RESEARCH ARTICLE

Removing chaos from confusion: assigning names to common human and animal pathogens in Neocosmospora

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Key words

eight new taxa Fusarium Neocosmospora pathogens phylogeny systematics

Abstract The genus Neocosmospora encompasses highly prevalent and aggressive human and animal fungal pathogens. Here we assign formal descriptions and Latin binomials to some of the most clinically relevant phylogenetic species of the genus. Three new species, named Neocosmospora catenata, N. gamsii and N. suttoniana (previously assigned to the informal names 'Fusarium' solani species complex (FSSC) lineages, FSSC 43, FSSC 7 and FSSC 20, respectively) are described on the basis of multilocus phylogenetic analyses (using *EF-1α*, ITS, LSU and RPB2 loci) and morphological characters. Lineage FSSC 9 is conspecific with the ex-type strain of Cylindrocarpon tonkinense, thus the new combination Neocosmospora tonkinensis is proposed. In addition, and based on the latest taxonomy for this generic complex, new combinations are introduced for four medically important taxa: Neocosmospora keratoplastica, N. lichenicola, N. metavorans and N. petroliphila. The most significant distinctive features for all the clinically relevant species treated here are compared and illustrated.

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INTRODUCTION

The genus Neocosmospora (as the 'Fusarium' solani species complex, FSSC) has been a highly renowned fungal group for more than 100 years, mainly because it contains significant plant pathogenic species, including agents of fruit-rot, root-rot and seedling damping-off, affecting diverse plant hosts (Leslie & Summerell 2006, Domsch et al. 2007, Nalim et al. 2011). In the last 50 years, however, this fungal group gradually and persistently became recognised as important in the clinical field. It is now known to contain some of the fungal species that are most clinically relevant as agents infecting immunocompetent hosts. This list of species includes the principal etiologic agents of fungal keratitis, which are often introduced via traumatic inoculation (De Hoog et al. 2000, Godoy et al. 2004, Shukla et al. 2008). In addition are the second most commonly isolated moulds in onychomycosis after the dermatophytes (Ghannoum et al. 2000, Scher et al. 2013). Species in Neocosmospora are also among the most significant pathogens associated with severe infections in transplant recipients and patients with haematological malignancies, persistent neutropenia or immunodepression caused by corticosteroid therapy (Lass-Flörl 2009, Torres & Kontoyiannis 2011, Guarro 2013, Slavin et al. 2015). Although fusarial infections are rare, nearly 50 % of these infections are attributed to Neocosmospora. The most commonly reported species correspond to 'F.' keratoplasticum, 'F.' petroliphilum, N. falciformis (syn. F. falciforme) and N. solani (syn. F. solani); plus several currently unnamed phylogenetic species. These organisms are recovered from diverse cutaneous and subcutaneous infections including arthritis, brain abscess, catheter-associated fungemia, disseminated infections, mycetoma, osteomyelitis, peritoneal dialysis-associated peritonitis and sinusitis, as well

Human pathogenic species in Neocosmospora are also among the most important fusarial agents of veterinary infections (Zhang et al. 2006, O'Donnell et al. 2008, 2010, 2016). Apart from N. solani, other species seem to show some degree of host specialisation. Neocosmospora falciformis has been repeatedly isolated from equine ocular infections, and has also been reported from canines and reptiles (O'Donnell et al. 2016), while 'F'. keratoplasticum and two currently unnamed phylogenetic species (FSSC 12 and FSSC 43) seem to have some adaptation to the marine environment, infecting mostly crustaceans, fish, marine mammals and reptiles (O'Donnell et al. 2016).

The generally high degree of antifungal resistance, variable in vitro susceptibility patterns and unpredictable response to antifungal compounds seen in Neocosmospora infections, coupled with the high virulence described in clinical reports and animal models of infection, are factors often associated with negative outcomes, placing these species among the most devastating fungal agents of human and animal disease (Sugiura et al. 2003, Azor et al. 2007, Araujo et al. 2015, Espinel-Ingroff et al. 2016, Taj-Aldeen et al. 2016).

Phylogenetic studies have shown that Neocosmospora solani, historically linked with human and veterinary disease, do not belong to a discrete taxon but rather represent an extensive evolutionary radiation comprising more than 15 phylogenetic species. With the exception of the four most commonly isolated species, N. falciformis, 'F.' keratoplasticum, 'F.' petroliphilum and N. solani, most of these phylospecies have not been formally described, thus are not linked to scientific names, in part because they are scarcely distinguishable by means of phenotypic comparison. Although phylogenetically well characterised, comprehensive morphological descriptions and diagnoses do not exist for these important lineages, which are currently identified following an informal haplotype nomenclatural system (Zhang et al. 2006, O'Donnell et al. 2008).

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as many other types of infections (Dignani & Anaissie 2004, Garcia et al. 2015, Hiebert et al. 2016).

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The use of Latin binomials is not a common feature for clades containing human and veterinary pathogens in *Neocosmospora*, mainly due to the conflicting taxonomy of the genus, the non-existence of nomenclatural types and the uncertainty of application of previously published names. Moreover, the name 'Fusarium' solani has been traditionally used by clinical microbiologists and plant pathologists as a wildcard to deal with isolates belonging to this complex when molecular tools are not available (Zhang et al. 2006, Nakamura et al. 2007, Bachmeyer 2007, O'Donnell et al. 2016). Meanwhile, new lineages not conforming to an existing haplotype designation are constantly being found (Guevara-Suarez et al. 2016, Melo et al. 2016).

Recently, Schroers et al. (2016) epitypified Neocosmospora solani (basionym: Fusisporium solani) linking this important plant and animal pathogen with clade 5 in FSSC. Al-Hatmi et al. (2018) formally proposed the name 'Fusarium' metavorans for FSSC 6, one of the most prevalent lineages in human disease, while FSSC 12, which includes important veterinary pathogens, is currently under study and will soon be formally described (Geiser pers. comm.). However, several unnamed clades are still in need of formal description, and those containing animal pathogens are of particular importance (O'Donnell et al. 2008, 2016). An accurate identification of pathogenic fusaria is essential for epidemiological purposes and for the prompt establishment of efficacious clinical treatment (Bachmeyer 2007). It is known that antifungal susceptibility in fusaria is variable among closely related taxa, and often isolate-dependant (Alastruey-Izquierdo et al. 2008, Tortorano et al. 2008). This phenomenon has not yet been reported in Neocosmospora (Azor et al. 2007, Bachmeyer 2007), and remains an understudied issue in the genus.

In the present study, we examine a set of isolates previously assigned to five of the most prevalent pathogenic clades in *Neocosmospora* (*'F.' metavorans*, FSSC 7, FSSC 9, FSSC 20 and FSSC 43), along with strains belonging to the most commonly encountered clinically relevant species mentioned above. Latin binomials, detailed illustrations, morphological descriptions and comparisons are provided in order to facilitate identification by clinical microbiologists.

MATERIALS AND METHODS

Strains

Forty-five isolates originally recovered from human and veterinary clinical specimens and belonging to the clades termed 'F.' metavorans, FSSC 7, FSSC 9, FSSC 20 and FSSC 43 as previously defined using multilocus phylogenetic data (Zhang et al. 2006, O'Donnell et al. 2008, 2016), were retrieved from the collections of the Agricultural Research Service, Peoria, IL, USA (NRRL) and the Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands (CBS). For morphological comparisons and phylogenetic analyses, cultures or DNA sequences from 88 additional isolates were included in the study; these isolates were obtained from the CBS, the personal collection of P.W. Crous (CPC) housed at CBS, the Fusarium Research Center housed in The Pennsylvania State University, State College, PA (FRC), the personal collection of Kerry O'Donnell (KOD), the University of Texas Health Science Center, San Antonio, TX (UTHSC), the American Type Culture Collection, Manassas, VA (ATCC), CABI Biosciences, Egham, Surrey, England (IMI) and NRRL (Table 1).

Morphology

Morphological observations and measurements of macro- and microscopic features were performed following the protocols of Aoki et al. (2003, 2005, 2013) with slight modifications as

described previously (Sandoval-Denis et al. 2018). Macroscopic characteristics of fungal growth were evaluated using cornmeal agar (CMA), oatmeal agar (OA) and potato dextrose agar (PDA) (recipes in Crous et al. 2009). Colony morphology, colour, odour and presence of diffusible pigments were recorded after cultures had grown 7 d at 25 °C in darkness, under continuous fluorescent light and using a 12/12 h cool fluorescent light/dark cycle. For growth rate experiments, cultures were made on PDA agar, by transferring 5 × 5 mm agar blocks from 7-d-old cultures growing on synthetic nutrient poor agar (SNA; Nirenberg 1976). These cultures were incubated in darkness at temperatures ranging from 6-40 °C in 3 °C intervals. Growth rates were recorded after 3 and 7 d by measuring the radial colonial size in at least four directions. The micromorphological examination was made using water as mounting medium, with material taken from cultures on SNA with and without sterilised pieces of carnation leaves, incubated at room temperature (Snyder & Hansen 1947, Fisher et al. 1982, Leslie & Summerell 2006) under a 12/12 h cool fluorescent light/dark cycle. Photographs and measurements were done using a Nikon Eclipse 80i microscope with Differential Interference Contrast (DIC) optics and a Nikon AZ100 stereomicroscope, both equipped with a Nikon DS-Ri2 high definition colour digital camera, and a Nikon SMZ1000 stereomicroscope equipped with a Nikon DS-Fi1 colour digital camera. Digital images were processed using the Nikon software NIS-elements D software v. 4.50. Measurements were taken for each structure from at least 30 randomly selected elements and the mean values, SD and maximum-minimum values were calculated. Line drawings were made from microphotographs using Adobe Illustrator CS5.1 v. 15.1.0.

DNA extraction, PCR amplification and sequencing

Isolates were grown for 7-10 d on malt extract agar (MEA) plates, incubated under continuous fluorescent light at room temperature. Total genomic DNA was isolated from fresh mycelium scraped from the agar surface using the Wizard® Genomic DNA purification Kit (Promega Corporation, Madison, WI, USA) according to the manufacturer's instructions. Four gene fragments, including the internal transcribed spacer region of the rDNA (ITS), a partial fragment of the large subunit of the rDNA (LSU) (spanning the variable domains D1 to D3), two fragments of the RNA polymerase's second largest subunit (RPB2) and a portion of the translation elongation factor 1-alpha ($EF-1\alpha$) were PCR amplified and sequenced according to previously published protocols (Sandoval-Denis et al. 2018) using the following primer pairs: ITS4/ITS5 for ITS (White et al. 1990), LR0R/LR5 for LSU (Vilgalys & Hester 1990, Vilgalys & Sun 1994), 5f2/7cr and 7cf/11ar for RPB2 (Liu et al. 1999, Sung et al. 2007) and EF-1/EF-2 for *EF-1* α (O'Donnell et al. 1998). Consensus sequences were assembled from forward and reverse sequences using Seqman Pro v. 10.0.1 (DNASTAR, Madison, WI, USA). All sequences newly generated in this study were uploaded to GenBank and the European Nucleotide Archive (Table 1).

Phylogenetic analyses

Alignments of sequences of the four individual loci were made using MAFFT v. 7 (Katoh & Standley 2013) under the European Bioinformatics Institute (EMBL-EBI https://www.ebi.ac.uk) framework (Li et al. 2015), visually checked and manually corrected if needed using MEGA v. 7 (Kumar et al. 2016). The best evolutionary model for each dataset (GTR+I+G) was calculated using MrModeltest v. 2.3 (Nylander 2004). Phylogenetic inferences were made using three independent algorithms, Maximum Likelihood (ML), Maximum Parsimony (MP) and Bayesian analysis (BA), for each locus. The individual gene trees were assessed for incongruence by checking their individual

Table 1 Origin, culture and DNA sequence accession numbers of the isolates included in this study.

| Species name | Lineage name ^a | Strain code ^b | Host/Sample | Country | GenE | 3ank/EMBL ac | GenBank/EMBL accession number ^c | er c |
|-------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | | | | ' | EF-1α | ITS | rsn | RPB2 |
| 'Fusarium' brasiliense 'Fusarium' euwallaceae | FSSC 36 | NRRL 31757 CBS 135854 ^T = NRRL 54722 NRPI 52626 | Glycine max Euwallacea sp. Erwallacea sp. | Brazil Israel | EF408409 JQ038007 KC691532 | EF408514 JQ038014 KC691560 | FJ919513 JQ038014 KC691560 | EU329565 JQ038028 K1171702 |
| 'Fusarium' solani f. sp. batatas 'Fusarium' solani f. sp. pisi | FSSC 23 FSSC 11 | NRRL 22400 NRRL 22278 | Ipomoea batatas Pisum sativum | USA USA | AF178343 AF178337 | AF178407 DQ094309 | DQ236345 DQ236351 | EU329509 EU329501 |
| 'Fusarium' solani f. sp. xanthoxyli 'Fusarium' striatum | FSSC 11 FSSC 22 FSSC 21 | NRRL 22820 NRRL 22277 NRRL 22101 | Glycine max Xanthoxylum sp. Cotton cloth | USA Japan Panama | AF178355 AF178336 AF178333 | DQ094310 AF178401 AF178398 | DQ236352 AF178370 AF178367 | EU329532 FJ240380 EU329490 |
| Geejayeesia atrofusca Geejayeesia cicatricum Neocosmospora catenata | FSSC 43 | NRRL 22099 NRRL 220316 CBS 125552 NRRL 54992 | Unknown Staphylea trifolia Buxus sempervirens Zebra shark multiple tissues | Unknown USA USA | JF 740782 AF 178361 HM626644 KC808213 | JF / 40905 AF178423 HQ728145 KC808255 | AF178392 MG189913 | JF / 4 1108 JX 17 1609 HQ7 28153 KC8 08354 |
| Neocosmospora croci | 7000 64 | NKKL 54933 NKKL 54933 CBS 142423 CPC 27187 | Lebra snark multiple tissues Potato Citrus sinensis Citrus sinensis | USA Germany Italy Italy | KC808214 JX435156 LT746216 LT746217 | KC808256 JX435206 LT746264 LT746265 | MG189914 JX435206 LT746264 LT746265 | KC808355 JX435256 LT746329 LT746330 |
| Neocosmospora cyanescens Neocosmospora falciformis | FSSC 27 FSSC 3+4 FSSC 3+4 FSSC 3+4 | CBS 518.82" = NRRL 37625 CBS 318.73 = NRRL 22660 CBS 475.67 ^T NPPI 54319 | Human foot Trichosanthes dioica Human soine | The Netherlands India Puerto Rico | FJ240353 JX435158 LT906669 HO401721 | EU329684 JX435208 MG18993 | EU329684 JX435208 MG189915 | EU329637 JX435258 LT960558 HO404723 |
| Neocosmospora gamsii | FSSSC 7 7 7 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 | CBS 217.53 = NRRL 22655 CBS 700.86 = NRRL 22236 CBS 130181 = NRRL 43502 CBS 143207 = NRRL 32323 CBS 143209 = NRRL 32770 | Plywood Unknown Human eye Human bronchoalveolar lavage fluid | Nigeria Brazil USA USA | DQ247637 DQ247624 DQ290488 DQ246951 DQ247083 | MG189936 DQ094763 DQ790532 DQ094420 DQ094544 | MG189916 MG189917 DQ790532 DQ236462 DQ236586 | LT960559 LT960560 DQ790576 EU329576 EU329615 |
| Neocosmospora haematococa Neocosmospora illudens Neocosmospora keratoplastica | T | CBS 149606T CBS 1196006T NRR 22090 CBS 490 63T NRR 43373 NDD1 43488 | Contain tuda namanen Dying tree Beilschmiedia tawa Human Contact lens | Sri Lanka Sri Lanka New Zealand Japan Malaysia Singapara | DQ247510 DQ247510 AF178326 LT906670 EF452920 | EF453072 | KM231664 AF178362 * EF453072 | LT960561 JX171601 LT960562 EF469959 |
| Neocosmospora lichenicola Neocosmospora macrospora | FSSC 16 FSSC 16 | NRRL 24230 NRRL 34123 NRRL 34123 CBS 142424 ^T CPC 28192 CPC 28192 | Human Human eye Citrus sinensis Citrus sinensis | Singapore Thailand India Italy Italy | DQ246877 DQ247192 LT746218 LT746219 | LO323030 DQ094355 DQ094645 LT746266 LT746267 | LU3236397 DQ236387 LT746281 LT746282 | EF470146 EF470146 EU329635 LT746331 LT746332 |
| Neocosmospora mahasenii Neocosmospora metavorans | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | CBS 119593 CBS 119594 CBS 11950400 = NRRL 42489 CBS 143194 = NRRL 22782 CBS 143195 = NRRL 22792 CBS 143199 = NRRL 28016 CBS 143199 = NRRL 28017 CBS 143201 = NRRL 28017 CBS 143202 = NRRL 28017 CBS 143202 = NRRL 28019 CBS 143219 = NRRL 32849 = FRC S-123 CBS 143216 = NRRL 3740 = UTHSC R-3564 CBS 143216 = NRRL 46737 CBS 143219 = NRRL 46708 = FMR 8634 F111 CBS 143219 = NRRL 46708 = FMR 8634 F111 NRRL 2853 NRRL 44904 NRRL 44904 | Citrus sinensis Dead branch of live tree Human corneal ulcer Human Human Human Human Human Human sye Human obne Human obne Human obst Human chest Human chest Human foot Unknown Unknown Unknown Unknown | Italy USA | LI/46220 DQ246854 DQ246854 DQ246873 DQ246873 DQ246875 DQ246876 DQ246876 DQ246876 DQ246876 DQ247094 DQ247155 FJ240356 FJ240356 FJ240356 GU170611 JF740852 | L1746268 JF433045 DQ790528 EU329671 EU329673 * EU329675 * EU329682 EU329685 EU329685 EU329686 EU329686 EU329686 EU329686 EU3296717 | L1746283 JF733045 DQ790528 EU329670 EU329673 FJ240359 FJ240360 FJ240371 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 EU329685 | LIT/46533 DQ790572 EU329528 EU329531 EF470141 EF470142 EF470143 EU329618 EU329618 EU329628 EU329628 EU329628 EU329628 EU32963 EU329643 EU32953 EU329666 |

| Table 1 (cont.) | | | | | | | | | 2 |
|-----------------------------|---------------------------|---------------------------------------------------------------------------------------|------------------------------------------------|----------------------|---------------------------------------|-----------------------|--------------------------------------------|-----------------------|------|
| Species name | Lineage name ^a | Strain code ^b | Host/Sample | Country | Genl | 3ank/EMBL ac | GenBank/EMBL accession number [°] | erc | |
| | | | | | ΕΕ-1α | ITS | rsn | RPB2 | |
| Neocosmospora petroliphila | FSSC 1 | NRRL 32315 = UTHSC 00-332 | Human groin ulcer | USA | DQ246943 | DQ094412 | DQ236454 | * | |
| | FSSC 1 | NRRL 46706 = FMR 8340 | Human blood | Qatar New Zeelend | AM412594 | EU329715 | EU329715 | EU329664 | |
| Neocosmospora pragramin | ECC 33 | NRRL 22032 CBS 341 03 - NBBI 53635 | norieria giabrata Human | New Zealand | AF 176354 | AF1/641/ | AF 175350 | JX171614 | |
| reocosmospora pseudensnorms | FSSC 33 | CBS 125729 ^T | Unknown dead tree | Sri Lanka | DQ247512 | XX433136 KC691584 | KC691584 | XX453246 KC691645 | |
| | FSSC 33 | NRRL 22354 | Bark | French Guiana | AF178338 | AF178402 | DQ236358 | EU329504 | |
| | FSSC 33 | NRRL 46517 = FRC S-1834 | Unknown | Unknown | KC691555 | KC691584 | KC691584 | KC691645 | |
| Neocosmospora solani | FSSC 5 | CBS 140079 ^{ET} = NRRL 66304 = FRC S-2364 | Solanum tuberosum | Slovenia | KT313611 | KT313633 | KT313633 | KT313623 | |
| | FSSC 5 | NRRL 32484 = FRC S-1242 | Human | NSA | DQ246982 | DQ094449 | DQ236491 | EU329583 | |
| | FSSC 5 | NRRL 43474 | Human eye | USA | EF452945 | EF453097 | EF453097 | EF469984 | |
| Neocosmospora sp. | FSSC 10 | NRRL 22098 | Cucurbit | USA | AF178327 | DQ094301 | DQ236343 | EU329489 | |
| | FSSC 10 | NRRE 22133 = A100 10039 CBS 143196 = NRRI 25392 = ATCC 32752 | Cucurbit American lobster | ranama USA | AF 176346 | DQ094502 F11329672 | F11329672 | EU329492 EU329537 | |
| | FSSC 12 | CBS 143203 = NRRL 32309 = UTHSC 00-1608 | Sea turtle | USA | DO246937 | DO094407 | DO236449 | EU329571 | |
| | FSSC 12 | CBS 143206 = NRRL 32317 = UTHSC 99-1886 | Treefish | USA | DQ246945 | DQ094414 | DQ236456 | EU329575 | |
| | FSSC 12 | CBS 143212 = NRRL 32821 = FRC S-1230 | Turtle eggs | NSA | DQ247128 | DQ094587 | DQ236629 | EU329625 | |
| | FSSC 12 | CBS 143220 = NRRL 54720 = UTHSC 10-3125 | Lined sea horse aquarium water | NSA | JQ743207 | JQ743209 | JQ743209 | JQ743211 | |
| | FSSC 12 | CBS 143221 = NRRL 54968 | Bonnet head shark | USA | LT906671 | MG189937 | MG189918 | KC808332 | |
| | FSSC 12 | CBS 143222 = NRRL 54970 = UTHSC 05-175 | Antler crab | USA 108 | KC808195 | MG189938 | MG189919 | KC808334 | |
| | FSSC 12 | CBS 143223 = NRRL 549/1 = UTHSC 05-2/74 CBS 143225 = NBBL 54074 = LITHSC 06-1538 | Reptile bronchus Honevcomb fish | NSA NSA | KC808196 KC808198 | KC808237 | MG189920 | KC808335 | |
| | FSSC 12 | CBS 143225 = INNNE 34974 = 011130 00=1330 CBS 143226 = NRRI 54979 = 117HSC 06-3660 | Kemps Ridley tirtle | A 67 | KC808202 | KC808239 | MG189921 | KC808337 | |
| | FSSC 12 | CBS 143227 = NRRI 54982 = UTHSC 07-3669 | Kemps Ridley turtle | USA | KC808205 | MG189939 | MG189923 | KC808345 | |
| | FSSC 12 | CBS 143230 = NRRL 62549 = UTHSC 08-1422 | Horseshoe crab | USA | KC808220 | KC808264 | MG189924 | KC808352 | |
| | FSSC 12 | NRRL 22642 = ATCC 38341 | Penaceous japonicus | Japan | DQ246844 | DQ094329 | DQ236371 | EU329522 | |
| | FSSC 12 | NRRL 22834 | Lobster | Australia | DQ247663 | * | * | FJ240382 | |
| | FSSC 12 | NRRL 46704 = FMR 7140 | Aquarium sand | Spain | * + | EU329713 | EU329713 | EU329662 | |
| | FSSC 12 | NRRL 46705 = FMR 7414 | Aquarium sand | Spain | * * * * * * * * * * * * * * * * * * * | EU329714 | EU329714 | EU329663 | |
| | FSSC 13 | NRRL 22161 NBBL 22163 = ATCC 48683 | Robinia pseudoacacia | Japan | AF1/8330 | DQ094311 | DQ236353 | EU329494 | |
| | FSSC 13 | NRRI 2258 = ALCC 18633 | Robinia pseudoacacia Robinia pseudoacacia | Japan | AF178353 | AF178416 | AF178385 | EU329516 | |
| | FSSC 14 | CBS 130177 = NRRL 22611 | Human cornea | USA | DQ246841 | DQ094326 | DQ236368 | EU329518 | |
| | FSSC 14 | NRRL 32705 = FRC S-0390 | Human skin | USA | DQ247025 | DQ094488 | DQ236530 | EU329594 | |
| | FSSC 15 | NRRL 28009 | Human | NSA | DQ246869 | DQ094351 | DQ236393 | EF470136 | |
| | FSSC 15 | NRRL 32792 = FRC S-1143 | Human cutaneous nodules | Japan | DQ247101 | DQ094561 | DQ236603 | EU329621 | |
| | FSSC 17 | NRRL 22157 = NRRL 22479 = ATCC 18689 | Morus alba | Japan | AF178359 | DQ094306 | DQ236348 | EU329493 | |
| | FSSC 17 | NRRL 22230 = ATCC 44934 | Morus alba | Japan | AF178358 | DQ094305 | DQ236347 | EU329499 | |
| | FSSC 18 | NRRL 31158 | Human | NSA | DQ246916 | DQ094389 | DQ236431 | EU329559 | |
| | FSSC 19 | CBS 571.94 = NRRL 36510 | Camelia sinensis | India | KC691530 | KC691558 | KC691558 | KC691619 | |
| | FSSC 19 | NRRL 20438 = IMI 296597 | Camelia sinensis | India | AF178332 | AF178397 | DQ236357 | JX171584 | |
| | FSSC 19 | NKKL 22346 OBS 447484 NIPPL 22380 | Camelia sinensis | India | FJZ40350 | EU329669 | EU329669 | EU329503 | |
| | FSSC 25 | CB3 11/46 = INNR 22363 CB3 102824 = NRRI 53598 | Lindellalon tanphera Leaf litter | Colombia | JX435147 | IX435197 | 1X435197 | LX435247 | |
| | FSSC 25 | CBS 130328 = NRRL 31169 | Human oral wound | USA | DQ246923 | DQ094396 | DQ236438 | KR673999 | |
| | FSSC 26 | NRRL 28541 = UTHSC 98-1305 | Human synovial fluid | USA | DQ246882 | EU329674 | EU329674 | EU329542 | |
| | FSSC 28 | CBS 109028 = NRRL 32437 | Human subcutaneous nodule | Switzerland | DQ246979 | DQ094446 | DQ236488 | EU329581 | |
| | FSSC 28 | NRRL 52705 | Unknown | Unknown | JF740787 | * | * | JF741113 | Р |
| | FSSC 29 | KOD253 | Unknown | Unknown | * 1 | (((* (| * [| | ers |
| | FSSC 29 | NRRL 28008 | Human | USA | DQ246868 | DQ094350 | DQ236392 | EF470135 | 001 |
| | FSSC 34 | NRRL 460390 NRRL 46703 = FMR 8281 | Olikilowii Nematode | Spain | GU170627 HM347126 | GU170647 FU329712 | GU170647 FU329712 | GU 170392 FU329661 | nia |
| | FSSC 35 | NRRL 46707 = FMR 8030 | Human | Brazil | HM347127 | EU329716 | EU329716 | EU329665 | – V |
| | FSSC 37 | NRRL 25137 | Diseased cocoa pods | New Guinea | JF740757 | JF740899 | JF740899 | JF741084 | olur |
| | FSSC 37 | NKKL 25138 NRRI 52782 | Ulseased cocoa pods Hypothenemis hambei adult | New Guinea Benin | UQ24/53/ * | JF740900 | JF740900 | JF 741085 IF741176 | ne - |
| | FSSC 38 | NRRL 52783 | Hypothenemus hampei adult | Uganda | JF740851 | * | * | JF741177 | 41, |
| | FSSC 40 | F1285 | Unknown | Unknown | * | * | * | * | 20 |
| | FSSC 45 | KOD614 | Unknown | Unknown | * | * | * | * | 18 |

| cont. | |
|-------|--|
| _ | |
| Table | |

| Species name | Lineage name ^a | Strain code ^b | Host/Sample | Country | Gen | GenBank/EMBL accession number ^c | ccession numb | er° |
|---------------------------|---------------------------|---------------------------------------------------|---------------------|-----------------|----------|--------------------------------------------|---------------|----------|
| | | | | | EF-1α | ITS | rsn | RPB2 |
| Neocosmospora suttoniana | FSSC 20 | CBS 124892 | Human nail | Gabon | JX435139 | JX435189 | JX435189 | JX435239 |
| | FSSC 20 | CBS 130178 = NRRL 22608 = UTHSC 93-1547 | Human | USA | DQ246838 | DQ236365 | DQ094323 | EU329517 |
| | FSSC 20 | CBS 143197 = NRRL 28000 | Human blood | USA | DQ246865 | DQ094347 | DQ236389 | EF470128 |
| | FSSC 20 | CBS 143204 = NRRL 32316 = UTHSC 00-264 | Human corneal ulcer | USA | DQ246944 | DQ094413 | DQ236455 | EU329574 |
| | FSSC 20 | CBS 143214 ^T = NRRL 32858 | Human wound | USA | DQ247163 | DQ094617 | DQ236659 | EU329630 |
| | FSSC 20 | CBS 143224 = NRRL 54972 = UTHSC 05-2900 | Equine eye | Unknown | KC808197 | MG189940 | MG189925 | KC808336 |
| | FSSC 20 | NRRL 28001 | Human skin | USA | DQ246866 | DQ094348 | DQ236390 | EF470129 |
| Neocosmospora tonkinensis | FSSC 9 | CBS 115.40 ^T = NRRL 53586 = IMI 113868 | Musa sapientum | Vietnam | LT906672 | MG189941 | MG189926 | LT960564 |
| | FSSC 9 | CBS 143038 | Human cornea | The Netherlands | LT906673 | MG189942 | MG189927 | LT960565 |
| | FSSC 9 | CBS 143208 = NRRL 32755 = FRCS-0452 | Turtle head lesion | USA | DQ247073 | DQ094534 | DQ236576 | EU329613 |
| | FSSC 9 | CBS 143217 = NRRL 43811 | Human cornea | USA | EF453053 | EF453204 | EF453204 | EF470092 |
| | FSSC 9 | FRC S-2484 | Unknown | Unknown | * | * | * | * |
| | FSSC 9 | FRC S-2540 | Unknown | Unknown | * | * | * | * |
| | FSSC 9 | NRRL 46615 | Unknown | Unknown | GU250543 | GU250666 | GU250666 | GU250728 |
| | FSSC 9 | NRRL 46676 | Unknown | Unknown | GU250546 | GU250669 | GU250669 | GU250731 |
| Neocosmospora vasinfecta | FSSC 8 | CBS 130182 = NRRL 43467 | Human | USA | EF452940 | EF453092 | EF453092 | EF469979 |
| | FSSC 8 | NRRL 34174 = UTHSC 03-1457 | Human | USA | * | * | * | EU329636 |

Following the clade nomenclature by O'Donnell et al. (2008)

large ribosomal subunit gene; RPB2 = partial fragment of the DNA-directed housed in The Pennsylvania State Uni-Peoria, IL, USA; All others = as named in K. O'Donnell's sequence database. Ex-type and Center FRC = Fusarium Research LSU = partial fragment of the large ribo and are not currently publicly available Spain; Reus, Universitat Rovira i Virgili, Surrey, UK; KOD = personal collection of Kerry O'Donnell; NRRL = collections of the Agricultural Research Service, r region of the rDNA; L 1 by Kerry O'Donnell a de Medicina, spacer FMR = Facultat Internal transcribed 1-alpha; ITS = Internal tran Sequences marked with * held at CBS; ITS= Crous, factor a **bold**. collection of Pedro W. EMBL= The European Molecular Biology Laboratory; EF-1a= partial fragment of the translation elongation for RNA polymerase II, second largest subunit. Accession numbers of sequences generated in this study are in Personal CPC = $1\alpha = \text{partial fragment of}$ versity Institute, Utrecht, The Notherlands; (
IMI = CABI Genetic Resource Collection, §
with ^T and ^{ET}, respectively.
at Biology Laboratory; EF-1a = partial fragm Molecular CBS = Westerdijk Fungal Biodiv versity, State College, PA, USA; ex-epitype strains are indicated ex-epitype strains are ir EMBL = The European

were provided

phylogenies for conflicts between clades with significant ML, MP and BA support, after which the four gene datasets were concatenated (Mason-Gamer & Kellogg 1996, Wiens 1998).

Maximum Likelihood and BA were run on the CIPRES Science Gateway portal (https://www.phylo.org/) (Miller et al. 2012) using RaxML v. 8.2.10 (Stamatakis 2014) and MrBayes v. 3.2.6 (Huelsenbeck & Ronquist 2001, Ronquist & Huelsenbeck 2003), respectively. For ML analyses the default parameters were used and BS was carried out using the rapid bootstrapping algorithm with the automatic halt option. Bayesian analyses included four parallel runs of 5000000 generations, with the stop rule option and a sampling frequency of 1000 generations. The burn-in fraction was set to 0.25, after which the 50 % majority rule consensus trees and posterior probability (PP) values were calculated. The resulting trees were plotted using FigTree v. 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree).

Maximum Parsimony analyses were carried out using PAUP v. 4.0b10 (Swofford 2002). Heuristic searches consisted of 1000 random stepwise addition replicates, with tree bisection and reconstruction (TBR) branch swapping. All characters were equally weighted and gaps were treated as missing data. Zero length branches were collapsed and all multiple, equally parsimonious trees were saved. Tree length, consistency index, retention index and rescaled consistency index (TL, CI, RI and RC, respectively) were calculated. Statistical support for the branches was evaluated using a bootstrap analysis (BS) of 1000 replicates.

RESULTS

Phylogenetic assessment of pathogenic clades in Neocosmospora

To show the current known diversity in Neocosmospora as well as the phylogenetic position and genealogical exclusivity of the most important lineages containing human and veterinary pathogens, an overview phylogeny was constructed based on the original alignments published by O'Donnell et al. (2008).

Individual gene phylogenies proved to be topologically consistent with each other, but showed different degrees of resolution for the most relevant pathogenic clades (data not shown, all trees available in TreeBASE). As evaluated on the basis of clade stability and MLBS values, RPB2 was the only locus unambiguously identifying all the clinically significant clades, including 'Fusarium' metavorans, FSSC 7, 9, 12, 20 and 43, as well as the important human and veterinary pathogens N. falciformis, 'F.' keratoplasticum, 'F.' petroliphilum and N. solani. Bootstrap values were between 93 and 100 %, except in the case of FSSC 9, where the BS was 76 %. The partitioned analysis of *EF-1α* resulted in moderate to highly supported monophyletic clades (BS = 75-100 %) for most of the pathogenic species with exception of FSSC 43. This analysis exposed considerable divergence among *EF-1α* sequences for strains within FSSC 7 and 'F.' keratoplasticum; the divergent strains formed sister lineages to the respective main clades. These subclades had low statistical support. The ITS phylogeny was able to clearly distinguish five of the most important lineages, N. falciformis, N. solani, 'F.' petroliphilum, FSSC 12 and FSSC 20, with BS = 76-99 %, while LSU allowed for the identification of only two pathogenic clades, FSSC 12 and FSSC 20, with BS = 71 and 92 %, respectively.

The final analysis included 3287 characters from four loci $(EF-1\alpha = 675, ITS = 491, LSU = 485, RPB2 = 1636)$ of 132 strains including the outgroup taxa 'F.' cicatricum = Geejayessia cicatricum and 'F.' staphyleae = G. atrofusca (Schroers et al. 2011, 2016). Of the characters used, 2 297 were variable $(EF-1\alpha = 395, ITS = 343, LSU = 441, RPB2 = 1118)$ and 742

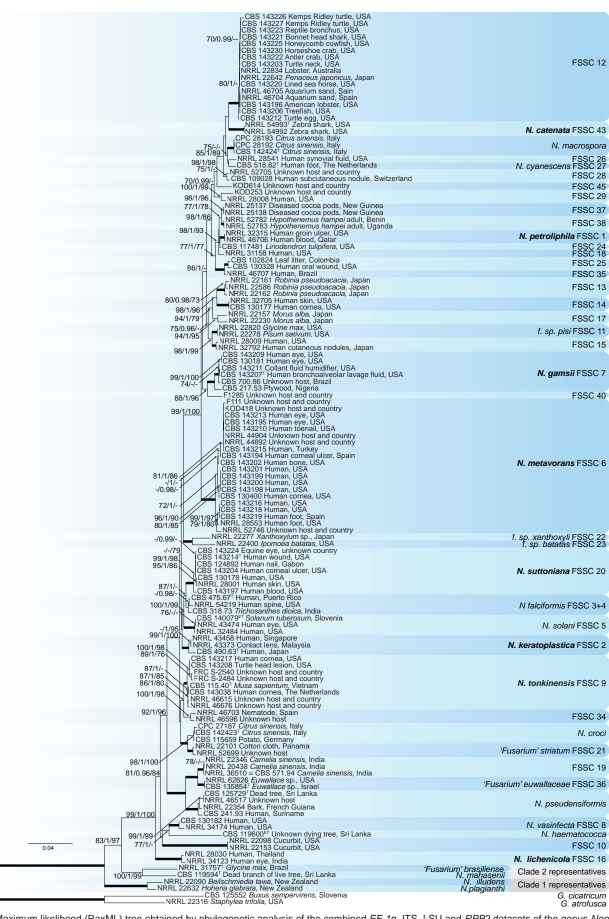


Fig. 1 Maximum likelihood (RaxML) tree obtained by phylogenetic analysis of the combined *EF-1α*, ITS, LSU and *RPB2* datasets of the genus *Neocosmospora*. Bootstrap support values from Maximum Likelihood (ML-BS), Maximum Parsimony (MP-BS) and Bayesian posterior probabilities (PP) above 70 % and 0.95, respectively, are indicated at the nodes. Nodes with full statistical support (ML-BS = 100, MP-BS = 100 and BS = 1) are indicated by **bold** branches. Names of new species and new combinations are in **bold**. *Geejayessia atrofusca* (CBS 125552) and *G. cicatricum* (NRRL 22316) were used as outgroup. CBS = Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands; CPC = Personal collection of Pedro W. Crous, held at CBS; FRC = Fusarium Research Center housed in The Pennsylvania State University, State College, PA, USA; KOD = personal collection of Kerry O'Donnell; NRRL = collections of the Agricultural Research Service, Peoria, IL, USA; All others = as named in O'Donnell's sequence database; ^{ET} = ex-(epi-)type strain; ^T = ex-(holo-)type strain.

were parsimony-informative (*EF-1* α = 207, ITS = 97, LSU = 33, RPB2 = 405). The ML search revealed a best tree with a lnL of -19162.599 (Fig. 1). The MP analysis produced 1000 equally parsimonious trees (TL = 2655 steps, CI = 0.489, RI = 0.830, RC = 0.406) highly congruent with that produced in ML. The BA lasted for 970 000 generations and the 50 % consensus tree and posterior probabilities were calculated from 728 trees (Fig. 1). The genus Neocosmospora received maximal statistical support (ML and MP BS = 100 % / 100 % and PP = 1). All human and veterinary pathogenic clades clustered within clade 3 of Neocosmospora sensu O'Donnell et al. (2008). All lineages containing clinically relevant unnamed phylogenetic species and currently known species resolved as monophyletic clades with strong statistical support (ML and MP BS = 100 % / 100 %and PP = 1) with exception of N. falciformis. This species lacked BS support in both ML and MP analyses, but had moderate BA support (PP = 0.98). The strain CBS 217.53, which showed a divergent $EF-1\alpha$ sequence, is provisionally retained here in clade FSSC 7 based on its morphological features. Clades FSSC 7, 20 and 43 are here described as the new species N. gamsii, N. suttoniana and N. catenata, respectively. The ex-type strain of Cylindrocarpon tonkinense (CBS 115.40) was found to cluster within FSSC 9, for which the new combination Neocosmospora tonkinensis is proposed. The recently described species 'F.' metavorans (Al-Hatmi et al. 2018), is here recombined in Neocosmospora and an emended description is provided.

Taxonomy and morphology

Based on the phylogenetic evidence and morphological observations compiled here, formal descriptions for the most clinically important unnamed clades in *Neocosmospora* are provided. In keeping with the current circumscription of the genus (Lombard et al. 2015), new combinations are needed for other clinically relevant species in *Neocosmospora*.

A summary of the main morphological features (Table 2), and a schematic overview comparison (Fig. 2) were produced to facilitate the distinction of the most frequently isolated pathogens within the genus.

Neocosmospora catenata Sandoval-Denis & Crous, sp. nov.— MycoBank MB822898; Fig. 3

Etymology. From Latin catena, meaning 'chain, succession'. Referring to the abundant chains of chlamydospores.

Type. USA, Georgia, *Stegostoma fasciatum* multiple tissues (CBS H-23225 – holotype; CBS 143229 = NRRL 54993 = UTHSC 09-1009 – culture ex-type).

Sporulation abundant from conidiophores formed directly on the substrate mycelium. Conidiophores up to 480 µm tall, erect, emerging from the agar surface as single phialides, unbranched or more commonly 1–3-times branched laterally bearing terminal monophialides; phialides subulate, subcylindrical to somewhat acicular, smooth- and thin-walled, (10.5–)32.5–55(–61.5)

Table 2 Main asexual morphological features of the most clinically relevant Neocosmospora species.

| Species name ^a | Aerial conidia | Sporodochial conidia (number of septa) | Chlamydospore diam |
|---------------------------|---------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------|
| N. catenata | (0(-1)-septate) (4.5-)6-9(-11) × (2.5-)3.5-4.5(-6) μm | N.A. | 5.5–9.5 μm, smooth-walled |
| N. falciformis#,† | (0-1-septate) 4.7-41.8 × 3.1-9.4 μm | (3-4-septate) Overall: 41.7–46.9 × 5.9–6.1 µm | 8–15 μm, rough-walled |
| N. gamsii | (0(-1)-septate) (5-)6.5-9.5(-11) × 2.5-3.5(-4.5) µm | $\begin{array}{l} ((3-)4-5(-7)\text{-septate}) \\ (3): 35.5-42.5\times5.5-6\ \mu\text{m} \\ (4): (36-)38.5-59(-63)\times5-5.5(-6)\ \mu\text{m} \\ (5): (50.5-)55-66(-71.5)\times(4.5-)5-6.5(-7)\ \mu\text{m} \\ (6): 67-77.5\times5.5-6.5\ \mu\text{m} \\ (7): 67.5-71\times6-7\ \mu\text{m} \\ \text{Overall: } (35.5-)51-68(-77.5)\times(4.5-)5-6(-7)\ \mu\text{m} \end{array}$ | 5.5–8(–9) µm, smooth-walled |
| N. keratoplastica# | (0-3-septate) 3.1-35.8 × 2.9-6.6 μm | ((1–)3–5-septate) Overall: $36.8-43.4 \times 5.3-5.7 \ \mu m$ | 6.0-8.0 µm, smooth- to rough-walled |
| N. metavorans | (0–1(–3)-septate) (4–)11–25.5(–35) × (2–)4–6(–7) μm | $\begin{array}{l} ((1-2-)3-5\text{-septate}) \\ (1): 22.5-25\times5-5.5\ \mu\text{m} \\ (2): 25.5-27.5\times6-7\ \mu\text{m} \\ (3): (30.5-)38-46(-47.5)\times(5-)5.5-6.5(-7.5)\ \mu\text{m} \\ (4): (43-)45-48.5\times(5.5-)6-7(-7.5)\ \mu\text{m} \\ (5): (46-)47-51.5(-53)\times(5.5-)6-7.5\ \mu\text{m} \\ \text{Overall: } (22.5-)38.5-50(-53)\times(5-)6-7(-7.5)\ \mu\text{m} \end{array}$ | 5–13.5 μm, smooth-walled |
| N. petroliphila# | (0(–1)-septate) 4.6–24.9 × 2.6–7.1 μm | (3–5-septate) Overall: 44–52.2 × 5.1–5.9 μm | smooth-walled |
| N. solani‡ | (0-3(-4-5)-septate) $(5.5-)13.5-43(-53)\times(2-)3-7(-8)\ \mu\text{m}$ | $\begin{array}{l} ((0-)3-4(-5)\text{-septate}) \\ (3): (24-)36-44(-48) \times (2-)4.5-6(-8) \ \mu\text{m} \\ (4): (31-)42-48(-52) \times (3-)4.5-6(-7.5) \ \mu\text{m} \\ (5): (41-)45-51(-56) \times (2.5-)4.5-6(-8) \ \mu\text{m} \\ \text{Overall: } (24-)34-52.5(-56) \times (2-)3-7.5(-8) \ \mu\text{m} \end{array}$ | 6.5–8.5 μm, rough-walled |
| N. suttoniana | (0-2(-3)-septate) $(6-)7.5-21(-31) \times (2.5-)3-5.5(-7.5) \mu m$ | $\begin{array}{l} ((3-)5-6\text{-septate}) \\ (3): 30.5-32.5\times7-7.5\ \mu\text{m} \\ (4): (49-)50-53.5\times6-6.5\ \mu\text{m} \\ (5): (30.5-)52-71(-77.5)\times(6-)7-8\ \mu\text{m} \\ (6): (75-)77-84.5(-86.5)\times(6.5-)7-8\ \mu\text{m} \\ \text{Overall: } (30.5-)50-75(-86.5)\times(6-)7-7.5(-8)\ \mu\text{m} \end{array}$ | (4.8–)6–8.5(–9.5) μm, verruculose |
| N. tonkinensis | (0-3(-4)-septate) $(6-)11-24(-37) \times (3.5-)4-6(-7) \ \mu\text{m}$ | $\begin{array}{l} ((1-)3-4(-5)\text{-septate}) \\ (1): 47-51 \times 6-7.5 \ \mu\text{m} \\ (3): (28-)32.5-42.5(-45.5) \times (5.5-)6-7.5 \ \mu\text{m} \\ (4): (40.5-)43-48(-49) \times 6-7.5 \ \mu\text{m} \\ (5): (40-)41.5-52 \times 6.9-7.3 \ \mu\text{m} \\ \text{Overall: } (27.5-)37-48(-50.5) \times (5.5-)6-7(-7.5) \ \mu\text{m} \end{array}$ | 6.5–10(–12) μm, smooth-walled |

¹ Conidial measurements from: # Short et al. (2013), † Chehri et al. (2015), [‡] Schroers et al. (2016).

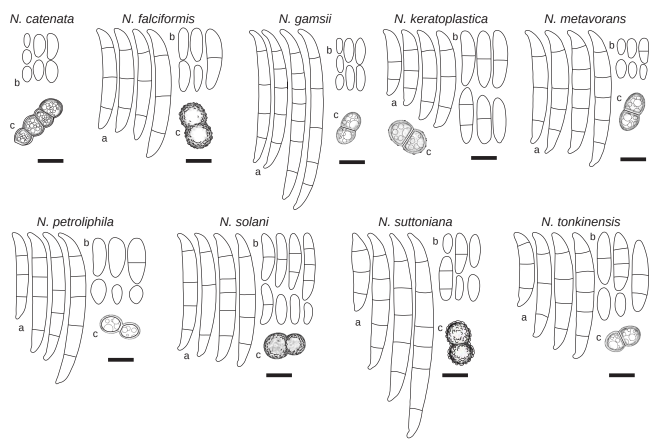


Fig. 2 Line drawings comparing the main conidial and chlamydospore features of the most clinically relevant species of *Neocosmospora*. a. Sporodochial conidia; b. aerial conidia; c. chlamydospores. — Scale bars = 10 μm.

 \times (1.5–)2.5–3.5(–4) µm, with distinct periclinal thickening and an apical flared collarette; conidia hyaline, obovate, ellipsoidal to reniform, commonly bent dorsoventrally, smooth- and thinwalled, 0(–1)-septate, (4.5–)6–9(–11) \times (2.5–)3.5–4.5(–6) µm, grouped on small false heads on the tip of monophialides. Chlamydospores abundant, subhyaline to pale brown, spherical to subspherical, 5.5–9.5 µm diam, solitary, in pairs, chains or clusters, terminal or intercalary, smooth- and thick-walled. Sporodochia and multiseptate conidia not seen.

Culture characteristics — Colonies on PDA growing in the dark with an average radial growth rate of 2.5-5 and 3.5-5.9 mm/d at 21 and 24 °C, respectively, reaching 74-82 mm diam in 7 d at 24 °C and occupying an entire 9 cm Petri dish in 7 d at 27 °C. Colony surface buff to rosy buff, flat, felty to velvety, radiate, with abundant aerial mycelium; colony margins irregular with abundant submerged mycelium. Reverse straw to buff coloured. Straw to pale sulphur yellow diffusible pigment produced between 21-27 °C, becoming ochreous to umber at 30-33 °C. Colonies on OA incubated at 24 °C in the dark reaching 80-90 mm diam in 7 d. Colony buff to honey, flat, membranous, becoming velvety with the production of short aerial mycelium; margins regular. Reverse buff to honey, without diffusible pigments. A hazel to isabelline pigment can be produced in incubation at 36 °C. On CMA incubated at 24 °C in the dark, cultures occupy an entire 9 mm Petri dish in 7 d. Colony colour sulphur yellow to straw, flat with abundant floccose aerial mycelium. Reverse sulphur yellow to straw without diffusible pigments.

Cardinal temperatures for growth — Minimum 12 °C, maximum 36 °C, optimal 24–27 °C.

Additional material examined. USA, Georgia, Stegostoma fasciatum multiple tissues (NRRL 54992 = CBS 143228 = UTHSC 09-1008).

Notes — *Neocosmospora catenata*, known from the zebra shark (*Stegostoma fasciatum*), is well-defined phylogenetically

as a fully-supported sister clade to FSSC 12, which is also known mostly from infections of marine animals. No single morphological feature exists allowing a quick phenotypic distinction of FSSC 12 from *N. catenata*, notwithstanding the tendency of the latter species to produce large, pigmented, catenate to clustered chlamydospores. The two strains studied here consistently failed to produce the characteristic falcate, multiseptate sporodochial conidia typical of the genus. Sporulation was abundant, but strictly microconidial. It is not clear if this phenomenon reflects strain degeneration or if it is a distinctive peculiarity of this clade. The two strains of *N. catenata* included in this study are, to our knowledge, the only material currently available in fungal collections. Additional isolates are needed to help in evaluating this potentially important differential morphological character.

Neocosmospora gamsii Sandoval-Denis & Crous, *sp. nov.* — MycoBank MB822899; Fig. 4, 5

Etymology. In honour and memory of Walter Gams, eminent mycologist and Fusarium researcher.

Type. USA, Pennsylvania, from human bronchoalveolar lavage fluid, D.A. Sutton (CBS H-23226 – holotype; CBS 143207 = NRRL 32323 = UTHSC 99-250 – culture ex-type).

Sporulation abundant from sporodochia and from conidiophores formed directly on the substrate mycelium. Conidiophores in the aerial mycelium up to 410 μm tall, irregularly or sympodially branched at various levels, bearing terminal monophialides; phialides subulate, subcylindrical or acicular, smooth- and thin-walled, $(37.5-)46.5-64(-78)\times(2-)2.5-4~\mu m$, with inconspicuous periclinal thickening; collarettes small and barely visible; conidia formed on aerial conidiophores hyaline, ellipsoidal to clavate, sometimes slightly and inequilaterally bent dorsoventrally, smooth- and thin-walled, 0(-1)-septate,



Fig. 3 Neocosmospora catenata. a. Colony on PDA; b. colony on OA; c. colony on CMA; d-g. conidiophores and phialides; h-i. tip of phialides showing apical collarettes; j-k. conidia; l-m. chlamydospores. — Scale bars: $h-i=5 \mu m$; all others = 10 μm .

 $(5-)6.5-9.5(-11) \times 2.5-3.5(-4.5) \mu m$, single or forming small false heads. Sporodochia at first cream coloured turning green to yellow-blue-green, formed abundantly on the surface of carnation leaves, rapidly confluent. Conidiophores in sporodochia, 23-47.5 µm tall, densely packed, irregularly or verticillately branched, terminal branches bearing 1(-2) monophialides; sporodochial phialides subulate to subcylindrical or doliiform, often slightly constricted or bent in the middle portion, $12-18.5(-24) \times (2.5-)3-3.5(-4) \mu m$, smooth- and thin-walled, often showing periclinal thickening and an evident flared collarette. Sporodochial conidia wedge-shaped, medium to robust, with an almost straight to slightly curved ventral line and a gentle, continuous dorsal curvature, tapering and becoming more pronouncedly curved towards the basal and apical levels, apical cell more or less equally sized than the adjacent cell, distinctly hooked with rounded ends and a notched to foot-like basal cell, (3-)4-5(-7)-septate, hyaline, thin- and smoothwalled. Three-septate conidia: $35.5-42.5 \times 5.5-6 \mu m$; 4-septate conidia: $(36-)38.5-59(-63) \times 5-5.5(-6) \mu m$; 5-septate conidia: $(50.5-)55-66(-71.5) \times (4.5-)5-6.5(-7) \mu m$; 6-septate

conidia: $67-77.5 \times 5.5-6.5 \mu m$; 7-septate conidia: 67.5-71 \times 6-7 µm; overall (35.5-)51-68(-77.5) \times (4.5-)5-6(-7) μm. Chlamydospores abundant, spherical to subspherical, $5.5-8(-9) \mu m$ diam, solitary or in pairs, terminal and intercalary, smooth- and thick-walled. Perithecia orange to dark brown-red, globose to pyriform, superficial, solitary or gregarious, coarsely warted, glabrous, $186-194 \times 138-156 \mu m$; warts $5-20 \mu m$ diam, 3.5-16 µm tall. Peridial wall composed of thick-walled cells of textura angularis, (7.5–)11.5–18(–20.5) µm diam. Asci clavate, unitunicate, with a broad and somewhat flattened and simple apex, $(70-)72-87.5(-97.5) \times (6.5-)7.5-9(-10) \mu m$, ascospores obliquely uniseriate or irregularly biseriate at the apex of the asci. Ascospores obovoid to subfusiform, 1-septate, $(9.5-)10.5-11.5(-12.5) \times (4.5-)5.0-6.5(-7.5) \mu m$, pale yellow-brown to golden yellow, thick-walled, longitudinally finely striated, often slightly constricted at the septum.

Culture characteristics — Colonies on PDA growing in the dark with an average radial growth rate of 2.5-4.6 and 3.3-5.7 mm/d at 21 and 24 °C, respectively, reaching 76-80 mm diam in 7 d at 24 °C. Colony surface pale luteous to rosy buff, flat,

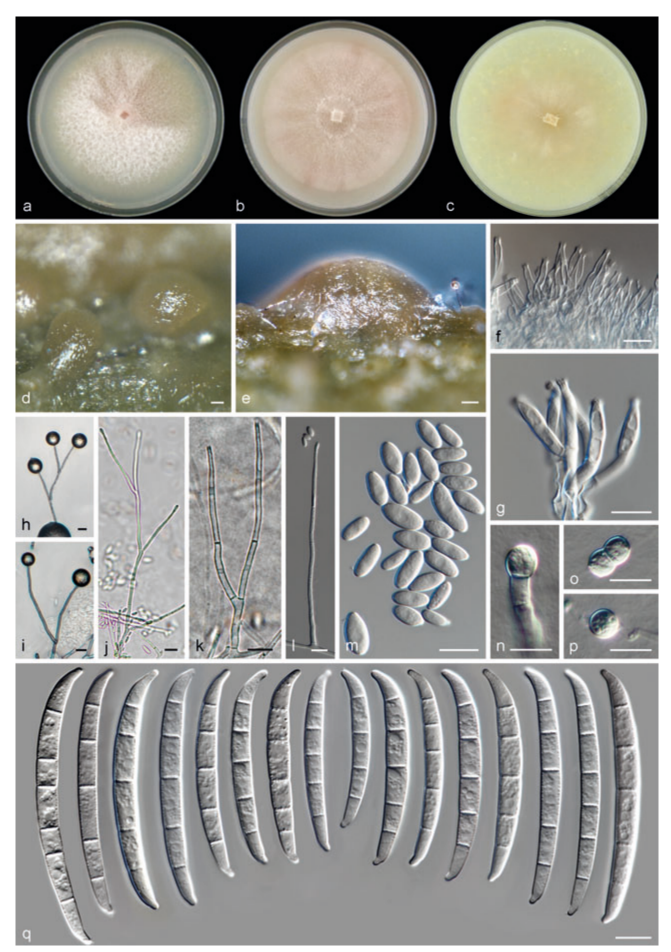


Fig. 4 Neocosmospora gamsii, asexual morph. a. Colony on PDA; b. colony on OA; c. colony on CMA; d-e. sporodochia formed on the surface of carnation leaves; f-g. sporodochial conidiophores and phialides; h-l. aerial conidiophores and phialides; m. aerial conidia; n-p. chlamydospores; q. sporodochial macroconidia. — Scale bars: $d-e = 20 \mu m$; all others = $10 \mu m$.

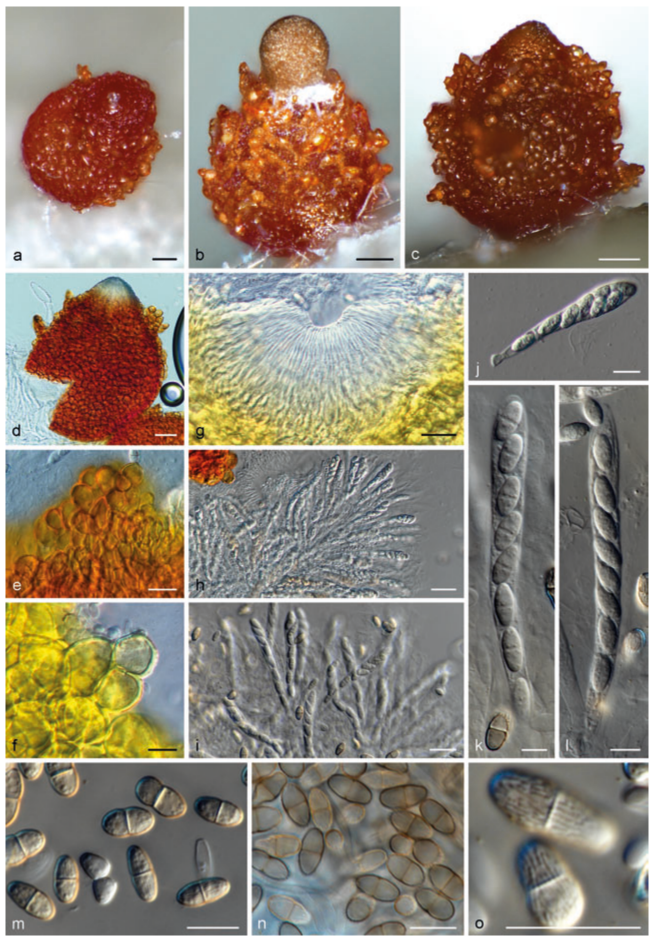


Fig. 5 Neocosmospora gamsii, sexual morph. a-c. Perithecia; d. perithecium showing a deep-red reaction on 3 % KOH; e. close view of perithecial warts (mounted on water); f. close view of perithecial warts showing a yellow reaction on lactic acid; g. ostiole and periphyses; h-l. asci and ascospores; m-n. ascospores; o. surface view of ascospores. — Scale bars: a-e, $g-i = 20 \mu m$; all others = 10 μm .

felty with velvety radial patches and abundant floccose white aerial mycelium; colony margins regular. Reverse pale luteous to orange or light scarlet toward the centre of the colony. Yellow to orange-yellow diffusible pigments can be formed at temperatures from 15 to 36 °C, becoming more intense as temperatures exceed 27 °C. Colonies on OA incubated at 24 °C in the dark reaching a maximum of 70-72 mm diam in 7 d. Colony surface pale rosy buff to pale rosy vinaceous, flat and radially folded, moist, bright and membranous, becoming felty to velvety or cottony with the production of abundant, short aerial mycelium often arranged in concentric rings, and becoming compact and restricted at 30-37 °C; margins regular. Reverse rosy vinaceous without diffusible pigments. On CMA incubated at 24 °C in the dark reaching a maximum of 35-40 mm diam in 7 d. Colony colour straw to pale buff with ochreous patches; colony surface flat with abundant submerged mycelium, and with rays of scant aerial mycelium. Reverse, straw to sulphur yellow without diffusible pigments.

Cardinal temperatures for growth — Minimum 9 °C, maximum 36 °C, optimal 24–30 °C.

Additional material examined. Brazil, substrate, date and collector unknown (CBS 700.86 = NRRL 22236). — Nigeria, from plywood, Feb. 1953, M.B. Schol-Schwarz (CBS 217.53 = NRRL 22655). — USA, Tennessee, from human eye, M. Brandt (CBS 130181 = NRRL 43502); Tennessee, from human eye (CBS 143209 = NRRL 32770 = FRC S-0524); New York, from humidifier coolant (CBS 143211 = NRRL 32794 = FRC S-1152).

Notes — This species was previously assigned to clade FSSC 7 in Neocosmospora. Morphologically N. gamsii resembles Fusarium eumartii, a known pathogen of potatoes (Solanum tuberosum) and tomatoes (Lycopersicon esculentum), for which also pathogenicity against pepper (Capsicum anuum) and eggplant (Solanum melongena) has also been demonstrated (Romberg & Davis 2006). Fusarium eumartii, however, has not been fully characterised phylogenetically and lacks authentic living strains for comparison. Two strains previously identified as F. eumartii, CBS 217.53 and CBS 700.86, were found to cluster within FSSC 7. The current concept of F. eumartii, however, based on morphology and host ranges, is polyphyletic, with isolates distributed among at least six monophyletic clades within Neocosmospora (unpubl. data). Neocosmospora gamsii can nonetheless be distinguished morphologically from the concept of F. eumartii, since it produces comparatively thin and short sporodochial conidia, which are also less frequently septate than conidia of F. eumartii and have a more pronounced apical curvature.

Among the clinically relevant species, *N. gamsii* stands out in its long, slender and highly septate (up to 7 septa) sporodochial conidia, comparable to those of *N. suttoniana*. The latter species, however, produces less frequently septate (up to 6 septa) sporodochial conidia with thick-walls and with a less pronounced overall curvature. It also produces rough-walled chlamydospores distinct from the smooth-walled chlamydospores seen in *N. gamsii*.

So far, this species is known mainly from human clinical specimens, causing mostly eye infections but also recovered from blood samples (Scheel et al. 2013). It was reported as one of many 'Fusarium' genotypes recovered from patients affected by a keratitis outbreak in the US (Chang et al. 2006).

Neocosmospora keratoplastica (Geiser et al.) Sandoval-Denis & Crous, *comb. nov.* — MycoBank MB822900

Basionym. Fusarium keratoplasticum Geiser et al., Fungal Genet. Biol. 53: 68. 2013.

Synonyms. Cephalosporium keratoplasticum T. Morik, Mycopathologia 2. 66. 1939, nom. nud. (fide Short et al. 2013).

Hyalopus keratoplasticum (T. Morik) M.A.J. Barbosa, Notarisia 19. 1941, nom. inval. (fide Short et al. 2013).

Fusarium solani (Mart.) Sacc. f. keratitis Y.N. Ming & T.F. Yu, Acta Microbiol. Sin. 12: 184. 1966.

Cylindrocarpon vaginae C. Booth, Y.M. Clayton & Usherw., Proc. Indian Acad. Sci. Pl. Sci. 94: 436. 1985.

Type. USA, Virginia, Winchester, from indoor plumbing, June 2009 (FRC S-2477 – holotype, metabolically inactive culture deposited at the Fusarium Research Center, ex-type strain: CBS 490.63 = NRRL 22661).

Description and illustrations — Short et al. (2013).

Notes — This cosmopolitan species is known almost exclusively from infected animals and from biofilms occurring in plumbing systems, including hospital water supplies (Short et al. 2013, 2014), but is also occasionally found in plant material and soil (Chehri et al. 2015, Shaffer et al. 2017). It is regarded as one of the most prevalent fusaria isolated from human disease worldwide, causing mostly corneal infections, but also isolated from blood, nails and skin (O'Donnell et al. 2008, Short et al. 2013). It is also a common species in animal infections, and has been reported from many different species, including mostly aquatic or aquatic-adapted animals such as the black spotted stingray (Taeniura melanopsila) (Fernando et al. 2015), grey seal (Halichoerus grypus) (O'Donnell et al. 2016), hammer-head sharks (Fernando et al. 2015), iguanas (O'Donnell et al. 2008, 2016), lung fish (O'Donnell et al. 2016) and shrimps including Penaeus japonicus and the California brown shrimp (O'Donnell et al. 2008, 2016). It causes extensive egg mortality in the green sea turtle, Chelonia mydas (Sarmiento-Ramírez et al. 2017), and, together with N. falciformis, represents a significant risk for the endangered hawksbill sea turtle. Eretmochelvs imbricata (Sarmiento-Ramírez et al. 2014). Terrestrial animals such as equines and Drymarchon corais, the indigo snake, may also be infected (O'Donnell et al. 2016).

Reported as having highly variable conidial morphology in culture (Short et al. 2013, 2014), *N. keratoplastica* frequently produces short, (1-2-)3-5-septate, arcuate sporodochial conidia somewhat reminiscent in shape of those seen in FSSC 12 (Short et al. 2013). However, the latter species produces 1–3-septate and much shorter and thinner sporodochial conidia (overall: $(19-)24.5-35(-41)\times5-6(-6.5)$ vs $13.2-60.1\times2.8-8.2$ in *N. keratoplastica*).

Interestingly, genetic analyses have demonstrated some significant degree of genetic transfer between *N. keratoplastica* and *N. tonkinensis*, as shown by perfect sequence matches between the nuclear rDNA regions in some isolates (Short et al. 2014).

Neocosmospora lichenicola (C. Massal) Sandoval-Denis & Crous, comb. nov. — MycoBank MB822901

Basionym. Fusarium lichenicola C. Massal., Ann. Mycol. 1: 223. 1903. Synonyms. Bactridium lichenicolum (C. Massal.) Wollenw., Fusaria autographica delineata 1: no. 456. 1916.

Monacrosporium tedeschii A. Agostini, Atti Ist. Bot. Lab. Crittog. Univ. Pavia. 4: 195. 1933.

Euricoa dominguiesii Bat. & H. Maia, Anais Soc. Biol. Pernambuco 13: 152. 1955.

Hyaloflorea ramosa Bat. & H. Maia, Anais Soc. Biol. Pernambuco 13: 155, 1955.

Mastigosporium heterosporum R.H. Petersen, Mycologia 51: 729. 1959. Cylindrocarpon lichenicola (C. Massal.) D. Hawksw., Bull. Brit. Mus. (Nat. Hist.). Bot. 6: 273. 1979.

Neocosmospora ramosa (Bat. & H. Maia) L. Lombard & Crous, Stud. Mycol. 80: 227, 2015.

non Fusarium lichenicola (Speg.) Sacc. & Trotter, Syll. Fung. 22: 1486. 1913. nom. Illeqit. (fide Hawksworth 1979).

Selenosporium lichenicola Speg., Anales Mus. Nac. Buenos Aires. 20: 459, 1910.

Type. ITALY, Verona, Tregnago, on Candelaria concolor, Nov. 1902, C. Massalongo (holotype PAD not seen, culture ex-type not known).

Description and illustrations — Wollenweber (1916), Petersen (1959), Hawksworth (1979), Summerbell & Schroers (2002).

Notes — This species is an infrequent agent of human disease, known from localised and invasive infections such as keratitis (Champa et al. 2013), onychomycosis (Guevara-Suarez et al. 2016), mycetoma (Chazan et al. 2004), intertrigo in warm climates, disseminated infection (Rodriguez-Villalobos et al. 2003) and peritonitis (Liu 2011). In addition, it is acknowledged as a phytopathogenic agent infecting *Camellia sinensis* (Shaw 1984), and causing corm rot of *Colocasia esculenta* (Usharani & Ramarao 1981) and fruit rot of pomelo (*Citrus maxima*) (Amby et al. 2015, Farr & Rossman 2017).

Morphologically, it is clearly recognisable in comparison with all other members of the genus in producing ellipsoidal, 0–3-septate aerial conidia that possess a short, truncate base, and that are not curved or pointed like the typical conidia of *Neocosmospora* species. Sporodochia are not produced. These distinctive features led to the species being transferred in the past to the genus *Cylindrocarpon* (Hawksworth 1979). Molecular evidence showed, however, it belongs in *Neocosmospora* (Summerbell & Schroers 2002).

Neocosmospora metavorans (Al-Hatmi et al.) Sandoval-Denis & Crous, comb. nov. — MycoBank MB823687; Fig. 6

Basionym. Fusarium metavorans Al-Hatmi et al., Med. Mycol. 56: S147. 2018.

Type. GREECE, Athens, from human pleural effusion, 2013, *M. Drogari* (CBS 135789 – holotype of *Fusarium metavorans*, maintained as metabolically inactive culture; CBS 135789 – culture ex-type).

Original description and illustrations — Al-Hatmi et al. (2018).

Emended description — Sporulation abundant from sporodochia and from conidiophores formed directly on the substrate and aerial mycelium. Conidiophores in the aerial mycelium up to 285 µm tall, unbranched, sympodial or irregularly branched up to three times at various levels, bearing terminal and single monophialides; phialides subcylindrical, smooth- and thinwalled, $(9-)14-45(-62) \times 4-7(-8)$ µm, with inconspicuous periclinal thickening and somewhat flared collarettes; conidia formed on aerial conidiophores hyaline, ellipsoidal, smooth- and thin-walled, 0-2(-3)-septate, $(4-)11-25.5(-35) \times (2-)4-6(-7)$ µm, single or forming small false heads. Sporodochia at first white, turning ochreous when mature, formed abundantly on the surface of carnation leaves and rarely on the agar surface, later clustering into dry pionnotes. Conidiophores in sporodochia 25-50 µm tall, verticillately branched, bearing 1-6 monophialides in terminal verticils; sporodochial phialides subulate to subcylindrical, $(11-)13.5-19(-22) \times 3-4.5 \mu m$, smooth- and thin-walled, with inconspicuous periclinal thickening and a short, evident, flared collarette. Sporodochial conidia medium to robust, with an almost straight, rarely bent ventral line and a continuous dorsal curvature, wider above the middle portion and tapering toward the basal cell; apical cell equally sized or smaller than the adjacent cell, blunt to slightly hooked with rounded tip; basal cell discretely notched, (1-2-)3-5-septate, hyaline, thin- and smooth-walled. One-septate conidia: 22.5-25 \times 5–5.5 µm; 2-septate conidia: 22.5–27.5 \times 6–7 µm; 3-septate conidia: $(30.5-)38-46(-47.5) \times (5-)5.5-6.5(-7.5) \mu m$; 4-septate conidia: $(43-)45-48.5 \times (5.5-)6-7(-7.5) \mu m$; 5-septate conidia: $(46-)47-51.5(-53) \times (5.5-)6-7.5 \mu m$; overall: $(22.5-)38.5-50(-53) \times (5-)6-7(-7.5) \mu m.$ Chlamydospores abundant, spherical to subspherical 5-13.5 µm diam, solitary or in pairs, terminal and intercalary, smooth- and thick-walled.

Culture characteristics — Colonies on PDA growing at 24 $^{\circ}$ C in the dark with an average radial growth rate of 6.3–7.1 mm/d, reaching 44–50 mm diam in 7 d. Colony surface at first white to

pale straw coloured, gradually turning pale brick to pale coral, flat, felty to cottony with abundant and short aerial mycelium often arranged in concentric rings; colony margins regular. Reverse white to pale yellow or rust coloured. Colonies on OA and CMA incubated at 24 °C in the dark reaching a maximum of 60–71 and 43–50 mm diam in 7 d, respectively. Colony surface white, pale straw to pale luteous or rust coloured, flat, radiated or radially folded, velvety to cottony with abundant white aerial mycelium; colony margins regular. Reverse at first white, then producing luteous or rust coloured pigments.

Cardinal temperatures for growth — Minimum 9 °C, maximum 36 °C, optimal 24–30 °C.

Additional material examined. Spain, from human corneal ulcer, 15 Mar. 1978 (CBS 143194 = NRRL 22782 = IMI 226114); from human foot, 14 July 2004, *F. Ballester* (CBS 143219 = NRRL 46708 = FMR 8634). — Turkey, from human (CBS 143215 = NRRL 37640 = UTHSC R-3564). — USA, Maryland, from human cornea, *M. Brandt* (CBS 130400 = NRRL 43489); San Francisco, from human eye, 14 Dec. 1970 (CBS 143195 = NRRL 22792 = IMI 153617); from human (CBS 143198 = NRRL 28016); from human (CBS 143199 = NRRL 28017); from human (CBS 143200 = NRRL 28018); from human (CBS 143201 = NRRL 28019); New England, from human bone, *A. Fothergill* (CBS 143202 = NRRL 28542 = UTHSC 98-1246); Maryland, from human toenail cancer (CBS 143210 = NRRL 32785 = FRC S-1123); Texas, from human eye (CBS 143213 = NRRL 32849 = FRC S-1355); Michigan, from human chest subcutaneous tissue, 2003, *M. Brandt* (CBS 143216 = NRRL 43717); Illinois, from human, *P. Kammeyer* (CBS 143218 = NRRL 46237).

Notes — One of the most prevalent clades isolated from human clinical specimens, *N. metavorans* is known to cause superficial and deep-seated or disseminated infections (O'Donnell et al. 2008). This species has been also recovered from insects (*Ceresa bubalus*, O'Donnell et al. 2012) and from plant material (Chen & Kirschner 2017, Al-Hatmi et al. 2018). It is also one of the few species in *Neocosmospora* for which a complete genome sequence is available (Coleman 2016, Herr et al. 2016).

This species shows a considerable similitude with N. solani and N. suttoniana in overall culture characteristics and the shape of the sporodochial conidia. However, sporodochial conidia in N. metavorans are slightly wider with conspicuously pedicellate basal cells. By contrast, foot cells are less evident in N. solani. Neocosmospora suttoniana can be differentiated by having much longer and septate sporodochial conidia (up 86.5 µm long and 6-septate) as well as by its verruculose chlamydospores (vs up to 53 µm long and 5-septate sporodochial conidia and smooth-walled chlamydospores in N. metavorans). The protologue of *N. metavorans* also points to a morphological similitude with N. solani s.str. The former species, however, is described as being distinct in the lack of sporodochial conidia and in having conidia in long chains. The ex-type strain of *N. metavorans* may not produce sporodochial conidia, but all the clinical isolates studied here were able to produce sporodochia and multiseptate conidia under standard culture conditions, while conidial chains, which are not an expected characteristic in this genus, were not observed. A re-examination of the ex-type culture is necessary to further evaluate its description. Moreover, we observed a much wider micromorphological variation among our isolates than was noted by Al-Hatmi et al. (2018), and hence, an emended morphological description and illustrations are provided.

Neocosmospora petroliphila (Q.T. Chen & X.H. Fu) Sandoval-Denis & Crous, *comb. nov.* — MycoBank MB822902

Basionym. Fusarium solani (Mart.) Sacc. var. petroliphilum Q.T. Chen & X.H. Fu, Acta Mycol. Sin., Suppl. 1: 330. 1987.

Synonyms. Fusarium solani (Mart.) Sacc. f. sp. cucurbitae W.C. Snyder & H.N. Hansen, Amer. J. Bot. 28: 740. 1941. Race 2.

Fusarium petroliphilum (Q.T. Chen & X.H. Fu) Geiser et al., Fungal Genet. Biol. 53: 69. 2013.

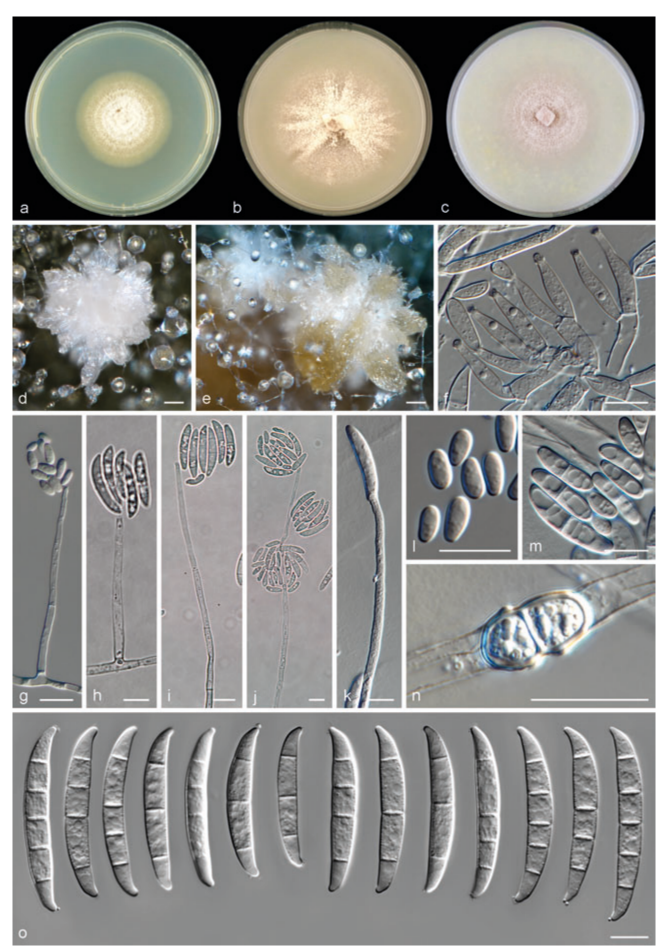


Fig. 6 Neocosmospora metavorans. a. Colony in PDA; b. colony in OA; c. colony in CMA; d-e. sporodochia formed on the surface of carnation leaves; f. sporodochial conidiophore and phialides; g-k. aerial conidiophores and conidia; l-m. aerial conidia; n. chlamydospores; o. sporodochial conidia. — Scale bars: $a-b=20~\mu m$; all others = 10 μm .

Type. China, from deteriorated petroleum (NF 4475, holotype of *F. solani* var. *petroliphilum*, metabolically inactive culture deposited at the Chinese Academy of Sciences Institute of Microbiology, Beijing, not seen; ex-type strain: NF4475 = NRRL 22268 = FRC S-2176).

Description and illustrations — Short et al. (2013).

Notes — Neocosmospora petroliphila and N. keratoplastica are the two most prevalent fusaria species in human clinical samples and are regarded as the most important agents of keratitis (Zhang et al. 2006, O'Donnell et al. 2007). Other known isolation sites of N. petroliphila from humans include blood (O'Donnell et al. 2008, Ersal et al. 2015), nails (Zhang et al. 2006, Guevara-Suarez et al. 2016), nasal mucosa and skin (Zhang et al. 2006, Ersal et al. 2015). Abiotic environments yielding this fungus include contact lens solution and ceiling plaster (O'Donnell et al. 2008). Neocosmospora petroliphila also occurs as the predominant species producing biofilms in plumbing systems together with N. keratoplastica (Mehl & Epstein 2008, Short et al. 2013). The species can infect animals, mostly those with aquatic habitats, such as cetaceans and fish (O'Donnell et al. 2016). It is a recognised agent of fruit rot on cucurbits (Toussoun & Snyder 1961, O'Donnell 2000).

Neocosmospora petroliphila was previously regarded as roughly distinguishable by forming 3–5-septate, falcate, robust sporodochial conidia, which on average were the largest such conidia occurring among the formally described, clinically relevant species known at that time – namely, *N. falciformis*, *N. keratoplastica* and *N. solani* (Short et al. 2013). Two species described here, *N. gamsii* and *N. suttoniana*, exhibit sporodochial conidia that are somewhat similar in shape and septation. Those of *N. petroliphila*, however, can be distinguished by being much shorter than those of *N. gamsii* and *N. suttoniana* (overall: 44–52.2 μm long), as well as markedly and regularly curved.

Neocosmospora suttoniana Sandoval-Denis & Crous, sp. nov. — MycoBank MB822903; Fig. 7

 $\label{eq:constraint} \textit{Etymology}. \ \ \text{In honour and memory of the clinical mycologist Deanna A. } \\ \text{Sutton.}$

Type. USA, Louisiana, from human (CBS H-23224 – holotype; CBS 143214 = NRRL 32858 = FRC S-1423 – culture ex-type).

Sporulation abundant from conidiophores formed directly on the substrate mycelium and less often from sporodochia. Conidiophores in the aerial mycelium erect, up to 250 µm tall, commonly solitary and simple, emerging from the agar surface or sporulating at the agar level, rarely 1-3-times branched laterally, bearing terminal monophialides; phialides subulate to subcylindrical, smooth- and thin-walled, $(6-)23.5-60.5(-63) \times (2-)3-3.5(-4)$ um, with conspicuous periclinal thickening and a minute. discreet collarette; conidia formed on aerial conidiophores, hyaline, obovoid, ellipsoidal, clavate to somewhat cylindrical, straight or curved dorsoventrally, smooth- and thin-walled, 0-2(-3)-septate, $(6-)7.5-21(-31) \times (2.5-)3-5.5(-7.5) \mu m$, single or grouped in false heads at the tip of monophialides. Sporodochia cream to rosy buff coloured, bright, formed scantly and tardily then clustering into dense masses on the surface of carnation leaves. Conidiophores in sporodochia 38-58 µm tall, densely packed, cushion-like, irregularly or verticillately branched, with terminal branches bearing 1–3 monophialides; sporodochial phialides subulate to subcylindrical, often curved near the middle portion, $(12-)13.5-19(-22.5) \times (2.5-)3-4(-5)$ µm, smooth- and thin-walled, without periclinal thickening and with an inconspicuous apical collarette. Sporodochial conidia falcate, widest at the central portion or right above it, gently tapering toward the basal part, robust, somewhat straight on both dorsal and ventral lines; dorsal curvature moderate and often not continuous, being more prominent in the apical and basal thirds; apical cell more or less equally sized or smaller than the adjacent cell, bluntly elongated or distinctly hooked; basal cell somewhat papillate to distinctly notched, (3-)5-6-septate, hyaline, thick- and smooth-walled. Three-septate conidia: $30.5-32.5\times7-7.5~\mu m;~4$ -septate conidia: $(49-)50-53.5\times6-6.5~\mu m;~5$ -septate conidia: $(30.5-)52-71(-77.5)\times(6-)7-8~\mu m;~6$ -septate conidia: $(75-)77-84.5(-86.5)\times(6.5-)7-8~\mu m;~overall~(30.5-)50-75(-86.5)\times(6-)7-7.5(-8)~\mu m.~Chlamydospores~abundant,~spherical~to~subspherical~(4.8-)6-8.5(-9.5)~\mu m~diam,~solitary~or~in~chains,~terminal~or~intercalary,~coarsely~roughened~to~verruculose-~and~thick-walled.$

Culture characteristics — Colonies on PDA growing in the dark with an average radial growth rate of 3.8-5.4 and 5-5.7 mm/d at 21 and 24 °C, respectively, reaching 65-85 mm diam in 7 d at 24 °C. Colony surface straw to olivaceous buff, flat, felty to velvety, aerial mycelium regular, white, formed in radial patches; colony margins regular. Reverse pale luteous to luteous. Pale sulphur yellow to straw diffusible pigments present at 18-36 °C. Colonies on OA and CMA incubated at 24 °C in the dark occupying an entire 9 cm Petri dish in 7 d. Colony colour sulphur yellow to straw, flat, felty to velvety, with rays of abundant aerial mycelium; margins regular. Reverse sulphur yellow to straw, without diffusible pigments.

Cardinal temperatures for growth — Minimum 12 °C, maximum 36 °C, optimal 24–33 °C.

Additional material examined. Gabon, from human nail, M. Kombila (CBS 124892). – USA, Massachusetts, from human, D.A. McGough (CBS 130178 = NRRL 22608 = UTHSC 93-1547); Georgia, from human blood (CBS 143197 = NRRL 28000); Florida, from human corneal ulcer, D.A. Sutton (CBS 143204 = NRRL 32316 = UTHSC 00-264); Florida, from equine eye (CBS 143224 = NRRL 54972 = UTHSC 05-2900).

Notes — Among the newly described species, *N. suttoniana*, previously assigned to clade FSSC 20 of Neocosmospora is the taxon that most closely resembles N. solani s.str. (Schroers et al. 2016), both species producing mostly 5-septate, robust sporodochial conidia. However, while N. solani produces 0-3-5-septate conidia, *N. suttoniana* produces much larger, more frequently septate (up to 6 septa) and more distinctly apically curved conidia, the conidial apex being also more elongated than in N. solani and somewhat hooked. In addition, sporodochia in N. suttoniana tend to develop belatedly, often after more than 10 d of incubation. Apical curvature is a common feature of sporodochial conidia among the clinically relevant species of Neocosmospora; however, it is much more noticeable in N. suttoniana and N. gamsii. The last two species are also distinguishable morphologically (see notes under N. gamsii). Comparable shape and degree of septation of the sporodochial conidia are also recorded for 'Fusarium' ensiforme which, however, produces overall smaller conidia and smooth-walled chlamydospores (Wollenweber & Reinking 1935) vs the verrucose chlamydospores of N. suttoniana. Other species producing rough-walled chlamydospores are 'Fusarium' ventricosum (currently classified as Rectifusarium ventricosum, Lombard et al. 2015) and 'F.' solani var. minus (Wollenweber & Reinking 1935), a species rarely reported as an etiologic agent of mycetoma (El-Zaatari & McGinnis 1993). 'Fusarium' solani var. minus forms mostly 3-septate sporodochial conidia (full range 3-5-septate vs (3-)5-6-septate in *N. suttoniana*), smaller $(20-41 \times 3.5-6 \mu \text{m vs} (30.5-)50-75(-86.5) \times (6-)$ 7–7.5(–8) µm in N. suttoniana) and more prominently curved conidia than those of N. suttoniana. In addition, N. suttoniana produces 0-2(-3)-septate aerial conidia (vs 0-septate in 'F.' solani var. minus). Neocosmospora suttoniana is an uncommon human pathogenic species, up to now reported from blood and causing eye infections in the USA and Africa (O'Donnell et al. 2008).



Fig. 7 Neocosmospora suttoniana. a. Colony in PDA; b. colony in OA; c. colony in CMA; d. sporodochia formed on the surface of carnation leaves; e-f. sporodochial conidiophores and phialides; g-j. aerial conidiophores, phialides and conidia; k-I. chlamydospores; m. sporodochial conidia. — Scale bars: $d-e=50 \mu m$; all others = 10 μm .

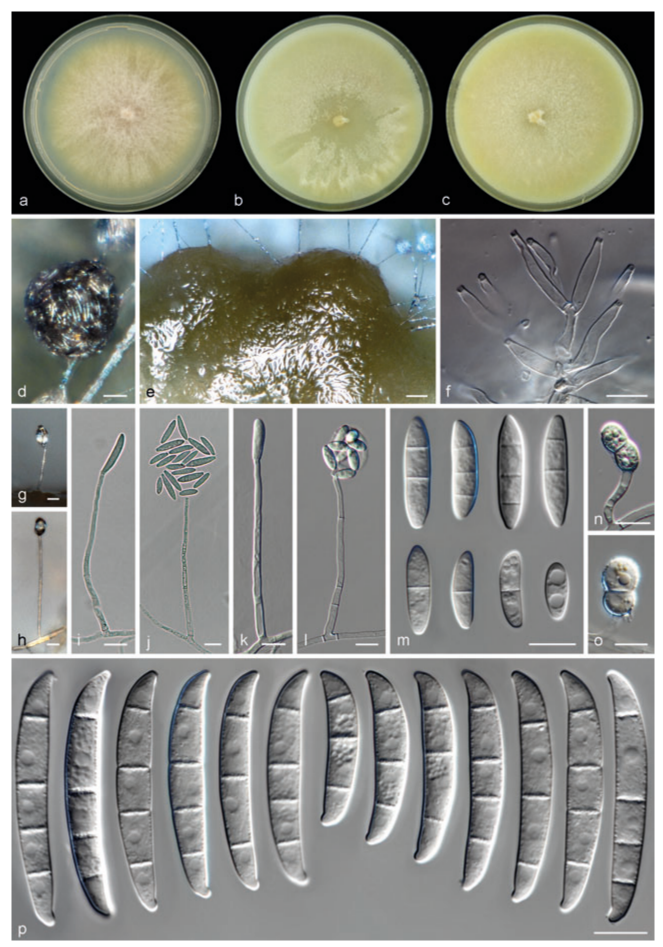


Fig. 8 Neocosmospora tonkinensis. a. Colony on PDA; b. colony on OA; c. colony on CMA; d-e. sporodochia formed on the surface of carnation leaves; f. sporodochial conidiophore and phialides; g-l. aerial conidiophores and phialides; m. aerial conidia; n-o. chlamydospores; p. sporodochial conidia. — Scale bars: $d-e=20~\mu m$; all others = 10 μm .

Neocosmospora tonkinensis (Bugnic.) Sandoval-Denis & Crous, *comb. nov.* — MycoBank MB822904; Fig. 8

Basionym. Cylindrocarpon tonkinense Bugnic., Encycl. Mycol. 11: 181.

Synonym. Fusarium ershadii Papizadeh et al., Eur. J. Pl. Pathol. doi: 10.1007/s10658-017-1403-6: 5 (2018) (nom. illegit., Art 52.1).

Type. VIETNAM, Tonkin, from *Musa sapientum*, 1936, *F. Bugnicourt* No 498 (IMI 113868 – holotype specimen; CBS 115.40 – ex-type culture of *Cylindrocarpon tonkinense*).

Sporulation abundant from sporodochia, and from conidiophores formed on the substrate and aerial mycelium, abundantly produced on hyphal ropes. Conidiophores in the aerial mycelium erect, up to 214 µm tall, simple or branched, branching irregular or verticillate, bearing terminal, long monophialides; phialides subulate to subcylindrical, straight, smooth- and thinwalled, $(42.5-)46.5-63.5 \times 3-4(-4.5) \mu m$, periclinal thickening and collarettes inconspicuous; conidia formed on aerial conidiophores hyaline, obovate, clavate to ellipsoidal, straight or slightly curved, smooth- and thin-walled, 0-3(-4)-septate, $(6-)11-24(-37) \times (3.5-)4-6(-7) \mu m$, single or forming small false heads on the tips of monophialides. Sporodochia at first citrine to hazel coloured turning dark bluish green, brown, vinaceous to purple slate, formed abundantly and clustering on the surface of carnation leaves and on the agar surface. Conidiophores in sporodochia, 22-34.5 µm tall, irregularly or verticillately branched; terminal branches bearing 1-4 monophialides; sporodochial phialides subulate, subcylindrical or somewhat ventricose, often swollen in the middle portion, tapering gently toward the apex $(15-)16-20(-21) \times (2.5-)3-4.5 \mu m$, smoothand thin-walled, with inconspicuous periclinal thickening, and a minute and short apical collarette. Sporodochial conidia wedgeshaped, robust, tapering toward the basal cell, with ventral line gently curved, almost straight between the second septum and the apical cell; dorsal curvature continuous, slightly more pronounced towards the apex; apical cell blunt and typically smaller than the adjacent cell; basal cell blunt to distinctly notched, (1-)3-4(-5)-septate, hyaline, thick- and smoothwalled. One-septate conidia: 47-51 × 6-7.5 μm; 3-septate conidia: $(28-)32.5-42.5(-45.5) \times (5.5-)6-7.5 \mu m$; 4-septate conidia: $(40.5-)43-48(-49) \times 6-7.5 \mu m$; 5-septate conidia: $(40-)41.5-52 \times 6.9-7.3 \mu m$; overall (27.5-)37-48(-50.5) \times (5.5–)6–7(–7.5) µm. *Chlamydospores* abundant, spherical to subspherical 6.5–10(–12) µm diam, hyaline to subhyaline, solitary or in pairs, chains or clusters, terminal or intercalary, smooth- and thick-walled.

Culture characteristics — Colonies on PDA growing in the dark with an average radial growth rate of 3.8–5.1 and 4.3–6 mm/d at 21 and 24 °C, respectively, reaching 76–84 mm diam in 7 d at 24 °C. Colony surface buff, honey with sulphur yellow periphery, flat, felty to floccose, radiated with abundant floccose white to yellow aerial mycelium; colony margins regular, fimbriate. Reverse sulphur yellow to brick coloured. Ochreous to fulvous pigments can be produced between 18–24 °C, a bright yellow pigment is formed between 27–30 °C becoming pale yellow to straw at 36 °C. Colonies on OA and CMA incubated at 24 °C in the dark occupying an entire 9 cm Petri dish in 7 d. Colony colour straw, sulphur to pure yellow, flat, felty, velvety to dusty with abundant short aerial mycelium, margins regular. Reverse sulphur yellow with abundant pure yellow diffusible pigment.

Cardinal temperatures for growth — Minimum 9 °C, maximum 36 °C, optimal 27–33 °C.

Additional material examined. Netherlands, Leiden, from human cornea, Oct. 2017, M.T. van der Beek (CBS 143038). – USA, Florida, from turtle head lesion (CBS 143208 = NRRL 32755 = FRC S-0452); Ohio, from human cornea (CBS 143217 = NRRL 43811).

Notes — *Neocosmospora tonkinensis*, previously known as FSSC 9, is known to include human pathogens, mostly isolated from corneal specimens (O'Donnell et al. 2008, Muraosa et al. 2017), as well as from animal infections (O'Donnell et al. 2008, 2016). Short et al. (2011) reported also the isolation of this species from water drains in the USA.

As already noted by Summerbell & Schroers (2002), the extype strain of C. tonkinense (CBS 115.40) clusters within this clade, but is distinctly separated and thus not congeneric with N. lichenicola as previously alleged (Hawksworth 1979). However, the former authors prevented any taxonomical changes arguing for a probable strain transposition since tapering, curved conidia were observed. During our examination of the ex-type culture, however, we also found the presence of multiseptate, almost cylindrical aerial conidia with more or less rounded apices. Although the observed conidia were slightly smaller and less septate than those reported in the protologue of C. tonkinense (Bugnicourt 1939) (overall from the original description 1-7-septate and 13-45 µm long vs 0-3(-4)-septate and (6-)11-24(-37) µm long in the ex-type); they were more similar in size and shape to those reported for the same strain by Booth (1966), thus a redescription and illustration of the species was provided. The observed differences may easily respond to the different culture conditions employed for the original description of C. tonkinense (slices of carrots and potatoes, beans and citrus twigs). Cylindrical aerial conidia of similar characteristic to those reported here were illustrated in the protologue of Fusarium ershadii, a superfluous name based on the ex-type culture of *C. tonkinensis* (Papizadeh et al. 2018). Similarly, while sporodochia and falcate multiseptate conidia were not observed in the ex-type, they were readily formed in the clinical isolates examined, phylogenetically shown to be conspecific with N. tonkinensis. Sporodochial phialides and conidia strongly resemble those of *N. metavorans*; however, these species are phylogenetically distant.

DISCUSSION

Neocosmospora is perhaps one of the best examples of a fungal genus undergoing fairly rapid speciation (Rossman et al. 1999). Molecular phylogenetic studies have revealed a hidden diversity of phylogenetic species in this genus. There are currently more than 60 recognised genealogically exclusive lineages, many of them showing pathogenic potential against plants, humans and diverse animals (O'Donnell 2000, Summerbell & Schroers 2002, O'Donnell et al. 2008, 2012, 2016, Sandoval-Denis et al. 2018). Our phylogenetic results were highly consistent with previous phylogenetic analyses (O'Donnell et al. 2008, 2016, Gräfenhan et al. 2011, Schroers et al. 2011, Lombard et al. 2015). Neocosmospora was found to be monophyletic, containing a surprisingly high diversity, with many species still needing a proper study and formal descriptions.

Achieving morphological species delimitation and identification in *Neocosmospora* and related genera is a difficult task, especially among human pathogenic species. Although morphological observations proved to be of great value when the appropriate morphological traits were evaluated under standardised culture conditions, we found notable interspecific differences in conidial dimensions, septation and shape for both aerial and sporodochial conidia. These differences, coupled with other features such as the chlamydospore surface texture, the overall cultural growth characteristics and the host of origin, can be of great value for presumptive identification of human and animal pathogenic species. However, considering that these organisms are highly variable in culture, molecular tools should always be applied, in order to ensure correct identification of the involved

species. The general recommendation for clinical microbiologists is to assess species level identification of these pathogens using $EF-1\alpha$ and RPB2 sequences, compared with curated reference sequences deposited in recognised databases as FUSARIUM-ID (http://isolate.fusariumdb.org, Geiser et al. 2004) and Fusarium MLST (http://www.westerdijkinstitute.nl/ Fusarium/) (O'Donnell et al. 2015, 2016). As also confirmed here, these two loci have high resolving power and allowed for a correct delimitation of the clinically relevant clades. This was especially true of RPB2, the only gene in our dataset able to identify all the pathogenic species with great certainty.

Sexual morphs are not usually found in culture. Only a third of the known *Neocosmospora* species, mostly plant-pathogenic taxa, have a known sexual morph (O'Donnell 2000, O'Donnell et al. 2008, Coleman 2016). Among the clinically relevant species, only *N. keratoplastica*, *N. petroliphila*, and an uncommon species, *N. pseudensiformis*, have been described with a sexual morph (Nalim et al. 2011, Short et al. 2013). Here, a sexual morph was described for *N. gamsii*. It was observed only in the ex-type strain and was produced homothallically, after prolonged incubation under standard culture conditions. However, given the infrequent occurrence of sexual structures in *Neocosmospora*, these features are not reliable in species delimitation (O'Donnell 2000).

Neocosmospora catenata was described here without sporodochial conidia, an important morphological feature for generic and, to some extent, specific classification. The lack of macroconidia is not uncommon in fresh Neocosmospora isolates, but in most cases, the production of such structures can be induced using carnation leaf agar or exposure to UV light; these techniques were ineffective in N. catenata. A failure to produce macroconidia should not be regarded as a potential differential character (Leslie & Summerell 2006). Caution is particularly suggested by the knowledge that other Neocosmospora species were originally based on concepts derived from isolates failing to produce macroconidia. For instance, N. falciformis, one the most prevalent fusarial human pathogens (O'Donnell et al. 2008, Guarro 2013) is based on Cephalosporium falciforme, originally described as producing only microconidia grouped in false heads on the tip of thin and elongated monophialides (Carrión 1951). This species was transferred to the genus Acremonium by Gams (1971), partly because of this morphology but also because the human-host-adapted ex-type isolate had a growth rate that fell below Gams' recognition standard for distinguishing Fusarium isolates. Molecular data, however, showed this species to cluster within the 'Fusarium' solani species complex, now Neocosmospora (Summerbell & Schroers 2002). Many fresh isolations of this species have later evidenced the production of distinctive multiseptate conidia, confirming its affinity with Neocosmospora (Edupuganti et al. 2011, Short et al. 2013, Chehri et al. 2015). Similarly, the recently described N. metavorans was characterised as lacking sporodochial conidia (Al-Hatmi et al. 2018). However, sporodochial conidia were readily produced by the large set of human-pathogenic isolates studied here, and the species was appropriately redescribed and illustrated.

The highly relevant clade FSSC 12, although included in our phylogenetic analyses, was not linked to a Latin binomial in this study. Members of this clade have been thoroughly evaluated and a formal description is being prepared in a different study (Geiser pers. comm.). Phylospecies FSSC 12 is known to cause lethal animal infections spanning a wide spectrum of host species, particularly aquatic animals held in captivity. Species affected include American lobster (Hamarus americanus) (Lightner & Fontaine 1975), antler crab (Manucomplanus varians), honeycomb cowfish (Acanthostracion polygonius), horseshoe crab, sea turtle (Lepidochelys kempii) (O'Donnell

et al. 2016), kuruma prawn (*Penaeus japonicus*) (Hatai et al. 1978), lined sea horse (*Hippocampus erectus*) (Salter et al. 2012), stingray (*Taeniura melanopsila*), scalloped hammerhead shark (*Sphyrna lewini*) (Fernando et al. 2015) and treefish (*Sebastes serriceps*) (O'Donnell et al. 2008). The species has also been found in water and sand from human-made aquatic habitats (O'Donnell et al. 2008).

The authors of this paper are aware that the generic placement of these taxa is controversial since, to date, two opposite views exist. However, while some researchers have indicated a preference for conserving the generic name *Fusarium* (= *Gibberella*) in a broad sense, to also include the genus Neocosmospora, no formal decision has yet been made to conserve the broad definition of Fusarium sensu Geiser et al. (2013), against morphologically and phylogenetically supported genera such as Neocosmospora (Gräfenhan et al. 2011, Schroers et al. 2011, Lombard et al. 2015). The concept espoused by Geiser et al. (2013) is broad and polyphyletic, encompassing an artificial arrangement of many distinct clades/genera with clearly different sexual morphologies. We have employed a taxonomical approach that, in our perspective is based on a sound and more natural classification, based not only in molecular phylogenetic exclusiveness, but also considering holomorphic morphological characters. Clinical microbiologists are encouraged to use up-to-date taxonomy and nomenclature for this fungal group and apply the generic name Neocosmospora, which embraces species demonstrated by substantial morphological and molecular evidence not to be congeneric with their closest relatives in Fusarium (Rossman et al. 1999, Gräfenhan et al. 2011, Schroers et al. 2011, Lombard et al. 2015, Sandoval-Denis et al. 2018).

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