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Improving ITS sequence data for identification of plant pathogenic fungi

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Summary Plant pathogenic fungi are a large and diverse assemblage of eukaryotes with substantial impacts on natural ecosystems and human endeavours. These taxa often have complex and poorly understood life cycles, lack observable, discriminatory morphological characters, and may not be amenable to *in vitro* culturing. As a result, species identification is frequently difficult. Molecular (DNA sequence) data have emerged as crucial information for the taxonomic identification of plant pathogenic fungi, with the nuclear ribosomal

internal transcribed spacer (ITS) region being the most popular marker. However, international nucleotide sequence databases are accumulating numerous sequences of compromised or low-resolution taxonomic annotations and substandard technical quality, making their use in the molecular identification of plant pathogenic fungi problematic. Here we report on a concerted effort to identify high-quality reference sequences for various plant pathogenic fungi and to re-annotate incorrectly or insufficiently annotated public ITS sequences from these fungal lineages. A third objective was to enrich the sequences with geographical and ecological metadata. The results – a total of 31,954 changes – are incorporated in and made available through the UNITE database for molecular identification of fungi (<http://unite.ut.ee>), including standalone FASTA files of sequence data for local BLAST searches, use in the next-generation sequencing analysis platforms QIIME and mothur, and related applications. The present initiative is just a beginning to cover the wide spectrum of plant pathogenic fungi, and we invite all researchers with pertinent expertise to join the annotation effort.

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Introduction

Plant pathogenic fungi are a large assemblage distributed across the fungal tree of life (Stajich et al. 2009). They share a nutritional strategy that adversely affects their plant hosts, sometimes in ways that have negative repercussions for human activities. Precise knowledge of the identity of the causal agent(s) of any given plant disease is the first step toward meaningful countermeasures and disease surveillance (Rossman and Palm-Hernández 2008; Kowalski and Holdenrieder 2009; Fisher et al. 2012). In addition, recent reports of emerging plant pathogens and their cross-kingdom infections to animals and immunocompromised humans accentuate the need for accurate and quick identification in potential outbreaks (Cunha et al. 2013; Gauthier and Keller 2013; Samerpitak et al. 2014). However, it is not always easy to identify plant pathogenic fungi to the species level, as they often lack discriminatory morphological characters or cultivable life stages (Kang et al. 2010; Udayanga et al. 2012). Molecular (DNA sequence) data have emerged as a key resource in the identification of plant pathogenic fungi and carry the benefit that all fungi, regardless of life stage, morphological plasticity, and degree of cultivability, can be analyzed (Shenoy et al. 2007; Sharma et al. 2013). As a result, recent years have seen substantial progress towards a comprehensive understanding of phytopathogenic fungi in terms of taxonomy, systematics, and ecology (Dean et al. 2012; Maharachchikumbura et al. 2012; Manamgoda et al. 2012; Woudenberg et al. 2013).

DNA data, however, are not a panacea for species identification. On the contrary, taxonomically and technically compromised DNA sequences are common in the international nucleotide sequence databases (Bidartondo et al. 2008; Kang et al. 2010). This makes their use as reference data for molecular species identification difficult, particularly because many users of newly generated sequence data may not be in a position to assess whether a proposed taxonomic affiliation is reliable. As a consequence, errors and mistakes propagate over time as users adopt incorrect species names and ecological properties retrieved from sequence similarity searches (Ko Ko et al. 2011; Nilsson et al. 2012). This is especially problematic for phytopathogens, where even closely related species may differ dramatically in terms of pathogenicity, host preference, and effective countermeasures (e.g., Barnes et al. 2004; Queloz et al. 2011). Although end users do have options to propose changes in the data and metadata in the public sequence databases, few users take action when they encounter compromised sequences (Pennisi 2008; Nilsson et al. 2012).

Molecular identification of fungi usually relies, at least in the first attempts, on sequencing the nuclear ribosomal internal transcribed spacer (ITS) region, the formal fungal barcode (Schoch et al. 2012). The largest database tailored for fungal ITS sequences is UNITE (<http://unite.ut.ee>; Abarenkov et al. 2010a). UNITE mirrors and curates the International Nucleotide Sequence Database Collaboration (INSDC:

GenBank, ENA, and DDBJ; Nakamura et al. 2013) for fungal ITS sequences and offers extensive capacities for analysis and third-party annotation of sequences to its users. It has been the subject of several annotation efforts (Tedersoo et al. 2011; Bengtsson-Palme et al. 2013; Kõljalg et al. 2013), but these have in part been biased towards basidiomycetes and mycorrhizal fungi. A similar effort for plant pathogenic fungi was initiated at the symposium “Classical and molecular approaches in plant pathogen taxonomy” (10–11 September 2013, Warsaw). In addition to several of the symposium participants, other experts on various fungal lineages known to harbour plant pathogens were invited as contributors through personal networking, email, and ResearchGate (<http://www.researchgate.net/>). Several experts on epiphytic and endophytic fungi also participated in the effort; while these fungi may not be plant pathogenic, they are often isolated alongside, or mistaken for, plant pathogenic fungi (Unterseher et al. 2013). Moreover, many fungi showing pathogenicity in certain plants represent common endophytes in other host plants (Delaye et al. 2013). This paper reports on the outcome of the annotation effort.

Materials and methods

Using third-party sequence annotation facilities provided by the PlutoF workbench (<http://plutof.ut.ee>, Abarenkov et al. 2010b), the participants examined fungal lineages and ecological groups of their respective expertise in UNITE for four parameters: (i) selection of representative sequences for species, (ii) improvement of taxonomic annotations, (iii)

addition of ecological metadata (chiefly host and country of collection), and (iv) compromised sequence data.

(i) Selection of representative sequences for species

UNITE clusters all public fungal ITS sequences to approximately the genus/subgenus level. A second round of clustering inside each such cluster seeks to produce molecular operational taxonomic units at approximately the species level; these are called *species hypotheses* (SHs; Fig. 1; Kõljalg et al. 2013). The species hypotheses are open for viewing and querying (<http://unite.ut.ee/SearchPages.php>) through uniform resource identifiers (URIs) such as “<http://unite.ut.ee/sh/SH158651.06FU>”. As a proxy for the species hypothesis, a representative sequence is chosen automatically from the most common sequence type in the species hypothesis. Through these representative sequences, UNITE assigns a unique, stable name of the accession number type – SH158651.06FU in its shortest form for the example above – to all species hypotheses to provide a means for unambiguous reference to species-level lineages even in the absence of formal Latin names. The representative sequences are also used for non-redundant BLAST databases for molecular identification in several next-generation sequencing analysis pipelines. Depending on the algorithm, including all available fungal ITS sequences in the reference database slows down sequence similarity searches significantly, and the use of downsized, non-redundant databases with only one sequence per taxon of interest is a common solution. The representative sequences of UNITE fulfill these criteria, since they comprise a single sequence from all fungal species hypotheses recovered to date

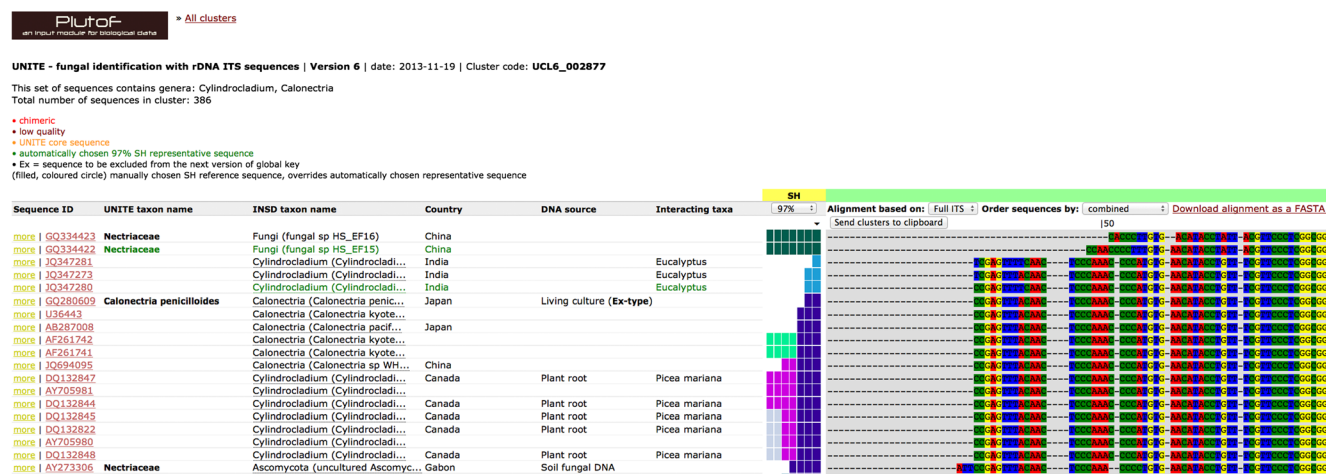


Fig. 1 A screenshot from the web-based PlutoF sequence management environment showing a *Nectriaceae* cluster, with the individual species hypotheses at different similarity levels indicated by the coloured vertical bars. Country of collection and host/interacting taxa are specified together with taxonomic re-annotations. Sequences from type material are indicated. For species hypotheses where no user has designated a reference

sequence, the clustering program chooses a sequence from the most common sequence type to represent that species hypothesis (shown in green font). The species hypotheses are mirrored by GenBank through a LinkOut function, making it possible to go from a BLAST search in GenBank to the corresponding species hypothesis in UNITE through a single click

through ITS sequences by the scientific community. However, there are situations where one would like to influence which sequence is chosen to represent a species hypothesis. In ideal cases, the type specimen or an ex-type culture has been sequenced. Such “type sequences” form the best possible proxy for the species hypothesis, as long as they are sufficiently long and of high technical quality.

To increase the proportion of plant pathology-related fungal taxa represented by sequences from types, we scanned the 27 largest journals in plant pathology (and 12 mycological journals known for an inclination towards plant pathology or fungi otherwise associated with plants) for descriptions of new (or typifications of existing) plant pathogenic or plant-associated species of fungi (Supplementary Item 1). For all descriptions where an ITS sequence was generated from the type specimen/ex-type culture by the original authors, we examined the sequence in the corresponding UNITE cluster for read quality and length. All type sequences deemed to be of high technical quality and sufficient length were designated as reference sequences for their respective species hypothesis.

(ii) Correction of taxonomic affiliations

Taxonomic misidentifications are rife in the public nucleotide sequence databases. Similarly, more than half of all public fungal ITS sequences are not annotated to the level of species, and most of these carry little or no taxonomic annotation save, e.g., “Uncultured fungus” (cf. Hibbett et al. 2011). This makes molecular identification difficult and can lead to an incorrect name or no name at all, even when full (e.g., *Colletotrichum melonis*) or partial (e.g., *Colletotrichum* sp. or Glomerellales) naming would have been possible. Clearly it is important to avoid the common mistake of over-estimating taxonomic certainty based solely on BLAST searches, which often yield many top hits with similar quality scores and can obscure sister-level relationships to the taxa represented in the top matches. BLAST results may also differ over time according to database content, and differ markedly when, e.g., the full ITS vs. partial ITS sequences or ITS sequences with non-trivial lengths of the ribosomal small and/or large subunits for the same strain are submitted to searches (U’Ren et al. 2009). Indeed, a substantive portion of misidentified sequences in public databases appear to have resulted from spurious applications of taxonomic names to sterile mycelia, environmental samples, or otherwise unknown strains, often being studied by non-taxonomists. However, careful evaluation of database matches can provide additional information about taxonomic placement that can be applied judiciously by experts to better serve the scientific community. In addition, sequences without taxonomic annotations (e.g., “Uncultured fungus”) are often unfairly disregarded in phylogenetic studies (Nilsson et al. 2011). Another reason to improve the taxonomic annotation of public ITS sequences is therefore to highlight their existence

and availability for use in phylogenetic and systematic studies. Such enhanced taxon sampling carries many advantages (Heath et al. 2008). We scanned our fungal lineages of expertise in UNITE to make sure the sequences carried the most accurate name possible, viz. the full species name for fully identified sequences, and the genus, family, order, class, or phylum name for sequences that could not be fully assigned.

(iii) Addition of geographical and ecological metadata

Although DNA sequences form the core of molecular identification of fungi, additional data are often needed for final, informed decisions on the taxonomic affiliation of newly generated sequences. For plant pathogenic fungi, the identity of the host and the geographical origin of the sequences are often critical information (Britton and Liebhold 2013). Yet these metadata are usually not included with sequence data in public sequence databases; Tedersoo et al. (2011) showed, for instance, that a modest 43 % of the public fungal ITS sequences were annotated with the country of origin. To the same effect, Ryberg et al. (2009) found that host of collection was reported for less than 25 % of all public fungal ITS sequences (although not all fungi necessarily have a host). We made sure that the sequences of our core expertise were as richly annotated as possible in UNITE through recursions to the original publications.

(iv) Technical quality of sequences

Detecting sequences of substandard quality in public databases is difficult because sequence chromatograms or other original data are not present for verification of nucleotide identity, and sequencing technologies have different error rates and types of errors (e.g., 454 pyrosequencing vs. Sanger sequencing). Standards also differ among researchers and computer programs with regard to quality thresholds and what is deemed acceptable for individual nucleotides or whole-sequence reads. The extent to which sequence depositors take measures to ensure that their sequence data are of satisfactory integrity also seems to differ markedly. To discriminate with full certainty among publicly deposited sequences of high and substandard quality is simply not possible in all situations (Nilsson et al. 2012). To remove all sequences that are putatively substandard is certain to lead to many instances of false-positive removals (i.e., removal of authentic albeit poorly known biodiversity), and in this study we settled for removing entries we could prove were compromised. We evaluated sequence quality on the basis of length, evidence of chimera formations or poor read quality, and mislabelling of the genetic marker that the data represent.

Results

The participants implemented a total of 31,954 changes, including 5,135 taxonomic re-annotations, 25,028 specifications of geographical and ecological metadata, 1,368 designations of reference sequences, and 401 exclusions of substandard sequences, distributed over some 48 fungal orders. The results were incorporated in UNITE for all its users. In addition, they are made publicly available through the UNITE release of all public fungal ITS sequences (<http://unite.ut.ee/repository.php>) for use in, e.g., local sequence similarity searches and sequence processing pipelines such as QIIME (Caporaso et al. 2010; Bates et al. 2013), mothur (Schloss et al. 2009), SCATA (<http://scata.mykopat.slu.se/>), CREST (Lanzén et al. 2012), and other downstream applications. UNITE also serves as one of the data providers for BLAST (Altschul et al. 1997) searches in the EUBOLD fungal barcoding database (<http://www.cbs.knaw.nl/eubold/>).

(i) Selection of representative sequences for species

The extraction of sequences from type material from the literature resulted in 965 designations of reference sequences (for as many species hypotheses and a total of 194 genera of fungi; Table 1). We also designated 403

additional reference sequences based on our expertise; 174 of these stemmed from type material and 229 were from other authentic material. The latter cases involved fungal taxa of our core expertise where we knew the type material was missing or too old for DNA sequencing and where we knew that the selected sequences were as close to the type as possible in terms of morphology, country, and/or substrate of collection. A total of 202 genera were designated with at least one reference sequence.

(ii) Correction of taxonomic affiliations

The process of verifying taxonomic names given to sequences resulted in a total of 5,135 changes (Table 1), notably for the orders Hypocreales (459 changes), Glomerellales (404 changes), and Botryosphaerales (393 changes). In addition, 22 ITS sequences were found to stem from kingdoms other than Fungi and were re-annotated accordingly.

(iii) Addition of geographical and ecological metadata

Our effort to complement the sequences with metadata from the literature resulted in a total of 14,478 specifications of host and 10,550 specifications of country of origin (Table 1).

Table 1 Summary of the changes made in the UNITE database. The 15 orders that saw the largest number of changes are specified separately; all other lineages are amalgamated into the “Others” category

Order	Taxonomic re-annotations	Country	Host	Reference sequences	Count
Hypocreales	459	3,751	2,960	118 (116)	7,288
Pleosporales	129	860	4,344	76 (76)	5,409
Capnodiales	200	960	1,696	181 (181)	3,037
Diaporthales	79	1,374	855	28 (28)	2,336
Glomerellales	404	814	824	148 (148)	2,190
Botryosphaerales	393	428	626	70 (67)	1,517
Mucorales	90	630	631	87 (63)	1,438
Eurotiales	420	411	226	168 (168)	1,225
Xylariales	90	225	823	19 (19)	1,157
Helotiales	333	301	290	108 (46)	1,032
Chaetothyriales	22	121	521	17 (17)	681
Puccinales	134	313	194	9 (1)	650
Agaricales	442	31	8	21 (21)	502
Pezizales	297	0	97	1 (1)	395
Erysiphales	143	55	66	129 (4)	393
Others	1,500	276	317	188 (183)	2,281

Taxonomic re-annotations = The number of taxonomic (re)annotations implemented. Country = The number of specifications of country of collection. A total of 94 different countries were added. Host = The number of host specifications added in the system. Reference sequences = The number of reference sequences designated through manual inspection (of which sequences from type material are indicated in parentheses). Count = Total number of changes

(iv) Technical quality of sequences

We detected a total of 363 sequences of substandard technical quality. These were marked as compromised, which precludes them from being used in molecular identification procedures while still keeping them open to direct searches in the system. This included 84 cases of chimeric sequences and 279 cases of low read quality. Another 38 sequences were annotated as ITS sequences by their submitters but were found to represent other genes and markers (notably the ribosomal small and large subunits) and were re-annotated accordingly.

Discussion

Fungal pathogens of agricultural, silvicultural, horticultural, and wild plants can compromise ecosystem health and cause considerable economic loss globally. Correct identification of these fungi and subsequent understanding of their biology and ecology are key elements in protecting their host plants (Rossman and Palm-Hernández 2008). However, identification of plant pathogenic fungi to the species level is relevant to more than just studies of plant pathology. Because of the ease and moderate cost at which large amounts of sequence data can be generated, fungi and fungal communities are now being studied by an increasing number of non-mycologists, notably soil biologists, molecular ecologists, and researchers in the medical sciences (e.g., Ghannoum et al. 2010; La Duc et al. 2012; Pautasso 2013). Phytopathogenic fungi also occur in these substrates and ecosystems in various life stages, including sterile mycelia, resting stages, and propagules. Although some plant pathogenic fungi have been studied in great detail, the biology of the majority of phytopathogenic fungi remains poorly known. Therefore, information stemming from non-mycological or non-pathological research efforts may increase our understanding of these taxa. As a consequence, it is important that all researchers, regardless of expertise and extent of mycological knowledge, can obtain reliable estimates of the taxonomic identity of plant pathogenic – and all other – fungi in whatever form they are recovered.

Molecular identification of plant pathogenic fungi can be challenging due to differing sequence and annotation quality of the available reference sequences. We have gone through a large number of plant pathogenic fungal groups within our collective expertise. A total of 31,954 changes in 48 fungal orders were implemented in UNITE for these groups (Table 1). However, not all plant pathogenic lineages of fungi – or, indeed, even the groups covered by the present effort – are satisfactorily resolved in UNITE. In addition, new sequences (of both known and unknown species) are continuously generated and deposited in the INSDC by the scientific community, such that a limited

group of people can never stay abreast of the data deposition. A community effort is clearly required. UNITE offers third-party annotation capacities to all its registered users. Registration is free, and contributions from all relevant scientific communities are most welcome. Even small edits – such as designating a reference sequence for a single species hypothesis, correcting and improving a handful of taxonomic annotations, or adding metadata that can be used for comparative studies (Supplementary Item 2) – will improve the database significantly and may be of substantial importance to other researchers. Going through the alignments and metadata for one's fungi of expertise in the web-based system is furthermore a good way to visualize and explore patterns in the data and identify new research questions.

Many of the corrections brought about by the present effort would have been unnecessary if the original sequence authors had taken the time to examine and annotate their sequences properly prior to submission. Lack of time and awareness of these issues are the presumed culprits. Guidelines on how to process newly generated sequences in a way to establish their integrity and maximize their usefulness to the scientific community are given in Seifert and Rossman (2010), Nilsson et al. (2012), Hyde et al. (2013), and Robbertse et al. (2014). In addition, to facilitate future assessments of sequence quality and other pursuits, we urge sequence depositors in INSDC to archive chromatograms and other relevant data in UNITE or in other resources that support long-term data storage and availability. The present initiative will contribute to more accurate molecular identification of plant pathogenic fungi for three sets of users: UNITE users, anyone using the ~350,000-sequence downloadable FASTA file of the UNITE/INSDC fungal ITS sequences (<http://unite.ut.ee/repository.php>) for local BLAST searches or similar, and researchers using any of the major next-generation sequencing analysis pipelines or the EUBOLD database to process newly generated fungal ITS datasets. In addition, following the data sharing history between UNITE and the INSDC, the results were made available to the INSDC to reach the widest possible audience. Fungal barcoding is in a state of constant development, but it should be clear that collaboration and data sharing among resources are necessary for the future development of the field. Mycology struggles for funding in competition with fields that are often deemed larger or more fashionable, and we simply cannot afford public fungal DNA sequences to remain in a suboptimal state. On the contrary, we hope mycologists will work together to make fungal sequence data as richly annotated and as easily interpreted as possible because, after all, many of the end users of those data will not be mycologists. The present study is a small step in that direction, and we hope that others will follow.

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