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FungalRoot: global online database of plant mycorrhizal associations

Nadejda A. Soudzilovskaia1†, Stijn Vaessen1*, Milagros Barcelo1*, Jinhong He2, Saleh Rahimlou3†, Kessy Abarenkov3†, Mark C. Brundrett4, Sofia I.F. Gomes1*, Vincent Merckx5,6 and Leho Tedersoo3†

1Environmental Biology Department, Institute of Environmental Sciences, CML, Leiden University, Einsteinweg 2, 2333 CC, Leiden, the Netherlands; 2South China Botanical Garden, Chinese Academy of Sciences, No.723 Xingke Road, Tianhe District, 510650 Guangzhou, China; 3Institute of Ecology and Earth Sciences, University of Tartu, Ulikooli 18, 50090 Tartu, Estonia; 4School of Biological Sciences, Faculty of Science, University of Western Australia, 35 Stirling Highway, Perth, WA 6009, Australia; 5Understanding Evolution Group, Naturalis Biodiversity Center, 2332 AA, Leiden, the Netherlands; 6Department of Evolutionary and Population Biology, Institute for Biodiversity and Ecosystem Dynamics, University of Amsterdam, 1098 XH Amsterdam, the Netherlands

Authors for correspondence:
Nadejda Soudzilovskaia
Tel: +31 (0)71 5277485
Email: n.a.soudzilovskaia@cml.leidenuniv.nl

Leho Tedersoo
Tel +372 737 5651
Email: leho.tedersoo@ut.ee

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Summary

• Testing of ecological, biogeographical and phylogenetic hypotheses of mycorrhizal traits requires a comprehensive reference dataset about plant mycorrhizal associations.
• Here we present a database, FungalRoot, which summarizes publicly available data about vascular plant mycorrhizal type and intensity of root colonization by mycorrhizal fungi, accompanied with rich metadata. We compiled and digitized data about plant mycorrhizal colonization in nine widespread languages.
• The present version of the FungalRoot database contains 36 303 species-by-site observations for 14 870 plant species, tripling the previously available compiled information about plant mycorrhizal associations. Based on these data, we provide a recommended list of genus-level plant mycorrhizal associations, based on the majority of data for species and careful analysis of conflicting data. The majority of ectomycorrhizal and ericoid mycorrhizal plants are trees (92%) and shrubs (85%), respectively. The majority of arbuscular and nonmycorrhizal plant species are herbaceous (50% and 70%, respectively).
• Our publicly available database is a powerful resource for mycorrhizal scientists and ecologists. It features possibilities for dynamic updating and addition of data about plant mycorrhizal associations. The new database will promote research on plant and fungal biogeography and evolution, and on links between above- and belowground biodiversity and ecosystem functioning.

Introduction

Mycorrhizal interactions with fungi represent one of the key innovations of terrestrial plants. A mycorrhiza is a mutualistic association between plant roots and fungi, where plants provide photosynthetically derived carbohydrates to fungi, and fungi deliver nutrients and water to plants and offer protection from abiotic and biotic stress (Smith & Read, 2008). Based on anatomy and partners’ identity, four principal types of mycorrhiza are recognized: arbuscular mycorrhiza (AM), ectomycorrhizal (EcM), ericoid mycorrhiza (ErM) and orchid mycorrhiza (OM) (Brundrett & Tedersoo, 2018). Although most vascular plant species form mycorrhizal symbiosis of only one type, AM-colonized roots are sometimes also co-colonized by EcM and ErM fungi (Smith & Read, 2008; Brundrett & Tedersoo, 2018). Many EcM fungi can form typical orchid mycorrhizal associations with mycoheterotrophic orchids (Imhof et al., 2013). EcM and ErM fungi also may grow endophytically in roots of other plants without forming any mycorrhiza-like structures (Kohout, 2017; Schneider-Maunoury et al., 2020).

Depending on mycorrhizal types and particular species, mycorrhizal fungi may build extensive mycelial networks that sustain nutrient acquisition and promote plant seedling establishment (Leake et al., 2004). Mycorrhizal types differ in their influence on plant nutrition and therefore affect plant carbon allocation strategies (Veresoglou et al., 2012b), litter quality (Cornelissen et al. (2001) cf. Koele et al. (2012), and decomposition (Cornelissen et al., 2001; Elumeeva et al., 2018), biogeochemical cycles (Veresoglou et al., 2012a; Soudzilovskaia et al., 2015a; Averill & Hawkes, 2016; Tedersoo & Bahram, 2019) and plant

*†These authors contributed equally to this work.
Plants also differ in the level of root colonization intensity by mycorrhizal fungi, which may have an effect on the efficiency of nutrient uptake (Karst et al., 2008; Hoeksema et al., 2010; Treseder, 2013) or protection against pathogens (Smith & Read, 2008). Much of the variation in colonization intensity seems to be related not only to plant and fungal identity, but also to seasonality and environmental conditions (Klironomos, 2000; Maltz et al., 2008). Furthermore, data about root colonization by mycorrhizal fungi provides insights into the level of intimacy of the plant–fungal interactions, linked to plant nutrition effectiveness and environmental drivers of plant communities (Soudzilovskaia et al., 2015b). Some plant species – so-called ‘facultatively mycorrhizal plants’ – may develop mycorrhizas in certain conditions but remain nonmycorrhizal in other conditions, depending on nutrient availability and neighbouring plants (Brundrett, 2017).

However, the type and intensity of root colonization by mycorrhizal fungi is unknown for the great majority of vascular plants and, when available, this information is scattered among multiple narrow-focused datasets, most of which cover specific geographical regions or mycorrhizal types. Sources of mycorrhizal records contain multiple errors, which have accumulated and passed on through literature reviews. Many of these errors are derived from alternative interpretations of mycorrhiza and mycorrhizal types, which are especially common in old literature (Brundrett & Tedersoo, 2019; Bueno et al., 2019). Unfortunately, these incorrect observations have been commonly used in traits-based case studies or meta-analyses without critical evaluation of the source reliability, which may have caused errors in interpretation and conclusions (Tedersoo et al., 2019). Furthermore, most data compilations lack geographical information and environmental metadata about the study sites. A substantial part of fundamental mycorrhizal research has been published in languages other than English, German or French, and thus has remained overlooked in reviews and datasets. Finally, virtually none of the existing data compilations distinguish between research focused on all mycorrhizal types detected for a particular plant species and specific mycorrhizal types (ignoring other types that may be present in the specimen, and therewith assigning a particular mycorrhizal type to the plant species under scrutiny, whereas that species might in reality feature double mycorrhizal colonization types).

Here, we present a global database FungalRoot, which accumulates information about plant mycorrhizal status and root colonization intensity. The FungalRoot database was assembled based on local databases and a large number of case studies as well as recent studies published in nine widespread languages. The database enables users to distinguish between reports of a presence of a particular mycorrhizal type, and reports where the plants were checked for all existing mycorrhizal types. In addition, our database provides information about the locality, ecosystem type, soil chemical data and the method of mycorrhizal assessment, enabling users to build more specific, local reference databases. FungalRoot offers possibilities to provide curator and third-party expert opinion regarding data reliability. Based on the current version of the database we provide a genus-level recommendation list for mycorrhizal type assignment of vascular plants, based on data quality, habitat information and phylogenetic relationships (Brundrett & Tedersoo, 2018), Supporting Information Table S2. These data considerably advance the previously available major check list of plant mycorrhizal status (Wang & Qiu, 2006; Akhmetzhanova et al., 2012) both in number of plant taxa considered, as well as in the accuracy of mycorrhizal trait diagnoses, because previous lists potentially contain many unmanaged errors (Brundrett & Tedersoo, 2019). We recommend using the corrected (genus-level) list of the mycorrhizal traits in comparative studies and meta-analyses, in cases where plant mycorrhizal types were not empirically determined, and when species-level information could not be found in the FungalRoot database.

Materials and Methods

Literature compilation

We combined data from 1775 sources of literature, including articles obtained through Google Scholar searches, large compilations of information on mycorrhizal colonization type and intensity in plants (Harley & Harley, 1987; Wang & Qiu, 2006; Akhmetzhanova et al., 2012; Hempel et al., 2013; Soudzilovskaia et al., 2015b; Gerz et al., 2016), and the authors’ personal literature collections, as well as unpublished data. For the Google Scholar search, we used a boolean search for ‘mycorrhiza’ AND ‘colonisation’ AND ‘name of each country’ in English and in other major languages (including German, Chinese, Farsi, French, Indonesian, Portuguese, Russian, Spanish). The articles were downloaded from the Internet, acquired from the authors or ordered using interlibrary loans. We focused mainly on papers with observations on at least five species or > 10 observations for a lower number of species separable by space or specific treatments. Observations in data compilations were traced back to the original references in order to add geographical and ecological metadata, check for methods and avoid transferring errors.

Presence of mycorrhizal status information of a plant species or genus was the minimum requirement to include observations in the database. In cases when the data on root colonization intensity by specific mycorrhizal types were reported, we included these data as well along with the information about a method used for colonization assessment. All collected references were carefully checked for information about geographical location, environmental conditions and habitat (see Supporting Information Table S1 for the lists of included variables and character states).

Data about site soil conditions were added to each record when available. Nitrate (NO$\text{_3}^-$/) or ammonium (NH$_4^+$) values were converted to N based on atomic mass (e.g. X mg NO$_3$/}
kg = X × 14/62 mg N kg⁻¹, as the atomic mass ratio between N and NO₃⁻ is 14:62. Likewise for NH₄⁺ with atomic mass ratio of 14:18 between N and NH₄⁺. Minimum and maximum values, or ranges, were included if available.

Assessment of mycorrhizal types

We followed the mycorrhiza definitions of Brundrett & Tedersoo (2018) that largely relied on Brundrett (2009). Because the associated fungi were rarely identified and their benefits to plants were not assessed in studies addressing mycorrhizal status or level of colonization of natural plants, in most cases we relied solely on the literature-reported morphological criteria (except Australian studies in 1980s and early 1990s that involved synthesis experiments). In brief, the presence of intracellular arbuscules, coils or pelotons was required to consider plants arbuscular mycorrhiza (AM), ericoid mycorrhiza (ErM) or orchid mycorrhiza (OM), respectively. For ectomycorrhizal (EcM), the presence of a Hartig net or a well-developed mantle (one or more hyphal layer) was required. Recently, Bueno et al. (2019) argued that arbuscular colonization is not required for functional AM symbiosis, and presence of other fungal structures, such as vesicles, could be considered as a diagnostic feature. However, the current version the FungalRoot is limited to the data corresponding to the mycorrhizal definitions of Brundrett & Tedersoo (2018). But the database is open to adding new definitions of mycorrhizas, and references based thereon; see below and in the S1 how to add new data to the database.

Misdiaognoses of mycorrhizal types are a common problem in scientific literature (Brundrett, 2017; Tedersoo & Brundrett, 2017; Brundrett & Tedersoo, 2019) and these could lead to errors in analyses based on uncritical use of literature compilations. We considered it important to report the original diagnosis of mycorrhizal type provided in the original publication. Simultaneously, we examined each record in our database against contemporary knowledge of plant species mycorrhizal types (consensus of records in this study; specific information in Merckx, 2013; Kohout, 2017; Tedersoo & Brundrett, 2017; Brundrett & Tedersoo, 2018, 2019). Based on this examination, in the current version of the database, we provide up to three expert opinions commenting on the reliability of the original diagnosis for each contradictory record (see subsection ‘Data records’ below). The dynamic setup of the database allows for the addition of more expert opinions. Based on the database records and the expert opinions, we prepared a recommendation list of mycorrhizal status at the plant genus level (Table S2). To compile this dataset, we considered individual studies of low reliability and excluded these from further comparisons if > 20% of records therein conflicted with other studies. We anticipate, however, that differences in nonmycorrhizal (NM) and AM habit may exist, especially in facultatively mycorrhizal plants and seasonally, depending also on age, environmental conditions and criteria used to assign mycorrhizal types (Brundrett & Tedersoo, 2019; Bueno et al., 2019). The records excluded during the compilation process of the recommendation list were still kept in the database.

Based on individual reports for species, we assigned mycorrhizal type or NM status to the entire genus if > 67% of reports (at least two observations) converged. The value of 67% was chosen because with an expectation of 100% consistency among the reports, the results in nearly all plant genera hosting arbuscular and ectomycorrhizal plants fall into the categories of EcM-AM or AM-NM. Besides the fact that such outcome is trivial, it also is misleading because of a considerable chance for at least some misdiagnosis cases. Therefore, we have opted a threshold of 2/3 of the observations (i.e. 67%) being consistent. The value of 2/3 (67%) was chosen arbitrarily, but is supported by the fact that in cases of detailed laboratory assessments being available for individual plant species, their conclusions are in the majority of cases consistent with a 60–75% of observations presented in the FungalRoot database. In putatively AM and NM groups, there were multiple genera that were reported to be either AM or NM in 33–67% of occasions. These facultatively arbuscular mycorrhizal taxa were encoded as AM-NM. In predominantly AM and EcM plant families, we considered a single positive report sufficient to consider the genus mycorrhizal. If there was a single NM report in these mycorrhizal families, the mycorrhizal status was considered unsettled, rather than prematurely nonmutualistic or unreliable, as recommended by Harley & Harley (1987). For genera that had no reports or single negative reports or two conflicting reports, we further relied on the list of putative NM plant families (Brundrett & Tedersoo, 2018) and EcM plant genera (Tedersoo, 2017), and studies considered unreliable in the first phase. Several aquatic and heterotrophic plant genera in the putatively NM plant families were considered as AM-NM because of multiple independent evidence for arbuscule formation. Given that the assignments of genus-level mycorrhizal type were conducted at 67% level of consistency, there was an obvious possibility for a mismatch in mycorrhizal types of individual species (e.g. Sun et al., 2019). Thus, the data presented in the Table S2 should, as such, never be used to detect misdiagnoses. Rather they should be used in ecological analyses in cases where more detailed information is not available. However, we plead for great caution in the use of these assignments and an adequate accounting for error propagation, when using these data.

According to our data compilation, 86 plant families lack information about mycorrhizal types (Table S3). Following Brundrett & Tedersoo (2018), we considered that Brassicales, Caryophyllales and Cyperales included multiple families with mostly NM or AM-NM species. In Brassicales, we relied partly on the distribution of mycorrhiza-related genes (Delaux et al., 2014). If such information was missing (Brassicales) and for other putatively AM-NM orders, we considered the mycorrhizal status of unstudied families unsettled. For orders that contain only AM genera, we considered unsampled families as putatively AM. We also included specific comments on species of EcM plants that have a larger group of congenic AM species (Ptisonia, Persicaria, Kobresia). For Australian Fabaceae, Goodeniaceae, Myrtaceae and Asteraceae, this was unfeasible because insufficient information. We identified only a single consistently NM plant species Astragalus alpinus within a mycorrhizal genus.
Technical validation

For correction and standardization of the species names included in the database, all observations were checked using the Taxonomic Name Resolution Service (TNRS) (http://tnrs.iplantcollaborative.org/). When partial matches were found, species names were corrected manually according to the best suggestion given by the TNRS. When no suggestions were given, the species name was checked in the original papers. If the species name could still not be corroborated, the record was removed, or, when possible, treated at the level of genus. The species names originally reported in the papers can be obtained from the authors upon request.

Data records

The data are organized into five categories: (1) observation identification; (2) location; (3) soil conditions; (4) host plant description; and (5) description of mycorrhizal colonization (Table S1). The fields for literature references refer to one particular study and include ‘publication_doi’ (for a Digital Object Identifier, DOI, of the citation) and ‘original_reference’ (full text citation in GoogleScholar APA format, necessary for older literature with no DOI or other alphabet). Chinese, Japanese, Persian, Arabic, Cyrillic and other alphabets also conform to this field, although sources in these languages (except Chinese) have been converted or translated into the main format during data management or within previous reviews for simplicity. The field ‘plot_name’ enables segregation of the study into smaller units by location but also by time, treatment or any custom difference. It is represented by the name of the locality or locality-by-treatment combination. All records within a plot have the same geographical coordinates. Identical plot names in different studies are not matched unless their coordinates match.

For the location category, ‘habitat naturality’ enables selection amongst eight possibilities (plus ‘other’ if none conform) that are related to the experimental conditions or habitat structure; ‘biome’ provides information about the overall climate and vegetation type; ‘country’ represents a user-selected field for countries (autonomous and overseas regions separately) following MIMARKS standards; ‘latitude’ and ‘longitude’ represent geographical coordinates, whereas ‘elevation’ represents altitude; ‘collection_date’ indicates date of observation.

In order to enable in-depth meta-analyses, we included 12 fields for soil chemical parameters that are commonly reported in mycorrhiza literature, along with the description of methods for their assessments. The fields ‘pH’, ‘pH_min’, ‘pH_max’, ‘pH_range’ and ‘pH_method’ denote the value and measurement method for determination of soil pH. The fields ‘total_organic_carbon’ and ‘total_organic_carbon_method’ are used to state the value (g/kg soil) and determination method for soil organic carbon content. Likewise, ‘total_nitrogen’, ‘total_extractable_nitrogen’ and ‘total_nitrogen_method’ indicate the value (g kg\(^{-1}\)) and method of determination for total soil nitrogen. The fields ‘total_phosphorus’, ‘total_available_phosphorus’ and ‘total_available_phosphorus_method’ indicate concentration of total phosphorus (mg kg\(^{-1}\) soil; method) or available phosphorus (mg kg\(^{-1}\) soil) and its method of measurement. ‘Potassium’, ‘calcium’ and ‘magnesium’ represent fields for K, Ca and Mg concentrations (mg kg\(^{-1}\) soil; method).

There are three fields for plant species. One of the most important fields is ‘species’, which represents the taxon studied. If no epithet is given, the taxon is identified to genus level. The field ‘number_of_individuals_studied’ represents the sample size of the original study: the total number of individuals of a given species assessed in the given study to establish a mycorrhizal type and/or level of colonization. The ‘host_age’ represents a selectable field of the age of particular individuals, ranging from < 1 month to > 10 yr.

Information about mycorrhizal type and colonization intensity and frequency are given in a suite of 13 fields due to data complexity. The field ‘mycorrhiza_type’ is a selection of 15 options indicating combinations from single mycorrhiza types to dual mycorrhizal colonization, including specification of whether other types have been addressed or not. We find these possibilities important to be considered in scientific analyses, as they allow more reliable negative results to be distinguished from negative reports that may be derived due to the lack of survey for other mycorrhiza types besides the focal AM or EcM. This field also includes suggestions for mycorrhiza-like associations in rootless plants, such as hepatics (levels ‘AM-like’, ‘EcM-like’, ‘ErM-like’ and ‘OrM-like’). The fields ‘AM_intensity’ and ‘AM_frequency’ indicate relative intensity (an estimate of colonization abundance within a root system, determined by one of the methods described in the ‘methods’ field) and frequency (% of plant individuals colonized), respectively. Analogous fields exist for EcM, ErM and OM. The field ‘AM_method’ enables 17 options to indicate the method and scale for determination of AM, whereas ‘EcM_method’, ‘ErM_method’ and ‘OM_method’ offer 10, seven and seven options, respectively.

The FungalRoot database contains six remarks fields. The ‘remark_mycorrhiza_type’ represents notes on reported mycorrhiza type or colonization determination method. Specially dedicated fields enable expert opinions about mycorrhizal type of each observation reported in the database. The fields contain name of the expert and the expert comment, so all expert opinions can be sorted with the data. The downloaded data could be sorted based on presence/absence of expert opinions. The expert opinion fields warn users against possible suspects for mycorrhizal type mis-assignments, which are common in the literature, while allowing the data reported by the original publication to be stored in the database. The current version of the database contains remarks of three experts: Leho Tedersoo, Mark Brundrett and Laura Martinez-Suz. However, the dynamic set-up of our database allows continuous data additions and editing, with a possibility to add new comments by external experts. We welcome scientists to collaborate by providing their expert opinion on the reliability of the database records.

The field ‘other_remarks’ provides additional information about methods, specific experimental treatments and other approaches used in each particular paper. Ecological and evolutionary analyses may be sensitive to such data.
In order to facilitate ease-of-use of the data and to enable efficient updating and versioning, the currently published version of the FungalRoot database is integrated within the online data management and publishing platform PlutoF (Abarenkov et al., 2010b). This structure enables management and editing of multiple fields, custom search by any field, and third-party annotations, such as comments or specification of missing details.

Results

FungalRoot database structure

Our data are freely available for the scientific community, upon citation of this manuscript. They are accessible through the PlutoF platform (https://plutof.ut.ee) for authenticated users, who also have the possibility to add new data and comment the existing data. Additionally the current version of the database is published through GBIF (https://www.gbif.org/dataset/744edc21-8dd2-474e-8a0b-b8c3d56a3c2d). When uploading the data to the PlutoF, the names were converted to the GBIF backbone taxonomy. Each new version of the data published on the PlutoF platform will be linked to the GBIF version of the database. Notes S1 provides detailed information on how to become a user of PlutoF. For versioning of the database we recommend referral to the DOI assigned to the data through GBIF system. All PlutoF platform users can join the PlutoF ‘Mycorrhiza Intensity’ workgroup and link new data records to the ‘Mycorrhiza Intensity project’. We invite experts to add new records to the database and to provide comments on the reliability of existing records, in terms of assignment of mycorrhizal types and other issues. We also invite users to add alternative definitions of mycorrhizal types, provided that records based on alternative definitions will be indicated as such. The database additions submitted will be regularly (e.g. monthly) processed by the PlutoF administrators and added to the database.

For data input, there are two principal ways: (1) using an upload file in a spreadsheet format or (2) direct data insertion over the web platform, which is analogous to the UNITE database system (Abarenkov et al., 2010a). Both the online data insertion and upload file contain the same data fields supplied with specific information. Some fields contain free text, whereas others enable a selection menu to secure consistent terminology. The scientific terminology generally follows MIMARKS standards that were supplemented with more detailed terms (such as mycorrhiza types and specific methods).

Mycorrhizal data

In total, our database contains 36,303 observations for 14,870 plant species. A total of 19,893 observations included in the database are linked to geographical coordinates (Fig. 1). Among the total number of observations, 45% and 2.5% include information about the intensity and frequency of mycorrhizal colonization, respectively. Of mycorrhiza types, studies

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**Fig. 1** Georeferenced records included in the FungalRoot database by colonization: (a) arbuscular mycorrhizal (AM), (b) ectomycorrhizal (EcM), (c) ericoid mycorrhizal (ErM) and (d) nonmycorrhizal (NM). Circle size reflects number of observations per site.
and observations about putatively AM plants prevail, followed by observations on EcM plants and NM plants (Fig. 2a). Among recorded habitats where mycorrhizal plants have been assessed, natural habitats prevail, being mostly forests and grasslands (Fig. 2b,c). Records are unequally distributed among plant species. Only 0.2% of the species had >40 records (Fig. S1). A large number of species (59%) had only one record, whereas 18% and 8% of species had two and three records, respectively.

Observations about mycorrhizal status were distributed unequally globally, with greatest density in North Europe and North America, and lowest density in Africa, Central Asia and Oceania (Fig. 1). This is related directly to historical and present development of mycorrhiza research in different regions. We found literature about mycorrhizal status of plants in nine languages that fitted our criteria for inclusion. Relevant literature in English language clearly dominated, followed by Chinese, Spanish, Portuguese, Russian and French. Among countries, Russia, China, USA and India are best represented (Fig. 2d).

Patterns in the colonization level data are presented in Fig. 3. For the majority of plant species featuring AM infection, the database contains information about AM colonization intensity (Fig. 3a,b). Likewise, for c. 65% of plant species featuring EcM infection and examined for all type of mycorrhiza, and for 49% of plants featuring ErM infection our database contains information about colonization intensity (Fig. 3d,f). Colonization intensity information for plant species featuring both AM and EcM infection, for species featuring EcM infection but not examined for other types of mycorrhiza, and for species featuring OM infection is scarce (Fig. 3c,e,f).

In order to examine how distinct mycorrhizal types are distributed across plant growth forms (trees, herbs, shrubs), we extracted the publicly available data from TRY (https://www.try-db.org/) (Kattge et al., 2011). In this analysis, we considered the mycorrhizal type to correspond to that in the original report. Among obligate AM plant species, 50% are herbaceous, 25% are trees, 10% are shrubs and the remaining 15% plant species are

![Fig. 2 Number of records in the FungalRoot database (a) per most common mycorrhizal type, (b) per habitat type, (c) per major biome type and (d) per country. In (a) the EcM-AM category refers to cases of mixed colonization by the two types of mycorrhizal fungi. The numbers of records for the types ‘ErM-AM’, ‘ErM-EcM’, ‘AM-like (nonvascular plants)’, ‘EcM-like (nonvascular plants)’, ‘ErM-like (nonvascular plants)’ and ‘OM-like (nonvascular plants)’ are 9, 14, 8, 22, 0 and 0, respectively. Owing to the small values these categories are not shown in the graph. In (c) the biome ‘Aquatic’ includes mangroves; ‘Anthrop.’ stands for ‘Anthropogenic’. In (d) the category ‘Former USSR’ refers to the records originated from the dataset of Akhmetzhanova et al. (2012), that are not assigned to Russia, but are assigned to other republics of USSR (now independent countries). Colonization types: AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; ErM, ericoid mycorrhizal; OM, orchid mycorrhizal.](image-url)
Fig. 3 Patterns of the colonization-level data showing number and percentage of species for which information about colonization level is available. (a) AM-only colonization, all other types of mycorrhizal colonization addressed. (b) AM-only colonization, all other types of mycorrhizal colonization not addressed. (c) Double colonization by EcM and AM mycorrhizal fungi. (d) EcM-only colonization, all other types of mycorrhizal colonization addressed. (e) EcM-only colonization, all other types of mycorrhizal colonization not addressed. (f) ErM colonization. (g) OM colonization. Colonization types: AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; ErM, ericoid mycorrhizal; OM, orchid mycorrhizal.
distributed across the other plant growth forms. Among facultatively AM (AM-NM) plants this ratio is 60 : 10 : 10 : 20. The great majority of ectomycorrhizal plants are trees and shrubs (92%) and most of ErM plants are shrubs (85%). Among NM plant species, 70% are herbaceous plants, 10% are trees and 20% belong to other growth forms (Fig. 4).
Finally, we examined the main patterns of variation in intensity of root colonization by mycorrhizal fungi. We calculated the coefficients of variation (CV) of the colonization intensity data for the 10 AM and 10 EcM species that had the highest number of observations of colonization intensity (Table S4). In order to examine whether plants show more variability in colonization intensity levels at the high or lower values of colonization intensity, we examined the CVs of the data points with colonization values of 0–50% and colonization values of 51–100% for each species (Table 1). Arbuscular mycorrhizal plants tend to have higher levels of variation in colonization intensity, although this pattern was only marginally significant ($P=0.07$; Table 1).

Assessment of colonization intensity values within the low (0–50%) and high (51–100%) ranges of colonization intensity revealed that AM plants had higher CVs of colonization intensity than EcM plants ($P=0.04$), only at low colonization intensity range. We also examined whether facultatively mycorrhizal plants had higher levels of variability in colonization intensity than obligately mycorrhizal plants. Due to the paucity of facultatively EcM plants, we conducted this assessment for AM plants only. We considered plants to be obligate AM if our database had >10 records for a plant species and all such records reported AM colonization >0%. For analysis consistency, we selected species that also had >10 records and AM colonization intensity between 0% and 100%. This pre-selection yielded 79 obligate AM species and 67 facultative ones. For each species we calculated the CV of colonization intensity and subjected these CVs to one-way ANOVA. This analysis indicated that facultative AM plant species have much higher variability in colonization level than obligate mycorrhizal (0.73 ± 0.04 vs 0.31 ± 0.02, mean ± SE, $P<0.001$, $F_{1,144} = 77.8$).

### Discussion

The FungalRoot database presented here provides species-by-site information about plant mycorrhizal associations and colonization intensity. Such data are urgently needed given the ultimate importance of such data for understanding functioning of terrestrial ecosystems. We have significantly advanced previous attempts of such data compilations by an exhaustive search for non-English literature, very old (>60 years) and recent literature, which resulted in a tripling of the number of species compared with the previously largest mycorrhizal type checklists (e.g. Wang & Qiu, 2006; Akhmetzhanova et al., 2012, Werner et al., 2018) that all contain records for approx. 3000 plant species (to a large extent overlapping).

The database enables summary of the contemporary information about the distribution of plant species per mycorrhizal type and distribution of mycorrhizal types per growth form. Our data confirm the earlier claims that the majority of mycorrhizal plants are arbuscular mycorrhizal (AM) (70% in our dataset), whereas ectomycorrhizal (EcM) plants, despite being widely distributed (Read, 1991), constitute only a tiny fraction of all vascular plant species (0.7% in our dataset). However, given the fact that our data represent the research efforts in mycorrhizal studies rather than the true distribution of mycorrhizal plant species, these numbers should be treated with caution. Our data suggest that only <5% of all approx. 308 312 vascular plant species (Christenhusz & Byng, 2016) have been examined for mycorrhizal type, with tropical plants being particularly understudied. Thus, further research is needed to obtain a truly quantitative understanding of patterns of mycorrhizal type distributions among vascular plants.

Despite the generally accepted view that the majority of EcM and ericoid mycorrhizal (ErM) plants are shrubs and trees, whereas AM and nonmycorrhizal (NM) type are more or less equally distributed among plant growth forms, quantitative analyses on the distribution of plant mycorrhizal types among growth forms has not been conducted until now. The data shown in Fig. 3 constitute a first attempt at quantitative exploration of thus far available information about mycorrhizal types of plant growth forms. The question of what aspects of plant and mycorrhizal fungal physiology have led to the overwhelming prevalence of woody forms among EcM and ErM plants is particularly intriguing. Further ecophysiological analyses of growth form preferences among plant mycorrhizal types will allow spatial patterns of plant growth form distributions to be linked to mycorrhizal traits. Given that the majority of ecological models of regional and global vegetation distribution, and ecosystem functioning are based on plant growth forms, this information will advance our understanding of the impacts of mycorrhizas on the functioning of terrestrial ecosystems.

By providing detailed information about the intensity of mycorrhizal root colonization of plant species across multiple sites, our database enables advanced analyses of the intriguing patterns of variation in this, so far, poorly understood plant trait. We detected that levels of ectomycorrhizal colonization exhibited lower variability than levels of AM colonization, especially in the lower range of colonization values (Tables 1, S4). Furthermore, our analyses suggest that obligate vs facultative habit of mycorrhizal colonization strongly affects variation in intensity of root mycorrhizal colonization, with
facultatively mycorrhizal plants having twice as large variation.

Erroneous mycorrhizal diagnoses, often provided in old literature, and their blind, uncritical use have resulted in biased or incorrect interpretations of effects of mycorrhizal types on evolutionary, biogeographic and ecophysiological processes. To overcome these issues, we compared the original records with expert opinions derived from the rest of the data and other publications to construct a recommendation list for plant mycorrhizal associations (Table S2). It must, however, be noted that using this list uncritically has the following limitations: (1) it provides insufficient information about individual species and the effect of edaphic and climatic effects on mycorrhizal status; and (2) it may offer erroneous assignments to facultatively mycorrhizal taxa in ecosystems that are early successional, or exhibit extreme levels of nutrients or climatic conditions, such as alpine, flooded or fertilized habitats. In such cases, we recommend considering species-level assignments, provided in the FungalRoot database, accompanied by edaphic data from specific regions or biomes, available as metadata in the FungalRoot database. For species and genera not covered in the FungalRoot database, we strongly recommend in situ determination of mycorrhizal types and mycorrhizal colonization.

In conclusion, the FungalRoot database features a number of unique characteristics, which will enrich the possibilities of scientific research based on the compiled metadata about locality, biome and edaphic conditions of the plant root sampling points. Such data enable quantitative analyses of drivers of mycorrhizal fungal colonization and distribution of mycorrhizal types, which are vital to help understand the impacts of mycorrhizal symbiosis on functioning of the human-affected ecosystems. Furthermore, the database records have been traced to original publications, which enabled us to eliminate duplicated data caused by combining information from multiple compilations. The thorough quality-check of the of mycorrhizal type data in the database, alongside the recommendations for the genus-level mycorrhizal colonization-type assignments (Table S2), have the potential to considerably reduce errors in scientific studies addressing mycorrhizal type effects. Therefore, our database can be readily used to assess the ecophysiological roles of mycorrhizal types in plant communities and ecosystem services and in comparative phylogenetics analyses targeting trait evolution. When coupled to other plant trait, ecological, evolutionary, soil and climate data, the FungalRoot database enables testing large-scale hypotheses about global processes, such as biogeochemical nutrient cycling, climate change impact, and the co-evolution of plants and fungi.

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Author contributions

NAS and LT planned and designed research and wrote the first draft of the manuscript; NS, LH, SV, MB, SR, MCB and JH performed the literature search and assembled the database; MCB, SG and VM contributed the data; LT and MCB enabled the data quality check; KA enabled embedding of the database into the PlutoF work-bench. All authors contributed to the manuscript writing and editing. NAS and LT contributed equally to this work. SV and MB contributed equally to this work, being the main data compilers.

ORCID

Kessy Abarenkov [1] https://orcid.org/0000-0001-5526-4845
Milagros Barcelo [1] https://orcid.org/0000-0003-0454-629X
Mark C. Brundrett [1] https://orcid.org/0000-0002-2501-9037
Sofia I.F. Gomes [1] https://orcid.org/0000-0001-7218-1531
Vincent Merckx [1] https://orcid.org/0000-0002-3959-8623
Saleh Rahimlou [1] https://orcid.org/0000-0003-0427-1329
Nadejda A. Soudzilovskaia [1] https://orcid.org/0000-0002-9584-2109
Leho Tedersoo [1] https://orcid.org/0000-0002-1635-1249
Stijn Vaessen [1] https://orcid.org/0000-0002-4481-6656

References


Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 Plant species that have highest number of records (>40) in the FungalRoot database.

Notes S1 Introduction to the use of PlutoF.

Table S1 Description of FungalRoot database fields.

Table S2 Recommended mycorrhizal status for plant genera, based on ≥67% consistency of species diagnosis for a given mycorrhizal type. It is a large Excel file submitted separately.
Table S3 Families not included in studies of mycorrhizal associations. It is a large Excel file submitted separately.

Table S4 Coefficients of variation (CV) of colonization intensity data for the 10 AM and 10 EcM species that have the highest number of observations of colonization intensity in the Fun- galRoot database.

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