Three new species of *Entoloma* subgenus *Pouzarella* from China based on morphological and molecular data

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Academic editor: O. Raspé | Received 11 September 2018 | Accepted 26 November 2018 | Published 13 December 2018

Citation: He X-L, Horak E, Wang D, Peng W-H, Gan B-C (2018) Three new species of *Entoloma* subgenus *Pouzarella* from China based on morphological and molecular data. MycoKeys 44: 1–18. https://doi.org/10.3897/mycokeys.44.24998

Abstract

In the present paper, three additional species of *Entoloma* subgenus *Pouzarella* viz. *E. erectoides*, *E. griseocarpum* and *E. rubropilosum* are described from China. *E. rubropilosum* is a typical species in section *Pouzarella*; *E. griseocarpum* and *E. erectoides* are members of sect. *Dysthales*. The taxa are further confirmed by ITS, RPB2, LSU and mtSSU analyses and phylogenetic relationships with other *Entoloma* subgen. *Pouzarella* species are also discussed. ITS sequence analysis showed that the sizes of the entire ITS region and ITS1 are remarkably divergent, while the ITS2 is conserved in length within *Entoloma* subgen. *Pouzarella*. Molecular analyses, based on the combined dataset, demonstrated that species diversity of subgen. *Pouzarella* in China is much higher than previously thought, in the present study twenty phylogenetic species from China are taken into consideration. On the other hand, morphological and molecular analyses suggested that classification of *Entoloma* subgen. *Pouzarella* probably has to be fundamentally re-adjusted based on additional data.

Keywords

Entolomataceae, systematics, taxonomy, multi-gene analyses

Introduction

*Pouzarella* Mazzer is a distinctive group of entolomatoid species that was accepted as a genus by some researchers (Horak 2008; Largent 1994; Mazzer 1976). Others consider it as a subgenus of *Entoloma* P. Kumm. (Noordeloos 1979, 1992, 2004; Noordeloos
and Gates 2012). Recent molecular phylogenetic studies (Co-David et al. 2009; Baroni and Matheny 2011; He et al. 2013; Largent and Bergemann 2015) based on multi-loci showed that *Pouzarella* actually represent a distinct monophyletic group separated from the other entolomatoid groups. In addition, taxa of *Pouzarella* are easily recognised both by macro- and micromorphological characters. However, here we still treat it as a subgenus of *Entoloma* s.l. because accepting *Pouzarella* as a separate genus would make *Entoloma* s.l. paraphyletic. Taxonomical revision of other well supported, distinct clades of *Entoloma* s.l. is therefore needed before a formal decision on the generic status of *Pouzarella* can be made. When such a revision is achieved, we predict, that *Pouzarella* will be a well-defined genus based on morphological and molecular evidence. In the field, species of *Entoloma* subgen. *Pouzarella* are often overlooked due to their small and dull-coloured basidiomes. However, the inconspicuous species are widespread and have been reported from tropical to temperate regions. So far, more than seventy species have been described worldwide (Mazzer 1976; Horak 1980, 1983, 2008; Desjardin and Baroni 1991; Noordeloos 1992, 2004; Noordeloos et al. 1992; Manimohan et al. 1995; Baroni and Ortiz 2002; Manimohan et al. 2006; Karstedt et al. 2007; Baroni et al. 2008; Largent et al. 2011; Baroni et al. 2012; Noordeloos and Gates 2012; He et al. 2013; Largent and Bergemann 2015; Raj and Manimohan 2017).

Basidiomes of members in subgen. *Pouzarella* are easy to recognise. However, many species have in common small basidiome size and greyish colours and, therefore, it is difficult to distinguish them to species by morphological characters only. Accordingly, both morphological and molecular data are needed to refine the species concept and understand the diversity of these small agarics in *Entoloma* subgen. *Pouzarella*.

In previous studies, seven species of *Entoloma* subgen. *Pouzarella* were reported to occur in China (Ying 1995; He et al. 2013). However, we believe that subgen. *Pouzarella* remains poorly understood in this region, rich in many diverse ecological habitats. In continuation of previous surveys, further field work was carried out in southwest and northeast China. More than 50 samples matching the concept of *Entoloma* subgen. *Pouzarella* were collected and many turned out to be different from the locally already known species. As a first step, three distinctive new taxa are described in the present study whereas the other specimens were shelved for the moment because of scarcity of material. To further confirm the three new taxa and infer the affinities amongst representative species of *Entoloma* subgen. *Pouzarella*, phylogenetic analysis was carried out based on the combination of ITS, RPB2, mtSSU and nLSU sequences.

**Materials and methods**

**Morphological descriptions**

Fresh basidiomes were photographed in the field and described macroscopically. Colour notations follow Kornerup and Wanscher (1978). Microscopic examination was done using a Leica DM5000B microscope. Basidiospores, basidia and pileipellis were
mounted and measured in 5% potassium hydroxide (KOH) and/or 1% Congo Red. Pigmentation of the micro-structures was observed in distilled water. Measurements of the basidiospores excluded hilar appendix (apiculus) and at least 30 basidiospores of each specimen were measured. Q represents the length to width ratio of a basidiospore in profile view; $Q$ represents the average Q of all basidiospores and is given ± standard deviation; $x$ represents the means of basidiospore length and width ± standard deviation. All cited collections, including the holotypes, are deposited at the Mycological Herbarium of Soil and Fertilizer Institute, Sichuan Academy of Agricultural Sciences (SAAS), Chengdu, China and the Herbarium ZT of ETH Zurich, Switzerland.

**DNA extraction, PCR amplification and sequencing**

Genomic DNA was extracted with Biospin Fungus Genomic DNA Extraction Kit following the manufacturer’s instructions. PCR amplification was performed using DreamTaq™ Green PCR Master qMix (2×), Fermentas. The primers for RPB2 amplification were rpb2-6F and rpb2-7R, rpb2-i6f and rpb2-i7r (Matheny 2005; Co-David et al. 2009). ITS regions were amplified with ITS5 and ITS4; LR0R and LR5, MS1 and MS2 were used for nLSU and mtSSU amplification, respectively (White et al. 1990; http://www.biology.duke.edu/fungi/mycolab/primers.htm).

**Sequence alignment and phylogenetic analyses**

Sequences used in phylogenetic analyses are listed in Table 1 and aligned in muscle 3.6 (Edgar 2004). The aligned sequences were manually modified where necessary in Mega 6.0 (Tamura et al. 2013). Phylogenetic analyses were based on the combined ITS, nLSU, RPB2 and mtSSU sequences. For the ITS region, only ITS1 and ITS2 were kept for further analyses. Conflicts between the ITS, nLSU, RPB2 and mtSSU datasets were evaluated by comparing the topologies resulting from the phylogenetic analysis of the single gene. As no conflict was detected amongst the well supported clades of the different trees, sequences of the four genes were combined for further analyses.

**Maximum likelihood analysis** – ML analysis was carried out by the web RAxML Version 8 (http://www.phylo.org/sub_sections/portal/) under the GTRGAMMAI model with 1000 bootstrap replicates (Stamatakis 2014). “Find best tree using maximum likelihood search” option was selected when analysis was undertaken.

**Maximum parsimony analysis** – MP analysis was performed using PAUP* version 4.0b10 (Swofford 2003). All characters were treated as unordered and of equal weight. Gaps were treated as missing data. Bootstrap values (BS) were obtained from 1000 replicates.

**Bayesian analysis** – Bayesian analysis was performed using MrBayes 3.2.6 (Ronquist and Huelsenbeck 2003). The best substitution models for each marker were selected using the Akaike Information Criterion (AIC) in jModelTest 2.1.7 (Darriba
### Table 1. A list of taxa, specimens and GenBank accession numbers of sequences used in this study.

| Taxa                | Collection No. | Origin          | GenBank accessions | Remarks                  |
|---------------------|----------------|-----------------|--------------------|--------------------------|
|                     |                |                 | ITS                | LSU                      | RPB2                     | mtSSU                     |                          |
| *Entoloma albostrigosum* | DL Largent 9641 | Australia: Queensland | –     | HQ876535 | HQ876513 | HQ876557 | GenBank ID: *Pouzarella albostrigos* |
|                     | DL Largent 9663 | Australia: Queensland | –     | HQ876536 | HQ876514 | HQ876558 | GenBank ID: *P. albostrigosa* |
| *E. annaeus*        | ME Noordeloos 200314 | China: Jilin | KC710056 | GQ289153 | GQ289225 | GQ289293 |                          |
| *E. berringtonense* | DL Largent 9901 (Holotype) | Australia: Queensland | –     | HQ876524 | HQ876543 | HQ876546 | GenBank ID: *P. berrington* |
| *E. changchunense*  | HMJAU 3886 (Holotype) | China: Jilin | –     | JQ933095 | –        | JQ993061 |                          |
| *E. cassinii*       | GDGM 28821 | China: Guangdong | KG78997 | JQ993085 | JQ993085 | JQ993058 |                          |
| *E. debile*         | DLL 99784 | Australia: Queensland | –     | HQ876528 | HQ876506 | HQ876550 | GenBank ID: *P. debile* |
| *E. dindenense*     | DL Largent 9623 | Australia: Queensland | –     | HQ876527 | HQ876505 | HQ876549 | GenBank ID: *P. fusca*  |
| *E. erectoides*     | SAAS 1232 (Holotype) | China: Sichuan | MHO20746 | KUS44255 | KUS44496 | –        |                          |
|                     | SAAS 945      | China: Sichuan | MHO20769 | KUS44299 | KUS44498 | –        |                          |
|                     | SAAS 1361 | China: Jilin | MHO20755 | –        | –        | KUS44484 | –        |
| *E. farinosum*      | DL Largent 9934 (Holotype) | Australia: Queensland | –     | HQ876516 | HQ876495 | HQ876538 | GenBank ID: *P. farinosa* |
|                     | DL Largent 9900 | Australia: Queensland | –     | HQ876515 | HQ876494 | HQ876537 | GenBank ID: *P. farinosa* |
| *E. furfunaceum*    | GDGM 28818 (Holotype) | China: Jilin | JX975293 | JQ993094 | JQ993084 | JQ993062 |                          |
|                     | SAAS 104       | China: Jilin | –     | –        | –        | –        |                          |
| *E. griseocarpum*   | SAAS 1230    | China: Tibet | MHO20753 | KUS44253 | KUS44500 | KUS44438 |                          |
|                     | SAAS 1328 (Holotype) | China: Tibet | MHO20766 | KUS44256 | KUS44501 | KUS44455 |                          |
|                     | SAAS 951      | China: Sichuan | MHO20770 | KUS44242 | KUS44499 | KUS44457 |                          |
| *E. lageniforme*    | DL Largent 9895 | Australia: Queensland | –     | HQ876523 | HQ876502 | HQ876545 | GenBank ID: *P. lageniformis* |
| *E. lasium*         | DL Largent 9662 | Australia: Queensland | –     | HQ876529 | HQ876507 | HQ876551 | GenBank ID: *P. lasia* |
|                     | DL Largent 9670 | Australia: Queensland | –     | HQ876530 | HQ876508 | HQ876552 | GenBank ID: *P. lasia* |
|                     | DL Largent 9807 | Australia: Queensland | –     | HQ876533 | HQ876511 | HQ876555 | GenBank ID: *P. lasia* |
|                     | DL Largent 9811 | Australia: Queensland | –     | HQ876534 | HQ876512 | HQ876556 | GenBank ID: *P. lasia* |
|                     | DL Largent 9729 | Australia: Queensland | –     | HQ876531 | HQ876509 | HQ876553 | GenBank ID: *P. lasia* |
|                     | DL Largent 9778 | Australia: Queensland | –     | HQ876532 | HQ876510 | HQ876554 | GenBank ID: *P. lasia* |
| *E. nodosporum*     | TENN:068582 | USA: Tennessee | KY744163 | MF797654 | –        | –        | GenBank ID: *P. nodosporum* |
| *E. pamiae*         | DL Largent 9794 | Australia: Queensland | –     | HQ876517 | HQ876496 | HQ876539 | GenBank ID: *P. pamiae* |
|                     | DL Largent 9834 | Australia: Queensland | –     | HQ876519 | HQ876498 | HQ876541 | GenBank ID: *P. pamiae* |
|                     | DL Largent 9808 | Australia: Queensland | –     | HQ876518 | HQ876497 | HQ876540 | GenBank ID: *P. pamiae* |
| *E. perbloxamii*    | MEN 2004071 (Holotype) | Australia: Tasmania | KC710117 | GQ289178 | GQ289249 | GQ289318 |                          |
| *E. pilocystidiatum*| DL Largent 9848 | Australia: Queensland | –     | HQ876520 | HQ876499 | HQ876542 | GenBank ID: *P. pilocystidiata* |
| Taxa | Collection No. | Origin                 | GenBank accessions      | Remarks                      |
|------|----------------|------------------------|-------------------------|------------------------------|
|      |                |                        | ITS | LSU | RPB2 | mtSSU |                               |
| E. pilocystidatum | DL Largent 9932 (Holotype) | Australia: Queensland | –  | HQ876521 | HQ876500 | HQ876543 | GenBank ID: P. pilocystidatum |
|      | DL Largent 9949 | Australia: Queensland | –  | HQ876522 | HQ876501 | HQ876544 | GenBank ID: P. pilocystidatum |
| E. prunuloides    | MEN 200340      | Slovakia               | KC710073 | GQ289184 | GQ289255 | GQ289324 |                               |
| E. rubropilosum   | SAAS 406 (Holotype) | China: Sichuan        | MH020761 | KU534218 | KU534488 | KU534439 |                               |
| E. rubropilosum   | SAAS 1112       | China: Tibet           | MH020767 | KU534252 | KU534502 | KU534454 |                               |
| E. setiforme      | DL Largent 9809 (Holotype) | Australia: Queensland | –  | HQ876526 | HQ876503 | HQ876547 | GenBank ID: P. setiformis     |
| E. setiforme      | DL Largent 9810 | Australia: Queensland | –  | HQ876526 | HQ876504 | HQ876548 | GenBank ID: P. setiformis     |
| E. silvanum       | K(M) 191739 (Holotype) | India: Kerala          | KJY643747 | KJY643724 | –        | –        |                               |
| E. sp. 1          | SAAS 894        | China: Sichuan         | MH020765 | KU534245 | KU534491 | KU534447 |                               |
| E. sp. 2          | SAAS 1088       | China: Jilin           | MH020749 | KU534246 | –        | KU534441 |                               |
| E. sp. 3          | SAAS 1210       | China: Jilin           | MH020752 | KU534248 | –        | KU534449 |                               |
| E. sp. 4          | SAAS 249        | China: Sichuan         | MH020759 | KU534243 | –        | –        |                               |
| E. sp. 5          | SAAS 1209       | China: Jilin           | MH020751 | –      | KU534492 | KU534448 |                               |
| E. sp. 6          | SAAS 291        | China: Jilin           | MH020760 | –      | KU534486 | KU534444 |                               |
| E. sp. 7          | SAAS 1290       | China: Jilin           | MH020754 | KU534249 | –        | KU534456 |                               |
| E. sp. 8          | SAAS 1360       | China: Jilin           | MH020756 | KU534258 | KU534493 | KU534450 |                               |
| E. sp. 9          | SAAS 1464       | China: Sichuan         | MH020747 | –      | –        | KU534442 |                               |
| E. sp. 10         | SAAS 100        | China: Sichuan         | MH020746 | –      | –        | KU534444 |                               |
| E. sp. 11         | SAAS 102        | China: Sichuan         | MH020745 | KU534254 | –        | –        |                               |
| E. strigosissimum | 152             | Italy                  | JF908004 | –      | –        | –        |                               |
| E. suvananense    | GDGM 28823 (Holotype) | China: Jilin          | JQ320113 | JQ410329 | –        | –        |                               |
| E. suvananense    | KA 12-1534      | South Korea            | KJ523135 | –      | –        | –        |                               |
| E. tenuissimum    | GDGM 28813      | China: Jilin           | JX975295 | JQ993097 | JQ993086 | JQ993059 |                               |
| E. violaceovillosum | GDGM 28814 (Holotype) | China: Heilongjiang   | JX975294 | JQ993096 | JQ993087 | JQ993060 |                               |
| E. yunnanense     | P. Manomohan 645 (Holotype) | India: Kerala         | –      | GQ289205 | GQ289273 | GQ289345 |                               |
| E. yunnanense     | GDGM 28815      | China: Yunnan          | JQ320108 | JQ320128 | –        | JQ993057 |                               |

Sequences in bold and marked with “KU” and “MH” are newly generated in this study. Sequences marked with “GQ” were from Co-David et al. (2009). Sequences marked with “HQ” were from Largent et al. (2011). KC710056, KC710073 and KC710117 were from Morgado et al. (2013). KC678996, KC678997 and sequences marked with “JQ” and “JX” were from He et al. (2013). KY643747 and KY643724 were from Raj and Manimohan (2017). JF908004 was from Osmundson et al. (2013). KJ523135 was from Kim et al. (2015). KY744163 and MF797654 are unpublished.
et al. 2012). GTR+I+G model was selected for nLSU, GTR+G for mtSSU and ITS and SYM+G for RPB2. Two runs of six Markov chains were run from random starting trees for 6 million generations and sampled every 100 generations. Every time the diagnostics were calculated, 25% of the samples from the beginning of the chain were discarded. Runs were stopped after the average standard deviation was below 0.01. Bayesian posterior probabilities (BPP) were determined after calculating a 75% majority rule consensus tree.

**Taxonomy**

*Entoloma erectoides* Xiao L. He & E. Horak, sp. nov.
MycoBank No: MB828692
Figs 1a, b, 2

**Diagnosis.** *E. erectoides* is distinguished by the greyish brown pileus covered with silvery fibrils, large basidiospores (13.5–17.5 × 8–9.5 µm) and presence of ovoid to sub-utriform cheilocystidia.

**Type.** CHINA. SICHUAN PROV.: Yajiang County, Gexigou National Nature Reserve, 29°33’N, 100°50’E, elevation ca. 2980 m, August 2014, He XL (SAAS 1232, holotype; ZT 14180, isotype).

**Etymology.** *Erectoides*, refers to the suberect to erect fibrils on the pileus.

**Description.** Pileus 5–15 mm broad, bluntly conic, convex or campanulate, dry, slightly hygrophanous, greyish-brown to brown (5C2–5D2), densely covered with suberect fibrils or minutely fibrillose squamules; fibrils silvery greyish, striate from entire margin to near centre. Lamellae sinuate, ventricose, distant, up to 3.5 mm wide, moderately thick, with two tiers of lamellulae, dark grey to brownish-grey, with entire and concolorous edge. Stipe 40–60 × 1–2.5 mm, central, cylindrical, equal, dry, concolorous with pileus, densely covered with grey to greyish fibrils, hollow, surface dry, with a pale yellow brownish to pale brownish strigose base. Context thin, concolorous with pileus. Odour and taste not distinctive.

Basidiospores (13–) 13.5–17.5 × 8–9.5 (–10.5) µm (x = 15.5 ± 0.5 × 8.8 ± 0.3 µm), Q = 1.50–1.94 (Q = 1.72 ± 0.03), heterodiametric, strongly angled in profile and face views with 6–9 facets, appearing nodulose, pale yellow brownish, thick-walled. Basidia 39–48 × 13–18 µm, subclavate or clavate, 4-spored. Aborted basidia scattered in the hymenium, often filled with dark brown amorphous cytoplasmic pigment. Lamellar trama dark brown, composed of parallel, cylindrical, heavily encrusted and thin-walled cells, 6–15 µm wide. Lamellar edge sterile. Cheilocystidia 33–90 × 12–33 µm, broadly ovoid to utriform (32–65 × 22–30 µm) or lageniform (70–90 × 16–20 µm), with pale brownish, intracellular pigment, slightly thick-walled. Pileipellis a trichoderm composed of clustered and suberect hyphae, walls externally encrusted with
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**Figure 1.** Basidiomata of the new species. **a, b** *E. erectoides* (SAAS 1361, SAAS 1232) **c, d** *E. griseocarpum* (SAAS 951, SAAS 1328) **e, f** *E. rubropilosum* (SAAS 406, SAAS 1112).
brown pigment; terminal cells 30–50 (–90) × 8–15 µm, cylindrical to slightly fusoid; subpellis composed of cylindrical, encrusted hyphae, up to 20 µm broad. Stipitipellis composed of thin-walled and pale yellowish-brown encrusted hyphae; terminal cells 40–80 × 9–15 µm, cylindrical to slender fusoid, walls encrusted with pale yellow-brown pigment. Oleiferous hyphae absent. Clamp connections absent.

**Habitat.** Scattered or gregarious on soil and amongst leaf litter in broadleaf forest dominated by *Quercus* or on soil amongst decaying leaves of *Betula*, *Pandus* and *Abies*.

**Additional collections examined.** CHINA. SICHUAN PROVINCE: CHINA: SICHUAN PROV. Yajiang County, Gexigou National Nature Reserve, 29°33’N, 100°50’E, elevation ca. 2980 m, 24 July 2013, He X.L. (SAAS 945). JILIN PROV.: Antu County, Changbai Mountains, 42°10’N, 127°55’E, elevation ca. 750 m, 25 August 2014, He X.L. (SAAS 1361).

**Comments.** Morphologically, *Entoloma erectoides* is a member of section *Dysthales*. In literature, a few species in section *Dysthales* are described having silvery fibrils or squamules on pileus and stipe. Accordingly, *E. erectoides* can be confused with the Argentinean *E. calileguense* Blanco-Dios (as *Pouzarella variabilis* T.J. Baroni, Albertó, Niveiro & B.E. Lechner in Baroni et al. 2012). Both species have silvery greyish-brown erect fibrils or squamules on the pileus and stipe. However, the latter species is easily separated by the much larger basidiospores (16–23.5 × 10–12 µm, Baroni et
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*E. farinosum* (Largent & Skye Moore) Noordel. & G.M. Gates, reported from Australia, differs by globose or nearly napiform cheilocystidia (Largent et al. 2011). In addition, this taxon is separated from *E. erectoides* by molecular evidence. *E. tenuissimum* T.H. Li & Xiao-Lan He, also recorded from China, is distinguished by the smaller and slimmer basidiomes and taxonomically is also distinctly different based on molecular analysis (He et al. 2013). *E. argenteolanatum* (T.J. Baroni, Perd.-Sánch. & S.A. Cantrell) Noordel. & Co-David was found on decaying leaves of tropical trees and shrubs in the Dominican Republic and is characterised by denser and longer silvery fibrils and the place of discovery in the Caribbean on the island of Hispaniola (Baroni et al. 2008). The other grey-brown species with silvery fibrils in section *Versatile* could be distinguished by the innately fibrillose pileus and stipe and colourless hymenial cystidia.

**Entoloma griseocarpum** Xiao L. He & E. Horak, sp. nov.
MycoBank No: MB828701
Figs 1c, d, 3

**Diagnosis.** *E. griseocarpum* is characterised by the greyish-brown pileus, large basidiospores (12.5–15.5 × 7.5–9 µm) and broadly clavate, ovoid to lageniform cheilocystidia.

**Type.** CHINA. TIBET: Linzhi, Lulang, 29°94’N, 94°79’E, elevation ca. 3800 m, 18 September 2014, He X.L. (SAAS 1328, holotype).

**Etymology.** *griseocarpum*, refers to the greyish-brown coloured basidiomes.

**Description.** Pileus 5–20 mm broad, hemispherical, convex, bluntly conic to broadly campanulate, dry, not hygrophanous, greyish-brown to brown (4D3–4E3), densely covered by suberect hispid or minutely squamulose overall, denser in centre; fibrils dark grey, pale grey brownish or concolorous with pileal surface (4D2–4D3), striate from entire margin to near centre. Lamellae sinuate with short decurrent tooth, ventricose, distant, moderately thick, up to 3 mm broad, with two tiers of lamellulae, dark grey to brownish-grey, with entire and concolorous edges. Stipe 20–50 × 0.7–1.5 mm, cylindrical, equal, dry, concolorous with pileus, densely covered with pale yellow brownish flocculose hairs, hollow, with a dirty yellowish to pale yellow brownish strigose base. Context thin, concolorous with pileus. Odour and taste not distinctive.

Basidiospores 12.5–15.5 (–17) × (6.5–) 7.5–9 (–9.5) µm (x = 13.8 ± 0.3 × 8.3 ± 0.3 µm), Q = 1.60–1.94 (Q = 1.71 ± 0.02), heterodiametric, strongly angled in profile and face view with 6–10 facets, appearing nodulose, pale yellow brownish, thick-walled. Basidia 35–55 × 11–13 (–15) µm, subclavate to clavate, 4-spored. Aborted basidia scattered in the hymenium, filled with dark brown amorphous cytoplasmic pigment. Lamellar trama dark brown, composed of parallel, cylindrical, heavily encrusted and thin-walled elements. Lamellar edges sterile. Cheilocystidia 23–50 × (10–) 12 20 µm, broadly clavate, ovoid to lageniform; with brown, intracellular pigment, slightly thick-walled. Pileipellis a trichoderm composed of yellow brown, suberect and multiseptate hyphae, walls heavily encrusted with brown.
Figure 3. Microscopic structures of *E. griseocarpum* (drawn from the holotype). a Basidiospores b Cheilocystidia c Pileipellis.

pigment; terminal cells 35–105 × 8–27 µm, cylindrical, subclavate or bullet-shaped, thin to moderately thick-walled; subpellis composed of cylindrical encrusted hyphae, up to 25 µm diam. Stipitipellis composed of yellow-brown encrusted hyphae; terminal cells 40–80 × 4–10 µm, slender cylindrical with obtuse apex, thin-walled, sparsely encrusted with pale yellowish-brown pigment. Oleiferous hyphae absent. Clamp connections absent.

**Habitat.** Scattered on soil amongst decaying litter in mixed conifer-broadleaf forest dominated by *Quercus, Betula, Rhododendron* and *Abies*.

**Additional collections examined.** CHINA. TIBET: Linzhi, Lulang, 29°94’N, 94°79’E, elevation ca. 3800 m, 18 September 2014, He X.L. (SAAS 1230, SAAS 1657, SAAS 1751, SAAS 1871). SICHUAN PROV.: Jiuzhaigou, 33°28’N, 103°59’E, elevation ca. 3000 m, 20 July 2013, He X.L. (SAAS 951).

**Comments.** The greyish-brown pileus covered by suberect hispid or minutely squamulose, the brown external encrustations on pileipellis and stipitipellis and the cylindrical terminal cells of pileipellis and stipitipellis indicate *E. griseocarpum* belongs to the sect. *Dysthales*. It is very similar to *E. albostrigosum* (Largent & Abell-Davis) Blanco-Dios and *E. lasium* (Berk. & Broome) Noordel. & Co-David (Largent et al. 2011). However, *E. albostrigosum* is distinguished by the white strigose base and
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*E. lasium* differs by the smaller basidiospores (8.9–14.5 × 5.1–8.7 µm, Largent et al. 2011). In addition, the two species are distant from *E. griseocarpum* following phylogenetic analysis. *E. puertoricense* Blanco-Dios (as *P. caribaea* T.J. Baroni & B. Ortiz in Baroni and Ortiz 2002) resembles *E. griseocarpum* by the brownish-grey coloured basidiomes but is separated by its broader basidiospores (12.5–16.5 × 8.3–11.3 µm, Q = 1.26–1.65, Baroni and Ortiz 2002). Moreover, *E. puertoricense* was discovered in a tropical habitat in Puerto Rico (Baroni and Ortiz 2002). The similar *E. japonicum* (Hongo) Hongo, described from Japan, is also reminiscent of *E. griseocarpum* in the brownish pileus but is distinguished by the much larger basidiospores (15–18.5 × 9–10.5 µm, Hongo 1959). The well-known *E. dythales* (Peck) Sacc. also differs by the larger basidiospores (14–20 × 7.5–10 µm, Mazzer 1976). *E. fulvolanatum* (Berk. & Broome) Blanco-Dios from Sri Lanka is not only separated by its type locality but also by the narrower basidiospores measuring 12–16 × 7–8 µm (Mazzer 1976). Two species in subgen. *Pouzarella*, recently described from geographically neighbouring India, viz. *E. peechiense* K. N. A. Raj & Manim. and *E. silvanum* K. N. A. Raj & Manim., have somewhat similar basidiomes as compared to *E. griseocarpum*; however, their ITS and LSU sequences are distinctly different (Raj and Manimohan 2017). The third Indian species *E. lomapadum* Manim., Joseph & Leelav. is readily recognised by the much smaller basidiospores measuring 11–13 × 6–9 µm (Manimohan et al. 1995). There were four other species in subgen. *Pouzarella* which showed some similarities to *E. griseocarpum*. *E. fibrillosipes* (Murrill) Noordel. & Co-David is distinguished by the much larger basidiospores (17–22 × 7.5–10 µm, Mazzer 1976). *E. subdeceptivum* Courtec. and *E. rotula* (Romagn.) Noordel. & Co-David are lignicolous (Mazzer 1976). *E. homomorphum* (Romagn.) Singer differs by the larger basidiospores (15–19 × 9–11.5 µm, Mazzer 1976).

**Entoloma rubropilosum** Xiao L. He & E. Horak, sp. nov.
MycoBank No: MB828699
Figs 1e, f, 4

**Diagnosis.** *E. rubropilosum* is distinct due to its reddish-brown coloured pileus and stipe, large basidiospores (13–17 × 7.5–9.5 µm), broadly clavate cheilocystidia, distinctive thick-walled setiform caulocystidia and terminal cells of the pileipellis hyphae.

**Type.** CHINA: SICHUAN PROV.: Yajiang County, Gexigou National Nature Reserve, 29°33’N, 100°50’E, elevation ca. 2950 m, 24 July 2013, He X.L. (SAAS 406, holotype).

**Etymology.** Rubropilosum, refers to the reddish coloured fibrils on the pileus.

**Description.** Pileus 7–20 mm broad, conical-convex, truncate conical to broadly campanulate, dark reddish-brown (8D2–8D3) at first, becoming greyish-orange to pale beige brownish (5B2–5C2), dry, slightly hygrophanous, densely covered by reddish-brown erect or suberect squamules and fibrils; fibrils much denser at disc, margin not striate or very slightly striate only. Lamellae adnate to sinuate, ventricose, up to 2.5 mm wide, relatively
thick, with two tiers of lamellulae, brownish-pink when mature, with concolorous and entire edges. Stipe 40–73 × 0.8–2 mm, central, cylindrical, hollow, densely covered with rust reddish hairs or fibrils, very dark brown strigose at base. Odour and taste not distinctive.

Basidiospores (12.5–) 13–17 × 7.5–9.5 µm (\( x = 15.2 \pm 0.5 \times 8.5 \pm 0.3 \) µm), \( Q = 1.53–1.98 \) (\( Q = 1.76 \pm 0.02 \)), heterodiametrical, with 6–8 facets in profile and face views, sometimes multi-angled to nodulose, pale brownish, thick-walled. Basidia (32–) 38–45 (–50) × 12–16 µm, clavate, 4-spored. Aborted basidia inconspicuous. Lamellar edges sterile. Cheilocystidia 25–50 × 12–18 µm, broadly clavate, with faintly pale brownish, intracellular pigment, slightly thick-walled. Pleurocystidia absent. Pileipellis a trichoderm composed of brown hyphae; terminal cells 23–110 × 6–18 µm (diameter was measured at the base), slender setiform, gradually tapering towards subacute apex, sometimes subfusoid to somewhat bullet-shaped, thick-walled, with intraparietal and intracellular brown pigment; subpellis composed of cylindrical, relatively thin-walled hyphae, encrusted with yellow-brown pigment. Stipitipellis composed of loosely entangled, rather slender hyphae; terminal cells 45–120 × 5–11 µm (diameter was measured at the base), distinctly setiform with obtuse or subacute apex, thick-walled, with intraparietal and intracellular brown pigment. Oleiferous hyphae absent. Clamp connections absent.

Habitat. Scattered on soil amongst decaying litter in broadleaf forest dominated by Quercus or in mixed forest with Quercus, Betula, Rhododendron and Abies, also on soil in bamboo forest.

Additional collections examined. CHINA. SICHUAN PROV.: Yajiang County, Gexigou National Nature Reserve, 29°33’N, 100°50’E, elevation ca. 2950 m, 24 July 2013, He X.L. (SAAS 765); 24 July 2013, He X.L. (SAAS 706); 3 August 2013, He X.L. (SAAS 706).
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1488, SAAS 1112, ZT 14179). TIBET: Linzhi, Lulang, 29°94’N, 94°79’E, elevation ca. 3800 m, 18 September 2014, He X.L. (SAAS 1618, SAAS 1087); Linzhi, Kadinggou, 29°50’N, 93°26’E, elevation ca. 2950 m, 24 September 2014, He X.L. (SAAS 1456).

**Comments.** The setiform terminal cells of pileipellis and stipitipellis place *E. rubropilosum* in sect. *Pouzarella*. It is readily recognised in the field. A few species of *Entoloma* subgen. *Pouzarella* with reddish-brown fibrils or squamules have been reported in literature (Mazzer 1976; Baroni et al. 2008). *E. ferreri* (T.J. Baroni, Perd.-Sánch. & S.A. Cantrell) Noordel. & Co-David is distinguished by dark blackish stains on the pileus caused from handling and non-setiform pileocystidia and caulocystidia (Baroni et al. 2008). *E. strigosissimum* (Rea) Noordel. is separated by the larger basidiospores [15–19 (23) × 8.5–10.5 (11.5) µm, Mazzer 1976]. *E. squamifolium* (Murrill) Singer might be confused with *E. rubropilosum* due to the ferruginous hairs on the stipe (Mazzer 1976). However, *E. rubropilosum* can be distinguished by the setiform pileocystidia and caulocystidia. Furthermore, type of *E. squamifolium* was collected in a tropical location. The recently described *E. wayanadense* K. N. A. Raj & Manim. from India, also discovered in a tropical area, is similar to *E. rubropilosum* in its greyish-orange pileus with long hairs and the setiform terminal cells of pileipellis, but differs by the absence of cheilocystidia. In addition, the partial ITS sequence (419 bp, KY 643748) of *E. wayanadense* is quite different from that of *E. rubropilosum* (Raj and Manimohan 2017).

### Key to the species of *Entoloma* subgen. *Pouzarella* described from China

1. Pileus reddish-brown or greenish-brown ...........................................2
   – Pileus greyish-brown ....................................................................3

2. Pileus covered with reddish-brown suberect fibrils .......... *E. rubropilosum*
   – Pileus greyish-brown with reddish tinge, zonate .......... *E. changchunense*

3. Pileus covered with appressed or suberect silvery fibrils ....................4
   – Pileus squamulose or covered with suberect brownish fibrils ..........5

4. Pileus covered with appressed silvery fibrils .......... *E. subaraneosum*
   – Pileus covered with suberect silvery fibrils ..................................6

5. Pileus fibrillo-squamulose or squamulose-tomentose, growing in tropical forest .................................................... *E. crassicystidiatum*
   – Pileus covered with erect or suberect fibrils .........................7

6. Pileus pale brownish with pinkish tinge, basidiospores larger, average (16.8 ± 0.5) × (10.8 ± 0.3) µm ................................................. *E. tenuissimum*
   – Pileus greyish-brown, basidiospores smaller, average (15.5 ± 0.5) × (8.8 ± 0.3) µm ................................................................. *E. erectoides*

7. Average spore length less than 13 µm .......................... *E. furfuraceum*
   – Average spore length more than 13 µm ..................................8

8. Pileus greyish-brown, striate from entire margin to near centre .......... *E. griseocarpum*
   – Pileus peach brown, not striate ............................................ *E. yunnanense*
Figure 5. Phylogenetic relationships of *Entoloma* subgen. *Pouzarella* species inferred from the combined ITS, LSU, mtSSU and RPB2 dataset (new species are in bold). Bayesian posterior probability values (BPP > 0.90) and MP BS support values (> 50%) are indicated above branches as BPP/BS; RAxML BS support values (> 50%) are listed below branches.

**Molecular analysis**

A total 76 sequences were generated in this study and they were deposited in GenBank. The combined dataset in the molecular analyses is composed of 62 specimens and 2925 aligned sites. MP, ML and Bayesian analyses produced almost the same topologies except for the unsupported branches and the MP tree is shown (Fig. 5).

For the ITS sequences used in the analyses, both the size of the entire ITS1-5.8-ITS2 region as well as for ITS1 and ITS2 were separately compared. It is remarkable
that the sizes of the entire ITS region and ITS1 were significantly divergent. In general, the total length of ITS sequences in subgen. *Pouzarella* ranged from 591 bp to 1086 bp. ITS1 was highly variable in length, while the length of ITS2 is relatively conserved within subgen. *Pouzarella*. ITS1 and ITS2 spacer varied from 229 to 690 bp and from 202 to 255 bp, respectively. However, it is noteworthy that two groups of the whole ITS region and ITS1 spacer could be partitioned in length. One group was varying from 591 bp to 709 bp and the other from 967 bp to 1086 bp. For the 5.8S region, 159–162 bp were yielded. Despite 5.8S is highly conserved, 4 indels and 11 nucleotide substitutes were found in this region. Regarding RPB2 sequences, their length was considerably conserved.

The three new species in this study were placed in different clades, showing they are quite different from each other. *E. rubropilosum* in relatively close to *E. strigosissimum* in the analyses, but similarity of their ITS sequences is only 84%. *E. griseocarpum* is grouped with *E. yunnanense* J.Z. Ying, but more than 150 different bases were observed in their ITS sequences. *E. erectoides* and *E. furfuraceum* nested in the same clade; however, more than 100 different bases were detected amongst their ITS sequences.

**Results and discussion**

In the present study, three new species of *Entoloma* subgen. *Pouzarella* viz. *E. griseocarpum*, *E. erectoides* and *E. rubropilosum*, are reported from southwest China. The description is based on morphological and molecular characters. Together with the seven afore-mentioned species, ten taxa of *Entoloma* subgen. *Pouzarella* are now recorded for China. Five of those have been discovered in the northeast of China, four in southwest China and only one was reported from southern China (Ying 1995; He et al. 2013).

In the phylogenetic analyses, 35 taxa in subgen. *Pouzarella* were included and twenty phylogenetic species from China were recovered, suggesting a high species diversity in this geographical region. Five distinct clades (Clades I–V) were observed and the three new species are phylogenetically separated from each other. Based on morphological characters, *Pouzarella* (as a genus or subgenus of *Entoloma* s.l.) was divided into three sections (*Dysthales*, *Pouzarella* and *Versatile*, Mazzer 1976; Noordeloos 1992). In our phylogenetic tree, Clade I corresponds to sect. *Pouzarella* while the taxa belonging to Clade II are accommodated in sect. *Versatile*. Except for several species of uncertain position, most members of Clade III, Clade IV and Clade V belong to the traditional sect. *Dysthales* morphologically. *E. rubropilosum* is nested in Clade I, which also includes *E. nodosporum* (G.F. Atk.) Noordel.and *E. strigosissimum*, as well as two still unknown species (*Entoloma* sp. 10 and *Entoloma* sp. 11) collected in China that morphologically fit the concept of sect. *Pouzarella*. These four species possess setiform pileocystidia and caulocystidia, somewhat reddish-brown or reddish fibrils on stipe and pileus. *E. erectoides* and *E. griseocarpum* are placed in Clade V.

The length of the ITS sequence of the species nested in Clade I, Clade II and Clade IV is relatively short, ranging from 591 bp to 709 bp. Available ITS sequences of *Entolo-
ma subgen. *Pouzarella* species in Clade V are recorded from 967 bp to 1086 bp. Unfortunately, no ITS sequences are available for comparison of the taxa belonging to Clade III. Eventual combination, referring both to morphological and molecular evidence, may in the future fundamentally change the classification of *Entoloma* subgen. *Pouzarella*.

**Acknowledgements**

We express special thanks to Ms. Bo Zhang (Jilin Agricultural University) and Dr. Hai-Xia Ma (Chinese Academy of Tropical Agricultural Sciences) for assistance during fieldwork. This study is funded by the National Natural Science Foundation of China (Nos. 31400023, 31770020) and the Sichuan Provincial Innovation Ability Promotion Engineering (2016ZYPZ-028).

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