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A new cystidiate variety of *Omphalina pyxidata* (Basidiomycota, tricholomatoid clade) from ItalyALFREDO VIZZINI^{1*}, MARIANO CURTI², MARCO CONTU³ & ENRICO ERCOLE¹¹*Dipartimento di Scienze della Vita e Biologia dei Sistemi - Università degli Studi di Torino
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ABSTRACT — A new variety of *Omphalina pyxidata*, var. *cystidiata*, is here described and illustrated based on morphological and molecular data. The new combination *Infundibulicybe lateritia* is introduced.

KEY WORDS — *Agaricomycetes*, *Agaricales*, omphalinoid fungi, *Contumyces*, taxonomy

Introduction

Within the omphalinoid fungi — small agarics with a convex to deeply umbilicate pileus, central stipe, thin context, decurrent lamellae, white spore-print, and thin-walled inamyloid smooth spores (Norvell et al. 1994, Lutzoni 1997, Redhead et al. 2002a,b) — taxa with cystidiate basidiomata are thus far known only in the hymenochaetoid clade (*Blasiphalia* Redhead, *Contumyces* Redhead et al., *Rickenella* Raithelth.; Moncalvo et al. 2000, 2002, Redhead et al. 2002b, Larsson et al. 2006, Larsson 2007).

During fieldwork focused on bryophilous *Galerina* species, we collected an omphalinoid fungus growing on mosses close to *Galerina discreta* E. Horak et al. We initially believed that these specimens represented a new species of *Contumyces* (Contu 1997, Redhead et al. 2002b) based on the presence of well-differentiated pileo-, caulo-, cheilo-, and pleurocystidia and an irregular hymenophoral trama. However an ITS-rDNA analysis implied they might represent an unknown as-yet undescribed variety of *Omphalina pyxidata* (Bull.) Quél., the type species of *Omphalina* Quél. (*Omphalina* s.s. = tricholomatoid clade sensu Matheny et al. 2006, Binder et al. 2010 and Vizzini et al. 2011a). This genus is characterized by bryophilous basidiomata with a reddish brown tinged

pileus and stipe, paler and well-developed lamellae, a smooth (not scaly) pileus, absence of hymenial and pileal cystidia, and presence of clamp-connections (Redhead et al. 2002a, Elborne 2008). We fully describe and illustrate the new taxon below.

Materials & methods

Morphology

The macromorphological descriptions follow the detailed field notes taken on fresh material. The micromorphological descriptions are based both upon study of fresh and herbarium material. Dried material was revived in 5% KOH and stained in Congo Red, Cotton Blue, and Melzer's reagent. The basidiospore range and means are based on measurements of 25 spores. In the macro- and microscopic descriptions Q = the quotient of spore length and width and Q_m = the average quotient; L = number of entire lamellae; l = number of lamellulae between each pair of entire lamellae. The basidial width was measured at the broadest part, and the length was measured from the apex (sterigmata excluded) to the basal septum. Colour codes in brackets (e.g., Se 45) follow Séguy (1936), hereafter referred to as (Se). Author citations follow the IPNI Authors and the Index Fungorum Authors of Fungal Names websites. Herbarium abbreviations follow Thiers (2011) with the exception of "EM" and "GT", referring to the personal herbaria of Enzo Musumeci and Gérard Trichies, respectively. The type collection is housed at TO (Herbarium of the Department of Plant Biology, University of Turin, Italy), and both name and Latin diagnosis of the new variety as well as the new combination are deposited in MycoBank (<http://www.mycobank.org>).

DNA extraction, PCR amplification, DNA sequencing

Genomic DNA was isolated from 1 mg of 3 herbarium specimens (TABLE 1) using the DNeasy Plant Mini Kit (Qiagen, Milan Italy). Universal primers ITS1f/ITS4 were used for the ITS region amplification (White et al. 1990; Gardes & Bruns 1993). Amplification reactions were performed in PE9700 thermal cycler (Perkin-Elmer, Applied Biosystems) in a 25 μ l reaction mixture using the following final concentrations or total amounts: 5 ng DNA, 1 \times PCR buffer (20 mM Tris/HCl pH 8.4, 50 mM KCl), 1 μ M of each primer, 2.5 mM MgCl₂, 0.25 mM of each dNTP, 0.5 unit of *Taq* polymerase (Promega). The PCR program was as follows: 3 min at 95°C for 1 cycle; 30 s at 94°C, 45 s at 50°C, 2 min at 72°C for 35 cycles, 10 min at 72°C for 1 cycle. PCR products were resolved on a 1.0% agarose gel and visualized by staining with ethidium bromide. PCR products were purified and sequenced by DiNAMYCODE srl (Turin, Italy). Sequence assembly and editing were performed using Geneious v5.3 (Drummond et al. 2010). The sequences are deposited in GenBank under the accession numbers given in TABLE 1.

Sequence alignment and phylogenetic analysis

Sequences included in the phylogenetic analyses were either generated in this study (TABLE 1) or retrieved from GenBank. Multiple sequence alignments for ITS fragments were generated using MAFFT (Katoh et al., 2002) with default conditions for gap opening and gap extension penalty. The alignment was slightly edited using MEGA 5.0 (Tamura et al. 2011). Phylogenetic analyses were performed using both Bayesian Inference (BI)

TABLE 1. *Omphalina pyxidata* collections used in this study.

COLLECTIONS	COLL. NO., COUNTRY, DATE, COLLECTOR	ITS Acc. No.
* <i>O. pyxidata</i> 1	GT99398, France, 02/12/1999, G. Trichies	JQ671000
* <i>O. pyxidata</i> 2	EM0434-05, Switzerland, 06/07/2005, E. Musumeci	JQ671001
<i>O. pyxidata</i> 3	TO AV98, Italy, 10/10/2010, A. Vizzini	JN944402
<i>O. pyxidata</i> 4	Lamoure L66-118hl4, culture (see Lutzoni 1997)	OPU66450
* <i>O. pyxidata</i> var. <i>cystidiata</i>	TO HG2512, Italy, 11/11/2010, M. Curti	JQ671002

* = collections newly sequenced in this study.

and Maximum Likelihood (ML) approaches. The BI was performed with MrBayes 3.1.2 (Huelsenbeck & Ronquist 2001) with four incrementally heated simultaneous Monte Carlo Markov Chains (MCMC) ran over 10 millions generations, under GTR+ Γ evolutionary model. Trees were sampled every 1000 generations resulting in an overall sampling of 10,001 trees; the first 2500 trees were discarded as “burn-in” (25%). For the remaining trees, a majority rule consensus tree showing all compatible partitions was computed to obtain estimates for Bayesian Posterior Probabilities (BPP). Branch lengths were estimated as mean values over the sampled trees. ML was performed with RAxML (Stamatakis 2006) under GTRGAMMA model and using thorough bootstrap with 20 runs and 1000 replicates. In both analyses a *Pseudoclitocybe cyathiformis* sequence (Genbank HM191730) was used as outgroup taxon, according to Binder et al. (2010) and Vizzini et al. (2011a). The ML consensus tree was used merely for comparison with the Bayesian tree and to support the analysis. However the BPP (Bayesian posterior probability) over 0.75 and the ML bootstrap (MLB) over 50% values are reported in the resulting tree. Pairwise % identity values of ITS sequences were calculated using MEGA 5.0 (Tamura et al. 2011).

Results

Molecular results

Bayesian and Maximum likelihood inferences were performed on a total of 18 samples, including 15 sequences available from GenBank and three newly sequenced specimens (TABLE 1). Final alignment length was 731 bp. Both Bayesian and Maximum likelihood analyses produced the same topology (FIG. 1). In both analyses our cystidiate collection clusters with four *O. pyxidata* sequences clearly forming a strongly supported clade (BPP 0.92 and MLB 85%). The five *O. pyxidata* sequences show a pairwise % identity value of 99.8 and could be considered conspecific.

The *O. pyxidata* sequence deposited in Genbank under the accession number JF908502 (MCVE 15669) was not used in the analysis because a Blastn search

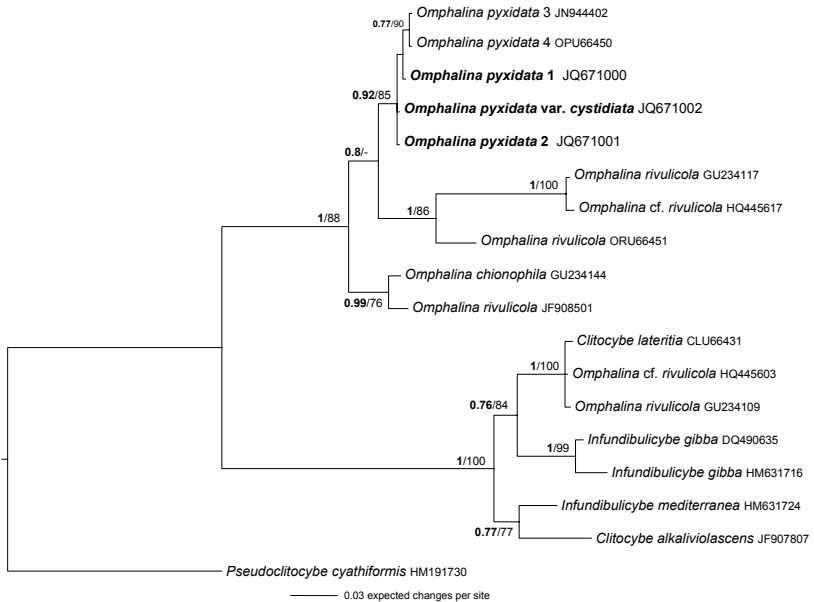


FIGURE 1. Bayesian phylogram obtained from the ITS sequence alignment. Support values for clades that are supported in either the Bayesian (Posterior Probabilities values – BPP) and Maximum likelihood (ML Bootstrap percentage – MLB) analyses are indicated. BPP above 0.75 and MLB above 50% are given above branches. Taxa in bold represent newly sequenced collections. Numbers (1–4) refer to the collections reported in TABLE I.

implies it represents a non-omphalinoid species in the *Pluteus cinereofuscus*/*P. nanus* complex.

Clitocybe lateritia, together with two *O. rivulicola* (J. Favre) Lamoure collections, clearly falls within *Infundibulicybe*. *Clitocybe alkaliviolascens* Bellù clusters close to *Infundibulicybe mediterranea* Vizzini et al.

Taxonomy

Omphalina pyxidata var. *cystidiata* M. Curti, Contu & Vizzini, var. nov. FIGS 2–3

MYCOBANK MB564484

A varietate typica differt praesentia cystidiorum in lamella, pileo, et stipite. Habitat: graegatim, ad muscos.

TYPE — Italy, Latium, prov. Rieti, com. Pozzaglia Sabina, Valle del Turano, 42.1596°N 12.9652°E, 878 m, 11.XI.2010, leg. M. Curti (TO HG2512 **holotype**).

ETYMOLOGY — the specific epithet refers to the presence of hymenial as well as of pileo- and caulocystidia.



FIGURE 2. *Omphalina pyxidata* var. *cystidiata*. Basidiomata. Bar = 20 mm.

PILEUS 3–15 mm broad, at first convex to applanate, later with a slightly depressed centre, not or slightly hygrophanous, minutely pubescent, dark brown-fulvous (Se 96, 121, 126, 146, 251, 252) fading to light brown (Se 133, 134, 173); margin at first involute then expanding, smooth to slightly striate in old basidiomes, whitish in dried basidiomes. LAMELLAE (L = 18–25; l = 1–2) decurrent, thick, distant with lamellulae occasionally interspersed, strongly forked and intervenose, whitish to light ochre (Se 199–200), with entire concolorous edge. STIPE 15–40 × 1–3 mm, cylindrical, slightly broadened at the base, sinuous-flexuous, solid then fistulous, cartilaginous, paler than pileus, light ochre-brown (Se 133, 174, 203–204), minutely pruinose to subglabrous. CONTEXT elastic, thin, whitish-ochre to light ochre-brown (Se 133, 134) in surface; smell and taste not distinctive. SPORE PRINT white.

SPORES (7–)7.5–9(–10) × 4.5–6(–6.5) μm , Q = 1.3–1.7, Qm = 1.51, largely ellipsoid to oblong in frontal view, lacrymoid to slightly amygdaliform in side-view, smooth, thin-walled, hyaline, inamyloid, mono-biguttulate, with a thick and evident hilum. BASIDIA 28–46(–60) × 6–9 μm , clavate, (two-) four-spored, sterigmata 3–7 μm long, usually clamped. CHEILOCYSTIDIA 23–45 × 3–6 × 2.5–5 μm , abundant, cylindrical, slender, apically tapered or slightly swollen, some laterally ventricose, hyaline, thin-walled, often with inner microguttules. PLEUROCYSTIDIA 25–50(–65) × 3.5–5.5 × 2.5–5(–7) μm , similar to cheilocystidia, but slightly longer. HYMENOPHORAL TRAMA irregular, consisting of 3.3–7 μm wide hyphae. PILEIPELLIS a cutis consisting of cylindrical, interwoven, 6–10

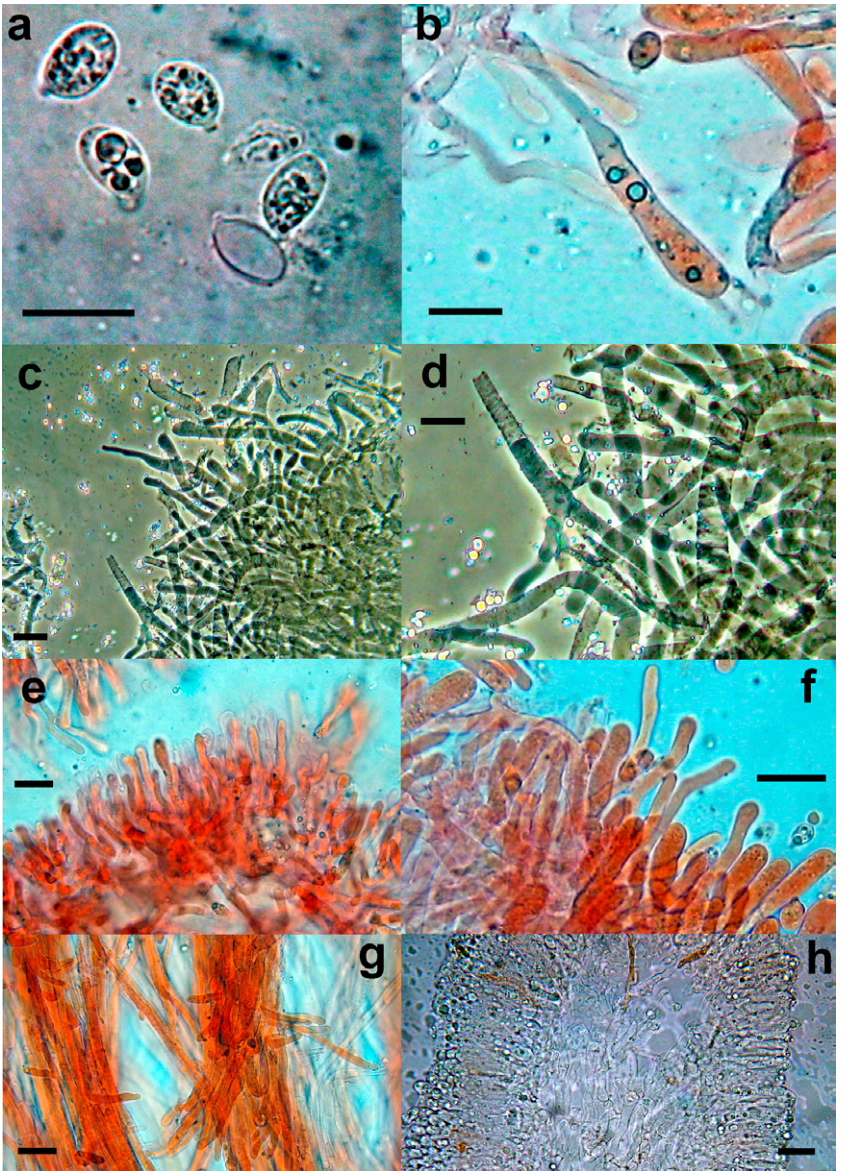


FIGURE 3. *Omphalina pyxidata* var. *cystidiata*. Microscopic features. a. Spores showing protruding hilum. b. Basidium with basal clamp. c. Pileipellis and pileocystidia. d. Encrusting pigment of pileipellis elements. e. Cheilocystidia (with inner microguttules). f. Pleurocystidia. g. Caulocystidia. h. Irregular hymenophoral trama. Bars: a–d = 10 µm; e–h = 20 µm.

µm wide hyphae; terminal elements with a subclavate 7–13 µm wide apex; subcutis made up of 3–7 µm wide hyphae; PIGMENT epiparietal, minutely to strongly encrusting; PILEOCYSTIDIA numerous, fusiform to lageniform, 35–60 × 3.5–9 × 3.5–5 µm. STIPITIPPELLIS a cutis made up, in 5% KOH, of ochre-yellow 3–8 µm wide hyphae bearing minute incrustations; CAULOCYSTIDIA similar to hymenial cystidia (30–65 × 4–5 × 3–5 µm). STIPITITRAMA non-sarcodimitic, consisting of hyphae up to 16 µm wide. CLAMP CONNECTIONS present at all septa. THROMBOPLEROUS HYPHAE (= oleiferous hyphae sensu Cléménçon 2004) present. APPRESSORIA on host surface not observed.

HABITAT — Gregarious, rarely subcaespitose, on *Tortula muralis* gametophytes (*Bryophyta*).

ADDITIONAL MATERIAL STUDIED — *Omphalina pyxidata* var. *pyxidata*: FRANCE, LORRAINE, MOSELLE, Moyeuve-Petite, 02 Dec 1999, on mosses, leg. G. Trichies (GT99398); SWITZERLAND, CANTON BASEL STADT, Lange Erlen, 06 Jul 2005, on mosses, 250 m asl, leg. E. Musumeci (EM0434-05); ITALY, PIEDMONT, CHISONE VALLEY, Pinerolo, Prà Martino, 10 Oct 2010, on mosses, 1000 m asl, leg. A. Vizzini (TO AV98).

Discussion

As highlighted by Vizzini et al. (2011a), *Omphalina* s.s. is sister to *Infundibulicybe* Harmaja, a genus recently segregated from *Clitocybe* (Fr.) Staude (Harmaja 2003). In the six-gene region sequence analyses by Matheny & al. (2006), *Infundibulicybe* falls outside the tricholomatoid clade, occupying an isolated position in the *Agaricales* (incertae sedis). However, recent analyses by Binder et al. (2010), Vizzini et al. (2011a) and Matheny (pers. comm.) place it with robust support as sister to the rest of the tricholomatoid clade. *Omphalina* and *Infundibulicybe* share the cream-reddish brown tinges of pileus and stipe, the long-decurrent lamellae, and strongly encrusting pigment (Harmaja 2003, Vizzini et al. 2011b).

Our molecular analysis (FIG. 1) and morphological comparison support the new taxon as a cystidiate variant of *O. pyxidata* thus far never described. *Omphalina pyxidata* var. *cystidiata* is unique within *Omphalina* s.s., where it is the only taxon thus far included that possesses well-developed cheilo-, pleuro-, pileo-, and caulocystidia (Bigelow 1970, 1985, Singer 1970, Lamoure 1974, 1975 1982; Cléménçon 1982, Bon 1997, Elborne 2008). Furthermore, we have found no cystidia in three examined European collections of *O. pyxidata* (see ADDITIONAL MATERIAL STUDIED). Morphologically, our new taxon resembles the omphalinoid bryophilous genera belonging to the hymenochaetoid clade (Redhead et al. 2002b, Antonín & Noordeloos 2004). However, *Rickenella* differs in having a regular hymenophoral trama (Contu 1997, Redhead et al. 2002b) and *Blasiphalia* by growing specifically on the *Blasia pusilla* gametophyte (*Marchantiophyta*) and forming appressoria on host surface (Larsson et al.

2006). Apart from molecular data that clearly support our new variety within the *Agaricales*, morphologically *Contumyces* (= *Jacobia* Contu, nom. illeg.; *Hymenochaetales*) might accommodate our taxon. *Contumyces* so far comprises only three species — *C. rosellus* (M.M. Moser) Redhead et al., *C. vesuvianus* (F. Brig.) Redhead et al., and *C. brunneolilacinus* (Contu et al.) Redhead et al. (Redhead et al. 2002b, Antonín & Noordeloos 2004) — but only the first named has been sequenced. *Contumyces rosellus* differs from *Omphalina pyxidata* var. *cystidiata* by the pink to pale salmon-pink colouration, white stipe, and longer and larger cystidia. *Contumyces vesuvianus* has orange to fulvous basidiomata, larger basidiospores, rarer differently shaped (mucronate/rostrate) cystidia and intracellular pigment in the pileipellis hyphae. As originally described (Contu 1989), *C. brunneolilacinus* differs in having a lilaceous-brown, sharply tomentose, hygrophanous pileus, pink-lilac lamellae, a *Pelargonium*-like smell, lageno-capitate cystidia, and intracellular pigment in the pileipellis hyphae. Finally, *C. brunneolilacinus* sensu Antonín & Noordeloos (2004) differs in having longer and larger cystidia and a non-tomentose pileal surface.

Our sequence analyses have also revealed that *Clitocybe lateritia* clusters in *Infundibulicybe* with two collections clearly misdetermined as *O. rivulicola* (FIG. 1). The three sequences are clearly conspecific (pairwise % identity value = 99.7). The two “*O. rivulicola*” sequences were derived from soil or ectomycorrhizal samples sampled from *Dryas octopetala* beds, as reported in two recent papers dealing with alpine-arctic fungi (Bjorbækmo et al. 2010, Geml et al. 2012). *Clitocybe lateritia*, a rare alpine-arctic species strictly associated with *Dryas octopetala* (*Rosaceae*), should be transferred to *Infundibulicybe* based on both sequence analyses and its *Infundibulicybe*-like features (reddish-brown colouration, strongly encrusting pigment in the pileipellis, partly lacrymoid spores) as inferred from Favre (1955), Gulden & Jennsen (1988), and Bon (1997). Consequently, we propose the following new combination:

Infundibulicybe lateritia (J. Favre) Vizzini & Contu, **comb. nov.**

MYCOBANK MB564485

= *Clitocybe lateritia* J. Favre, *Ergebn. wiss. Unters. schweiz.*

Natn. Parks, n.s. 5(33): 54, 199 (1955).

TYPE: Switzerland, Munt la Schera, 2400 m a.s.l., in *Dryas* beds, 21.08.1941, leg. J. Favre (Lectotype designated here, G-K16433; specimen illustrated in Favre 1955: Fig. 34; Pl. IV, Fig. 11).

Finally, *Clitocybe alkaliviolascens*, a species of the *Infundibulicybe gibba* complex recently described from Mediterranean areas and characterized by a dark brown pileus surface turning pink-violet with a 30% KOH drop, and spores reaching 10 µm in length (Bellù 1995, 1996), also belongs in *Infundibulicybe*. Bellù will propose the new combination in a forthcoming paper (Bellù, pers. comm.).

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