New *Fusarium* species from the Kruger National Park, South Africa

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**Abstract**

Three new *Fusarium* species, *F. convolutans*, *F. fredkrugeri*, and *F. transvaalense* (Ascomycota, Hypocreales, Nectriaceae) are described from soils collected in a catena landscape on a research supersite in the Kruger National Park, South Africa. The new taxa, isolated from the rhizosphere of three African herbaceous plants, *Kyphocarpa angustifolia*, *Melhania acuminata*, and *Sida cordifolia*, are described and illustrated by means of morphological and multilocus molecular analyses based on sequences from five DNA loci (CAL, EF-1\(\alpha\), RPB1, RPB2 and TUB). According to phylogenetic inference based on Maximum-likelihood and Bayesian approaches, the newly discovered species are distributed in the *Fusarium buharicum*, *F. fujikuroi*, and *F. sambucinum* species complexes.

**Keywords**

Natural parks, phylogeny, fungi, multigene, morphology, diversity

**Introduction**

Fungi are common colonisers of the plant rhizobiome and endosphere, where they play a key role in modulating the interactions between plant roots and soil (Zachow et al. 2009; Visioli et al. 2014). The direct and indirect interaction between fungal growth in the rhizosphere and its effect on plant growth and health is well docu-
mented (Havlicek and Mitchell 2014; Hargreaves et al. 2015; Lareen et al. 2016). Such effects include either a positive feedback by producing plant growth promoting factors, solubilising and stimulating nutrient uptake by plant roots or by inhibiting the growth of concomitant pathogenic organisms (Schippers et al. 1987; Mommer et al. 2016). Conversely, deleterious effects have also been observed, either related to the presence of pathogenic fungal species or caused by fungal-induced modifications of plant root functions, impeding root growth or negatively altering nutrient availability (Schippers et al. 1987; Mommer et al. 2016). Likewise, plants can select and harbour a particular fungal community on its roots via root exudates (Lareen et al. 2016; Sasse et al. 2018), while abiotic influences including water availability, climate and season, soil type, grazers and other animals, orchestrate the development of a unique fungal diversity (Philippot et al. 2013; Havlicek and Mitchell 2014; Hargreaves et al. 2015; Lareen et al. 2016).

The genus *Fusarium* Link (Hypocreales, Nectriaceae) includes a vast number of species, commonly recovered from a variety of substrates including soil, air, water and decaying plant materials; being also able to colonise living tissues of plants and animals, including humans; acting as endophytes, secondary invaders or becoming devastating plant pathogens (Nelson et al. 1994). In addition to their ability to colonise a multiplicity of habitats, *Fusarium* is a cosmopolitan genus, present in almost any ecosystem in the world, including human-made settings such as air and dust in the indoor environment or even in hospitals (Perlroth et al. 2007; Aydogdu and Asan 2008; Pinheiro et al. 2011).

Being common inhabitants of plant root ecosystems, fusaria and, particularly *Fusarium graminearum* Schwabe, *F. proliferatum* (Matsush.) Nirenberg ex Gerlach & Nirenberg, *F. verticillioides* (Sacc.) Nirenberg (Syn. *F. moniliforme* J. Sheld.), *F. oxysporum* Schltdl., as well as species recently segregated from *Fusarium*, including *Neocosmospora phaseoli* (Burkh.) L. Lombard & Crous (Syn. *Fusarium phaseoli* Burkh.) and *N. virguliforme* (O’Donnell & T. Aoki) L. Lombard & Crous (Syn. *F. virguliforme* O’Donnell & T. Aoki), have been regularly studied for their interactions with the rhizobiome, motivated mainly by the importance of these organisms as soil-borne plant pathogens and the need to develop effective control mechanisms (Larkin et al. 1993; Hassan Dar et al. 1997; Pal et al. 2001; Fravel et al. 2003; Idris et al. 2006; Díaz Arias et al. 2013). Similarly, abundant data is available regarding the ecology and distribution of plant-associated fusaria, particularly related to pathogenic species or commonly isolated endophytes (Leslie and Summerell 2006). Little attention has however been given to the occurrence of non-pathogenic fungal species, including *Fusarium* spp. in root microbial communities (Zakaria and Ning 2013; Jumpponen et al. 2017; LeBlanc et al. 2017), while comprehensive DNA sequence-based surveys have been directed mostly to the study of highly relevant and abundant rhizosphere fungal genera such as *Trichoderma* Pers., *Verticillium* Nees or mycorrhizal fungi (Zachow et al. 2009; Bent et al. 2011; Ruano-Rosa et al. 2016; Saravanakumar et al. 2016).
The Kruger National Park (KNP) in South Africa is one of the largest natural reserves in Africa, encompassing a number of non-manipulated landscapes, with almost no human alteration (Carruthers 2017). Recently, four research “supersites” have been identified and established in KNP, each of these supersites representing unique geological, ecological and climatic features of the park (Smit et al. 2013). A multidisciplinary study was conducted in KNP aimed to determine functioning and interaction between abiotic and biotic components, as well as soil properties, hydrology and other processes that determine the structure, biodiversity and heterogeneity of a catena or hill slope ecosystem on one of these “supersites”, located deep inside the KNP (data not published). In order to assess the microbial soil population and community dynamics, mainly focused on bacteria, several rhizosphere samples were obtained from diverse African plants on one of these exceptional protected savannah landscapes. From these collections, interesting fusaria were isolated from the root ecosystem of three native African herbaceous plants i.e. *Kyphocarpa angustifolia* (Moq.) Lopr. (Amaranthaceae), *Melhania acuminata* Mast. (Malvaceae) and *Sida cordifolia* Linn. (Malvaceae). According to their unique morphological traits and clear phyllogenetic delimitations, these isolates are described here as three new *Fusarium* species.

**Methods**

**Study site and sampling**

During March 2015, rhizosphere soil from three herbaceous plants was collected in the Southern Granites “supersite” catena (Stevenson-Hamilton supersite) in the KNP, between 25°06’28.6S, 31°34’41.9E and 25°06’25.7S, 31°34’33.7E (Fig. 1). A catena consists of different soil types observed from a crest to a valley bottom with a wetland or drainage exhibiting different water retention capabilities due to the slope or aspect (topography) and the depth of underlying geological rocks (Brown et al. 2004, Van Zijl and Le Roux 2014). The main characteristics of the Stevenson-Hamilton supersite are described in detail by Smit et al. (2013). Briefly, in this site, a single catena landscape covers approximately 1 km from top to bottom and consists of a hill slope, a sodic site (or grazing lawn), a riparian and floodplain area and a dry drainage line. Three species of plants were selected for sampling occurring at the two extremes of the catena. Two of these species (*Kyphocarpa angustifolia* and *Sida cordifolia*) occurred at both top and bottom sites while *Melhania acuminata* only occurred at the top site. The soil (100 mm depth) at the top of the slope is Clovelly with a high percentage of sand (90%) and a low cation exchange capacity (CEC) (mean sodium concentration of 1062 mg/kg) and pH (mean 5.85). The soil at the bottom of the slope is of the Sterkspruit type, with higher clay content thus higher CEC (mean sodium concentration of 3802 mg/kg) and higher pH (mean...
Figure 1. Map of the Kruger National Park (KNP) in South Africa. The arrows indicate the location of the four research “supersites” (adapted from Smit et al. 2013). Sampling site is indicated with a black star. The inset shows the location of the KNP within South Africa, indicated by a grey box.

6.4). Rhizosphere soil of 10 plants of the same species occurring at each top or bottom site was sampled using a core soil sampler. A total of 50 samples consisting of ca. 200 g of soil from the roots of each plant were taken, deposited in zip-lock plastic bags and kept on ice in a cool bag at approximately 5 °C until analysed in the laboratory.

Isolation of *Fusarium* strains

Soil samples were mixed thoroughly and sieved to remove large elements. Fine soil particles were uniformly spread and distributed over the surface of pentachloronitrobenzene agar (PCNB; also known as the Nash-Snyder medium, recipe in Leslie and Summerell 2006) supplemented with streptomycin (0.3 g/l) and neomycin sulphate (0.12 g/l) and malt-extract agar (MEA; recipes on Crous et al. 2009) on 9 mm Petri dishes and incubated at 24 °C for 10 d under a natural day/night photo-period. Each soil sample was processed in duplicate. Fungal growth was evaluated daily and growing colonies were transferred to fresh Potato Dextrose Agar (PDA;
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Colonies were evaluated for their macro- and microscopic characteristics and a total of 19 fungal cultures showing features typical of *Fusarium* were subjected to single spore isolation as described previously (Sandoval-Denis et al. 2018). Single spore isolates were finally transferred and maintained in Oatmeal Agar plates and slants (OA; recipe in Crous et al. 2009). Fungal strains isolated in this study were deposited in the collection of the Westerdijk Fungal Biodiversity Institute (CBS; Utrecht, the Netherlands), the working collection of Pedro W. Crous (CPC), held at CBS (Table 1); and voucher specimens were deposited in The South African National Collection of Fungi (NCF) (Mycology Unit, Biosystematics Division, Plant Protection Institute, Agricultural Research Council, Pretoria, South Africa).

**Morphological characterisation**

*Fusarium* isolates were characterised morphologically according to procedures described elsewhere (Aoki et al. 2013; Leslie and Summerell 2006, Sandoval-Denis et al. 2018). Colonial growth rates and production of diffusible pigments were evaluated on PDA, colony features were also recorded on corn-meal agar (CMA; recipe in Crous et al. 2009) and OA. Colour notations followed those of Rayner (1970). For the study of micro-morphological features, cultures were grown for 7–10 d at 24 °C, using a 12 h light/dark cycle with near UV and white fluorescent light. Aerial and sporodochial conidiophores and conidia and formation of chlamydospores were evaluated on Synthetic Nutrient-poor Agar (SNA; Nirenberg 1976) and on Carnation Leaf Agar (CLA; Fisher et al. 1982). Measurements and photomicrographs were recorded from a minimum of 30 elements for each structure, using sterile water as mounting medium and a Nikon Eclipse 80i microscope with Differential Interference Contrast (DIC) optics and a Nikon AZ100 dissecting microscope, both equipped with a Nikon DS-Ri2 high definition colour digital camera and the Nikon software NIS-elements D software v. 4.30.

**DNA isolation, amplification and sequencing**

Isolates were grown for 7 d on MEA at 24 °C using the photoperiod described above. Fresh mycelium was scraped from the colony surface and subjected to total DNA extraction using the Wizard® Genomic DNA purification Kit (Promega Corporation, Madison, WI, USA), according to the manufacturer’s instructions. Fragments of five DNA loci were amplified using primers and PCR conditions described by O’Donnell et al. (2009) for calmodulin (*CAL*), O’Donnell et al. (2010) for the RNA polymerase largest subunit (*RPBI*) and second largest subunit (*RPB2*), O’Donnell et al. (1998) for the translation elongation factor 1-alpha (*EF-1α*) and Woudenberg et al. (2009) for
### Table 1. Origin, strain and GenBank/ENA accession number of strains and DNA sequences included in this study.

| Species name          | Strain\(^t\) | Country     | Host                           | Sequence accession number\(^b\) |
|-----------------------|--------------|-------------|--------------------------------|---------------------------------|
| *Fusarium agapanthi*  | NRRL 54463\(^t\) | Australia   | *Agapanthus* sp.               | KU900611 KU900630 KU900620 KU900625 KU900635 |
| *Fusarium ananatum*   | CBS 118516\(^t\) | South Africa | *Ananas comosus* fruit         | LT996175 LT996091 LT996188 LT996137 LT996112 |
| *Fusarium andiyazi*   | CBS 119857\(^t\) = NRRL 31727 | South Africa | *Sorghum bicolor* soil debris | LT996176 LT996092 LT996189 LT996138 LT996113 |
| *Fusarium anthophilum*| CBS 737.97 = NRRL 13602 | Germany     | *Hippeastrum* sp.              | LT996177 LT996093 LT996190 LT996139 LT996114 |
| *Fusarium armениcum*  | NRRL 6227    | USA         | Fescue hay                     | JX171446 JX171560               |
| *Fusarium asiaticum*  | CBS 110257 = NRRL 13818 | Japan       | Barley                         | JX171459 JX171573               |
| *Fusarium batridioides* | NRRL 20476  | USA         | Cronartium conigenum           | AF158343 AF160290 Not public Not public U34434 |
| *Fusarium begoniae*   | CBS 403.97\(^t\) = NRRL 25300 | Germany     | Begonia elatior hybrid         | AF158346 AF160293 LT996191 LT996140 U61543 |
| *Fusarium buharicum*  | CBS 796.70 = NRRL 13371 | USSR        | *Gossypium* rotting stem base  | LT996112 JF741086 JF741086 U61548 |
| *Fusarium bulbicola*  | CBS 220.76\(^t\) = NRRL 13618 | Germany     | Nerine bowdendi                | KF466327 KF466415 KF466394 KF466404 KF466437 |
| *Fusarium brachygiabum* | NRRL 13829  | Japan       | *Pinus radiata*                | JM931393 JM931943 JM931951 JM932080 JM932080 |
| *Fusarium circinatum* | CBS 405.97\(^t\) = NRRL 25331 | Australia   | *Coxis gasteenii*              | LT996178 KP083251 KP083269 KP083274 LT996115 |
| *Fusarium concentricum* | CBS 450.97\(^t\) = NRRL 25181 | Costa Rica  | *Musa sapientum* fruit         | AF158335 AF160282 LT996192 JF741086 JF741086 U61548 |
| *Fusarium continuum*  | F201128      | China       | *Zanthoxylum bungeanum* stem   | KM236720 KM520389 KM236780     |
| *Fusarium convolutans* | CBS 144207\(^t\) = CPC 33733 | South Africa | *Kypocarpa angustifolia* thizophore | LT996094 LT996193 LT996141 |
| *Fusarium culmorum*   | CBS 417.86 = NRRL 25475 | Denmark     | Moldy barley kernel            | JX171515 JX171628               |
| *Fusarium denticulatum* | CBS 735.97 = NRRL 25302 | USA         | *Ipomoea batatas*              | AF158322 AF160269 LT996195 LT996143 U61550 |
| *Fusarium dlaminii*   | CBS 119860\(^t\) = NRRL 13164 | South Africa | Soil debris in cornfield       | AF158330 AF160277 KU171681 KU171701 U34430 |
| *Fusarium fracticadum* | CBS 137234\(^t\) | Colombia    | *Pinus maximonoii* stem        | LT996179 KJ541059 LT996196 LT996144 KJ541051 |
| *Fusarium fractiflexum* | NRRL 28852\(^t\) | Japan       | *Cymbidium* sp.                | AF158341 AF160288 Not public LT575064 AF160315 |
| *Fusarium fredkrugeri* | NRRL 26152   | Niger       | Unknown                        | AF160306 AF160321               |
| *Fusarium fredkrugeri* | CBS 144209\(^t\) = CPC 33747 | South Africa | *Melhania acuminata* thizophore | LT996181 LT996097 LT996199 LT996147 LT996117 |
| *Fusarium gidami*     | CBS 144210 = NRRL 26061 | Madagascar  | *Striga hermonthica*           | AF158356 AF160303 LT996197 LT996145 AF160319 |
| *Fusarium gidami*     | CBS 144495 = CPC 33746 | South Africa | *Melhania acuminata* rhizosphere | LT996180 LT996096 LT996198 LT996146 LT996116 |
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| Species name | Strain | Country | Host | Sequence accession number |
|--------------|--------|---------|------|----------------------------|
| *Fusarium* fujikuroi | CBS 423.97 = NRRL 13566 | China | Oryza sativa | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* globosum | CBS 423.97 = NRRL 13566 | South Africa | Zea mays | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* goulgardi | NRRL 66250 | Australia | Xanthorrhoea glauca | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* graminearum | CBS 423.97 = NRRL 13566 | USA | Corn | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* konzum | CBS 423.97 = NRRL 13566 | USA | Sorghum bicolor | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* kyushuense | NRRL 25349 | Japan | Triticum aestivum | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* lactis | CBS 423.97 = NRRL 13566 | USA | Ficus carica | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* langsethiae | NRRL 66250 | Norway | Oats | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* lateritium | NRRL 13622 | USA | Ulmus sp. | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* longipes | NRRL 13368 | Australia | Soil | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* mangiferae | NRRL 25226 | Australia | Mangifera indica | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* nigrum | CBS 423.97 = NRRL 13566 | USA | Pinus patula | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* palustre | NRRL 13368 | USA | Spartina alterniflora | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* parvisorum | CBS 423.97 = NRRL 13566 | Colombia | Pinus patula | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* phyllophilum | CBS 423.97 = NRRL 13566 | USA | Dracaena deremensis | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* poae | NRRL 13714 | Unknown | Unknown | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* proliferatum | CBS 423.97 = NRRL 13566 | USA | Cattleya pseudobulb, hybrid | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* pseudocircinatum | CBS 423.97 = NRRL 13566 | Ghana | Solanum sp. | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* pseudograminearum | CBS 423.97 = NRRL 13566 | Australia | Hordeum vulgare | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* pseudonygamai | CBS 423.97 = NRRL 13566 | Nigeria | Pennisetum typhoides | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* poae | CBS 423.97 = NRRL 13566 | USA | Pseudotsuga menziesii | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| *Fusarium* poae | CBS 423.97 = NRRL 13566 | USA | Pseudotsuga menziesii | AF158333, AF160279, KF466417, KF466396, KF466406, KF466320, KF466459 |
| Species name | Strain[^1] | Country | Host | Sequence accession number[^5] |
|--------------|------------|---------|------|------------------------------|
| Fusarium ramigenum | CBS 418.98[^1] = NRRL 25208 | USA | Ficus carica | KF466335 KF466423 KF466401 KF466412 KF466445 |
| Fusarium sacchari | CBS 223.76 = NRRL 13999 | India | Saccharum officinarum | AF158331 AF160278 JX171466 JX171580 U34414 |
| Fusarium sambucinum | NRRL 22187 = NRRL 20727 | England | Solanum sp. | AF158331 AF160278 JX171466 JX171580 U34414 |
| Fusarium sarcochroum | CBS 745.79 = NRRL 20472 | Switzerland | Viscum album | JX171472 JX171586 |
| Fusarium sibiricum | NRRL 53430[^1] | Russia | Avena sativa | LT996184 KJ541067 LT996206 LT996153 KJ541057 |
| Fusarium sororuli | CBS 137242[^1] | Colombia | Pinus patula stems | LT996184 KJ541067 LT996206 LT996153 KJ541057 |
| Fusarium sp. | NRRL 66179 | USA | Hibiscus moscheutos | KX302913 KX302921 KX302929 |
| Fusarium sporotrichioides | NRRL 66180 | USA | Hibiscus moscheutos | KX302914 KX302922 KX302930 |
| Fusarium sterilhypomon | NRRL 66181 | USA | Hibiscus moscheutos | KX302915 KX302923 KX302931 |
| Fusarium stilboideus | NRRL 66182 | USA | Hibiscus moscheutos | KX302916 KX302924 KX302932 |
| Fusarium subglutinans | NRRL 66183 | USA | Hibiscus moscheutos | KX302917 KX302925 KX302933 |
| Fusarium sublunatum | NRRL 66184 | USA | Hibiscus moscheutos | KX302918 KX302926 KX302934 |
| Fusarium sp. | CBS 201.63 = NRRL 36351 | Portugal | Anacis hypogaea stored nut | GQ915484 |
| Fusarium sudanense | CBS 219.76 = NRRL 13613 | Germany | Succisa pratensis flower | LT996185 LT996207 LT996154 U34419 |
| Fusarium torreyae | CBS 483.94[^1] | Australia | Soil | U34418 |

[^1]: Strain number[^1] corresponds to the culture collection number.
[^5]: Sequence accession numbers correspond to the nucleotide database.
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### Table: New Fusarium species from the Kruger National Park, South Africa

| Species name                  | Strain Information                                                                 | Host Information                                      | Sequence accession number | CAL     | EF-1α | RPB1 | RPB2 | TUB     |
|-------------------------------|-------------------------------------------------------------------------------------|--------------------------------------------------------|---------------------------|---------|-------|------|------|---------|
| Fusarium transvaalense        | CBS 144212 = CPC 30932                                                              | Sida cordifolia rhizosphere                            | LT996109                  | LT996100|       |      |      |         |
| Fusarium transvaalense        | CBS 144213 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996119                  | LT996118|       |      |      |         |
| Fusarium transvaalense        | CBS 144214 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996101                  | LT996102|       |      |      |         |
| Fusarium transvaalense        | CBS 144215 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996103                  | LT996104|       |      |      |         |
| Fusarium transvaalense        | CBS 144216 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996105                  | LT996106|       |      |      |         |
| Fusarium transvaalense        | CBS 144217 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996107                  | LT996108|       |      |      |         |
| Fusarium transvaalense        | CBS 144218 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996109                  | LT996110|       |      |      |         |
| Fusarium transvaalense        | CBS 144219 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996111                  | LT996112|       |      |      |         |
| Fusarium transvaalense        | CBS 144220 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996113                  | LT996114|       |      |      |         |
| Fusarium transvaalense        | CBS 144221 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996115                  | LT996116|       |      |      |         |
| Fusarium transvaalense        | CBS 144222 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996117                  | LT996118|       |      |      |         |
| Fusarium transvaalense        | CBS 144223 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996119                  | LT996120|       |      |      |         |
| Fusarium transvaalense        | CBS 144224 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996121                  | LT996122|       |      |      |         |
| Fusarium transvaalense        | CBS 144225 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996123                  | LT996124|       |      |      |         |
| Fusarium transvaalense        | CBS 144226 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996125                  | LT996126|       |      |      |         |
| Fusarium transvaalense        | CBS 144227 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996127                  | LT996128|       |      |      |         |
| Fusarium transvaalense        | CBS 144228 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996129                  | LT996130|       |      |      |         |
| Fusarium transvaalense        | CBS 144229 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996131                  | LT996132|       |      |      |         |
| Fusarium transvaalense        | CBS 144230 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996133                  | LT996134|       |      |      |         |
| Fusarium transvaalense        | CBS 144231 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996135                  | LT996136|       |      |      |         |
| Fusarium transvaalense        | CBS 144232 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996137                  | LT996138|       |      |      |         |
| Fusarium transvaalense        | CBS 144233 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996139                  | LT996140|       |      |      |         |
| Fusarium transvaalense        | CBS 144234 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996141                  | LT996142|       |      |      |         |
| Fusarium transvaalense        | CBS 144235 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996143                  | LT996144|       |      |      |         |
| Fusarium transvaalense        | CBS 144236 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996145                  | LT996146|       |      |      |         |
| Fusarium transvaalense        | CBS 144237 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996147                  | LT996148|       |      |      |         |
| Fusarium transvaalense        | CBS 144238 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996149                  | LT996150|       |      |      |         |
| Fusarium transvaalense        | CBS 144239 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996151                  | LT996152|       |      |      |         |
| Fusarium transvaalense        | CBS 144240 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996153                  | LT996154|       |      |      |         |
| Fusarium transvaalense        | CBS 144241 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996155                  | LT996156|       |      |      |         |
| Fusarium transvaalense        | CBS 144242 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996157                  | LT996158|       |      |      |         |
| Fusarium transvaalense        | CBS 144243 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996159                  | LT996160|       |      |      |         |
| Fusarium transvaalense        | CBS 144244 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996161                  | LT996162|       |      |      |         |
| Fusarium transvaalense        | CBS 144245 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996163                  | LT996164|       |      |      |         |
| Fusarium transvaalense        | CBS 144246 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996165                  | LT996166|       |      |      |         |
| Fusarium transvaalense        | CBS 144247 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996167                  | LT996168|       |      |      |         |
| Fusarium transvaalense        | CBS 144248 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996169                  | LT996170|       |      |      |         |
| Fusarium transvaalense        | CBS 144249 = CPC 30932                                                              | Mahonia acuminata rhizosphere                           | LT996171                  | LT996172|       |      |      |         |

**Strain Information**: CBS: Westerdijk Fungal Biodiversity Institute. CPC: Collection of Pedro W. Crous, held at CBS. **Host Information**: Soil, rhizosphere, trunk. **Sequence accession number**: LT996099 – LT996134. **CAL**: Calmodulin. **EF-1α**: Translation elongation factor 1-alpha. **RPB1**: RNA polymerase largest subunit. **RPB2**: RNA polymerase second largest subunit. **TUB**: Tubulin. **Country**: Australia, Brazil, Germany, China, Ivory Coast. **Host**: Sida cordifolia, Melhania acuminata, Kyphocarpa angustifolia, Mangifera indica, Lactarius pubescens, Zoë mopy, Coffea sp., mang., Winter wheat. **Sequence information**: New sequences are shown in **bold**. Sequences marked as “Not public” were obtained from Kerry O’Donnell’s alignment datasets.
beta-tubulin (*TUB*). Sequencing was made in both strand directions using the same primer pairs as for PCR amplification on an Applied Biosystems, Hitachi 3730xl DNA analyser (Applied Biosystems Inc., Foster City, California, USA). Consensus sequences were assembled using Seqman Pro v. 10.0.1 (DNASTAR, Madison, WI, USA). All DNA sequences generated in this study were lodged in GenBank and the European Nucleotide Archive (ENA) (Table 1).

**Molecular identification and phylogenetic analyses**

A first analysis was based on pairwise alignments and blastn searches on the *Fusarium* MLST (http://www.westerdijkinstitute.nl/fusarium/) and NCBI (https://blast.ncbi.nlm.nih.gov/Blast.cgi) databases, respectively, using *EF-1α* and *RPB2* sequences in order to resolve the position of the KNP isolates amongst the different species complexes recognised in *Fusarium* (O’Donnell et al. 2013). Sequences from individual loci were aligned using MAFFT (Katoh and Standley 2013), on the web server of the European Bioinformatics Institute (EMBL–EBI; http://www.ebi.ac.uk/Tools/msa/mafft/) (Li et al. 2015).

Phylogenetic analyses were based on Maximum-likelihood (ML) and Bayesian (B) analyses, both algorithms run on the CIPRES Science Gateway portal (Miller et al. 2012). Evolutionary models were calculated using MrModelTest v. 2.3 using the Akaike information criterion (Nylander 2004; Posada and Crandall 1998). For ML, RAxML-HPC2 v. 8.2.10 on XSEDE was used (Stamatakis 2014), clade stability was tested with a bootstrap analysis (BS) using the rapid bootstrapping algorithm with default parameters. The B analyses were run using MrBayes v. 3.2.6 on XSEDE (Ronquist and Huelsenbeck 2003) using four incrementally heated MCMC chains for 5M generations, with the stop-rule option on and sampling every 1000 trees. After convergence of the runs (average standard deviation of split frequencies below 0.01) the first 25% of samples were discarded as the burn-in fraction and 50% consensus trees and posterior probabilities (PP) were calculated from the remaining trees.

Phylogenies were first made individually for each locus dataset and visually compared for topological incongruence amongst statistically supported nodes (ML-BS ≥ 70% and B-PP ≥ 0.95) (Mason-Gamer and Kellogg 1996, Wiens 1998), before being concatenated for multi-locus analyses using different locus combinations according to strains and DNA sequences currently available in public databases, in addition to previously published phylogenies (O’Donnell et al. 2000, 2013; Herron et al. 2015; Lupien et al. 2017; Moussa et al. 2017, Sandoval-Denis et al. 2018). A further 232 sequences representing 72 taxa were retrieved from GenBank and included in the phylogenetic analyses, while an additional 58 DNA sequences were obtained from 24 fungal strains requested from the CBS and NRRL (Agricultural Research Service, Peoria, IL, USA) culture collections (Table 1). All alignments and trees generated in this study were uploaded to TreeBASE (https://treebase.org).
Results

Phylogenetic analyses

Pairwise DNA alignments and BLAST searches using EF-1α and RPB2 sequences showed that the 19 isolates from KNP belonged to three different species complexes of the genus *Fusarium* i.e. the *F. buharicum* Jacz. ex Babajan & Teterevn.-Babajan species complex (FBSC; two isolates), the *F. fujikuroi* Nirenberg species complex (FFSC; two isolates) and the *F. sambucinum* Fuckel species complex (FSAMSC; 15 isolates). According to these results, sequences of related taxa and lineages were retrieved from GenBank and incorporated into individual phylogenetic analyses for each species complex.

Multi-locus analyses were carried out in order to further delimit the KNP *Fusarium* isolates amongst the known diversity in their respective species complexes. With the exception of the FFSC, the topologies observed from ML and B analyses of single and multi-locus datasets were highly congruent, with only minor differences affecting unsupported nodes on the trees (all trees available in TreeBASE). The characteristics of the different alignments and tree statistics for all the species complexes are shown in Table 2.

The analysis of the FBSC included sequences of EF-1α, RPB1 and RPB2 loci from 18 isolates representing 10 taxa, including members of the *Fusarium torreyae* T. Aoki, J.A. Sm., L.L. Mount, Geiser & O’Donnell species complex (FTYSC) and *Fusarium lateritium* Nees species complex (FLSC) as outgroup (Fig. 2). The four ingroup taxa resolved with high statistical support. Two KNP isolates from *K. angustifolia* obtained from the bottom site of the catena (CBS 144207 and 144208) clustered in a sister relationship with the clade representing *Fusarium sublunatum* Reinking, but were genetically clearly delimited.

The phylogeny of the FFSC included sequences of CAL, EF-1α, RPB1, RPB2 and TUB loci from 48 strains and 44 taxa, including two outgroups (*F. oxysporum* CBS 716.74 and 744.97) (Fig. 3). The phylogeny showed a clear delimitation between the biogeographic clades recognised in this species complex (African, American and Asian clades *sensu* O’Donnell et al. 1998). Both American and Asian clades were shown as monophyletic with high ML-BS and B-PP support; in contrast, the African clade was resolved as polyphyletic, comprising two distinct and highly supported lineages. A terminal, speciose clade (African A) encompassing 17 taxa and a basal clade (African B), close to the American clade which included the ex-type of *Fusarium dlaminii* Marasas, P.E. Nelson & Toussoun (CBS 119860) and a sister terminal clade (ML-BS=100, B-PP=1) comprising two KNP isolates from *M. acuminata* (CBS 144209 and 144495) and two unidentified African *Fusarium* isolates (CBS 144210 and NRRL 26152). From the loci used here, only TUB resolved both African clades as sister groups; however, its monophyly was not supported by clade stability measurements (data not shown). Conversely, individual CAL, EF-1α and RPB2 phylog-
Table 2. Characteristics of the different datasets and statistics of phylogenetic analyses used in this study.

| Analysis† | Locus‡ | Number of Sites§ | Evolutionary model¶ | Number of trees sampled in B | Maximum-likelihood statistics | Tree length |
|-----------|--------|------------------|---------------------|-----------------------------|-------------------------------|-------------|
|           | Total  | Conserved | Phylogenetically informative | B unique patterns | | |
| Fusarium buharicum SC | EF-1a | 495 | 300 | 119 | 198 | GTR+G | 414 | -11313.23702 | 0.598675 |
|           | RPB1  | 930 | 682 | 203 | 211 | SYM+G | 357 | -11313.23702 | 0.598675 |
|           | RPB2  | 1663 | 1251 | 330 | 310 | GTR+I+G | 364 | -11313.23702 | 0.598675 |
| Fusarium fujikuroi SC | CAL | 545 | 423 | 67 | 167 | SYM+G | 282 | -20603.30043 | 0.567054 |
|           | EF-1a | 677 | 428 | 127 | 295 | GTR+I+G | 282 | -20603.30043 | 0.567054 |
|           | RPB1  | 1534 | 1219 | 185 | 137 | SYM+I+G | 282 | -20603.30043 | 0.567054 |
|           | RPB2  | 1551 | 1211 | 227 | 315 | GTR+I+G | 282 | -20603.30043 | 0.567054 |
|           | TUB   | 488 | 351 | 66 | 336 | SYM+G | 282 | -20603.30043 | 0.567054 |
| Fusarium sambucinum SC | RPB1  | 854 | 594 | 201 | 213 | SYM+I+G | 241 | -9871.793718 | 0.740271 |
|           | RPB2  | 1580 | 1128 | 346 | 396 | GTR+G | 241 | -9871.793718 | 0.740271 |

† SC: Species complex.
‡ CAL: Calmodulin. EF-1α: Translation elongation factor 1-alpha. RPB1: RNA polymerase largest subunit. RPB2: RNA polymerase second largest subunit. TUB: Tubulin.
§ B: Bayesian inference.
¶ G: Gamma distributed rate variation among sites. GTR: Generalised time-reversible. I: Proportion of invariable sites. SYM: Symmetrical model.

Figure 2. Maximum-likelihood (ML) phylogram obtained from combined EF-1α, RPB1 and RPB2 sequences of 18 strains belonging to the Fusarium buharicum (FBSC), Fusarium tricinctum (FTSC) and Fusarium lateritium (FLSC) species complexes. Numbers on the nodes are ML bootstrap values above 70% and Bayesian posterior probability values above 0.95. Branch lengths are proportional to distance. Ex-type strains are indicated with T. Strains corresponding to new species described here are shown in **bold**.
New Fusarium species from the Kruger National Park, South Africa

enies resolved African B as basal to the ingroup, while RPB1 allocated this clade as basal to the American clade. Nonetheless, all the individual phylogenies, in addition to the combined dataset, clearly demonstrated genealogical uniqueness of the terminal clade encompassing KNP isolates.

Figure 3. Maximum-likelihood (ML) phylogram obtained from combined CAL, EF-1α, RPB1, RPB2 and TUB sequences of 48 strains belonging to the Fusarium fujikuroi (FFSC) and Fusarium oxysporum (FOSC) species complexes. Numbers on the nodes are ML bootstrap values above 70% and Bayesian posterior probability values above 0.95. Branch lengths are proportional to distance. Ex-type, ex-neotype and ex-paratype strains are indicated with T, NT and PT, respectively. Strains corresponding to new species described here are shown in bold.
The FSAMSC was studied using combined RPB1 and RPB2 sequences. The phylogeny included 35 isolates from 20 taxa, including the two outgroups Fusarium circinatum Nirenberg & O’Donnell (CBS 405.97) and Fusarium fujikuroi Nirenberg (NRRL 13566) (Fig. 4). Fifteen KPN Fusarium isolates from the three sampled plant species (three isolates from K. angustifolia, four isolates from M. acuminata and eight isolates from S. cordifolia), all obtained from the top site of the catena, clustered with an unidentified Fusarium isolate (NRRL 31008) in a distinct clade (ML-BS=100, B-PP=1), close to Fusarium brachygibbosum Padwick (strain NRRL 13829).

**Figure 4.** Maximum-likelihood (ML) phylogram obtained from combined RPB1 and RPB2 sequences of 35 strains belonging to the *Fusarium sambucinum* (FSAMSC) and *Fusarium fujikuroi* (FFSC) species complexes. Numbers on the nodes are ML bootstrap values above 70% and Bayesian posterior probability values above 0.95. Branch lengths are proportional to distance. Ex-type strains are indicated with T. Strains corresponding to new species described here are shown in bold.
New Fusarium species from the Kruger National Park, South Africa

The clades including KNP isolates and corresponding to previously undisclosed lineages of *Fusarium* are described in the taxonomy section as the three novel species, *F. convolutans*, *F. fredkrugeri* and *F. transvaalense*.

**Taxonomy**

*Fusarium convolutans* Sandoval-Denis, Crous & W.J. Swart, *sp. nov.*

MycoBank: MB825102

*Fig. 5*

**Diagnosis.** Different from *F. circinatum*, *F. pseudocircinatum* O’Donnell & Nirenberg and *F. sterilhyposum* Britz, Marasas & M.J. Wingf. by the absence of aerial conidia (microconidia) and the presence of chlamydospores. Different from *F. buharicum* Jacz. ex Babajan & Teterevn.-Babajan and *F. sublunatum* by its shorter, less septate and less curved conidia and by the presence of sterile hyphal coils.

**Type.** South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06′33.9″S, 31°34′40.9″E, from rhizosphere soil of *Kyphocarpa angustifolia*, 23 Mar 2015, W.J. Swart, holotype CBS H-23495, dried culture on OA, ex-holotype strain CBS 144207 = CPC 33733.

**Description.** Colonies on PDA growing in the dark with an average radial growth rate of 2.1–4.8 mm/d, 4.4–5.8 mm/d and 4.6–6.3 mm/d at 24, 27 and 30 °C, respectively; reaching 11–28 mm diam. in 7 d at 24 °C and a maximum of 23–37 mm diam. in 7 d at 30 °C. Minimum temperature for growth 12 °C, maximum 36 °C, optimal 27–33 °C. Colony surface white to cream coloured, flat and highly irregular in shape, velvety to felty, with scant and short aerial mycelium; colony margins highly irregular to rhizoid, with abundant white to grey submerged mycelium. Reverse white, straw to yellow diffusible pigment produced between 21–33 °C, scarcely produced and turning luteous to orange at 36 °C. Colonies on CMA and OA incubated in the dark reaching 40–48 mm diam. in 7 d at 24 °C. Colony surface white to cream coloured, flat or slightly elevated at the centre, velvety to dusty; aerial mycelium abundant, short and dense, concentrated on the colony centre; margins membranous and regular, buff to honey coloured, without aerial mycelium. Reverse ochreous without diffusible pigments. Sporulation scant from conidiophores formed on the aerial mycelium, sporodochia not formed. *Conidiophores* on the aerial mycelium straight or flexuous, smooth- and thin-walled, simple, mostly reduced to conidiogenous cells borne laterally on hyphae or up to 50 μm tall, bearing terminal single or paired monophialides; *phialides* subulate to subcylindrical, smooth- and thin-walled, 15.5–22 μm long, (3.5–)4–5 μm at the widest point, with inconspicuous periclinal thickening and a short-flared collarette; *conidia* clustering in discrete false heads at the tip of monophialides, lunate to falcate, curved or somewhat straight, tapering gently toward the basal part, robust; apical cell often equal in length or slightly shorter than the adjacent cell, blunt
Figure 5. *Fusarium convolutans* sp. nov. A–D Colonies on PDA, SNA, OA and CMA, respectively, after 7 d at 24 °C in the dark E–I Conidiophores, phialides and conidia J–M Chlamydosporas N–P Sterile hyphal projections Q Conidia. Scale bars: 20 μm (E, F); 5 μm (G–I); 10 μm (J–Q).
New Fusarium species from the Kruger National Park, South Africa

Chlamydospores abundantly formed, globose to subglobose, smooth- and thick-walled, (9.5–)11–13.5(–14) μm diam.; terminal or intercalary in the hyphae or conidia, often borne laterally at the tip of elongated, cylindrical, stalk-like projections, solitary or in small clusters. Sterile, coiled, sometimes branched hyphal projections abundantly formed laterally from the substrate and aerial mycelium.

Distribution. South Africa.

Etymology. From Latin, “convolutans”, participle of convolutare, coiling, in reference to the abundant sterile, coiled lateral hyphal projections.

Additional isolate examined. South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06’33.9”S, 31°34’40.9E, from rhizosphere soil of Kyphocarpa angustifolia, 23 Mar 2015, W.J. Swart, CBS 144208 = CPC 33732.

Notes. The main morphological feature of *F. convolutans*, namely the production of sterile, coiled hyphal projections, grossly resembles other *Fusarium* species producing similar structures i.e. *F. circinatum*, *F. pseudocircinatum* and *F. sterilihyphosum*. The three latter species, however, are genetically unrelated to *F. convolutans*, being allocated in the FFSC; and are also easily differentiable by the characteristics of the aerial conidia (typical *Fusarium* microconidia are absent in the new species) and the lack of chlamydospores (present in the new species) (Leslie and Summerell 2006). *Fusarium convolutans* can be easily differentiated morphologically from their phylogenetically closely related species, *F. buharicum* and *F. sublunatum*. It has relative simple conidiophores and shorter, less septate and markedly less curved conidia (up to 38.5 μm long and 1–3-septate vs. up to 87 and 81 μm long, 0–8-septate in *F. buharicum* and *F. sublunatum*, respectively) (Gerlach and Nirenberg 1982). *Fusarium buharicum* and *F. sublunatum* also lack sterile hyphal coils.

_Fusarium fredkrugeri_ Sandoval-Denis, Crous & W.J. Swart, sp. nov.

MycoBank: MB825103

Fig. 6

Diagnosis. Differs from _Fusarium dlaminii_ Marasas, P.E. Nelson & Toussoun by producing only one type of aerial conidia, shorter sporodochial conidia and the absence of chlamydospores.

Type. South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06’48.6”S, 31°34’36.5”E, from rhizosphere soil of Melhania acuminata, 23 Mar 2015, W.J. Swart, holotype CBS H-23496, dried culture on OA, culture ex-holotype CBS 144209 = CPC 33747.

Description. Colonies on PDA growing in the dark with an average radial growth rate of 4.7–5.8 mm/d and reaching 22–35 mm diam. in 7 d at 24 °C, filling an entire 9 cm Petri dish in 7 d at 27 and 30 °C. Minimum temperature for growth 12 °C, maxi-
Figure 6. *Fusarium fredkrugeri* sp. nov. A–D Colonies on PDA, SNA, OA and CMA, respectively, after 7 d at 24 °C in the dark E–G Sporodochia formed on the surface of carnation leaves H–N Aerial conidiophores, phialides and conidia O, P Aerial conidia Q Sporodochial conidiophores and phialides R Sporodochial conidia. Scale bars: 100 μm (E–G); 10 μm (H–R).
New Fusarium species from the Kruger National Park, South Africa

New Fusarium species from the Kruger National Park, South Africa

Maximum 36 °C, optimal 27–30 °C. Colony surface at first white to cream coloured, later turning bay to chestnut with pale luteous to luteous periphery; flat, felty to cottony with abundant erect- aerial mycelium forming white patches; colony margins regular and filiform with abundant submerged mycelium. Reverse pale luteous, a blood sepioid pigment is scarcely produced at 24 °C, pigment production is markedly enhanced at 27–30 °C, becoming greyish-sepia at 33 °C. Colonies on CMA and OA incubated at 24 °C in the dark reaching 65–67 mm diam. or occupying an entire 9 cm Petri dish in 7 d, respectively. Colony surface pale bay coloured, flat, felty to velvety, aerial mycelium scant, forming white to cream patches; margins regular. Reverse pale bay to pale vinaceous. Sporulation abundant from conidiophores formed on the substrate and aerial mycelium and from sporodochia. Conidiophores on the aerial mycelium straight or flexuous, erect or prostrate, septate, smooth- and thin-walled, often appearing rough by accumulation of extracellular material, commonly simple or reduced to conidiogenous cells borne laterally on hyphae or up to 200 μm tall and irregularly branched at various levels, branches bearing lateral and terminal monophialides borne mostly single or in pairs; phialides subulate, ampulliform, lageniform to subcylindrical, smooth- and thin-walled, (8.5–)9.5–17.5(–24.5) μm long, 2–3(–3.5) μm at the widest point, without periclinal thickening, collarettes inconspicuous; conidia formed on aerial conidiophores, hyaline, obovoid, ellipsoidal to slightly reniform or allantoid, smooth- and thin-walled, 0-septate, (4.5–)5–8.5(–12.5) × (1.5–)2–3.5(–6) μm, clustering in discrete false heads at the tip of monophialides. Sporodochia pale orange to pink coloured, often somewhat translucent, formed abundantly on the surface of carnation leaves and on the agar surface. Conidiophores in sporodochia 26–46 μm tall, densely aggregated, irregularly and verticillately branched up to three times, with terminal branches bearing 2–3 monophialides; sporodochial phialides doliiform to subcylindrical, (9–)11.5–15.5(–18.5) × (2.5–)3–4(–4.5) μm, smooth- and thin-walled, with periclinal thickening and an inconspicuous apical collarette. Sporodochial conidia falcate, tapering toward the basal part, robust, moderately curved and slender; basal cell more or less equally sized than the adjacent cell, blunt to slightly papillate; basal cell papillate to distinctly notched, (1–)3–4-septate, hyaline, thin- and smooth-walled. One-septate conidia: 13–17(–18) × (2.5–)3–4 μm; two-septate conidia: 15 × 4.5 μm; three-septate conidia: (16–)28.5–39(–45) × (3–)4–5(–5.5) μm; four-septate conidia: 39.5–40(–41) × 4.5–5 μm; overall (13–)27.5–39.5(–45) × (3–)3.5–5.5 μm. Chlamydospores absent.

**Distribution.** Madagascar, Niger and South Africa.

**Etymology.** In honour and memory of Dr. Frederick J. Kruger, pioneer of forest hydrology, fynbos ecology and invasive species and fundamental for the collections included in this study.

**Additional isolates examined.** Madagascar, from Striga hermonthica, unknown date, A.A. Abbasher, CBS 144210 = NRRL 26061 = BBA 70127. South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06′48.6″S, 31°34′36.5″E, from rhizosphere soil of Melhania acuminata, 23 Mar 2015, W.J. Swart, CBS 144495 = CPC 33746.

**Notes.** This species is genetically closely related to *F. dlaminii*, both species having similar colonial morphology, optimal growth conditions and biogeography. Moreo-
ver, both species exhibit relatively short aerial phialides producing conidia in heads, somewhat resembling those produced by *F. oxysporum* rather than most members of the FFSC (Leslie and Summerell 2006; Marasas et al. 1985). However, besides exhibiting much faster growth rates, *F. fredkrugeri* presents clearly distinctive morphological features such as the production of only one type of aerial conidia (vs. two types in *F. dlaminii*; allantoid to fusiform and 0-septate; and napiform 0–1-septate); orange to pink sporodochia, produced on carnation leaves but also abundantly on the agar surface (vs. orange sporodochia, produced only on the surface of carnation leaves in *F. dlaminii*) (Leslie and Summerell 2006). Additionally, *F. fredkrugeri* produces shorter and less septate sporodochial conidia ((1–)3–4-septate and up to 45 μm long in the latter species vs. mostly 5-septate and up to 54 μm long in *F. dlaminii*) while chlamydospores are not produced. The latter feature, coupled with the somewhat more complex conidiophores also clearly differentiates *F. fredkrugeri* from *F. oxysporum.*

**Fusarium transvaalense** Sandoval-Denis, Crous & W.J. Swart, sp. nov.  
MycoBank: MB825104  
Fig. 7

**Diagnosis.** Different from most species in FSAMSC by its slender sporodochial conidia with tapered and somewhat rounded apex; its smooth- to tuberculate, often pigmented chlamydospores and the formation of large mycelial tufts on OA.

**Type.** South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06’45.5"S, 31°34’35.0"E, from rhizosphere soil of *Sida cordifolia*, 23 Mar 2015, W.J. Swart, holotype CBS H-23497, dried culture on SNA, culture ex-holotype CBS 144211 = CPC 30923.

**Description.** Colonies on PDA growing in the dark with an average radial growth rate of 8.5–9.3 mm/d, reaching 34–37 mm diam. in 7 d at 24 °C, filling an entire 9 cm Petri dish in 7 d at 27–33 °C. Minimum temperature for growth 12 °C, maximum 36 °C, optimal 27–30 °C. Colony surface at first white, turning coral to dark vinaceous with white periphery and abundant yellow hyphae at the centre; flat, velvety to woolly, with abundant aerial mycelium and erect hyphal strings reaching several mm tall; colony margins regular and filiform. Reverse with yellow, coral or dark vinaceous patches, coral diffusible pigments strongly produced between 15–30 °C, turning scarlet to orange at 33–36 °C. Colonies on CMA and OA incubated at 24 °C in the dark occupying an entire 9 cm Petri dish in 7 d. Colony surface coral, rust to chestnut coloured in irregular patches, flat, felty to woolly, aerial mycelium scarce on CMA, mostly as radially dispersed white patches, on OA aerial mycelium abundant, especially on the periphery of the colony, forming dense, pustule-like, white mycelial tufts, formed by abundant intermingled hyphae and chlamydospores, 1–1.5 cm tall, with flesh to coral coloured stipes; margins on CMA and OA regular. Reverse pale luteous with red to coral periphery. Sporulation abundant from conidiophores formed on the aerial mycelium, at the agar level and from sporodochia. **Conidiophores** on the aerial mycelium straight or flexuous, septate, smooth- and thin-walled, up to 150 μm tall, sometimes
Figure 7. *Fusarium transvaalense* sp. nov. A–D Colonies on PDA, SNA, OA and CMA, respectively, after 7 d at 24 °C in the dark E Pustule-like growth on OA F, G Sporodochia formed on the surface of carnation leaves H–L Aerial conidiophores phialides and conidia M Aerial conidia N, O Chlamydospores P Sporodochial conidiophores and phialides Q Sporodochial conidia. Scale bars: 2 mm (E); 20 μm (F–J); 5 μm (K); 10 μm (L–Q).
emerging from irregular, swollen, pigmented and rough-walled cells on the hyphae; simple or sparingly and irregularly branched, branches bearing terminal, rarely lateral monophialides or reduced to conidiogenous cells borne laterally on hyphae; phialides on the aerial conidiophores short ampulliform, subulate to subcylindrical, smooth- and thin-walled, (7–)9–14(–15) μm long, (3–)4–5 μm at the widest point, without periclinal thickening and with a minute, inconspicuous collarette; conidia formed on aerial conidiophores of two types: a) hyaline, obovoid, ellipsoidal to clavate, smooth- and thin-walled, 0–1-septate, 2–14 × 2–4 μm; b) lunate to short falcate with a pointed apex and a somewhat flattened base, smooth- and thin-walled, 3–5-septate. Three-sep
tate conidia: (16–)18–27(–29) × 5–6 μm; four-septate conidia: 21–24(–25) × 5–6 μm; five-septate conidia: (25–)27–33 × 5–6 μm. Sporodochia cream to orange coloured, formed abundantly on the surface of carnation leaves and rarely on the agar surface, at first very small and sparse later becoming aggregated. Conidiophores in sporodochia 22–31 μm tall, irregularly branched, bearing clusters of 3–6 monophialides; sporodochial phialides doliform to ampulliform, (5–)9–14(–18) × (3–)4–5 μm, smooth- and thin-walled, with periclinal thickening and a short apical collarette. Sporodochial conidia falcate, wedge-shaped, tapering towards both ends, markedly curved and robust; apical cell longer than the adjacent cell, pointed; basal cell distinctly notched, sometimes somewhat extended (1–)3–5(–6)-septate, hyaline, smooth- and thick-walled. One-septate conidia: 19 × 4 μm; three-septate conidia: 20–27(–28) × 5–7 μm; four-septate conidia: (29–)30–32 × 5–7 μm; five-septate conidia: (26–)29–41(–53) × 4–5(–6) μm; six-septate conidia: 36 × 7 μm; overall (19–)25.9–40(–53) × (3.5–)4–6(–7) μm. Chlamydospores abundant, hyaline or pigmented, smooth- to rough-walled or tuberculate, 7–8 μm diam., terminal or intercalary, solitary, in chains or in clusters.

**Distribution.** Australia and South Africa

**Etymology.** After Transvaal, the name of a former colony and Republic located between the Limpopo and Vaal rivers, currently a province of South Africa and where this species was found. From Latin *trans* meaning “on the other side of” and Vaal a South African river.

**Additional isolates examined.** South Africa, Kruger National Park, Skukuza, Granite Supersite, 25°06’48.6”S, 31°34’36.5”E, from rhizosphere soil of *Melhania acuminata*, 23 Mar 2015, W.J. Swart, CBS 144224 = CPC 30928, CBS 144212 = CPC 30929); 25°06’45.6”S, 31°34’37.7”E, CBS 144496 = CPC 33750, CBS 144213 = CPC 33751; 25°06’48.8”S, 031°34’36.6”E, from rhizosphere soil of *Sida cordifolia*, 23 Mar 2015, W.J. Swart, CBS 144214 = CPC 30946; 25°06’45.7”S, 31°34’35.1”E, CBS 144215 = CPC 33723; 25°06’45.5”S, 31°34’35.0”E, CBS 144216 = CPC 30918, CBS 144217 = CPC 30919, CBS 144218 = CPC 30922, , CBS 144219 = CPC 30926, CBS 144220 = CPC 30927); 25°06’51.4”S, 31°34’37.5”E, from rhizosphere soil of *Kyphocarpa angustifolia*, 23 Mar 2015, W.J. Swart, CBS 144221 = CPC 33740; 25°06’51.8”S, 31°34’38.1”E, CBS 144222 = CPC 30939, CBS 144223 = CPC 30941.

**Notes.** *Fusarium transvaalense* exhibits a sporodochial conidial morphology typical of members of FSAMSC with marked dorsiventral curvature and tapered ends. Several species in FSAMSC form comparable conidia in culture i.e. *F. crookwellense*
L.W. Burgess, P.E. Nelson & Toussoun, *F. sambucinum*, *F. sporotrichioides* Sherb., *F. venenatum* Nirenberg and *F. culmorum* (Wm.G. Sm.) Sacc. However, with the exception of *F. sporotrichioides*, the conidia of most species above-mentioned, differ by being more robust and often more pointed apically. *Fusarium transvaalense* differs from *F. sporotrichioides* by the absence of pyriform aerial conidia.

Two strains NRRL 13829 and NRRL 31008, previously identified as *F. brachygibbosum* Padwick showed different degrees of genetic similitude with the new species. While NRRL 31008 clustered within *F. transvaalense*, NRRL 13829 formed a clearly delimited sister lineage. Morphologically, *F. transvaalense* exhibits significant differences allowing its separation from *F. brachygibbosum*. Both species produce sporodochial conidia with similar septation and sizes; however, *F. brachygibbosum* commonly exhibits a bulge in the middle portion of the conidia (Padwick 1945), a feature not present in *F. transvaalense*. In addition, the latter species produces comparatively larger sporodochial conidia, when elements with the same degree of septation are compared; its chlamydospores are smaller, smooth-walled to markedly tuberculate and pigmented (7–8 μm vs. 10.7–15.3 μm, smooth-walled and hyaline in *F. brachygibbosum*) and has a distinctive colonial growth on OA, forming large, pustule-like hyphal tufts, a feature not reported for *F. brachygibbosum* (Padwick 1945).

**Discussion**

In this study, three new *Fusarium* spp. were introduced, isolated from rhizosphere soils of three native African shrubs in a protected savannah ecosystem deep inside the Kruger National Park, South Africa.

Some remarkable differences were noted regarding the distribution of the novel fungal species and their respective hosts on this particular site. For instance, *F. transvaalense*, which exhibited the greatest relative abundance, was found in high quantities from the rhizospheres of the three hosts sampled, showing a considerable genetic diversity. Interestingly, this species was only on the top of the catena, even when two of its hosts, *K. angustifolia* and *S. cordifolia*, were found and sampled either at the top and bottom sites. Similarly, *F. fredkrugeri* was recovered only from soils under *M. acuminata*, a host species which occurred only at the top location. In contrast, *F. convolutans* was found in the rhizosphere of *K. angustifolia*, occurring only at the bottom of the catena, while none of the three fungal species was found associated with *S. cordifolia* at the bottom of the site. Nevertheless, not being an objective of this work, it was not possible to categorically assign these new species to specific hosts or locations. Likely, these fungi could be in low abundance and thus not detectable using the current methods. However, plant species composition varies considerably through a catena ecosystem, in relation to the different soil characteristics, pH gradient and water availability, which also greatly influence microbial and animal biodiversity (Lareen et al. 2016; Mohammadi et al. 2017). However, the full patterns of variation between locations on this particular catena still need to be systematically assessed and compared. As evidenced
here, certain differences do exist between the soils at the upper and bottom locations of the Stevenson-Hamilton supersite, which might explain the fungal diversity variation observed here. The cation exchange capacity (CEC; capacity of a soil to hold exchangeable cations) varies considerably between sampling sites, basically depending on the proportion of sand versus clay content of each soil type (Ketterings et al. 2007; Van Zijl and Le Roux 2014). It is known that CEC greatly impacts the soil's ability to retain essential nutrients and prevents soil acidification (Ketterings et al. 2007). Nutrient content also increased from the top to the bottom of the slope which is consistent with the increase in CEC. Nutrient poor soils are also a driver of biological diversity and most likely influenced fungal diversity in these particular locations (Havlicek and Mitchell 2014, Mapelli et al. 2017).

The three *Fusarium* species, described here, were not associated with any visible symptomatology on their hosts. However, they cannot be ruled out as pathogens since they were not assessed for pathogenicity against the sampled plants nor any other putative host species at the same locations. Likewise, it is unknown if these fungi exert any beneficial or deleterious effect on their ecosystems. These are important unsolved questions that need further evaluation. However, as shown by phylogenetic analyses, each of the three new species was in close genetic proximity with well-known plant pathogenic *Fusarium* spp. on their respective species complexes, which could suggest a potential pathogenic role. *Fusarium convolutans* clustered within the FBSC, together with three known plant pathogenic *Fusarium* spp. i.e. *F. buharicum*, a pathogen of *Hibiscus cannabinus* L. and *Gossypium* L.; *F. sublunatum*, known to affect banana and *Theobroma cacao* L. in Central America (Gerlach and Nirenberg 1982, Leslie and Summerell 2006) and a newly discovered although unnamed phylogenetic species causing wilt, crown and root rot of *Hibiscus moscheutos* L. (Lupien et al. 2017). *Fusarium transvaalense* belonged to the FSAMSC, a genetically diverse group common in temperate and subtropical zones (Leslie and Summerell 2006). *Fusarium sambucinum*, the conserved type species of the genus (Gams et al. 1997) being an aggressive plant pathogen and one of the most important agents of potato dry rot (Peters et al. 2008); while the latter species and several others in the complex have been reported causing disease on diverse crops, including many cereals and fruits (Leslie and Summerell 2006).

*Fusarium fredkrugeri* is here recognised and formally proposed as a new species. Although the clade representing this taxon had already been identified as a distinct unnamed phylogenetic species by O’Donnell et al. (2000), it had not been given a formal description pending the collection of additional isolates. Two other African isolates previously determined to belong to this clade i.e. CBS 144210 from *Striga hermonthica* (Del.) Benth. in Madagascar and NRRL 26152 from an unknown substrate in Niger, were incorporated into the analyses, although the latter strain is not viable anymore (NRRL, pers. comm.), thus not available for morphological assessment. Strain CBS 144210, however, is known as a pathogen of the ‘purple witchweed’, a parasite plant common to sub-Saharan Africa and known to devastate *Sorghum bicolor* (L.) Moench and *Oryza sativa* L. plantations (O’Donnell et al. 2000; Yoshida et al. 2010). As previously demonstrated by O’Donnell et al. (2000), our phylogenetic results showed that...
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the clade comprising *F. fredkrugeri* and its sister species *F. dlaminii* does not cluster within the main African core of species in the FFSC. Thus, despite the African origin of our isolates, the predicted biogeographic patterns did not match the observed phylogeny. It has been hypothesised that this should not be the result of genetic markers tracing different phylogenies, but the consequence of losing the phylogenetic signal due to saturated sites and introns (O’Donnell et al. 2000). However, the inclusion in our analysis of additional, highly informative and slowly evolving loci such as *RPB1* and *RPB2* yielded similar results, which points out the need to re-evaluate the phylogeographic arrangement of this important species complex including the vast new data generated during the last 20 years that challenges the established assumptions (Kvas et al. 2009; Walsh et al. 2010; O’Donnell et al. 2013; Laurence et al. 2015). Nevertheless, although rather unlikely, alternative factors such as anthropogenic dispersion of *F. fredkrugeri*, its host or additional invasive alternative hosts, cannot be rejected as an explanation for the discordance between biogeography and phylogenetic results. However, these scenarios are difficult to imagine given the characteristics of the sampled site, not being an agroecosystem but a protected, isolated zone, with minimal human intervention (Smit et al. 2013).

This study is a new example of how easily new *Fusarium* spp. can be found when mycological studies are directed to neglected natural ecosystems of minimal anthropogenic disturbance (Phan et al. 2004; Leslie and Summerell 2011; Summerell et al. 2011; Burgess 2014, Laurence et al. 2015). Although irrelevant for some researchers, finding and properly describing new species, regardless of whether they have little or no pathogenic or mycotoxigenic potential, is of utmost importance to improve our understanding on the diversity, biogeographic and phylogeographic patterns of such a complex and heterogeneous genus as *Fusarium*. In addition, this study remarks on the significance and need to further stimulate the exploration of conserved, non-manipulated natural environments (supersites) and their potential impact on biodiversity research on the fungal kingdom.

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