**Fusarium volatile**, a new potential pathogen from a human respiratory sample

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- clinical samples
- FFSC
- French Guiana
- fungal taxonomy
- phylogeny

**Abstract:** We describe the isolation and characterization of *Fusarium volatile* from a bronchoalveolar lavage (BAL) sample of a female patient living in French Guiana with underlying pulmonary infections. Phylogenetic analysis of fragments of the calmodulin (*cmdA*), translation elongation factor (*tef1*), RNA polymerase second largest subunit (*rpb2*), and β-tubulin (*tub*) loci revealed that strain CBS 143874 was closely related to isolate NRRL 25615, a known but undescribed phylogenetic species belonging to the African clade of the *Fusarium fujikuroi* species complex. The fungus differed phylogenetically and morphologically from related known species, and is therefore described as the new taxon *Fusarium volatile*. Antifungal susceptibility testing suggested that the new species is resistant to echinocandins, fluconazole, itraconazole with lower MICs against amphotericin B, voriconazole and posaconazole.

**INTRODUCTION**

Species of the *fujikuroi* species complex (FFSC) of the genus *Fusarium* have been extensively studied in view of their ability to cause infections in plants and to produce toxins that may lead to food poisoning (Chilaka et al. 2017). Moreover, some members of the FFSC have repeatedly been reported from opportunistic infections in humans, which may be mild or local in the case of onychomycosis and keratitis, or invasive and severe in individuals with extended burn wounds and bone marrow transplant recipients. In addition, systemic and disseminated infections occur in severely immunocompromised patients (Guarro et al. 2013, Al-Hatmi et al. 2016b, de Hoog et al. 2019).

The FFSC is one of the larger groups within the genus *Fusarium* and contains species with diverse ecologies (Nirenberg & O’Donnell 1998, O’Donnell et al. 2000). Species of the FFSC are characterised by forming yellow, orange or purple colonies on potato dextrose agar (PDA); globose, oval, napiform or clavate macroconidia are borne in chains and false heads on mono- and polyphialides, while different combinations of microconidial morphologies and phialide types can coexist; the macroconidia are thin-walled, almost straight to slightly curved, with a well-developed pedicellate basal cell. Chlamydospores are rarely formed in the FFSC, although they can be present in some species, formed mostly intercalarily on the hyphae, grouped in chains or clusters (Nirenberg & O’Donnell 1998, Leslie & Summerell 2006).

Molecular studies suggested that at least 50 phylogenetic lineages may be recognized within FFSC. Three major clades with limited biogeographic distribution can be distinguished, and these are termed the African, American and Asian clades (O’Donnell et al. 1998). Thus far 16 species of the FFSC have been reported to cause human infections, namely *F. acutatum*, *F. ananatum*, *F. anayazi*, *F. anthophilum*, *F. fujikuroi*, *F. guttiforme*, *F. musae*, *F. napiforme*, *F. nygamai*, *F. proliferatum*, *F. ramigenum*, *F. sacchari*, *F. subglutinans*, *F. temperatum*, *F. thapsinum* and *F. verticillioides* (Al-Hatmi et al. 2016a, de Hoog et al. 2019).

Identification to species level in FFSC is often difficult because of the high morphological diversity and intraspecific variation of microscopic features (O’Donnell et al. 2015). Currently known species of the FFSC are morphologically and genetically very similar and often can only be reliably separated using multilocus molecular analyses (Geiser et al. 2004). However, considering that some members of the FFSC are relevant human opportunistic pathogens or important toxin producers, their correct identification is crucial. The present paper describes a species of FFSC, based on isolate CBS 143874 collected from a human patient specimen in Cayenne, French Guiana, and characterised by morphological and phylogenetic methods. A morphological identification key is provided to identify the novel species and additional FFSC species known from human clinical specimens.
CASE REPORT

The patient was a 22-yr-old woman, native of Brazil, but has lived in French Guiana for 8 yr. She presented at the Internal Medicine Department of the Andrée Rosemon hospital centre in Cayenne (French Guiana) with a 4-mo history of chronic asthenia at the moment of clinical examination. A right cervical, subangulomaxillary adenophlegmon had appeared 1 mo earlier. She did not have any other clinical signs and had no cough or weight loss and denied night sweats. She had antecedents of disseminated lupus erythematosus diagnosed a year before and was treated with an association of hydroxychloroquine and corticoids. Corticotherapy had been interrupted 4 mo earlier.

Chest computed tomography revealed the presence of a left apical pulmonary cavern with alveolar opacity. Computed tomography of the head and neck highlighted the presence of several cervical necrotic adenomegalies on the right side associated with an abscessed subcutaneous collection. Inflammatory syndrome was mild (C-reactive protein = 7 mg/L). A chest radiography suggested the presence of an aspergilloma, with an image compatible with an Aspergillus ball into the pulmonary cavern.

Presence of bacteria and fungi was assessed by culturing a broncho-alveolar fluid (BAL) sample. Cultures were negative with an image compatible with an inflammatory syndrome was mild (C-reactive protein = 7 mg/L). A chest radiography suggested the presence of an aspergilloma, with an image compatible with an Aspergillus ball into the pulmonary cavern.

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MATERIALS AND METHODS

Strains

A Fusarium sp. strain was isolated from BAL fluid specimen of the patient in French Guiana and submitted to the WI, under accession number CBS 143874. Ex-type and reference strains spanning the known diversity of the FFSC were selected based on phylogenetic and morphological similarity to the strain under study (Table 1). The clinical isolate was grown on PDA slants for 7 d at 25 °C, and was maintained as working culture or stored in 20 % (v/v) glycerol at -80 °C for prolonged use.

Table 1. GenBank accession numbers of Fusarium species included in this study.

| Species          | Collection | Source               | Country     | cmdA         | rpb2         | tef1         | tub          |
|------------------|------------|----------------------|-------------|--------------|--------------|--------------|--------------|
| F. agapanthi     | NRRL 54463 | Agapanthus praecox   | Australia   | KU900611     | KU900625     | KU900630     | KU900635     |
| F. ananatum      | CBS 118516 | Ananas comosus fruit | South Africa| LT996175     | LT996137     | LT996091     | LT996112     |
| F. andiyazi      | CBS 119857 | Sorghum bicolor soil debris | South Africa | LT996176 | LT996138 | LT996092 | LT996113 |
| F. anthophilum   | CBS 737.97 | Hibiscus sp.         | Germany     | LT996177     | LT996139     | LT996093     | LT996114     |
| F. bactridioides | CBS 20476  | Cronartium conigenum | USA         | AF158343     | -            | AF160290     | U34434       |
| F. begoniae      | CBS 403.97 | Begonia elatior hybrid | Germany     | AF158346     | LT996140     | AF160293     | U61543       |
| F. bulbicola     | CBS 220.76 | Nerine bowdennii     | Germany     | KF466327     | KF466404     | KF466415     | KF466437     |
| F. circinatum    | CBS 405.97 | Pinus radiata        | USA         | KF466327     | KF466404     | KF466415     | KF466437     |
| F. coecis        | NRRL 66233 | Coix gasteenii       | Australia   | KM231393     | KM068354     | KM231943     | KM232080     |
| F. ficirescens   | CBS 125181 | Ficus carica fruit   | Iran        | KU603958     | KT154002     | KU604452     | KP662896     |
| F. lactis        | CBS 411.97 | Ficus carica         | USA         | AF158325     | LT996149     | AF160272     | U61551       |
| F. mexicanum     | NRRL 31630 | Capsicum anuum      | Belgium     | FR870301     | FR870313     | FR870289     | FR870325     |
| F. nygamai       | NRRL 53580 | Mangifera indica  | Mexico      | GU737389     | -            | GU737416     | GU737308     |
| F. oxysporum     | CBS 716.74 | Vicia faba           | Germany     | AF158366     | JX171583     | AF008479     | U34435       |
| F. phyllophilum  | CBS 744.97 | Pseudotsuga menziesii | USA         | AF158365     | LT750605     | AF160312     | U34424       |
Table 1. (Continued).

| Species                  | Collection | Source                  | Country          | GenBank/ENA accession number |
|--------------------------|------------|-------------------------|------------------|-----------------------------|
| *F. pseudocircinatum*    | CBS 449.97 | Solanum sp.             | Ghana            | AF158324 LT996151 AF160271  |
| *F. pseudonymgamaei*     | CBS 417.97 | Pennisetum typhoides    | Nigeria          | AF158316 LT996152 AF160263  |
| *F. ramigemum*          | CBS 418.98 | Ficus carica            | USA              | KF466335 KF466412 KF466423  |
| *Fusarium* sp.          | NRRL 25346 | Pine pitch canker       | USA              | AF158349 - AF160296 U61642   |
| *F. subglutinans*        | CBS 747.97 | Zea mays                | USA              | AF158342 JX171599 AF160289  |
| *F. sudanense*          | CBS 454.97 | Striga hermonthica      | Sudan            | LT996185 LT996155 KU711697  |
| *F. temperatum*         | NRRL 25622 | Zea mays                | South Africa     | AF158354 Not public AF160307 |
| *F. terricola*          | CBS 483.94 | Soil                    | Australia        | KU603951 LT996156 KU711698  |
| *F. thapsinum*          | CBS 733.97 | Zea mays                | South Africa     | LT996186 XJ171600 AF160270  |
| *F. tjaetaba*           | NRRL 66243 | Sorghum interjectum     | Australia        | LT996187 KPO83275 KPO83263  |
| *F. udum*               | NRRL 22949 | Lactarius pubescens     | Germany          | AF158328 LT996172 AF160275  |
| *F. verticillioides*     | CBS 734.97 | Zea mays                | Germany          | AF158315 EF470122 AF160262  |
| *F. xylarioides*         | CBS 115135 | Human                   | Sweden           | KU603944 KU604217 KU604384  |
| *F. volatile*           | CBS 143874 | Human bronchoalveolar lavage fluid | French Guiana | MK984595 LR596006 LR596007 |
| *F. xylarioides*         | CBS 143874 | Human bronchoalveolar lavage fluid | French Guiana | MK984595 LR596006 LR596007 |
| *F. xylarioides*         | NRRL 25615 | Oryza sativa seed       | Nigeria          | AF158357 - AF160304 AF160348 |
| *F. werrikimbe*          | CBS 125535 | Sorghum leiacadum       | Australia        | - - EF107131 EF107133 |
| *F. xylarioides*         | F19361     | Sorghum leiacadum       | Australia        | - - EF107131 EF107133 |
| *F. xylarioides*         | NRRL 25486 | Coffea trunk            | Ivory Coast      | - HM068355 AY707136 AY707118 |

1 CBS: Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands. F: University of Sydney, Sydney, New South Wales, Australia. NRRL: Agricultural Research Service Culture Collection, National Center for Agricultural Utilization Research, USDA, Peoria, IL, USA. 2 ENA: European Nucleotide Archive. cmdA: calmodulin. rpb2: RNA polymerase second largest subunit. tef1: translation elongation factor–alpha. tub: β-tubulin. Sequences marked as Not public are available on the sequence datasets published by Edwards et al. (2016).

Morphology

CBS 143874 was characterised morphologically following procedures described elsewhere (Aoki et al. 2013, Leslie & Summerell 2006, Sandoval-Denis et al. 2018). Colony growth rates and production of diffusible pigments were evaluated on PDA, colony features were also recorded on malt extract agar (MEA) and oatmeal agar (OA). Colour notations followed those of Rayner (1970). Micro-morphological features were studied from cultures grown for 7–10 d at 24 °C, using a 12 h light/dark cycle with near-UV and white fluorescent light. Features of the mycelium scraped from the surface of 7-d-old cultures on MEA were carried out using three independent algorithms: Maximum-Likelihood (ML), Maximum Parsimony (MP) and Bayesian

DNA extraction, amplification and sequencing

The Wizard® Genomic DNA purification Kit (Promega, Madison, WI, USA) was used to extract total genomic DNA from fresh mycelium scraped from the surface of 7-d-old cultures on MEA at 24 °C. Partial fragments of four loci were PCR-amplified following previously published protocols using the following primer pairs: BT-2a/BT-2b for the β-tubulin gene (tub) (Glass & Donaldson 1995), CL1/CL2 for the calmodulin gene (cmdA) (O’Donnell et al. 2009), EF-1/EF-2 for the translation elongation factor–alpha gene (tef1) (Glass et al. 2010), and RPB2-5f2/7cr plus RPB2-7cf/11ar for two non-contiguous fragments of the RNA polymerase second largest subunit (rpb2) (Liu et al. 1999, Sung et al. 2007). Sequencing was done in both directions using the respective PCR primers on an Applied Biosystems 3730xl DNA Analyzer (Life Technologies, Carlsbad, CA, USA). The DNA sequences were analysed and consensus sequences were assembled using SeqMan Pro v. 13 (DNASTar, Madison, WI, USA).

Phylogenetic analyses

Phylogenetic analyses of single loci and the combined dataset were carried out using three independent algorithms: Maximum-Likelihood (ML), Maximum Parsimony (MP) and Bayesian

Cardinal growth temperatures were determined on MEA and PDA plates incubated in the dark for 2 wk at temperatures of 18–40 °C at intervals of 3 °C; with two replicates for each isolate.
Table 2. MIC values of clinical isolate CBS 143874 (µg/mL).

| Strain   | AMB | FLC | ITC | VOR | POS | ISA | ANI | MICA |
|----------|-----|-----|-----|-----|-----|-----|-----|------|
| CBS 143874 | 1   | >64 | >16 | 1   | 0.5 | 4   | >8  | >8   |

AMB: amphotericin B. FLC: fluconazole. ITC: itraconazole. VOR: voriconazole. POS: posaconazole. ISA: isavuconazole. ANI: anidulafungin. MICA: micafungin.

inference (BI). Both ML and BI were run on the CIPRES Science Gateway portal (Miller et al. 2012). Evolutionary models were calculated with MrModelTest v. 2.3 using the Akaike information criterion (Nylander 2004). For ML, RAxML-HPC2 v. 8.2.10 on XSEDE was used (Stamatakis 2014), with a bootstrap analysis (BS) based on default parameters. The BI analyses were run using MrBayes v. 3.2.6 on XSEDE (Ronquist & Huelsenbeck 2003) using four incrementally heated MCMC chains for 5 M generations and a sample frequency of every 1 000 trees. The 50 % consensus trees and posterior probabilities (PP) values were calculated after discarding the first 25 % of samples as burn-in. Maximum-parsimony analyses were run using PAUP v. 4.0b10 (Swofford 2003). Heuristic searches included 1 000 random stepwise addition replicates, with tree bisection and reconstruction (TBR) branch swapping; all characters were equally weighted and gaps treated as missing data. Branches of zero length were collapsed and all multiple, equally parsimonious trees were saved. Tree statistics [tree length (TL), consistency index (CI), retention index (RI) and rescaled consistency index (RC)] were calculated. Clade stability was evaluated using a bootstrap analysis (BS) of 1 000 replicates.

Antifungal susceptibility

Antifungal susceptibility testing of CBS 143874 was performed by the CLSI broth microdilution as described in the CLSI document M38-A2 (Clinical and Laboratory Standards Institute 2008) with modifications according to Al-Hatmi et al. (2015). The following drugs were used: amphotericin B (Sigma-Aldrich), fluconazole (Pfizer, Groton, CT, USA), itraconazole (Janssen Pharmaceutica, Tilburg, The Netherlands), voriconazole (Pfizer), posaconazole (Merck), isavuconazole (Basilea Pharmaceutica, Basel, Switzerland), micafungin (Astellas, Ibaraki, Japan), and anidulafungin (Pfizer). Three reference strains (Paeilomyces variotii ATCC 22319, Candida krusei ATCC 6258, and Candida parapsilosis ATCC 22019) were included as quality controls.

RESULTS

Molecular analyses

Topologies obtained from the analyses of the majority of individual gene datasets were congruent, differing only in the positions of unsupported nodes. Individual analyses of the cmdA, rpb2, tef1 and tub loci consistently resolved Fusarium sp. CBS 143874 as a member of the African clade sensu O’Donnell et al. (1998) (Fig. 1). Phylogenies based on cmdA, rpb2 and tef sequences showed that strain CBS 143874, together with an unidentified isolate (NRRL 25615) formed a genetically exclusive lineage, phylogenetically related to F. coicus and F. verticilloides. In contrast, the tub phylogeny failed to unambiguously identify most of the Fusarium species included in this study, showing mostly marginal statistical support values that did not allow unequivocal separation of lineages. Nevertheless, this locus confirmed the phylogenetic position of CBS 143874 and NRRL 25615 as close relatives of F. coicus. With the exception of tub, single locus phylogenies did not support the monophyly of the African clade of FFSC, which resolved as polyphyletic.

The final, combined alignment encompassed a total of 3 305 positions (cmdA 545, rpb2 1 591, tef1 680, tub 489), of which 696 sites were variable (cmdA 86, rpb2 295, tef1 194, tub 121) and 452 were phylogenetically informative (cmdA 61, rpb2 204, tef1 119, tub 68). The combined phylogeny confirmed the results of the individual phylogenetic analyses, Fusarium sp. strains CBS 143874 and NRRL 25615 formed a highly-supported, genetically exclusive group, closely related to but different from F. coicus and F. verticilloides (Fig. 2); consequently, the above-mentioned phylogenetic clade is proposed here as the new species Fusarium variotii.

Antifungal susceptibility testing

Antifungal susceptibility testing according to CLSI M38A (Clinical and Laboratory Standards Institute 2008) demonstrated that Fusarium variotii had a low MIC of 1 µg/mL against amphotericin B, voriconazole (1 µg/mL), and posaconazole (0.5 µg/mL), whereas the fungus had high MICs for fluconazole (>64 µg/mL), itraconazole (>16 µg/mL), isavuconazole (4 µg/mL), anidulafungin (>8 µg/mL), and micafungin (>8 µg/mL) (Table 2).

Taxonomy

Fusarium variotii Al-Hatmi, Sand.-Den., S.A. Ahmed & de Hoog, sp. nov. MycoBank MB831243. Figs 3, 4.

Etymology: Named after its supposedly airborne entry in the human patient.

Typus: French Guiana, Cayenne, bronchoalveolar lavage (BAL) effusion of patient with lung infection, 2017, M. Demar (holotype CBS H-24004, culture ex-type CBS 143874).

Mycelium consisting of hyaline, smooth, branched, 1.5–5.5 μm diam hyphae. Aerial conidiophores erect, often reduced to conidiogenous cells, borne laterally on hyphae, less commonly irregularly or verticillately branched, up to 80 μm long, rarely proliferating, bearing terminal single mono- and polyphialides; aerial phialides subulate to subcylindrical, smooth- and thin-walled, (7–)11.5–21(–29) × 2–4.5 μm (av. 16.2 × 3.3 μm), commonly bearing a single conidiogenous locus, rarely polyphialidic, lacking noticeable periclinal thickening or collarettes; aerial conidia ellipsoid, allantoid, club-shaped to somewhat cylindrical, often with a flattened base, straight or gently curved, smooth- and thin-walled, 0–2(–3)-septate, (4.5–)5.5–19(–30) × (1.5–)2–4(–5.5) μm (av. 12.4 × 3.1 μm), grouped in moderately long, straight or flexuous chains, quickly collapsing to form discrete conidial heads; mycrocyst conidiation present. Sporodochia produced infrequently in the aerial mycelium, rarely on the surface of carnation leaves, tan
Fusarium volatile from bronchoalveolar lavage

Fig. 1. Maximum Parsimony (MP) trees obtained from the individual phylogenetic analyses of the cmdA, rpb2, tef1 and tub datasets of representative isolates of the *Fusarium fujikuroi* species complex. Numbers on the nodes are MP and Maximum-Likelihood (ML) bootstrap values (BS) above 70 % and Bayesian posterior probability values (PP) above 0.95. Thickened branches indicate full statistical support (MP-BS, ML-BS = 100 % and PP = 1). Coloured branches indicate the African (red) and American (blue) clades according to O’Donnell et al. (1998). The clinical isolate is highlighted in **bold**. Ex-type and ex-neotype strains are indicated with T and NT, respectively. The trees are rooted with *Fusarium oxysporum* CBS 744.97 and CBS 716.74.

**TL** = tree length, **CI** = consistency index, **RI** = retention index, **RC** = rescaled consistency index.

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to pale orange; *sporodochial conidiophores* simple or sparingly irregularly branched, 16–21(–22) × (3–)3.5–4.5 μm (av. 18.6 × 3.8 μm), bearing terminal and lateral single monophialides or terminal whors of up to 4 monophialides; *sporodochial phialides* doliiform to subcylindrical, smooth- and thin-walled, (9.5–)11–15(–19) × (2.5–)3–4(–4.5) μm (av. 12.9 × 3.6 μm),
Fig. 2. The first of 16 most parsimonious trees obtained from the combined cmdA, rpb2, tef1 and tub sequences of 43 strains belonging to the *Fusarium fujikuroi* species complex (FFSC). Numbers on the nodes are MP and Maximum-Likelihood (ML) bootstrap values (BS) above 70 % and Bayesian posterior probability values (PP) above 0.95. Thickened branches indicate full statistical support (MP-BS, ML-BS = 100 % and PP = 1). Coloured branches indicate the African (red) and American (blue) clades according to O’Donnell et al. (1998). Isolates and name of the new species are highlighted in **bold**. Ex-type and ex-neotype strains are indicated with T and NT, respectively. The tree is rooted with *Fusarium oxysporum* CBS 744.97 and CBS 716.74. TL = tree length, CI = consistency index, RI = retention index, RC = rescaled consistency index.

Fig. 3. *Fusarium volutile* (ex-type CBS 143874). A–C. Colonies (left obverse, right reverse) on MEA, PDA and OA, respectively, after 14 d at 24 °C. D–H. Aerial conidiophores and chains of conidia. I–L. Aerial phialides. M. Sporodochial phialides. N. Aerial conidia showing microcyclic conidiation. O, P. Aerial conidia. Q. Sporodochial conidia. Scale bars: D–H = 20 µm, I–L = 5 µm, all others = 10 µm.
sporodochial conidia falcate, straight or dorsiventrally curved, curvature often more distinctly pronounced on the dorsal line, tapering toward the basal part; apical cell conical and slightly hooked; basal cell foot-shaped, (0–)1–3-septate, hyaline, thin- and smooth-walled. Aseptate conidia (14–)15–20.5 × 3–4 μm (av. 17.8 × 3.5 μm), 1-septate conidia (17–)19–24.5(–27) × (2–)3–4 μm (av. 21.7 × 3.5 μm), 2-septate conidia (26–)27–30(–31) × 3.5–4 μm (av. 28.4 × 3.9 μm), 3-septate conidia (27–)29–34.5(–36.5) × (3–)3.5–4(–4.5) μm (av. 31.7 × 4 μm); overall (14–)21–32.5(–36.5) × (2–)3–4.5 μm (av. 26.6 × 3.8 μm). Chlamydospores not observed.

Colonies growing in the dark after 7 d at 24 °C. On MEA reaching 50–68 mm diam, white, salmon to peach; colony surface raised to slightly umbonate, velvety felty; margin undulate to filiform; reverse orange to luteous with pale luteous periphery. On OA reaching 65–74 mm diam, saffron to rosy buff, turning pale vinaceous toward the periphery, membranous; margin entire with abundant submerged mycelium; reverse saffron to salmon, pale orange at the centre. On PDA with an average radial growth rate of 4.7–5.7 mm/d, reaching 60–80 mm diam, salmon, saffron to pale ochraceous with peach centre; flat with slightly raised centre, felty to cottony; margin irregular, undulate to lobate; reverse pale orange to ochraceous.

Cardinal growth temperatures: optimal development at 27–33 °C, minimum 18 °C, maximum 37 °C. The species was still able to grow at 37 °C, but not at 40 °C.

Notes: Fusarium volatile is phylogenetically closely related to F. coicus and F. verticillioides. The three mentioned species share the common morphological features attributed to the FFSC, such as the lack of chlamydospores, formation of oval to clavate microconidia and presence of monophialides, while sporodochia are not commonly produced. However, F. coicus and F. volatile differ significantly from F. verticillioides by having up to 3-septate microconidia (usually aseptate in F. verticillioides; Laurence et al. 2015).

Fig. 4. Fusarium volatile (ex-type CBS 143874). A–E. Aerial conidiophores. F. Sporodochial conidiophore. G. Sporodochial conidia. H. Aerial conidia. Scale bar = 10 µm.
### Key to species of the *Fusarium fujikuroi* species complex known from human clinical specimens

| 1. Polyphialides present | 2. |  
|--------------------------|---|---
| 1. Polyphialides absent | 6. |
| 2. Microconidial chains present | 3. |
| 2. Microconidial chains absent | 8. |
| 3. Sporodochia orange | 4. |
| 3. Sporodochia tan to pale orange | 10. |
| 4. Chlamydospores present | \( F. \text{ngamai} \) |
| 4. Chlamydospores absent | 5. |
| 5. Pyriform conidia present | \( F. \text{fujikuroi} \) |
| 5. Pyriform conidia absent | \( F. \text{ramigenum} \) |
| 6. Napiform conidia present | 7. |
| 6. Napiform conidia absent | 11. |
| 7. Chlamydospores present | \( F. \text{napiforme} \) |
| 7. Chlamydospores absent | \( F. \text{thapsinum} \) |
| 8. Conidiophores mainly prostrate, rarely branched | 9. |
| 8. Conidiophores mainly erect, branched | 13. |
| 9. Chlamydospores present, polyphialides rare | \( F. \text{acutatum} \) |
| 9. Chlamydospores absent, polyphialides abundant and proliferating extensively | \( F. \text{sacchari} \) |
| 10. Microconidia 0-septate; macroconidia 3–5-septate, straight or almost so | \( F. \text{proliferatum} \) |
| 10. Microconidia 0–2–3-septate; macroconidia 0-3-septate, gently curved | \( F. \text{volatilae} \) |
| 11. Microconidia in chains only | \( F. \text{verticillioides} \) |
| 11. Microconidia in head and chains | 12. |
| 12. Macroconidia present, 3–6-septate | \( F. \text{andiyazi} \) |
| 12. Macroconidia absent | \( F. \text{musae} \) |
| 13. Globose microconidia present | 14. |
| 13. Globose microconidia absent | 15. |
| 14. Colonies on PDA orange | 16. |
| 14. Colonies on PDA purple or violet | \( F. \text{anamnatum} \) |
| 15. Macroconidia formed only on aerial mycelium, sporodochia not produced | \( F. \text{temperatum} \) |
| 15. Macroconidia formed on sporodochia | \( F. \text{guttiforme} \) |
| 16. Microconidia 0-septate, macroconidia rare or absent | \( F. \text{subglutinans} \) |
| 16. Microconidia 0–1-septate, macroconidia abundant |

### DISCUSSION

Fusariosis is usually acquired by inhalation of conidia or after trauma, skin burns, or sometimes through central venous access, or at lower incidence the gastrointestinal tract after consumption of contaminated food (Carneiro et al. 2011, Muhammed et al. 2011). *Fusarium* species can affect humans either by infection (Al-Hatmi et al. 2016b) or by mycotoxicosis (Marasas et al. 1984). Clinically, a disseminated fusariosis is characterised by persistent sepsis despite broad-spectrum antibiotic therapy. Although all organs may be concerned, cutaneous involvement is predominant, followed by pulmonary infection. Given that they are widespread in natural and human-made environments, *Fusarium* species may contaminate laboratory specimens and yield false-positive responses. The interpretation of *Fusarium* growth from clinical materials strongly depends on the clinical context (Nucci & Anaissie 2007). However, repeated isolation of the fungus and culture from sinus aspirate or deep respiratory secretions in severely immunocompromised hosts should always be considered as diagnostic of fusariosis (Nucci & Anaissie 2007).

In our study, we isolated a novel *Fusarium* species from broncho-alveolar lavage (BAL) aspirate. We cannot definitively differentiate between a fungal colonisation or an environmental contamination as the novel species was isolated in a single BAL sample and as the patient did not have any pulmonary clinical symptoms and evolved positively without antifungal treatment. Nevertheless, the chest radiography suggested the presence of an aspergilloma, given a suspicious image compatible with a Monod sign into the pulmonary cavern that could have been attributed to the presence of *Fusarium*, as deep infections by *Fusarium* and *Aspergillus* spp. may be confused. These opportunists present similar radiologic results, share comparable histologic appearances with hyaline, septate, branched hyphae and can cause similar clinical syndromes (Hayden et al. 2003). It is, therefore, important to make the correct diagnosis to optimise treatment and improve prognosis. Similarly, the correct species-level identification for *Fusarium* infections is crucial for a positive outcome since different infectious species may present marked differences in their antifungal susceptibility patterns, at least in vitro (Al-Hatmi et al. 2015).

Our phylogenetic analysis based on a four gene dataset (cmdA, rpb2, tef and tub) showed that *F. volatile* belongs to the FFSC, where it formed a genetically exclusive, strongly supported monophyletic clade, phylogenetically related to *F. coicus* and *F. verticillioides*. The phylogenetic clade representing *F. volatile* had already been recognised as a distinct species in FFSC, however, it was not formally described as such (O’Donnell et al. 2000). The FFSC is a species-rich group, currently comprising more than 50 taxa, including opportunists on humans, economically relevant plant pathogens, and mycotoxin producers. Most of these species have been recognised based on phylogenetic analyses (Kvas et al. 2009). A large number of cryptic, unnamed
phylogenetic species remain to be formally described. Based on phylogenetic relationships, O’Donnell et al. (1998) organised the FFSC into three main clades (African, American and Asian clades), each encompassing numerous species with similar biogeographic patterns. Our results showed that, despite being isolated in South America, *F. volatile* clusters in the African clade, which matches with the African origin of a genetically identical strain (NRRL 25615) from Nigeria. Interestingly, our individual phylogenies do not support the current delimitation of the African clade of FFSC, which was found to be polyphyletic using *cmdA*, *rpb2* and *tef1* markers. The phylogenies support the results previously reported by other authors using the same and additional phylogenetic markers (Kvas et al. 2009, Walsh et al. 2010, O’Donnell et al. 2013, Laurence et al. 2015, Sandoval-Denis et al. 2018), which shows that the biogeographic clade distribution of the FFSC needs further re-evaluation.

The voucher strain CBS 143874 of *F. volatile* clusters as a sister clade to *F. coicis* and *F. verticillioides*, the latter species being known to be capable of causing human infection (Al-Hatmi et al. 2016b). *Fusarium verticillioides* is an important producer of mycotoxins, including fumonisins (Leslie & Summerell 2006, Rosa Junior et al. 2019); these toxins are highly detrimental to animals and are suspected to be responsible for acute and chronic human diseases (Leslie & Summerell 2006). In contrast, *F. coicis* is known only from *Coix gasteeni* (*Poaceae*), a rare Australian grass species, while no human infection or toxin production has been reported. These clinical data suggest that the new species *F. volatile* is unlikely a primary human pathogen but rather an opportunist that takes advantage of the host’s compromised immune response. Production of mycotoxins, however, was not tested in *F. volatile* and remains to be studied.

Among the FFSC species known to occur on humans (de Hoog et al. 2019), *F. volatile* is morphologically similar to *F. fujikuroi*, *F. nygamai*, *F. proliferatum* and *F. ramigenum*. All the latter species are characterised by obovate to clavate microconidia formed in false heads and chains from mono- and polyphialides, and by, except *F. nygamai*, absence of chlamydospores. *Fusarium volatile* can be recognised by its ellipsoid to cylindrical microconidia (vs. the pyriform microconidia of *F. fujikuroi* and *F. proliferatum*, and the obovate microconidia of *F. ramigenum*), its less-septate macroconidia (up to 3-septate vs. up to 5-septate in all the species listed above), and its tan to pale orange sporodochia (orange in *F. fujikuroi* and *F. nygamai*) (Leslie & Summerell 2006). Two additional opportunistic species on humans, *F. acutatum* and *F. sacchari*, produce microconidia only on false heads, but might also be confused with *F. volatile* because of their similar morphology and septation of macroconidia. *Fusarium volatile* differs from *F. acutatum* by its 0–3-septate microconidia (0-septate in the latter species) and its less curved macroconidia. It can be separated from *F. sacchari* by the microconidial shape (oval in *F. sacchari*), while polyphialides are rarely seen in *F. volatile* (common in *F. sacchari*).

Antifungal susceptibility testing demonstrated that *F. volatile* had reduced MICs for posaconazole (0.5 µg/mL), followed by amphotericin B and voriconazole (1 µg/mL), whereas MIC values for fluconazole, isavuconazole, itraconazole and the echinocandins were elevated (Table 2). In the present case, no antifungal therapy was given. In general, *Fusarium* spp. show a remarkably high degree of intrinsic resistance to a wide spectrum of clinically available antifungal drugs. Prior to the voriconazole era, the initial approach of treating invasive fusariosis consisted on the administration of high-dose (>5 mg/kg/d) liposomal amphotericin B (Al-Hatmi et al. 2017). After FDA approval in 2002, voriconazole has become the first-line treatment because of its lower toxicity and higher clinical efficacy against fusariosis (Walsh et al. 1998, Stempel et al. 2015). The European Fungal Infection Study Group and the European Confederation of Medical Mycology recommended a lipid formulation of amphotericin B or voriconazole for treating invasive fusariosis (Tortorano et al. 2014).

In conclusion, we described the new species *F. volatile*, belonging to the FFSC. This new taxon was found in a BAL sample from a patient with non-haematological predisposing conditions. Further studies are required to determine natural ecology, transmission routes and the potential pathogenic role of this new species.

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