**Penicillium simile** sp. nov. revealed by morphological and phylogenetic analysis

Domenico Davolos,1 Biancamaria Pietrangeli,1 Anna Maria Persiani2 and Oriana Maggi2

1Department of Productive Plants and Interaction with the Environment (DIPIA), National Institute of Occupational Safety and Prevention (ex-ISPESL-INAIL), Via Urbana, 167 – 00184 Rome, Italy
2Department of Environmental Biology, Sapienza University of Rome, P.le A. Moro, 5 – 00185 Rome, Italy

The morphology of three phenetically identical Penicillium isolates, collected from the bioaerosol in a restoration laboratory in Italy, displayed macro- and microscopic characteristics that were similar though not completely ascribable to *Penicillium raistrickii*. For this reason, a phylogenetic approach based on DNA sequencing analysis was performed to establish both the taxonomic status and the evolutionary relationships of these three peculiar isolates in relation to previously described species of the genus *Penicillium*. We used four nuclear loci (both rRNA and protein coding genes) that have previously proved useful for the molecular investigation of taxa belonging to the genus *Penicillium* at various evolutionary levels. The internal transcribed spacer region (ITS1–5.8S–ITS2), domains D1 and D2 of the 28S rDNA, a region of the tubulin beta chain gene (*benA*) and part of the calmodulin gene (*cmd*) were amplified by PCR and sequenced. Analysis of the rRNA genes and of the *benA* and *cmd* sequence data indicates the presence of three isogenic isolates belonging to a genetically distinct species of the genus *Penicillium*, here described and named *Penicillium simile* sp. nov. (ATCC MYA-4591T = CBS 129191T). This novel species is phylogenetically different from *P. raistrickii* and other related species of the genus *Penicillium* (e.g. *Penicillium scabrosum*), from which it can be distinguished on the basis of morphological trait analysis.

**INTRODUCTION**

In recent years, the accurate and rapid detection of microfungal species present in the air of indoor workplaces has received a remarkable amount of attention due to the spread of diseases (e.g. allergic responses) caused by airborne moulds and yeasts (Keswani et al., 2005). The recognition and characterization of microfungal species is increasingly being performed by means of a molecular approach based on DNA sequencing analysis, as opposed to methods based exclusively on the observation of morphological traits, which yield poorer results (Taylor et al., 2000; Haugland et al., 2004; Davolos & Pietrangeli, 2007).

Taxa belonging to the genus *Penicillium* have frequently been isolated from bioaerosols. However, although the toxin profiles and allergic reactions of these filamentous fungi differ considerably, they have rarely been identified at the species level (Ramirez et al., 1980). In this study, three phenetically identical *Penicillium* isolates, collected recently during a project conducted to investigate the bioaerosol in a restoration laboratory in Italy, which morphologically resemble *Penicillium raistrickii* but can be distinguished from the latter on the basis of macro- and microscopic characteristics, were subjected to a phylogenetic investigation to obtain a better taxonomic resolution and to gather information on their evolutionary relationships. As molecular markers related to the rRNA genes, we used the internal transcribed spacers (ITS1, ITS2) including the 5.8S rDNA gene, and the domains D1 and D2 of the 28S rDNA gene, together known as the ID region. In addition, two protein coding genes, the tubulin beta chain gene (*benA*) and the calmodulin gene (*cmd*), were examined because they have previously proved to be useful for the molecular investigation of taxa belonging to the genus *Penicillium* at different taxonomic levels (e.g. Samson et al., 2004; Wang & Zhuang, 2007; Serra et al., 2008).

The main goal of our study was to assess the taxonomic status and the phylogenetic position of these three peculiar *Penicillium* isolates by comparing their micro- and macromorphological characteristics and their nucleotide sequence divergence levels (using a multilocus approach).
with those of other known species belonging to the genus *Penicillium*.

**METHODS**

**Strains.** *P. simile* sp. nov. strains 414², 415 and 416 were isolated from the air in three different places in a cultural heritage restoration laboratory at Castel Gandolfo (Rome, Italy) in May 2006. *P. raistrickii* G Smith strains A2a–8 and 3Ba–AP–1 were isolated from the bioaerosol in the crypt of the Capuchin friars in Rome (Italy) in May 2007. All microfungal strains were isolated using a volumetric air sampling surface air sampler, with 5.5 cm RODAC plates containing Sabouraud dextrose agar plus chloramphenicol.

**Media and growth conditions.** According to standardized descriptions, the isolates were examined phenotypically then cultured in Czapek yeast extract (CYA), Blakeslee’s malt extract (MEA) and G2SN agar in 9 cm polystyrene Petri dishes and incubated at 25, 5 and 37 °C for 7 days in the dark, as described by Pitt (1979). The capitalized colour names and codes used to describe the colony colours refer to Kornerup & Wanscher (1967).

**Micromorphological features.** Micromorphological structures (conidia, phialides, metulae and stipes) were photographed and measured using a Leica DM 5000B digital microscope equipped with a Leica DFC420 C camera, differential interference contrast (DIC; also known as Nomarski optics) and Leica LAS image analysis software.

**DNA extraction, PCR amplification and DNA sequencing.** Samples of pure cultures from 3-day-old CYA agar plates were added to 1.5 ml tubes containing TE buffer [10 mM Tris/HCl (pH 8.0), 1 mM EDTA (pH 4.5)], gently homogenized and then lysed by either sonication or enzymic digestion. For physical disruption, tubes were placed in a sonicator (ultrasonic bath LBS2; Falc Instruments) at 60 kHz for 5–3 min then heated at 100 °C for 5–10 min. Samples were centrifuged in a micro-centrifuge for 2 min at 14 000 r.p.m. at room temperature, and the supernatant extracts containing unpurified fungal DNA were then stored at −20 °C for subsequent PCR analysis. Enzymic digestion was instead initially performed with lyticase (20 mg ml⁻¹; Sigma) at 40 °C for 1 h, and then by incubating with proteinase K (20 mg ml⁻¹; Sigma) at 50–60 °C for several hours followed by phenol/chloroform extraction and alcohol precipitation to yield PCR-amplifiable genomic DNA; the resulting DNA pellets were dissolved in 50–100 µl TE buffer and stored at −20 °C.

For the 5.8S rDNA, ITS1, ITS2 and the D1 and D2 regions of the 28S rDNA gene, together known as the ID region, approximately 1100 base pairs (bp), PCR was performed by using primers ITS1, ITS4 (White et al., 1990) and NL–4 (O’Donnell, 1993). In addition, a region of approximately 900 bp of the tubulin beta chain (β-tubulin) gene (*benA*) was amplified by using primer benA1 (Geiser et al., 1998) and the new reverse primer benA3 (5’-GAAGGGAACGATGTGACGAC-3’; Davolos & Pietrangeli, 2007) based on conserved regions of the fungal β-tubulin gene sequence available from GenBank (NCBI). The region of the β-tubulin monomer examined encodes the N-terminal domain [that includes the first 205 residues in which parallel β-strands (B) alternate with α-helices (H)] and helices H6 and H7 (residues 206–240) of the intermediate domain (Nogales et al., 1998). Furthermore, a 550–600 bp-long region of the calmodulin gene (*cmd*), flanking parts of the second and fifth exons, was amplified and sequenced using the primers cmd-f (5’-GATTTCCTCCACCCAGAAGCACA-3’; present study, a modification of the cmdD3 primer (Wang & Zhuang, 2007) based on the *cmd* sequence from the genome of *Penicillium chrysogenum* Wisconsin 54–1255 (van den Berg et al., 2008; Supplementary Table S1) and cmdA1 (5’-GCCYCAGCGGATCATCTCGTC-3’; Wang & Zhuang, 2007), and for *P. simile* sp. nov., the new primers cmd2–f (5’-AGTGTCGGA-GTACAAAGGAGGC-3’; present study) and cmd-r (5’-ATTCGTCGCACTTCTGTGTC-3’; present study) designed according to the *cmd* gene sequences of *P. scabrosum* strains available from GenBank (Supplementary Table S1) and of strains A2a–8 and 3Ba–AP–1 of *P. raistrickii* newly sequenced in this study (Supplementary Table S1).

PCR amplification was performed using Taq DNA polymerase with aliquots of each cell lysate suspension or genomic DNA in TE buffer as a template. Conditions for PCR amplification were: 2 min at 95 °C, followed by 35 cycles, each consisting of 95 °C for 40s, 50–52 °C for 40s, and 1–3 min (according to the examined region size) at 72 °C. The final PCR extension step was 10 min at 72 °C. PCR products were analysed on a 1 % agarose gel and visualized with specific dyes for DNA. Purified PCR products were sequenced (using BigDye terminator v3.1; Applied Biosystems) on both strands using the same PCR primers. Cycle-sequencing reactions were sequenced on an Applied Biosystems ABI Prism 3100 DNA Sequencer.

**Gene sequence and phylogenetic analysis.** Nucleotides and inferred amino acid residues were compared with accessible data from GenBank databases using NCBI’s Blast server. Sequence alignments were performed by using the CLUSTAL X v.1.8 program. Phylogenetic analyses were conducted by using the neighbour-joining (NJ) method, and the maximum-parsimony (MP) method with unweighted codon positions and the close-neighbour-interchange option. The substitution model for the nucleotide sequences was the Kimura two-parameter with complete deletion of gaps; both transitions and transversions were included in the analysis. Phylogenetic analyses performed by means of the NJ and MP methods were conducted using MEGA 4 (Tamura et al., 2007) calculating 1000 bootstrap replications. As the 28S rRNA and *cmd* genes for some of the species examined were not available from GenBank (Supplementary Table S1), concordance was evaluated only for the ITS1–5.8S–ITS2 (known as the ITS region) and the tubulin beta chain gene datasets with the partition homogeneity test implemented with PAUP*4.0b10, using 1000 replicates and the heuristic general search option (Swofford, 2003). The partition homogeneity test revealed no significant difference between the ITS region and the tubulin beta chain gene (*P=0.62*); we also performed a phylogenetic analysis by pooling the ITS and *benA* sequences.

Bayesian analyses were performed in MrBayes v.3 (Ronquist & Huelsenbeck, 2003). The number of substitution types was set to six, and the evolutionary model to the generalized time-reversible model with a gamma–distributed rate variation across sites and a proportion of invariable sites (GTR + Gamma + I); all priors were left default to allow estimation of the parameters from the data. Two independent Markov chain Monte Carlo analyses were run simultaneously for 5 × 10⁶ generations and sampled every 100 generations. The first 25 % of the generations were discarded as burn-in and a 50 % majority rule consensus tree was calculated from the remaining trees.

Sequences of the ID region and the *benA* and *cmd* genes from *P. simile* sp. nov. and *P. raistrickii* obtained in the present study were deposited in the NCBI database (Supplementary Table S1). In addition, since some species of the genus *Penicillium*, including *P. raistrickii*, were traditionally classified in the subgenus *Furcatum* but can be inserted within the subgenus *Penicillium* (Petersen, 2000; Petersen et al., 2003; Frisvad et al., 2006), the ITS1–5.8S–ITS2, 28S rDNA, and *benA* and *cmd* gene sequences obtained from single *Penicillium* lineages phylogenetically representing both the subgenus *Furcatum* (*Penicillium citrinum* and *Penicillium decatum*; Supplementary Table S1) and the subgenus *Penicillium* (*Penicillium brevicepsum*, *P. chrysogenum* strains G6 and Wisconsin 54–1255, and *Penicillium olsonii*; Supplementary Table S1) were included in the phylogenetic analysis and used for the multiple gene comparison.
RESULTS

**Morphology**

The *Penicillium* strains 414T, 415 and 416 displayed identical colony and microscopic morphologies. On the basis of their micro- and macromorphological features, these three *Penicillium* isolates appeared to be most closely related to *P. raistrickii* G. Smith. The similarities we observed included: a velvety, or nearly velvety, colony surface on CYA that was radiately sulcate, with white floccose aerial hyphae; roughened conidiophore walls; the presence of bi- and terverticillate penicilli; spheroidal conidia with smooth, or nearly smooth, walls; well-developed globose to subglobose sclerotia. Other morphological characteristics were, however, distinguishable. The most noticeable differences between these three *Penicillium* isolates and *P. raistrickii* were: on CYA, a colony surface that was radiately sulcate with few furrows and reversed colouring (from light yellow to brown); the presence of light brown exudates; on MEA, the presence of the centre overlaid with sectors of shallow radiating white sterile hyphae; penicilli that were often terverticillate, yielding conidiophores that were less rugose with shorter stipes, metulae and phialides and with bigger conidia.

*Penicillium* strains 414T, 415 and 416 also shared some morphological similarities with *P. scabrosum* (Frisvad et al., 1990), including a velutinous texture and white and/or yellow mycelium on CYA and occasionally reddish-brown exudates; however, our three isolates differed from *P. scabrosum* in the following ways: slower mycelial growth rate both on CYA (26–32 mm diam. after 1 week at 25 °C) and on MEA (21–31 mm diam. after 1 week at 25 °C), colony reverse strongly coloured red–brown in conspicuous concentric zones, colour often diffusing into the agar even on MEA; noticeably roughened and often encrusted conidiophores, ramus at an angle of approx. 45°, much longer metulae and phialides, rough-walled conidia and absence of sclerotia.

The micro- and macromorphological differences observed in *Penicillium* strains 414T, 415 and 416 in comparison with *P. raistrickii*, *P. scabrosum* and other related species, e.g. *Penicillium soppii*, belonging to Section *Ramosum*, series *Lanosa* (Stolk & Samson, 1985; see Frisvad et al., 2006), suggested that they should be transferred to *Penicillium simile* sp. nov., a novel species described here (Figs 1 and 2).

**Molecular and phylogenetic analysis**

For the *P. simile* sp. nov. strains 414T (holotype, see below), 415 and 416, and for the *P. raistrickii* strains A2a–8 and 3Ba–AP–1, the primers for the rRNA regions successfully amplified PCR products of the expected size, a total of approximately 1100 nt (Supplementary Table S1). The sequences obtained from the ID region were aligned with sequences extracted from GenBank (NCBI). *P. simile* 414T (holotype) and the two additional isolates (415 and 416) appeared invariant in the ID region sequences examined. The ITS1–5.8S–ITS2 sequences (approx. 535 nt) of *P. simile* were similar to those of the *P. raistrickii* strains deposited in the NCBI database (Supplementary Table S1), exhibiting three different nucleotides in ITS1 (a transversion, C→A) and ITS2 (two transversions, T→A and A→T). BLAST analysis of the examined 615 nt region of the 28S rDNA gene of *P. simile* compared with those available for *P. raistrickii* (Supplementary Table S1) revealed only a transition (G→A) yielding a 99.8% identity. A marked identity emerged in comparison with *Penicillium swiecickii* NRRL 918T (Supplementary Table S1) and *P. soppii* NRRL 2023T (=IBT 18220T; Frisvad et al., 2006; GenBank no. AF033488), 99.3 and 99%, respectively. The ITS1–5.8S–ITS2 and 28S rDNA trees (data not shown)

![Fig. 1. Colonies of strain ATCC MYA-4591T: (a, b) grown on CYA (left) and MEA (right) at 25 °C, after 7 days, front (a) and reverse (b) views; (c) colony showing radial sulcation and brown exudates.](http://ijs.sgmjournals.org)
demonstrated the presence of two main clades: one group corresponded to species belonging to the subgenus *Furcatum* (*P. citrinum* and *P. decaturense*; Supplementary Table S1), while the other clade was formed by all the remaining species examined, including those chosen here to represent the subgenus *Penicillium* (e.g. *P. brevicompactum* and *P. olsonii*; Supplementary Table S1); within the latter partition, *P. simile* sp. nov. was most closely related to *P. raistrickii*.

Furthermore, for the *P. simile* sp. nov. strains 414T (holotype), 415 and 416, and for *P. raistrickii* A2a–8 and 3Ba–AP–1 (Supplementary Table S1), a section (approx. 900 bp) of the *benA* gene was amplified and sequenced by the primers *benA*1 and *benA*3. An exon–intron structure, assigned by comparing DNA sequences with homologous sequences extracted from GenBank (NCBI), indicated that the introns were bounded by canonical GT–AG splice sites. From the partial *benA* sequence analysis, *P. simile* 414T (holotype) and the two isolates 415 and 416 appeared isogenic. *P. simile* and *P. raistrickii* A2a–8 and 3Ba–AP–1 had the same intron size, though a single deletion was observed in the former. A certain degree of divergence emerged in the *benA* gene sequences (four–eight base pair differences) of previously studied strains (including IBT 21396T) of *P. raistrickii*. *P. raistrickii* A2a–8 and 3Ba–AP–1 displayed identical *benA* gene sequences and, when compared with other strains of *P. raistrickii*, yielded a 98–100 % nucleotide identity (the nucleotide changes occurred in the non-coding regions alone). The *benA* sequence divergence value between *P. simile* and *P. raistrickii* was approximately 3.4 %; the divergence was mainly located in the regions of introns, while their exon sequences only yielded eight different nucleotides. The nucleotide changes in the protein coding region occurred mainly as silent changes, though a single non–synonymous transition (GUU for valine → AUC for isoleucine in α-helix H5) was found in *P. raistrickii* (strains BLSA11t, A2a–8 and 3Ba–AP–1, for which a comparable 900 nt sequence length has been obtained; Supplementary Table S1).

Phylogenetic analysis conducted using the NJ and MP methods revealed the same tree topology. The NJ cladogram of the partial β-tubulin gene produced by using bootstrap analysis in Fig. 3 shows that the species of the genus *Penicillium* examined generally resolved into well supported clades. A clear separation emerged for the clade formed by the two species belonging to the subgenus *Furcatum* (Supplementary Table S1). The *P. raistrickii* strains A2a–8 and 3Ba–AP–1 clustered together in a clade that formed with other conspecific taxa (Fig. 3). *P. simile* sp. nov. was most closely related to *P. raistrickii*, but was clearly distinct from the latter species (bootstrap value of 99 %; Fig. 3). *P. scabrosum* appeared to be a sister species (bootstrap value of 51 %) in relation to the clade formed by *P. simile* and *P. raistrickii* (Fig. 3). Bayesian analysis of the *benA* gene sequences (data not shown) generally confirmed the results yielded by the NJ phylogenetic analysis (Fig. 3), including the observation that *P. simile* and *P. raistrickii* were two distinct evolutionary entities (1.00 posterior probability) and that *P. scabrosum* was recognized as their basal lineage (posterior probability of 0.94).
The NJ tree of the combined sequences of the ITS region and benA gene yielded the topology shown in Supplementary Fig. S1. *P. simile* sp. nov. and *P. raistrickii* were well separated (bootstrap value of 97%; Supplementary Fig. S1). *P. scabrosum* emerged as a sister species (bootstrap value of 94%) to the clade formed by *P. simile* and *P. raistrickii* as well as to another clade (89%) including *Penicillium lanosum*, *P. swiecickii*, *Penicillium jamesonlandense* and *Penicillium ribium* (Supplementary Fig. S1).

In addition, cmd gene sequences (approx. 600 bp) were obtained for *P. simile* sp. nov. (holotype; Supplementary Table S1) and for the *P. raistrickii* strains A2a–8 and 3Ba–AP–1 (Supplementary Table S1). The introns were bounded by GT–AG splice sites; the two isolates of *P. raistrickii* yielded identical cmd sequences. A BLAST analysis showed that the cmd gene sequences of *P. simile* displayed 86% sequence similarity with the cmd gene from strains of *P. scabrosum*. In the cmd region we examined, *P. simile* displayed unique cmd gene sequences that distinguished it from *P. raistrickii*. In particular, the *P. simile* and *P. raistrickii* strains (Supplementary Table S1) had different nucleotides (sequence divergence value of approx. 6.5%) in the intron regions (the third and the fourth intron were slightly different in length due to the presence of indels) while their exon sequences were nearly identical, exhibiting seven different nucleotides that occurred as synonymous changes. Three transitions that occurred in *P. raistrickii* appeared to be apomorphic substitutions in relation to the cmd gene sequences of *P. simile* and *P. scabrosum*. The topology of the NJ tree yielded by the cmd sequences (Supplementary Fig. S2) also identified *P. simile* and *P. raistrickii* as two clearly separate species (100% bootstrap support); *P. scabrosum* emerged as the basal taxon (bootstrap support of 93%) to the clade formed by these two species (Supplementary Fig. S2). The Bayesian inference on the cmd sequences (data not shown) complemented the information yielded by the NJ phylogenetic analyses (Supplementary Fig. S2) and confirmed the genetic separation between *P. simile* and *P. raistrickii* (1.00 posterior probability).

**DISCUSSION**

The increasing amount of attention being dedicated to microfungal species in indoor environments in recent years has enhanced the need for detailed investigations of airborne moulds and yeasts. In particular, a thorough knowledge of airborne microfungal diversity sheds light on the role such microfungi play in respiratory problems, such
as allergies (Scott et al., 2004; Davolos & Pietrangeli, 2007). In this regard, the analysis of gene sequences has proved to be an important means of understanding evolutionary relationships at different taxonomic levels for genera such as *Penicillium*, whose identification on the basis of morphological features has recently posed a challenge (Peteron, 2004; Peterson et al., 2004; Samson et al., 2004). Molecular data have already indicated the genetic isolation of several taxa in the genus *Penicillium*, thereby allowing recent evolutionary speciation among morphologically closely related species to be evaluated (Samson et al., 2004; Kwaśnja & Nirenberg, 2005; Serra & Peterson, 2007; Sonjak et al., 2007a, b) and geographically restricted taxa to be identified (Scott et al., 2004).

In the present study, the two isolates A2a–8 and 3Ba–AP–1, which were isogenic at each of the loci examined, displayed identical ID region sequences and very similar *benA* gene sequences (98.5–100% identity) to those of *P. raistrickii* deposited in GenBank (at the time of writing, *cmd* sequences for *P. raistrickii* were not available in public databases, thus preventing a phylogenetic comparison using this gene), and are identified as such (Supplementary Table S1; Fig. 3; Supplementary Fig. S1); future studies are warranted to determine whether they are biogeographically rare or widespread strains. It is noteworthy, however, that the strains of *P. raistrickii* displayed a certain degree of morphological variability, which points to infraspecific genetic variation (Fig. 3). Future taxonomic studies are needed to redefine the specific structures of this species on the basis of all the information available on taxa belonging to the species.

Although the features of *P. simile* sp. nov., the novel species of the genus *Penicillium* being proposed here, appear to be highly similar to those of *P. raistrickii*, these two species are morphologically distinguishable. Indeed, the differences between *P. simile* (Figs 1 and 2) and *P. raistrickii* can easily be observed by means of both macromorphological (mycelia, hyphae, exudates) and micromorphological (penicilli, metulae, phialides and conidia) investigation.

As regards the molecular data we obtained, the *P. simile* sp. nov. isolates were isogenic at the loci examined, as might be expected given that all three were collected from the same locality. Although the ID region sequences of *P. simile* are similar to those of *P. raistrickii* (Supplementary Table S1), *P. simile* is clearly distinct, as the statistical results unequivocally indicate, from *P. raistrickii* in both the β-tubulin and calmodulin gene regions analysed (Fig. 3, Supplementary Figs S1 and S2), which confirms the phylogenetic importance of such protein coding genes in the study of closely related species of the genus *Penicillium*. Indeed, the phylogenetic position of *P. simile* sp. nov. is distinct within a strongly supported clade based on *benA* gene sequences of *P. jansonii*, *P. lanosum*, *P. raistrickii*, *P. ribiurum*, *P. scabrosum*, *P. soppii* and *P. scitwickii* (Fig. 3), which are traditionally classified in the subgenus *Furcatum* but can, according to recent molecular studies (see Peterson, 2000; Peterson et al., 2003; Frisvad et al., 2006), be inserted within the subgenus *Penicillium*. Our results are in agreement with those of recent studies in which the β-tubulin gene sequences were found to be more variable than those of the ITS region and other genes, often allowing closely related *Penicillium* lineages to be discriminated (Seifert et al., 2007; Peterson & Horn, 2009; Houbraken et al., 2010, 2011). Indeed, the β-tubulin gene divergence value found between *P. simile* and *P. raistrickii* (approx. 3.4%) can be ascribed to the range of genetic variation data between species of the genus *Penicillium* (0.5–67.2%) previously obtained from homologous sequences (Samson et al., 2004; Seifert et al., 2007; O’brien et al., 2008; Houbraken et al., 2010, 2011). Intriguingly, the substitution of the highly conserved valine residue for isoleucine (both non-polar amino acids) in α-helix H5 (see Nogales et al., 1998) found in *P. raistrickii* may be considered an apomorphic molecular trait if compared with related species including *P. simile* (Fig. 3).

In addition, in the *cmd* gene region examined, *P. simile* sp. nov. and *P. raistrickii* appear as two distinct species, as the statistical data clearly indicate (Supplementary Fig. S2); their *cmd* gene sequence similarity is 92.3%, a value that is lower than those reported for closely related species of the genus *Penicillium* (Wang & Zhuang, 2007; Peterson & Horn, 2009). Moreover, some transitions found in the *cmd* regions of *P. raistrickii* appear to be apomorphic substitutions, while *P. simile*, *P. scabrosum* and other related species of the genus *Penicillium* retained the plesiomorphic sequences of the gene.

At present, the information available for the two protein coding loci we examined in this study indicates that *P. scabrosum* may be representative of an early evolutionary stage of *P. simile* sp. nov. and *P. raistrickii* (Fig. 3, Supplementary Fig. S2), thereby suggesting that these three species of the genus *Penicillium* share a common ancestor. The analysis based on the ITS sequences alone reveals few phylogenetically informative sites due to the low variability of this rRNA region (including the identical 5.8S rDNA gene sequences) at that evolutionary scale (see Frisvad et al., 2006). However, the very high bootstrap proportions and Bayesian posterior probabilities suggest, even when the ITS and *benA* gene sequences are pooled (Supplementary Fig. S1), that *P. simile* has an independent evolutionary lineage.

To sum up, our molecular data suggest that *P. simile* sp. nov. and *P. raistrickii* have evolved, as the morphological differences between these species suggest, and should be treated as two closely related, though distinct entities (Fig. 3, Supplementary Figs S1 and S2). These two species of the genus *Penicillium* are likely to have evolved recently from a common ancestor as suggested by low genetic divergence values. In particular, analyses of the β-tubulin and calmodulin gene coding regions (similar exon sequences) point to a recent speciation event within this monophyletic group (Fig. 3, Supplementary Figs S1 and S2), in which *P. raistrickii* appears to exhibit derived molecular traits.
**Taxonomy**

**Latin diagnosis of *Penicillium simile* Davolos, Pietrangeli, Persiani & Maggi sp. nov.**

Coloniae in agaro CYA crescentes, post 7 dies 25 °C, 35 mm diam, radialiter sulcatae et centraliter umbonatae, textura superficie velutinosa ad moderate floccose, margines regulares, moderate profundae, mycelium album ad pallen (M. 1–2A3) usque ad moderate brunneum (M. 5D6–8); conidiogenesis diffusa quondam moderata, concolor callaidis (M. 24A–B6); sclerotia mycelio circumdata, alba ad pallentia (M. 3A3); exudatum moderate brunneum adest (M. 6D1–2); pigmentation dissoluble abest; facies reversa flavam ad brunneam (M. 7D–E7). Coloniae in agaro MEA crescentes, post 7 dies 25 °C, 45 mm diam, plane, radialiter hyphae albae crescentes; sclerotia adsunt post 14 dies, alba ad flavida (M. 3A6–7); exudatum abest; conidiogenesis griseo-viridis (M. 25C4) ad profunde viridem (M. 25E4); facies reversa pallen ad luteam (M. 4A–B8). In agaro G25N coloniae 13–15 mm diam, plane ad radialiter sulcatae, conidiogenesis moderata, facies reversa pallens. Incubata ad 5 °C post 7 dies coloniae 3–5 mm diam, mycelium album. Incubata ad 37 °C in CYA incrementum nullum post 7 dies. Conidiophora portata in aeresc et sub superficie orientibus hyphis, stipes fere longus et latus 280–380 × 3.8–4.7 μm, rugosa, portantes penicillos biverticillatos et terverticillatos, rami rugosi, 20–23 × 2.8–3.8 μm; metulae 3–5 verticillatae, 9.5–11.5 × 2.8–3.8 μm, in apice inflatae vel clavatae, 5.7–6 μm, aliquando rugosae, phialides 8–10 verticillatae, ampulliformes, 6.6–7 × 3.5–3.8 μm; conidia globosa, 2.3–3.3 μm, levia, portata in catenis divaricatis. Sclerotia 130–260 μm diam; Status teleomorphosis ignotus. Typus MB509645.

**Description of *Penicillium simile* Davolos, Pietrangeli, Persiani & Maggi sp. nov. (Figs 1 and 2)**

Colonies grown for 7 days on CYA at 25 °C attain 35 mm diameter, are deeply radially sulcate, centrally umbonate, with a velutinose to moderately floccose texture; moderately deep to deep margins, which are entire (Fig. 1a, left); usually white to pale yellow (M. 1–2A3) or light brown (M. 5D6–8) mycelium; conidiogenesis is often sparse and confined to margin, sometimes moderate and coloured turquoise blue to greyish turquoise (M. 24A–B6); sclerotia usually enveloped by mycelium, white to pale yellow (M. 3A3) with sclerotoid texture (Fig. 2e); exudates usually present (Fig. 1c), light brown (M. 6D1–2); soluble pigment absent; reverse (Fig. 1b, left) from light yellow to brown (M. 7D–E7). Colonies grown for 7 days on MEA at 25 °C attain 45 mm diameter, are plane, not furrowed, with the centre overlaid in sectors by shallow radiate white sterile hyphae, obscuring the otherwise heavily sporulating areas (Fig. 1a, right); strictly velutinous surface texture, margins subsurface, entire, white mycelium; sclerotia not visible after 7 days but visible after 14 days, white to yellow (M. 3A6–7); conidiogenesis moderate from greyish green (M. 25C4) to dull green (M. 25E4); exudates absent; soluble pigment absent; reverse pale in localized sectors corresponding to the overgrowth of vegetative hyphae (Fig. 1b, right), up to orange yellow (M. 4A–B8). On G25N medium, colonies attain 13–15 mm diameter, are plane or lightly radially sulcate, consisting of dense white mycelium with low to deep margins, which are entire; moderate conidiogenesis, absent exudates and soluble pigment; reverse pale. At 5 °C, colonies usually attain 3–5 mm diameter composed of white mycelium. When incubated at 37 °C, no growth is observed in 7 days.

Conidiophores borne from subsurface mycelium as well as from aerial hyphae, with stipes usually 280–380 × 3.8–4.7 μm, occurring at marginal areas, thick and rugose wall, terminating in biverticillate and terverticillate penicilli (Fig. 2a, b, c) with rough-walled rami, 20–23 × 2.8–3.8 μm; terminal metulae in divergent verticils of 3–5, apically inflated or clavate, to 5.7–6 μm at the apices, measuring 9.5–11.5 × 2.8–3.8 μm, sometimes rough-walled; phialides in verticils of 8–10 per metula, ampulliform, 6.6–7 × 3.5–3.8 μm; conidia spheroidal, 2.3–3.3 μm, smooth-walled (Fig. 2d), borne in divergent columns; sclerotia with diameter 130–260 μm (Fig. 2e); not known to produce a teleomorphic state.

**Holotype**

Strain 414T has been deposited (as a herbarium specimen) in the Department of Environmental Biology, Sapienza University of Rome, Rome, Italy (two additional strains, designated 415 and 416, are located in the same herbarium, in the American Type Culture Collection (ATCC) with the accession number ATCC MYA-4591T, and in the Centraalbureau voor Schimmelcultures (CBS) Fungal Biodiversity Centre with the accession number CBS 129191T. Mycobank accession no. MB509645.

**Etymology**

*simile* (L. neut. adj.) from the Latin word *similis* which means ‘similar, related’ and reflects the close morphological similarity of the novel species to *P. raistrickii*.

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