Genotypic variation in Norway spruce correlates to fungal communities in vegetative buds

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Abstract
The taxonomically diverse phyllosphere fungi inhabit leaves of plants. Thus, apart from the fungi’s dispersal capacities and environmental factors, the assembly of the phyllosphere community associated with a given host plant depends on factors encoded by the host’s genome. The host genetic factors and their influence on the assembly of phyllosphere communities under natural conditions are poorly understood, especially in trees. Recent work indicates that Norway spruce (Picea abies) vegetative buds harbour active fungal communities, but these are hitherto largely uncharacterized. This study combines internal transcribed spacer sequencing of the fungal communities associated with dormant vegetative buds with a genome-wide association study (GWAS) in 478 unrelated Norway spruce trees. The aim was to detect host loci associated with variation in the fungal communities across the population, and to identify loci correlating with the presence of specific, latent, pathogens. The fungal communities were dominated by known Norway spruce phyllosphere endophytes and pathogens. We identified six quantitative trait loci (QTLs) associated with colonization by the spruce needle cast pathogen Lirula macrospora or the cherry spruce rust (Thekopsora areolata) in asymptomatic tissues were detected. The identification of the nine QTLs shows that the genetic variation in Norway spruce influences the fungal community in dormant buds and that mechanisms underlying the assembly of the communities and the colonization of latent pathogens in trees may be uncovered by combining molecular identification of fungi with GWAS.

KEYWORDS
cherry spruce rust, peroxidase, phenology, Picea abies, Rhizosphaera kalkhoffii, Sydowia polyspora
At any given time during their life cycle, trees are colonized by a wide range of microbes, including fungi. These fungi may reside both epiphytically, on the surface of the tissue, and endophytically, within the host tissue without causing any visible symptoms (Arnold, Henk, Eells, Lutzoni, & Vilgalys, 2007; Rajala et al., 2013; Rodríguez, White, Arnold, & Redman, 2009; Sieber, 2007). The epi- or endophytic fungi associated with a tree have varying ecological roles, such as being mutualistic symbionts (latent) pathogens or facultative saprotrophs (Rajala, Velmala, Vesala, Smolander, & Pennanen, 2014; Saikkonen, 2007). Pathogenic fungi cause damage to the host that can reduce host growth and fitness, such as by decreasing photosynthetic capacity, and causing premature leaf shed or lesion formation (Hanso & Drenkhan, 2012; Pan et al., 2018). Highly diverse fungal communities have been reported in studies on current-year Norway spruce needles, that is those that flushed during the sampling season (Menkis, Marciulytės, Gedminas, Lyniškienė, & Povilaitienė, 2015; Nguyen, Boberg, Ihrmark, Stenström, & Stenlid, 2016). It has been shown that the needle endophyte community in Norway spruce (*Picea abies* L., Karst) varies among genotypes (Rajala et al., 2013, 2014). This would indicate that apart from the presence of fungi with the capacity to colonize the host, environmental and climatic factors (Eusemann et al., 2016; Menkis et al., 2015; Møller & Ahu, 2018; Nguyen et al., 2016; Rodriguez et al., 2009), the assembly of the phyllosphere community associated with a given tree will also depend on factors encoded by the host’s genotype (Cordier, Robin, Capdevielle, Desprez-Loustau, & Vacher, 2012; Rajala et al., 2013, 2014). Although it is hypothesized that the plant-inhabiting community is synergistically determined by environmental and host genetic factors (Horton et al., 2014; Rajala et al., 2013; Terhonen, Blumenstein, Kovaluč, & Asiegbu, 2019), the role of specific host genetic variation under natural conditions is poorly understood. Reports on specific host genetic variation that affects the phyllosphere community composition come from annual plant species and deal primarily with bacterial colonization (Horton et al., 2014; Roman-Reyna et al., 2019; Wallace, Kremling, Kovar, & Buckler, 2018). In a study of 196 accessions of *Arabidopsis thaliana*, Horton et al. (2014) showed that both the presence/absence and abundance of fungal species in the associated phyllosphere communities are influenced by the *Arabidopsis* genotype, but for only the more abundant species in the study. Genome wide association study (GWAS) results have implicated a role of both defence responses and cell-wall integrity in the assembly of *Arabidopsis* phyllosphere fungal communities (Horton et al., 2014). However, hitherto there have been no reports on specific genetic variants affecting phyllosphere fungal community composition in perennial plant species, such as trees. These have a more complex architecture (relative to most annual plants) and are exposed to their changing biotic and abiotic environment over many consecutive growth cycles. These interactions between genetics, environment and time are likely to significantly influence the structure of the phyllosphere community of a host tree.

Conifers dominate the boreal forests in the Northern Hemisphere (Farjon & Page, 1999). Owing to their often large population sizes, outbreeding mating systems and efficient gene flow (wind pollination), conifers are characterized by high levels of heterozygosity and intraspecific diversity, which is reflected in low levels of genetic differentiation between populations (Savolainen, Pyhäjärvi, & Knürr, 2007). Norway spruce is one of the most important conifer species in Europe, both ecologically and economically. Together with Scots pine (*Pinus sylvestris* L.), it essentially makes up the continuous boreal forests of the continent. The fungal community composition of Norway spruce needles has been reported to change along a latitudinal gradient on a continental scale as well as between individual genotypes of Norway spruce within a stand (Nguyen et al., 2016; Rajala et al., 2013, 2014), a pattern which may be considered consistent with horizontal transfer of the fungi, and possibly also with the high intraspecific diversity in conifers (Prunier, Verta, & MacKay, 2016).

The development of high-throughput sequencing (HTS) methods paved the way for the generation of the first draft assembly of the Norway spruce genome (Nystedt et al., 2013). The availability of the genome sequence for Norway spruce has opened new possibilities for the development of genetic markers to produce highly resolved genotypes of an individual tree (Vidalis et al., 2018) and to conduct GWAS (Baison et al., 2019). Similarly, sequencing of the ITS (internal transcribed spacer region) with HTS methods has allowed mycologists to describe fungal communities and community dynamics in various ecosystems in greater detail than before, advancing functional understanding of various ecological processes and phylogenetic relationships (Clemmensen et al., 2013; Kubartová, Ottosson, Dahlberg, & Stenlid, 2012; Rosling et al., 2011; Seena & Monroy, 2016; Tedersoo et al., 2014; Vorišková & Baldrian, 2012). The combination of ITS sequencing of phyllosphere fungi with the recently available genotyping resources in Norway spruce in an association study may provide insights into how tree phyllosphere communities assemble, through the identification of specific fungal taxa and with genetic variants associated with general shifts, in the tree phyllosphere communities. For instance, the communities of seemingly healthy needles often include known (e.g., *Lirula macrosora* or *Rhizosphaera kalkhoffii*) or suspected (e.g., *Phoma herbarum* and *Sydowia polyspora*) needle pathogens (Menkis et al., 2015; Nguyen et al., 2016; Rajala et al., 2013, 2014). Furthermore, in a recent metatranscriptomics study of Norway spruce tissue samples, similar frequencies of fungal transcripts were found both in needle and in bud samples (Delhomme et al., 2015), suggesting that vegetative buds as well as needles harbour active fungal communities, but the study provided no insights into the composition of the bud community.

Here we report the results of a study in the perennial conifer *Norway spruce* that combined ITS sequencing of the fungal communities associated with dormant buds with GWAS of 478 individuals to (a) describe the fungal community associated with dormant buds in a large population of unrelated trees and thus describe the abundance of possible latent pathogen colonizations of asymptomatic tissues, (b) suggest loci in the genome of *Norway spruce* that correlate with variation in the fungal communities across the studied
population, and (c) identify loci in the genome of Norway spruce that correlate with the presence of specific, latent, pathogens.

2 MATERIALS AND METHODS

2.1 Amplification, sequencing and analysis of phyllosphere fungal community

Healthy looking vegetative buds were sampled from 518 trees in the southern Swedish Norway spruce breeding archives, located at Ekebo and Maltesholm. The trees in the breeding archive were planted approximately 35 years ago with 7 m between each tree and 5 m between rows. Buds were collected by hand from exposed branches about 2 m above ground. The collected buds were placed into labelled Ziploc plastic bags. In the field, the samples were stored in sterox boxes filled with cooling blocks. After each day of field work the samples were transferred to −20°C for long-term storage. Total genomic DNA was extracted from approximately five buds per tree using the Qiagen Plant DNA extraction kit (extraction details as described by Baison et al., 2019).

DNA samples were amplified separately using the primer pair gITS7 (5′-xxxxxxxxGTGARTCATCGARTCTTG-3′) and ITS4 (5′-xxxxxxxxXCCTTCGCTTATTTGATATG-3′) (Ihrmark et al., 2012) containing 8-bp sample identification barcodes denoted by x, as previously described (Clemmensen, Ihrmark, Durling, & Lindahl, 2016). Each sample was amplified using unique barcode combinations. The gITS7–ITS4 amplicon amplifies the ITS2 region, which provides good species resolution capacity and can be sequenced throughout the entire length with available HTS technologies (Clemmensen et al., 2016; Ihrmark et al., 2012).

Prior to the amplification of the ITS2 region, DNA quantification was performed using the Qubit ds DNA Broad Range Assay Kit (ThermoFischer) and samples were diluted when needed ensuring sample concentration in the range of 1–10 ng/µl. PCR amplifications were done according to Clemmensen et al. (2016). In brief 50-µl reactions with a final concentration of 1× DreamTaq Green Buffer, 200 µM dNTPs, 750 µM MgCl₂, 1.25 µM DreamTaq polymerase (ThermoFischer Scientific), 0.3 µM Tagged gITS7/ITS4 primer mix and approximately 2.5 ng of template DNA were run with the following cycle conditions: 5 min at 94°C, 27 cycles of 30 s at 94°C, 30 s at 56°C and 30 s at 72°C, and a final 7 min at 72°C; blanks were run as negative controls. The resulting amplicons were purified with the AMPure kit (Beckman Coulter). The concentration of purified PCR products was determined fluorometrically using the Qubit ds DNA Broad Range Assay Kit (ThermoFischer), and equimolar mixes of PCR products from 88 samples, including the blanks, were pooled, creating six sample pools that were used for PacBio SMRT circular consensus sequencing using eight SMRT cells per pool. Construction of the sequencing library and sequencing were carried out by NGI SciLifeLab.

The sequences generated were subjected to quality control and clustering in the SCATA NGS sequencing pipeline (http://scata.mkyop.at.slu.se). Quality filtering of the sequences included the removal of short sequences (<200 bp), sequences with low read quality (any base in the sequence that has a PHRED score < 10) and primer dimers; homopolymers were collapsed to 3 bp before clustering. Sequences that were missing a tag or primer were excluded. The primer and sample tags were then removed from the sequence, but information on the sequence association with the sample was stored as metadata. The sequences were then clustered into different operational taxonomic units (OTUs) that essentially correspond to the species level by single-linkage clustering based on 98.5% similarity. The most common genotype (real read) for clusters was used to represent each taxon. For clusters containing two sequences, a consensus sequence was produced. The fungal taxa were taxonomically identified using the UNITE database version 7.2 (https:// unite.ut.ee/index.php) and the BLASTN algorithm. The criteria used for identification were as follows: sequence coverage > 80%, similarity to species level 98%-100% and similarity to genus level 94%-97%. Sequences not matching these criteria were considered unidentified. To obtain further information on abundant yet unidentified sequences (i.e., suspected Norway spruce clusters), a secondary search was performed in GenBank (NCBI) using the BLASTN algorithm. After removal of singletons and nonfungal sequence reads, the relative abundances of each fungal cluster (OTU) in each sample were calculated. The data set was tested for PCR (polymerase chain reaction) contamination by analysing the PCR blanks, and confounding factors such as variation between sequencing libraries, sites and site characteristics. From the original population samples, 473 trees growing at the Maltesholm site were selected for subsequent analysis.

The 1% most abundantly sequenced OTUs were analysed with multivariate ordination methods in PAST 3.20 (Hammer, Harper, & Ryan, 2001). To identify the main drivers in the data set, a principal component analysis (PCA) was made and the first six eigenvectors from the PCA were analysed in the subsequent GWAS.

OTUs including more than 2% of the reads, corresponding to known conifer pathogens and with a presence in at least 35% of the Norway spruce samples, were selected for targeted GWAS of latent pathogens (Figure S3). Separate files with either relative abundance or presence/absence data were prepared for each of the OTUs that met the criteria for GWAS of latent pathogens and were used in trait-association mapping.

2.2 Norway spruce genotyping and SNP annotation

Generation and evaluation of Norway spruce exome capture is described elsewhere (Vidalis et al., 2018). In brief, 478 samples from a subset of 9,000 maternal trees on which sequence capture was performed using 40,018 previously evaluated diploid probes and samples, were sequenced to an average depth of 15×. Illumina sequencing compatible libraries were amplified with 14 cycles of PCR with the probes being hybridized to a pool comprising 500 ng of eight equimolarly combined libraries following Agilent's SureSelect
Target Enrichment System (Agilent Technologies) protocol. These enriched libraries were then sequenced on an Illumina HiSeq 2500 using the 2 × 100-bp sequencing mode.

Read mapping and initial variant calling is described in detail by Baison et al. (2019). Basically, the sequence reads were aligned to the Norway spruce genome using the Burrows–Wheeler Aligner (BWA; Li & Durbin, 2010) and variant detection utilized the Genome Analysis Software Kit (GATK; DePristo et al., 2011; McKenna et al., 2010) best practice pipeline with a training subset created for application in the Variant Quality Score Recalibration (VQSR) method. Only bi-allelic single nucleotide polymorphisms (SNPs) with a minor allele frequency (MAF) and “missingness” of <0.05 and >20%, respectively, were removed. For the selected set of trees a total of 178,101 SNPs passed the filtering and were used for downstream analysis. Annotation was performed using default parameters of snPEff 4 (Cingolani et al., 2012) and local Norway spruce genome annotated database. Ensembl general feature format (GTF, gene sets) information was utilized to build the Picea abies snPEff database.

2.3 | Trait association mapping

Loadings on the first six axes from the PCA of the relative abundance data of the 1% largest OTUs were used for the GWAS, with each individual axis explaining at least 5% of the variance in the detected phyllosphere communities. Subsequently, the relative abundance or presence/absence data for the OTUs (OTU_5, OTU_9, OTU_15 and OTU_19) on each host genotype were used for the trait-association mapping.

The statistical LASSO model as described by Li et al. (2014) was applied to the traits associated with the detected phyllosphere communities and phyllosphere pathogens.

The LASSO model is:

\[
\min_{(\alpha_0, \alpha)} \frac{1}{2n} \sum_{i=1}^{n} \left( y_i - \alpha_0 - \sum_{j=1}^{p} x_{ij} \alpha_j \right)^2 + \lambda \sum_{j=1}^{p} |\alpha_j|,
\]

where \( y_i \) is the phenotypic value of an individual \( i \) \( i = 1, ..., n; n \) is the total number of individuals, \( \alpha_0 \) is the population mean parameter, \( x_{ij} \) is the genotypic value of individual \( i \) and marker \( j \) coded as 0, 1 and 2 for three marker genotypes AA, AB and BB, respectively, \( \alpha_j \) is the effect of marker \( j \) \( j = 1, ..., n; n \) is the total number of markers, and \( \lambda >0 \) is a shrinkage tuning parameter. A fundamental idea of LASSO is to utilize the penalty function to shrink the SNP effects toward zero, and only keep a small number of important SNPs which are highly associated with the trait in the model.

The stability selection probability (SSP) of each SNP being selected by the model was determined to identify significant SNPs (Gao et al., 2014; Li & Sillanpää, 2015). For a marker to be declared significant, an SSP inclusion ratio (Frequency) was calculated for all selected SNPs for each trait and a minimum inclusion frequency of 0.56 was chosen as the most prudent cut-off. The set of markers with nonzero effects was recorded; after bootstrapping, this provides an approximation of a p-value. The SSP threshold used for defining significant SNPs was estimated as suggested by Meinshausen and Bühlmann (2010). The threshold is calculated based on the number of SNPs, number of individuals included in the subset and expected number of false positives. A lambda of 250 with 1,000 bootstraps and a false positive cutoff of five (5) were applied to the entire association analysis. Population structure was accounted for in all analyses by including the first five components from a PCA of the genotypic data as covariates in the LASSO model and the total variance explained by the five PCs was 19.3%. Finally, an adaptive LASSO approach (Zou, 2006) was used to determine the percentage of phenotypic variance (PVE; \( H^2_{\alpha j} \)) of all quantitative trait loci (QTLs). The analyses were all performed with glmnet in RStudio, R version 3.4.0 (Team, 2015), and the codes used can be found at https://github.com/RosarioGarciaLab/Norway-Spruce-Association-Mapping.

Information on putative candidate genes associated with the QTLs and the expression pattern of the candidate genes in the Norway spruce clone Z4006 were collected from the publicly available Norway spruce genome portal and P. abies exAtlas (https://www.congenie.org). The position of the detected QTLs in the Norway spruce genome was estimated by searching an ultradense genetic map (Bernhardsson et al., 2019) for markers derived from the same probes from which the SNP markers holding the QTLs originated.

3 | RESULTS

3.1 | Norway spruce buds are colonized by well-known phyllosphere endophytes and pathogens

After quality control, 676,375 ITS sequence reads remained. At 98.5% similarity, these sequences clustered into 4,899 OTUs, excluding 3,357 singletons. Fungal taxa were identified using the UNITE database. However, the dominant cluster (85% of reads, a spruce ITS sequence [AJ243167] and 10 other nonfungal clusters were removed from the data set leaving 4,888 OTUs [104,768 sequence reads]). After removal of the plant clusters the median read count over the samples in the data set was 148 reads. Relative abundances in each sample were calculated on the remaining OTUs. After testing and removal of confounding factors and of samples with poor amplification/sequencing success (total number of remaining fungal reads <20) the analyses were limited to the largest clusters only (Horton et al., 2014), leaving 473 Norway spruce samples and 49 OTUs (top 1%, including 80% of the fungal reads, Supporting Information S1) in the subsequent analyses. The majority of these belonged to Ascomycota, 56% belonged to Pezizomycotina alone and two OTUs (4%) represented Taphrinomycotina (Figure S2). The basidiomycete orders Agaricomycotina (25%), Pucciniomycotina (13%) and Ustilagomycotina (2%) were among the most abundant OTUs (Figure S2). Several of the most abundant OTUs represented yeasts (e.g., OTU_2 and OTU_1), and correspond to the basidiomycete yeasts Curvibasidium cygneicollum and Filobasidium wieringae,
| Cluster ID | Cluster size | Samples (%) | Reference | Accession | Score | E-value | % |
|-----------|--------------|-------------|-----------|-----------|-------|---------|---|
| OTU_2     | 10,385       | 81.5        | *Curvibasidium cygneicollum* | KY102972 | 573   | 3.00E-162 | 100 |
| OTU_1     | 9,573        | 78.3        | *Filobasidium wieringae* | KY103446 | 608   | 8.00E-173 | 100 |
| OTU_4     | 8,116        | 85.6        | Fungi     | KP897174 | 460   | 2.00E-128 | 100 |
| OTU_3     | 6,942        | 96.1        | *Cladosporium herbarum* | MF326605 | 449   | 4.00E-125 | 100 |
| OTU_5     | 6,855        | 81.7        | *Sydowia polyspora* | KU837235 | 473   | 2.00E-132 | 100 |
| OTU_6     | 5,878        | 81.3        | *Aureobasidium pullulans* | MF687197 | 460   | 2.00E-128 | 100 |
| OTU_8     | 4,796        | 76.5        | *Vishniaczyma victoriae* | MF927673 | 433   | 4.00E-120 | 100 |
| OTU_10    | 4,018        | 42.6        | Fungi     | KP897236 | 473   | 2.00E-132 | 100 |
| OTU_7     | 3,716        | 71.6        | *Itersonilia pannonica* | KY103613 | 470   | 3.00E-131 | 100 |
| OTU_11    | 2,604        | 49.5        | Ceramothyrium sp. | KC978733 | 427   | 2.00E-118 | 97.3 |
| OTU_12    | 2,337        | 61.9        | Xylariales | KP897169 | 494   | 2.00E-138 | 100 |
| OTU_13    | 2,259        | 63.9        | Perusta inaequalis | AY560007 | 473   | 2.00E-132 | 100 |
| OTU_19    | 2,200        | 39.8        | *Lirula macropora* | HQ902159 | 392   | 6.00E-108 | 99.1 |
| OTU_9     | 2,183        | 41.4        | *Thekopsora areolata* | DQ087231 | 575   | 8.00E-163 | 100 |
| OTU_14    | 2,178        | 64.1        | Tris traticolus sp. | KT696338 | 451   | 1.00E-125 | 97.0 |
| OTU_15    | 2,057        | 57.6        | *Rhizosphaera kalkhoffii* | KY003236 | 475   | 6.00E-133 | 100 |
| OTU_18    | 1,813        | 77.5        | *Sporobolomyces roseus* | MF102879 | 555   | 9.00E-157 | 100 |
| OTU_17    | 1,813        | 71.0        | Melampsoraceae | KCI83505 | 593   | 2.00E-168 | 99.7 |
| OTU_24    | 1,190        | 72.8        | Botrytis cinerea | MF996364 | 444   | 2.00E-123 | 100 |
| OTU_23    | 1,084        | 65.9        | *Ramularia vaccinnicola* | KY979777 | 435   | 1.00E-120 | 100 |
| OTU_20    | 995          | 60.0        | Rhodosporidobolus colorit | KY104695 | 556   | 3.00E-157 | 100 |
| OTU_22    | 908          | 60.0        | Exobasidium sp. | EU692770 | 366   | 4.00E-100 | 96.4 |
| OTU_25    | 824          | 60.4        | Cystofilobasidiales | HM240834 | 571   | 1.00E-161 | 100 |
| OTU_37    | 807          | 61.1        | Leotiomycetes | KMS19282 | 433   | 4.00E-120 | 99.2 |
| OTU_29    | 785          | 64.5        | Taphrina carpini | AY239215 | 536   | 3.00E-151 | 99.7 |
| OTU_21    | 745          | 53.8        | Fungi     | KP891308 | 654   | 0       | 100 |
| OTU_30    | 726          | 65.5        | *Phoma herbarum* | KT355016 | 460   | 2.00E-128 | 100 |
| OTU_34    | 690          | 56.0        | Helotiales | AY971722 | 377   | 2.00E-103 | 99.5 |
| OTU_36    | 647          | 56.6        | Dothiora sp. | KU728514 | 396   | 5.00E-109 | 94.3 |
| OTU_39    | 584          | 54.6        | *Cyphellophora sessilis* | KPS00571 | 518   | 1.00E-145 | 100 |
| OTU_43    | 547          | 55.8        | Fungi     | KPS97199 | 442   | 6.00E-123 | 100 |
| OTU_44    | 539          | 58.2        | Dothideomycetes | FR773250 | 433   | 4.00E-120 | 100 |
| Cluster ID | Cluster size | Samples (%) | Reference | Accession | Score | E-value | % |
|-----------|--------------|-------------|-----------|-----------|-------|---------|---|
| OTU_41    | 459          | 47.7        | Phaffia sp. | HF558647  | 590   | 3.00E-167 | 97.2 |
| OTU_45    | 432          | 48.3        | Trichomium sp. | KP004468 | 444   | 2.00E-123 | 97.0 |
| OTU_50    | 408          | 48.5        | Bannozyma sp. | KY558343  | 551   | 3.00E-156 | 99.7 |
| OTU_53    | 383          | 53.8        | Cyphellophora | AM902002  | 484   | 1.00E-135 | 100  |
| OTU_49    | 375          | 50.7        | Chaetothyriales | GQ999538  | 518   | 1.00E-145 | 99.7 |
| OTU_55    | 366          | 51.9        | Fungi      | KP897431  | 529   | 5.00E-149 | 100  |
| OTU_56    | 362          | 52.3        | Botryosphaeriales | KM216350  | 427   | 2.00E-118 | 99.2 |
| OTU_57    | 355          | 55.4        | Dioszegia aurantiaca | KY103346  | 407   | 2.00E-112 | 100  |
| OTU_54    | 353          | 52.1        | Genolevuria sp. | KF036585  | 460   | 2.00E-128 | 97.4 |
| OTU_47    | 348          | 45.6        | Cystofilobasidium capitatum | KY103158  | 593   | 2.00E-168 | 100  |
| OTU_72    | 301          | 21.6        | Naeva sp.  | JN120378  | 429   | 5.00E-119 | 98.8 |
| OTU_61    | 300          | 23.0        | Alternaria infectoria | Y17066   | 468   | 1.00E-130 | 100  |
| OTU_62    | 292          | 18.6        | Taphrinia padi | KU134843  | 564   | 2.00E-159 | 99.7 |
| OTU_60    | 286          | 18.8        | Fungi      | KP897203  | 477   | 2.00E-133 | 100  |
| OTU_77    | 267          | 23.0        | Symmetrospora sp. | EU002876  | 507   | 3.00E-142 | 97.0 |

*a*Cluster ID is the identity of the assembled OTU.

*b*Cluster size is the number of reads that are associated with the OTU.

*c*Samples (%) is the frequency of the OTU presence in the 493 samples.

*d*The taxonomic assignment where “Reference” is the taxonomic assignment based on BLAST searches in UNITE, “Accession” is the best hit, “score” and “E-value” are the BLAST score and E-values, and “%” is the percentage identity in the alignment.
respectively while OTU_6 corresponds to the common yeast-like ascomycete *Aureobasidium pullulans* (Table 1). The abundant OTU_3 is probably a member of the *Cladosporium herbarum* complex (Table 1). Other OTUs comprise well-known needle endophytes, such as *Ceramothyrium* (OTU_11) and *Perusta inaequalis* (OTU_17) (Table 1). OTU_4, one of the most abundant OTUs, is apparently a hitherto undescribed ascomycete fungus (Table 1), and is identical to the relatively abundant sequence cluster “Unidentified sp. 2168.9” reported from the Norway spruce phyllosphere (Menkis et al., 2015). Several of the abundant OTUs represent needle pathogens: OTU_9 (*Thekopsora areolata*), OTU_15 (*Rhizospherea kalkhoffii*) and OTU_19 (*Lirula macrospora*), or latent pathogens: OTU_5 (*Sydowia polyspora*), OTU_24 (*Botrytis cinerea*) and OTU_30 (*Phoma herbarum*) (Table 1).

To reduce the number of variables for the subsequent GWAS analysis, the relative abundance data of the OTUs/species with the highest number of reads were used in a PCA to identify the most prominent drivers in the data set. The first two principal components (PCs) explained 14.9% and 12.8% of the total variance respectively in the phyllosphere community. The first and second PCs were strongly influenced by the abundance data of the undescribed ascomycete (OTU_4) and *C. cygneicollum*, while the second axis was also driven by *C. herbarum*, *S. polyspora* and OTU_10 (unclassified fungus; Figure 1a; Table 1). In addition to several of the taxa shaping the first two PCs, *F. wieringae* appeared to be an important driver on the third, and *T. areolata* on the fourth (Figure 1b; Table 1). PCs 5 and 6 explained 7.8% and 6.3% of the variance, respectively but the axes were shaped by mostly different OTUs than the first axis, such as *A. pullulans*, *T. areolata*, *R. kalkhoffii*, *Ceramothyrium* sp. and *L. macrospora* (Figure 1c; Table 1).

### 3.2 Trait association-mapping

Using the loadings on the first six PCs of the PCA (Figure 1) for GWAS, six QTLs associated with the fungal community that determine the PCs were detected (Table 2). The contigs of four of these QTLs could be identified on different linkage groups in an ultradense Norway spruce genetic linkage map (Bernhardsson et al., 2019; Table S4), indicating that the QTLs are located at different positions in the genome. The three SNPs that were significantly associated with PC3 are positioned in or adjacent to four Norway spruce gene models: MA_24477g0010, MA_31029g0010, MA_10428833g0010 and MA_10428833g0020 (Table 3). The gene ontology (GO) terms GO:0007264 (intracellular protein transport) and GO:0006886 (small GTPase mediated signal transduction) and GO:0006886 (intracellular protein transport) are associated with the gene model MA_24477g0010 that is similar to the *Arabidopsis* ADP-ribosylation factor A1F gene (Table 3). MA_10428833g0020, encoding a putative ubiquitin-protein ligase, is associated with the terms GO:0006817 (phosphate ion transport), GO:0016036 (cellular response to phosphate starvation), GO:0010337 (regulation of salicylic acid metabolic process), GO:0080021 (response to benzoic acid) and GO:0009626 (plant-type hypersensitive response). Neither of the other two gene models have any functional information connected to them in the Norway spruce genome portal. However, MA_31029g0010 is homologous to AT3G59340, a putative solute carrier family 35 protein which is reported to be located in *Arabidopsis* chloroplasts. Inspection of the expression patterns of MA_24477g0010, MA_31029g0010, MA_10428833g0010 and MA_10428833g0020 reported in the *Picea abies* exAtlas (Nystedt et al., 2013) suggest that the candidate genes may be highly expressed in bark phloem, cambium and primary xylem (samples Z4006TR12, Z4006TR24, Z4006TR25 and Z4006TR19) and that relatively high levels of MA_31029g0010 transcript accumulate in vegetative shoots, developing buds, pine apple galls and immature female cones (Figure 2). Interestingly, the gene model MA_208236g0010 associating with PC5 showed an expression pattern reminiscent of MA_31029g0010 with relatively high transcript expression accumulating in vegetative shoots and developing buds (Figure 2). There is no functional information for MA_208236g0010 in the Norway spruce genome portal but the putative *Arabidopsis* orthologue AT2G47820 encodes an arginine-glutamic acid dipeptide repeat protein, and PLAZA Homology enrichment analysis ([https://bioinformatic.s.ps.ugent.be/plaza/](https://bioinformatic.s.ps.ugent.be/plaza/)) suggests that the molecular function of GO:0003677 (DNA binding) is associated with the product of the gene model.

The candidate gene MA_10433886g0010 associated with PC4 is linked with the GO-term GO:007067 (mitosis) in the *P. abies* exAtlas (Nystedt et al., 2013) and its peak expression is found in flushing vegetative shoots although it is also expressed in, for example, buds. MA_19950g0010, the other candidate gene associated with PC4, appears to act in signal perception or transduction as its *Populus* orthologue and best *blastx* hits both encode WD-40 repeat family proteins (Table 3). This gene model has a contrasting expression pattern between early- and late-season buds (Figure 2). The gene models associated with PC3 have very low levels of expression in vegetative shoots that have just began to flush (Z4006TR02), while the gene models associated with PC4 and PC5 show relatively high expression in this tissue (Figure 2).

The presumed latent pathogen *S. polyspora*, the cherry spruce rust fungi (*T. areolata*) and the species causing needle cast diseases (*L. macrospora* and *R. kalkhoffii*; Horst, 2013; Kaitera, 2013; Pan et al., 2018; Rajala et al., 2013) met the criteria for targeted GWAS of specific, latent, pathogens. Trait-association mapping with presence/absence or relative abundances of *S. polyspora* or *R. kalkhoffii* yielded no QTLs, whereas two QTLs for *L. macrospora* presence/absence in the host phyllosphere communities and one for *T. areolata* presence/absence were identified (Table 2). None of these QTLs was shared with the QTLs identified with the 1% most abundant OTUs; these traits were associated on independent SNPs, and appeared to be located at different positions in the genome (Table S4). The *L. macrospora* presence/absence in the host phyllosphere communities was associated with SNPs positioned in or adjacent to the Norway spruce gene models MA_97571g0020 and MA_10432519g0010 and the gene model MA_10g0010 was associated with *T. areolata* presence/absence (Table 3). No functional information is associated with MA_97571g0020 in the Norway spruce genome portal,
but the PFAM ID PF03801 (HEC/Ndc80p family) is associated with MA_10432519g0010; high relative expression levels of this gene model are found in young vegetative shoots (Z4006TR01) and in immature male cones in the _P. abies_ exAtlas (Figure 2). MA_10g0010, associated with _T. areolata_, encodes a previously undescribed class III peroxidase. This candidate gene has a very distinct expression pattern, high expression levels of this gene model are found in young vegetative shoots, developing buds, wood and girdled twigs (Figure 2), and very high relative expression is found in immature female cones (Figure 2).
TABLE 2 Significant association in the GWA study

| Trait  | SNP             | Allele | SNP feature | Frequency | PVE  |
|--------|-----------------|--------|-------------|-----------|------|
| PC3    | MA_24477_24501  | T*C    | downstream_gene | 0.649     |      |
|        | MA_31029_9337   | C*G    | missense    | 0.633     |      |
|        | MA_10428833_21190 | C*T  | upstream_gene | 0.552     |      |
| PC4    | MA_19950_16139  | G*C    | upstream_gene | 0.533     |      |
|        | MA_10433886_12255 | T*A  | upstream_gene | 0.682     | 1.2% |
| PC5    | MA_208236_3389  | A*G    | stop_retained | 0.565     |      |
| L. macrospora | MA_97571_20468 | C*A    | upstream_gene | 0.566     |      |
|        | MA_10432519_8378 | C*A  | synonymous   | 0.566     |      |
| T. areolata | MA_10_25927    | G*A    | upstream_gene | 0.993     |      |

The trait upon which the marker associates, PC3–PC5 indicate the associations with loadings on the respective PC, and L. macrospora and T. areolata specify associations with the presence/absence of data in fungi among the samples.

The SNP name consists of the contig (MA_number) and SNP position on the contig. For example, the first SNP MA_24477_24501 was located on contig MA_24477 at position 24,501 bp.

Allelic variation associated with the SNP.

Stability selection probability inclusion ratios for markers declared significant.

Phenotypic variance explained (only values larger than 1.0% are displayed).

4 | DISCUSSION

Foliar fungi of conifers are commonly predicted to be horizontally transmitted and the phyllosphere community of a given tree would be expected to represent a sample of the fungi present in the environment (Millberg et al., 2015; Rodríguez et al., 2009; Terhonen et al., 2019). In angiosperm tree species, the genetic distance between host tree individuals may have a critical impact on assembly of the fungal community (Ahlholm, Helander, Henriksson, Metzler, & Saikkonen, 2002; Cordier et al., 2012). The impact of the tree genotype is unclear in conifers where some studies report an effect of the host genotype on community composition (Rajala et al., 2013), and others find no such correlation (Eusemann et al., 2016). In this study, combining ITS sequencing and GWAS techniques, we provide further evidence that host genotype influences the composition of the phyllosphere community in the significant marker trait-association mapping of the variation in communities associated with dormant Norway spruce buds.

To our knowledge, no studies of conifer phyllosphere communities have used vegetative buds as a template. However, in a transcriptomic study of various Norway spruce tissues the highest frequency of fungal transcripts was found in needle and bud samples (Délhomme et al., 2015), suggesting that dormant vegetative buds, like needles, may be colonized by phyllosphere fungi. The results from our study further support this, as the taxonomic identities of many of the most heavily sequenced OTUs show affinity to previously reported members of the Norway spruce phyllosphere fungal community. For instance, *Phialophora sessilis*, *Taphrina carpini*, *Ceramothyrium* sp., *Phoma herba-rum*, *Aureobasidium pullulans*, *Sydowia polyspora*, *Rhizosphaera kalkhoffii* and *Lirula macrospora* are species which have been reported from studies on Norway spruce needle communities (Menkis et al., 2015; Nguyen et al., 2016; Rajala et al., 2013, 2014). The frequencies and abundances of phyllosphere fungi differ depending on the species present in the local environment (Nguyen et al., 2016; Rodríguez et al., 2009), physiological status of the host tree (Menkis et al., 2015; Rajala et al., 2014), tissue sampled and sample handling. The current study is an illustration of this, as the fraction of yeast-like fungi was higher here compared to earlier studies. The presence of a large fraction of yeast species may be due to the use of nonsurface-sterilized explants for DNA extraction, unlike in some previous studies (Nguyen et al., 2016; Rajala et al., 2013). This indicates a need for a similar marker trait association approach with both surface sterilized and nonsterilized explants and perhaps with the amplification and sequencing of the ITS2 region from several single buds from each tree instead of from a pooled sample. However, extraction and community sequencing of nonsurface-sterilized material as in our study, and the study by Menkis et al. (2015), indicates that the Norway spruce phyllosphere may have quite abundant, yet undescribed and possibly specific, epiphytes as illustrated by OTU_4. This unidentified taxon was one of the most common fungi in the phyllosphere community in the current study and the study by Menkis et al. (2015). It dominated on the first two PCA axes together with the basidiomycete yeast *Curvibasidium cygneicollum*, the common outdoor mould *Cladosporium herbarum*, to some extent the weak pathogen *S. polyspora*, and *Fibobasidium wirtingiae*, suggesting that these axes may be driven primarily by highly frequent and highly abundant phylloplane fungi. It has previously been reported that tree genotypes can influence the occurrence of leaf epiphytes (Bálint et al., 2013; Cordier et al., 2012), but no marker-trait associations were detected with the two first PCs in this study. Possibly a much larger number of trees would be needed to pick up any association with the community on dormant buds. Other PCs were driven more strongly by well-known needle and wood colonizers such as *S. polyspora*, the biotrophic cherry spruce rust fungus *Thekopsora areolata* (primarily PC3 and PC4) and *L. macrospora* (PC5 and PC6).

Several, probable latent pathogens were identified among the most abundant species: *T. areolata*, *R. kalkhoffii*, *L. macrospora*,
| Trait | Candidate gene | GO terms | Orthologues | Best BlastX hit |
|-------|----------------|-----------|-------------|-----------------|
| **PC3** | MA_24477g0010 | GO:0005215—transporter activity, GO:0005525—GTP binding, GO:0005794—Golgi apparatus, GO:0006471—protein ADP-ribosylation, GO:0006886—intracellular protein transport, GO:0007264—small GTPase mediated signal transduction, GO:0016192—vesicle-mediated transport, GO:0016787—hydrodrome activity | Potri.002G191400.1, AT1G0630.1 | XP_024995498.1, ADP-ribosylation factor 2 isoform X1 [Cynara cardunculus var. scolymus] |
| | MA_31029g0010 | N/A | Potri.014G151400.1, AT1G10630.1 | XP_024995498.1, ADP-ribosylation factor 2 isoform X1 [Cynara cardunculus var. scolymus] |
| | MA_10428833g0010 | N/A | N/A | ABR18169.1, unknown [Picea sitchensis] |
| | MA_10428833g0020 | GO:0004842—ubiquitin-protein ligase activity, GO:0005634—nucleus, GO:0005739—mitochondrion, GO:0005829—cytosol, GO:0007067—mitosis, GO:0009507—chloroplast | Potri.016G02860.1, ABR18169.1, unknown [Picea sitchensis] |
| **PC4** | MA_19950g0010 | N/A | Potri.014G027200.1 | XP_006826575.1, protein ROOT INITIATION DEFECTIVE 3 isoform X2 [Amborella trichopoda] |
| | MA_10433886g0010 | GO:0005634—nucleus, GO:0005739—mitochondrion, GO:0005829—cytosol, GO:0007067—mitosis, GO:0009507—chloroplast | Potri.006G263300.2, AT5G47690.3 | XP_003635522.2, PREDICTED: sister chromatid cohesion protein PDS5 homolog A-like [Vitis vinifera] |
| **PC5** | MA_208236g0010 | N/A | Potri.008G210200.1, AT2G47820.2 | XP_020082821, uncharacterized protein LOC109706424 isoform X2 [Ananas comosus] |
| *Lirula macrospora* | MA_97571g0020 | N/A | N/A | XP_006350818, PREDICTED: acyl-coenzyme A thioesterase 13-like [Solanum tuberosum] |
| | MA_97571g0010 | GO:0003863—3-methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transfering) activity, GO:0005739—mitochondrion, GO:0008152—metabolic process | Potri.005G185400.1, AT5G09300.2 | XP_020524476.1, 2-oxoisovalerate dehydrogenase subunit alpha 2, mitochondrial isoform X2 [Amborella trichopoda] |
| | MA_10432519g0010 | N/A | Potri.011G208700.1, AT3G54630.1 | XP_021663255.1, kinetochore protein ndc80 [Hevea brasiliensis] |
| *Thekopsora areolata* | MA_10g0010 | GO:0008152—metabolic process, GO:0016491—oxidoreductase activity, GO:0016772—transferase activity, transferring phosphorus-containing groups, GO:0046872—metal ion binding | Potri.007G074700.1, AT5G47000.1 | XP_006847800.1, peroxidase 31 [Amborella trichopoda] |

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**Notes:**

- **a** The trait upon which the candidate gene associates; PC1–PC6 indicate the associations with loadings on the respective PC; and *L. macrospora* and *T. areolata* indicate associations with the presence/absence data of these fungi among the samples.
- **b** GO terms associated with the gene model in the Norway spruce v1.0 genome portal.
- **c** Angiosperm orthologues to the Norway spruce gene models reported in the Norway spruce v1.0 genome portal.
- **d** The best BlastX hit recovered when querying the NCBI nonredundant protein database with the Norway spruce gene model (E-value cut-off E < 1.0E-10).
Botrytis cinerea, P. herbarum and S. polyspora (Hennon, 1990; Horst, 2013; Kaitera, 2013; Pan et al., 2018; Rajala et al., 2013). With the exception of T. areolata, these species have been reported from Norway spruce phyllosphere communities previously (Menkis et al., 2015; Nguyen et al., 2016; Rajala et al., 2013, 2014). The latent pathogens were, however, much more frequent in our study, found in up to 82% of the samples. It is possible that a surface sterilization treatment would have reduced the frequency of these taxa, but S. polyspora, L. macrospora, T. areolata and R. kalkhoffii were relatively highly abundant (File S3). Thus, our interpretation is that these four pathogens had colonized asymptomatic buds, acting as latent pathogens.

The interactions between trees and phyllosphere fungi are diverse and not fully understood. It has been reported that endophytic fungi are more diverse and abundant than pathogens within the phyllosphere community (Rodriguez et al., 2009; Terhonen et al., 2019), something that is reflected also in our study. Endophytic fungi in the phyllosphere have the potential to both enhance and reduce tree growth and fitness through various mechanisms (Rodriguez et al., 2009; Terhonen et al., 2019), one of them being the capacity of phyllosphere pathogens to act as both repressors and enablers of disease (Ridout & Newcombe, 2015).

A recent study on Populus and Melampsora established that the interaction of Melampsora with endophytic disease repressors is governed by hierarchical contingency rules that determine when and where antagonists in the fungal community reduce plant disease severity, but that this interaction is forestalled by genetic variation for Melampsora resistance in Populus (Busby, Crutsinger,
Barbour, & Newcombe, 2019). If this is a general rule in the interaction between phyllosphere endophytes, pathogens and their hosts, identifying host genetic variation that restricts the colonization by pathogenic fungi or covariates with particular phyllosphere endophyte communities may help to increase our understanding of the interaction between trees and their phyllosphere fungal communities. In this study we were successful in identifying nine such QTLs, highlighting various potential mechanisms.

The gene models MA_10428833g0010 and MA_10428833g0020, associated with PC3, appear to either belong to the same gene as indicated by the shared best BLASTX hit, ABR18169.1, or possibly represent a tandem duplication of Ubiquitin E3 ligase genes. Angiosperm orthologues of ABR18169.1 have been shown to be linchpins in the responses to nitrate and phosphate starvation (Park, Seo, & Chua, 2014; Peng et al., 2008) and to regulate plant immune responses (Yao & Iba, 2008). The potential connection to nitrate and phosphate starvation responses makes this a highly interesting locus as the nutrient status of Quercus macrocarpa and Pinus monticola leaves is one of the factors that influence their phyllosphere fungal community (Jumpponen & Jones, 2010; Larkin, Hunt, & Ramsey, 2012). Furthermore, a recent study has shown that an impaired phosphate accumulation capacity in Arabidopsis led to a strong shift in the phyllosphere fungal community (Finkel et al., 2019).

Several of the potential orthologues of the Norway spruce candidate genes associated with PC3 and PC4 indicate that shoot development and morphogenesis are important factors in shaping genotype-specific phyllosphere communities in this study. The poplar orthologue of MA_19950g0010, Potri.014G027200.1, encodes a WD-40 repeat family protein with similarity to ROOT INITIATION DEFECTIVE 3 (RID3). RID3 has been functionally characterized and the RID3 locus controls cell division and shoot apical meristem formation in Arabidopsis (Tamaki et al., 2009), acting as a negative regulator of key genes. In this context it is noteworthy that the expression of MA_19950g0010 is reported in the P. abies exAtlas to be very low during the early phases of Norway spruce bud development just after growth termination when the buds go through a phase of rapid needle initiation and changes in starch and tannin distribution patterns (Hejnówicz & Obarska, 1995). The sister chromatin cohesion proteins PDS5s, the Arabidopsis orthologues of MA_104333886g0010 which is associated with PC3, are involved in mitosis, and depletion of this protein lead to severe effects on development, among other traits (Pradillo et al., 2015). Notably, MA_104333886g0010 is most highly expressed in the early phases of bud flush, which is characterized by high mitotic activity and also by changes in both morphology and starch and tannin distribution (Hejnówicz & Obarska, 1995). Interestingly, the candidate genes associated with PC3 show a concomitant downregulation in vegetative shoots that have just begun to flush (Z4006TR02), compared to earlier and later phases of shoot development; the genes are differentially expressed between samples taken from needles and stem, indicating that the cellular processes the candidate genes are associated with are not ubiquitous in the phyllosphere and that their absence could be important for colonization success for certain fungal taxa. The observations associated with PC3 and PC4 suggest that bud flush, and the changes in distribution of energy and defensive metabolites that the tissue transitions through, may be a critical phase in the colonization of Norway spruce by the studied phyllosphere fungal community and their later occurrence in dormant buds. This is in line with the work by Eusemann et al. (2016) who postulated that tree phenology was one of the drivers between assembly of the phyllosphere community in their study. Phenology (timing of bud burst) is under very strong genetic control in Norway spruce, and it has a narrow sense heritability of approximately 0.8 (Hannerz, Sonesson, & Ekberg, 1999). The function and expression patterns of each of the identified Norway spruce candidate genes, and perhaps the role of phenology, in the interaction between phyllosphere fungi and Norway spruce, will need to be tested in future experiments.

One of the candidate genes associated with the presence/absence of L. macrospora in the communities, MA_10432519g0010, has similarity to the HEC/Ndc80p family proteins. These proteins are part of the kinetochoore complex, which provides an attachment site for spindle fibres in the centromere of chromosomes, and thus are important for cell division (Shin, Jeong, Park, Kim, & Lee, 2018). Disruption of the kinetochoore complex may affect, for example, morphogenesis (Du & Dawe, 2007; Lermontova et al., 2011; Shin et al., 2018). It may be difficult to reconcile a potential role of MA_10432519g0010 in development and morphogenesis with control of L. macrospora colonization as disease symptoms are commonly seen on second-year needles (Butin, 1995). However, it has been suggested that L. macrospora infects flushing tissues in the spring with a latency period of about 1 year until symptoms are visible (Hennon, 1990), which would connect the peak transcriptional activity of the candidate gene with the crucial infection phase.

MA_10g0010, which harbours an SNP associated with the presence of T. aureolata in the phyllosphere fungal community is a previously undescribed member of the class III peroxidase family. The gene model shows an expression pattern strongly associated with differentiating and lignifying tissues in the P. abies exAtlas, particularly with young female cones. Clearly the candidate gene is active in tissues susceptible to the pathogen, and basidiospores of T. aureolata are thought to infect Norway spruce cones and young shoots in the spring (Hietala, Solheim, & Fossdal, 2007; Kuporevich & Transhel, 1957). Thus, it is not far-fetched to imagine that MA_10g0010 may be associated with processes in these tissues that control their vulnerability to colonization, possibly through cell-wall enforcement (Elfstrand et al., 2001; Fagerstedt, Kukkola, Koistinen, Takahashi, & Marjamaa, 2010; Kärkönen, Warinowski, Teeri, Simola, & Fry, 2009; Marjamaa et al., 2006) or processes controlling the timing of bud break and bud flush; overexpression of the class III peroxidase SPI2 in Norway spruce plants led to significant delays of these process compared to in wild-type Norway spruce plants (Clapham, Häggman, Elfstrand, Aronen, & Arnold, 2003; Elfstrand et al., 2001).

Combining molecular identification and barcoding of phyllosphere fungi with GWAS provides insight into host-encoded factors
focusing the assembly of phyllosphere communities and the coloniza-
tion of needle pathogens. Taken together, GWAS results suggest
that processes in the morphogenesis and flush of the Norway spruce
shoot exert a strong influence on the dominant players in the phyllo-
sphere community detected in dormant buds.

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J.S.; execution of the study: M.E., J.B., L.Z. and M.R.G.G.; drafting
manuscript: M.E., J.B. and L.Z.; all authors read and
approved the final version.

DATA AVAILABILITY STATEMENT
ITS7/ITS4 raw read data are stored at the NCBI Sequence Read
Archive in the Bioproject PRJNA590990. Contact M.Rosario.
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REFERENCES
Ahlholm, J. U., Helander, M., Henriksson, J., Metzler, M., & Saikkonen, K.
(2002). Environmental conditions and host genotype direct genetic di-
versity of Venturia ditricha, a fungal endophyte of birch trees. Evolution,
56, 1566–1573. https://doi.org/10.1111/j.0014-3820.2002.tb01468.x
Arnold, A. E., Henk, D. A., Eells, R. L., Lutzon, F., & Vilgalys, R.
(2007). Diversity and phylogenetic affinities of foliar fungal en-
dophytes in loblolly pine infected by culturing and environmen-
tal PCR. Mycologia, 99, 185–206. https://doi.org/10.1080/15572
536.2007.11832578
Baison, J., Vidalis, A., Zhou, L., Chen, Z.-Q., Li, Z., Sillanpää, M. J., …
Garcia-Gil, M. R. (2019). Genome-Wide Association Study (GWAS)
identified novel candidate loci affecting wood formation in Norway
spruce. The Plant Journal, 100, 83–100. https://doi.org/10.1111/
tpj.14429
Bálint, M., Tiffin, P., Hallström, B., O’Hara, R. B., Olson, M. S., Fankhauser,
J. D., … Schmitt, I. (2013). Host genotype shapes the foliar fungal
microbiome of balsam poplar (Populus balsamifera). PLoS ONE, 8,
e53987. https://doi.org/10.1371/journal.pone.0053987
Bernhardsson, C., Vidalis, A., Wang, X., Scofield, D. G., Schiffthaler, B.,
Baison, J., … Ingvarsson, P. K. (2019). An ultra-dense haploid ge-
netic map for evaluating the highly fragmented genome assembly
of Norway spruce (Picea abies). G3: Genes Genomes Genet, 9, 1623–
1632. https://doi.org/10.1534/g3.118.200840
Busby, P. E., Crutsinger, G., Barbour, M., & Newcombe, G. (2019).
Contingency rules for pathogen competition and antagonism in a
genetically based, plant defense hierarchy. Ecology and Evolution, 9,
6860–6868. https://doi.org/10.1002/ece3.5253
Butin, H. H. (1995). Tree diseases and disorders: Causes, biology, and control
in forest and amenity trees. Oxford: Oxford University Press.
Cingolani, P., Platts, A., Wang, L. L., Coon, M., Nguyen, T., Wang, L., …
Ruden, D. M. (2012). A program for annotating and predicting the
effects of single nucleotide polymorphisms, SnpEff. Fly, 6, 80–92.
https://doi.org/10.4161/fly.19695
Clapham, D. H., Hägghman, H., Elfstrand, M., Aronen, T., & von Arnold,
S. (2003). Transformation of Norway spruce (Picea abies) by particle
bombardment. In J. F. Jackson & H. F. Linskens (Eds.), Genetic trans-
formation of plants (pp. 127–146). Berlin: Springer.
Clemmensen, K. E., Bahr, A., Ovaskainen, O., Dahlberg, A., Ekblad, A.,
Wallander, H., … Lindahl, B. D. (2013). Roots and associated fungi
drive long-term carbon sequestration in boreal forest. Science, 339,
1615–1618. https://doi.org/10.1126/science.1231923
Clemmensen, K. E., Ihmark, K., Durling, M. B., & Lindahl, B. D. (2016).
Sample preparation for fungal community analysis by high-through-
put sequencing of barcode amplicons. In F. Martin & S. Uroz (Eds.),
Microbial environmental genomics (MEG) (pp. 61–88). New York:
Springer.
Cordier, T., Robin, C., Capdevielle, X., Desprez-Loustau, M.-L., & Vacher,
C. (2012). Spatial variability of phyllosphere fungal assemblies:
Genetic distance predominates over geographic distance in a
European beech stand (Fagus sylvatica). Fungal Ecology, 5, 509–520.
https://doi.org/10.1016/j.fusco.2011.12.004
Delhomme, N., Sundström, G., Zamani, N., Lantz, H., Lin, Y.-C., Hvidsten,
T. R., … Street, N. R. (2015). Serendipitous meta-transcriptomics:
The fungal community of Norway spruce (Picea abies). PLoS ONE, 10,
e0139080. https://doi.org/10.1371/journal.pone.0139080
DePristo, M. A., Banks, E., Poplin, R., Garimella, K. V., Maguire, J. R.,
Hartl, C., … Daly, M. J. (2011). A framework for variation discovery
and genotyping using next-generation DNA sequencing data. Nature
Genetics, 43, 491–498. https://doi.org/10.1038/ng.806
Du, Y., & Dawe, R. K. (2007). Maize NDC80 is a constitutive feature of
the central kinetochore. Chromosome Research, 15, 767–775. https://
doi.org/10.1007/s10077-007-1160-z
Elfstrand, M., Fossdal, C., Sitbon, F., Olsson, O., Löneborg, A., & von
Arnold, S. (2001). Overexpression of the endogenous peroxidase-like
gene spi 2 in transgenic Norway spruce plants results in increased
total peroxidase activity and reduced growth. Plant Cell Reports,
20, 596–603. https://doi.org/10.1007/s002990100360
Eusemann, P., Schnittler, M., Nilsson, R. H., Jumpponen, A., Dahl, M.
B., Würth, D. G., … Unterseher, M. (2016). Habitat conditions and
phenological tree traits overrule the influence of tree genotype in
the needle mycobiont–Picea glauca system at an arctic treeline ec-
tonite. New Phytologist, 211, 1221–1231. https://doi.org/10.1111/nph.13988
Fagerstedt, K. V., Kukkola, E., Koistinen, V. V. T., Takahashi, J., & Marjamaa,
K. (2010). Cell wall lignin is polymerised by class iii secretable plant
peroxidases in Norway spruce. Plant Cell Reports, 29, 186–194. https://
doi.org/10.1007/s00299-010-0928.x
Farjon, A., & Page, C. N. (1999). Conifers. Status survey and conservation
action plan. Gland, Switzerland: IUCN.
(SPO76) during meiosis and DNA repair in Arabidopsis thaliana. Frontiers Plant Science, 6. https://doi.org/10.3389/fpls.2015.01034
Prunier, J., Verta, J.-P., & MacKay, J. J. (2016). Conifer genomics and adaptation: At the crossroads of genetic diversity and genome function. New Phytologist, 209, 44–62. https://doi.org/10.1111/nph.13565
Rajala, T., Velmala, S. M., Tuomivirta, T., Haapanen, M., Müller, M., & Pennanen, T. (2014). Endophyte communities vary in the needles of Norway spruce clones. Fungal Biology, 117, 182–190. https://doi.org/10.1016/j.funbio.2013.01.006
Rajala, T., Velmala, S. M., Vesala, R., Smolander, A., & Pennanen, T. (2014). The community of needle endophytes reflects the current physiological state of Norway spruce. Fungal Biology, 118, 309–315. https://doi.org/10.1016/j.funbio.2014.01.002
Ridout, M., & Newcombe, G. (2015). The frequency of modification of Dothistroma pine needle blight severity by fungi within the native range. Forest Ecology and Management, 337, 153–160. https://doi.org/10.1016/j.foreco.2014.11.010
Rodriguez, R. J., White, J. F. Jr, Arnold, A. E., & Redman, R. S. (2009). Fungal endophytes: Diversity and functional roles. New Phytologist, 182, 314–330. https://doi.org/10.1111/j.1469-8137.2009.07277.x
Roman-Reyna, V., Pinili, D., Borjaa, F. N., Quibod, I., Groen, S. C., Mulyaningsih, E. S., & Oliva, R. (2019). The rice leaf microbiome has a conserved community structure controlled by complex host-microbe interactions. bioRxiv, 615278
Rosling, A., Cox, F., Cruz-Martinez, K., Ihrmark, K., Grelet, G.-A., Lindahl, B. D., ... James, T. Y. (2011). Archaeorhizomycetes: Unearthing an ancient class of ubiquitous soil fungi. Science, 333, 876–879. https://doi.org/10.1126/science.1206958
Saikkonen, K. (2007). Forest structure and fungal endophytes. Fungal Biology Reviews, 21, 67–74. https://doi.org/10.1016/j.fbr.2007.05.001
Savolainen, O., Pyhäjärvi, T., & Knürr, T. (2007). Gene flow and local adaptation in trees. Annual Review of Ecology, Evolution, and Systematics, 38, 595–619. https://doi.org/10.1146/annurev.ecolsys.38.091206.095646
Seena, S., & Monroy, S. (2016). Preliminary insights into the evolutionary relationships of aquatic hyphomycetes and endophytic fungi. Fungal Ecology, 19, 128–134. https://doi.org/10.1016/j.faneco.2015.07.007
Shin, J., Jeong, G., Park, J.-Y., Kim, H., & Lee, I. (2018). MUN (MERISTEM UNSTRUCTURED), encoding a SPC24 homolog of NDC80 kinetochores complex, affects development through cell division in Arabidopsis thaliana. The Plant Journal, 93, 977–991.
Sieber, T. N. (2007). Endophytic fungi in forest trees: Are they mutualists? Fungal Biology Reviews, 21, 75–89. https://doi.org/10.1016/j.fbr.2007.05.004
Tamaki, H., Konishi, M., Daimon, Y., Aida, M., Tasaka, M., & Sugiyama, M. (2009). Identification of novel meristem factors involved in shoot regeneration through the analysis of temperature-sensitive mutants of Arabidopsis. The Plant Journal, 57, 1027–1039. https://doi.org/10.1111/j.1365-313X.2008.03750.x
Team, R. (2015). RStudio: integrated development for R. Boston: RStudio Inc. Retrieved from http://www.rstudio.com
Terhonen, E., Blumenstein, K., Kovalchuk, A., & Asiegbu, F. O. (2019). Forest tree microbiomes and associated fungal endophytes: Functional roles and impact on forest health. Forests, 10, 42. https://doi.org/10.3390/f10010042
Vidalis, A., Scofield, D. G., Neves, L. G., Bernhardsson, C., Garcia-Gil, M. R., & Ingvarsson, P. K. (2018). Design and evaluation of a large sequence-capture probe set and associated SNPs for diploid and haploid samples of Norway spruce (Picea abies). bioRxiv, https://doi.org/10.1101/291716
Voříšková, J., & Baldrian, P. (2012). Fungal community on decomposing leaf litter undergoes rapid successional changes. The ISME Journal, 7, 477. https://doi.org/10.1038/ismej.2012.116
Wallace, J. G., Kremling, K. A., Kovar, L. L., & Buckler, E. S. (2018). Quantitative genetics of the maize leaf microbiome. Phytobiomes Journal, 2, 208–224. https://doi.org/10.1094/PBIOMES-02-18-0008-R
Yaeno, T., & Iba, K. (2008). BAH1/NLA, a RING-type ubiquitin E3 Ligase, regulates the accumulation of salicylic acid and immune responses to Pseudomonas syringae DC3000. Plant Physiology, 148, 1032–1041.
Zou, H. (2006). The adaptive lasso and its oracle properties. Journal of the American Statistical Association, 101, 1418–1429.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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