Phylogeny and taxonomic revision of *Kernia* and *Acaulium*

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The genera *Kernia* and *Acaulium* comprise species commonly isolated from dung, soil, decaying meat and skin of animal. The taxonomy of these fungi has been controversial and relies mainly on morphological criteria. With the aim to clarify the taxonomy and phylogeny of these fungi, we studied all the available ex-type strains of a large set of species by means of morphological and molecular phylogenetic analyses. Phylogenetic analysis of the partial internal transcribed spacer region (ITS) and the partial 28S rDNA (LSU) showed that the genera *Kernia* and *Acaulium* were found to be separated in two distinct lineages in Microascaceae. Based on morphological characters and multilocus phylogenetic analysis of the ITS, LSU, translation elongation factor 1α and β-tubulin genes, the species in *Kernia* and *Acaulium* were well separated and two new combinations are introduced, i.e. *Acaulium peruvianum* and *Acaulium retardatum*, a new species of *Kernia* is described, namely *Kernia anthracina*. Descriptions of the phenotypic features and molecular phylogeny for identification are discussed for accepted species in two genera in this study.

*Kernia* was erected by Nieuwland¹ for a group of fungi with cleistothecial, with *Kernia nitida* (Saccardo) Nieuwland as type species firstly and subsequent species have all been characterized by fascicled hair-like ascocarp appendages and reddish-brown to brown ascospores. In 1971, the ascomycete genus *Kernia* is emended to revised concepts by Malloch². Five species, *K. bifurcotricha* Saxena & Mukerji, *K. hippocrepida* Malloch & Cain, *K. nitida* (Sacc.) Nieuwland, *K. hyalina* Malloch & Cain and *K. pachypleura* (Ames) Benjamin and *K. geniculotricha* Seth, are placed in synonymy with *K. nitida*. Besides, *K. bartlettii* (Massee & Salmon) Benjamin, *K. furcotricha* Tandon & Bilgrami, and *K. spirotricha* Benjamin, are excluded from *Kernia*. And some new species and combinations were added to it later as *K. ovata* (Booth) Malloch & Cain, *K. fintara* Udagawa & T. Muroi, *K. setadispersa*, *K. cauquensis*, *K. irregularis*, *K. peruviana* Udagawa & Furuya, *K. columnaris* (H.J. Swart) Woudenberg & Samson³–¹⁰. In recently, *K. hyalina* is excluded from *Kernia* by phylogenetic analysis based on a combined LSU and ITS sequence dataset and morphological characteristics¹¹. Although 11 species are accepted in *Kernia*, many described species are of doubtful identity because their type materials are lost and their protologues are uninterpretable.

The genus *Acaulium* was established as the sexual morph and the type species is *Acaulium albonigrescens* Sopp¹², and this genus was considered as congeneric with *Microascus*¹³–¹⁵. *Acaulium* is characterised by anamelic conidiogenesis, guttulate conidia and mycelium forming abundant hyphal fascicles and has generally been considered a synonym of *Scopulariopsis* but recently was re-instated as an accepted genus of Microascaceae with three species as *A. acremonium* (Delacr.) Sandoval-Denis, Guarro & Gené, *A. albonigrescens* Sopp, Skr. Vidensk.-Selsk. and *A. caviariforme* (Malloch & Hubart) Sandoval-Denis, Guarro & Gené. *Acaulium albonigrescens*, formerly known as *Doratomyces putredinis*, is transferred to *Acaulium* and redescribed by Woudenberg¹⁰ based on morphological, physiological and molecular phylogenetic analyses. In addition, *A. pannemaniae* Sandoval-Denis is introduced in this genus by morphological and phylogenetic analyses of LSU¹⁷. Five species are currently accepted at present¹⁰.

*Kernia* currently comprises species that are commonly isolated from the dung of animal¹⁰, ¹²–¹⁶, except two species *K. retardata* and *K. peruviana*¹⁰, ¹⁷, which isolated from soil. *Acaulium* species have been reported from a variety of environments such as skin of a horse, decaying meat, soil and so on¹⁰, ¹⁶. From beginning, the genera of *Kernia* and *Acaulium* have been controversial and rely mainly on morphological criteria. Recent molecular

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Table 1. Strains and sequence accession numbers included in this study. CBS: CBS Fungal Biodiversity Centre, Utrecht, The Netherlands. Sequences newly generated in this study are indicated in bold. ‘T’ represents type strain.

| Species | Strain no. | Isolation source | Location | GenBank accession no. |
|---------|------------|------------------|----------|-----------------------|
|         |            |                  |          | ITS TUB LSU TEF       |
| Acaulium acremonium | CBS 290.38T | Skin of a horse | Denmark  | LM652456 LN851108 LN851001 HG380362 |
| Acaulium albogriseus | CBS 109.69T | Litter, treated | Japan    | KY852469 LN851111 KY852480 LN851058 |
| Acaulium albnum | CBS 539.85T | Hair in dung in pole cat | Netherlands | MN991960 MN982419 MN991968 MN982411 |
| Acaulium caviariforme | CBS 536.87T | Decaying meat | Belgium  | LM652392 LN851112 LN851005 LN851059 |
| Acaulium pansennanum | CBS 145025T | Soil | Netherlands | LS999990 LS999993 LS999991 LS999992 |
| Acaulium perverum | CBS 320.91T | Soil | Peru     | MN991959 MN982418 MN991966 – |
| Acaulium retardatum | CBS 707.82T | From paddy soil | Japan    | MN991961 – MN991969 MN982412 |
| Cephalotrichus asperulum | CBS 582.71T | Soil | Argentina | LN850960 LN851114 LN851007 LN851061 |
| Cephalotrichus brevistipitatum | CBS 157.57T | Solanum tuberosum | Netherlands | LN850984 LN851138 LN851031 LN851084 |
| Cephalotrichus dendrocephalum | CBS 528.85T | Cultivated soil | Iraq     | LN850966 LN851120 NG_059041 LN851041 |
| Cephalotrichus microsporum | CBS 523.63T | Wheat field soil | Germany  | LN850967 LN851121 LN851014 LN851068 |
| Gamsia columbina | CBS 233.66T | Sandy soil | Germany  | LN850990 LN851147 LN851039 LN851092 |
| Graphium penicillioides | CBS 102632T | Populus nigra | Czech Republic | KY825474 – KY825485 – |
| Kernia anthracina | CGMCC 3.19001T | Dung of marmot | Beijing, China | MK773539 MK773545 MK773542 MK773568 |
| Kernia anthracina | CGMCC 3.19002 | Dung of marmot | Beijing, China | MK773540 MK773546 MK773543 MK773569 |
| Kernia anthracina | CGMCC 3.19003 | Dung of marmot | Beijing, China | MK773541 MK773547 MK773544 MK773570 |
| Kernia columnaris | CBS 159.66T | Dung of hare | South Africa | MN991957 MN982416 MN991962 MN982409 |
| Kernia geniculosatrica | CBS 599.68T | On dung of Orezytolagus canculus | Germany  | MN991956 MN982414 MN991964 MN982408 |
| Kernia hippocrepida | CBS 774.70T | On dung of Erethizon dorsata | Ontario, Canada | MN991954 MN982413 – MN982406 |
| Kernia nitida | CBS 282.52T | Chrysolina sanguinolenta | France  | MN991955 MN982415 MN991963 MN982407 |
| Kernia pachyleura | CBS 776.70T | Dung of Loxodonta africana | Uganda  | MN991958 MN982417 MN991965 MN982410 |
| Microascus cirratus | CBS 217.31T | Leaf of Prunus sp. | Italy     | KX923838 – AF400860 – |
| Microascus longirostris | CBS 196.61T | Wasp's nest | USA: Maine | LM652421 LM652634 LN851043 LN652566 |
| Microascus senegalensis | CBS 277.74T | Mangrove soil | Senegal  | KS923929 – AF400867 – |
| Petriella massapora | CBS 745.69 | On rotten wood of Populus grandidentata | Ontario, Canada | MH1859407 – AF027663 – |
| Petriella setifera | CBS 390.75 | Skin lesion in Turpisops truncatus | Netherlands | AY882355 – AF027664 – |
| Petriellopsis africana | CBS 311.72T | Brown sandy soil | Namibia  | AJ888425 – EF151331 – |
| Pseudallescheria ellipsospora | CBS 418.73T | Soil | Tajikistan | EF151323 MH217617 AF027671 – |
| Wardomyces anomalus | CBS 299.61T | Air cell of egg | Canada: Ontario | LN850992 LN851149 LN851044 LN851095 |
| Wardomyces inflatus | CBS 367.62T | Greenhouse soil | Belgium  | LN850994 LN851153 AF400886 LN851099 |
| Wardomyces pulvinatus | CBS 112.65T | Salt-marsh | England, UK | LN850997 LN851156 LN851051 LN851102 |

studies have demonstrated that the Microascaceae contains several closely related genera and difficult to separate morphologically16. Multilocus phylogenetic analysis have considerably improved our understanding of species concepts in many fungal groups18–25, but the studies for revising the genera of Kernia and Acaulium are relatively limited. Besides, during our investigation of intestinal fungi in animals in China, three particular Kernia isolates from the dung of marmot were isolated. The present work also aims to clarify the taxonomic position of these strains as putative new species using the genealogical concordance analysis24. We provide a multigene (ITS, LSU, TEF, TUB) phylogeny of Kernia and Acaulium and related fungi based on a large set of strains, which includes all available ex-type cultures and well-identified reference strains from international culture collections.

Results

Generic circumscription. DNA sequences determined in this study are deposited in GenBank, and accession numbers are listed in Table 1. To delineate generic boundaries, we conducted a phylogenetic analysis using the combined LSU and ITS datasets including 29 currently accepted species belonging to nine genera of Microascaceae and one species of the family Graphiaceae. Graphium penicillioides were selected as outgroup (Fig. 1). The final alignment consisted of 31 strains and contained 1,385 characters (LSU 796, ITS 589). Figure 1 shows the ML tree including ML bootstrap values (bs) and posterior probabilities (pp) values. The trees obtained from ML and Bayesian analyses of the individual loci and the combined analysis showed congruent topologies. The phylogenetic inferences (Fig. 1) showed that Kernia and Acaulium were monophyletic, the species of Kernia and Acaulium clustered into a single, well-supported lineage (bs = 100%/pp = 100%), respectively. Figure 2 is demonstrated that the colonies of K. peruviana CBS 320.91, K. retardata CBS 707.82 and A. albnum CBS 539.85 can form white to pale grey colonies with dense hyphal fascicles. Therefore, K. peruviana and K. retardata were identified as the Acaulium species in this study. However, other type Kernia species grow slowly and form compact, brown
Figure 1. Maximum likelihood (ML) tree obtained from the combined LSU and ITS sequences of 31 representative taxa of Microascaceae and Graphiaceae. Numbers on the branches are ML bootstrap values (bs) above 75%, followed by Bayesian posterior probabilities (pp) above 95%. A dash (–) indicates support value lower than 75% bs or 95% pp. Branch lengths are proportional to distance. Ex-type strains are indicated with T. The tree was rooted to Graphium penicillioides (CBS 102632).

Figure 2. The colony morphology of the species in Kernia and Acaulium was growing on PDA, CMA and OA after 35 days, respectively.
to dark brown colonies apparently different with the *Acaulium* species. Among with the *K. geniculotricha* and *K. nitida* were located in the same clade with the value of (bs = 94%/pp = 91%) and combined with morphological characters, they could be identified as the same species in this research. Three *Kernia* strains which were isolated from the dung of marmot, were clustered with the species of *K. hippocrepida* and have a well-supported value (bs = 100%/pp = 100%).

**Phylogeny of two genera and genealogical concordance analysis.** The second dataset was composed of 19 taxa (including the outgroup) with the following four loci combined: *ITS* (1–434), *LSU* (435–1,163), *TEF* (1,164–2,024), *TUB* (2,025–2,401). The phylogenetic tree was constructed and the branch support values (≥ 50%) and the Bayesian posterior probabilities (≥ 95%) were indicated (Fig. 3). The phylogenetic tree grouped 19 strains into three clades comprising *Kernia* (bs = 100%/pp = 100%) and *Acaulium* (bs = 98%/pp = 100%) subclades with high bootstrap values. Coupled with morphological characteristics (Fig. 2), two new combination species *A. peruvianum* and *A. retardatum* are proposed. Besides, Three *Kernia* strains were clustered with the species of *K. hippocrepida* in independent group and have a well-supported value (bs = 100%/pp = 100%).

BLAST searches of GenBank using the *ITS* sequences of three *Kernia* strains isolated from the dung of marmot revealed that three strains showed 94.7% similarity to *K. nitida* CBS 282.52T (KY852476). Multiple sequence alignment and sequence polymorphism analysis were further carried out in the *ITS* region. Compared with *K. hippocrepida* CBS 774.70T, had only two variable positions that is transition and indel in the *ITS* region. However, sequence polymorphism analysis in the *TUB* region indicated that strain CGMCC 3.19001T have 35 variable positions, including 12 transitions, 7 transversions, and 16 indels, indicating a lower similarity (93.2%) to *K. hippocrepida* CBS 774.70T (Supplementary Table 1). In genealogical concordance analysis, *K. hippocrepida* and *K. anthracina* strains were divided into different statistically supported subclades in *TUB* and *TEF* tree with a high bootstrap value (bs = 100%/pp = 100%) (Fig. 4). In addition, *K. geniculotricha* CBS 599.68T and *K. nitida* CBS 282.52T were clustered together and have a high similarity (99%) from four different gene tree.

**Taxonomy.** Based on the results of the above multilocus sequence analysis and a morphological analysis, the species of the genera *Acaulium* and *Kernia* have been reassessed accordingly. Their current circumscription is revised and several new taxa and combinations are proposed as follows:

*Acaulium peruvianum* (Udagawa & Furuya) L. Su *comb. nov.* Fig. 5.

MycoBank: MB 834193.

*Basionym:* *Kernia peruviana* Udagawa & Furuya, *Mycotaxon* 33: 295. 1988.

*Hyphae* hyaline to subhyaline, smooth-walled, 1–4 μm (x̅ = 2.7 μm) wide. *Conidiophores* often arising from the substratum or from the aerial mycelium, branched or unbranched, septate, smooth, cylindrical, 9–25 × 2–4 μm (x̅ = 14.0 × 3.0 μm). *Conidiogenous cells* solitary or more commonly united into synnemata, percurrent in
Conidiophores or produced on hyphae in laterally, flask-shaped, subhyaline and smooth-walled, 6–11 × 2–3.5 μm (x̅ = 8.9 × 2.7 μm). Conidia ellipsoidal to fusiform, with a truncate base and rounded or bluntly pointed apex, subhyaline, smooth and slightly thick-walled, 3.5–7 × 1–3 μm (x̅ = 5.0 × 2.2 μm). Sexual morph observed. Cleistotheccia superficial, non-ostiolate, dark brown to black, globose, 119–160 μm (x̅ = 143.9 μm) diam., glabrous at maturity except for a few hyphal attachments. Asci 8-spored, globose to ovoid, evanescent. Ascospores irregularly arranged, pale yellowish brown to brown, broadly ovoid to fusiform, 3–5 × 2–4 μm (x̅ = 4.0 × 2.8 μm).

Colonies on PDA reaching 17 mm diameter after 10 days at 20 °C, planar, finely felty with tufts of mycelium in center, white to cream. On SDA reaching 21 mm diameter, planar, finely felty with tufts of mycelium in center, white to cream. On CMA reaching 16 mm diameter, planar, subhyaline. On OA reaching 15 mm diameter, planar to low convex, white to creamcoloured centre, margin discrete.

Specimens examined. Peru, Tamshiyacu, near Iquitos, T. Akiyama, from soil, 1987, S. Udagawa (culture ex-type CBS 320.91 = NHL 2,985).

Notes. This species was originally placed in *Kernia* based on morphological features of the well developed sexual morph. In our phylogenetic analysis, the ex-type culture of *Acaulium peruvianum* grouped with high statistical support with species of *A. album*. *A. peruvianum* is morphologically different with *A. album*, *A. album*,
Acaulium peruvianum produces sexual and asexual morphs in culture, besides, most of conidiogenous cells directly produced from hyphae. However, A. album has abundant monoverticillate, irregularly biverticillate and terverticillate, or reduced to single conidiogenous cells. Conidiogenous cells of A. album are smaller (6–8.5 × 2.5–3 μm) than A. peruvianum.

Acaulium retardatum (Udagawa & T. Muroi) L. Su comb. nov.

MycoBank: MB 834194.

Basionym: Kernia retardata Udagawa & T. Muroi, Trans. Mycol. Soc. 22(1): 18. 1981.

Hyphae hyaline to subhyaline, thin- and smooth-walled, 1–4 μm (x̄ = 2.6 μm) wide. Conidiophores branched or unbranched, septate, cylindrical. Conidiogenous cells flask-shaped to nearly cylindrical, subhyaline and smooth-walled, terminal or lateral in hyphae or hyphae coil, 8.5–23 × 2–5 μm (x̄ = 13.3 × 3.0 μm). Conidia ellipsoidal to fusiform, with a truncate base, subhyaline, smooth and slightly thick-walled, 4–10.5 × 3–6 μm (x̄ = 7.1 × 4.9 μm). Sexual morph observed. Cleistothecia superficial, non-ostiolate, dark brown to black, globose, 106.5–154 μm (x̄ = 129.1 μm) diam., glabrous at maturity except for a few hyphal attachments. Asci 8-spored, globose to subglobose, evanescent. Ascospores irregularly arranged, grey to pale yellow, broadly ovoid to ellipsoidal, 4–8 × 3–5 μm (x̄ = 5.6 × 3.7 μm).

Colonies on PDA reaching 5 mm diameter after 10 days at 20 °C, slow growing, raised centrally, with flat and irregular margin, white. On SDA reaching 9 mm diameter, moderately growing, raised centrally, aerial mycelium absent or sparse, white to cream. On CMA reaching 9 mm diameter, moderately growing, planar, white, margin discrete. On OA reaching 15 mm diameter, planar, white.

Specimens examined. Japan, Nishinasuno-machi, Nasu-gun, Tochigi, Udagawa, S, from rice-field soil, 1988, S. Udagawa (culture ex-type CBS 707.82 = NHL 2,879).

Notes. This species was originally placed in Kernia based on sexual morphological features. The phylogenetic analysis shows that the ex-type culture of Acaulium retardatum grouped with statistical support with species of
A. *albonigrescens* and *A. caviariforme*. *A. retardatum* is morphologically similar to *A. caviariforme*; both species produce sexual and asexual morphs in culture. However, *A. caviariforme* has fusiform, pale orange to copper-red ascospores, and brown, obvoid to ellipsoidal conidia; ascospores of *A. retardatum* are smaller, broadly ovoid to ellipsoidal.

**Kernia anthracina** L. Su, H. Zhu & C. Qin, sp. nov. Fig. 7.  
MycoBank: MB 830661. 
Etymology: Referring to the coal-black colony.  
Holotype: HMAS 255463.  
*Hyphae* septate, branched, catenate, hyaline to subhyaline, mostly 2–4.5 μm (x̅ = 3.1 μm) wide. *Conidiophores*, with scopulariopsis-like branching pattern, produce acrospores. *Conidia* formed in slimy heads at the apex of the scopulariopsis-like branch, broadly clavate to ellipsoid with a slightly apiculate base, smooth to finely roughened, 6.5–14 × 1–5 μm (x̅ = 10.8 × 3.2 μm). *Cleistothecia* abundant in CMA, gregarious, superficial, non-ostiolate, glabrous at maturity, black, globose to subglobose, 55–106 μm (x̅ = 77.9 μm) diameter; peridium with a textura intricata. *Asci* ampulliform. *Ascospores* irregularly arranged, pale yellowish brown to straw coloured, ovoid to fusiform or ellipsoidal, 5–8 × 3–6 μm (x̅ = 6.9 × 4.8 μm). Optimal growth temperatures are 25–30 °C, no growth at 40 °C.  
Colonies on PDA reaching 6 mm diameter after 20 days at 30 °C, 4 mm on SDA, 5 mm on CMA and 13 mm on OA. Colonies on PDA, black in obverse, compact, reverse black, raised centrally, aerial mycelium absent or sparse.  
Specimens examined. China, Beijing, Fangshan District, in north center for experimental animal resources, Institute of medical laboratory animal science, Chinese academy of medical sciences, 116°13′ E, 39°48′ N, 58 m above sea level, from fresh dung samples of healthy *Marmota monax*, 7 December 2017, collected and isolated by L. Su (HOLOTYPE: HMAS 255463, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China; dried culture of ex-type CGMCC 3.19001T on PDA), living cultures, CGMCC 3.19001, CGMCC 3.19002, CGMCC 3.19003.

Notes. Strains of *K. anthracina* have the typical features of *Kernia* such as compact growth, nonostiolate, ascospores, and ovoid to ellipsoidal ascospores. *K. anthracina* not only has a scopulariopsis-like asexual morph, but is also supported by phylogenetic tree based on the combined four genes dataset (Fig. 3) and genealogical concordance analysis (Fig. 4). Although *K. anthracina* is closed to *K. hippocrepida* CBS 774.70 with ITS sequences, they apparently differs from morphological characters as *K. hippocrepida* produced reniform ascospores and conidiophores produced from coiled or irregularly twisted (Fig. 8A–D) while *K. anthracina* production of scopulariopsis-like conidiophores and ovoid to ellipsoidal ascospores.

**Kernia nitida** (Saccardo) Nieuwland. Amer. Midland Natur. 4: 379. 1916. Figure 8E–J.  
*Basionym*: *Magnisia nitida* Saccardo. Michelia, 1: 123. 1878.  
*Synonym*: *Kernia geniculotricha* Seth. Acta Bot. Neerl. 17: 481. 1968.  
Description and illustrations: Seth (1968).

Specimens examined. Germany, near Hamburg, dung of rabbit, 1968, H.K. Seth (culture ex-type CBS 599.68 = ATCC 18529).

Notes. Although Malloch and Cain placed the species of *K. geniculotricha* as the synonym of *K. nitida* only based on numerous drawings and aquarels which are thus proposed, the isolate CBS 599.68 studied and proposed here conforms to the morphological characteristics of descriptions and phylogeny of *K. nitida*. The isolate of CBS 599.68 forms compact colonies on PDA, simple or branched, hyaline to light brown conidiophores, and bearing a cluster of annellophores directly at the apex or repeatedly and compactly branching to form a dense penicillus, 10–20.5 × 1–3.5 μm (x̅ = 14.4 × 2.0 μm), produced in clusters of two to three at the tips of the conidiophores or metulae, rarely solitary, flaskshaped to nearly cylindrical, 7–15 × 1–4 μm (x̅ = 9.7 × 2.5 μm); conidia...
ovoid to ellipsoidal, 3.65 ± 2.5–4.5 μm (x̄ ± 5.1 × 3.2 μm). Besides, The isolate has a sexual morph characterised by abundant cleistothecia on CMA, gregarious, superficial, non-ostiolate, black, opaque, ovoid, 143–323.5 μm (x̄ = 203.9 μm) diam., hairs emerging as two opposing or triangle symmetrical on the cleistothecium, dark brown to black. Asci 8–spored, globose to ampulliform, 8–18 μm (x̄ = 11.2 μm) diam., evanescent. Ascospores irregularly arranged, pale brown, smooth, broadly ovoid to globose, 3.5–6 × 2.5–5.5 μm (x̄ = 4.9 × 3.6 μm). In addition, *K. geniculotricha* is a well-circumscribed species described from on dung of Oryctolagus cuniculus in Germany. All these characters are similar to the species of *K. nitida*. Combined with phylogeny and genealogical concordance analysis, we identified the *K. geniculotricha* as the synonym of *K. nitida*.

**Identification keys.** According to the morphological features, identification keys were constructed for the different genera including all the phylogenetic species recognised in this study (Supplementary information 1).

**Discussion**

The family Microascaceae was established by Luttrell, comprising saprobic and plant pathogenic species. Some species of Microascaceae are opportunistic pathogens and show intrinsic resistance to antifungal agents. Recent molecular studies have demonstrated that the Microascaceae contains several closely related genera that are difficult to separate morphologically including *Microascus*, *Scedosporium* and *Scopulariopsis*. Recently, three of the most debated genera of the family, *Microascus*, *Scopulariopsis* and *Pithoasces* were revised by morphology and multigenie phylogeny. As a result, several taxa were excluded from these genera and appeared as a new lineage within the Microascaceae as *Acaulium*.

In this study we have reviewed the taxonomic circumscription of species in the genera of *Kernia* and *Acaulium*, traditionally referred to as sexual and asexual morphs, respectively, and two genera using a polyphasic approach based on the genealogical concordance analysis, phylogeny and morphological data. These results show that *Kernia* and *Acaulium* constitute two phylogenetically distant lineages, combining the results of phenotypic data, delineate the accepted species of the two new combination species, proposing a new species, which are clarifying the identity of species as *Kernia nitida* and *K. geniculotricha*, reclassifying the white synnematous species as *Acaulium peruvianum* and *A. retardatum*, and describing a new species *K. anthracina* isolated from the dung of marmot. The species of *Kernia* are mainly isolated from dung of various animals, while the species of *Acaulium* have a worldwide distribution and are mainly isolated from dung, litter, soil, skin of a horse and decaying meat.

Sandoval-Denis et al. was firstly attempts to clarify phylogenetically the relationships among the different genera of the Microascaceae by the use of partial LSU and ITS sequences. Subsequently, Microascaceae was revised by Sandoval-Denis et al. based on morphological, physiological and molecular phylogenetic analyses using DNA sequence data of four loci (ITS, LSU, TEF and TUB). These studies demonstrated that several genera of Microascaceae raised questions concerning correct positions of several members of the family and their generic circumscriptions, suggesting a possible subdivision of *Microascus* and *Scopulariopsis* into several smaller genera as *Kernia* and *Acaulium*. Our results based on the phylogenetic reconstructions of two loci (LSU, ITS) indicated that *Kernia* and *Acaulium* fall into two groups (Fig. 1). Besides, it also shows that *Kernia* and *Acaulium* species can be well separate by phylogenetic analysis of four loci (LSU, ITS, TEF and TUB). As known that *Acaulium* is characterised by the formation of pale colonies with dense hyphal fascicles and the presence of abundant oil drops in the mycelium, conidia and ascospores, showing a guttulate appearance. The new combination species *A. peruvianum* and *A. retardatum* clustered in *Acaulium* group, in which the species produce white and pale grey colonies, and have a wide isolation source. A new species was clustered in *Kernia* group, which form compact dark brown or black colonies, and mainly isolated from dung (Figs. 2, 3). In addition, species delineation was also assessed in the genus of *Kernia* as the closed species of *K. anthracina* and *K. hippocrepida* under the genealogical concordance analysis using DNA sequence data of four loci (Fig. 4).

The absence of clear diagnostic morphological characters can be used to identify species which belonging to the *Kernia* and *Acaulium* species. The species of *K. geniculotricha* and *K. nitida* have been identified two different species according to morphologically characters, but molecular data can easily identified them as the same species, using any of the four genes studied here. Some species as *A. peruvianum* and *A. retardatum* isolates were initially identified as *K. peruvianum*, *K. retardate* at CBS based on their morphology. Combined the molecular data, the group of species that would previously have been included in *Kernia* and *Acaulium* are easily recognized. Our phylogeny demonstrates that, although *Kernia* and *Acaulium* share similar morphological and ecological traits, they are in fact genetically distant. The phylogenetic data is supported by relevant morphological differences, such as the color of colonies, the shape of ascospores or conspicuously hairy ascomata.

The new species, *K. anthracina* and *K. hippocrepida*, are very similar in ITS but easily distinguished by TUB and TEF sequences. All *Kernia* species can be well separated with TUB and TEF partial gene sequences. Based on ITS alone, *K. anthracina* and *K. hippocrepida* cannot be distinguished (Fig. 4), but morphology and TUB and TEF sequences clearly differentiate them. The lack of the isotype herbarium specimens examined here prevented us from conclusively characterizing live of the other described species *K. bifurcotrecha*, *K. setadispersa*, *K. caquenesis*, *K. irregularis*, *K. oviata*, leaving them as nomina dubia. From our study, we found that it is easy to identify the species of *Kernia* and *Acaulium* by polyphasic approach.

The delimitation of the two genera in this study contributes to an integrated phylogeny of the family Microascales. The two monophyletic genera currently accepted are statistically supported in the four-locus phylogeny (Fig. 3). There are seven species included in *Acaulium* by our revision, while ten species in *Kernia*. It is regret that some species of *Kernia* absent holotype material and unavailable for these species. Therefore, further studies are needed to establish a comprehensive modern classification of the *Kernia* and to give better insight into the evolutionary relationships among the species in the genus.
Materials and methods

Eight Kernia and Acaulium ex-type strains were obtained from the CBS culture collection (CBS) housed at the Westerdijk Fungal Biodiversity Institute (WI), Utrecht, the Netherlands. More isolates potentially related to the obtained Acaulium strains were selected based on a preliminary phylogenetic analysis of LSU + ITS sequences from GenBank, as well as several cultures of the Kernia, which isolated from the feces of Marmota monax\(^2\), maintained in China General Microbiological Culture Collection Center (CGMCC) in China. All the strains used in this study are listed in Table 1. The strains were incubated on different media such as Potato dextrose agar (PDA), Malt extract agar (MEA), Sabouraud Dextrose Agar (SDA), Corn meal agar (CMA), and Oatmeal agar (OA) (Becton, Dickinson & Co.) at 20 °C. Colony morphology and microscopic characteristics were examined, measured and photographed after incubation for 10 days with the methods of Su et al.\(^2\). Means and standard deviations (SD) were calculated from at least 50 measurements. The ex-type living cultures were deposited in the China General Microbiological Culture Collection Center (CGMCC). The dried culture and microscope slide were deposited in Herbarium Mycologicum, Academia Sinica, Beijing, China (HMAS). Nomenclatural novelties and descriptions were registered in MycoBank (https://www.MycoBank.org).

DNA extraction, PCR amplification and sequencing. Total genomic DNA was extracted from mycelia grown on PDA or OA plates using the protocol of Guo et al.\(^2\). Primers ITS1 and ITS4 were used to amplify the ITS region of the nuclear rRNA gene\(^2\), primers LROR/LR5 primers were used for the partial 28S rDNA (LSU)\(^3\), primers 983F and 2218R\(^4\) for the elongation factor 1-a gene (TEF), and primers Bt2a and Bt2b\(^5\) for the partial β-tubulin gene (TUB). PCR was performed in a 25 μL reaction volume containing 1.0 μL DNA template, 1.0 μL of each forward and reverse primers, 12.5 μL 2 × MasterMix (Tiangen Biotech Co. Ltd., Beijing, China) and 10.5 μL ddH2O with the following cycling parameters: 94 °C for 40 s; 35 cycles at 94 °C for 40 s, annealing temperature specific for the gene amplified (52 °C for LSU, 55 °C for TEF and ITS, 58 °C for TUB) for 40 s and 72 °C for 120 s; and a final extension at 72 °C for 10 min. The PCR products were sequenced by Beijing Sunbio-tech Co. Ltd. (Beijing, China). Sequences were compared with accessions in the GenBank database via a BLASTn search to determine the most likely taxonomic designation.

Phylogenetic analysis. Sequence data of the four loci were aligned with Clustal X\(^3\). Reference sequences were retrieved from GenBank and the accession numbers indicated in Table 1. Manual editing of sequences was performed in MEGA6\(^4\). The concatenated sequences (LSU + ITS) or (LSU + TUB + TEF + ITS) were assembled using SeaView\(^5\) and alignments were deposited in TreeBASE (www.treebase.org, submission no.: S25764). The combined dataset of two or four loci was analyzed phylogenetically using Bayesian MCMC\(^3\) and Maximum Likelihood\(^7\) respectively. For the Bayesian analyses, the models of evolution were estimated by using MrModeltest 2.3\(^8\). Posterior probabilities (PP)\(^9,10\) were determined by Markov Chain Monte Carlo sampling (MCMC). Six simultaneous Markov chains were run for 2,000,000 generations and trees were sampled every 100th generation (resulting in 20,000 total trees). The first 4,000 trees represented the burn-in phase of the analyses and were discarded and the remaining 16,000 trees were used for calculating PP in the majority rule consensus tree. For the ML analysis in RAxML\(^3\), the GTRGAMMA model was used for all partitions, in accordance with recommendations in the RAxML manual against the use of invariant sites. Analyses were performed using the CIPRES web portal\(^11\). Trees were visualised in TreeView 1.6.6\(^12\).

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Author contributions
L.S. performed all the experimental work, culturing the samples, isolating in pure culture the fungi and performing their phenotypic characterization, as well as the DNA extraction and purification, gene sequencing and data processing for phylogenetic analysis, being one of the major contributors of this manuscript. H.Z., and C.Q., supervised all steps of the experimental work by L.S., and reviewing of the draft several times. Y.N. helped buy processing for phylogenetic analysis, being one of the major contributors of this manuscript. H.Z., and C.Q., supervised all steps of the experimental work by L.S., and reviewing of the draft several times. J.G. and L.Z. gave useful suggestions to write the manuscript and reviewed several times the draft.

Competing interests
The authors declare no competing financial interests.

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