fr., *Russula badia* Beeli, and *Tricholoma virgatum* (Fr.) Kumm. In contrast, bud-bursting trees were associated with *Russula illota* Romagn., *Russula ochroleuca* Pers., *Russula* sp. Pers., *Tomentella subtilacina* (Ellis and Holw.) Wakef., and *Tylospora fibrillosa* (Burt) Donk. Moreover, *Lactarius tabidus* Fr. and *Amphimema byssoidei* (Pers.) J. Erikss. were the only ectomycorrhizal OTUs that were significantly associated with trees with fully expanded needles. (Figure 3).
Significant differences in ectomycorrhizal OTUs abundances along the transition between time periods were further confirmed for sampling year 2017. Dormant trees were associated significantly with *Elaphomyces granulatus* Fr., *Lactarius lignyotus* Fr., *Russula badia* Beeli, and *Tricholoma virgatum* (Fr.) Kumm. In contrast, bud-bursting trees were associated with *Russula ilioata* Romagn., *Russula ochroleuca* Pers., *Russula* sp. Pers., *Tomentella sublilacina* (Ellis and Holw.) Wakef., and *Tylospora fibrillosa* (Burt) Donk. Moreover, *Lactarius tabidus* Fr. and *Amphinema byssoides* (Pers.) J. Erikss. were the only ectomycorrhizal OTUs that were significantly associated with trees with fully expanded needles. (Figure 3).

**Figure 2.** Mean and maximum abundances of selected ectomycorrhizal OTUs at species level in sampling year 2016. Ectomycorrhizal OTUs with abundances below 2% were excluded from the plot. Letters mark significantly different result.

In 2016, the ectomycorrhizal community of bud-bursting trees significantly differed from dormant trees and from trees with needles that were fully expanded. However, there was an overlap between the ectomycorrhizal community of dormant trees and trees with full needle expansion. Similar patterns were observed in 2017, where the ectomycorrhizal community of dormant trees significantly differed from bud-bursting trees as well as from trees with needles that were fully expanded. In sampling year 2017, we observed a dominance of different ectomycorrhizal fungi compared to 2016, which could be due to the dry

**Figure 3.** Mean and maximum abundance of selected ectomycorrhizal OTUs at species level in sampling year 2017. Ectomycorrhizal OTUs with abundances below 2% were excluded from the plot. Letters mark significantly different results.
4. Discussion

Using sequential soil coring and ectomycorrhizal community assessment, 35 different ectomycorrhizal OTUs were identified during three different time periods. The number of identified OTUs is comparable to other published studies in which authors used Sangar sequencing for identifying ectomycorrhizal symbionts of silver fir (varies from 15–42) [1,2,38,39].

By comparing ectomycorrhizal community composition between two sampling years and through three different time periods, which were linked to tree phenology, significant shifts in the community were confirmed. As some authors have already suggested regarding the mycorrhizal effect on host phenology [11–13], we tried to link observed shifts in ectomycorrhizal community composition with silver fir tree phenology, suggesting a possible functional role of ectomycorrhizal fungal species during different phenological stages, especially during the carbon expensive stage, such as bud break. However, other factors, such as fine root dynamic and environmental conditions, were also taken into consideration.

In 2016, the ectomycorrhizal community of bud-bursting trees significantly differed from dormant trees and from trees with needles that were fully expanded. However, there was an overlap between the ectomycorrhizal community of dormant trees and trees with full needle expansion. Similar patterns were observed in 2017, where the ectomycorrhizal community of dormant trees significantly differed from bud-bursting trees as well as from trees with needles that were fully expanded. In sampling year 2017, we observed a dominance of different ectomycorrhizal fungi compared to 2016, which could be due to the dry winter in 2016 and the overall drier year, with a smaller amount of precipitation compared to sampling year 2017 [40].

Overall, through observed findings we confirmed the expected shift in the ectomycorrhizal community between phenological stages. However, in contrast to the findings of Hupperts et al. [41], we did not confirm statistically significant differences in the ectomycorrhizal community between all three phenological stages. Namely, there was an overlap either between dormant trees and trees with fully expanded needles (2016) or between bud-bursting trees and trees with fully expanded needles (2017).

Estimating the role of ectomycorrhizal root tips in nutrient and carbon cycling requires a quantification of their numbers and turnover rates [42]. Observed differences in ectomycorrhizal OTUs abundance between the phenological stages in our study could thereby be a result of the production of new root tips and/or their turnover. The mortality of fine roots is greater during winter [43] as root growth declines substantially over winter when water requirements are at their lowest [44]. In contrast, the initiation of fine root growth in the forest floor coincides with leaf expansion in the forest canopy, and is most rapid in early summer [45]. A significant difference in ectomycorrhizal community composition during bud break and full needle expansion in sampling year 2017 could thereby be linked with fine root seasonal dynamics. However, an overlap between dormancy and full needle expansion in sampling year 2016 is difficult to explain through only fine root dynamics.

Although there have been several attempts to link ectomycorrhizal root tip turnover rates to needle or leaf biomass [42], so far, there are no published data about the possible relation between ectomycorrhizal root tip turnover and needle/leaf phenology. However, due to a relatively long ectomycorrhizal root tips’ lifespan (minimum reported is 129 days and maximum is 675 days) [46–50], the turnover rate in our study can hardly be linked to only fine root dynamics and ectomycorrhizal root tip turnover. Thus, the time-lapse between two sampling periods may depend partly on fine root seasonal dynamics and ectomycorrhizal root tip turnover; other factors, such as host tree phenology, should also be taken into consideration.

The observed changes in the ectomycorrhizal community between dormant and bud-bursting silver fir trees can thereby be linked to the functional role of ectomycorrhizal fungal species. As already reported, ectomycorrhizal symbiosis can help host plants through a timely provision of carbon supply, particularly when demand is high [11,51,52]. Several
studies have pointed to the possible degradative abilities of ectomycorrhizal fungi [53,54], which may have retained and can, under specific conditions, express genes for decomposing enzymes [53,55–57]. For members of genera *Tomentella* and *Tylospora*, Read and Perez-Moreno [58] suggested the species’ ability to decompose carbohydrates. For ectomycorrhizal fungi *Tylospora fibrillosa*, the presence of manganese peroxidase, which is one of the key enzymes responsible for lignin degradation, was confirmed in the study by Cairney et al. [59] as well as in the study by Burke and Cairney [59]. Moreover, Chambers et al. [60] confirmed the extracellular activity of manganese peroxidase in part of a gene identified in the ectomycorrhizal fungus *Tylospora fibrillosa*, which additionally supports the records about capable production of manganese peroxidase by this ectomycorrhizal fungus. Along with *Tylospora fibrillosa*, for members of genera *Russula*, the presence of *laccase* enzymes has already been confirmed [56,57,59]. In our study, ectomycorrhizal fungi *Russula illota*, *Russula sp.*, *Russula ochroleuca*, *Tomentella sublilacina*, and *Tylospora fibrillosa* were the most abundant ectomycorrhizal fungi associated with bud-bursting trees, with significantly higher abundances during bud burst. Thereby, we could potentially link higher abundances of the mentioned species to their functional role and possible degradative abilities during carbon-expensive phenophase, such as bud-brake. In contrast, Hupperts et al. [41] confirmed only higher activity of β-glucosidase during dormancy and a weak positive relation between host phenology and overall ectomycorrhizal community enzymatic activity. Although several ectomycorrhizal fungi are predicted to be able to decompose organic matter, we must take into consideration that they are not able to survive solely as saprotrophs as their ability to decompose lignocellulose and degrade cellulose is much lower compared to wood-decaying fungi and/or litter saprotrophs and insufficient for growth [61].

Among others, the observed ectomycorrhizal fungal occurrence pattern and differences in overall ectomycorrhizal communities between phenological stages could be linked with seasonal variations and the change in environmental conditions. There are quite a few studies in which authors confirmed the significant impact of seasonal variations on ectomycorrhizal community compositional changes [11,14,62–65]; however, direct comparison of our results to published ones would not be relevant due to the upgraded soil sampling in our study, which was performed more frequently (at a monthly basis), rather than once per individual season. Precipitation, soil moisture, and temperature are some of the most important environmental variables affecting ectomycorrhizal community composition and richness [66–69]. The significant impact of precipitation on the fungal community was also confirmed by Hawkes et al. [70], who observed a shift in fungal community in the months when the rainfall during the month before sampling was extremely low. Based on our previously published study within a PhD thesis [40], observed community shifts could thereby be further linked with monthly precipitation. In sampling year 2016, a lower amount of precipitation was recorded in April compared to February and May (Table S3); therefore, a significantly different ectomycorrhizal community during bud break (May) compared to dormancy (March) and full needle expansion (June) could be a result of a change in precipitation. However, in sampling year 2017, a lower amount of precipitation was recorded for February and March; therefore, shifts in the community during dormancy (March) cannot be exclusively linked with precipitation as a similar community was observed during bud break (April) and full needle expansion (May). Shifts in the ectomycorrhizal community could also be reflected by significant changes in temperature [71,72]; however, in our PhD thesis [40], significant changes in temperature were not confirmed.

A possible mechanism for temporal shifts in the ectomycorrhizal community could thereby be a combination of various biotic and abiotic factors, e.g., host trees’ physiology, root turnover, ectomycorrhizal lifespan, and environmental conditions.

5. Conclusions

In this study, we found statistically significant ectomycorrhizal community compositional shifts between three different time periods linked to silver fir needle phenology.
Moreover, we recorded significant differences in ectomycorrhizal fungal OTUs abundances associated with different time periods by which we indicated a potential link between ectomycorrhizal community composition and host tree physiology, especially during bud-burst, which is known to be a highly C-demanding growth phase. The observed results most likely reflect both the influence of host tree physiology and the change in environmental conditions. However, further analyses of ectomycorrhizal fungal enzymatic activities are needed to link shifts in ectomycorrhizal community composition with host trees’ physiology and for a better understanding of the functional ecology of ectomycorrhizal fungal symbionts.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/f12121719/s1. Table S1: General linear hypothesis (GLHT) with Tukey contrast based on generalized linear models (GLMs) with Poisson regression of selected ectomycorrhizal taxa (n.s = p > 0.05; p < 0.001 ***; p < 0.01 **; p < 0.05 *). Ectomycorrhizal taxa with abundances lower than 2% were excluded from statistical analyses; ‘absent’ in the table means that the specified ECM taxa did not appear in one of two compared phenological stages, therefore, comparison with GLHT was not possible (for detailed information look at Table S2). Table S2: Ectomycorrhizal taxa presence (+) or absence (−) during individual phenological stage with number of root tips, separated by sampling year (2016, 2017). Table S3: Monthly precipitation per sampling time and sampling year.

Author Contributions: T.U.N., H.K. and T.G. conceived and designed the experiments; T.U.N. and R.D. performed the experiments; T.U.N. and R.D. analyzed the data; T.U.N. wrote the paper. All authors have read and agreed to the published version of the manuscript.

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Data Availability Statement: The datasets generated during and/or analyzed during the current study are available in the GenBank repository, under accession numbers MK820070-MK820175.

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