Molecular systematics of *Keratinophyton*: the inclusion of species formerly referred to *Chrysosporium* and description of four new species

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**Abstract**

Four new *Keratinophyton* species (*Ascomycota, Pezizomycotina, Onygenales*), *K. gollerae*, *K. lemmensii*, *K. straussii*, and *K. wagneri*, isolated from soil samples originating from Europe (Austria, Italy, and Slovakia) are described and illustrated. The new taxa are well supported by phylogenetic analysis of the internal transcribed spacer region (ITS) region, the combined data analysis of ITS and the nuclear large subunit (LSU) rDNA, and their phenotype. Based on ITS phylogeny, within the *Keratinophyton* clade, *K. lemmensii* is clustered with *K. durum*, *K. hubeiense*, *K. submersum*, and *K. siglerae*, while *K. gollerae*, *K. straussii* and *K. wagneri* are resolved in a separate terminal cluster. All four new species can be well distinguished from other species in the genus based on phenotype characteristics alone. Ten new combinations are proposed for *Chrysosporium* species which are resolved in the monophyletic *Keratinophyton* clade. A new key to the recognized species is provided herein.

**Keywords:** *Chrysosporium*, Keratinophilic fungi, Keratinolysis, One fungus = one name, New taxa

**INTRODUCTION**

*Keratinophyton* is a genus of microscopic fungi (*Ascomycota, Onygenales, Onygenaceae*) comprising species that live mostly on the remains of hair and feather in soil as saprotrophs (Cano and Guarro 1990; Hubka et al. 2016; Sutton et al. 2013; Vidal et al. 2000). Formerly, they were classified in *Aphanoascus* mainly based on the presence of ascomata (cleistothecia) composed of a membranous peridium (Cano and Guarro 1990; Cano et al. 2002). In a review employing a phenotypic and phylogenetic approach, Cano et al. (2002) accepted 18 *Aphanoascus* species which all have sexual morphs. Only recently, the polyphyletic status of *Aphanoascus s. lat.* has been resolved by Sutton et al. (2013) who established the genus *Keratinophyton* encompassing and redisposing six species, namely *K. durum*, *K. hispanicum*, *K. multisporum*, *K. punsolae*, *K. saturnoideum* and the type species *K. terreum*. Ascospores of *Keratinophyton* species are characterized by a conspicuous equatorial rim and pitted wall, while *Aphanoascus* species have reticulate ascospores without a rim (Sutton et al. 2013). Within *Keratinophyton*, only *K. multisporum* is related to a *Malbranchea* asexual morph (Cano and Guarro 1990), while the remaining known species have a *Chrysosporium* asexual morph. In addition to the above mentioned species, the monophyletic *Keratinophyton* clade currently encompasses at least 11 species known only as asexual morphs (Cano and Guarro 1990; Hubka et al. 2016; Sharma and Shouche 2017; Liang et al. 2009; van...
Oorschot 1980; Vidal et al. 2000; Vidal et al. 2002; Zhang et al. 2016; Zhang et al. 2017). Sharma and Shouche (2017) introduced a new species, Keratinophyton turgidum, based on the morphology of its chrysosporium-like aleuric formations and ITS locus phylogenetics. The same authors stated that all species in this clade have a Chrysosporium asexual morph require redisposing in the genus Keratinophyton.

The presence of this large group of ubiquitous and keratinolytic species is rather common especially in areas with high animal activity that results in transfer of the keratinous material (fur, hairs, etc.) to the soil (Papini et al. 1998; Vidal et al. 2000). The following reports confirm their world-wide distribution and occurrence in different habitats usually associated with soil environments, e.g. soil in city parks (Papini et al. 1998; Vidyasagar et al. 2005), flower pots (Singh et al. 2009), sand in children’s sandpits (Labuda et al. 2008), mud (Zaki et al. 2005), poultry farms (Anbu et al. 2004; Cano and Guarro 1990), marshy meadows, salt pans, desert, cultivated or uncultivated soils (Cano and Guarro 1990; Chmel and Vlăciciková 1977; Deshmukh 2004; Deshmukh et al. 2008; Han et al. 2013; Javorekova et al. 2012; Zhang et al. 2016; Zhang et al. 2017) and river sediments (Ulfig et al. 1997; Vidal et al. 2000; Vidal et al. 2002). In general, these fungi are rarely reported as animal pathogens, and in fact, only two species C. echinulatum and C. pan-nicola (formerly known as C. evolceanui) have been involved in mycoses (Hajsig et al. 1974; Cabanes et al. 2014; Hubka et al. 2016).

During a microbiological survey of environmental samples (soil and compost) in July 2019, several interesting Chrysosporium asexual morphs were isolated. These isolates were phenotypically similar to those previously isolated from the same samples in August 2015 by one of us (R.L.). These isolates were designated BiMM-F76, BiMM-F77 (also strain RL-07, isolated in July 2019), BiMM-F78 (also strains RL-05 and RL-06, isolated in July 2019), and BiMM-F250. All strains were further characterized in terms of morphology, physiology, and molecular phylogeny. Phylogenetically informative sequences were obtained from the internal transcribed spacer (ITS) region of the nuclear large subunit (LSU) rDNA. Overall, the resulting data revealed that these isolates represent novel species of the genus Keratinophyton, and they were illustrated for the first time in this paper.

MATERIALS AND METHODS

Sample collection and isolation of the fungi

A sample of a garden soil in Vieste (Italy) was collected in July 2004, one of a forest soil in Tatranská Lomnica (The Slovak Republic) in August 2011, and one of compost from an agricultural base at the Institute of Agrobiotechnology (IFA Tulln, Austria) in August 2015. All three samples were taken from the surface layer (3–5 cm deep), dried, and stored in plastic bags in a fridge (5–8 °C) until the time of analysis (August 2015 and July 2019). Isolation of the keratinophilic fungi was performed as described previously (Javorekova et al. 2012). Each sample was divided into 10 subsamples. The subsamples (20 g each) were poured into Petri dishes and soaked with antibiotic solution containing 0.5 g cycloheximide and 0.1 g chloramphenicol. Sterile defatted horse hair fragments (10 pieces of ca 2.0 cm per plate) were used as baits. The Petri dishes were then incubated at laboratory temperature (23–25 ± 1 °C), under ambient daylight, for a period of 2–3 months and remoistened with sterile deionized water when necessary. The Petri dishes were checked weekly for the presence of fungi, and isolates were cultured on Sabouraud 4% dextrose agar (SDA; Merck, Darmstadt, Germany) supplemented with 0.5 g cycloheximide and 0.05 g chloramphenicol. Pure cultures were then transferred onto potato dextrose agar (PDA; Van Waters and Rogers (VWR) International, Leuven, Belgium). The preliminary identification of the resulting keratinophilic fungi was carried out based on their phenotypic characteristics according to van Oorschot (1980) and Vidal et al. (2000, 2002).

Morphological analysis

For phenotypic determination, the strains were transferred (three-point inoculation with a needle) to PDA, Malt Extract Agar (MEA; Merck, Darmstadt, Germany), and SDA, and incubated for 14 d in the dark at 25 °C. Christensen’s urea agar (Sigma-Aldrich, St Louis, MO, USA) was used for additional physiological and biochemical characteristics (25 °C, 14 d, in the dark). Corn Meal Agar (CMA; Oxoid, Basingstoke, UK), Potato Carrot Agar (PCA) (Samson et al. 2010) and Emerson YpSs agar (Atlas 1946) were used for stimulation of sexual reproduction (at 20 °, 25 °, and 28 °C, for up to 3 months in the dark).

Colony size (mm), colony structure and characteristics were noted after 14 d (on PDA, MEA, SDA, PYE, YpSs, CMA, and PCA). However, the cultivation was extended up to 3 months to observe and record changes in pigmentation of the colonies as well as to determine the onset of sexual reproduction. In order to determine the optimal and minimum/maximum temperatures for growth, PDA, MEA and SDA plates were incubated at 5 °, 8 °, 10 °, 12 °, 15 °, 18 °, 20 °, 25 °, 28–32 °, 35 °, and 37 °C, and the growth rate was measured on the 14th day of cultivation. For comparative descriptions of the macroscopic and microscopic characteristics, PDA was used according to Vidal et al. (2002), Hubka et al. (2016) and Sharma and Shouche (2017).
For determination of microscopic traits, PDA was used after 14–18 d. Conidiophore and conidia formation were observed in situ under low magnification (50–100x). Details of conidiophores, conidia (aleurioconidia) and other microscopic structures, such as width of hyphae, were observed in Melzer’s reagent and lactic acid with cotton blue. Photomicrographs were taken in Melzer’s reagent and lactic acid with cotton blue using phase and Nomarski contrast optics on an Olympus BX51 microscope with Olympus DP72 camera and QuickPHOTO Micro 3.0 software. Photographs of the colonies were taken with a Sony DSC-RX100.

Scanning electron microscopy (SEM) was performed on a JEOL JSM-6380 LV microscope (JEOL, Tokyo, Japan). Fungal samples were prepared according to a simplified method (Samson et al. 1979). Pieces of colonies (ca. 3 × 5 mm) growing on PDA were fixed in 6% glutaraldehyde overnight in the refrigerator (ca. 20 h), then dehydrated in 2-methoxyethanol for 10 min. This was followed by critical point drying and gold coating in a BAL-TEC SCD 050 Sputter Coater. The samples were observed with spot size 35–39 and accelerating voltage 20–23 kV.

Dried fungarium specimens deposited as holotypes in the collections of the Mycological Department, National Museum in Prague, Czech Republic (PRM); ex-type cultures were deposited in the Bioactive Microbial Metabolites (BiMM) Fungal Collection, UFT- Tulln in Austria and in the Culture Collection of Fungi in Prague (CCF).

Keratinolytic activity
Keratinolytic activity was tested by placing a few sterilized blond hairs of a 5 y old child on a PDA plate 1 cm away from the point of inoculation (van Oorschot 1980). Ability to digest keratin was observed after 21 d of incubation at 25 °C in the dark. In addition, a hair perforation test was also performed following de Hoog et al. (2020) using 25 mL water containing 2–3 drops 10% yeast extract (YEW). The hairs were examined microscopically after 14 and 21 d of the inoculation at 25 °C in the dark. At the end of the incubation period, a few pieces of hair were taken out from the testing media (PDA and YEW). The overgrowing fungus was deactivated with 70% ethanol and then removed from the hair surface mechanically in a stream of a tap water. The degree of hair digestion-degradation (keratinolytic activity) was assessed in the light microscope under 100x and 400x magnification. For the observation and microphotography of the hairs, water was used as mounting fluid. Intensity of degradation of the hair was estimated on a scale of 0 to 4 (Marchisio et al. 1994): 0 = no degradation; 0–1 = light degradation on the cuticle; 1 = moderate degradation on the cuticle and/or rare formation of boring hyphae; 2 = degradation of cuticle and cortex, with about 20% degradation of the hair; 3 = degradation of cuticle and cortex, with about 50% degradation of the hair; 4 = degradation of cuticle and cortex, with about 80% degradation of the hair. The photomicrographs of the hairs were taken using a Motic BA 310 microscope with Motic Image Plus 3.0 software. The final microscopic pictures were black-and-white inverted.

DNA extraction, PCR amplification and sequencing
DNA was extracted using a standard cetyltrimethyl ammonium bromide (CTAB) procedure, as described previously (Doyle and Doyle 1987). The internal transcribed spacer (ITS) region was amplified with primers ITS1-F (Gardes and Bruns 1993) and ITS4 (White et al. 1990) using Taq-polymerase. The D1/D2 domains of the large-subunit (28S) rRNA gene (LSU) were amplified and sequenced using the primer pair ITS1/TW14 (White et al. 1990; Mori et al. 2000). All reactions were performed in an Eppendorf Gradient MasterCycler (Eppendorf, Hamburg, Germany). Conditions for amplification of ITS and LSU domains: 95 °C for 5 min; 35 cycles of 95 °C for 30 s, 54 °C for 30 s, 72 °C for 90 s, and finally 5 min at 72 °C. The PCR products were sequenced with the same primers used for the PCR amplifications (Micosynth, Balgach, Switzerland). All sequences obtained in this study were deposited in GenBank nucleotide database (Table 1).

Phylogenetic analysis
For phylogenetic analysis, sequences were aligned with ClustalX (Larkin et al. 2007). Phylogenetic analysis based on ITS locus was performed using GTR + I + G4 + F model with 1000 bootstrap replicates on IQ-TREE web server (Trifinopoulos et al. 2016) and ITS-LSU combined data phylogeny was constructed using MRBAYES v3.2.7a dev (Ronquist and Huelsenbeck 2003) with default settings on the CIPRES portal (http://www.phylo.org/). Ctenomyces serratus (type species CBS 187.61) was used as an outgroup. TREEVIEW v1.6.6 (Page 1996) and iTOL v6 (Letunic and Bork 2019) were used to display and edit phylogenetic trees.

RESULTS

Phylogenetic analyses and keratin degradation
The results of the morphological analyses are given for each novel species under the Taxonomy section below. Temperature dependent growth of the new Keratinophyton species on PDA, MEA and SDA after 14 d are provided in Table S1a–c. Briefly, K. lemmensii grew better than the other three new species on the same type of media and at the same incubation temperatures. All species showed good growth at 20–25 °C on all three media.

Ability to digest keratin after 21 d was observed in all four new species on both testing media (PDA and
YEW). However, a value of attack intensity on the hair according to the scale of Marchisio et al. (1994) differed substantially amongst the species. It was very strong in *K. gollerae* and *K. straussii* (=4), moderate in *K. wagneri* (=2), and weak in *K. lemmensii* (=0–1) (Fig. 10).

**Phylogenetical analysis**

The phylogenetic tree of ITS dataset (*n* = 32) was 551 bp in length which had 286 variable and 200 parsimony-informative sites. ITS phylogeny indicated the presence of six terminal clusters in the monophyletic *Keratinophytin* clade with high bootstrap support and low interspecific sequence divergence (Fig. 1a). *Keratinophytin saturnoideum* and *K. minutisporosum* formed a basal branch to the clade. Isolate BiMM-F76 (*K. lemmensii* sp. nov.) was close to *K. durum* (with 99% ITS and 95% LSU similarity) and clustered also with *K. hubeiense* and *K. gollerae* specific sequence divergence (Fig. 1a).

**Table 1** List of the strains included in the study

| Species name | Strain* | Source | GenBank accession numbers |
|--------------|---------|--------|---------------------------|
| *A. canadensis* | UAMH 4574 | Carnivore dung, Canada | AJ439435 |
| *A. clathratus* | IMI 320400 | Arable soil, Spain | AJ439436 |
| *A. cubensis* | FMR 4220 | Soil of tobacco field, Cuba | AJ439432 |
| *A. foetidus* | CBS 453.75T | Myomys daitoni coat, Nigeria | KT155907 |
| *A. fulvescens* | NBRC 30411 | Soil of rice paddy field, Japan | JN943432 |
| *A. keratinophilus* | IFM 55159T | Pasture land soil, Papua New Guinea | NR165936 |
| *A. mephitis* | IMI 151084T | Dung of wolf, Canada | AJ439439 |
| *A. orissae* | CBS 340.89 | Soil in animal husbandry, Kuwait | AJ390393 |
| *A. pinea* | FMR 4221 | Forest soil, Cuba | AJ439433 |
| *A. reticuliporus* | CBS 392.67T | Soil, New Zealand | MH859002 |
| *A. verrucosus* | NBRC 32381T | Arable soil, Spain | NR131309 |
| *K. clavisporum* (C. clavisporum) | G80.1T | Plant root soil, China | KY026601 |
| *K. durum* | CBS 118.85T | Soil, Nepal | MH61856 |
| *K. echinulatum* (C. echinulatum) | CCF 4652T | Soil of the foot, Czechia | LT548276 |
| *K. fluviatile* (C. fluviatile) | FMR 6005T | River sediments, Spain | AJ005367 |
| **K. gollerae** | BiMM F250 | Forest soil, Slovakia | MN633084 |
| *K. hispanicum* | CBS 456.90T | Beach soil, Spain | KT155910 |
| *K. hubeiense* (C. hubeiense) | EM6661T | Soil under the chicken feather, China | KJ849227 |
| **K. lemmensii** | BiMM F76 | Compost soil, Austria | MN633082 |
| *K. linfense* (C. linfense) | GZAC H31T | Rhizosphere soil, China | NR182829 |
| *K. minutisporosum* (C. minutisporosum) | IMI 379912T | River sediments, Spain | KT155616 |
| *K. pannicola* (C. pannicola) | CBS 116.63T | Soil, India | AJ005368 |
| *K. punsolae* | IMI 334198T | Arable soil, Spain | AJ439440 |
| *K. qinghaiense* (C. qinghaiense) | GZUIFR Chry 11T | Farmland soil, China | JX868607 |
| *K. saturnoideum* | CBS 628.88T | Arable soil, Spain | NR077135 |
| *K. sigleae* (C. sigleae) | UAMH 6541T | Garden soil, Spain | AJ31684 |
| **K. straussii** | BiMM F78 | Garden soil, Italy | MN633081 |
| *K. submersum* (C. submersum) | CBS 101575T | River sediments, Spain | NR157445 |
| *K. teneum* | CBS 342.64T | Lawn soil, India | KT155876 |
| *K. turgidum* | CBS 142956T | Barber shop soil, India | KY290503 |
| **K. wagneri** | BiMM F77 | Forest soil, Slovakia | MN633083 |
| *Ct. serratus* | CBS 187.61T | Soil, Australia | NR144890 |

*BiMM, Bioactive Microbial Metabolites Unit, UFT-Tulln, Austria; UAMH, University of Alberta Microfungus Collection and Herbarium; IMI, CAB International Biosciences, Egham, UK; FMR, Facultad de Medicina in Ciencias de la Salud, Reus, Spain; CBS (Westerdijk Fungal Biodiversity Institute), Utrecht, The Netherlands; NBRC, NITE Biological Resource Centre, Japan; IFO, Institute for Fermentation, Osaka, Japan; G, EM, and GZUIFR strains, The Institute of Fungus Resource, Guizhou University, China; A, Aphanoascus; K, Keratinophyton; C, Chrysosporium; Ct, Cenomyces; T, ex-type culture. Data in bold generated in the present study.
gollerae sp. nov., and K. wagneri sp. nov., represented by the ex-type cultures BiMM-F78, BiMM-F250 and BiMM-F77, respectively, were resolved in a separate terminal cluster-lineage. A concatenated phylogeny of ITS and LSU sequences \( (n = 22) \) was 1094 bp length and included 354 variable and 224 parsimony-informative sites. According to a combined data set analysis, four clusters were found in the Keratinophyton clade with K.
saturnoideum as a basal branch (Fig. 1b). Differently from the ITS phylogeny, K. durum was placed in a different cluster from K. submersum and K. lemmensis in the concatenated loci phylogeny (Fig. 1b).

**TAXONOMY**

The phylogenetic analyses strongly supported the recent distinct classification of the species previously classified as Chrysosporium and only known from asexual morphs into two phylogenetically different genera, Aphanoascus and Keratinophyton (Sharma and Shouche 2017; Sutton et al. 2013). Species described in Chrysosporium which were resolved in a monophyletic clade with Keratinophyton are therefore combined into Keratinophyton in the present paper and provided together with four new Keratinophyton species. The main distinguishing phenotypic characteristics of the four new species were compared with those in the other members of the genus that are also unable to produce ascomata (Table 2).

*Keratinophyton clavisperum* (Zhang, Han & Liang) Labuda & Bernreiter, **comb. nov.**

MycoBank: MB833653

*Basionym: Chrysosporium clavisperum* Y.W. Zhang, Y.F. Han & Z.Q. Liang - *Phytotaxa* 303: 177; 2017.

*Type:* GZUIFR-G80.1; isolated from plant root soil by Y. Luo, China. For detailed description of the species, see the Zhang et al. (2017).

*Keratinophyton echinulatum* (Hubka, Mallátová, Čmoková & Kolařík) Labuda & Bernreiter, **comb. nov.**

MycoBank: MB833636

*Basionym: Chrysosporium echinulatum* Hubka, Mallátová, Čmoková & M. Kolařík - *Persoonia* 36: 410; 2016.

*Type:* CCF 4652 = CBS 141178 = UAMH 11824; from sole of the foot by N. Mallátová, Czechia. For detailed description of the species, see the Hubka et al. (2016).

*Keratinophyton fluviale* (Vidal & Guarro) Labuda & Bernreiter, **comb. nov.**

MycoBank: MB833637

*Basionym: Chrysosporium fluviale* Vidal & Guarro - *Mycol. Res.* 104: 245; 2000.

*Type:* CBS 100809 = FMR 6005 = IMI 378764, isolated from river sediments, by P. Vidal, Spain. For detailed description of the species, see the Vidal et al. (2000).

*Keratinophyton gollerae* Labuda, Bernreiter, Kubátová, Schüller & Strauss, **sp. nov.**

(Figs. 2 and 3)

MycoBank: MB833633

*Etymology:* Named in honour of Sabine Strauss-Goller, Department of Applied Genetics and Cell Biology, Fungal Genetics and Genomics Laboratory, University of Natural Resources and Life Sciences, Vienna (BOKU), Austria, an expert in the fungal genetics and indoor mould analyses.

*Type:* Slovak Republic: Tatranská Lomnica, from forest soil, Jul. 2019, R. Labuda (PRM 952499 – holotype; BiMM-F250 = CCF 6360 – ex-type cultures). ITS sequence, GenBank MN633084; LSU sequence, GenBank MT874997.

*Description:* Sexual morph not observed on any of the media used. Asexual morph on PDA. Vegetative mycelium of hyaline, septate, smooth-walled, sparsely to pronouncedly branched hyphae, often at right angles, 1.0–5.0 μm diam. Racquet hyphae present. Conidia (aleurioconidia), hyaline, white in mass, thin-walled, mostly smooth to finely roughened, some also verrucose (light microscope) and irregularly ornamented with minute warts (SEM). Terminal and lateral conidia born on main fertile hyphae or from side branches of variable length, sessile or on short protrusions, occasionally only very slightly swollen and of variable length, solitary, 1–3 (5) per conidiogenous cell, obovate to clavate, mostly 1-celled, (3.5–)5.0–7.0(–10.0) x (1.5–)2.0–2.5(–3.0) μm (mean = 5.2 ± 0.9 x 2.2 ± 0.2 μm, n = 120). Intercalary conidia not observed. Chlamydospores not observed.

*Culture characteristics:* Colonies on PDA 20–22 mm diam at 25°C, after 14 d, powdery to downy (mealy), with abundant sporulation, white to creamy, flat, umbonate at the centre, with regular colony margin submerged into agar, reverse white to slightly yellowish, no pigment or exudate produced. At 30°C, no growth (germination only). Colonies on SDA 23–25 mm diam at 25°C, after 14 d, morphology similar to when on PDA with more floccose colony margin and more yellowish colonies, with dark yellow reverse. At 30°C, no growth (no germination). Colonies on MEA 14–16 mm diam at 25°C, after 14 d, morphology similar to PDA with more floccose colonies and with yellow reverse. At 30°C, no growth (no germination). Colonies on CMA and PCA attaining 15–20 mm diam at 25°C, after 21 d, white, granular, with good sporulation, reverse yellowish. No ascomata observed after prolonged incubation (3 months). The optimum temperature for growth on PDA, SDA and MEA 15–25°C (Table S1a–c). Minimum growth (microcolonies to 1 mm in diam) at 10°C. Germination of the conidia observed at 8°C. The maximum temperature for growth on PDA 29°C, while 27°C and 28°C on MEA and SDA, respectively (microcolonies to 1 mm diam). Keratinolytic activity very strong (Fig. 10b), with hair attack intensity = 4. Urease activity negative (after 14 d of incubation).

*Diagnosis:* Keratinophyton gollerae molecularly can be distinguished from other Keratinophyton species by ITS locus analysis. Combination of the following phenotypic features can be used to differentiate this fungus from other species in the genus: (1) obovoid-clavate and smooth to finely roughened conidia, (2) No growth at
Table 2: Comparison of the key phenotypic characteristics of *Keratinophyton* species

| Species | Growth at 30 °C on PDA | Colony color, growth/ reverse on PDA at 25 °C, after 14d | Conidial shape | Conidial dimensions (μm) | Conidial surface | Intercalary conidia | References |
|---------|------------------------|----------------------------------------------------------|----------------|--------------------------|-----------------|---------------------|------------|
| *K. gollerae* sp. nov. | None | White to creamy, 20–22 mm/white to yellowish | Obovoid to clavate | 5.0–70 × 20–2.5 | Smooth to finely roughened | Absent | This study |
| *K. lemmensii* sp. nov. | Present (good) | White, 28–35 mm/lemon yellow | Clavate to filiform | 3.0–40 μm (1- to 2-celled) | Smooth | Present | This study |
| *K. straussii* sp. nov. | Present (good) | White to creamy, 24–28 mm/white to yellowish | Obovoid to clavate | 4.5–50 × 25–3.0 | Verrucose | Absent | This study |
| *K. wagneri* sp. nov. | Present (restricted) | White to yellowish, 25–30 mm/white to yellowish | Obovoid to clavate | 4.0–80 × 25–4.0 | Verrucose | Absent | This study |
| *K. clavisporum* | Present (restricted) | White, 53 mm (26 °C)/red-brown | Clavate to long-ellipsoidal | 5.0–10 × 2.5–5.0 | Smooth | Absent | Zhang et al. 2017 |
| *K. echinulatum* | Present (good) | Yellow to pale orange yellow, 28–45 mm/orange yellow | Obovoid to clavate | 4.5–70 × 2.5–4.0 | Echinulate | Present | Hubka et al. 2016 |
| *K. fluviatile* | Present (good) | White to yellowish white, 60–70 mm (30 °C)/brownish orange | Obovate, clavate, nearly ellipsoidal or pyriform | 3.5–15 x 2.0–3.0 (1- and 2-celled) | Verrucose | Present (very rare) | Vidal et al. 2000 |
| *K. qinghaiense* | Present (good) | White to yellowish, 30 mm (7 days)/yellowish | Clavate to cylindrical | 3.6–13 × 1.8–3.6 | Smooth | Present | Han et al. 2013 |
| *K. hubeiense* | Present (restricted) | Grey white to white, 65–66 mm/reverse yellowish | Obovoid to ellipsoidal | 2.2–43 × 1.6–3.2 | Smooth | Absent | Zhang et al. 2016 |
| *K. linfenense* | Present (good) | White to cream, 72 mm (30 °C)/white to light yellow | Ellipsoidal to fusiform, also clavate | 3.2–54 × 1.4–22 | Smooth | Absent | Liang et al. 2009 |
| *K. minutisporosum* | Present (good) | White to yellowish white, 55–70 mm/white | Pyriform or subglobose, also clavate | 3.0–40 (−11) × 1.5–3.5 | Verrucose | Present (very rare) | Vidal et al. 2002 |
| *K. pannicola* | Present (good) | White to pale yellow, 20–38 mm/pale brown | Obovoid to clavate | 6.0–11 × 3.5–4.5 | Verrucose | Present (less abundant) | van Oorschot 1980 |
| *K. siglerae* | Present (good) | Griseous orange, 15–20 mm (21 d)/pale brown | Cylindrical to clavate | 5.0–30 × 2.0–3.5 (1- and 2-celled) | Smooth to slightly verrucose | Present | Cano and Guarro 1994 |
| *K. submersum* | Present (restricted) | Yellowish white, 50–60 mm/yellowish white | Clavate, also pyriform, obovoid and subglobose | 4.0–35 × 2.5–5.0 (1- to 4-celled) | Smooth to verrucose-thick-walled | Present (in old cultures) | Vidal et al. 2002 |
| *K. turgidum* | Present (good) | White, 50–55 mm (SGA at 28 °C)/pale brown | Pyriform to oval | 5.0–70 × 3.5–5.0 | Smooth | Present | Sharma and Shouche 2017 |

*If not stated other medium  
*If not stated otherwise  
PYE: Phytone yeast extract agar  
Yanfeng Han personal communication
Notes: Based on a search of NCBI GenBank nucleotide database, the closest hit for *K. gollerae* using the ITS sequence is *K. minutisporosum* (as *Chrysosporium minutisporosum* CBS 101577; GenBank acc. KT155616), with identity = 487/543 (90%) and gaps 11/543 (2%). Phenotypically, *K. gollerae* can be readily distinguished from the *K. minutisporum* by its smooth to finely roughened larger conidia (5–7 × 2–2.5 μm vs. 3–4 × 1.5–3.5 μm), dark yellow colony reverse at 25°C on PDA. Based on ITS phylogeny (Fig. 1a), *K. gollerae* formed a cluster together with *K. straussii* and *K. wagneri*, and it can be differentiated by its inability to grow at 30°C, narrower and mostly smooth to finely roughened conidia, and its slower growth at 25°C on PDA. Moreover, in comparison with *K. straussii*, *K. gollerae* grows substantially faster at 15°C (on PDA and SDA) and its conidia germinate at 8°C (see Table S1a–c).

*Keratinophyton hubeiense* (Zhang, Han & Liang) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833638
Basionym: *Chrysosporium hubeiense* Yan W. Zhang, Y.F. Han & Z.Q. Liang - Phytotaxa **270**: 213; 2016.
Type: GZAC EM66601, isolated from soil under the chicken feather by Y.R. Wang, China. For detailed description of the species, see the Zhang et al. (2016).

*Keratinophyton lemmensii* Labuda, Bernreiter, Kubášová & Schüller, **sp. nov.**
(Figs. 4 and 5)
MycoBank: MB833632

Etymology: Named in honour of Marc Lemmens, Department of Plant Protection, University of Natural Resources and Life Sciences, Vienna (BOKU), Austria, an expert in fungal plant pathology.

Type: Austria: Tulln and der Donau, from compost soil at IFA Tulln, Aug. 2015, R. Labuda (PRM 952498 – holotype; BiMM-F76 = CCF 6359 – ex-type cultures). ITS sequence GenBank MN633082; LSU sequence GenBank MT874998.

Description: Sexual morph not observed on any of the media used in the present study. Vegetative mycelium consisting of hyaline, smooth-walled, septate, sparsely branched hyphae, 1.5–5.0 μm diam. Racquet hyphae present. Conidia aleuroconidia, hyaline, white in mass, thin-walled, smooth to sparsely irregularly ornamented with minute warts (SEM); terminal and lateral conidia born on main fertile hyphae as sessile or on short protrusions, solitary, 1–3 (–5) per conidiogenous cell, obovate to clavate, 1-celled, (3.0–)4.5–6.5(–7.5) x (1.5–)2.0–2.5(–4.0) μm (mean = 4.9 ± 0.8 x 2.4 ± 0.4 μm, n = 120), and filiform, often sinusoidal, 1- to 2-celled, 25–35(–40) μm long conidia also present. Intercalary conidia (arthroconidia) present, 10–15 μm long. Chlamydospores not observed.
**Culture characteristics:** Colonies on PDA 28–35 mm diam at 25 °C, after 14 d, floccose, with good sporulation, white, flat, slightly elevated (umbonate) at the centre, with irregular margin, reverse lemon yellow, soluble pigment bright yellow, a few small clear to yellow-orange exudate droplets produced. At 30 °C, 38–45 mm diam after 14 d, white, flat, floccose and radially sulcate with good sporulation only at the centre, and with lemon yellow reverse. Colonies on SDA 28–35 mm diam at 25 °C, after 14 d, morphology similar to PDA, without exudate and with pale-yellow reverse. Colonies on MEA 20–25 mm diam at 25 °C after 14 d, morphology similar to PDA, exudate absent, and pale-yellow reverse. At 30 °C, 18–20 mm diam after 14 d, white, floccose and radially sulcate, with good sporulation, and with pale yellow reverse. Colonies on CMA and PCA, 45–50 mm diam at 25 °C, after 21 d, white, flat and spread with poor sporulation, reverse white. No ascomata observed after prolonged incubation (3 months). The optimum temperature on PDA, SDA and MEA 25–30°C (Table S1a–c). Minimum
growth (1–2 mm diam) at 8 °C. The maximum temperature for growth 32 °C (microcolonies to 1 mm diam). Keratinolytic activity very weak (Fig. 10a), with hair attack intensity = 0–1. Urease activity positive (after 3 d of incubation).

Diagnosis: This species molecularly can be distinguished from other Keratinophyton species by ITS locus analysis. Phenotypically, K. lemmensii is unique and differs from the relatives in the same clade based on ITS phylogeny (K. durum, K. hubeiense, K. submersum, and K. siglerae) by the combination of the following features: (1) presence of long filiform often sinusoidal uni- to bicellular conidia (to 40 μm), (2) white, moderately fast growing colonies (28–35 mm diam, on PDA at 25 °C), (3) production of lemon yellow pigment on PDA at 25 °C, (4) minimum 8 °C and maximum 32 °C growth temperature, (5) very weak keratin digestion after 21 d. Presence of filiform often sinusoidal conidia and

Fig. 5 Line drawing of micromorphology of Keratinophyton lemmensii (BIMM-F76). a, b Conidiophores with young and mature aleurioconidia, including arthroconidia on PDA (after 14 d). a Branched conidiophore. b Unbranched conidiophore with sessile aleurioconidia. Bar = 10 μm
abundant arthroconidia, production of bright yellow pigment on PDA and good growth at 30 °C.

Notes: Based on a search of NCBI GenBank nucleotide database, the closest hit for *K. lemmensii* using the ITS sequence was *K. durum* (FMR5651; GenBank acc. AJ439434; identities = 568/577 (98%), gaps 0/577 (0%). However, *K. lemmensii* can be directly distinguished from *K. durum* by its asexual morph also by the presence of numerous arthroconidia which are completely missing in the latter species (Cano and Guarro 1990; Currah 1985).

**Keratinophyton linfenense** (Liang, Liang & Han) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833639

*Basionym*: *Chrysosporium linfenense* Z.Q. Liang, J.D. Liang & Y.F. Han - *Mycotaxon* 110: 67; 2009.

*Type*: GZUXIFR H31, isolated from rhizosphere soil by G. Don, China. For detailed description of the species, see Liang et al. (2009).

**Keratinophyton minutisporosum** (Vidal & Guarro) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833640

*Basionym*: *Chrysosporium minutisporosum* P. Vidal & Guarro - *Stud. Mycol.* 47: 205; 2002.

*Type*: CBS 101577 = IMI 379912 = FMR 6096 isolated from river mouth sediment by P. Vidal, Spain. For detailed description of the species, see Vidal et al. (2002).

**Keratinophyton pannicola** (Corda) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833643

*Basionym*: *Capillaria pannicola* Corda - *Icon. Fung.* 1: 10; 1837.

≡ *Sporotrichum pannicola* (Corda) Rabenh. - *Deutschl. Krypt.-Fl.* 1: 78; 1844.

≡ *Chrysosporium pannicola* (Corda) Oorschot & Stalpers - *Stud. Mycol.* 20: 43; 1980.

*Synonym*: *Trichophyton evolceanui* H.S. Randhawa & R.S. Sandhu - *Mycopath. Mycol. Appl.* 20: 232; 1963.
≡ *Chrysosporium evolceanui* (Randhawa & Sandhu) Garg - *Sabouraudia* 4: 262; 1966.

*Type*: CBS 116.63 = ATCC 22400 = IHEM 4436 = IMI 147545 = NCPF 489 = RV 26475 = UAMH 1275, isolated from soil by Randhawa & Sandhu, India.

**Keratinophyton siglerae** (Cano & Guarro) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833641

*Basionym*: *Chrysosporium siglerae* Cano & Guarro - *Mycotaxon* 51: 75; 1994.

*Type*: UAMH 6541 = FMR 3066 = IMI 336467, isolated from garden soil, Spain. For detailed description of the species, see Cano and Guarro (1994).

**Keratinophyton submersum** (Vidal & Guarro) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833642

*Basionym*: *Chrysosporium submersum* P. Vidal & Guarro - *Stud. Mycol.* 47: 200; 2002.

*Type*: CBS 101575 = IMI 379911 = FMR 6088, isolated from river mouth sediment by P. Vidal, Spain. For detailed description of the species, see Vidal et al. (2002).

**Keratinophyton straussii** Labuda, Bernreiter, Kubátová & Schüller, **sp. nov.**
(Figs. 6 and 7)

*MycoBank*: MB833634

*Etymology*: Named in honour of Joseph Strauss, Head of the Department of Applied Genetics and Cell Biology, founder of the Fungal Genetics and Genomics Laboratory, University of Natural Resources and Life Sciences, Vienna (BOKU), Austria, and an expert in fungal genetics, epigenetics and functional genomics.

*Type*: *Wiste*, from garden soil, Aug. 2015, *R. Labuda* (PRM 952500 – holotype; BiMM-F78 = CCF 6361 – ex-type cultures). ITS sequence, GenBank MN633081; LSU sequences, GenBank MT874996.

*Description*: *Sexual morph* not observed on any of the media used. *Asexual morph* on PDA. Vegetative mycelium of hyaline, septate, smooth-walled, sparsely to pronunciably branched hyphae, usually at right angles, 1.5–4.0 μm diam. *Racquet hyphae* present. *Conidia* (aleurioconidia), hyaline, white to yellowish in mass, thin-walled and regularly ornamented with minute warts (SEM) and coarsely roughened (light microscope). Terminal and lateral conidia born on main fertile hyphae or from side branches of variable length, sessile or on short protrusions, commonly slightly swollen, length variable, solitary, 1–3 (5) per conidiogenous cell, obovate to clavate, 1-celled, (3.5–)4.5–5.0 (–6.5) x (2.0–)2.5–3.0 (–3.5) μm (mean = 4.9 ± 0.4 x 2.6 ± 0.2 μm, n = 120), very rarely 2- to 3-celled, to 12 μm large aaleurioconidia also present. *Intercalary conidia* not observed. *Chlamydospores* not observed.

*Culture characteristics*: Colonies on PDA 24–28 mm diam at 25°C, after 14 d, powdery to downy (mealy), with abundant sporulation, white to very slightly creamy yellowish, flat, slightly elevated (umbonate) remaining powdery at the centre, with irregular margin, reverse white with slightly yellowish centre, no pigment or exudate produced. At 30°C, 15–20 mm diam after 14 d, white to creamy yellowish, flat, powdery to downy (mealy) with very good sporulation, and with white to yellowish reverse. Colonies on SDA 16–20 mm diam at 25°C, after 14 d, morphology as on PDA with dark yellow reverse. In age (after 5 wk) yellow pigment produced and colony reverse becoming bright reddish yellow to orange. At 30°C, 15–20 mm diam after 14 d, white to creamy yellowish, umbonate, with strong sporulation, and with yellowish reverse. Colonies on MEA 18–20 mm diam at 25°C, after 14 d, morphology as on PDA with more floccose and yellowish. At 30°C, 5–10 mm diam
after 14 d, slightly umbonate, floccose to granular, with very good sporulation white to yellowish, and with yellow reverse. Colonies on CMA and PCA 18–20 mm diam at 25 °C, after 21 d, white, granular, good sporulation, reverse yellowish. No ascomata observed after prolonged incubation (3 months). The optimum temperature for growth on PDA, SDA and MEA 20–25 °C (Table S1a–c). Minimum growth (microcolonies to 1–2 mm diam) at 10 °C. No germination of the spores at 8°C. The maximum temperature for growth 32 °C (microcolonies to 1–2 mm diam). Keratinolytic activity very strong (Fig. 10c), with hair degradation intensity = 4. Urease activity negative (after 14 d of incubation).

Diagnosis: Keratinophyton straussii molecularly can be distinguished from other Keratinophyton species by ITS locus analysis. Phenotypically, it can be differentiated by combination of the ability to grow at 30 °C, white to creamy colonies with white to yellowish reverse at 25 °C on PDA and conidia morphology (obovoid to clavate and verrucose) (Table 2).

**Additional material examined:** Italy: Vieste, from garden soil, isolated from different sub-samples, July 2019,
R. Labuda RL-05 ITS sequence, MT898644; LSU sequence, MT898648); ibid., RL-06 (ITS sequence, MT898645; LSU sequence, MT898649).

Notes: Based on a search of the NCBI GenBank nucleotide database, the closest hit for *K. straussii* using the ITS sequence was *K. minutisporosum* (as *Chrysosporium minutisporosum* CBS 101577; GenBank acc. KT155616), with identity = 489/543 (90%) and gaps 10/543 (1%). Two species can be differentiated from each other based on growth rate and colony reverse at 25 °C on PDA (Table 2). Additionally, *K. straussii* differs from *K. wagnerii* by its ability to grow at 30 °C and strong keratinolytic activity. For the morphological differences between *K. gollerae* and *K. straussii*, see under *K. gollerae*. Additional strains RL-05 and RL-06 grew relatively better (to 5 mm larger diam) than the ex-type culture at 30 °C.

*Keratinophyton qinghaiense* (Han, Liang & Liang) Labuda & Bernreiter, **comb. nov.**
MycoBank: MB833655

**Basionym:** *Chrysosporium qinghaiense* Y.F. Han, J.D. Liang & Z.Q. Liang - *Mycosystema* **32:** 607, 2013.
Type: GZAC GZUIFR-Chry 11, from farmland soil by, Y.F. Han, China.

*Keratinophyton wagneri* Labuda, Bernreiter, Kubáňová & Schüller, **sp. nov.**
MycoBank: MB 833635.

Etymology: Named in honour of Martin Wagner, Head of the Unit for Food Microbiology and Head of Institute for Food Safety, Food Technology and Veterinary Public Health, University of Veterinary Medicine, Vienna (Austria), an expert in veterinary microbiology.

Type: Slovak Republic: Tatranská Lomnica, from forest soil, Aug. 2015, R. Labuda (PRM 952501 – holotype; BiMM-F77 = CCF 6362 – ex-type cultures). ITS sequence, GenBank MN633083; LSU sequence, GenBank MT874999.

Description: Sexual morph not observed on any of the media used. Asexual morph on PDA. Vegetative mycelium hyaline, septate, smooth-walled, sparsely to pronouncedly branched hyphae, 2.0–6.0 μm diam. Racquet hyphae present. Conidia (aleurioconidia), hyaline, white to yellowish in mass, thin-walled and regularly ornamented with minute warts (SEM) and coarsely roughened (light microscope). Terminal and lateral conidia born on

Fig. 8 Keratinophyton wagneri (BiMM-F77). a Colonies on PDA (after 14 d) at 20°C, 25°C and 30°C. b Conidiophores with aleurioconidia. c Aleurioconidia (on PDA, after 14 d). d, e Scanning electron microscopy (SEM) of conidiogenous cells and aleurioconidia (on PDA, after 14 d). Bars = 20 μm (b), 10 μm (c), 5 μm (d), 2 μm (e)
main fertile hyphae or from side branches of variable length, sessile or on short protrusions, occasionally swollen and of variable length, solitary, 1–4 (−10) per conidiogenous cell, obovate to clavate, single celled, (4.0–) 5.5–6.5 (−8.0) × (2.5–) 3.0–3.5 (−4.0) μm (mean = 5.7 ± 0.4 × 3.2 ± 0.2 μm, n = 120), rarely 2-celled, up to 12 μm large ones also present. *Intercalary conidia* not observed. *Chlamydospores* not observed.

*Culture characteristics:* Colonies on PDA 25–30 mm diam at 25°C, after 14 d, powdery to downy (mealy), with abundant sporulation, white to slightly yellowish, flat, slightly elevated (umbonate) and more floccose at the centre, margin irregular, reverse white with slightly yellowish centre, no pigment or exudate produced. At 30°C, 4–8 mm diam after 14 d, white, floccose with poor sporulation, and with yellowish reverse. Colonies on

![Fig. 9](image_url) Line drawing of micromorphology of *Keratinophyton wagneri* (BIMM-F77). a, b Conidiophores with young and mature aleurioconidia on PDA (after 14 d). a Branched conidiophore. b Unbranched conidiophore with sessile aleurioconidia. Bar = 10 μm
SDA 14–18 mm diam at 25 °C, after 14 d, morphology similar to PDA. In age, yellowish brown (amber) pigment produced and colony reverse becoming dark reddish brown (after 4 wk). At 30 °C, no growth or only microcolonies. Colonies on MEA 18–22 mm diam at 25 °C, after 14 d, morphology as on PDA but more yellowish. At 30 °C, no growth or only micro-colonies produced. Colonies on CMA and PCA 20–25 mm diam at 25 °C, after 21 d, white to yellowish, granular, good sporulation, reverse yellowish. Pinkish pigment after 3–4 wk. on PCA (in both tested strains). No ascomata observed after prolonged incubation (3 months). The pti-mum temperature for growth on PDA, SDA and MEA 20–25 °C (Table S1a–c). Minimum growth (1–2 mm diam) at 10 °C, and germination of a majority of the conidia at 8 °C. The maximum temperature for growth 31 °C (1–3 mm diam). Keratinolytic activity weak to moderate (Fig. 10d), with hair attack intensity = 2. Urease activity negative (after 14 d of incubation).

Diagnosis: K. wagneri molecularly can be distinguished from other Keratinophyton species by ITS locus analysis. Phenotypically, it can be differentiated by combination of the growth rate at 30 °C and conidia size (4.0–8.0 × 2.5–4.0 μm) and morphology (obovoid to clavate, verrucose) (Table 2).

Additional material examined: Slovak Republic: Tatranská Lomnica, from forest soil, isolated from a different sub-sample, July 2019, R. Labuda, RL-07 (RL; ITS sequence, MT903275; LSU sequence, MT903309).

Notes: Based on a search of the NCBI GenBank nucleotide database, the closest hit for K. wagneri using the ITS sequence was K. minutisporosum (as Chrysosporium minutisporosum CBS 101577; GenBank: KT155616); with identity = 486/541 (90%) and gaps 11/541 (2%). Morphologically, K. wagneri can be separated from C. minutisporosum by its larger conidia (4.0–8.0 × 2.5–4.0 μm vs. 3.0–4.0 × 1.5–3.5 μm) and growth rate at 25 °C on PDA after 14d (25–30 mm vs. 55–70 mm). Keratinophyton straussii and K. wagneri seem to be very similar, however, they can be distinguished by: (1) size of conidia (av. = 4.9 × 2.5 μm vs. 5.7 × 3.2 μm), (2) growth at 30 °C on PDA (15–20 mm vs 3–4 mm diam), (3) morphology of conidiogenous cells (commonly vs. non–to occasionally swollen), (4) colony pigmentation on SDA after prolonged incubation (bright orange vs. dark brown), and (5) keratinolytic ability after 3 wk. (very strong vs moderate). In addition, the production of a pinkish pigment on PCA after 3–4 wk. (at 20 °C and 25 °C) has been observed only in K. wagneri. Moreover, conidia of this species are more coarsely roughed (warty) than those in K. straussii (Fig. 8c–e).

All four new species are readily distinguished from the other taxa in the genus Keratinophyton, based on phenotypical characteristics such as growth at high...
temperature and/or conidial morphology (Cano and Guarro 1990; Cano and Guarro 1994; Currah 1985). The most important species-specific phenotypic distinguishing characteristics are found as morphology of conidia (shape, surface and dimensions) and growth rate at 30 °C after 14 d on PDA.

**KEY TO SPECIES OF KERATINOPHYTON**

This key is modified from that of Cano et al. (2002). The given data for source and origin represent the type strains of the related species.

| Key | Description | Source | Characteristics |
|-----|-------------|--------|-----------------|
| 1   | Ascomata developed |  |  |
| 2   | Ascomata not developed |  |  |
| 3   | Ascospores smaller |  |  |
| 4   | Ascospore with broad equatorial rim |  |  |
| 5   | Ascospore with narrow equatorial rim |  |  |
| 6   | Ascospores discoid; daily growth 3–4 mm at 28 °C on PYE agar and reverse uncoloured; from beach soil, Spain |  |  |
| 7   | No or restricted (<1 cm in diam) growth at 30 °C on PDA; intercalary conidia absent |  |  |
| 8   | Conidia smooth; racquet hyphae present |  |  |
| 9   | Conidia echinulate; obvoid to clavate; colony reverse white at 25 °C on PDA; from soil, India |  |  |
| 10  | Conidia obvoid to ellipsoidal, 2.2–4.3 × 1.6–3.2 μm; reverse yellowish on PDA at 25 °C; from soil under the chicken feeder, China |  |  |
| 11  | Conidia smooth, ellipsoidal or fusiform; colony reverse white to slightly yellowish at 25 °C on PDA; from forest soil, Slovakia |  |  |
| 12  | Conidia smooth, or smooth to verrucose |  |  |
| 13  | Conidia lenticular, 5–6 × 2.5–3.5 μm; pronounced radial ridges at 37 °C on PYE agar and reverse uncoloured; from lawn soil, India |  |  |
| 14  | Conidia smooth; racquet hyphae absent; colony reverse yellowish at 25 °C on PDA; from farmland soil, China |  |  |
| 15  | Conidia smooth; racquet hyphae present; colony reverse lemon yellow at 25 °C on PDA; from compost soil, Austria |  |  |
| 16  | Conidia clavate to cylindrical; racquet hyphae absent; colony reverse orange yellow at 25 °C on PDA; from river sediments, Spain |  |  |
| 17  | Conidia clavate to long-ellipsoidal; colony reverse brown in centre and light yellow in margin at 25 °C on PDA; from plant root soil, China |  |  |
| 18  | Conidia clavate; colony reverse white to slightly yellowish at 25 °C on PDA; from forest soil, Slovakia |  |  |
| 19  | Conidia smooth, or smooth to verrucose |  |  |
| 20  | Conidia obvoid to clavate; colony reverse white with slightly yellowish centre at 25 °C on PDA; from garden soil, Italy |  |  |
| 21  | Conidia larger |  |  |
| 22  | Conidia smaller |  |  |
| 23  | Conidia more than 3 μm wide; colony reverse brown at 25 °C on PYE agar; from soil, India |  |  |
| 24  | Conidia up to 3 μm wide; colony reverse brownish orange at 25 °C on PYE agar; from river sediments, Spain |  |  |
| 25  | Conidia pyriform to oval, 5–7 × 3.5–5 μm |  |  |
| 26  | Conidia larger |  |  |
| 27  | Conidia clavate to cylindrical; racquet hyphae absent; colony reverse yellowish at 25 °C on PDA; from farmland soil, China |  |  |
| 28  | Conidia obovoid to clavate; colony reverse white to slightly yellowish at 25 °C on PDA; from forest soil, Slovakia |  |  |
| 29  | Conidia smooth, or smooth to verrucose |  |  |
| 30  | Conidia obvoid to clavate; colony reverse white at 25 °C on PDA; from river sediments, Spain |  |  |

**DISCUSSION**

**Phylogeney**

Phylogenetic reconstruction using ITS sequences resulted in clustering of a new species, Keratinophyton lemensis, with *K. durum* (as Aphanoascus durus; Cano and Guarro 1990), *K. hubeiense* (as Chrysosporium hubeiense; Zhang et al. 2016) and *K. submersum* (as Chrysosporium submersum; Vidal et al. 2002), and forming a sister clade to *K. siglerae* (as Chrysosporium siglerae; Cano and Guarro 1994). The other three novel species, *K. gollerae*, *K. straussii*, and *K. wagneri*, were resolved in a separate terminal clade (Fig. 1a). Its sister clade encompasses *K. clavisporum* (as Chrysosporium
clavisporum; Zhang et al. 2017), K. quinghaense (as Chrysosporium quinghaense; Han et al. 2013), K. linfenense (as Chrysosporium linfenense; Liang et al. 2009), and K. turgidum (Sharma and Shouche 2017). Based on the phylogeny and as a result of the abandoning of separate names for morphs of the same fungus (May et al. 2019), species previously described in Chrysosporium require redispersing in the genus Keratinophyton. In our study we confirmed ten species required transfer. The monophyletic genus Keratinophyton is now extended and includes 25 species including ten species known from sexual morphs (Sutton et al. 2013; and this paper) and 15 species which are currently known only from asexual morphs (including the recently described K. turgidum (Sharma and Shouche 2017). The species known only from the asexual morphs can be distinguished by particular combinations of their morphological traits (colony colour and growth rate, growth response at higher/lower temperatures, as well as morphology of conidia) and differences in the ITS regions (Fig. 1a, Table 2).

Ecology and distribution
Almost all known Keratinophyton species have been isolated from soil or soil-like substrates, such as river sediments, compost and sand (Table 1; Cano and Guarro 1990; Sharma and Shouche 2017; Labuda et al. 2008; Liang et al. 2009; van Oorschot 1980; Vidal et al. 2000; Vidal et al. 2002). Hubalek (2000) provided a list of keratinolytic fungi associated with free-living mammals and birds of which Keratinophyton pannicola (as Chrysosporium evolceanui) has been isolated from a variety of animals, different species of rodents in Australia, Czechia, Germany, the UK, and the former Yugoslavia; a rabbit in Canada; and from birds in Australia (Queensland), Czechia, India, and the former Yugoslavia. Keratinophyton durum (as Aphanoascus durus) has been isolated from a hedgehog in Ivory Coast, and K. terreum (as Aphanoascus terreus) has been found associated with a variety of rodents in Czechia, Germany, India, Nigeria, Romania, and the former Yugoslavia and, and further birds in Australia (Queensland) Czechia, India, the USA, and the former Yugoslavia (Hubalek 2000). To the best of our knowledge, there is only a single report of a human clinical isolate belongs to K. echinulatum (CCF 4652 = CBS 141178) from the sole of the foot of a 35-year-old woman in Czechia (Hubka et al. 2016). However, Hubka et al. (2016) indicated that the etiological significance of this fungus was unclear, and they concluded that the infection was actually caused by another dermatophyte, which was not isolated or was overgrown by K. echinulatum. A few other cases have been published in a small range of animals including Keratinophyton pannicola (as Chrysosporium pannicola) from skin of a dog in former Yugoslavia (Hajsig et al. 1974; van Oorschot 1980) and from a case of keratomycosis in a horse (Grahn et al. 1993).

In her review on Chrysosporium and related genera in Onygenaceae, Sigler (2003) stated that some reports concerning Chrysosporium species as etiological agents must be viewed with caution, in case the isolated fungus has neither been identified to species level nor documented well enough to confirm the aetiology. In the follow-up list of medically relevant species provided by Sigler (2003), no species is mentioned as being currently affiliated within the genus Keratinophyton, while K. pannicola (as C. pannicola) is included in the Atlas of Clinical Fungi (de Hoog et al. 2020) as a concern in skin infections. Even though the keratinophilic fungi were considered as potential pathogens by several researchers (Rippon 1982; Papini et al. 1998); they rarely cause infections. Therefore, soil is proposed as an epidemiological and probably also an evolutionary link, that relates geophilic, zoophilic, and anthropophilic keratinophilic fungi (Papini et al. 1998). Interestingly, during a mycological investigation of the soil samples in the present study, a high prevalence of geophilic dermatophytes such as Nannizzia gypsea from Italy (collected in 2004), a co-occurrence of Arthroderma uncinatum with Aphanoascus keratinophilus (as Chrysosporium keratinophilum) from the Slovak Republic (collected in 2011), and Arthroderma terrestrre along with abundant A. uncinatum from Austria (collected in 2015) were noted (data not shown).

As the members Keratinophyton are considered as typical soil-borne fungi (Cano and Guarro 1990; Cano et al. 2002; Sutton et al. 2013) and there is no solid evidence of pathogenicity, it is likely that previously reported animal-associated cases reflect environmental transmissions from soil to the animals during activities in contact with soil. The ability of these fungi to persist and survive in the soil was observed also during the present study, as in case of K. straussii, the type strain was isolated 11 years after sampling in 2004, and two more strains (RL-05 and RL-06) representing the same taxon were isolated in a repeated study even 15 years after the sampling. Likewise, a second strain (RL-07) used for the description of K. wagneri and the type of K. gollerae (BiMM-F250) were both isolated 8 years after the samples were collected.

The degree of keratin degradation by the novel strains described here varied. It was very strong in both K. gollerae and K. straussii compared to other tested strains, attacking the cuticle and cortex of hairs with about 50–80% degradation. In addition to keratin degradation, keratinolytic fungi share common properties with dermatophytes (Marchisio et al. 1994; Mitola et al. 2002). Even though some of these fungi can
grow at 37 °C (Fig. 1a), potential pathogenicity to homeothermic vertebrates (mammals and birds) by these fungi seems highly unlikely because of some presumably missing pathways in their metabolism. Instead, their strong keratolytic ability might be providing a competitive advantage in the nature to acquire nutrients from hair and may have potential in industry for the production of proteolytic enzymes to degrade keratinous materials (hairs, fur, feathers, etc.). Furthermore, these fungi represent a yet unexplored possible source of new bioactive compounds as there is not much known of these properties in the genus (Kushwaha and Guarro 2000).

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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