Two new species of the *Fusarium fujikuroi* species complex isolated from the natural environment

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Abstract Two new species in the *Fusarium fujikuroi* species complex (FFSC) are introduced. One of these, represented by strain CBS 454.97 was isolated from plant debris (*Striga hermonthica*) in the Sudan, while the second, represented by strains CBS 119850 and CBS 483.94, which originated from soil in Australia. Molecular analyses were performed including *TEF1* spanning 576 bp region, 860 bp region of *rPB2*, and 500 bp *BT2* region. Phylogenetic trees based on these regions showed that the two species are clearly distinct from all known taxa in the *F. fujikuroi* species complex. Based on phenotypic, physiological characters and molecular data, we introduce *Fusarium sudanense* and *Fusarium terricola* as novel species in the complex.

Keywords *Fusarium* · Saprobe · Morphology · Molecular phylogeny

Introduction

*Fusarium* is a large and variable genus with nearly 300 recognized species occurring worldwide in a diversity of habitats. Particularly in plant pathology, species have extensively been studied because of their opportunism on numerous hosts, among which are economically important crops. For example, many *formae speciales* have been reported in *F. oxysporum* and relatives (Ordonez et al. 2015) as etiologic agents of plant diseases. Some species seem to have a narrow host range or may even be host-specific, such as...
Fusarium ficicrescens that has as yet only been found on figs (Al-Hatmi et al. 2016a). Members of the genus are increasingly observed as agents of human infection (Al-Hatmi et al. 2016b). A further significant property is their production of mycotoxins, especially in Fusarium species that occur in association with farm animals receiving cereal-based diets (de Nijs et al. 1997).

Typically, most species are soil-borne, causing diseases in seedlings or weakened plants (Watanabe 2013). Fusarium is a common mould in the environment and different environmental factors, such as moisture, temperature, nutrients and other ones appear to be of great importance for colonization of a wide diversity of substrates and ecological niches (Smith 2007). Geographical factors including climate are of prime importance for the diversity of Fusarium species (Summerell et al. 2010; Karim et al. 2016). Strictly saprobic Fusarium have received less attention, though they are widely distributed in natural habitats, notably in soil, where they might have a role in the turnover of organic matter (Karim et al. 2016). However, saprobic strains may become opportunistic upon availability of a susceptible host (Rep et al. 2005). Furthermore, given the widespread occurrence of Fusarium in the environment, it seems reasonable to hypothesize also that pathogenic forms of Fusarium may have evolved from non-pathogenic ancestors (Alves-Santos et al. 1999). Thus, many Fusarium species with importance to environment, agriculture and human health have a reservoir in soil, and their infections in a wide range of plants (Wakelin et al. 2008), animals (O’Donnell et al. 2016) and humans (Al-Hatmi et al. 2016b) are regarded to be of an opportunistic nature.

The Fusarium fujikuroi species complex (FFSC) is one of the larger groups within the genus Fusarium with various ecologies (Nirenberg and O’Donnell 1998; O’Donnell et al. 2000; Al-Hatmi et al. 2015). Studies suggested that with the use of molecular data more than 50 phylogenetic species within the fujikuroi complex might be recognized (O’Donnell et al. 2015). Recently, Herron et al. (2015) described eight more species in the fujikuroi complex from stem cankers and branches of Pinus plants. Laurence et al. (2015) added three additional species from Australian natural forests, Al-Hatmi et al. (2016a) described F. ficicrescens from figs in Iran and Edwards et al. (2016) published F. agapanthi as a novel plant pathogen from Australia and Italy.

Recent and historical ecosystem surveillance in Australia has resulted in the discovery of novel Fusarium species including F. aywerte, F. babinda, F. beominforme, F. burgessii, F. coicus, F. gaditjirri, F. goolgardi, F. iyarnte, F. mundagurra, F. nurragi, F. newnesense, F. nygamai, F. tjaetaba, F. tjaynera and F. werrikimbe (Laurence et al. 2015). This number has increased to 16 species with the recent description of F. agapanthi above (Edwards et al. 2016). In the present study, the taxonomic status of all available strains of the F. fujikuroi species complex was verified using a polyphasic approach. The resultant data show that some isolates represent two new Fusarium species, for which we propose the names Fusarium terricola for a species isolated from Australia and Fusarium sudanense that was isolated in Sudan.

Materials and methods

Strains

Three strains in the reference collection of Centraalbureau voor Schimmelcultures (housed at Westerdijk Fungal Biodiversity Institute), previously identified morphologically as F. nygamai, were analyzed and compared with all available members of the F. fujikuroi species complex. Two of these strains (CBS 119850 and CBS 483.94) were isolated from soil in Australia, while an additional strain (CBS 454.97) originated from Striga hermonthica. The latter strain was included in a multilocus molecular phylogenetic analysis as Fusarium sp. NRRL 26793 as a distinct clade (Herron et al. 2015; Laurence et al. 2015).

Morphology

Colony characteristics and growth morphology were studied by inoculating the isolates onto plates of Malt Extract Agar (MEA; Oxoid, U.K.), Oatmeal Agar (OA; home-made at CBS), Potato Dextrose Agar (PDA; Oxoid), Synthetic Nutrient Agar (SNA; CBS) (Nirenberg 1976) and carnation leaf agar (CLA; CBS) (Leslie and Summerell 2006). Cultures were grown under 12 h light–dark (l/d) cycles with UV and daylight colour fluorescent lights at 24 °C. Morphological characters examined included the shape and size of macroconidia produced in sporodochia on Carnation Leaf Agar (CLA) (Fisher et al. 1982), the...
shape and mode of formation of microconidia on CLA and SNA (Nirenberg 1976), the production of chlamydospores on Potato Dextrose Agar (PDA). Microscopic slides were prepared for each isolate by mounting structures in lactic acid and the slides were made from cultures grown on CLA plates which were observed after 5 days of incubation at 24 °C. Slides were examined with a Nikon Eclipse 80i light microscope, and pictures were taken using a camera attached to the microscope (Nikon; digital-sight DS-5M). A minimum of 10 measurements per structure were taken and the average was calculated.

Growth rate
Cardinal growth temperatures were determined on MEA and PDA plates incubated in the dark for 2 weeks at temperatures of 18–40 °C at intervals of 3 °C; with two replicates for each isolate. Average growth rates per species were calculated and expressed as diametric growth per 24 h.

DNA amplification and sequencing
The following partial genes were amplified directly from genomic DNA for multilocus sequence typing: elongation factor 1 alpha (TEF1) (O’Donnell et al. 2010), the second largest subunit of RNA polymerase (rPB2) (Reeb et al. 2004), and β-tubulin (BT2). PCR amplification and sequencing were performed according to the protocol applied by Al-Hatmi et al. (2016a).

Phylogenetic inference
To confirm the identity of our presumed new Fusarium species, we evaluated their position in Bayesian phylogenetic and RAxML trees of the following individual gene markers (BT2, TEF1 and rPB2). In these analyses, our sequences, together with sequences retrieved from GenBank were analysed (Table 1). Sequences were aligned with MAFFT (www.ebi.ac.uk/Tools/msa/mafft/), followed by manual adjustments with MEGA v6.2 and BioEdit v7.0.5.2. A single alignment was constructed for TEF1 and BT2 and rPB2. The analysis included 58 sequences for TEF1, 50 sequences for BT2 and 32 sequences for rPB2. The best-fit model of evolution, determined by MEGA v6.2, was used to infer the appropriate substitution model that would best fit the model of DNA evolution for each sequence data set. Maximum likelihood (ML) and Bayesian inference (BI) analyses were used to estimate phylogenetic relationships. ML analysis was performed with RAxML-HPC v7.0.3 (Stamatakis et al. 2005; Stamatakis 2006) with a K2+G model of evolution for TEF1, BT, rPB2 and the combined data. Nodal support was determined by nonparametric bootstrapping (BS) with 1000 replicates. BI analysis was performed in a likelihood framework as implemented in MRBAYES v3.0b4 to reconstruct phylogenetic trees (Huelsenbeck and Ronquist 2001). Multiple Bayesian searches using Metropolis-coupled Markov chain Monte Carlo sampling were conducted. One cold and three heated Markov chains were used in the analysis. Analyses were run for 10 million generations, with trees sampled every 1000 generations. The first 25% of the trees, which represented the burn-in phase of the analysis, were discarded. The remaining trees were used for calculating posterior probabilities (PP) of recovered branches (Larget and Simon 1999) in the 50% majority rule consensus tree. Sequences included in this study were supplemented with those from GenBank Fusarium oxysporum was used as outgroup and the GenBank accession numbers for the three strains are shown in Table 1.

Results
Using the BLAST similarity search (performed on January 15 2017), the TEF1 region of the strain CBS 454.97 showed 99% (546/547 bp) similarity to F. andiyazi strain F16 (JX307409.1) which appears to be wrongly labeled in GenBank. Another closely related strain was Fusarium sp. NRRL 26793 with 99% similarity. Further comparison using the FUSARIUM ID database (http://isolate.fusariumdb.org) (Geiser et al. 2004) revealed Gibberella fujikuroi species complex (GFSC) NRRL 26793 with 99.83% identity, while the Fusarium MLST database (http://www.cbs.knaw.nl/fusarium) (O’Donnell et al. 2010) yielded F. nygamai with 99.82% similar to NRRL 26793 (AF160309). CBS 119850 and CBS 483.94 showed a similarity of 100% with F. andiyazi strain F16 (JX307409.1) in GenBank, G. fujikuroi species complex (GFSC) with 98.93% similarity in FUSARIUM ID, and NRRL 26793 Fusarium sp. with 98.9% similarity in Fusarium MLST.
Table 1 GenBank accession numbers of the *F. fujikuroi* species complex used in phylogenetic analysis of *F. terricola* and *F. sudanense*

| Species           | Collection | β-tubulin | TEF1α       | RPB2                      | Reference                                      |
|-------------------|------------|-----------|-------------|---------------------------|------------------------------------------------|
| *F. acutatum*     | NRRL 13308 | U34431    | AF160276    | (CBS402.97)/KT154005      | Scauflaire et al. (2011), Al-Hatmi et al. (2016a) |
| *F. agapanthi*    | NRRL 54465 | KU9006361 | KU9006311   | KU9006261                 | Edwards et al. (2016)                          |
| *F. andiyazi*     | CBS 119857 | KP662894  | KP662901    | CBS 119857/KT154004       | Al-Hatmi et al. (2016a)                         |
| *F. anthophilum*  | NRRL 13602 | U61541    | AF160292    | (CBS222.76)/KT154006      | Scauflaire et al. (2011), Al-Hatmi et al. (2016a) |
| *F. bactridioides*| NRRL 20476 | U34434    | AF160290    | –                         | Scauflaire et al. (2011)                        |
| *F. begoniae*     | NRRL 25300 | U61543    | AF160293    | –                         | Scauflaire et al. (2011)                        |
| *F. brevicatenulatum* | NRRL 25446 | U61623.1  | AF160265    | –                         | Scauflaire et al. (2011)                        |
| *F. bulbicola*    | NRRL 13618 | U61546    | AF160294    | KF466404                  | Scauflaire et al. (2011), Proctor et al. (2013) |
| *F. cinctinum*    | NRRL 25331 | U61547    | AF160295    | JX171623                  | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. coicus*       | RBG 5368   | –         | KP083251    | KP083274                  | Laurence et al. (2015)                         |
| *F. concentricum* | NRRL 25181 | U61548    | AF160282    | –                         | Scauflaire et al. (2011)                        |
| *F. denticulatum* | NRRL 25302 | U34453.1  | AF160271    | –                         | Scauflaire et al. (2011)                        |
| *F. dlaminii*     | NRRL 13164 | U34430    | AF160277    | –                         | Scauflaire et al. (2011)                        |
| *F. fiscicrescens*| CBS 125178 | KP662896  | KP662899    | KT154002                  | Al-Hatmi et al. (2016a)                         |
| *F. fracticaudum* | CMW: 25245 | –         | –           | –                         | Herron et al. (2015)                            |
| *F. fractiflexum* | NRRL 28852 | AF160315  | AF160288    | –                         | Scauflaire et al. (2011)                        |
| *F. fujikuroi*    | NRRL 13566 | U34415    | AF160279    | EF470116                  | Scauflaire et al. (2011), O’Donnell et al. (2007) |
| *F. globosum*     | NRRL 26131 | U61557    | AF160285    | KF466406                  | Scauflaire et al. (2011), Proctor et al. (2013) |
| *F. guttiforme*   | NRRL 22945 | U34420    | AF160297    | JX171618                  | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. inflexum*     | NRRL 20433 | U334435   | AF8479      | JX171583                  | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. konzum*       | MRC 8544   | EU220234  | EU220235    | –                         | Scauflaire et al. (2011)                        |
| *F. lactis*       | NRRL 25200 | U61629    | AF160272    | KM582794                  | Scauflaire et al. (2011), Triest et al. (2015)  |
| *F. mangiferae*   | NRRL 25226 | U61561    | AF160281    | JX171622                  | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. marasasianum* | CMW: 25261 | KJ541054  | KJ541063    | –                         | Herron et al. (2015)                            |
| *F. madagurra*    | RBG 5717   | –         | KP0832561   | KP0832761                 | Laurence et al. (2015)                         |
| *F. musae*        | NRRL 28893 | FNS45374  | FNS552092   | FNS552114                 | Van Hove et al. (2011)                          |
| *F. napiforme*    | NRRL 13604 | U34428    | AF160266    | EF470117                  | Scauflaire et al. (2011), O’Donnell et al. (2007) |
| *F. nygamai*      | NRRL 13448 | U34426    | AF160273    | EF470114                  | Scauflaire et al. (2011), O’Donnell et al. (2007) |
| *F. parvisorum*   | CMW: 25267 | KJ541055  | KJ541060    | –                         | Herron et al. (2015)                            |
| *F. pininemorale* | CMW: 25243 | KJ541049  | KJ541064    | –                         | Herron et al. (2015)                            |
Using a BLAST similarity search, the rPB2 region of strain CBS 454.97 (KU604266) showed 99% (791/794 bp) similarity to *F. nygamai* (FRC M-7492 = KF466410), the next closest taxon was a strain of *F. nygamai* (PUF025 = HQ423219.1) with 99% similarity (788/794 bp). The rPB2 sequence of CBS 483.94 and CBS 119850 (= KU604268) shared 99% similarity (785/791 bp) with *F. nygamai* (FRC M-7492 = KF466408.1) in GenBank, *G. fujikuroi* species complex (GFSC) with 98.74% similarity in FUSARIUM ID, and *F. nygamai* (CBS 749.97) with 98.74% similarity in *Fusarium* MLST. The different indication of the species complexes, either with *Gibberella* or with *Fusarium*, is due to the use of either the name of the sexual or the asexual morph, respectively; at present the name *Fusarium* is preferred over *Gibberella* and hence the same species complex is now known as FFSC.

For further understanding of relations between species, a phylogenetic tree was constructed for each locus separately, i.e. TEF1, BT2, and rPB2. In each single tree of BT2, rPB2 and TEF1 separately, strains CBS 119850 and CBS 483.94 from soil in Australia, and an additional strain CBS 454.97 from plant debris in Sudan were found to form a monophyletic clades supported by a high bootstrap values (Figs. 1, 2, 3).

The TEF1 dataset comprising 58 sequences consisted of 53 taxa with 576 characters, from which 202 were variable, 111 parsimony-informative and 91 were singletons. Phylogenetic analyses of 50

| Species                  | Collection | β-tubulin | TEF1α | rPB2 | Reference                  |
|--------------------------|------------|-----------|-------|------|----------------------------|
| *F. phyllophilum*        | NRRL 13617 | U34432    | AF160274 | KF466410 | Scauflaire et al. (2011), Proctor et al. (2013) |
| *F. proliferatum*        | NRRL 22944 | U34416    | AF160280 | JX171617 | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. pseudoanthophilum*   | NRRL 2520  | U61631    | AF160264 | –     | Scauflaire et al. (2011) |
| *F. pseudocircinatum*    | NRRL 22946 | U34453    | AF160271 | –     | Scauflaire et al. (2011) |
| *F. pseudonygamai*       | NRRL 13592 | U34421    | AF160263 | –     | Scauflaire et al. (2011) |
| *F. ramigenum*           | NRRL 25208 | U61632    | AF160267 | KF4664121 | Scauflaire et al. (2011) |
| *F. sacchari*            | NRRL 13999 | U34414    | AF160278 | JX171580 | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. sororula*            | CMW: 40578 | KJ541057  | KJ541067 | –     | Herron et al. (2015) |
| *F. subglutinans*        | NRRL 22016 | U34417    | AF160289 | JX171599 | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. succisae*            | NRRL 13613 | U34419    | AF160291 | –     | Scauflaire et al. (2011) |
| *F. sudanense*           | CBS 454.97 | KU603909  | KU711697 | KU604266 | This study |
| *F. sterilihyphosum*     | CML 283    | DQ445780  | DQ452858 | –     | Scauflaire et al. (2011) |
| *F. temperatum*          | MUCL 52436 | HM067692  | HM067684 | –     | Scauflaire et al. (2011) |
| *F. terricola*           | CBS 483.94 | KU603908  | KU711698 | KU604267 | This study |
| *F. terricola*           | CBS 119850 | KU603907  | KU711699 | KU604268 | This study |
| *F. tjaetaba*            | RBG 5361   | –         | KP083263 | KP083275 | Laurence et al. (2015) |
| *F. thapsinum*           | NRRL 22045 | U34444    | AF160270 | JX171600 | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *F. udum*                | NRRL 22949 | U34433    | AF160275 | –     | Scauflaire et al. (2011) |
| *F. verticillioides*      | NRRL 22172 | U34413    | AF160262 | EF470122 | Scauflaire et al. (2011), O’Donnell et al. (2013) |
| *Fusarium* sp.           | NRRL 26756 | –         | AF1603071 | –     | O’Donnell et al. (2000) |
sequences of BT2 resolved the phylogenetic positions of the two novel taxa in relation to the currently recognised monophyletic species in the *F. fujikuroi* species complex used in the current analysis (Figs. 1, 2). The BT2 dataset comprising 50 sequences consisted of 48 taxa with 500 characters, from which 129 were variable, 70 parsimony-informative and 58 were singletons. In our study, we were able to cover all taxa which have *rPB2* sequences retrieved from the GenBank. We used 32 sequences retrieved from GenBank representing 28 species of the *fujikuroi* complex. Ribosomal polymerase B2 (*rPB2*) is one of the most informative gene fragments and resolves taxonomy at or near the species-level in *Fusarium*, but its drawback is that fewer sequences are available in GenBank. The alignment of *rPB2* sequences had a length of 800 nucleotides when the outgroup was included; 175 were variable, 104 parsimony-informative and 71 were singletons.

The combined TEF1 and *rPB2* alignment for 28 species consisted of 32 sequences each with 1411 characters; the ML/BI tree is shown in Fig. 4. The
analysis indicated that the isolates (CBS 454.97) and (CBS 119850 and CBS 483.94) form distinct clades separated from other species of *fujikuroi* complex and these two clades have support (75% BS and 0.8 PP); for CBS 454.97, and (99% BS and 1 PP) support for (CBS 119850 and CBS 483.94 respectively) (Fig. 4). Bayesian and maximum likelihood phylogenetic trees constructed with *rPB2* sequences of available strains appeared well-resolved. All clades had statistical support between 60–100% and all species were well separated. Intraspecific polymorphism within the species clusters was observed with *BT2*, *TEF1* and
Overall topologies of the trees were similar to those described previously for the FFSC (Al-Hatmi et al. 2016c).

**Taxonomy**

*Fusarium terricola* Al-Hatmi, S.A. Ahmed and de Hoog, sp. nov.—Fig. 5. MycoBank MB 816188.

Etymology: terricola means soil-loving, referring to the fungus’ apparently preferred habitat.

Holotype: dried specimen in herbarium CBS H-22548; living ex-type strain CBS 483.94, isolated from desert soil, Queensland, Australia.

Description based on CBS 483.94 on MEA and CLA growing in the dark at 27 °C after 7 days. Colonies growing rapidly, attaining 50 mm diam. Obverse aerial mycelium cottony, initially white and later becoming pinkish to purple on MEA (Fig. 5). Reverse pinkish-orange to darker purple. Sporodochia seen after 7 days of incubation as pale orange spots on pieces of carnation leaf placed on CLA. Sporulation on SNA starting early in aerial mycelium and later on agar surface. Aerial conidiophores in darkness mostly prostrate, simple to sparsely branched, but some erect and branching sympodially or verticillately, resulting in a complex tree-like morphology (Fig. 5d). Conidiophores 90–100 μm; conidiogenous cells are mostly polyphialidic. Conidia produced mostly on phialides formed directly on substrate hyphae (Fig. 5g, h). Monophialides 10.0–14.5 × 3–5 μm, ellipsoidal, tapered towards the apex with minute basal frill. Microconidia ovoidal, 5.7–4.2 × 1.8–2.4 μm. Macroconidia abundant, 24.0–31.9 × 5.6–6.0 μm, 2–5 septate, falcate, with a beaked apical cell and a foot-like basal cell (Fig. 5). Chlamydospores absent.

*Fusarium sudanense* S.A. Ahmed, Al-Hatmi and de Hoog, sp. nov.—Fig. 6. MycoBank MB 816189.
Etymology: named after the country of isolation, Sudan.

Holotype: dried specimen in herbarium CBS H-22547; living ex-type strain CBS 454.97, from plant debris (Striga hermonthica), Sudan.

Description based on CBS 454.97 on MEA and CLA growing in the dark at 27 °C after 7 days. Colonies expanding, attaining 45 mm diam. Aerial mycelium cottony, initially white and later becoming light pinkish, reverse pink-orange (Fig. 6a, b). Hyphae 1.9–2.9 μm, smooth-walled, hyaline, branched, septate. Conidiophores phialidic with mostly monophialides, rarely polyphialides (Fig. 6g–i). Monophialides 13.0–17.4 × 2.0–3.0 μm, elongate-ampulliform or subcylindrical and tapered at the apex, or short ossiform, wider at the base. Microconidia abundant, sub spherical or ovoidal, 3.5–10.5 × 2.7–1.7 μm (Fig. 6j) Macroconidia not seen. Chlamydospores appearing after 1 week of incubation, single or in chains, consisting of enlarged, thick-walled vegetative cells within hyphae (intercalary) or at hyphal tips (terminal), 8–13 μm diam (Fig. 6c, d).

Cardinal growth temperature tests showed that all cultures evaluated in this study had their optimal development at 27–33 °C, with growth abilities ranging between 18 °C the lowest temp tested and 40 °C as the highest. All strains were still able to grow at 37 °C, but not at 40 °C.

Discussion

This study was initiated to characterize Fusarium strains held at the CBS reference collection at Utrecht,
The Netherlands using polyphasic approaches. Phylogenetic analyses of a 3-gene dataset strongly supported the genealogical exclusivity of *F. terricola* and *F. sudanense* (Taylor et al. 2000). Both species received strong monophyletic bootstrap support in the individual analysis of each gene (Figs. 1, 2, 3) and combined (Fig. 4). Despite phylogenetic differences, *F. terricola* and *F. sudanense* isolates are morphologically similar to the remaining species in the *F. fujikuroi* species complex, however, there are several morphological differences between both species. The morphological description was based on two strains and therefore the phenotypic variability of the described species cannot be predicted. Morphological species concepts are regarded to be unreliable at the species level in *Fusarium* taxonomy (Al-Hatmi et al. 2016d). Diagnostic morphological characteristics between species are not easily observed due to intraspecific variation and because *Fusarium* species over longer phylogenetic distances may look very similar. The biological species concept in the genus is rudimentary due to lack of sexual recombination in several species groups and because the concept may be complicated by parasexuality, hybridization and horizontal gene transfer (Park 2013). For this reason genealogical concordance and absence of recombination between lineages is therefore mostly applied for species delimitation (Taylor et al. 2000).

To overcome possible problems due to phenotypic overlapping, we applied multigene phylogenies to recognize species boundaries. The *TEF1* alpha, is the recommended barcoding region for clinical *Fusarium* spp. (Stielow et al. 2015; Al-Hatmi et al. 2016c). The grouping of the *F. terricola* and *F. sudanense* was clear based on *TEF1* data. *Fusarium terricola* and *F. sudanense* were seen as a sister clade, closely related to undescribed species KU508366.1 *Fusarium* sp. strain T6.1 (Fig. 1). Additional *BT* and *rPB2* sequences data, however, significantly improved resolution and confirmed *F. terricola* and *F. sudanense* as two clades distinct from *F. fujikuroi* complex, closely related to *F. nygamai* (Figs. 2, 3). MLH-BI analyses of the *TEF1*–*α*, *BT* and *rPB2* loci strongly supported a sister group relationship between *F. terricola* and *F. sudanense* and maintained their status as independent evolutionary lineages (Figs. 1, 2, 3).

Based on the phylogenetic species concept, molecular diagnostics using available genetic marker sequences have played an important role in understanding the systematics of the *Fusarium* (Geiser et al. 2004; O’Donnell et al. 2010). The selected marker sequences *TEF1*, *BT* and *rPB2* still have limitations such as incongruent topologies among single gene trees and lack of resolution needed to distinguish species boundaries. For example, our *TEF1* tree (Fig. 1) shows different species (NRRL 25200, *F. lactis* and T6.1 *Fusarium* sp.) as being closest relatives of the proposed taxa, while *BT*, *rPB2* and the concatenated trees indicate *F. nygamai* as being closely related (Figs. 2, 3, 4). In the *Fusarium fujikuroi* complex several genes such as *TEF1*, *rPB2* and *BT* have been used in the construction of the species phylogeny due to their highly conserved regions and the reasonable degree of variation among multiple taxa. However, our results show incongruency among these genes. For example, the molecular phylogeny based on sequenced *TEF1* is incongruent with the *rPB2* and *BT* as a single gene. This might be due to some recombinations going on within the clade in *TEF1*.

A lack of concordance between molecular markers such as *TEF1*, *rPB2* and IGS within the *F. oxysporum* complex has been reported by O’Donnell et al. (2009). Incongruency between single gene phylogenies above species level can be caused by a combination of analytical and biological factors, the analytical factors including taxon sampling, outgroup selection, criteria of optimality, and modeling of sequence evolution in phylogeny construction (Rokas et al. 2003). As biological factors, some studies considered natural selection, recombination and genetic drift of *Fusarium* species (Rokas et al. 2003; Taylor et al. 1999). This might tell us that the *Fusarium* taxonomy has a fundamental flaw due to ongoing evolution and incomplete lineage sorting.

In the present study we characterized two novel *Fusarium* species recovered from soil and plant debris as *F. terricola* and *F. sudanense*. Further research is needed to determine the relation between opportunism on plants or on humans, because both species had an optimum growth around 27 °C and were still able to grow at 37 °C, but not at 40 °C. They thus potentially
might be able to cause infections in humans and plants, but invasion of living organisms has as yet not been observed.

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