

Molecular Phylogeny of the *Pseudallescheria boydii* Species Complex: Proposal of Two New Species†

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Pseudallescheria boydii (anamorph *Scedosporium apiospermum*) is the species responsible for human scedosporiosis, a fungal infection with a high mortality rate and which is difficult to treat. Recently, it has been demonstrated that high genetic variation exists within this species. We have performed a morphological and molecular study involving numerous strains of clinical or environmental origins and from different countries. The analysis of partial sequences of the β -tubulin (two loci) and calmodulin genes and the internal transcribed spacer region of the rRNA gene has demonstrated that *P. boydii* is a species complex. The combined analysis of the sequences of the four loci of 60 strains has showed the presence of 44 haplotypes in the ingroup. Three species morphologically related to *P. boydii* sensu stricto, i.e., *Pseudallescheria angusta*, *Pseudallescheria ellipsoidea*, and *Pseudallescheria fusioidea*, which had previously been considered synonyms, could be differentiated genetically from *P. boydii* in our study. It is relevant that two of the three strains now included in *P. ellipsoidea* have caused invasive infections. The species *Pseudallescheria minutispora* and *Scedosporium aurantiacum* are clearly phylogenetically separated from the other species studied and are here proposed as new. Morphological features support this proposal. All the strains included in *S. aurantiacum* species have a clinical origin, while those included in *P. minutispora* are environmental. Further studies are needed to demonstrate whether all the species included in the *P. boydii* complex have different clinical spectra and antifungal susceptibility.

Pseudallescheria boydii (anamorph *Scedosporium apiospermum*) is a ubiquitous ascomycetous fungus that causes a wide array of human infections that can affect practically all the organs of the body (8). These infections have been known for a long time, but in recent years, a marked increase in severe invasive infections has been noticed, mainly in immunocompromised hosts. The treatment of these infections has not yet been resolved, and the mortality rate is very high (3, 17). One of the most typical features of this species, which is very rare in other pathogenic fungi, is its ability to develop sexual structures on routine culture media. The presence of spherical ascospores (cleistothecia) and fusiform or ellipsoidal ascospores allows easy identification of this species and its differentiation from the other species of *Scedosporium*, *Scedosporium prolificans*, whose sexual state still remains unknown.

On the basis of nuclear DNA-DNA reassociation, some studies have proved that important genetic variation exists in *P. boydii*. Gueho and de Hoog (10) found three infraspecific ecological and clinical groups. Rainer et al. (16) reported the existence of five different small-subunit rRNA gene sequence lengths. Random amplified polymorphic DNA studies also demonstrated that numerous and very different genotypes can be found (7). Other authors have reported considerable differences with respect to growth and sporulation (4, 5, 9). In addition, a high variability in antifungal susceptibility of the different isolates and in their clinical response has been observed (1, 2). All these data seem to suggest that *P. boydii* is

probably a species complex. In recent years, application of the phylogenetic species concept in different biological species of pathogenic fungi has revealed phylogenetic lineages that reflected species divergence (12, 13) and the existence of cryptic species. These putative cryptic species in *P. boydii* can show different pathological behavior and different antifungal susceptibility, so their delimitation and characterization are key in order to choose the appropriate treatment of the severe infections caused by these fungi.

This paper reports the results of a combined phenotypic and phylogenetic study of numerous clinical and environmental strains, including several fresh isolates, of the *P. boydii* species complex and the description of two new species.

MATERIALS AND METHODS

Fungal isolates. Sixty isolates of *Pseudallescheria boydii* and relatives from environmental or clinical sources were included in the study (Table 1). Clinical isolates were provided by different reference culture collections (Centraalbureau voor Schimmelcultures, Utrecht, The Netherlands [CBS], Facultat de Medicina i Ciències de la Salut, Reus, Spain, [FMR], The BCCM/IHEM Biomedical Fungi and Yeasts Collection, Brussels, Belgium [IHEM], Mycothèque de l'Université Catholique de Louvain, Belgium [MUCL], Collection of the Institute for Tropical Medicine, Antwerp, Belgium [RV], and Robert Koch Institute, Berlin, Germany [RKI]) or different physicians. Environmental isolates were generally fresh isolates recovered by the authors from soil samples from different geographical regions, and others were also provided by different culture collections. In addition, reference strains of each of the species *Pseudallescheria angusta*, *Pseudallescheria ellipsoidea*, and *Pseudallescheria fusioidea* were included in the study. Isolates were stored at 4 to 7°C until morphological or molecular studies were performed.

Isolation from soil. Soil samples were collected mainly from the superficial layer of soil (mainly garden soils) by using sterilized polyethylene bags. These bags were closed by rubber bands and labeled. Suspensions of this material were cultured on the selective medium Dichloran Rose-Bengal chloramphenicol agar (Oxoid, United Kingdom) with benomyl added at a final concentration of 10 μ g/ml and incubated at room temperature. When typical colonies of *P. boydii* were observed, we tried to isolate them in pure culture.

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TABLE 1. Isolates included in the study and their origins

Isolate	Source	GenBank accession no.			
		BT2	TUB	CAL	ITS
FMR 4072	River sediment, Tordera River, Spain	AJ889592	AJ890122	AJ890216	AJ888384
FMR 4167	Otitis, Valladolid, Spain	AJ888952	AJ889580	AJ890152	AJ888385
FMR 6694	Cerebral abscess, Barcelona, Spain	AJ888953	AJ889581	AJ890153	AJ888386
FMR 6697	Sputum, Madrid, Spain	AJ888954	AJ889582	AJ890154	AJ888387
FMR 6918	Garden soil, Barcelona, Spain	AJ888955	AJ889583	AJ890155	AJ888433
FMR 6919	Garden soil, Barcelona, Spain	AJ888956	AJ889985	AJ890156	AJ888388
FMR 6920	Garden soil, Barcelona, Spain	AJ888957	AJ889986	AJ890157	AJ888434
FMR 6921	Garden soil, Barcelona, Spain	AJ888958	AJ889987	AJ890158	AJ888389
FMR 6922	Garden soil, Barcelona, Spain	AJ888959	AJ889988	AJ890159	AJ888390
FMR 7884	Transplant, Madrid, Spain	AJ888959	AJ890126	AJ890209	AJ888391
FMR 7885	Pleural liquid, Madrid, Spain	AJ888960	AJ889989	AJ890160	AJ888392
FMR 8521	Forest soil, Montsia, Spain	AJ888953	AJ890007	AJ890178	AJ888430
FMR 8522	Forest soil, Deltebre, Spain	AJ888954	AJ890008	AJ890179	AJ888431
FMR 8530	Cultivated soil, Deltebre, Spain	AJ888957	AJ890011	AJ890182	AJ888406
FMR 8532	Cultivated soil, Montsia, Spain	AJ888958	AJ890012	AJ890183	AJ888407
FMR 8534	Cultivated soil, Montsia, Spain	AJ888959	AJ890013	AJ890185	AJ888408
FMR 8535	Soil, Buenos Aires, Argentina	AJ888960	AJ890014	AJ890186	AJ888409
FMR 8537	Soil, Buenos Aires, Argentina	AJ888961	AJ890110	AJ890187	AJ888410
FMR 8538	Soil, Buenos Aires, Argentina	AJ888962	AJ890111	AJ890188	AJ888443
FMR 8539	Soil, Buenos Aires, Argentina	AJ888963	AJ890112	AJ890189	AJ888411
FMR 8540	Soil, Buenos Aires, Argentina	AJ888964	AJ890113	AJ890190	AJ888412
FMR 8541	Soil, Buenos Aires, Argentina	AJ889605	AJ890128	AJ890215	AJ888413
FMR 8619	Keratitits, Brazil	AJ888958	AJ890115	AJ890192	AJ888416
FMR 8620	Keratitits, Brazil	AJ889585	AJ890116	AJ890193	AJ888417
FMR 8621	Cystic fibrosis, Barcelona, Spain	AJ888958	AJ890117	AJ890194	AJ888418
FMR 8622	Foot skin, Barcelona, Spain	AJ889587	AJ890118	AJ890195	AJ888419
FMR 8623	Leukemic patient, Barcelona, Spain	AJ888959	AJ890125	AJ890210	AJ888427
FMR 8625	Leukemic patient, Zaragoza, Spain	AJ889588	AJ890119	AJ890196	AJ888420
FMR 8630	Ulcer of ankle, Santiago de Compostela, Spain	AJ888959	AJ890133	AJ890219	AJ888440
RV 43605	Human, Zaire	AJ888951	AJ889579	AJ890341	AJ888383
IHEM 14263	Cystic fibrosis, patient 1, Angers, France	AJ888961	AJ889990	AJ890161	AJ888436
IHEM 14268	Cystic fibrosis, patient 4, Giens, France	AJ888962	AJ889991	AJ890162	AJ888393
IHEM 14354	Cystic fibrosis, patient 7, Giens, France	AJ888963	AJ889992	AJ890163	AJ888437
IHEM 14358	Cystic fibrosis, patient 9, Tours, France	AJ888964	AJ889993	AJ890164	AJ888438
IHEM 14451	Cystic fibrosis, patient 3, Giens, France	AJ888965	AJ889994	AJ890165	AJ888394
IHEM 14462	Cystic fibrosis, patient 8, Tours, France	AJ888966	AJ889995	AJ890166	AJ888395
IHEM 14464	Cystic fibrosis, patient 8, Tours, France	AJ888967	AJ889996	AJ890167	AJ888396
IHEM 14467	Cystic fibrosis, patient 9, Tours, France	AJ888968	AJ889997	AJ890168	AJ888397
IHEM 14638	Cystic fibrosis, patient 1, Angers, France	AJ888969	AJ889998	AJ890169	AJ888398
IHEM 14754	Cystic fibrosis, patient 8, Tours, France	AJ888970	AJ889999	AJ890170	AJ888399
IHEM 14758	Cystic fibrosis, patient 1, Angers, France	AJ888984	AJ890000	AJ890171	AJ888400
IHEM 15144	Cystic fibrosis, patient 8, Tours, France	AJ888987	AJ890001	AJ890172	AJ888401
IHEM 15149	Cystic fibrosis, patient 5, Giens, France	AJ888988	AJ890002	AJ890173	AJ888402
IHEM 15458	Cystic fibrosis, patient 6, Giens, France	AJ889599	AJ890135	AJ890221	AJ888441
IHEM 15579	Cystic fibrosis, patient 2, Angers, France	AJ889600	AJ890136	AJ890222	AJ888439
IHEM 15642	Cystic fibrosis, patient 4, Giens, France	AJ888949	AJ890003	AJ890174	AJ888403
MUCL 6106	Forest soil, Haasrode, Belgium	AJ888950	AJ890004	AJ890175	AJ888404
MUCL 8302	Soil, Germany	AJ888955	AJ890009	AJ890180	AJ888442
MUCL 8522	Humic soil, Baarn, the Netherlands	AJ889589	AJ890120	AJ890208	AJ888421
MUCL 14009	Forest soil, Yangambi, Zaire	AJ888951	AJ890005	AJ890176	AJ888422
MUCL 14092	Forest soil, Yangambi, Zaire	AJ889602	AJ890130	AJ890213	AJ888429
MUCL 18784	Treated wood (<i>Coelocarpon preussi</i>), Ivory Coast	AJ888956	AJ890010	AJ890181	AJ888405
MUCL 20263	Greenhouse soil, Herverlee, Belgium	AJ888952	AJ890006	AJ890177	AJ888423
MUCL 29258	Fuel oil, Antwerpen, Belgium	AJ889593	AJ890123	AJ890217	AJ888424
RKI 2782/95	Trauma and sepsis, Hamburg, Germany	AJ889598	AJ890134	AJ890220	AJ888432
RKI 2956/93	Bronchoalveolar lavage fluid, Berlin, Germany	AJ888965	AJ890114	AJ890191	AJ888415
CBS 101.22	Mycetoma, Texas	AJ889590	AJ890121	AJ890207	AJ888435
CBS 106.53	Goat dung, Aligarh, India	AJ889601	AJ890131	AJ890212	AJ888428
CBS 254.72	Half-digested sewage tank, Ohio	AJ889604	AJ890129	AJ890214	AJ888414
CBS 311.72	Brown sandy soil, Tsintsabis, Namibia	AJ889603	AJ890132	AJ890218	AJ888425
CBS 418.73	Soil, Tadjikistan	AJ889595	AJ890124	AJ890211	AJ888426
FMR 7294	Blood, Australia	AJ888959	AJ890127	AJ890223	AJ888444

DNA extraction, amplification, and sequencing. DNA was extracted and purified directly from fungal colonies according to the Fast DNA kit protocol (Bio101, Vista, Calif.), with a minor modification that consisted of the homogenization step repeated three times with a FastPrep FP120 instrument (Thermo Savant, Holbrook, N.Y.). The DNA was quantified with GeneQuant *pro* (Amersham Pharmacia Biotech, Cambridge, England). The internal transcribed spacer (ITS) region of the nuclear rRNA gene was amplified with the primer pair ITS5 and ITS4 (21), a fragment of the nuclear gene calmodulin (CAL) was amplified with the degenerated primer pair CL1 and CL2A (14), and two regions within the β -tubulin gene, BT2 and TUB, were amplified using the degenerated primer pair BT2-F (5'-GG(CT)AACCA(AG)AT(ATC)GGTGC(CT)GC(CT)-3') and BT2-R (5'-ACCTC(AG)GTGTAGTGACCCTTGGC-3') and TUB-F/TUB-R (6), respectively.

The PCR mixture (25 μ l) included 20 to 60 ng of fungal DNA template, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, and 2.5 mM MgCl₂ (10 \times Perkin-Elmer buffer II plus MgCl₂ solution; Roche Molecular Systems, Branchburg, N.J.), 100 μ M each deoxynucleoside triphosphate (Promega, Madison, Wis.), 1 μ M of each primer, and 1.5 U of AmpliTaq DNA polymerase (Roche). The amplification program included an initial denaturation step at 94°C for 5 min followed by 35 cycles of denaturation at 95°C for 30 s, annealing for 1 min at 50°C (ITS), 55°C (CAL and TUB), or 60°C (BT2), and extension for 1 min at 72°C. A final extension step at 72°C for 7 min was included at the end of the amplification. After PCR, the products were purified with a GFX™ PCR DNA purification kit (Pharmacia Biotech, Cerdanyola, Spain) and stored at -20°C until they were used in sequencing.

The protocol for sequencing was the Taq DyeDeoxy Terminator Cycle Sequencing kit (Applied Biosystems, Gouda, The Netherlands). Reactions were run with a 310 DNA sequencer (Applied Biosystems). The consensus sequences were obtained using the Autoassembler program (Applied Biosystems).

Phylogenetic analysis. The sequences were aligned using the Clustal X (version 1.8) computer program (19) followed by manual adjustments with a text editor. Most-parsimonious tree (MPT) analyses were performed by using PAUP* version 4.0b10 (18). One hundred heuristic searches were conducted with random sequence addition and tree bisection reconnection branch-swapping algorithms, collapsing zero-length branches and saving all minimal-length trees (MulTrees) on different data sets. *Scedosporium prolificans* (FMR 7294) and *Pseudallescheria africana* (CBS 311.72) were chosen as the outgroup. Regions of sequences with ambiguous alignments were excluded from all analyses (ITS, positions 58 and 59; BT2, positions 90 to 126), and gaps were treated as missing data. Support for internal branches was assessed using a heuristic parsimony search of 500 bootstrapped data sets. The combined data set was tested for incongruence with the partition homogeneity test (PHT), as implemented in PAUP*. To avoid detecting incongruence that is expected within lineages, partition homogeneity tests were restricted to data sets containing only 20 individuals that represented the main lineages (CBS 254.72, FMR 4072, FMR 4167, FMR 6694, FMR 6697, FMR 6920, FMR 6921, FMR 7884, FMR 8532, FMR 8540, FMR 8541, FMR 8623, FMR 8625, FMR 8630, IHEM 14268, IHEM 14467, IHEM 15458, MUCL 14009, RKI 2956/93, RKI 2782/95, and RV 43605).

To test alternative phylogenetic relationships, the Kishino-Hasegawa maximum-likelihood ratio test (11) was performed, as implemented in PAUP*.

Morphological study. The fungi were subcultured on potato dextrose agar (PDA; Difco Laboratories, Detroit, Mich.) for macroscopic examination and growth rates at 25, 37, 40, 42, 45, and 50°C in darkness. For the study of microscopic characteristics, they were cultivated on oatmeal agar (OA) (30 g oat flakes, 1 g MgSO₄ · 7H₂O, 1.5 g KH₂PO₄, 15 g agar, 1 liter tap water). The microscopic features were determined by making wet mounts with lactic acid, which were then examined under a light microscope (Leitz Dialux 20).

Nucleotide sequence accession numbers. All the sequences obtained were deposited in the GenBank database. Accession numbers are shown in Table 1.

RESULTS

Phylogeny. Sixty isolates of *P. boydii* and relatives were chosen to examine species limits and evolutionary relationships among them. With the primers used, we were able to amplify and sequence 522 bp, 419 bp, 549 bp, and 570 bp of the ITS, BT2, TUB, and CAL loci, respectively. Of the 2,060 nucleotides sequenced, 386 characters (18.73%) were parsimony informative in the different *P. boydii* isolates. The lowest number was 42 in the ITS region, and the highest was 160 in CAL

region. Sequences of the four region genes were analyzed phylogenetically as separate and combined data sets.

Phylogenetic analysis of the 419-bp BT2 data set yielded 12 MPT, which resulted in a total of 25 haplotypes (Fig. 1). There were 246 constant, 93 parsimony-informative, and 77 variable parsimony-uninformative characters in this fragment. The type strains of *P. fusioidea*, *P. ellipsoidea* and *P. angusta* were interspersed with the isolates of *P. boydii*. Four main highly supported clades (100%) were shown: the basal one comprising 4 European clinical isolates; two other small clades made up of 5 and 2 almost exclusively environmental isolates; and the biggest one, which comprised the 49 remaining isolates. However, inside the latter, another six terminal branches were present, each of them supported by a 100% bootstrap.

Parsimony analysis of the TUB data set yielded 420 MPT with 203 steps in length, in which 18 nodes received 100% bootstrap support. There were 420 constant, 90 parsimony-informative, and 39 variable parsimony-uninformative characters in this set. These trees resulted in a total of 22 different haplotypes (Fig. 2). Although the tree topology was slightly different from that of the previously mentioned locus, the three small, most-basal clades formed were also maintained here. Two of these clades, formed by four clinical and two environmental isolates, respectively, were the most phylogenetically distant.

Analysis of CAL and ITS sequences yielded a single MPT of 377 steps in length and 5,000 MPT of 97 steps in length, respectively. The numbers of haplotypes observed were 21 in the CAL tree (Fig. 3) and 19 in the tree based on ITS sequences (Fig. 4). Both trees showed similar topologies to that of the BT2 tree. The two above-mentioned basal clades were also placed here away from the other isolates. Overall, the ITS rRNA gene data set is considerably less informative for phylogenetic reconstruction than the other three markers.

The result of the partition homogeneity test showed that the sequence data sets for the four loci were congruent ($P = 0.07$) and could therefore be combined. A total of 2,496 MPT were produced from a heuristic search using the combined data set of 2,060 characters from the four loci (Fig. 5). From these characters, 1,440 were constant, 386 were parsimony informative, and 234 were variable parsimony noninformative. Clustering was similar to that observed in the particular trees of the different genes analyzed. A total of 44 haplotypes were shown. Most nodes in the combined analysis showed increased clade support as measured by bootstrapping (20 nodes with 100% bootstrap support). As within the ITS, TUB, and CAL gene trees, two clades were identified as the basal-most lineages (clades 1 and 2), each of them with a bootstrap support of 100%. Phylogenetic analysis of the remaining monophyletic ingroup taxa (bootstrap, 100%) identified a basal clade (clade 3) and two bigger clades (clades 4 and 5), all them with 100% bootstrap support. The type strains of *P. boydii*, *P. ellipsoidea*, *P. fusioidea*, and *P. angusta* were placed in clade 5.

Morphology. All the 60 isolates that constituted the ingroups in the different trees obtained in our phylogenetic analyses were clearly identified by the presence of a characteristic *Scedosporium* anamorph. Although no relevant morphological differences were observed among them, the isolates that constitute clade 1 showed narrower conidia (2 to 5 μ m wide) than the rest (3 to 6 μ m wide). In addition, the members of this

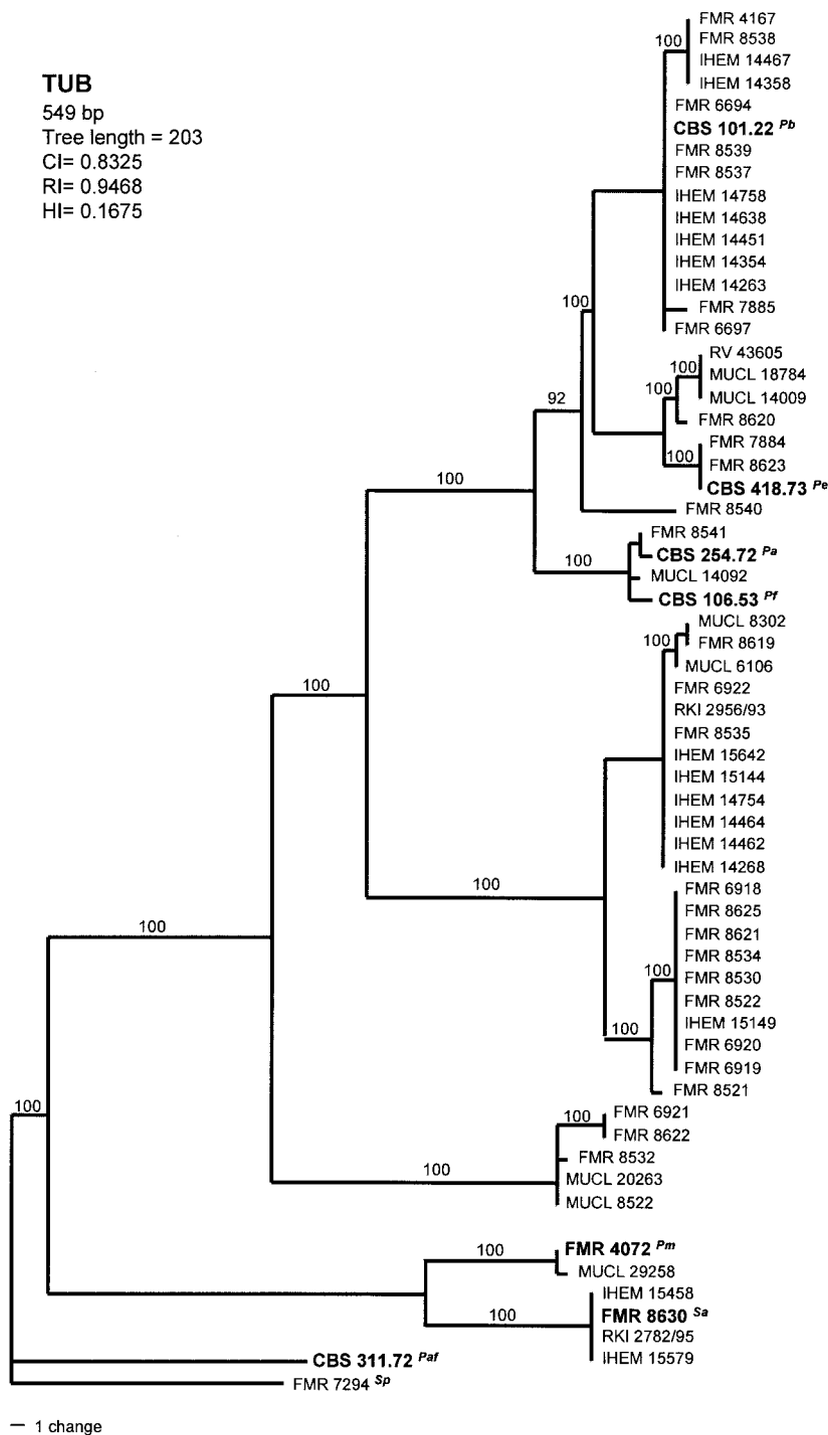


FIG. 2. One of the 420 most-parsimonious trees obtained from heuristic searches based on TUB sequence. Bootstrap support values above 90% are indicated at the nodes. Type strains are indicated with boldface type. *P. africana* and *S. prolificans* were used as outgroups. *Pb*, *P. boydii*; *Pe*, *P. ellipsoidea*; *Pf*, *P. fusioidea*; *Pa*, *P. angusta*; *Pm*, *P. minutispora*; *Sa*, *S. aurantiacum*; *Paf*, *P. africana*; *Sp*, *S. prolificans*; CI, consistency index; RI, retention index; HI, homoplasy index.

(Table 2). Another typical feature, common to the half of the isolates included in the study, was the development of a second type of anamorph, namely, *Graphium* sp. *Graphium* is characterized by the production of synnemata terminated in a slimy head of conidia. The size of the synnemata was very variable

(80 to 750 μm long) and depended on the culture medium used and the age of the culture. The production of *Graphium* was not exclusive of any clade since the isolates that produced them were distributed in all the five clades (Fig. 5). However, it was most common in clade 4 (in 14 of the 22 isolates) and in the

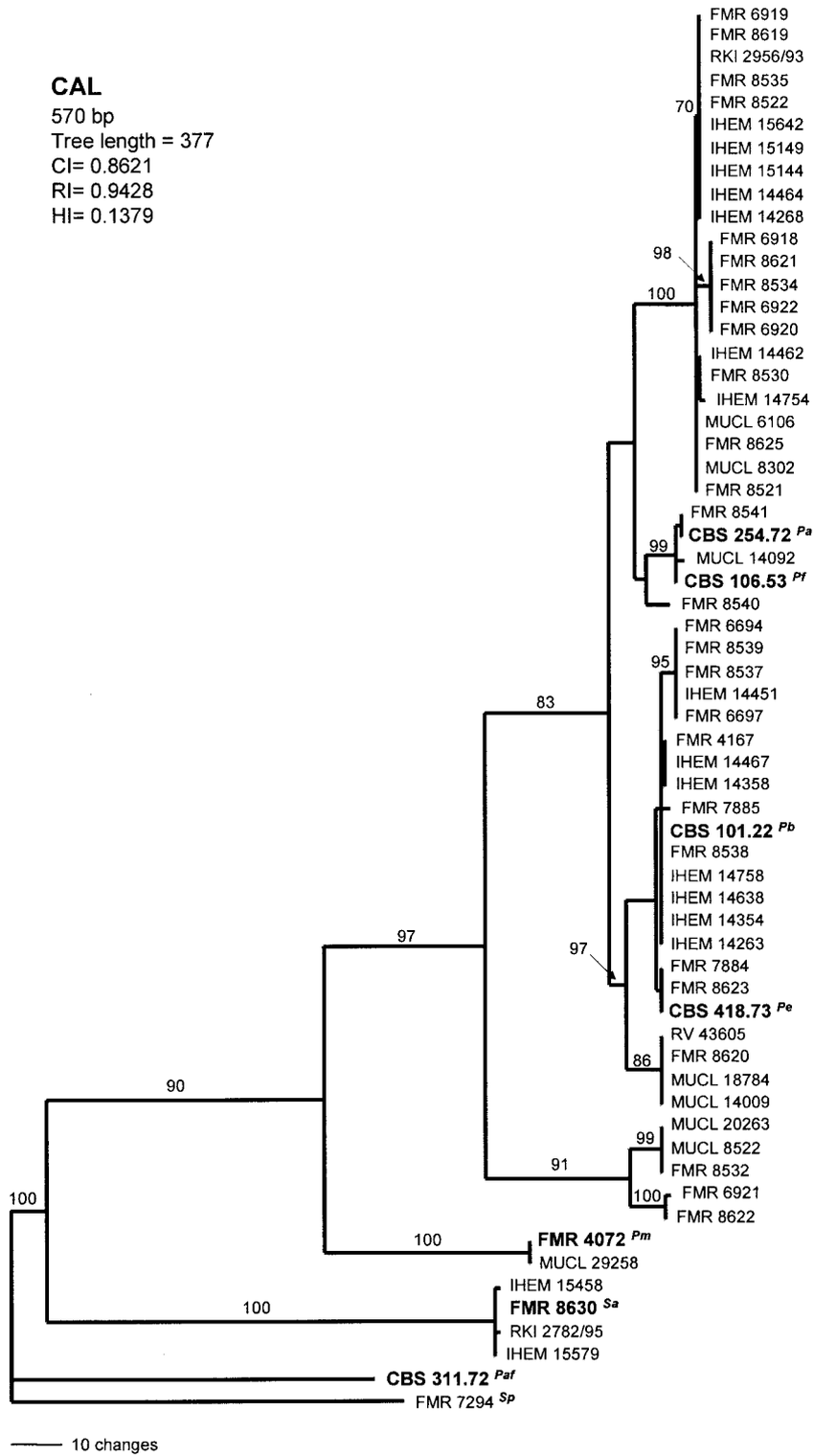


FIG. 3. The single most-parsimonious tree obtained from heuristic searches based on CAL sequence. Bootstrap support values above 70% are indicated at the nodes. Type strains are indicated with boldface type. *P. africana* and *S. prolificans* were used as outgroups. *Pb*, *P. boydii*; *Pe*, *P. ellipsoidea*; *Pf*, *P. fusoidea*; *Pa*, *P. angusta*; *Pm*, *P. minutispora*; *Sa*, *S. aurantiacum*; *Par*, *P. africana*; *Sp*, *S. prolificans*; CI, consistency index; RI, retention index; HI, homoplasy index.

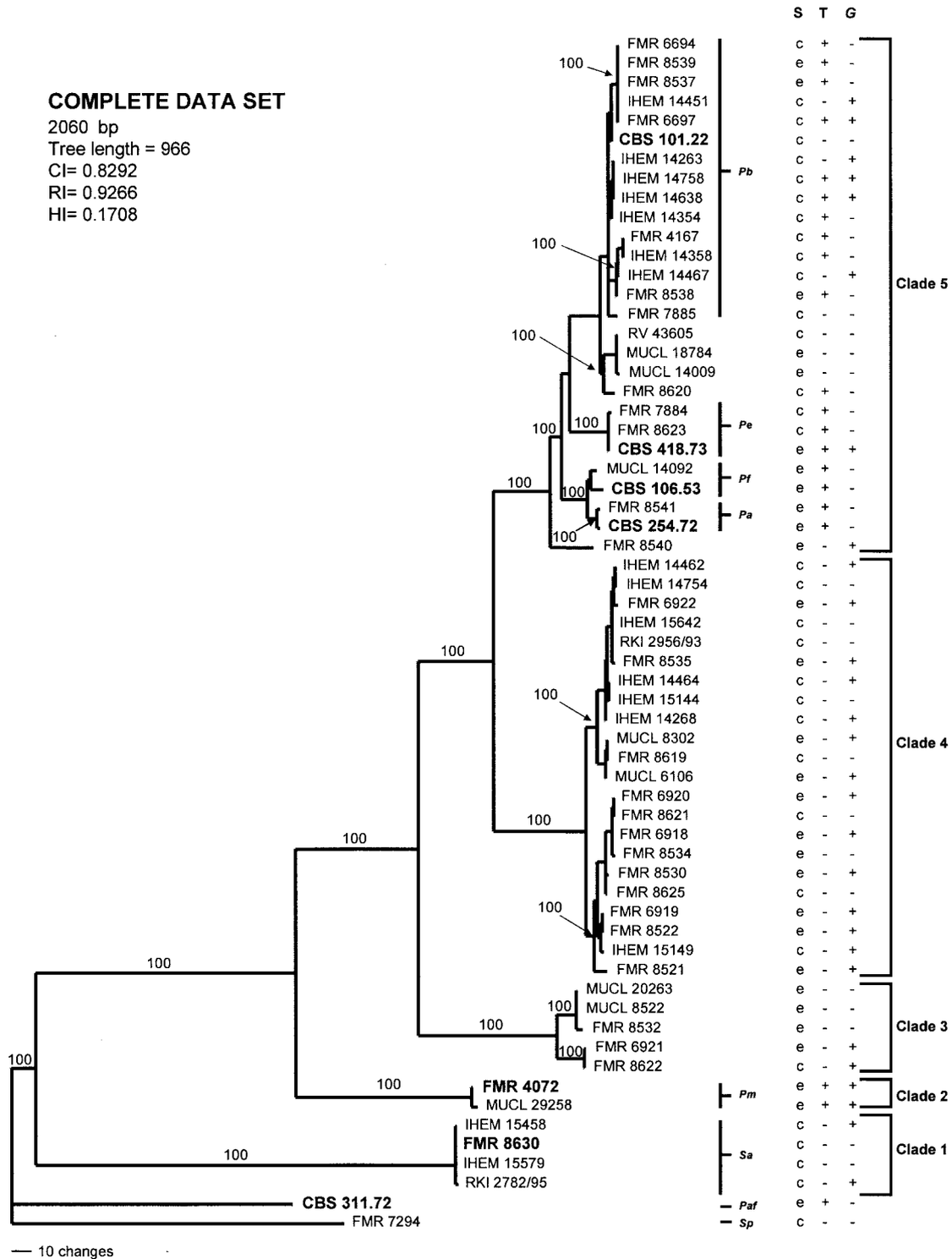


FIG. 5. One of the 2,496 most-parsimonious trees obtained from heuristic searches based on analysis produced from the combined data set. Bootstrap support values of 100% are indicated at the nodes. Type strains are indicated with boldface type. *P. africana* and *S. prolificans* were used as outgroups. *Pb*, *P. boydii*; *Pe*, *P. ellipsoidea*; *Pf*, *P. fusioidea*; *Pa*, *P. angusta*; *Pm*, *P. minutispora*; *Sa*, *S. aurantiacum*; *Paf*, *P. africana*; *Sp*, *S. prolificans*; S, source; T, teleomorph; G, *Graphium* anamorph; CI, consistency index; RI, retention index; HI, homoplasy index; c, clinical; e, environmental; +, presence; -, absence.

TABLE 2. Relevant features to differentiate the clinical species of *Pseudallescheria/Scedosporium* spp.

Species	Diffusible pigment on PDA at 25°C	Maximum growth temp (°C)	Teleomorph		Anamorph	
			Development of ascomata	Shape of ascospores	Shape of conidiogenous cells	Most common shape of conidia borne on vegetative hyphae
<i>P. boydii</i>	–	40	±	Broadly fusiform	Cylindrical	Globose
<i>P. ellipsoidea</i>	–	40	+	Ellipsoidal	Cylindrical	Ellipsoidal
<i>S. aurantiacum</i>	+ (yellow)	45	–		Cylindrical or slightly flask-shaped	Obovoid
<i>S. prolificans</i>	–	40	–		Flask-shaped	Globose

that clades 1 and 2 represent two different species from those up to now accepted in *Scedosporium* and *Pseudallescheria*, respectively, and are consequently here proposed as new.

***Scedosporium aurantiacum*.** Gilgado, Cano, Gené, et Guarro, sp. nov. (Fig. 6).

Etym.: referred to the yellow color of the diffusible pigment of the colonies. Coloniae in agar dense, plerumque gossypinae, luteolus vel brunneolus griseae; reversum brunneolus aurantiacum. Pigmentis flavis in culturis formantis. Conidiophora solitaria vel synnemata. Synnemata erecta, 330 to 750 µm alta, cum caule atro griseo, cylindrico, 7.5 to 17.5 µm lato, et capitulo mucoso usque ad 60 µm alto, 70 to 140 µm lato. Conidia obovoidea, subcylindrica vel claviformia, 5 to 14 by 2 to 5 µm. Conidia sessilis copiosae, laterales, unicellularia, brunnea, plerumquam obovoidea, 6 to 10 by 3 to 5 µm. Teleomorphosis ignota. Holotypus, IMI 392886, ex mycosis humanum (cultura viva, FMR 8630, IHEM 21147, CBS 116910).

The colonies on PDA attained a diameter of 40 to 50 mm after 14 days at 25°C. They were dense and usually cottony, but in some isolates they were lanose, especially at the center, frequently showing a concentric growth of aerial mycelium of different colors, yellowish gray combined with brownish gray areas, usually with a whitish, irregularly lobate and fimbriate margin, and the reverse was brownish orange at the center and brown to colorless towards the periphery. All isolates produced a light yellow diffusible pigment on PDA and OA after a few days of incubation. Conidiophores were solitary on aerial mycelium or grouped to form synnemata (*Graphium*) mainly on the agar surface. Solitary conidiophores were often reduced to a conidiogenous cell growing laterally on undifferentiated mycelium or branched, usually bearing verticils of two to three conidiogenous cells. Synnemata were present only in the isolates IHEM 15458 and RKI 2782/95. They were erect, 330 to 750 µm long, consisting of a cylindrical stipe 7.5 to 17.5 µm wide, dark gray, smooth-walled, and slightly roughened apically, and they terminated in a slimy head of conidia, up to 60 µm long by 70 to 140 µm wide. The conidiogenous cells were percurrent, lateral, or terminal, subhyaline, smooth-walled, cylindrical, or slightly flask shaped, 10 to 37 µm long by 1.5 to 2.5 µm wide, less frequent intercalary as a lateral projection on hyphae, and up to 5 µm long by 2 µm wide. There were three types of conidia: (i) those produced on solitary conidiophores were subhyaline, smooth-walled, obovoid, or subcylindrical, and 5 to 14 µm by 2 to 5 µm; (ii) those produced on synnemata were predominantly cylindrical or claviform, 6 to 12 µm by 3 to 5 µm with a wide truncate base; (iii) those developed mainly from the undifferentiated hyphae of the substrate were sessile

or on short protrusions, solitary, lateral, brown, smooth, and thick-walled, usually obovoid, 6 to 10 µm long by 3 to 5 µm wide. The latter were abundantly produced by all isolates. Teleomorph was unknown.

The optimum growth temperature was from 37°C to 40°C with colonies on PDA attaining a diameter up to 60 to 67 mm after 14 days. Maximum growth was at 45°C. The fungus did not grow at 50°C.

***Pseudallescheria minutispora*.** Gilgado, Gené, Cano, et Guarro, sp. nov. (Fig. 7).

Etym.: referred to the small size of the ascospores. Coloniae in agar dense, gossypinae vel lanosae, aurantium griseae vel brunneolus griseae; reversum incoloratum. Pigmentis in culturis non formantis. Ascomata solitaria, non ostiolata, globosa vel subglobosa, 50 to 150 µm, luteolus vel brunneolus grisea, cum peridium ex textura epidermoidea. Asci octospori, globosi vel subglobosi, 12 to 15 by 10 to 13 µm, evanescentes. Ascosporeae unicellulares, subhyalinae vel dilute brunneae, laeves, tenuitunicatae, ellipsoideae, 5 to 7 by 3 to 4 µm. Conidiophora solitaria vel synnemata. Synnemata erecta, 180 to 300 µm alta, cum caule atrobunneo, cylindrico, 7.5 to 17.5 µm lato, et capitulo mucoso, 60 to 100 µm alto et 80 to 170 µm lato. Conidia obovoidea, ellipsoidea, cylindrica vel claviformia, 5 to 14 by 2.5 to 4.5 µm. Conidia sessilis parvasae, laterales, unicellularia, subhyalina, plerumquam obovoidea, 7 to 10 by 2.5 to 5 µm. Holotypus, IMI 392887, ex sedimentis fluvialibus (cultura viva, FMR 4072, IHEM 21148, CBS 11691).

The colonies on PDA attained a diameter of 50 to 57 mm after 14 days at 25°C. They were dense, cottony to lanose, orange gray combined with brownish gray areas, with a whitish, lobate or irregular, and fimbriate margin; the reverse was colorless. Diffusible pigment was absent. All isolates developed abundant ascomata on OA. The ascomata were solitary, non-ostiolate, globose to subglobose, and 50 to 150 µm in diameter, with a peridium of textura epidermoidea, yellowish gray to brownish gray, and often covered with brown, thick-walled septate, 2.2 to 3 µm wide. The asci were eight-spored, globose to subglobose, and 12 to 15 µm long by 10 to 13 µm wide with evanescent walls. Ascospores were unicellular, subhyaline to light brown, smooth and thin-walled, ellipsoidal, and 5 to 7 µm long by 3 to 4 µm wide, with a germ pore at each pole and usually with oil drops. Both isolates developed the two typical anamorph simply or scarcely branched conidiophores, up to 35 µm long. Its conidia were subhyaline to light brown, smooth-walled, obovoid, ellipsoidal or subclaviform, and 6 to 11 µm long by 3 to 4 µm wide. The *Graphium* anamorph produced synnemata which were erect and 180 to 300 µm long, with a

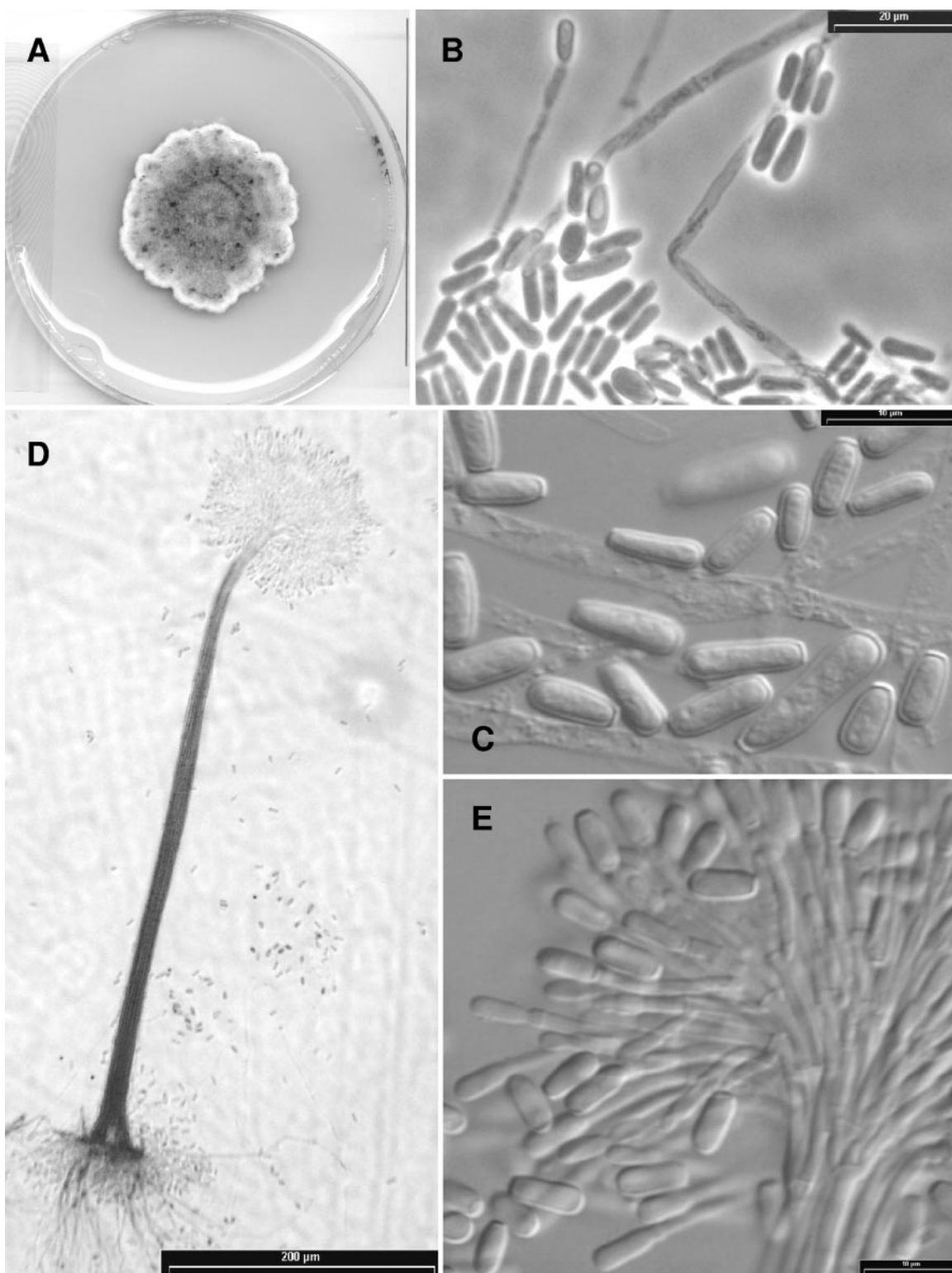


FIG. 6. *Scedosporium aurantiacum* isolates (A, D, E) IHEM 15458 and (B, C) FMR 8630. (A) Colony growing on PDA after 14 days of incubation at 25°C. (B, C) A conidiogenous cell and conidia from solitary conidiophores. (D) A synnema of the *Graphium* anamorph. (E) Apical part of a synnema producing conidia.

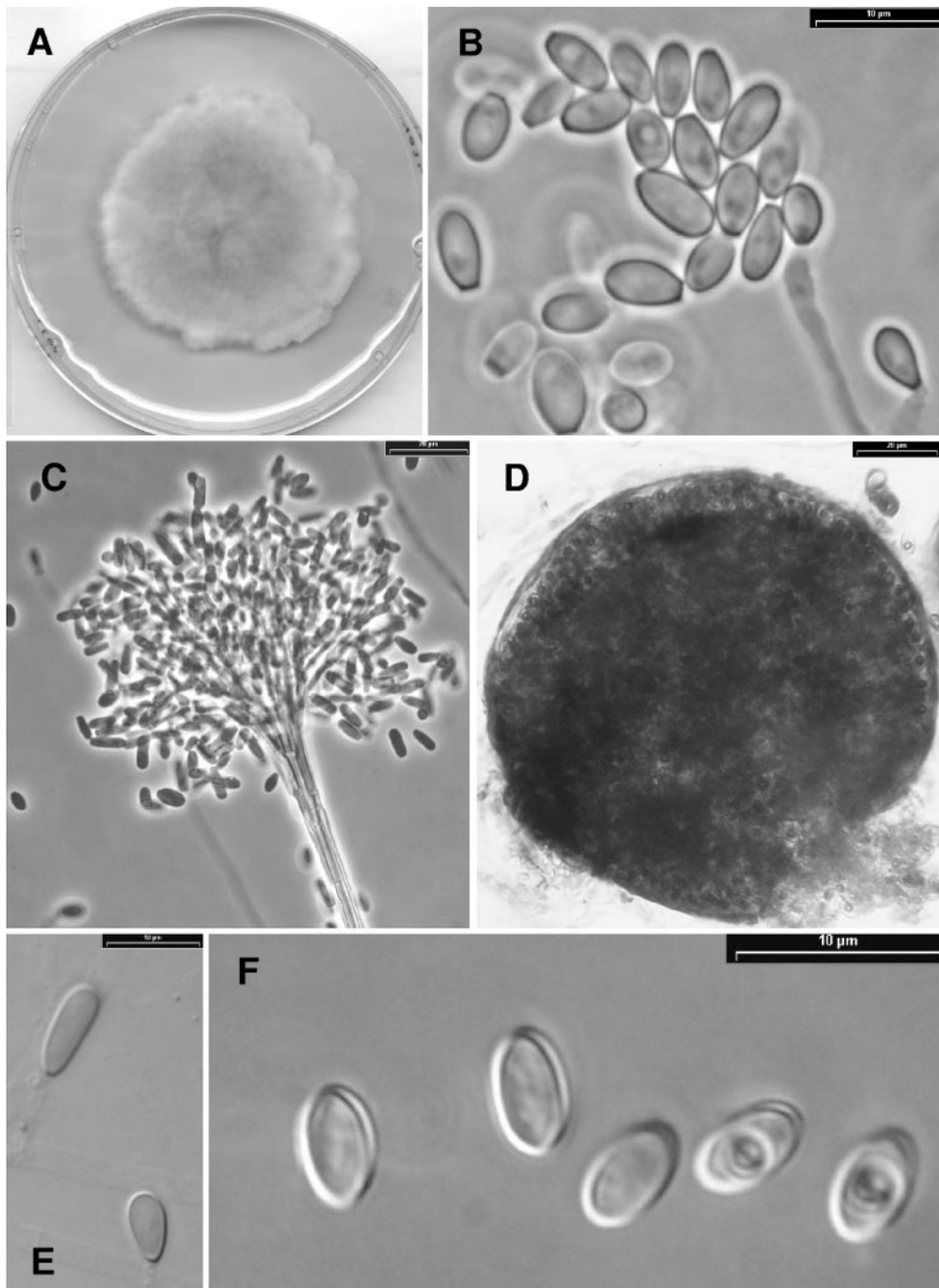


FIG. 7. *Pseudallescheria minutispora* strain FMR 4072. (A) Colony growing on PDA after 14 days of incubation at 25°C. (B) Conidiogenous cells and conidia of the *Scedosporium* anamorph. (C) Apical part of a synnema of the *Graphium* anamorph producing conidia. (D) Ascoma. (E) Conidia borne on undifferentiated hyphae. (F) Ascospores.

cylindrical stipe that was 7.5 to 17.5 µm wide, and they were smoke brown, smooth-walled, slightly roughened apically, slightly inflated at the base, and up to 25 µm wide and terminated in a slimy head of conidia that was 60 to 100 µm long by 80 to 170 µm wide. The conidia were predominantly cylindrical or claviform, 5 to 14 µm long by 2 to 4.5 µm wide, with a wide truncate base. The conidiogenous cells were percurrent, lateral or terminal, subhyaline, smooth-walled, usually cylindrical, 10 to 35 µm long by 1.5 to 2 µm wide, and less frequently inter-

calary as a lateral projection on hyphae, up to 6 µm long by 2 µm wide. Conidia from undifferentiated hyphae were scarcely produced. They were lateral, usually sessile, subhyaline, smooth and thick-walled, ellipsoidal to obovoid, and 7 to 10 µm long by 3.5 to 5 µm wide.

The optimum growth temperature was from 25°C to 30°C with colonies on PDA attaining a diameter up to 50 to 61 mm after 14 days. Maximum growth was at 40°C. The fungus was unable to grow at 42°C.

DISCUSSION

DNA sequences from four loci were analyzed to investigate phylogenetic relationships and species limits within the *P. boydii* species complex. Until recently, the genus *Pseudallescheria* was considered to comprise the following seven species (20): *P. africana*, *P. angusta*, *P. boydii*, *P. desertorum*, *P. ellipsoidea*, *P. fimeti*, and *P. fusioidea*. All of these species are morphologically very similar, and the main distinction among them is based on the size of the cleistothecia and ascospores (20). Recently, Rainer et al. (16), using different molecular techniques, concluded that *P. ellipsoidea*, *P. fusioidea*, and *P. angusta* are probable synonyms of *P. boydii*. In our study, we have included the type strains of these three species, and the results confirmed that all of them are genetically and morphologically different from *P. boydii*. We have also detected a high number of phylogenetic species, but only two of them can be clearly recognized morphologically.

The information provided by the four loci analyzed was very similar, which proved to be excellent phylogenetic markers for species level systematics within *Pseudallescheria*. The less informative locus was ITS, which only resolved 9 phylogenetically distinct species, whereas CAL, BT2, and TUB resolved 12, 14, and 15 species, respectively. Apart from *P. africana*, which was the outgroup of the present analysis, the two phylogenetic species most clearly separated were the two proposed as new, i.e., *S. aurantiacum*, represented by four clinical isolates, and *P. minutispora*, represented by two environmental isolates. However, we have recently studied another isolate of clinical origin from Germany (RKI 866/94) that genetically and morphologically matches the features of the latter species. The two clades formed by these species were highly supported in all the phylogenetic trees. Up to now, practically all the described species of *Pseudallescheria*, with the exception of *P. boydii*, *P. fusioidea*, and *P. ellipsoidea*, were monotypic; i.e., they are known by only one isolate. In the present study, the type strain of *P. angusta* nested with another soil isolate (FMR 8541) from Argentina. Both isolates showed identical ITS sequences, and those of the other loci studied were only different in 1 or 2 nucleotides (BT2, 1 nucleotide; TUB, 1 nucleotide; CAL, 2 nucleotides). The type strain of *P. fusioidea* nested with an environmental isolate from Zaire in three of the four loci analyzed. Both isolates also showed identical ITS sequences and differed in a few nucleotides in the other genes (BT2, 8 nucleotides; TUB, 4 nucleotides; CAL, 2 nucleotides). Our phylogenetic study revealed that *P. fusioidea* and *P. angusta* were phylogenetically very close. They only differed in 22 bp in the combined data set. However, *P. angusta* showed smaller ascomata (up to 110 μm in diameter versus up to 160 μm in diameter for *P. fusioidea*) and narrower ascospores (3 to 3.5 μm wide versus 4 to 4.5 μm wide for *P. fusioidea*). In addition, the isolates of *P. fusioidea* grew faster than those of *P. angusta* (69 to 70 mm versus 52 to 54 mm at 14 days on PDA at 25°C). The teleomorphs developed by the isolates of *P. angusta*, *P. ellipsoidea*, and *P. fusioidea* were consistent with the morphological features for the species described previously by von Arx et al. (20). However, the isolates that nested with the type strain of *P. boydii* (CBS 101.22) showed larger ascospores (6 to 9 by 5 to 6 μm) than those described previously by von Arx et al. (20) for such species (6 to 7 by 3.5 to 4 μm). Unfortunately, in our

study, the type strain of *P. boydii* only produced the *Scedosporium* anamorph. It is an old strain that has probably lost the ability to develop the sexual state. *P. ellipsoidea* was the species that was genetically and morphologically closest to the group of isolates that nested in the same branch as the type strain of *P. boydii*. However, *P. ellipsoidea* can be distinguished by its ellipsoidal ascospores, while the ascospores of the members of such groups of isolates are broadly fusiform. Moreover, in the former, the conidia from vegetative hyphae are ellipsoidal to obovoid and scarce, while those of the *P. boydii* branch are abundant and predominantly globose to subglobose (Table 2). Further studies of this group of isolates are required in order to define the morphological features of *P. boydii* sensu stricto.

Up to now, *P. boydii* had been considered the only pathogenic species of the genus *Pseudallescheria*, but this study has demonstrated that other phylogenetic species of the *P. boydii* complex also included clinical isolates. However, the clinical strains are not homogeneously distributed in the different clades, and some correlation between the clades and the clinical origin of the strains could be observed instead. Clinical strains were mainly concentrated in three of the five clades. *Scedosporium aurantiacum* (clade 1) grouped European clinical strains exclusively. Clades 4 and 5 were the biggest ones and included numerous strains each. However, 72% of the isolates of clade 5 were clinical, while only 50% of the isolates of clade 4 had such origin. It is noteworthy that two of the three strains included in the *P. ellipsoidea* group (CBS 418.73, FMR 7884, and FMR 8623) had caused disseminated infections, which emphasizes the clinical relevance of this species.

Most of the terminal branches that grouped more than one isolate included clinical and environmental isolates. This seems to demonstrate that any environmental strain can cause infection under the appropriate conditions. Using a murine model of invasive infection by *S. prolificans*, Ortoneda et al. (15) proved that there are no virulence differences between environmental strains and those that caused colonization or infection.

As expected, these results have demonstrated that *P. boydii* does not represent a single species. It encompasses a high number of phylogenetic species, although only a few of them can be recognized morphologically. One of the most important findings of this work is to provide phenotypic features useful for the distinction of some of these species (Table 2). Considering that not all the hospitals have facilities for molecular diagnosis and that not all these species are equally involved in human infections, these results can be especially useful for clinical microbiologists or laboratorians in order to identify these fungi. Judging by the high clinical relevance of this fungal group, further investigation is expected in the near future. It is especially important to determine if these species, and perhaps others, that could be identified in the future using similar approaches and involving more isolates from different sources and geographical regions are equally pathogenic to humans. Knowledge of the degree of virulence of these species and their response to the antifungal drugs may also be very useful in order to choose the appropriate treatment of the severe and refractory infections attributed to *P. boydii* sensu lato. Furthermore, taking into account that many of these species can only be reliably separated through molecular phylogenetics of DNA

sequences, finding morphological apomorphies for their laboratory identification would also be very valuable.

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