High throughput sequencing combined with null model tests reveals specific plant-fungi associations linked to seedling establishment and survival

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Abstract

1. Plant-fungal interactions are important for plant community assembly, but quantifying these relationships remains challenging. High throughput sequencing of fungal communities allows us to identify plant-fungal associations at a high level of resolution, but often fails to provide information on taxonomic and functional assignment of fungi.

2. We transplanted seeds of *Pinus cembra* across an elevational gradient (1850–2250 m a.s.l.) and identified environmental factors and known fungal associates important for seedling establishment and survival. We then applied null model tests to identify taxonomically unassigned fungi associated with pine recruitment.

3. Early seedling establishment was determined by abiotic environmental factors, while seedling survival was predominantly affected by biotic environmental factors (i.e., the abundance of a fungal pathogen known from literature and the distance to adult trees). Null model tests identified known mycorrhizal partners and a large number of unknown operational taxonomic units (OTUs) associated with seedling survival, including saprotrophic and pathogenic species. These results highlight that unknown fungal OTUs, which are usually discarded from analyses, could play a crucial role for plant survival.

4. Synthesis. We conclude that high throughput metabarcoding paired with null model tests, is a valuable approach for identifying hidden plant-fungal associations within large and complex DNA metabarcoding datasets. Such an approach can be an important tool in illuminating the black box of plant-microbe interactions, and thus understanding ecosystem dynamics.

Keywords

DNA metabarcoding, elevational gradient, fungal pathogens, ITS, mycorrhiza, plant regeneration, plant-soil interaction, transplant experiment

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1 | INTRODUCTION

Plant-soil interactions play an important role in plant community assembly (Bever et al., 2010; Klironomos, 2002; Mangan et al., 2010; Wardle, Bardgett, Klironomos, & Seta, 2004). Positive or negative interactions between plants and soil microbiota can accelerate or retard community succession (e.g., Fukami & Wardle, 2005; Wardle et al., 2004), promote species coexistence (e.g., Connell, 1971; Hersh, Vilgalys, & Clark, 2012; Janzen, 1970; Packer & Clay, 2000) and influence the spread of alien plant species (e.g., Chun, Kleunen, & Dawson, 2010; Urcelay, Longo, Geml, Tecco, & Nouhra, 2017). Positive interactions, such as mutualisms between plants and fungi, promote the fitness of both partners (Lambers & Teste, 2013; Peay, 2018; Smith & Read, 2008; Waller, Felten, Hiiesalu, & Vogt-Schilb, 2018). In contrast, negative plant-soil interactions, often associated with the accumulation of fungal pathogens, can have wide-ranging consequences in plant communities, such as preventing the dominance of species (Bever, Mangan, & Alexander, 2015; Malcolm, Kuldau, Gugino, & Jimenez-Gasco, 2013; Tucker & Talbot, 2001). Fungal pathogens can cause seed decay, necrosis of roots and ultimately seedling mortality (Bever et al., 2015; Packer & Clay, 2000). Virulence of the pathogens may depend on environmental factors (Bever et al., 2015; Packer & Clay, 2000), including soil nutrient availability and temperature (Alexander, 2010; Bever et al., 2015). Such effects can be further modified by mycorrhizal fungi, which can provide protection against disease, greater access to soil nutrients and resistance to climate extremes (Jung, Martinez-Medina, Lopez-Raez, & Pozo, 2012). It is also known that different aspects of plant-soil interactions affect different life stages of plants (van der Putten et al., 2013). In the early stages of recruitment, e.g. during germination, plants are especially vulnerable to abiotic factors (e.g. water availability), whilst seedlings may be more affected by the abundance of fungal pathogens and mutualists, as well as other biotic factors (such as competition with neighbouring plants or herbivory; Hersh et al., 2012; Packer & Clay, 2000; van der Heijden & Horton, 2009).

To date, most knowledge of plant-soil interactions has been obtained from lab and pot experiments, which have revealed the relative importance of positive and negative interactions between entire microbial communities associated with a particular plant species or plant community (Kardol, Deyn, Laliberté, Mariotte, & Hawkes, 2013). These experiments often only exhibit the overall net balance of positive and negative interactions (Klironomos, 2002; van der Putten et al., 2013), and the role of individual soil microbial species is not identified. Accordingly, conclusions from these studies remain broad and lack mechanistic detail. Further, experimental settings often over-simplify the complexity of real ecosystems, although it is known that the environmental context affects the outcome of plant-soil interactions (Kardol, Martijn Bezemer, & Putten, 2006; Manning, Morrison, Bonkowski, & Bardgett, 2008; van der Putten et al., 2013, 2016). The quantification of plant-soil interactions during different plant life-history stages across ‘real world’ environmental gradients has been challenging (Kardol et al., 2013). This is mainly due to methodological constraints, including difficulties in characterizing complex microbial communities, and difficulties in understanding the ecological roles of recorded species, given that only about 7% of an estimated 1.5 million fungal species are taxonomically and functionally assigned (Blackwell, 2011; Tedersoo et al., 2014).

High throughput metabarcoding has advanced our understanding of fungal diversity (Tedersoo et al., 2014), distribution (Bahram et al., 2018), community composition (Mucha et al., 2017), ecological roles of members of the community (de Vries et al., 2018) and species interactions (Bogar et al., 2018). This method allows microbial taxa to be grouped into ecologically meaningful molecular operational taxonomic units (OTUs; Taberlet, Coissac, Hajibabaei, & Rieseberg, 2012), and to assign putative ecological functions such as saprotrophs, pathogens or mutualists (Nilsson et al., 2018). However, the identification of fungal diversity relies on curated databases (e.g. UNITE; Koljalg et al., 2005) and specific pipelines (e.g. FUNGuild; Nguyen et al., 2016), and the sparsity of data in these databases severely limits the taxonomic and functional assignment of taxa detected by metabarcoding. As a result, a large fraction of OTU-level diversity is often excluded from downstream ecological analyses (Blackwell, 2011; Tedersoo et al., 2014). For example, in a previous study on the spatial occurrences of plant-associated fungi (Merges, Bâlint, Schmitt, Böhning-Gaese, & Neuschem, 2018) we obtained a dataset of over 1,000 distinct fungal OTUs, indicating a highly diverse fungal community. However, BLAST searches of the acquired fungal OTUs against reference sequence databases (i.e. UNITE and GenBank; Koljalg et al., 2005) allowed us to assign only 58% of the OTUs to genus level. Only these taxa could be included in a literature research to identify candidate OTUs associated with our focal plant species, and the remaining 42% of unassigned OTUs were omitted from further analysis (Merges et al., 2018). This phenomenon has occurred in many similar studies where only a small fraction of the acquired OTUs were included in the final analyses (e.g. Glynou, Nam, Thines, & Macià-Vicente, 2017; Mundra et al., 2015; Schmidt et al., 2017). It is unlikely that the problem of OTU identification due to incomplete databases will disappear soon given the sheer diversity of fungi (Blackwell, 2011; Tedersoo et al., 2014) and our limited biological knowledge of many taxa, especially those which are only known by internal transcribed spacer (ITS) sequences from environmental samples. Nevertheless, some unassigned OTUs could be major players in ecological dynamics (Toju et al., 2017; Toju, Yamamoto, Tanabe, Hayakawa, & Ishii, 2016), and analysing all community members, regardless of taxonomic or functional assignment status, could greatly facilitate our understanding of ecosystem dynamics.

Detecting influential species in a community, so-called keystone taxa, is important for understanding ecosystem function (Banerjee, Schlaeppi, & van der Heijden, 2018) and ecosystem responses to disturbance (Stinson et al., 2006). Different approaches have been proposed to pinpoint species with putative key ecological functions in natural communities including microbial network analysis (Banerjee et al., 2018; Berry & Widder, 2014). Network analysis has been applied to visualize co-occurrences between microbial species and statistically determine ‘keystone’ taxa (i.e. highly connected taxa, with a strong effect on the network structure (Banerjee et al., 2018; Berry...
null model approach, (Gotelli, Ulrich, & Maestre, 2011), which can detect whether the presence, absence or abundance of a ‘species’ is associated with a target variable, such as an ecosystem function or performance variable (Gotelli et al., 2011; Soliveres et al., 2016; Ulrich, 2010; Ulrich, Piwczyński, Maestre, & Gotelli, 2012). Null model randomization tests provide the opportunity to link a functional response to the abundance of species, functional groups or OTUs (Gotelli et al., 2011; Soliveres et al., 2016; Ulrich et al., 2012). Further advantages are that (a) randomization tests are distribution free and well-suited to the non-normal distributions of OTU data; (b) potential species interactions are taken into account by maintaining the community structure during modelling (Gotelli et al., 2011; Ulrich et al., 2012); (c) null model randomization tests have a clear cause-and-effect hypothesis, where species affect the functional response. This underlying hypothesis is advantageous where a directional hypothesis is reasonable, e.g. for investigating effects of species on ecosystem functions (Gotelli et al., 2011; Soliveres et al., 2016). Null model approaches may allow the identification of species that influence other species or ecosystem functions (Gotelli et al., 2011; Soliveres et al., 2016; Ulrich, 2010), but their application to OTU-based microbial community data remains to be tested.

In this study, we used DNA metabarcoding to assess which fungal species were linked to the performance of Swiss stone pine (Pinus cembra) at two life stages, establishment and survival, across the pine’s natural distributional range. The elevational distribution of the pine, ranging from 1,850 to 2,250 m a.s.l., exhibits pronounced changes in both abiotic and biotic conditions, such as microbial community composition (Merges et al., 2018; Neuschulz, Merges, Bollmann, Gugerli, & Böhning-Gaese, 2018; Roll-Hansen, 1989). Seedling establishment of P. cembra is affected by abiotic factors, such as light and water availability (Neuschulz, Mueller, Bollmann, Gugerli, & Böhning-Gaese, 2015), but also by biotic factors, such as competition or plant-soil interactions (Barbeito, Dawes, Rixen, Senn, & Bebi, 2012; Neuschulz et al., 2015). While P. cembra seedlings are thought to need several mycorrhizal species to efficiently acquire nutrients and water, seedlings are also severely affected by needle pests, such as the snow blight fungus Gremmenia infestans (Roll-Hansen, 1989; Smith & Read, 2008).

We hypothesized that seedlings experience different feedback effects from fungal communities during a) seedling establishment (0–4 months) than b) later stage survival (5–15 months). We predicted that initial seedling establishment is primarily vulnerable to abiotic environmental stresses (Kueppers et al., 2016; Walck, Hidayati, Dixon, Thompson, & Poschlod, 2011), while seedling survival will be more strongly affected by the availability of mycorrhizal partners and the presence of known fungal pathogens (Bardgett, Bowman, Kaufmann, & Schmidt, 2005). Furthermore, we predicted that a null model approach could help to identify fungal associates important for seedling establishment and survival that are so far taxonomically unassigned. We tested these hypotheses by conducting seed translocation experiments and DNA metabarcoding of fungal communities along replicated altitudinal gradients that span the entire elevational distribution of P. cembra and reach beyond the tree’s upper distribution limit. We build this present study on that of Merges et al. (2018), which explored the occurrences of a P. cembra-associated subset of taxonomically assigned fungi in relation to environmental factors and adult conspecific trees. First, we used the full DNA metabarcoding dataset containing known plant associates (Merges et al., 2018) and tested the association of known plant associates with seedling establishment and survival of P. cembra. Second, we applied null model randomization tests to (a) reveal whether this approach can detect the same known fungal associates as used in the first analysis and (b) identify previously taxonomically unassigned fungi that are associated with seedling establishment and survival.

2 | MATERIALS AND METHODS

2.1 | Study area and design

We conducted this study in the Central Alps within the core of P. cembra’s distributional range. We selected two elevational gradients near Davos, Switzerland, one in the Fluela valley (46°48′.0.25″N 09°54′.15.38″E) and one in the Sertig valley (46°44′.0.76″N 9°51′.3.5″E). In both valleys, the lowest elevational belts (about 1,850 m a.s.l.) are covered by mixed coniferous forests, mainly comprised of European larch (Larix decidua Mill.) and Norway spruce (Picea abies (L.) H. Karst). Pinus cembra has a unimodal abundance distribution from 1,850 m a.s.l. up to 2,150 m a.s.l., where P. cembra trees (>3 m tall, Harsch, Hulme, McGlone, & Duncan, 2009) form the tree line. Smaller P. cembra individuals can be found up to 2,200 m a.s.l., but none are present at and over 2,250 m a.s.l. (Neuschulz et al., 2018).

2.2 | Seed translocation experiments

To study the effects of plant-fungal associations on plant establishment and survival in an environmental context, we conducted seed translocation experiments in both valleys in 2014 and 2015. We divided each valley into nine elevational belts with 50 m elevational intervals ranging from 1,850 to 2,250 m a.s.l. Following a random stratified sampling design we installed ten seed bags in 2014 and 20 seed bags in 2015 at each elevational belt (Figure 1). Each seed bag was made of 1.5 mm wire-mesh (to prevent loss of seeds) and contained five P. cembra seeds. Seed bags were placed 4 cm deep in the soil and fixed by metal pins. To break dormancy the seeds were placed in a wet clay-sand mixture and exposed to temperature shifts between 5 and 25°C for 22 weeks (simulated seasonal variation, G. Reiss, pers. comm.). Seed bags were distributed over five microhabitat types (a) under ericaceous vegetation cover, (b) close (0.05–1 m) to adult P. cembra individuals, (c) open soil, (d) rocky habitat, (e) microsite covered by snow [i.e. late snow lie areas] at each elevational belt (Figure 1). At 2,250 m a.s.l. (treeless, high-elevational belt) the microhabitat ‘close to P. cembra’ was substituted by matgrass (Nardus stricta...
In total, we installed 180 seed bags in 2014 (i.e. two replicates per microhabitat per elevational belt) and 360 in 2015 (i.e. four replicates per microhabitat per elevational belt). We installed the experiment at the beginning of the growing season end of May and evaluated whether seedlings had established before the end of the growing season at the end of September (Figure S1). We further monitored the survival of seedlings until the end of their second growing season in the following year (Figure S1).

2.3 | Environmental factors

We tested the effects of environmental factors (i.e. light availability, temperature, soil moisture, vegetation cover and distance to conspecific adults) and fungal occurrences on the establishment and survival of *P. cembra* seedlings. We focused on these early life stages, since they are most vulnerable, and thereby the bottleneck of plant regeneration (Vitasse et al., 2012). Light availability was measured as canopy openness above each seed bag with a spherical densitometer. Ericaceous vegetation cover was recorded by estimating the percentage cover of dominant ground flora species: *Loiseleuria procumbens* (L.) Desv., *Vaccinium* spp. L. and *Rhododendron ferrugineum* L. within 1 m² of each seed bag (Braun-Blanquet, 1964). Ericaceous plants compete for resources with *P. cembra* seedlings but also act as nurse plants by ameliorating harsh environmental conditions (Bardgett & Wardle, 2010; Castro, Zamora, Hódar, & Gómez, 2002). Distance to conspecific adults was measured as the distance from each seed bag to the closest adult *P. cembra* by estimation and using a laser range finder (Nikon 800S) for distances over 10 m. The distance to conspecific adults can be an important factor in plant recruitment, as species-specific pathogens and herbivores accumulate close to conspecific adult plants and negatively affect seedling survival (Janzen-Connell Hypothesis; Connell, 1971; Janzen, 1970).

Temperature was measured for 16 of the 30 seed bags per elevational belt per valley using iButton data loggers (Maxim). Soil surface temperatures were recorded every 4 hr over the duration of the study. We calculated the mean of daily maximum temperatures of the hottest three months (MMaxST), since extreme maximum temperatures can induce rapid drying of soils, and thereby induce desiccation of seedlings (Andrus, Harvey, Rodman, Hart, & Veblen, 2018; Kueppers et al., 2016; Tingstad, Olsen, Klanderud, Vandvik, & Ohlson, 2015). We also calculated the mean daily minimum temperatures of the coldest three months (MMinWT), since extreme frost events can lead to high seedling mortality in high-elevation ecosystems (Kueppers et al., 2016; Lenoir, Gégout, Marquet, Ruffray, & Brisse, 2008). We measured soil moisture under dry weather conditions in September by averaging five tensiometer (Theta-Kit version 3) measurements for each seed bag. Water availability is especially important during early establishment, where root systems are barely developed and desiccation can rapidly occur (Kueppers et al., 2016).

2.4 | Soil fungal communities

*Pinus cembra* is obligately mycorrhizal, i.e. it requires an ectomycorrhizal (ECM) mutualist for survival in field conditions (e.g. to acquire soil nutrients; Smith & Read, 2008). In contrast, the presence of a known pathogen, the snow blight fungus (*G. infestans*), severely limits *P. cembra* survival (Barbeito, Brücker, Rixen, & Bebi, 2013; Roll-Hansen, 1989), mostly by infecting needles covered by snow in winter. To assess the impact of these and other fungi on seedling
performance we re-analysed a DNA metabarcoing dataset of the soil fungal communities that have been recorded at each elevational belt (Merges et al., 2018). Occurrence data for pine-associated fungi obtained from this dataset are published in Merges et al. (2018). This previous study focussed on occurrence patterns of pine-associated fungi in relation to environmental factors and their host, but the effect of fungal occurrences on seedling establishment and survival was not addressed (Merges et al., 2018). Soil fungal communities were recorded by collecting soil samples near the eight seed bags planted per elevational belt following the stratified-random sampling design, focusing on the two most distinct microhabitats (i.e. ericaceous vegetation cover and close to adult *P. cembra* individuals, Figure 1; Merges et al., 2018). Although soil microbial communities are known to be very stable across years, they often show high within-year seasonality (Lipson & Schmidt, 2004; Rudolph, Schleuning, & Piepenbring, 2018; Schadt, Martin, Lipson, & Schmidt, 2003). Therefore, soil was sampled twice (May and September 2015) around each seed bag, to account for seasonality, resulting in a total of 288 soil samples (Merges et al., 2018). ECM roots of *P. cembra* were collected to establish a reference database for *P. cembra*‐relevant ECM fungi (Merges et al., 2018). Roots of saplings and adult trees were sampled at 1,850, 2,050 and 2,200 m a. s. l. in both valleys in May 2015, of which 100 ectomycorrhizal root tips per elevation were collected for DNA extraction. DNA extraction, amplification and sequencing are described in Merges et al. (2018). In brief, 300 mg of each root and soil sample were used for DNA extraction (Cubero & Crespo, 2002). During polymerase chain reaction (PCR) the ITS2 region was amplified. An Illumina MiSeq was used for paired-end sequencing (2 × 300 bp) at Fasteris SA, Plan-les-Ouates, Switzerland (Merges et al., 2018). Default options for the Illumina pipeline developed by Bálint, Schmidt, Sharma, Thines, and Schmitt (2014) were applied. Fungi were identified by blasting the OTU representative sequences against UNITE database and GenBank nucleotide database using a 97% similarity threshold (Merges et al., 2018). Rare OTUs from potential erroneous sequences were excluded, following the recommendation of Bokulich et al. (2013).

Where possible, the OTU reads were classified into pathogens and mutualists (Merges et al., 2018). For pathogens, a systematic search of peer-reviewed journal articles was conducted (Merges et al., 2018). For ECM, the *P. cembra* ECM root samples taken at the study site were used to identify a list of candidate ECM OTUs, which resulted in 35 species known to be ECM mutualists (Table S1, Merges et al., 2018). For further details on the classification of pathogens and mutualists of *P. cembra* see Merges et al. (2018).

### 2.5 Statistical analyses

We fitted two models describing the determinants of establishment (i.e. seed germination and seedling establishment within the first growing season) and survival (i.e. survival from the end of the first growing season to the end of the second growing season). First, we used generalized linear mixed models with a binomial error distribution in the `r` package ‘lme4’ (Bates, Maechler, Bolker, & Walker, 2015). The predictor variables were environmental factors (i.e. light availability, temperature, soil moisture, vegetation cover and distance to conspecific adults), the diversity of *P. cembra*‐associated mycorrhiza (OTU antilogarithm of the Shannon diversity) and the abundance of pathogenic fungal OTUs, as identified by literature research (see Merges et al., 2018). Abundance data were calculated at the plot level based on the number of soil cores in which an OTU was detected (i.e. between 0–8, Figure 1). We assume that OTUs with high plot abundances have a higher likelihood to interact with seedlings growing on the plots. Shannon diversity was chosen to represent the frequency of potential positive interactions since *P. cembra* is known to be associated with a large number of mutualists, whereas abundance was chosen for pathogens, since there are only few fungal pathogens reported to be explicitly associated with *P. cembra* (Barbeito et al., 2012; Rainer et al., 2015). We also included plot ID, region and year as random effects to account for spatial and temporal autocorrelation. Observational level random factors were included in the seedling establishment model to account for overdispersion (Albrecht et al., 2015; Bates et al., 2015). We selected the models by adding explanatory variables in a stepwise manner based on a hypothesized ‘hierarchy of controls’ using the Akaike information criterion correcting for small sample sizes (AICc; Burnham & Anderson, 2002) and likelihood ratio deletion tests to find the most parsimonious models explaining either establishment or survival (Figure S2; Table S1; Diaz et al., 2007; Manning et al., 2015). The first step consisted of a model only containing abiotic environmental factors (i.e. maximum summer temperature, minimum winter temperature, mean soil moisture, light availability), which we hypothesized to be the underlying ultimate cause for distribution of all subsequently added explanatory variables in the following steps. In the second step, we added biotic factors, such as vegetation cover and distance to conspecific adults, since these proximate causes are potentially shaping the fungal community variables added in the following step. Finally, the third set of terms was fungal community data: Antilogarithm of Shannon diversity of ectomycorrhizal communities and the abundance of the two pathogenic OTUs (282, 1,198). We calculated marginal and conditional $R^2$ to explore the variance explained by the models (Nakagawa & Schielzeth, 2013). Marginal $R^2$ describes the variance explained by the fixed effects, whereas conditional $R^2$ is the variance explained by both fixed and random effects (Nakagawa & Schielzeth, 2013).

Second, to detect associations of potential mycorrhizal and pathogenic fungal OTUs with seedling establishment and survival that are not described in the existing literature, we applied null model randomization tests on the full OTU table (Gotelli et al., 2011; Soliveres et al., 2016). These tests are based on a null model approach where one linear regression is performed between the response and each given species (Soliveres et al., 2016; Ulrich et al., 2012). The observed regression slope is then compared to 1,000 random permutations of the species’ values and a standardized effect size (SES) is calculated for each species according to: $S_{\text{sim}} = (S_{\text{obs}} - S_{\text{sim}})/SD$; where $S_{\text{obs}}$ is the observed regression slope $S_{\text{sim}}$ is the average of the 1,000 simulated regression slopes and
**RESULTS**

In the seed translocation experiments, 411 (15%) of the 2,700 planted seeds established as seedlings within the first growing season (i.e. the period between May and September in 2014 and 2015; Figure S1). Of these seedling cohorts 68 (17%) survived to the end of the following growing season (i.e. the period from September to September of the following year; Figure S1).

We identified a total of 1,074 OTUs across the whole study (Merges et al., 2018). Of these, we found two pathogenic OTUs (assigned to the snow blight fungus *G. infestans*) in 83 (29%) soil samples (Table S2). In *P. cembra* root samples, we found 35 ECM OTUs and 184 (64%) of the soil samples contained at least one of these (Table S2).

**TABLE 1** Final generalized linear mixed model showing the effect of environmental factors (i.e., temperature, soil moisture, light availability and their interactions on Swiss stone pine (*Pinus cembra*) seedling establishment). Observational level, plot ID, region and sampling season were included as random effects. Significance (*p* < .05) is indicated in bold.

| Variable | Parameter estimate | SE  | ΔAIC<sub>c</sub> | p value<sup>a</sup> |
|----------|--------------------|-----|-----------------|---------------------|
| Intercept | −1.568             | 0.432 |                 | <.001               |
| Mean summer maximum temperature (MMaxST) | −0.384 | 0.111 | −19.3 | <.001 |
| Soil moisture (SM) | 0.257 | 0.089 | −10.4 | <.001 |
| Light availability (LA) | 0.285 | 0.092 | −17.5 | <.001 |
| MMaxST × SM | 0.305 | 0.096 | −7.9 | .002 |
| MMaxST × LA | −0.240 | 0.097 | −3.8 | .017 |
| SM × LA | −0.270 | 0.076 | −10.4 | <.001 |

Note: *N* = 539, explained variance = 6.0% (marginal R<sup>2</sup>)/22% (conditional R<sup>2</sup>). Abbreviations: LA, light availability, MMaxST, mean daily maximum temperature for the hottest three months; SM, soil moisture.

<sup>a</sup>Assessed with a likelihood ratio deletion test.
Seedling establishment was negatively associated with maximum summer temperature and positively associated with light availability and soil moisture (Figure 2a, Table 1). However soil moisture was negatively associated with the establishment of seedlings at sites with high light availability (significant soil moisture x light availability interaction; Figure 2a, Table 1). In contrast, ectomycorrhizal diversity (i.e. antilogarithm of Shannon diversity) and pathogen abundance did not affect the establishment of seedlings, i.e. these variables were not selected for the most parsimonious model (i.e. Table 1). Seedling survival was significantly positively associated with light availability and light availability at long distances to conspecific adults (significant light availability x distance to conspecific adults (Figure 2b, Table 2). In contrast to seedling establishment, survival of seedlings was negatively associated with the abundance of the pathogenic snow blight fungus, *G. infestans*, and the distance to conspecific adults (Figures 2b and 3, Table 2). Survival of seedlings was estimated to be 16% lower when *G. infestans* (OTU 282) was present in one soil core per plot and 69% lower when present in six soil cores per plot (i.e. highest abundance) relative to seedling survival at plots without *G. infestans* (Figure 3).

For seedling establishment, the null model randomization test identified 29 (3%) of the 1,074 OTUs as significantly positively associated with established seedlings and 32 (3%) as negatively associated (Figure 4, Table S2). Here, 82% of the OTUs were unassigned, 11% could be assigned to ectomycorrhizal fungi, 2% to (predominately) animal pathogen-saprotroph taxa, 3% could be assigned as lichenized and 2% as root endophyte. For seedling survival, the null model randomization test showed 296 (28%) of the 1,074 OTUs as significantly positively associated with survival of seedlings and 217 (20%) as negatively associated (Figure 4, Table S2). Here, 87% of the OTUs were unassigned, 10% were ectomycorrhizal fungi, 1% belonged to (predominately) animal pathogen-saprotroph taxa, 1% was assigned as lichenized and 1% as root endophyte. Twenty-two of 35 ECM OTUs (63%) present in our ECM reference database were identified to be significantly associated with the survival of *P. cembra* seedlings. For seedling establishment, the null model randomization test in which environmental factors had been accounted for revealed that, of the 1,074 OTUs, two (0.2%) were positively associated and eight (0.7%) were negatively associated with established seedlings. Here, 80% were unassigned, 10% were assigned as lichenized and 10% as root endophytes. Eight of 35 ECM OTUs (23%), which were present in our ECM reference database, were significantly associated with the survival of *P. cembra* seedlings.

![Figure 3](image-url) **Figure 3** Generalized linear mixed effects model showing the effects of fungal pathogen abundance operational taxonomic unit (OTU 282) (min = 0 soil cores per plot, max = 6 soil cores per plot) on the probability of Swiss stone pine (*Pinus cembra*) seedling survival. Models were fitted with binomial error distribution. Points indicate jittered raw data. Lines show the model fit (p < .01) with standard error added as grey shadow. Predictions for OTU 282 are plotted for the mean values of the other significant explanatory variables included in the model.

### Table 2

| Variable                        | Parameter estimate | SE  | ΔAIC <a> | p value <b> |
|---------------------------------|--------------------|-----|----------|-------------|
| Intercept                       | −2.60              | 0.85|          |             |
| Light availability (LA)         | 1.44               | 0.73| −2.42    | .041        |
| Distance to adult (DA)          | −2.05              | 1.51| −10.24   | <.001       |
| Fungal pathogen abundance (OTU 282) | −0.44              | 0.16| −7.63    | .003        |
| LA x DA                         | 2.55               | 1.41| −2.97    | .026        |

Note: N = 302, explained variance = 49% (marginal R²)/53% (conditional R²).
Abbreviations: LA, light availability; DA, distance to conspecific adult *P. cembra*.

<sup>a</sup>Assessed with a likelihood ratio deletion test.

<sup>b</sup>Significance (p < .05) is indicated in bold.
out of 1,074 OTUs. Tests on data uncorrected for the environment showed 61 significant OTU-establishment relationships, and the tests on data corrected for the environment showed 10 significant OTU-establishment relationships. In contrast, for seedling survival, the number of significant associations with OTUs was eight to ten times higher than what was expected by chance. Tests on uncorrected data showed 513 significant OTU-survival relationships, and the tests on data corrected for the environment showed 432 significant OTU-survival relationships. These results are consistent with our other results in indicating a weak role of soil fungi in establishment, but a strong role in determining survival.

4 | DISCUSSION

We found clear evidence that the very early life stages of juvenile trees (i.e. establishment of seedlings during the first four months after translocation) were determined by abiotic environmental factors, whereas the later stages of recruitment (i.e. survival of seedlings until the age of 15 months) were predominantly affected by biotic environmental factors (i.e., the abundance of a known fungal pathogen and the distance to adult trees). Using null model randomization tests, we revealed patterns of association between unassigned fungi and the establishment and survival of seedlings. This demonstrates the general potential of this method for identifying microbial species involved in the plant-microbe interactions, which drive plant community assembly.

Our results concur with previous studies demonstrating that young seedlings are especially vulnerable to abiotic factors, but that the later stages of recruitment depend more on the biotic environment (van der Heijden & Horton, 2009; Hersh et al., 2012; Packer & Clay, 2000). The early establishment of *P. cembra* was fostered by high light availability and soil moisture and limited by high maximum temperature. These findings are in accordance with a previous study on coniferous subalpine tree species in North America that showed similar positive responses to light availability (Kroiss, Hillerislambers, & D’Amato, 2015). Several studies have shown that warm and dry conditions during seedling establishment limit seedling survival rates (Andrus et al., 2018; Kueppers et al., 2016; Tingstad et al., 2015). Extreme temperature events are often related to summer drought stress, as the soil surface dries out rapidly and establishing seedlings with poorly developed root systems could fail to assimilate sufficient water for physiological processes (Andrus et al., 2018; Brodersen, Germinot, & Smith, 2018). The survival of *P. cembra* seedlings was reduced at sites with a high abundance of the pathogenic snow blight fungus *G. infestans*, and by increasing distance to conspecific adults. Snow blight fungi, such as *G. infestans*, are known to infect pine needles mostly under winter snow cover, where infected needles are killed (Burdon, Wennstrom, Ericson, Muller, & Morton, 1992; Roll-Hansen, 1989). Negative distance dependency of seedling survival is a well-known mechanism in the Janzen-Connell framework (Connell, 1971; Janzen, 1970) in which pathogens and herbivores accumulate close to adult plants, thereby creating negative above- and below-ground feedbacks for conspecific seedlings (Bell, Frecleton, & Lewis, 2006; Liang et al., 2016; Merges et al., 2018; Packer & Clay, 2000). However, in this study there was a decrease of seedling survival with increasing distances to conspecific adults. This could be explained by more favourable conditions for snow blight fungus *G. infestans* above the tree line (i.e. higher density snowpack), harsher abiotic conditions away from adult trees and missing positive interactions with the mycelia of suitable ectomycorrhizal partners.

We could not detect any effects of ground vegetation cover on the establishment and survival of seedlings, although several previous studies found significant relationships between these factors (e.g. Andrus et al., 2018; Bardgett & Wardle, 2010; Castro et al., 2002; Kueppers et al., 2016; Tingstad et al., 2015). Ectomycorrhizal diversity was also not associated with the establishment and survival of seedlings. This could be explained by the fact that mycorrhizal diversity was generally high, with a minimum of 16 ECM species being present at each elevational belt. This high ECM diversity could have provided ample inoculum, as well as a suitable mutualistic partner.
should the species identity of the fungal partner matter. Potential benefits for recruitment of seedlings mediated through only a few suitable mutualistic partners were indicated by the results from the environment-controlled null model approach, where only 23% of the ECM OTUs in our ECM reference database were shown as significantly associated with seedling survival.

The null model approach revealed that fungi, which are currently taxonomically and functionally unassigned, might be associated with the establishment and survival of plant seedlings. Interestingly, when we compared the results of the model that linked known fungal associates with the associations detected by the null model randomization tests, we found evidence from both approaches that early seedling establishment was less affected by microbial interactions. First, the number of fungal OTUs associated with seedlings during establishment was eight to 43 times lower than the number of fungal OTUs associated with the survival of seedlings. Second, the number of OTUs significantly associated with seedling establishment was comparable to or lower than what we had expected by chance, whereas the number of OTUs significantly associated with seedling survival was eight to ten times higher than expected. Such variation in the strength and number of associations between different seedling life stages could be linked to variability in potential benefits gained from mutualists such as mycorrhizal fungi (Bardgett et al., 2005). For example, the very early life stages of large seeded plants like *P. cembra* (i.e. germination and early establishment) are independent of nutrient supply by fungal mutualist, whereas for survival the formation of mycorrhiza is obligate (Smith & Read, 2008). In the second null model randomization test, where we used the residual variance of the models to account for environmental factors, less than one percent of fungal OTUs were significantly associated with establishment and 40% with survival of seedlings. Fungal OTUs filtered out by this analysis may share the same environmental niche as the establishing and surviving seedlings, whereas those retained fungal OTUs may be more intimately linked to the establishment and survival of seedlings.

Apart from the ECM OTUs present in our reference database that were yielded in the null model tests as significantly associated with seedling survival, the majority of fungi identified were not present in the UNITE database (Koljalg et al., 2005) or the fungal community dataset FUNGuild (Nguyen et al., 2016). However, those few fungal species that were assignable belonged mostly to the ecological guild of mycorrhiza, thereby representing a group of fungi known to interact with plants (Table S1). The identification of additional mycorrhizal fungi, so far unknown to be associated with *P. cembra*, supports the validity of the null model randomization tests for identifying plant-fungal associations. Accordingly, our results suggest that a null model approach can be a valuable technique for reducing the complexity of DNA metabarcoding datasets (e.g., the reduction of very large number of OTUs into a far smaller number of candidate OTUs), as it allows the identification of potential plant-fungal associations with OTUs, independent from their functional and taxonomical assignment. Ideally, results of null model randomization tests are confirmed by checking whether at least some of the candidate taxa are known to be associated with the function or plant species of interest, thus demonstrating credibility. We are aware that by linking plot level abundance to seedling performance we might have missed some interactions with fungi that are only found in the rhizosphere (Genney, Anderson, & Alexander, 2006; Lindahl et al., 2007). Nevertheless, we were able to identify the effect of a known pathogen, *G. infestans*, on seedling survival, which illustrates that associations can be detected at this scale. The null model approach suggested here can be an important tool in illuminating the black box of plant-microbe interactions, e.g. by generating a shortlist of microbial species that may be important in driving plant performance and thus should be investigated in further detail. For example, the unassigned OTUs identified here could potentially be located on our field site, cultured and used in more controlled studies of seedling recruitment. Previous studies have demonstrated a scattered distribution of fungi across elevational (Merges et al., 2018; Van Nuland, Bailey, & Schweitzer, 2017) and latitudinal (van der Putten, 2012; Tedersoo et al., 2014) gradients and have predicted consequences for the movement of tree lines through plant seedling survival in response to climate change. Here we provide evidence that the abundance of certain fungal OTUs can be significantly associated with survival rates of plant seedlings. Our findings suggest that if fungi have a lower capacity to disperse and migrate under climate change than plants, potential tree line advances in response to altered climatic conditions could be slowed down or accelerated depending on the balance of pathogens to mutualists and their presence and dispersal beyond the current tree line. Such limitation may be even stronger in Arctic ecosystems, where dispersal limitation of micro-organisms could be far greater due to the greater distances involved.

## 5 | CONCLUSIONS

Identifying microbial species involved in plant-soil interactions is a major challenge in ecology, particularly in field-based studies. Here we present one of the first field studies to identify a subset of functionally distinct members of the soil fungal community that are likely to affect plant establishment and survival. Our results show that the proportion of taxonomically and functionally unknown fungi that are associated with seedling performance is particularly high during early plant establishment, which may hint at previously unknown roles of fungi in seed germination and survival. The results presented here outline future research directions in above- and below-ground species interactions, for instance testing whether individual soil microbial species can accelerate or retard plant community succession (e.g., Fukami & Wardle, 2005; Wardle et al., 2004) or promote species coexistence (e.g., Connell, 1971; Hersh et al., 2012; Janzen, 1970; Packer & Clay, 2000). We demonstrate that DNA metabarcoding coupled with an ecologically relevant classification of ITS sequences and experimental linkage
to plant life stages is a promising approach to unravel potential plant-fungal associations at a previously unattainable resolution. Furthermore, this approach may also be applied to a wide range of other poorly understood soil taxa, which determine plant performance (e.g. oomycetes and bacteria). In the past, incomplete reference databases and limitations in taxonomically assigning fungal OTUs have led researchers to disregard a substantial part of the observed biodiversity. Thus, only a small fraction of plant-fungal interactions are known, and ecologically important relationships remain hidden. Our study reveals that the combination of community barcoding and a null model approach has the potential to overcome some of these constraints. For example, applying our approach in other systems could help to identify ‘keystone’ OTUs with an important role in structuring plant communities over a wide range of habitats and ecosystem types.

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AUTHORS’ CONTRIBUTIONS

D.M., M.B., I.S and E.L.N. conceived and designed the project. D.M. and M.B. collected the data. D.M. performed the analyses with input from M.B., P.M. and E.L.N. D.M., and E.L.N. led the writing of the manuscript. All authors contributed to various drafts and gave final approval for publication.

DATA AVAILABILITY STATEMENT

Data from this paper are deposited in the Dryad Digital Repository: https://doi.org/10.5061/dryad.qh5js47 (Merges, Bálint, Schmitt, Manning, & Neuschulz, 2019).

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