Internal transcribed spacer primer evaluation for vascular plant metabarcoding

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Abstract

The unprecedented ongoing biodiversity decline necessitates scalable means of monitoring in order to fully understand the underlying causes. DNA metabarcoding has the potential to provide a powerful tool for accurate and rapid biodiversity monitoring. Unfortunately, in many cases, a lack of universal standards undermines the widespread application of metabarcoding. One of the most important considerations in metabarcoding of plants, aside from selecting a potent barcode marker, is primer choice. Our study evaluates published ITS primers in silico and in vitro, through mock communities and presents newly designed primers. We were able to show that a large proportion of previously available ITS primers have unfavourable attributes. Our combined results support the recommendation of the introduced primers ITS-3p62plF1 and ITS-4unR1 as the best current universal plant specific ITS2 primer combination. We also found that PCR optimisation, such as the addition of 5% DMSO, is essential to obtain meaningful results in ITS2 metabarcoding. Finally, we conclude that continuous quality assurance is indispensable for reliable metabarcoding results.

Key Words

barcoding, DMSO, internal transcribed spacer, Spermatophyta, mock community, PCR, Tracheophyta

Introduction

Globally, one million species are threatened by extinction in the near future and 68% of monitored populations are declining (IPBES 2019; Grooten et al. 2020). An estimated two out of five plant species are threatened with extinction (Antonelli et al. 2020). Accurate monitoring is vital to understand and alleviate the driving forces behind the unprecedented biodiversity decline (IPBES 2019; Grooten et al. 2020). The term metabarcoding, which describes the analysis of complex DNA samples with the aim of taxonomic identification, has the potential to provide a scale and accuracy in biodiversity surveys that was previously unattainable for many taxonomic groups (Deiner et al. 2017; Ruppert et al. 2019). However, the increase in technical complexity (Piper et al. 2019), compared to most other monitoring methods (Marsh and Trenham 2008; Prosek et al. 2020), also implies a higher susceptibility to errors and therefore requires stringent quality control (Deiner et al. 2017; Ruppert et al. 2019; Thalinger et al. 2020). The objective validation of metabarcoding methods can most efficiently be implemented through mock communities, since the composition of environmental DNA (eDNA) samples is unknown (Bjørnsgaard et al. 2017; Elbrecht and Leese 2017; Smith et al. 2017; Zhang et al. 2018; Braukmann et al. 2019; Thalinger et al. 2020). eDNA studies can be performed either on an amplicon basis (metabarcoding) or on a genome basis (metagenomics). Although genomic methods bypass most PCR biases (Porter and Hajibabaei 2018; Piper et al. 2019), their efficiency currently is not on par with metabarcoding (Braukmann et al. 2019; Ruppert et al. 2019). In metabarcoding studies, the internal transcribed spacer 2 (ITS2) is widely used because it has a high success rate in species-level identification across the plant kingdom (see Kolter and Gemeinholzer 2020 for a detailed discussion of different plant markers). ITS2 also has one of the largest number of reference sequences in public DNA
sequence libraries amongst the most common plant barcode markers (Kolter and Gemeinholzer 2020). Universal criticism, based on the multi-copy nature of ITS2, can be countered by the fact that Song et al. (2012) discovered that 97% of all ITS2 variants in their analysis could only be found within a single species. Song et al. (2012) furthermore reported that intra-genomic distances between variants are smaller than intra-specific or inter-specific distances. Therefore, ITS2 remains to be an important tool in metabarcoding, phylogeny and many other applications (Kay et al. 2006; Feliner and Rosselló 2007; Cheng et al. 2016; Alanagreh et al. 2017; Liu et al. 2019).

Aside from marker choice, primer choice has repeatedly been identified as one of the key factors to facilitate accurate recovery of taxa in a sample (Krehenwinkel et al. 2017; Elbrecht et al. 2019; Hajibabaei et al. 2019; Kelly et al. 2019; Piñol et al. 2019; Li et al. 2020). However, despite the availability of multiple ITS primer sets (White et al. 1990; Gu et al. 2013; Cheng et al. 2016; Moorhouse-Gann et al. 2018; Tremblay et al. 2019), we have identified persistent problems with the amplification success of ITS in multiple recent large-scale barcoding studies (Braukmann et al. 2017; Gill et al. 2019; Jones et al. 2021). The lack of a side-by-side evaluation of current ITS primers makes it impossible to identify and solve the underlying issues of low ITS amplification success rates, compared to other barcode markers.

Our study evaluates ITS primers, based on an in silico and in vitro analysis. The in vitro analysis, performed by using two mock communities, aims to compare the uniformity of amplification achieved with different primers and identify primer-specific amplification biases. The in silico analysis identified mismatches of common ITS primers in Spermatophyta (Cycadopsida, Gnetopsida, Pinopsida, Liliopsida and Magnoliopsida) and led to the design of five new ITS primers with improved universality. Our primer design was focused towards Spermatophyta, as this plant taxon is well represented in public sequence repositories compared to other plant taxa. However, we also reported mismatches in Bryophyta, Fungi, Polyplodiopsida and Lycopodiopsida if adequate taxonomic representation were available. Furthermore, due to the high guanine-cytosine (GC) content in a substantial number of ITS2 sequences, we investigated the impact of dimethyl sulphoxide (DMSO) on mock community species retrieval success.

Material and methods

De novo primer design

Spermatophyta sequences containing the ITS region, used as a template to generate primers, were downloaded from GenBank in April 2018 as described in Kolter and Gemeinholzer (2020). Degenerate consensus ITS sequences on family level of the nrDNA LSU (large subunit of the nuclear ribosomal DNA) and nrDNA SSU (small subunit of the nuclear ribosomal DNA) regions were used to identify conserved flanking regions and, subsequently, suitable primer locations in a single step.

To screen for potential primer sequences in the 5.8S nrDNA region, a consensus sequence of all Spermatophyta plant sequences was established (Suppl. material 1: Suppl. file 1). All nucleotides with more than 2% abundance at a specific position were taken into consideration and represented by the respective IUPAC code. Following this, all 24-mers, extracted from the aforementioned single consensus sequence with a maximum of eight IUPAC ambiguity codes and ending in a C or G, were identified as primer candidates. Primer candidates with hairpin structures with an average melting temperature above 50 °C and those with a GC content below 40% or above 80% were filtered out. Primer candidates forming self-dimers or hetero-dimers with any reverse primer (created in this work), with a Gibbs free energy (AG) higher than one fourth of the maximum Gibbs free energy were discarded or modified. The remaining primer candidates were aligned with 5.8S nrDNA consensus sequences on family level and mismatches were manually resolved by adding a degenerate nucleotide code, if possible. Some mismatches were specific for certain plant families and, therefore, could not be resolved without overly inflating the overall primer degeneracy (Suppl. material 1: Suppl. file 2). The three forward primers have been named in accordance with White et al. (1990) followed by the position (p) within the 5.8S nrDNA region, the specificity (pl = plant, un = universal), the orientation (F = forward, R = reverse) and a revision number (e.g. ITS-3p53pF1). Previously published primers located in the 5.8S nrDNA region played no role in primer positioning or design.

In silico primer evaluation

Primer statistics were calculated using the online tool OligoAnalyzer 3.1 (Owczerzary et al. 2008) with the following settings: monovalent cations (K+) 50 mM, divalent cations (Mg2+) 2.5 mM, dNTP 0.05 mM and primer 0.2 µM. Primer melting temperatures were calculated using the method of Allawi and SantaLucia (1997).

We selected 22 frequently used ITS primers from literature to be analysed alongside the five newly designed primers (Table 1). Primers were validated with Tracheophyta sequences downloaded from GenBank in September 2020 and subsequently processed as described in Kolter and Gemeinholzer (2020). Additional sequence alignment filter steps on family level included the removal of columns with more than 95% gap characters and alignment columns that were supported by less than three species. The sequence coverage of the LSU and SSU region varied from 22,574 to 86,024 (SSU) and from 25,845 to 90,319 (LSU) due to the fact that regions located upstream of the most common ITS primers were less covered in GenBank (Appendix 1). Primer BEL-3, designed by Chiou et al. (2007) and referred to as S3R by
Chen et al. (2010), was not evaluated in silico as its distance to the ITS2 region resulted in very poor sequence availability. To evaluate the specificity towards Fungi and Bryophyta, sequences were extracted from Cheng et al. (2016). The specificity analyses of Polypodiopsida and Lycopodiopsida has been limited to the 5.8S nrDNA region (for a detailed sequence list with taxonomic information, see Suppl. material 1: Suppl. file 1).

Primers were compared to the DNA sequences using the R packages ShortRead and Biostrings (Morgan et al. 2009; Pagès et al. 2020). To give each species the same weight in primer evaluation, the sum of mismatches per sequence has been divided by the number of sequences of that respective species. Adding these numbers up on a family level and dividing them by the total number of species per respective family, resulted in a mismatch score, given in percentages (Suppl. material 1: Suppl. file 3). Only mismatch scores above 30%, per primer position, were reported (for an unfiltered list, see Suppl. material 1: Suppl. file 2). Figures were created using ggplot (Wickham 2016). Higher taxonomic names (above the rank of family) have been retrieved by the R package rgbif from the Global Biodiversity Information Facility (GBIF) backbone taxonomy and are meant to be descriptive only (Chamberlain and Boettiger 2017).

Mock community design

We extracted DNA from 58 herbarium specimen by the use of silica-coated ferric beads and the tissue protocol by Sellers et al. (2018). The two mock communities (mix 1, mix 2) were constructed by considering: 1) DNA concentration, 2) individual amplification success, 3) taxonomic diversity and 4) inclusion of samples with high GC content. We did not use known mismatches as a criterion for inclusion or exclusion. Non-Spermatophyta species were added to investigate primer specificity. Mix 1 (Appendix 4) was created from 23 different Spermatophyta plus four Lycopodiopsida species (Selaginella kraussiana, Selaginella denticulata, Huperzia carinata and Equisetum arvense) and two fungal species (Aspergillus chevalieri and Pseudogymnoasus pannorum). Mix 2 (Appendix 5) was created from 20 different Spermatophyta plant species plus two fungal species (Talaromyces wortmannii and Aureobasidium pullulans). The mixtures contained equal amounts of DNA from each species (1 ng), as determined by Qubit v.4.0 (Invitrogen, dsDNA HS Assay Kit Q32854). This resulted in two mock communities with a taxonomic spread of 16 Spermatophyta plant orders and a difference in GC content of ~25% (Appendices 5 and 6). This intentionally resulted in mock communities with a wide variety of ITS copy numbers per species.

Sequencing primer design

Primers contained a part of the Illumina TruSeq read primer in addition to the target primer to act as a linker between PCR number one (target amplification) and PCR number two (Illumina indexed adapter being added), which results in the following forward primer: 5’-CACACGTGTGCTCTCAGATCT [optional spacer] [target primer]-3’ (reverse: 5’-CTACACGACGTCTTCGCATCT [optional spacer] [target primer]-3’). A spacer that is non-complementary to the target sequence was
added to: 1) prevent more than three identical consecutive nucleotides, 2) stop the TruSeq sequence from interfering with primer binding if it showed the potential to be partially complementary to the target sequence and, 3) to act as a mini-barcode with a length of 3 bp to facilitate pooling of samples with otherwise identical primers after the first round of PCR (termed internal index by Glenn et al. (2019)). The second round of PCR targeted the primer linker added before, which also acts as a part of the read primer in the upcoming sequencing reaction and added the barcoded Illumina indexes (i7 and i5) and the p5 and p7 sequencing adapters (Appendix 3).

**PCR setup**

PCRs were conducted in a 12.5 µl reaction mix containing: 3.125 µl Trehalose (20%), 1.25 µl reaction buffer (10×), 0.625 µl MgCl₂ (50 mM), 1.25 µl DMSO (50%), 0.3 µl bovine serum albumin (BSA) (0.01 mg/ml), 0.25 µl each forward and reverse primer (5 µM), 0.3125 mM dNTP (2 mM), 0.06 µl Platinum Taq (Invitrogen) polymerase (5 U/µl), 1 µl DNA template and 4.0775 µl ddH₂O (modified from Fazekas et al. (2012)).

PCR cycling conditions were 3 minutes at 95 °C for an initial denaturation, followed by 30 cycles (each: 30 sec denaturation at 95 °C, 30 sec annealing at 50 °C, 45 sec elongation at 72 °C) and followed by a final extension at 72 °C for 6 minutes. PCRs were set up in nine technical replicates and three non-template controls (Appendix 8). Five µl of three PCRs with different primer regions were pooled and subsequently treated with Exo I (Thermo Scientific, EN0582). Samples were sent to LGC Genomics GmbH (Berlin, Germany) where they were sequenced on a MiSeq (2x300bp) after an additional 12 PCR cycles (second round of PCR) to add to the indexed Illumina adapter. The second round of PCRs was performed by an initial three cycles at low annealing temperature (each: 15 sec denaturation at 96 °C, 30 sec annealing at 50 °C, 90 sec elongation at 70 °C) and followed up by nine cycles with increased annealing temperature (each: 15 sec denaturation at 96 °C, 30 sec annealing at 58 °C, 90 sec elongation at 70 °C) using MyTaq Red Mix (Bioline, BIO-25044) polymerase. The sequencing library concentrations were adjusted to approximately meet the target of 100,000 total reads per sample which translates to 50,000 paired reads and approximately 16,670 paired reads per PCR (first round of PCR). The sequencing library included all PCR control reactions which totalled approximately 25% of all samples (Suppl. material 1: Suppl. file 4).

**PCR optimisations**

Due to the relatively high GC content of ITS2 amplicons, we optimised PCR conditions in a pre-trial and concluded that the additive DMSO at a concentration of 5% enables amplification of ITS2 from genomic plant templates (Suppl. material 1: Suppl. file 1). To assess the impact of DMSO on mixed PCR templates, like the mock communities used in this study, we compared 0% DMSO to 5% DMSO using the ITS-3p62pF1 + ITS-4unR1 primer combination. All other parameters were identical to the protocols mentioned earlier.

**Sequence data analysis**

Sequencing data was processed with R (Suppl. material 1: Suppl. file 3) and VSEARCH (Rognes et al. 2016; R Core Team 2020). All filtering steps were applied to individual reads instead of read pairs (paired forward and reverse reads) to retain reads in which one of the sequencing directions did not produce any or not enough data. Such reads were granted permission to bypass the merge step if they had a minimum length of 150 bp after primer removal and quality control. In case both reads passed the quality filters and length requirement, but could not be merged, the longer read was retained and the corresponding paired read was discarded. The first step of filtering raw reads was performed by R and included the removal of sequences showing errors in the primer sequence or sequences that were shorter than 30 nucleotides. Subsequent filtering steps by vsearch removed all reads with any undetermined nucleotides (N) and truncated the reads after the continuously accumulated chance of an erroneously assigned nucleotide reached one (maximum expected errors). The merge step allowed only reads to pass (for exceptions, see above) which showed a minimum overlap of 20 nucleotides with a maximum of five differences and which produced a merged sequence with a minimum length of 60 nucleotides. Merged reads were deduplicated by vsearch at 100% identity in a reversible manner and identified by SINTAX (Edgar 2016) using a database by Ankenbrand et al. (2015). Obvious misidentifications (i.e. due to missing reference sequences) were manually corrected. Mock communities were analysed on a genus level.

**Sequence data derived metrics**

To assess the successful detection of taxa in the mock communities, we calculated its read abundance for each taxon and primer combination. We define the read abundance for each taxon in each replicate as the proportion of reads for a given taxon relative to 1000 reads. If the median read abundance of all replicates of one taxon of a specific primer combination was above 0.1 and the taxon was detected (≥ one read) in all replicates of the respective primer combination, the taxon was classified as present (Table 3; Appendices 4 and 5). We set a minimum of more than one read in 10,000 per replicate as the detection threshold as, based on the read depth, this requires a taxon to be not represented by singletons only. We furthermore calculated a retrieved taxa score by including partial detections (taxon not detected in all replicates). The most optimal outcome would be for all taxa within one mock community to be detected in all replicates. The retrieved taxa score expresses how many of those detec-
tions, in relation to the maximum number of possible detections (i.e. mix 1: 21 * 9 replicates), were successful by a specific primer combination.

The required read depth for each primer combination to detect all but one taxon with a confidence of 95% within a mock community was calculated separately for mix 1 and mix 2 in multiple steps. First, instead of assigning an arbitrary penalty score to missing values (taxon not detected in some replicates), we limited the calculation to taxa which could be detected in all replicates in all primer combinations (Appendices 4 and 5). Second, we subsampled the reads of each replicate for each primer combination in increments of 100 reads, each 100 times. If the specific taxon was found (≥ one read) in at least 95 of the 100 subsamples per primer combination (Table 3). If the number of reads of any primer combination were not sufficient to achieve 95% detection probability for some difficult to detect taxa with low read abundances, we linearly interpolated the read abundances to increase the sample size.

Results

In silico primer evaluation

We tested a total of 26 primers (Table 1), located in three distinct regions (SSU, 5.8S rDNA and LSU). Primer melting temperatures of ITS primers analysed in this study ranged from 55.4 °C to 73.0 °C (Table 2). With four exceptions (ITS1, ITS-D, ITS-p3 and 58SPL), the hairpin melting temperature was below the melting temperature of the primer itself (Table 2). Some of the primer variants of primers featuring ambiguous nucleotides (i.e. UniPlantF and ITS-3p53plF1), also form hairpin structures with a melting temperature higher than their respective template in our database (Supp. file 5). We listed the plant families that are potentially negatively impacted by hairpin structures with a melting temperature higher than 50 °C (Suppl. material 1: Suppl. file 6).
GC primer content ranged from 36% to 80% with a 3’ terminal GC stretch of zero to five (Table 2). The UniPlantR primer variant (CCCGCCTGACCTGGGGTCGC) that matches with the most (~72%) plant template sequences in our database, has a GC content of 80% (Suppl. material 1: Suppl. file 5). The primer lengths varied between 19 bp and 24 bp and resulted in a maximum ∆G between -44.32 and -35.49 kcal per mole (Table 2). The self-dimer ∆G ranged from -13.74 to -3.61 kcal per mole (Table 2). The smaller the ratio between maximum ∆G and self-dimer ∆G, the less likely it is that the primer forms troublesome dimers. The number of mononucleotide and dinucleotide repeats ranged from two to four (Table 2). We identified four primer hotspots within the 5.8S nrDNA region (Fig. 1). Their central motifs are approximately located at the positions: 5–20 bp (ITS-D, ITS-p3, ITS-p2, ITS-2plR1), 30–50 bp (ITS3, ITS-u3, ITS-3p34unF1), 60–75 bp (ITS-S2F, UniPlantF, ITS-3p53plF1, ITS-3p62plF1) and 100–108 bp (ITS-u2, 58SPL).

The total amplicon length of each primer combination can be calculated by adding the primer lengths, the distance to the ITS region of interest and the length of the Table 3. Mock community key attributes.

| Primer combination     | Mix 1 (n = 21) | Mix 2 (n = 18) |
|------------------------|----------------|----------------|
|                        | Forward        | Reverse        | Fungi reads (%) | Lycopodiopsida reads (%) | missed taxa (n) | retrieved taxa score (%) | min. required read depth (incl. singletons) | min. required read depth (excl. singletons) | Fungi reads (%) | missed taxa (n) | retrieved taxa score (%) | min. required read depth (incl. singletons) | min. required read depth (excl. singletons) |
| ITS-3p62plF1           | ITS-4unR1      | <1             | 5              | 3                        | 92                     | 2,000           | 3,300                      | 1                             | 3              | 94                     | 1,300                     | 2,000                      |
| UniPlantF              | UniPlantR      | <1             | 0              | 7                        | 81                     | 5,500           | 9,600                      | <1                           | 5              | 81                     | 2,400                     | 3,400                      |
| UniPlantR              | ITS-4unR1      | 41             | 5              | 6                        | 85                     | 3,200           | 5,100                      | 47                           | 5              | 86                     | 2,400                     | 5,600                      |
| ITS-3p62plF1           | UniPlantR      | <1             | 0              | 5                        | 83                     | 5,300           | 9,400                      | <1                           | 4              | 83                     | 1,900                     | 2,700                      |
| 58SPL                  | ITS-4unR1      | 10             | <1             | 6                        | 81                     | 4,800           | 7,600                      | 16                           | 5              | 80                     | 1,300                     | 2,100                      |
| ITS-u3                 | ITS-u4         | 57             | 13             | 6                        | 90                     | 6,600           | 10,800                     | 74                           | 4              | 87                     | 6,600                     | 9,800                      |
| ITS-p4                 | ITS-p4         | <1             | 34             | 6                        | 83                     | 2,400           | 4,000                      | <1                           | 5              | 83                     | 1,800                     | 2,700                      |
| ITS-S2F                | Bel-3          | <1             | 26             | 8                        | 83                     | 6,300           | 11,200                     | <1                           | 3              | 89                     | 5,200                     | 7,600                      |
| ITS3                   | ITS4           | <1             | 6              | 7                        | 75                     | 8,800           | 14,300                     | 87                           | 6              | 77                     | 9,100                     | 14,900                     |

Note: The top three most optimal values in each column were highlighted (bold) until three values were chosen and the next higher or lower value is different from the previous one. Lycopodiopsida read proportions were not highlighted, as it depends on the particular study whether they are negatively (non-target taxon) or positively (target taxon) evaluated. A taxon was defined as missing if the read abundance were lower than 1 in 10000 reads or if it was not represented in all technical PCR replicates. The required read depth only considers taxa that could be fully recovered (present in all replicates) in all primer combinations. Values were rounded, but never to zero, if at least one read could be detected.

Figure 1. Position of previously published and newly introduced ITS primers in the 5.8S nrDNA region (5’ → 3’). Primer positions may be shifted by ± 2 bp in comparison to previously published alignments due to different annotation software being used to identify the 5.8S region. Primers usually used to amplify the ITS1 region (Table 1) have been reverse-complemented to fit the 5’ → 3’ plus strand orientation of the figure. A black arrowhead indicates the direction of amplification of the original primer sequence (Table 1). Primers introduced in this work are marked by an asterisk.

5.8S nrDNA position [bp]

https://mbmg.pensoft.net
ITS marker (Table 1). Most ITS2 sequences have a length between 183 bp (0.5% percentile) and 271 bp (99% percentile), with a median length of 220 bp (Kolter and Gemeinholzer 2020).

**In silico primer evaluation of the SSU region**

Regardless of their position, the primers located in the SSU region, flanking the ITS1 region (ITS1, ITS-A, ITS-u1, ITS-p5), have five or fewer mismatches with the exception of ITS5 (Tables 1 and 2; Suppl. material 1: Suppl. file 2 and Suppl. material 1: Suppl. file 3). However, a stable hairpin structure, with a melting temperature above the annealing temperature, is formed by a 7 bp long self-complementary stretch of the ITS1 primer (Table 2).

**In silico primer evaluation of the 5.8S nrDNA region (reverse)**

The ITS-2plR1 primer displays the lowest number of mismatches (Table 2, Suppl. material 1: Suppl. file 3) in comparison to other primers in the 5.8S region that are generally used as reverse primers for the ITS1 region. The ITS-2plR1 primer is similar to the ITS-p2 primer, the introduction of ambiguous base pairs reduces the number of Spermatophyta families with mismatches from 44 to 8 (Table 2). The remaining eight primer mismatches can be found in the plant families Potamogetonaceae, Apiaceae, Plumbaginaceae, Thismiacae, Siparunaceae, Melanthiaceae, Eriocaulaceae and Juncaceae (Suppl. material 1: Suppl. file 3). The primer with the next lowest number of mismatches, ITS2, with mismatches in 26 families (Table 2), features a stretch of eight self-complementary bases located near the 3' end and, therefore, bears a high risk of producing unwanted by-products.

**In silico primer evaluation of the 5.8S nrDNA region (forward)**

The primers located in the 5.8S nrDNA region that are usually used to serve as forward primers to amplify the ITS2 region can be split into two major groups by the number of Spermatophyta families with mismatches (Table 2). Group one (ITS3, ITS-D, ITS-S2F, ITS-u3 and ITS-p3) does not match in a minimum of 22 families, while group two (UniPlantF, ITS-3p* and 58SPL) has a maximum of 11 mismatched families (Table 2). Most mismatches were found in families outside of Magnoliopsida, like Orchidaceae, Plumbaginaceae, Thismiacae, Siparunaceae and Typhaceae (Suppl. material 1: Suppl. file 3).

**In silico primer evaluation of the LSU region**

Primers located in the LSU region, except for UniPlantR, are overlapping each other at a distance of 35–49 bp to the ITS2 region (Table 1). A modification of the ITS4 primer, ITS-4unR1, displays half the amount of mismatched Spermatophyta families when compared to other primers in the LSU region (Table 2, Suppl. material 1: Suppl. file 3). The remaining mismatches in the families Gnetaceae, Araucariaceae, Cupressaceae, Pinaceae and Junaceae can be eliminated by adding one additional ambiguous nucleotide at 11T>D. (Suppl. material 1: Suppl. file 3).

**Mock community sequence analysis**

The sequencing yielded 3,196,249 paired raw reads (NCBI BioProject PRJNA740294). The filtering step that removed most reads on average (~30%) was to require an exact primer fit at the start of the sequence. A total of 6.5% of all reads failed to merge. As the overall read quality was very high, other filters (combined) removed, on average, less than 5% of the total reads. This results in a total of 2,128,039 merged reads to enter the analysis. There was no contamination detected in the blanks that could have affected the results. Rarefaction curves show a flat slope, except for the primer combination ITS-p3 + ITS-p4, as it yielded less reads than the other primer combinations (Suppl. material 1: Suppl. file 4). All primer combinations are represented by nine replicates, with the exception of ITS-S2F + BEL-3 which is represented by three replicates only. There is a negative Spearman correlation of -0.7 and -0.67 between the GC content and the number of reads recovered for each taxon in mix 1 and mix 2, respectively (\( p < 0.01, \text{df} = 16 \)). This is expected, as species with a relatively high GC content were intentionally included (Appendices 4 and 5). The GC content of taxa ranged from 52% to 77% in mix 1 and from 52% to 79% in mix 2 (Appendices 4 and 5). We recovered no correlation between the number of reads for a specific taxon and the number of mismatches to its respective primer pair. The number of reads from unexpected taxa (reads from species that were not fungi and not included in the mock communities) were negligible with less than 1 in 1,000 reads.

**Mock community primer tests**

*Philodendron angustisectum* could not be detected by any primer combination targeting the ITS2 region in mix 1 and *Sassafras albidum* could not be detected in mix 2, most probably due to DNA degradation. Both will be excluded from the following analysis. The number of missed taxa per mix ranges from 3 to 9 out of 21 for mix 1 and from 3 to 6 out of 18 for mix 2 (Table 3). The number of retrieved taxa is highest with the ITS-3p62plF1 + ITS-4unR1 primer combination and lowest with ITS3 + ITS4 (Table 3). The same pattern in reverse was observed for the required read depth for both analyses, with and without singletons (Table 3). Primer combinations (mix 1 and mix 2) that contain the reverse primer UniPlantR generally recovered very low reads of Liliopsida, especially

https://mbmg.pensoft.net
Gagea graeca, which becomes nearly undetectable with this primer (Appendices 4 and 5).

The 58SPL primer has a similar number of mismatches compared to 3p62plF1 (Table 2). Yet, in direct comparison, we find the key metrics clearly in favour of the 3p62plF1 primer (Table 3). Taking a closer look, the taxa showing a read abundance greater than 200 in the 3p62plF1 + ITS-4unR1 primer combination are increasing in read abundance in the 58SPL + ITS-4unR1 primer combination, while all other taxa, with the exception of Ericales, are decreasing in read abundance (Appendices 4 and 5).

The number of fungi sequences varies from < 1% to 80% of all reads in mix 1 and from < 1% to 88% in mix 2 (Table 3). Abundances are associated with the average number of mismatches in fungi (Tables 2 and 3). The primer combinations UniPlantF + UniPlantR, ITS-3p62plF1 + UniPlantR, ITS-p3 + ITS-p4 and ITS-S2F + BEL-3 are all expected to have three or more mismatches in fungi which resulted in less than 1% of reads to be classified as fungi (Tables 2 and 3). The primer combinations UniPlantF + ITS-4unR1, ITS-u3 + ITS-u4 and ITS3 + ITS4 have, on average, less than one mismatch in fungi and produce fungal read abundances of 41% to 87% (Tables 2 and 3). The Lycopodiopsida in mix 1 are represented by 0 to 345 reads per 1,000 reads (Table 3; Appendices 4 and 5).

Mock community PCR optimisations

Addition of 5% DMSO to the PCR mix has a threefold effect: 1) In most cases, it reduces the number of reads necessary to detect the species with a GC content of ≥ 62% (Fig. 2). 2) It reduces the number of taxa that have at least one failed replicate from nine to three for mix 1 and from six to three in mix 2 (Fig. 2). 3) It enables Lindera (mix 1) and Asimina (mix 2) to be detected (Fig. 2). Taxa like

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**Figure 2.** Impact of DMSO on mock community representation. The detection chance (colour) per genus (black lines) was tracked per replicate (horizontal coloured lines) by subsampling the reads 100 times randomly in steps of 100 from 100 to 10000 (x-axis). The detection chance was defined as the number of subsamples where the respective genus could be detected by at least one read.
Smilax (mix 1) and Quercus (mix 2) that did not show a substantial improvement compared to 0% DMSO, had an overall very low number of reads (Appendices 4 and 5).

Discussion

Selecting and refining: one possible approach to plant metabarcoding studies

Although universal ITS primers have been proposed in the past (White et al. 1990; Blattner 1999; Cheng et al. 2016; Moorhouse-Gann et al. 2018), the increasing availability of publicly available ITS sequences allows for an improvement in universality that was not previously possible. This is demonstrated by comparing the number of sequences (Cycadopsida, Gnetopsida, Pinopsida, Liliopsida and Magnoliopsida) used for primer design by Cheng et al. (2016) and this study (55,700 and 187,522 nrDNA sequences, respectively). Moorhouse-Gann et al. (2018) used a geographically restricted (Mauritius and United Kingdom) database of less than 10,000 nrDNA sequences for primer design.

A plant-specific primer with a low number of mismatches, weak secondary structures and a uniform amplification of a complex DNA mixture has the highest chance to deliver representative results when used in combination with an unknown eDNA sample (Tables 2 and 3). The detailed mismatch lists in this study furthermore allow primers to be customised to fit the exact needs of a given study (Suppl. material 1: Suppl. file 2 and Suppl. material 1: Suppl. file 3). Especially studies including Orchidaceae may find that their unique sequence characteristics warrant extra attention (Suppl. material 1: Suppl. file 3). With the help of the family-level alignments (Suppl. material 1: Suppl. file 3), an orchid-specific primer can be synthesised and added to the degenerate primer mix in a relative quantity of 1/n (n = total number of primer variations of the respective degenerate primer).

In silico primer evaluation

Finding the balance between the elimination of primer mismatches and the number of total primer variants was one of the main design goals of this study. The five primers we generated (ITS-2pIR1, ITS-3p* and ITS-4unR1) achieved less mismatches than most of the previously published ITS primers (Table 2). This results are a significant improvement, as it is rather difficult to predict whether or not a mismatch is critically affecting subsequent amplification or may be of minor importance. The impact of primer mismatches on the overall assay success are determined by parameters that include, but are not limited to, salt concentration, annealing temperature, mismatch position, mispaired nucleotide and template concentrations (Ayyadevara et al. 2000; Waterfall et al. 2002; Sipos et al. 2007; Wu et al. 2009; Lefever et al. 2013; Rejali et al. 2018). Studies by Lefever et al. (2013) show that the impact of (non-3’-terminal) primer mismatches on primer melting temperature is hard to predict, ranging from 0–8 °C for one mismatch and 2–20 °C for two mismatches. Despite the fact that PCR performance has been reported to decrease fundamentally in severity if the mismatch is located more than 8 bp (Rejali et al. 2018) or more than 12 bp (Wu et al. 2009) away from the 3’ terminal position, multiple studies suggest that mismatches towards the 5’ end of primers may not be completely neutral (Sipos et al. 2007; Boyle et al. 2009; Lefever et al. 2013). For these reasons, up-to-date primer development with a minimal number of mismatches is a prerequisite for any successful DNA-based environmental assay, as it increases the number of true positive detections.

Although the study by Cheng et al. (2016) shows a drastic reduction of mismatches of more than 80% in Angiosperms in the ITS-u3 primer versus the ITS3 primer, our results indicate only a reduction of approximately 15% (Table 2). This can be attributed to the way mismatches have been counted. This study counts mismatches and summarises them on a family level, while Cheng et al. (2016) counts mismatches on a sequence-based level. This leads to a scenario where 15% of the included Angiospermb plant families represent 50% of the informative nature of the analysis. We believe that our method of analysis is better suited for a wide breadth of applications, as families with little researched taxa are given the same weight as plant families with a better sequence coverage.

An additional advantage of our method is that it reveals mismatch patterns more easily. If a mismatch occurs consistently in a large number of families and if some of these families are only represented by few sequences, the underlying pattern becomes very hard to catch, if the analysis is based on sequence level only. This can be seen in the ITS-p4 and ITS-u2 primer (Suppl. material 1: Suppl. file 3). The majority of mismatches in these primers could have been eliminated by introducing a single ambiguous nucleotide (ITS-p4: 10G > K and ITS-u2: 10G > R). Making three modifications in the ITS-p3 primer (i.e. 8C>Y, 10G > R, 14C > Y) eliminates most of its mismatches. However, unfortunately, this would introduce a hairpin structure with a melting temperature above the primer melting temperature (Suppl. material 1: Suppl. file 5). In addition to the positioning of the ITS-p3 primer, which adds 140 bp of minimally informative nucleotides to the amplicon length, this limits the potential of the ITS-p3 primer.

The threshold we report of a 30% frequency of mismatch in the corresponding plant family prevents outdated taxonomic assignments (i.e. a genus is placed in the wrong family) and misidentified sequences from introducing noise, which would result in primer design with an unnecessarily high number of ambiguous nucleotides. However, there are rare cases of mismatches of the same nucleotide and at the same primer position that occur at very low frequency within a noticeable number of plant families (Suppl. material 1: Suppl. file 2). One example shown in our data is the UniPlantF and the (partially over-

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and nrDNA copy number variation between 150 and > 100,000 per genome (Prokopowich et al. 2003; Wang et al. 2019) sequencing results of the mock community should be treated qualitatively and not quantitatively. Therefore, we scored the number of missed taxa instead of their relative abundance (read counts) as a measure of primer fitness. The minimally-required read depth provides an indication of how evenly amplification occurred (Table 3). We purposefully did not correct for the differences in nrDNA copy numbers to create mock communities in which some taxa are under-represented by at least one order of magnitude.

Having these aforementioned restrictions in mind, selection of the best suited primers is, in general, the most important tool in minimising additional biases during PCR and library preparation (Schirmer et al. 2015). Although previous studies have focused on optimising existing ITS primers or generating new ones (Morgan et al. 2009; Cheng et al. 2016; Moorhouse-Gann et al. 2018), none of them evaluated primer performance using a mock community. We effectively address this knowledge gap by the first primer efficacy assessment, based on two plant mock communities. The importance of this approach is underlined by the fact that some primers selectively disfavour a certain set of taxa, even without any mismatch being present. Other possible explanations for primer specific taxon bias are secondary structures or stretches with high or low GC content close or at the primer binding site and different binding characteristics of the variants of a primer with ambiguous nucleotides. This may be the case for Gagea graeca (Appendices 4 and 5), as it is nearly absent from sequencing runs using the UniPlantR primer, which has no mismatch in Gagea graeca. In addition to its bias, the mismatches of the UniPlantR primer in other Liliopsida families warrant caution when using this primer in combination with unknown eDNA samples (Suppl. material 1: Suppl. file 3).

One possible explanation for the mixed performance of the 58SPL primer could be that the relatively high primer hairpin melting temperature makes a large proportion of the 58SPL primer unavailable to anneal to its intended template sequence, disfavouring amplicons with a rare prevalence (Table 2). If this is the case, increasing the primer concentration or raising the annealing temperature might alleviate this issue.

The ITS3 + ITS4 primer combination, included in this study, was originally designed to amplify fungi (White et al. 1990). Due to its confirmed high preference towards fungi, as well as the high missed taxa rate, we discourage the use of this primer pair for plant metabarcoding. In spite of this, the ITS3 + ITS4 primer combination still holds its place in recent literature (Gresty et al. 2018; Kamo et al. 2018; Besse 2021; Cámara et al. 2021).

The ITS-u3 + ITS-u4, ITS-p3 + ITS-p4 and ITS-S2F + BEL-3 primer combinations have some performance metrics in their favour (Table 3). Despite this, the relatively high number of plant families with mismatches, as shown by the in silico analysis (Table 2), warrants careful evaluation before using them with eDNA. These results

Mock community primer test

Due to PCR stochasticity (Kebschull and Zador 2015) and nrDNA copy number variation between 150 and
emphasise that, despite the usefulness of mock communities, in silico evaluation can provide valuable additional information on primer universality. While the universality of aforementioned primers could be improved by adding additional ambiguities, the overlap with already existing or already optimised primers, introduced in this paper, indicates that most of them can be replaced with updated versions (Fig. 1).

To our knowledge, this is the first mock community-based primer comparison in the context of metabarcoding in the plant kingdom. Although our mock communities only reflect a fraction of the genetic diversity within plants, we have demonstrated that there are differences between different ITS primer combinations and that these differences are not necessarily based on primer mismatches. The differences not originating from primer mismatches cannot be detected by in silico analyses only, further illustrating the need for mock community studies to verify the results of metabarcoding programmes. In contrast, the universality of the primers can be better assessed by the more comprehensive evaluations made during the in silico analysis. Should the need arise, in essence, we recommend an integrative approach to evaluate primers by combining in silico and mock community analyses. The composition of the mock community should ideally be connected to the respective study area. Considering that we thoroughly screened the whole 5.8S nrDNA region for potential primer sequences, we advise using one of the primers presented in this paper as a starting point for further refinement, if needed.

Mock community PCR optimisations

Varadharajan and Parani (2021) mentions 65% GC content as the threshold for which additives are impacting the PCR success dramatically; similarly, our results indicate 62% GC as the threshold where DMSO starts to improve the sequencing result of a diverse mock community (Fig. 2). In accordance with Varadharajan and Parani (2021), we also show that concentrations below 5% DMSO did not suffice to maximise amplification success (Suppl. material 1: Suppl. file 1). Of the ITS2 plant sequences available in public repositories in 2018, 28.5% had a GC content of 65% or higher (Kolter and Gemeinholzer 2020). The lack of PCR optimisation for high GC regions, as plants with high GC levels are eliminated from the PCR success dramatically; similarly, our results in combination with the UniPlantR primer, offers the shortest possible ITS2 amplicon. However, secondary structures and mismatches could negatively affect PCR efficiency and universality.

The past has shown that ITS-based studies have struggled with amplification success (Braukmann et al. 2017; Gill et al. 2019; Jones et al. 2021). However, this work eliminates the most pressing issues, namely the lack of PCR optimisations and the lack of a comprehensive primer evaluation. In contrast to other plant markers, the ITS region now combines high informative and the potential for high amplification success.

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Ethics statement

The authors declare no conflict of interest.

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Appendix 1

Table A1. Taxonomic breakdown of sequences used for the respective silico primer evaluation.

| nrDNA region | sequences (min/max) | species (min/max) | families (min/max) | order (min/max) |
|--------------|---------------------|------------------|-------------------|----------------|
| SSU          | 22,574 / 63,024     | 15,180 / 35,113  | 95 / 149          | 38 /44         |
| 5.8S*        | 187,522             | 85,362           | 210               | 53             |
| LSU          | 25,845 / 90,319     | 14,663 / 43,795  | 109 / 176         | 42 / 49        |

* partial 5.8S nrDNA sequences filtered prior to analysis.

Appendix 2

Table A2. PCR and sequencing primers used in MiSeq sequencing (5‘→3’).

| location | orientation | unmodified primer name | modified primer name | primer tail | spacer | primer sequence |
|----------|-------------|-------------------------|----------------------|-------------|--------|-----------------|
| SSU      | forward     | ITS3                    | ITS-3T7F1            | CATACGCTGTCCTCCGGACTCT | GCATTGATGAAAGACGCACGCTATCCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | ACCTAGCTGTCGCCGAGCTGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| LSU      | reverse     | UniPlantF               | UniPlantR-T5         | CATACGACGCTGTCCTCCGGACTCT | AGCGTCTGCCTTACCGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| LSU      | reverse     | ITS-3p62F1              | ITS-3T7p62F1         | CATACGACGCTGTCCTCCGGACTCT | ACGTCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| SSU      | forward     | ITS-4u4F1               | ITS-4u4F1            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCCGGACTCT | TACCGCTGTCCTCCGGACTCT |
| SSU      | reverse     | ITS-4u4R1               | ITS-4u4R1            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCCGGACTCT | TACCGCTGTCCTCCGGACTCT |
| LSU      | reverse     | ITS-4u4T5               | ITS-4u4T5            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| LSU      | reverse     | ITS-4u4T5               | ITS-4u4T5            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| LSU      | reverse     | ITS-4u4T5               | ITS-4u4T5            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |
| LSU      | reverse     | ITS-4u4T5               | ITS-4u4T5            | CATACGACGCTGTCCTCCGGACTCT | GCAGTCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT | TACCGCTGTCCTCGCTGATAGATCTGACGACGCTGTCCTCCGGACTCT |

Note: The primer tail is a part of the TruSeq read primers. Throughout the paper primers will be addressed by their unmodified names.

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Appendix 3

Figure A1. Sequencing primer and two-step PCR layout. Note: The template specific primers used in this example are for demonstration purposes only and vary in each unique PCR setup. Additional template (x) strands bending away from the primer sequence in the 1st PCR round demonstrate their non-complementarity.
### Table A3. Mix 1 read abundances and GC values.

| classification | order family | genus species | GC content [%] | read abundances (per 1,000 reads, median of replicates) per primer combination and presence/ absence per replicate |
|----------------|--------------|---------------|----------------|----------------------------------------------------------------------------------------------------------------------------------|
| Liliopsida     |              |               |                |                                                                                                                                  |
| Asparagales    | Asparagaceae | Bellesidia trifoliata | 70 | 3.85 | 0.30 | 0.73 | 0.91 | 1.00 | 1.26 | 1.67 | 0.75 |
| Liliales       | Liliaceae    | Gagea graeca  | 65 | 182 | 0.57 | 71.1 | 0.64 | 43.3 | 61.5 | 169 | 25.3 |
| Poales         | Poaceae      | Brita maxima  | 64 | 0.39 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.09 | 0.00 |
| Poales         | Poaceae      | Tordylium apulum | 56 | 22.5 | 7.85 | 9.32 | 9.09 | 5.18 | 6.06 | 10.8 | 6.05 |
| Poales         | Poaceae      | Arabis verna  | 52 | 163 | 429 | 95.5 | 435 | 166 | 23.1 | 115 | 91.7 |
| Poales         | Poaceae      | Arbutus spec. | 59 | 222 | 413 | 134 | 410 | 305 | 59.0 | 135 | 208 |
| Magnolipsida   | Apiales      | Tordylium apulum | 56 | 22.5 | 7.85 | 9.32 | 9.09 | 5.18 | 6.06 | 10.8 | 6.05 |
| Laurales       | Lauraceae    | Lindera obtusiloba | 77 | 0.07 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| Malpighiales   | Euphorbiaceae | Mercurialis annua | 58 | 46.0 | 8.22 | 27.1 | 7.82 | 34.4 | 12.1 | 23.1 | 16.5 |
| Piperales      | Aristolochiaceae | Aristolochia guichardii | 76 | 0.85 | 0.41 | 0.50 | 0.36 | 0.38 | 0.18 | 0.69 | 0.19 |
| Rosales        | Rosaceae     | Prunus dulcis  | 65 | 9.0 | 11.2 | 3.92 | 12.6 | 3.45 | 3.01 | 8.30 | 5.35 |
| Sapindales     | Anacardiaceae | Pistacia lentiscus | 55 | 32.2 | 57.0 | 18.6 | 58.8 | 34.7 | 6.84 | 19.0 | 12.8 |
| Poales         | Fabaceae     | Anthyllis circinnata | 55 | 232 | 53.9 | 163 | 53.5 | 300 | 60.4 | 136 | 336 |
| Pinopsida      | Pinaceae     | Pinus sylvestris | 59 | 0.22 | 0.54 | 0.10 | 0.63 | 0.08 | 0.29 | 0.16 | 0.00 |

Note: * This GC content is the average of *Equisetum* (73% GC), *Huperzia* (72% GC) and *Selaginella* (57% GC). # Equal parts of *Arbutus unedo* and *Arbutus andrachne*. The ITS-S2F + BEL-3 primer combination yielded only 3 replicates. The ITS-p3 + ITS-p4 primer combination yielded less reads than the other combinations. The highest three read abundance values in each column, including fungi, are highlighted (bold). Absence of a taxon in one or more replicates is represented by a grid pattern where each white cell represents one replicate the respective taxa could not be detected in. *Philodendron angustisectum* could not be detected.

### Appendix 4
### Appendix 5

Table A4. Mix 2 read abundance and GC values.

| classification | order family | genus species | GC content [%] | read abundances (per 1,000 reads, median) per primer combination and presence / absence per replicate |
|----------------|--------------|---------------|----------------|---------------------------------------------------------------------------------|
| **Asparagales** | Asparagaceae  | Maianthemum bifolium | 74 | 0.07 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.07 | 0.00 | 0.00 |
| Liliopsida     | Liliales     | Gagea graeca    | 64 | 183 | 0.62 | 64.3 | 0.92 | 45.6 | 64.4 | 234 | 34.7 | 32.1 |
| Poales         | Poales       | Schoenus nigricans | 70 | 2.48 | 0.16 | 1.52 | 0.16 | 0.49 | 0.46 | 2.70 | 0.38 | 0.23 |
| Apiaceae       | Apiaceae     | Yordanum spicatum | 56 | 28.9 | 9.94 | 10.1 | 10.8 | 6.67 | 7.39 | 19.4 | 8.99 | 3.03 |
| Asteraceae     | Asteraceae   | Geropogon hybridus | 56 | 86.0 | 171 | 40.2 | 173 | 32.6 | 19.2 | 98.2 | 64.8 | 8.99 |
| Brassicas      | Brassicas    | Arabis verna    | 52 | 281 | 560 | 101 | 535 | 210 | 25.2 | 187 | 158 | 18.0 |
| Caryophyllales | Caryophyllales | Silene colorata | 60 | 39.3 | 67.1 | 12.8 | 83.5 | 4.56 | 11.7 | 55.6 | 15.0 | 6.96 |
| Caryophyllales | Polygonacea  | Polygonum arenstrum | 77 | 4.75 | 16.5 | 2.10 | 23.8 | 3.47 | 0.96 | 4.27 | 2.19 | 0.51 |
| Ericaceae      | Ericaceae    | Calluna vulgaris | 62 | 1.32 | 3.94 | 1.04 | 3.70 | 2.15 | 0.33 | 0.71 | 0.38 | 0.14 |
| Fabaceae       | Fabaceae     | Tordylium apulum | 56 | 1.39 | 5.39 | 0.73 | 5.33 | 1.38 | 0.28 | 0.89 | 0.38 | 0.23 |
| Fagales        | Fagaceae     | Quercus ithaburensis | 68 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| Lauraceae      | Lauraceae    | Laurus nobilis | 75 | 0.29 | 0.00 | 0.09 | 0.04 | 0.00 | 0.00 | 0.06 | 0.04 | 0.18 |
| Magnoliopsida  | Magnoliopsida | Asimina triloba | 79 | 0.15 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| Malpighiales   | Malpighiales | Euphorbia annua | 58 | 51.8 | 11.0 | 26.3 | 8.83 | 43.6 | 14.2 | 36.6 | 30.8 | 13.8 |
| Rosales        | Rosaceae     | Prunus dulcis | 65 | 11.0 | 14.9 | 4.00 | 16.7 | 4.45 | 3.09 | 12.4 | 9.70 | 1.32 |
| Solanaceae     | Solanaceae   | Convolvulus tricolor | 58 | 25.7 | 30.3 | 8.60 | 37.8 | 4.29 | 7.06 | 35.3 | 17.2 | 3.84 |
| Pinales        | Pinales      | Pinus strobos | 59 | 0.35 | 0.65 | 0.21 | 0.74 | 0.05 | 0.27 | 0.34 | 0.15 | 0.00 |

**Note:** The ITS-S2F + BEL-3 primer combination yielded only 3 replicates. The ITS-p1 + ITS-p4 primer combination yielded less reads than the other combinations. The highest three read abundance values in each column, excluding fungi, are highlighted (bold). Absence of a taxon in one or more replicates is represented by a grid pattern where each white cell represents one replicate the respective taxa could not be detected in. *Sassafras albidum* could not be detected.
Appendix 6

Table A5. Minimal required read depth for mix 1.

| classification | genus species | GC [%] | ITS-3p62F1 + ITS-4unR1 | UniPlantR | UniPlantF + ITS-4unR1 | ITS-3p62F1 + UniPlantR | UniPlantF + ITS-4unR1 | 58SPL + ITS-u4 + ITS-p4 + BEL-3 | ITS-u3 + ITS-p4 + ITS-5S2F | ITS-u3 + ITS-p4 + BEL-3 | ITS3 + ITS4 |
|----------------|--------------|--------|-------------------------|---------|------------------------|------------------------|------------------------|---------------------------------|---------------------------------|---------------------------------|-----------|
| Liliopsida     | Asparagales  | Asparagus | 70 | 1.3 | NA | 2.8 | 7 | 5 | 5.2 | 3.7 | 2.5 | 6 |
| Liliales       | Liliaceae    | Gagea   | 65 | 0.1 | 7.5 | 0.1 | 6 | 0.2 | 0.1 | 0.1 | 0.2 | 0.2 |
| Liliales       | Smilacaceae  | Smiles  | 71 | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Poales         | Cyperaceae   | Schoenus | 70 | 1.7 | NA | 3.7 | 22.2 | 8.3 | 8.1 | 2.9 | 16.9 | 15.1 |
| Poales         | Poaceae      | Brica   | 64 | 12.2 | NA | 21.7 | NA | NA | NA | 13.1 | 10.5 | 12.6 | NA |
| Apiales        | Apiaceae     | Tordylium | 56 | 0.3 | 0.6 | 0.5 | 0.5 | 0.9 | 0.7 | 0.5 | 0.7 | 1.7 |
| Brassicaceae   | Brassicae    | Arabis  | 52 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.3 |
| Dipsacaceae    | Caprifoliaceae | Valerianella | 72 | 0.3 | 1.5 | 0.6 | 1.2 | 0.4 | 0.8 | 0.4 | 0.2 | 2.2 |
| Eriales        | Eriaceae     | Arabis  | 59 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.2 |
| Eriales        | Eriaceae     | Erica   | 56 | 4.1 | 1.1 | 5.5 | 0.9 | 2.9 | 14.5 | 4.6 | 28.9 | NA |
| Fabales        | Fabaceae     | Anthyllis | 55 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.2 |
| Gentianales    | Rubiaceae    | Sherardia | 62 | 1.5 | 1.1 | 3.7 | 0.8 | 20.7 | 3.4 | 2.1 | NA | 8.7 |
| Laurales       | Lauraceae    | Cinnamomum | 76 | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Malpighiales   | Euphorbiaceae | Mercurialis | 58 | 0.1 | 0.6 | 0.2 | 0.6 | 0.2 | 0.4 | 0.3 | 0.4 | 0.4 |
| Piperales      | Aristolochiaceae | Aristolochia | 76 | 5.5 | 11.2 | 8.1 | 11.9 | 12.4 | 17.7 | 6.6 | 18.9 | 22.4 |
| Rosales        | Rosaceae     | Prunus  | 65 | 0.5 | 0.5 | 1.2 | 0.4 | 1.3 | 1.5 | 0.6 | 0.9 | 3.4 |
| Sapindales     | Anacardiaceae | Pistacia | 55 | 0.2 | 0.1 | 0.3 | 0.1 | 0.2 | 0.7 | 0.3 | 0.4 | 1.8 |
| Solanales      | Solanaceae   | Solanum | 77 | 8.3 | 22.1 | NA | 12.1 | 20.3 | NA | 33.9 | NA |
| Pinaceae       | Pinus         | Pinus   | 59 | 16.4 | 8.4 | NA | 6.6 | NA | NA | NA | NA |

Note: The lowest three values of each row have been highlighted (bold).

Appendix 7

Table A6. Required read depth for mix 2.

| classification | genus species | GC [%] | ITS-3p62F1 + ITS-4unR1 | UniPlantR | UniPlantF + ITS-4unR1 | ITS-3p62F1 + UniPlantR | UniPlantF + ITS-4unR1 | 58SPL + ITS-u4 + ITS-p4 + BEL-3 | ITS-u3 + ITS-p4 + ITS-5S2F | ITS-u3 + ITS-p4 + BEL-3 | ITS3 + ITS4 |
|----------------|--------------|--------|-------------------------|---------|------------------------|------------------------|------------------------|---------------------------------|---------------------------------|---------------------------------|-----------|
| Liliopsida     | Asparagales  | Asparagus | 74 | NA | NA | NA | NA | NA | NA | NA | NA |
| Liliales       | Liliaceae    | Gagea   | 64 | 0.1 | 6.5 | 0.1 | 5 | 0.1 | 0.1 | 0.1 | 0.2 |
| Poales         | Cyperaceae   | Schoenus | 70 | 2 | NA | 2.9 | 22.3 | 9.6 | 9.4 | 1.7 | 11.7 | 16.8 |
| Apiales        | Apiaceae     | Tordylium | 56 | 0.2 | 0.5 | 0.5 | 0.5 | 0.8 | 0.6 | 0.3 | 0.5 | 1.4 |
| Brassicaceae   | Brassicae    | Arabis  | 56 | 0.1 | 0.1 | 0.1 | 0.1 | 0.2 | 0.3 | 0.1 | 0.1 | 0.5 |
| Dipsacaceae    | Caprifoliaceae | Geropogon | 56 | 0.1 | 0.1 | 0.1 | 0.1 | 0.2 | 0.3 | 0.1 | 0.1 | 0.3 |
| Eriales        | Eriaceae     | Silene  | 60 | 0.2 | 0.1 | 0.4 | 0.1 | 1.1 | 0.4 | 0.1 | 0.4 | 0.7 |
| Eriales        | Eriaceae     | Caltha  | 62 | 3.3 | 1.2 | 4.4 | 1.2 | 2.2 | 13.9 | 7.1 | 11.4 | NA |
| Fabales        | Fabaceae     | Anthyllis | 55 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 | 0.2 |
| Fabales        | Fabaceae     | Quercus | 68 | NA | NA | NA | NA | NA | NA | NA | NA |
| Laurales       | Lauraceae    | Laurus  | 75 | 19 | NA | NA | NA | NA | NA | NA | 25.8 |
| Magnoliales    | Annonaceae   | Asimina | 79 | NA | NA | NA | NA | NA | NA | NA | NA |
| Malpighiales   | Euphorbiaceae | Mercurialis | 58 | 0.1 | 0.5 | 0.2 | 0.5 | 0.1 | 0.4 | 0.2 | 0.2 | 0.4 |
| Rosales        | Rosaceae     | Prunus  | 65 | 0.5 | 0.4 | 1.2 | 0.3 | 1.1 | 1.5 | 0.4 | 0.5 | 3.5 |
| Solanales      | Convolvalaceae | Convolvulus | 58 | 0.2 | 0.2 | 0.6 | 0.2 | 1.1 | 0.6 | 0.2 | 0.3 | 1.3 |
| Pinaceae       | Pinus         | Pinus   | 59 | 16.2 | 6.8 | NA | 6.1 | NA | 15.7 | NA | 31.2 | NA |

Note: The lowest three values of each row have been highlighted (bold).

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Appendix 8

Figure A2. Experimental workflow and study design.
Supplementary material 1

Supplementary files
Author: Andreas Kolter, Birgit Gemeinholzer
Data type: zip. archiv

Explanation note: Suppl. file 1. Lists a taxonomic breakdown of the in silico primer mismatch testing. It furthermore contains the sequence files used for primer evaluation and the DMSO trial experiment figure. Suppl. file 2. Lists the result of the primer in silico testing and visualises them without any thresholds. Suppl. file 3. Contains family level alignments of the SSU, LSU and 5.8S nrDNA regions. It also contains the custom R scripts used for in silico testing, as well as mismatch figures on family level with a threshold of 30% in various formats. Suppl. file 4. Contains read number information and rarefaction curves of the mock communities. It also contains a graphical representation of the detection chance per genus per read number. Suppl. file 5. Contains information about the primers with ambiguities used in this study and a breakdown to all possible primer variants.

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