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OF RESEARCH ON TAXONOMY & NOMENCLATURE OF FUNGI & LICHENS

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MARCELLE LE GAL: A REMINISCENCE

RICHARD P. KORF

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Ithaca, New York 14853 USA

Marcelle Le Gal, born Marcelle Choquart on February 14th, 1895 at Amiens, France, died in the same city on June 23, 1979. Her influence on mycology, and particularly on the development of the taxonomy of discomycetes, has been immense, even though she only entered the field when she was nearly 40 years of age. Her early publications dealt primarily with discomycetes and showed an exceptional attention to detail and clear deliniation of characters such that I, young and wholly inexperienced, immediately accepted her publications as authoritative and as models of procedure. My own work on discomycetes had only begun in 1944, and in 1947 she sent me her monumental "Recherches sur les ornementations sporales des discomycètes operculés," her just-published doctorate thesis (which was to receive the French Academy's coveted "Prix Montagne."). I was quickly drawn into a lively correspondence with her, and despite the difference in our ages, she cheerfully commented upon and examined problem discomycetes that I sent her over the years, never stinting of her time and advice.

It was 1949 before we were to meet in person, on the occasion of a trip to European herbaria when I was searching for type specimens of species of the Arachnopezizeae for my own thesis studies. At the Laboratoire de Cryptogamie of the Muséum National d'Histoire Naturelle in Paris I was cordially greeted by Professor Roger Heim and his staff, but above all by Mme Le Gal herself. The memory is one I shall ever cherish, for she devoted time and energy to showing me her collections, and the museum's discomycete holdings, including those of her idol (and mine), the eminent Émile Boudier. Quiet-spoken, ever helpful, Marcelle Le Gal was from that moment on to be a major influence in my professional life. Her devotion to the study of fungi,

her insight and mastery of the fungi she knew, remain an inspiration to all who knew her. An unexpected surprise to me as a young graduate student was that she entertained me with a sumptuous meal in her Paris apartment, where I was to meet her greatest joy, her husband, Étienne. They had married in 1922 on her return from the United States where she had earned her M.A. at Columbia University.

On several later occasions I was able again to partake of Mme Le Gal's hospitality and kindness during visits to Paris. The last of these was sad beyond any expectation, for when I visited her in 1973 her beloved husband had only recently died. Her grief had caused her to withdraw from duties at the museum to her home in Amiens, and to stop work on the important monograph of the genus Scutellinia that had been occupying her for over a decade. When she learned that I was to visit the Paris museum, she made the unprecedented effort of coming to the museum to talk with me about our taxonomic problems, and offered me the loan of several of her specimens of a critical genus we discussed. Her sadness over the loss of her husband was profound, and had clearly devastated her emotionally, affecting both her personal and professional life. She was soon to turn her back on mycology, perhaps even on humanity, as the result of an unfortunate accident she suffered in Amiens and of her grief over the loss of her constant companion-husband. Mycology's great loss is that her Scutellinia monograph will apparently remain unpublished and incomplete.

Marcelle Le Gal is one of the very few to have markedly affected my development as a scientist. From her I learned patience and devotion to detailed study of minute structures as keys to relationships. Not always was I to be in her good graces; particularly because of my insistence on following the Code of Nomenclature and the principle of priority, I became a "Peck's bad boy" in her view. For years I licked the wounds inflicted by her vitreolic attack on "le jeune mycologue américain" in her masterful diatribe against the Code of Nomenclature (Le Gal, 1958). That she eventually forgave my youthful exuberance was one of my great joys.

I shall miss Marcelle Le Gal's cautionary council more than many other mycologists will; our interests coincided closely, and it is in her footsteps that I have trod most often, secure in the knowledge that her obser-



MARCELLE LE GAL (1895-1979)

Photographed by the author in the gardens outside the
Muséum National d'Histoire Naturelle, Paris, during
the 1954 International Botanical Congress

vations were dependable and her sense of taxonomic direction unerring. A kinder, more loving spirit has never before, perhaps, graced mycology.

For those who read French, I commend a touching tribute to her by Patrick Joly (1980), that captures the essence of this most wondrous scientist. Yet another revealing tribute to her contains a complete listing of her mycological papers, the article by her long-time associate in the field and laboratory, Henri Romagnesi (1980).

REFERENCES CITED

- JOLY, P. 1980. Marcelle Le Gal (1885-1979). *Cryptogamie, Mycol.* 1: 93-96.
- LE GAL, M. 1958. Petite promenade à travers le maquis de la nomenclature. *Rev. Mycol. (Paris)* 23: 121-126.
- ROMAGNESI, H. 1980. Mme Marcelle Le Gal (1895-1979). *Bull. Soc. Mycol. France* 96: 125-131.

MYCOTAXON

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April-June 1981

DISCOMYCETES EXSICCATI, FASC. IV

RICHARD P. KORF AND SUSAN C. GRUFF

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Ithaca, New York 14853 USA*

Twenty-five additional numbers comprise this fourth fascicle of *Discomycetes Exsiccati* [for distribution of sets, see the note on page 15].

We express particular thanks to Dr. Henry Dissing and to Dr. Sigmund Sivertsen for providing us with holotype material of *Boudiera dennisii* Diss. & Siv. in Dissing to issue here as ISOTYPES (No. 77); a second collection of this species was collected by the senior author of this paper in their company and constitutes AUTHENTIC material from a previously unreported locality (No. 78).

As previously noted (Dissing & Korf, 1980), No. 84 in this fascicle constitutes ISONEOTYPE material of *Ruhlandiella berolinensis* Hennings, the type material of which is presumed to have been destroyed in Berlin during the Second World War.

An examination of Rehm's type specimen of *Humaria gregaria* Rehm showed that it has warted ascospores, not smooth as is usually assumed. A smooth-spored variant is issued here (No. 90) as *Trichophaea gregaria* (Rehm) Boud. f. *laevispora* Korf & Gruff, f. nov., the specimens issued being ISOTYPES:

Trichophaeae gregariae f. *gregariae similis*, sed ascosporis perfecte laevibus differt. HOLOTYPE: CUP-MJ 611 (ISOTYPI in Korf & Gruff, Disc. Exs. 90 dispersi). PARATYPE: CUP-MJ 41. On clay bank. Cinchona Botanical Garden, elev. 4750 ft., St. Thomas Parish, Jamaica. Leg. R.P.Korf, leader; J.R.Dixon, K.P.Dumont, R.W.Erb, D.H.Pfister, D.R.Reynolds, A.Y.Rossman & G.J.Samuels. 8.I.1971.

The typical, warted-spored form is also issued (No. 89) for comparison, as *Trichophaea gregaria* f. *gregaria*.

By far the most abundant of Durand's collections of *Sarcosoma cyttarioides* Rehm in Dur., his No. 1305 (= CUP-A 12278), is

formally designated here as the LECTOTYPE of that species, and the specimens issued (No. 79) as *Plectania cyttarioides* (Rehm in Dur.) Korf are thus ISOLECTOTYPES. [All packets of this collection in the Durand (CUP-D) and Atkinson (CUP-A) herbaria in CUP bear the data for collecting site as "Glen Mary," while the original description (Durand, 1903) states, "Most abundant in Glen Burney." This latter name does not appear on any of the *Sarcosoma* packets in these herbaria.]

Mr. G. Beaton, of Australia, who has consistently provided us with exciting discomycete finds, has generously given us TOPOTYPE material of the elegant species *Underwoodia beatonii* Rifai for issue here as No. 91. Dr. Peter Milan Petersen, of the University of Copenhagen, has contributed the material from Greenland of *Sarcoleotia globosa* (Sommerf. : Fr.) Korf issued as No. 100.

Taxonomic and nomenclatural notes again appear on some of the labels: (79) on *Plectania cyttarioides* as the type species of *Plectania* sect. *Plicosporae*; (87) on Karsten's use of the combination *Scutellinia scutellata*, which while the earliest is not validly published since the generic name was not yet validly published; (89) on Karsten's similarly first use of the combination *Sepultaria gregaria*, likewise not validly published since its generic name had not yet been validly proposed, and on the taxonomic synonymy of *Trichophaea gregaria* var. *intermedia* with *T. gregaria* f. *gregaria*; (95) on an aberrant collection or unnamed variant of *Arachnopeziza cornuta* having only 1-celled ascospores; (96) on adoption of *Arachnopeziza leonina* instead of *A. candido-fulva*; (97) on discarding the group "Anomalae" of *Arachnopeziza* since the "granulations" on the hairs of *A. trabinelloides* (sole representative of that group) seem to be more like the resinous excretions sometimes evident in the group "Typicae" than like the granulations on the hairs in *Dasyscyphus*.

ACKNOWLEDGEMENTS

Particular thanks are due the US National Science Foundation, which under a series of grants to the senior author has funded the collecting in Jamaica and the Canary Islands, and to the Fulbright Commission that supported the collecting in Japan. The Okinawa collection was by virtue of support from the US Office of Education, the US Civil Administration of the Ryukyu Islands, and the University of the Ryukyus. Collections from Finland and Norway were financed by the University of Copenhagen and the University of Trondheim. Dr. William Dress, of the Bailey Hortorium, Cornell University, kindly checked the Latin diagnosis.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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76. *Anthracobia macrocystis* (Cooke) Boudier, Hist. classific. discomyc. Europe, p. 65. 1907.

On a burnt *Eucalyptus* sp. clear-cut.

At km mark 14, Bosque de la Esperanza, near Pico de las Flores, Tenerife, Canary Islands.

Leg: R.P.K., W.C.Denison, L.M.Kohn & M.A.Sherwood 8.I.1976
Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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77. *Boudiera dennisii* Dissing & Sivertsen in Dissing, Kew Bull. 31: 755. 1977. ISOTYPE

Near a small stream, Nordland, between Berget and Fisksjømoen, 22 km NW of Mo in Rana, Norway.

Leg: H.Dissing, S.Sivertsen & T.Schumacher 6.IX.1975
Det: H.Dissing & S.Sivertsen

DISCOMYCETES EXSICCATI

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78. *Boudiera dennisii* Dissing & Sivertsen in Dissing, Kew Bull. 31: 755. 1977. AUTHENTIC

On mud flat at mouth of brook, among *Equisetum*, *Calamagrostis neglecta*, and *Lamprospora ovalispora*.

Loevvaiäkka, near Levajok, along Tana River, Finnmark Fylke, Norway.

Leg: S.Sivertsen, H.Dissing & R.P.K. 21.VIII.1978
Det: S.Sivertsen & H.Dissing

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79. *Plectania cyttarioides* (Rehm in Durand) Korf, Mycologia
49: 110. 1957. [ISOLECTOTYPE of *Sarcosoma cyttarioides*
Rehm in Durand]

On dead twigs, leaves, etc.

Blue Ridge Mountains, Glen Mary, Blowing Rock, North
Carolina.

NOTES: This is the type species of *Plectania* Fuckel sect. *Plicisporae* Korf (*loc. cit.*). R.P.K.

Leg: E.J.Durand (1305)

30.VIII.1901

Det: E.J.D.

DISCOMYCETES EXSICCATI

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80. *Pseudoplectania nigrella* (Pers. : Fr.) Fuckel,
Jahrb. Nassauischen Vereins Naturk. 23-24: 324. 1870.

In clay soil on roots of *Dicranopteris linearis* and rarely
of *Pleioblastus linearis*.

University of Ryukyus Recreation Area, Oku, Kumigami-son,
Okinawa, Japan.

Leg: R.P.K., K.P.Dumont, Z.Koja & K.Kuroshima

17.III.1966

Det: R.P.K.

DISCOMYCETES EXSICCATI

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81. *Ptychoverpa bohémica* (Kromb.) Boudier, Hist. classif.
discomyc. Europe, p. 34. 1907.

On ground in mixed woods.

Lloyd-Cornell Preserve, McLean, New York.

Leg: Mycology Class

2.V.1955

Det: R.P.K.

DISCOMYCETES EXSICCATI

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82. *Pulvinula globifera* (Berk. & Curt. in Berk.) Le Gal,
Prodr. Flore Mycol. Madagascar 4: 94. 1953.

On soil.

At 1860 ft. elev. in rain forest, El Yunque, Puerto Rico.

Leg: G.Abawi

27.I.1969

Det: R.P.K.

DISCOMYCETES EXSICCATI

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83. *Pulvinula niveo-alba* Moravec, Česká Mykol. 23: 231.
1969.

On duff.

Juuma-Jäkälävuoma, Kuusamo, Finland.

Leg: S.Sivertsen

25.VIII.1978

Det: S.S.

DISCOMYCETES EXSICCATI

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84. *Ruhlandiella berolinensis* Hennings, Hedwigia 42: (24).
1903. ISONEOTYPE, designated by Dissing & Korf,
Mycotaxon 12: 295. 1980.

On soil.

Burn site along road in *Eucalyptus* grove at km mark 14,
Bosque de la Esperanza, Tenerife, Canary Islands.

Leg: R.P.K., R.Fogel, G.L.Hennebert & L.M.Kohn 29.XII.1976

Det: H.Dissing & R.P.K.

DISCOMYCETES EXSICCATI

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- 85.
- Scutellinia erinaceus*
- (Pers. : Fr.) O. Kuntze, Revis.
-
- gen. pl. 2: 869. 1891.

On mossy wood.

Michigan Hollow, Danby, New York.

Leg: I.J.Gamundí, K.S.Thind, W.C.Denison, 9.VIII.1960
R.T.Moore, V.P.Tewari & R.P.K.

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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- 86.
- Scutellinia pennsylvanica*
- (Seaver) Denison, Mycologia
-
- 51: 619. 1961 ('1959').

On old wood.

"Big Woods," SW of Ann Arbor, Michigan.

Leg: L.E.Wehmeyer, R.P.K. & al. 19.VII.1948
Det: W.C.Denison

DISCOMYCETES EXSICCATI

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- 87.
- Scutellinia scutellata*
- (L. : Fr.) Lambotte, Fl. mycol.
-
- belge, Suppl. 1: 299. 1887.

On *Fagus* sp.

Lloyd-Cornell Preserve, Slaterville, New York.

NOTES: Neither a generic name "*Scutellinia* (Cooke) Karst." nor the combination
"*S. scutellata* (L.) Cooke" was validly published by Karsten (Medd. Soc. Fauna
Fl. Fenn. 11: 145. 1884) [ICBN, Arts. 41, 43]. R.P.K.Leg: M.A.Rosinski II.1952
Det: W.C.Denison

DISCOMYCETES EXSICCATI

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88. *Scutellinia umbrosum* (Fr.) Lambotte, Fl. mycol. belge, Suppl. 1: 300. 1887.

On soil.

Montezuma Wildlife Preserve, Seneca County, New York.

Leg: I. Müller & R.P.K.

20.VI.1962

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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89. *Trichophaea gregaria* (Rehm) Boud. f. *gregaria*, Hist. classif. discomyc. Europe, p. 60. 1907.

On soil and rotting plant matter.

Peck Foray, Ashokan, New York.

NOTES: Neither a generic name "*Sepultaria* (Cooke) Karst." nor the combination "*S. gregaria* (Rehm) Karst." was validly published by Karsten (Medd. Soc. Fauna Fl. Fenn. 11: 145. 1884) [ICBN, Arts. 41, 43]. Le Gal's *T. gregaria* var. *intermedia* Le Gal [Rev. Mycol. (Paris) 2: 214. 1937] is a synonym, since Rehm's type specimen also has warted ascospores, not smooth as she assumed. R.P.K.

Leg: D. Malloch, R.P.K. & al.

10.IX.1972

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
RICHARD P. KORF & SUSAN C. GRUFF, EDITORS

90. *Trichophaea gregaria* (Rehm) Boud. f. *laevispora* Korf & Gruff, Mycotaxon 13: 5. 1981. ISOTYPE

On clay soil bank.

Trail between Freetown and Wag Water River, near Hardwar Gap, St. Andrew Parish, Jamaica.

Leg: R.P.K., J.R. Dixon, K.P. Dumont, R.W. Erb,
D.H. Pfister, D.R. Reynolds, A.Y. Rossman
& G.J. Samuels

18.I.1971

Det: R.P.K. & S.C.G.

DISCOMYCETES EXSICCATIDEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
RICHARD P. KORF & SUSAN C. GRUFF, EDITORS

91. *Underwoodia beatonii* Rifai, Verh. Kon. Ned. Akad. Wetensch., Afd. Natuurk., Tweede Sect., 57(3):.69. 1968. TOPOTYPE

In sandy soil and debris under *Melaleuca lanceolata* beside cliff walk, Anglesea camping ground, Australia.

Leg: G. Beaton

23.VIII.1968

Det: G.B.

DISCOMYCETES EXSICCATIDEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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92. *Urmula craterium* (Schw. : Fr.) Fr., Summa veg. Scand., sect. post., p. 364. 1849.

On soil and wood.

Woods near Westhaven Road, Town of Ithaca, Tompkins County, New York.

Leg: R.J. & K. Smith, K.T.Korf & R.P.K.

V.1960

Det: R.P.K.

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93. *Wynnea americana* Thaxter, Bot. Gaz. 39: 246. 1905.

On soil in woods.

2.5 miles east of Obi, at crest of hill, elev. 2200 ft., Alleghany County, New York.

Leg: G. & M. Smith, J. Bluhm, L.M. Kohn & R.P.K.

2.IX.1976

Det: G. & M.S.

DISCOMYCETES EXSICCATI

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RICHARD P. KORF & SUSAN C. GRUFF, EDITORS

94. *Arachnopeziza aurelia* (Pers. : Fr.) Fuckel, Jahrb.
Nassauischen Vereins Naturk. 23-24: 303. 1870.

On bark and acorns of *Quercus* sp.

France Brook, Alleghany State Park, New York.

Leg: R.P.K. & al.

9.VI.1961

Det: R.P.K.

DISCOMYCETES EXSICCATI

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95. *Arachnopeziza cornuta* (Ellis) Korf, Lloydia 14: 158.
1952 ('1951'). [var.?]

On rotted wood.

Lloyd-Cornell Preserve, Ringwood, New York.

NOTES: Apparently lacking the 2- and 3-septate ascospores by which one usually recognizes this species. R.P.K.

Leg: G.L.Hennebert (3096) & R.P.K.

17.V.1962

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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96. *Arachnopeziza leonina* (Schw. : Fr.) Dennis, Kew Bull.
17: 351. 1963.

On a very rotted log.

East end of Lake Shikotsu, Ibari Pref., Hokkaido, Japan.

NOTES: This specimen was reported by Korf (Bull. Natl. Sci. Mus. 4: 392. 1959) as *A. candido-fulva* (Schw.) Korf. Dennis's transfer is correct, since Schweinitz described *Peziza leonina* earlier than *Peziza candido-fulva*, and Fries accepted both. R.P.K.

Leg: S.Imai, S.Kamei, Y.Otani, R.P.K. & al.

20.V.1958

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
RICHARD P. KORF & SUSAN C. GRUFF, EDITORS

- 97.
- Arachnopeziza trabinelloides*
- (Rehm) Korf, Lloydia 14:
-
169. 1952 ('1951').

On wood.

Margaretville, Delaware County, New York.

NOTES: The "granulations" on the hairs are more like resinous excretions than the granulations of *Dasyocypha* spp. of the "Typicae" group. This species belongs close to *A. cornuta* in *Arachnopeziza* group "Typicae," and should not be segregated in a group "Anomaliae" as was done by Korf (*loc. cit.*). R.P.K.

Leg: R.T.Pennoyer & R.P.K.

9.V.1959

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
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- 98.
- Parachnopeziza miniopsis*
- (Ellis) Korf, Mycotaxon 7:
-
469. 1978.

On bark.

France Brook, Alleghany State Park, New York.

Leg: R.P.K. & al.

9.VI.1961

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
RICHARD P. KORF & SUSAN C. GRUFF, EDITORS

- 99.
- Pezoloma laricina*
- (Ell. & Everh.) Korf, Phytologia 21:
-
205. 1971.

On duff, mostly *Tsuga* needles and cones.

Peck Foray XIV, Twin Valleys, Lewis, New York.

Leg: G.Abawi, D.H.Pfister & R.P.K.

15.VI.1968

Det: R.P.K.

DISCOMYCETES EXSICCATI

DEPARTMENT OF PLANT PATHOLOGY, CORNELL UNIVERSITY
RICHARD P. KORF & SUSAN C. GRUFF, EDITORS100. *Sarcoleotia globosa* (Sommerf.: Fr.) Korf, Phytologia
21: 206. 1971.On *Sphagnum fuscum*.

Sydprøven (60°21'N, 45°34'W), Greenland.

Leg: P.Milan Petersen (71.162)

30.IX.1971

Det: R.P.K.

DISTRIBUTION OF SETS

The first fascicle of this series was deposited in 11 institutions (Korf, 1958), with 2 additional ones listed by Korf and Gruff (1978) as receiving fascicles II and III. By error they listed "BP = Museum of Natural History, Budapest," whereas the set is actually on deposit at BPU = Institute of Plant Taxonomy and Ecology of Eötvös L. University. A 14th set is now on deposit at Copenhagen, also beginning with fasc. II. The complete list of deposit herbaria is thus:

BPI (Beltsville)	FH (Cambridge)*	NY (New York)
BPU (Budapest)*	K (Kew)	PC (Paris)
C (Copenhagen)*	LAH (Lahore)	PR (Prague)
CUP (Ithaca)	MICH (Ann Arbor)	UC (Berkeley)
DAOM (Ottawa)		UPS (Uppsala)

*lacking fascicle I.

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ALTERNARIA THEMES AND VARIATIONS

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Amherst, Massachusetts 01003

PROLOGUS

Being the initiation of a series of observations on *Alternaria* species, in this instance on some whose nonbeaked conidia very rarely occur in chains (*A. molesta* sp. nov., *A. mouchaccae* nom. nov.) and others whose erostrate conidia proliferate in chains by means of interpolated conidiophores (*A. chlamydospora*, *A. limaciformis* sp. nov., and two commonly misrepresented phytopathogens, *A. triticina* and *A. longipes*).

I. EROSTRATAE, SOLITARIAE

Most species of *Alternaria* produce conidia in chains, some readily and abundantly under conditions of either nature or culture (e. g., *A. tenuissima* (Kunze in Nees & Nees ex Persoon) Wiltshire), others sparsely and perhaps only tardily after manipulation of culture medium, light quality, and temperature. This latter reluctantly catenulate group includes, among others, common phytopathogens whose conidia are thin walled and filamentously rostrate (e. g., *A. zinniae* M. B. Ellis) as well as others whose conidia are relatively thick walled and erostrate (e. g., *A. radicina* Meier, Drechsler & Eddy).

Alternaria species of a third group produce two conidia in a chain so rarely, under any condition reported or observed thus far, that any chain can be considered little more than an aberration. Two notable members of this group are *A. chrysanthemi* Simmons & Crosier (Simmons 1965) and *A. helianthi* (Hansford) Tubaki & Nishihara (1969). Both of these species are phytopathogens; the nonbeaked conidia of both are relatively large for the genus, with lengths up to 120 μ m and 110 μ m respectively (Fig. 1).

An additional member of this group with erostrate, solitary conidia is known in the form of a small-spored species isolated from a skin lesion of a captive harbour porpoise (by S. Andersen, Odense, Denmark; transmitted by G. A. de Vries, Baarn, The Netherlands, as CBS M 339.77). It is reported here as novel on the basis of examination of axenic cultures, primarily on potato-carrot agar, hay extract agar, and 20% V-8 juice agar (hereafter as PCA, Hay, V-8; see Stevens 1974), in a diurnal fluorescent light/dark cycle, at 22 C.

1. *Alternaria molesta* Simmons, sp. nov. (Fig. 1)
 [molesta = troublesome (to both *Phocaena* and Simmons, though in different ways)]

*Ex culturis in agaro PCA descripta. Coloniae griseo-brunneae, lanuginosae, postea atrobrunneae, applanatae. Mycelium initio ex hyphis septatis, ramosis, subhyalinis vel pallide stramineis, levibus, 3.5-5.0µm crassis compositum. Conidiophora simplicia, singulatim ex lateribus hypharum oriunda, recta vel flexuosa, cylindrica, pallide flavo-brunnea, levia, 0-10 septata, plerumque 4-5µm crassa, usque ad 100 (plerumque 5-20)µm longa, apice rotundato et uniporoso, raro geniculata. Conidia solitaria, subcylindrica, ovoidea, obclavata vel ellipsoidea, septis transversalibus plerumque 3-6, longitudinalibus nullis vel paucis, ad septa definite constricta, pallide straminea, levia, 15-38 x 7-12 (plerumque 27 x 12)µm. Origo typi: ex laesione pellis *Phocaena phocaenae*, Odense, Daniae. Typus: partes ex Simmons 32-075 (CBS M 339.77) desiccatae et in BPI, CBS, DAOM, IMI, NY conservandae.*

The type isolate is stable and predictable in its growth and sporulation characteristics under specified culture conditions. It grows well on PCA and V-8 (15mm radially in 4 da), producing colonies that are dark olive-brown with a shallow aerial layer of paler woolly hyphae. Conidium production occurs initially on hyphae near the substrate surface but spreads gradually to the intertwined aerial elements. After the agar surface of an entire culture plate has been overgrown, the colony becomes almost opaque because of the very dark color of both the submerged and the aerial hyphae. Although the mycelium of colonies on Hay is scarcely visible, it gives rise to conidia readily and abundantly.

Conidia of *A. molesta* are produced one per conidiophore tip; conidiophore tip proliferation (geniculation) is uncommon. Among many hundreds of conidia viewed in slide

preparations, only half-a-dozen had an apical 1-cell extension with a terminal scar, and thus presumably had served as the basal unit in a chain of two conidia. The color of conidium and conidiophore walls is so dilute that septa and disjunction scars appear in bold contrast in transmitted light. A very high percentage of conidia mature as ellipsoid to ovoid units with three transverse septa and at a size of about $27 \times 12 \mu\text{m}$. The conidium population is, for an *Alternaria* species, remarkably restricted in morphological variation, with very little tendency to produce longitudinally asymmetrical or excessively cellular entities. Fig. 1 illustrates conidiophores and a representative range of *A. molesta* conidium shape, size, and septation. Conidium outlines for *A. chrysanthemi* (Fig. 1, lower left) and *A. helianthi* (Fig. 1, lower right) are included for purposes of comparison.

2. *Alternaria mouchaccae* (Fig. 2)

Arid and semidesert soils harbor remarkable numbers of phaeodictyosporic hyphomycete species (Durrell & Shields 1960; Kuehn 1960; Ranzoni 1968; States 1978). New taxa almost inevitably have been unearthed and described in *Ulocladium*, *Embellisia*, and *Alternaria*. Pertinent to the present discussion is a species published as *Ulocladium chlamydosporum* Mouchacca (1971), which was described as new on the basis of several isolates from arid and newly cultivated soil of the New Valley region of Egypt. A second dematiaceous entity, often associated with *U. chlamydosporum* in nature and morphologically similar to it, later was published as a new species *Alternaria chlamydospora* Mouchacca (1973). Study of representative strains of these two species (IMI 156434 and IMI 156427 respectively; EGS 31-061 and 31-060) reveals that they are congeneric in *Alternaria*. Their morphological similarity and natural association in the soil tempts one to speculate that they also may be conspecific although culturally in different phases of progressive degeneration (a condition noted by Mouchacca, 1971). However, relying to the greatest extent possible on comparisons of 1-conidium reisolates on PCA and Hay agars, of conidiogenesis, and of conidium development before typical cell distortion begins (Figs. 2 & 3), I conclude that there are two distinguishable species, *Alternaria chlamydospora* Mouchacca (discussed below as species no. 3) and *Alternaria mouchaccae* Simmons, *nom. nov.* [\equiv *Ulocladium chlamydosporum* Mouchacca, *Revue de Mycologie* 36(2): 120, 1971; *non A. chlamydospora* Mouchacca, 1973].

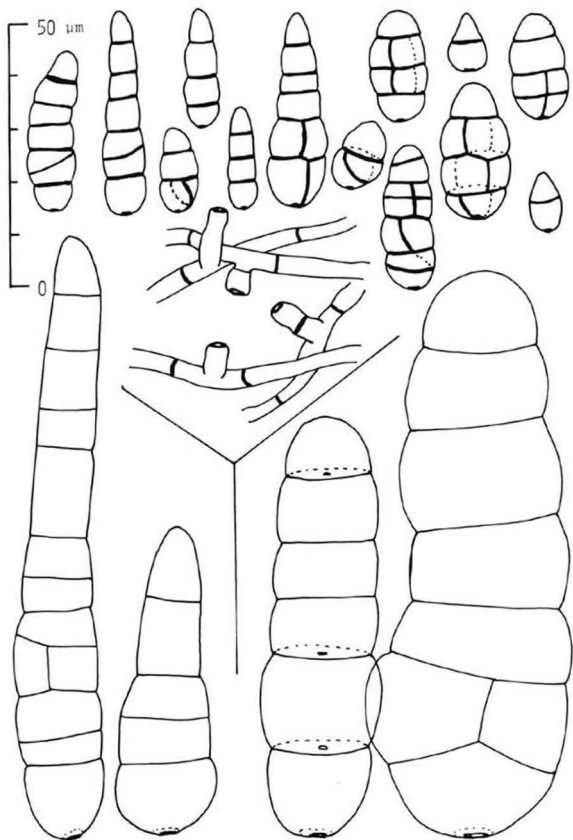


Fig. 1. *Alternaria molesta* ex Type (at top), from culture on PCA; *A. chrysanthemi* (lower left); *A. helianthi* (lower right).

A. mouchaccae is yet another example of a species that produces nonbeaked solitary conidia when grown on media of low nutrient content. False conidial beaks (secondary conidiophores) are extremely rare under such conditions; only two were found among hundreds of conidia examined. A few young conidium initials might be misinterpreted as ulocladoid on the basis of a narrow isthmian attachment to the conidiophore. However, most young conidia are spherical or ovoid from the beginning and have bases in close contact with conidiogenous cells, which themselves exhibit conspicuously pigmented areas around the terminal disjunction scars — characters more nearly typical of *Alternaria* than of *Ulocladium*. Conidium basal scars appear always to have been pushed into an eccentric position following the division of any basal cell by a vertical or oblique septum. Enlarging conidia often become conspicuously swollen and distorted, especially when produced in colonies on V-8 agar. The strain studied, "*cultura typica*" (Mouchacca 1971), exhibits abundant nonsporulating hyphal overgrowth, although conidium production on nonaerial mycelium remains good as long as only conidia are used as inoculum in serial transfers.

Considerable experience with moderately unstable isolates of this sort suggests that hyphal chlamydospores, stromatic cellular masses, and progressive degeneration originally described for this material, and still present, may be characteristic of the typical culture rather than of the species *in toto*, and that focusing attention on the morphology of conidia before they become distorted may permit recognition of *A. mouchaccae* among future isolates, whether or not accessory structures are present.

II. EROSTRATAE, PROLIFICANTES

The character "conidium with beak," i. e., a conidium with an apical portion morphologically distinguishable from the septate sporebody, strongly influences the commonly held generic concept of *Alternaria*. Neergaard (1945) discussed the nature of the conidial beak, emphasizing its species-diagnosis utility and making a strong point of differentiating between "true beak (eurostrum)" as an integral part of the conidium and "false beak (pseudorostrum), formed like the conidiophore of the species concerned." The nature of the *Alternaria* false beak remains poorly understood, however, if we judge on the basis of much of the available descriptive literature and posited ranges of conidium length. Evaluation of this literature, a complex

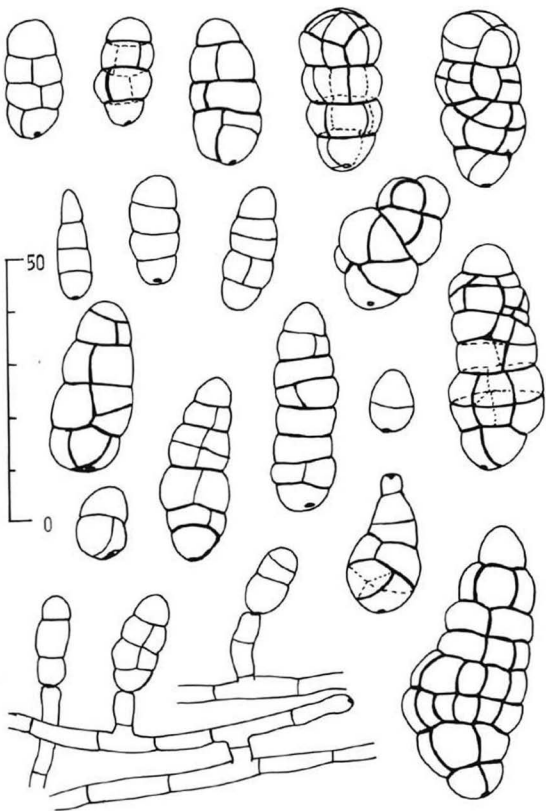


Fig. 2. *Alternaria mouchaccae* ex Type, from PCA culture.

task, is needed in aid of correct *Alternaria* determinations, especially for pathogens that figure largely and too often with incorrect names in the international literature.

Conidia of four species discussed in Section I above typically have no portion that can be distinguished objectively as a beak, true or false (an extremely rare pseudo-rostrum being exceptional). Conidia of a second, relatively large group of species are characterized as being non-beaked when produced but having a marked ability to produce chains through the agency of secondary conidiophores (false beaks), readily and sometimes even exuberantly.

3. *Alternaria chlamydospora* Mouchacca (Fig. 3)

The kinds of conidium enlargements that characterize *A. chlamydospora* (Mouchacca 1973; his Figs. 2 & 3) could support a suggestion that mycotaxonomists ignore the fungus and its name as representing a cultural monstrosity. A high percentage of conidia become grossly swollen and distorted; conidiogeny may become so disorganized that lateral hyphal branches, recognizable as potential conidiophores, sometimes enlarge abruptly terminally into quite recognizable *Alternaria* conidioids (Mouchacca, l. c., Fig. 3) instead of arising in the tretic mode of conidium ontogeny believed to be typical of the genus. Most definitions of "chlamydospore" would encompass such cellular oddities, even though some of them, at least, are able to produce typical apical conidiophores and, presumably, secondary conidia.

Eliminating this organism from taxonomic consideration would be inappropriate for several reasons, two important ones being (1) that the conidiogeny apparatus and the morphology of a respectable portion of conidia are distinctive and predictable up to the point at which individual cells become aberrant and (2) that recognizable chlamydosporic, conidioid isolates (as contrasted to conidium verum strains) are not uncommon among isolates of dematiaceous hyphomycetes and for practical purposes, including medical and phytopathological, cannot be ignored.

As for *A. chlamydospora* as a recognizable species, its individual conidia vera lack a definable beak. However, apical and even lateribasal secondary conidiophore production is common, as is subsequent chain formation. It would be idle to speculate whether or not this species exists in nature in a nonchlamydosporic state. Nevertheless, it would be a fascinating exercise (1) for an experimental systematist to attempt to isolate this fungus from whatever aberrational influence or factor is present and (2) for a

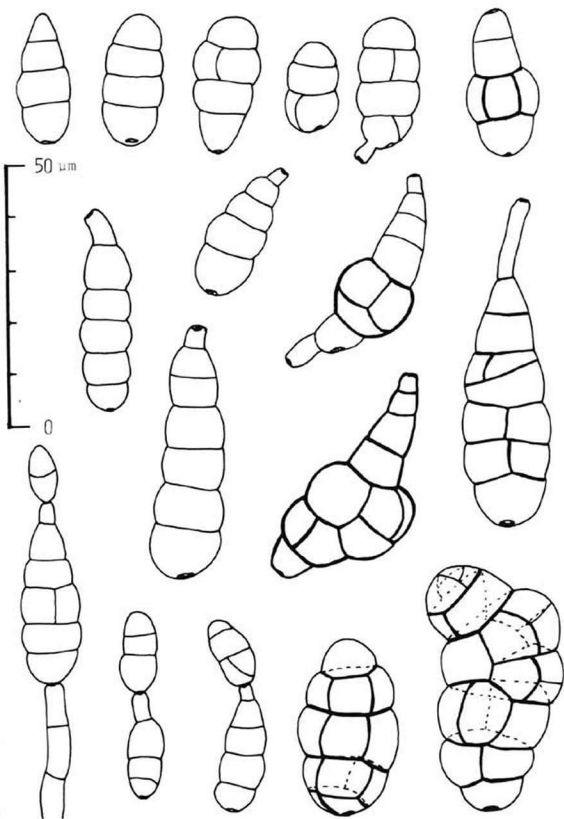


Fig. 3. *Alternaria chlamydospora* ex Type, from PCA culture.

mycopathologist to characterize the factor itself.

4. *Alternaria limaciformis* (Fig. 4)

A singular isolate received from the Commonwealth Mycological Institute, Kew, in 1954 has remained nameless since that time (along with several other such offerings) while preconceived ideas on *Alternaria* were being tested (Simmons 1967, 1978) and somewhat similar genera *Ulocladium* and *Embellisia* (Simmons 1967, 1971) were being segregated. Conidia of this isolate are solitary for the most part (well over 90%) and have no definable beak. However, secondary apical and lateribasal conidiophores are fairly common, with chains of two conidia equally common and of three rare but easily detectable in each colony scanned. Conidia often exhibit one or two minor arcs of curvature, even at the very young stage preceding septation, thus appearing somewhat vermiform. As transverse, longitudinal, and oblique septa are formed, especially in the central two quarters of enlarging conidia, the curvature often becomes accentuated two- or even three-dimensionally. The somewhat stubby, slightly swollen, sigmoid outline of many conidia bears a fancied resemblance to that of *Limax* garden slugs.

Alternaria limaciformis Simmons, sp. nov.

Ex culturis in agaris descripta. Coloniae in PCA planae, griseo-brunneae, atrantes; in V-8 planae, atro-brunneae, rubescentes; in Hay inconspicuae attamen conidiogena. Mycelium ex hyphis dilute brunneis, 3-4 μ m diam., necnon atrobrunneae, 6-7 μ m diam., in funiculis radiantibus compositum. Conidiophora recta vel acclivis, simplicia vel ramosa, ex hyphis submersis et aeriis oriunda, usque 5 μ m diam. et plerumque 150-200 μ m longa, ad quinque genicula sporifera praebentia. Conidia plerumque solitaria, aliquando brevicatenulata; initio ovoidea vel plerumque brevivermicularia, hyalina, levia; denique late ovoidea, ellipsoidea, flexa vel limaciformia, straminea, 30-45 x 9-18 μ m; septis transversalibus plerumque tribus ad sex et longitudinalibus obliquisve nullis vel in quoque cellula centrali uno ad duobus. Habitatio typi: solum prope Goole, Yorks., Britannia. Typus: partes ex Simmons 7-086 (IMI 52976, leg. A. R. Cottrell 307.2) desiccatae et in BPI, CBS, DAOM, IMI, NY conservandae.

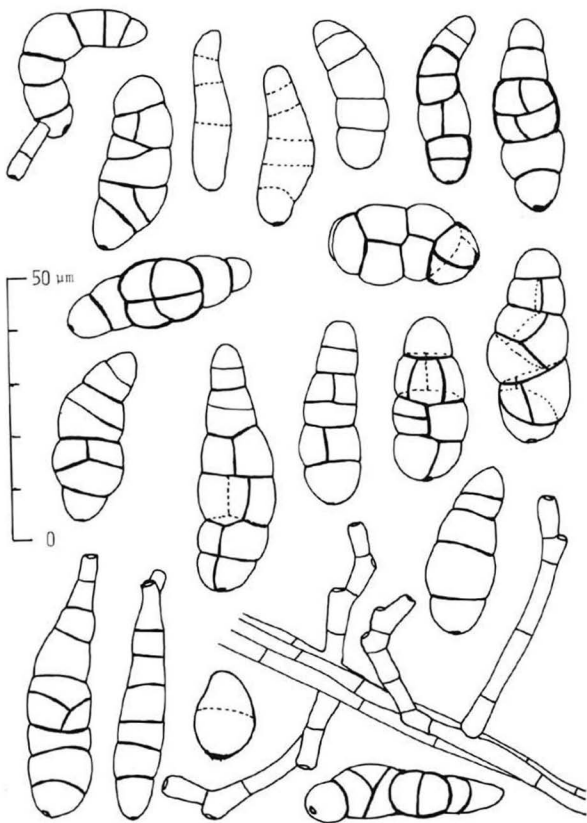


Fig. 4. *Alternaria limaciformis* ex Type, from PCA culture.

5. *Alternaria triticina* (Fig. 5)

A. triticina Prasada & Prabhu (1963), a striking species originally described from wheat leaves and, fortunately, represented in culture by an isolate from the type material, is another example of an *Alternaria* with nonbeaked conidia that readily produce apical outgrowths and, subsequently, short chains of two and three conidia. In my experience the great bulk of spotted wheat leaves that reach laboratories in both tropical and temperate regions as *Alternaria* specimens bear members of the taper-beaked *Alternaria alternata* (Fries) Keissler and *A. tenuissima* groups and, commonly, the *Alternaria* state of *Pleo-spora infectoria* Fuckel. *A. triticina* specimens are so rare in major *Alternaria* reference collections, including my own, that it seems likely that the species itself either is rare (for such a common and intensively cultivated host) or that it does not sporulate readily in nature, and thus escapes discovery on most diseased specimens.

A representative isolate (EGS 17-061, ex ITCCF 1186 G. C.) from the type specimen remains in good sporulating condition almost 20 years after its isolation. Some of its distinctive characteristics deserve review, in view of the circumstance that its original description lacks illustrations and is misleading about conidium surface ornamentation and that the influential published illustrations of M. B. Ellis (1976) appear either to be based on atypical material or to emphasize features (e. g., conidia gradually tapering distally and with smooth walls) that are not typical of the species.

Conidia of typical *A. triticina* terminate distally in an obtusely conical cell. Conidia produced in nature and in culture seldom taper into a rostrate structure, although the terminal cell often renews growth abruptly and produces a morphological and functional conidiophore. Conidium size typically does not exceed 60 x 25 μ m. Ornamentation of the conidium wall is shallow and inconspicuous, though present, in the type specimen; in the type isolate verrucae are so abundant on the conidium surface (6-8 per 10 μ m) and so conspicuous as to make observation of secondary conidial septa difficult. Young conidia are ovoid, becoming ellipsoid. A prominent, perhaps useful diagnostic characteristic of enlarging conidia is the tendency for them to become distinctly inequilateral.

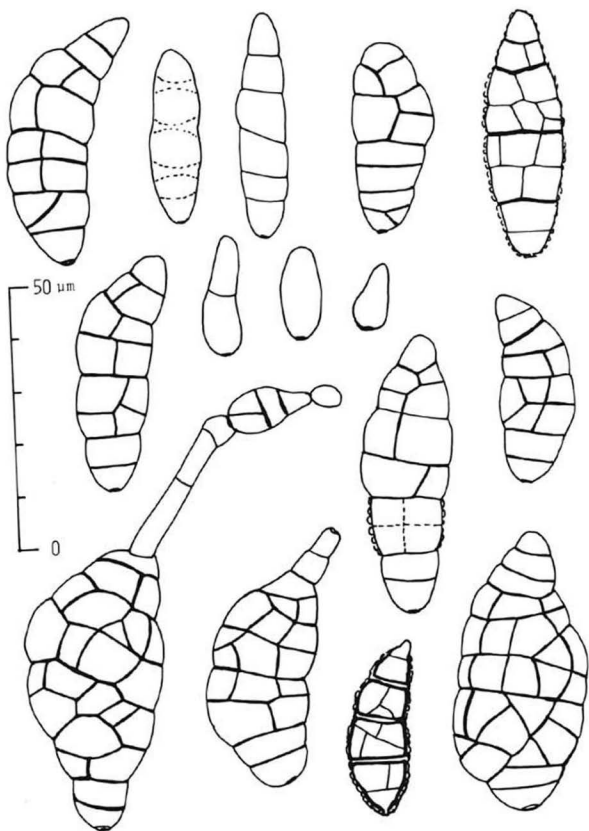


Fig. 5. *Alternaria triticina* ex Type, from PCA culture.

6. *Alternaria longipes* (Fig. 6-8)

The fungus species named *Alternaria longipes* (Ellis & Everhart) Mason has suffered considerable indignity at the hands of mycosystematists, nomenclaturists, phytopathologists, and tobacco farmers. Amends are due here as an introduction to this further example of a species with beakless conidia that readily generate apical conidiophores.

A collection of diseased tobacco leaves was received by J. B. Ellis with a letter dated Oct. 7, 1891, from Gerald McCarthy, North Carolina Agricultural Experiment Station, Raleigh. This material typifies the description of *Macrosporium longipes* Ell. & Ev. (1892); it still exists (NY), packeted in McCarthy's original letter with a scrap of paper carrying one of Ellis's characteristic sketches of a conidium and notes: "Brown spot"; "35-45 μ " (length of conidium body), "15-30 μ " (length of basal "stipe," a common misinterpretation at the time of an apical conidiophore or beak).

E. W. Mason (1928) transferred the Ellis and Everhart epithet to *Alternaria*. His decision to do so was based on a J. C. F. Hopkins isolate from Rhodesian tobacco of "an *Alternaria* of the *A. solani* type," which large-spored species the isolate later was considered to be. Mason soon recognized his misdetermination, inscribing a copy of his 1928 paper sent to John A. Stevenson: "Alas I got the wrong one. Tisdale is preparing to pounce on me over this." The error was unfortunate in its influence on phytopathologists working with tobacco diseases at the time; it was corrected in the Appendix of Hopkins's *Tobacco Diseases* (1956, pp. 163-164), along with comments on three, perhaps four distinguishable species of *Alternaria* associated with tobacco leaf diseases.

W. B. Tisdale and R. F. Wadkins (1931) also proposed a new combination *A. longipes* after making a detailed experimental study of the brown spot disease in Florida and of the fungus considered to be causal. The evidence of their descriptions and illustrations suggests that they were working with the same fungus that Ellis and Everhart had described. The work of Tisdale and Wadkins, because of its experimental approach, also was influential, but the correct attribution of the name in *Alternaria* remains *A. longipes* (Ell. & Ev.) Mason by priority of publication.

North Carolina, which supplied the type material of *A. longipes*, has continued to support interest and research in many aspects of this commercially important disease and

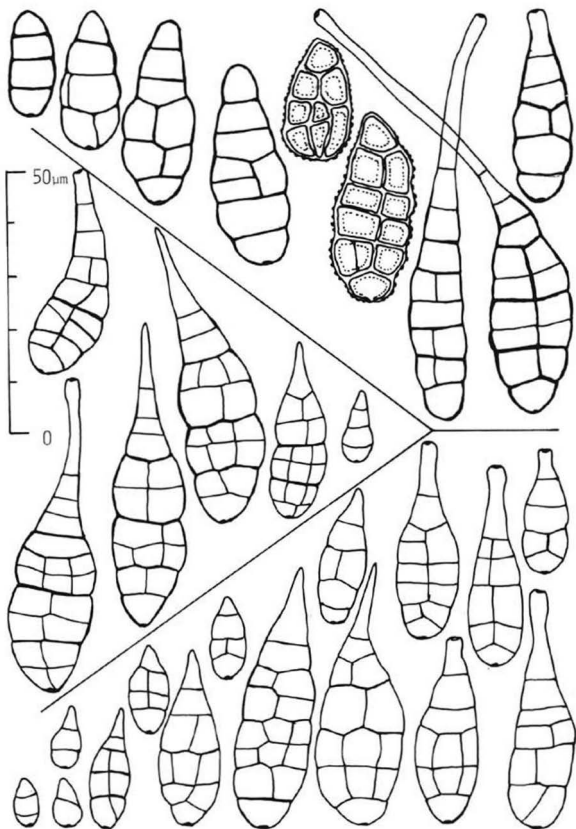


Fig. 6. Comparison of conidia selected from type specimens of *Alternaria longipes* (at top), *A. tenuissima* (left center) and *A. alternata* (bottom).

associated fungi for about 90 years. Reports on much of the tobacco brown spot work of this period reached publication using the name *A. longipes* for whatever small-spored species was/were present. The practical difficulty of distinguishing *A. longipes* from other species commonly present on tobacco led G. B. Lucas (1971) to suggest that *A. longipes* and genericotypical *A. alternata* are the same species. The suggestion is seductive in its ease of application to determinations of field specimens; it is unwarranted, however, on the basis of the literature on the subject. To the extent that tobacco disease work of the past decade is bracketed with the single fungus name *A. alternata*, the work requires reconsideration with respect to the identity of the fungus species actually involved.

Alternaria alternata (the name legitimized at present over the earlier *A. tenuis* Nees) may or may not be associated with brown spot lesions of tobacco in North Carolina; the species *A. longipes*, however, certainly has such a relationship with much pertinent material, not only in the U.S.A. but also in tobacco production areas of other parts of the world. Many field specimens and isolated strains among scores that have reached me are readily identifiable morphologically as *A. longipes*, and many of the latter have been characterized to me by the donors as being virulent pathogens under experimental conditions.

Examples of reputedly virulent strains that are morphologically typical *A. longipes* include EGS 30-101 (ex C. E. Main M-74, recd 1973); EGS 16-135 (ex G. B. Lucas R-66, recd 1963); EGS 30-033 (ex J. R. Stavely A-5, recd 1971); EGS 30-034 (ex J. R. Stavely A-3, recd 1971); EGS 30-080 (ex 1971 Fungicide Oxford #163 (slow), recd 1971); EGS 17-179 & 17-180 (ex D. C. M. Corbett, recd 1966 via IMI 123399 & 123400).

The distinctness of *A. longipes* from *A. alternata* and from the commonly encountered species *A. tenuissima* is illustrated (Fig. 6) with outlines of a range of conidia from the type specimen of each species. An excellent modern field collection of *A. longipes* (representing several received in 1973 from E. K. Sobers, Tifton, Georgia) is EGS 33-095, illustrated in Fig. 7.

In culture *A. longipes* produces conidia successively and so rapidly in chains that the transition region between conidium body and new conidiophore tends to be poorly defined. Nevertheless, most conidia reveal the site of the change from conidium to beak through a slight outline constriction and by reductions in wall ornamentation, color, and thickness. Conidium illustrations in Fig. 8 emphasize

several features of cultured material: the extraconidial nature of the conidiophore/beak and the fundamentally erostrate nature of *A. longipes* conidia; the somewhat reduced size of conidia produced in culture; and the remarkable thickness of conidium walls, which is observable in some field material and which is accentuated in conidia produced in culture on PCA. The comparatively restricted growth of *A. longipes* on V-8 agar, stressed by Sobers and Douplik (1969) in their studies of the species, has proved a particularly useful differential diagnostic feature of pure, stable isolates of the species.

The voluminous international literature and the unpublished virulence/aggressivity information shared with me

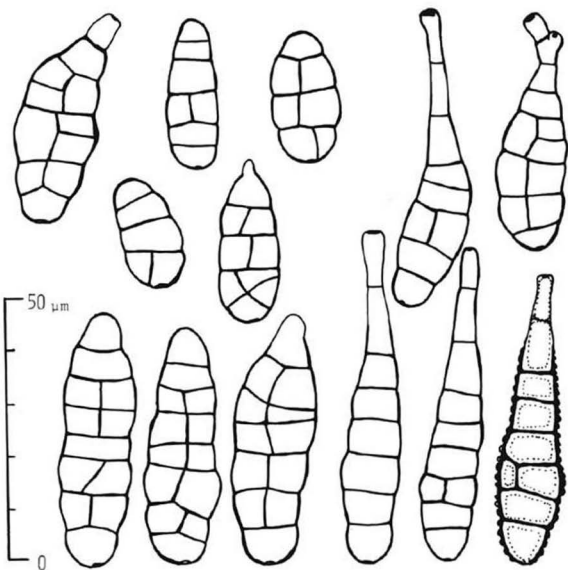


Fig. 7. *Alternaria longipes*, representative conidia from a modern field collection (Sobers, 1973; EGS 33-095).

by several phytopathologists suggest that many strains of morphologically distinct *A. longipes* are pathogenic to tobacco plants, that others have degenerated in pathogenicity during their years in culture, that other distinguishable species of *Alternaria* very commonly are found on and isolated from tobacco lesions, and that the total picture, insofar as species of *Alternaria* are involved, needs review.

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Continued support by the several herbaria and mycological institutions noted by abbreviations in the text, by the National Science Foundation (DEB79 04189), and by the U. S. Army Natick Research and Development Command is gratefully acknowledged.

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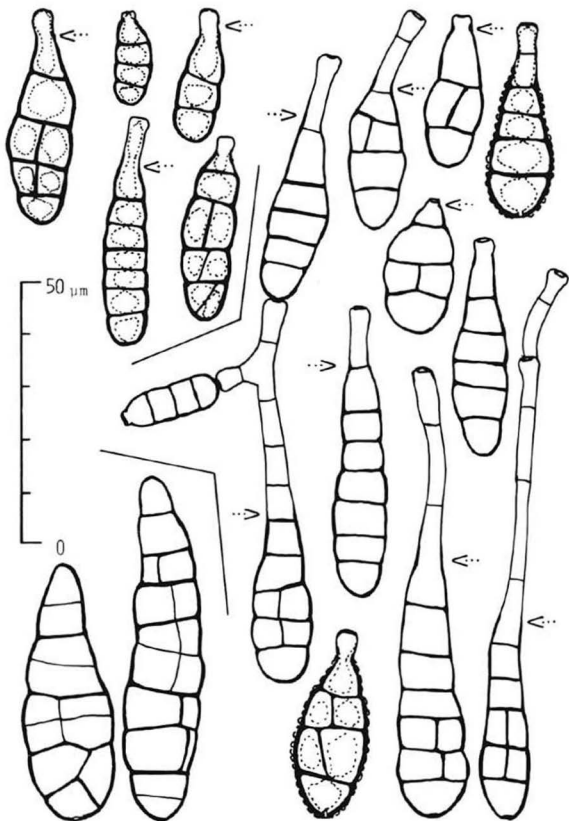


Fig. 8. *Alternaria longipes* conidia: large erostrate from nature (lower left), smaller pseudorostrate from Hay (right) and PCA (upper left); arrows indicate conidium/conidiophore transition zone. Conidium walls usually ornamented, thick.

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STUDIES IN TROPICAL CORTICIACEAE (BASIDIOMYCETES) III. TWO NEW SPECIES OF LAXITEXTUM

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SUMMARY

Laxitextum incrustatum nov. sp. and L. lutescens nov. sp. are described from Africa. A key to the known species in the genus is provided.

The genus Laxitextum was described by Lentz (1955) in a rather wide sense, but was later emended and restricted by Boidin (1958). The type species L. bicolor is widespread and is illustrated in Eriksson and Ryvar den (1976). Talbot (1951) reported L. bicolor from South Africa, Nigeria, Kenya and Uganda, while Boidin (1960) reported it from Zaire. During our work with African corticioid fungi, we came across two new species in the genus which are described below.

Key to species

1. Hymenium and pileus cream to deep yellow, gloeocystidia distinctly amyloid, spores 3,5-4 x 3-3,5 um.....L. lutescens
1. Hymenium white to pale cream, pileus if present, dark brown, gloeocystidia non-amyloid or only very weakly so, spores 4-5 um long.....2
2. Trama and pileus, if present, dark brown, tramal hyphae brown and smooth, spores oblong ellipsoid, 4-5 x 2,5-3 um.....L. bicolor
2. Trama cream, pileus not known, tramal hyphae pale yellowish and encrusted, spores broadly ellipsoid 4-5 x 3-3,5 um.....L. incrustatum

LAXITEXTUM INCRUSTATUM Hjortst. & Ryv. nov. spec.

Fructificatio mollis, plerumque resupinata, late effusa, 0.2 - 0.4 mm crassa; tramate distincto evoluto; hymenium leve, cremeum vel obscure ochraceum, leniter rimosum; systema hyphale monomiticum; hyphis cum fibulis, rectis, luteobrunneis, reniformibus, plerumque incrustatis, 3.5 - 4 um latis, aliquot oleosis, amyloidibus; gloeocystidia cum schizopapillis, fusiformia, subulata vel obtusa, 70 - 80 x

5 μ m, vulgo non-amyloidibus; basidia anguste clavata, 15 - 25 x 4 - 5 μ m, 4 - sterigmatibus; sporis subglobois vel ellipsoidibus, echinulatis, (4-)4.5(-5) x 3-3.25 μ m, valde amyloidibus.

HOLOTYPUS: Africa. Tanzania. Arusha Prov. Arusha Nat. park. Mt. Meru E. slope, road to the crater alt. 1800-2300 m.

8. Feb. 1973. L. Ryvarden 10108 (0).

PARATYPI: do. L. Ryvarden 10048; Tanzania: Kilimanjaro Prov. Mt. Kilimanjaro W slope, W. Kilimanjaro Forest Sta. alt. c. 1800 m. 10.-11. Feb. 1973 L. Ryvarden 10198; Tanga Prov. Lushoto distr., Usambara Mts. Magamba c. 4 km N of Lushoto, alt. 1600-2000 m. 21.-22. Feb. 1973. L. Ryvarden 10750. All in 0.

USA: Louisiana, St. Martinsville, no date, Leg. A.B. Langlois. (TRTC), do. Sept. 32. 1889, Leg. A.B. Langlois 2095 (TRTC). Without locality, Sept. 1913, on oak branches lying on the ground, C.G. Lloyd 46777 (TRTC).

Fruitbody soft, resupinate and widely effused, but loosening at the margin which is striate or fibrillose, about 0.2-0.4 mm thick with the trama distinctly developed, hymenium smooth, creamish to dull ochraceous, more or less cracked. Hymenial layer somewhat more dense than in the other species of the genus. Hyphal system monomitic with loosely interwoven tramal hyphae, golden yellow, straight and encrusted. The hyphal wall is thickened and the diameter may vary from 3,5 to 6 μ m, often with adventitious septa between each clamped cell. Subhymenial hyphae more irregular, pale coloured to hyaline. Oleiferous hyphae occur more or less abundant, usually with a slight amyloid reaction. All hyphae with clamps. Gloeocystidia arising from the tramal layer, fusiform, subulate or with obtuse apex, sometimes with schizopapilles, generally 70-80 x 5 μ m, but the length varies considerably and they may sometimes be up to 100 μ m or more, mostly without amyloid reaction, negative in benzaldehyde. Basidia narrowly clavate, 15-25 x 4-5 μ m, with four sterigmata. Spores subglobose to ellipsoid, echinulate, (4-)4.5(-5) x 3-3.25 μ m, strongly amyloid.

Remarks. The species is somewhat similar to L. lutescens (see below), but is easily distinguished by its slightly larger spores, encrusted hyphae and paler hymenium. Furthermore, all known specimens are resupinate. The hymenial layer is denser and the gloeocystidia are non-amyloid or only very weakly so. Because of its well developed subiculum and negative reaction in benzaldehyde the species seems to belong in Laxitextum and is related to L. bicolor. However, it also has similarities to a group of species in Gloeocystidiellum such as G. propinquum (Jacks.) Parm., G. sibiricum Parm., and Gloeocystidium lacticolor Bres. (type examined). The main differences are the distinct subiculum in Laxitextum and that oleiferous hyphae are lacking in these species and the sulfovanilline reaction is positive, (except in G. lacticolor?). Another

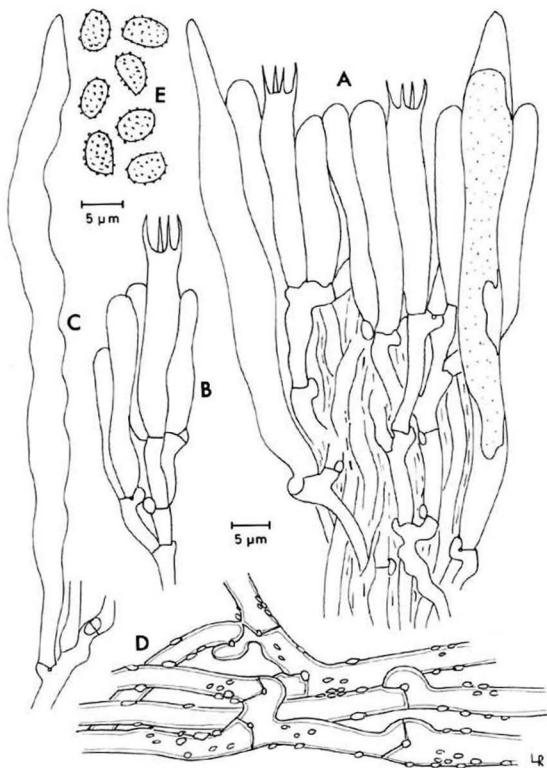


Fig. 1. Laxitextum incrustans. a) section through the hymenium b) basidium c) gloeocystidium d) subicular hyphae e) spores. From the holotype.

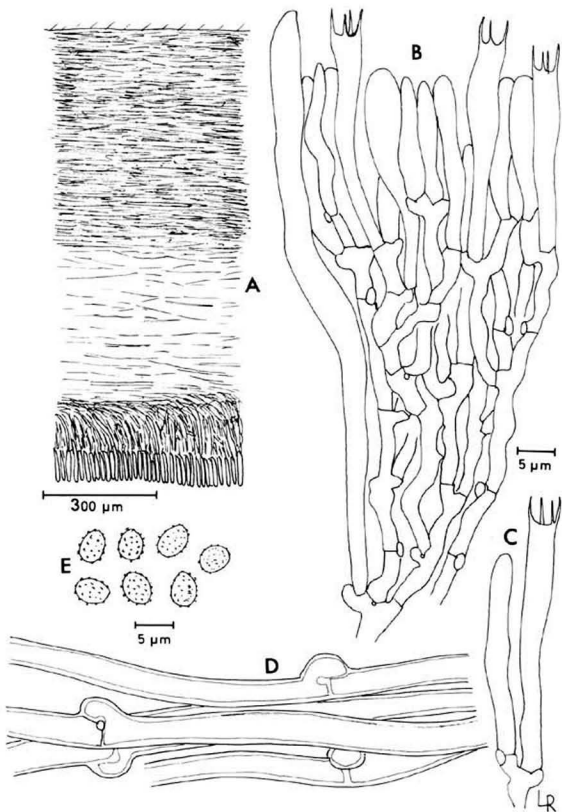


Fig. 2. *Laxitextum lutescens*. a) section through the fruit-body b) section through the hymenium c) basidium d) subicular hyphae e) spores. From the holotype.

species which also comes close to the group is Gloeocystidiellum furfuraceum (Bres.) Donk, but it lacks clamps.

We have also compared the new species with Scytinostromella humifaciens (Burt) Freeman & Petersen because of its somewhat similar spores and gloeocystidia. However, in that species the spores are smaller and the gloeocystidia react positively in benzaldehyde. Furthermore, like all species in Scytinostromella, also S. humifaciens, it is distinctly dimittic with skeletal hyphae.

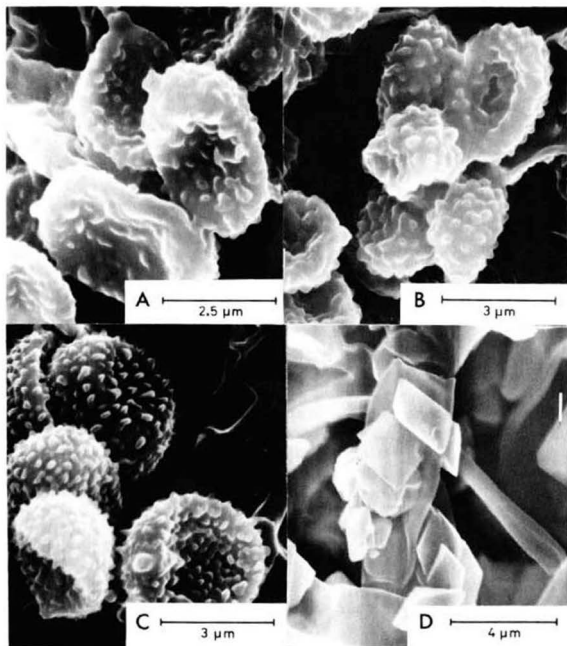


Fig.3. Spores of a) Laxitextum bicolor Coll. Ryv. 12092 b) L. lutescens, holotype c) L. incrustans, holotype d) en-crustated hyphae from L. incrustans, holotype. SEM by L. Ryvarden.

LAXITEXTUM LUTESCENS Hjortst. & Ryv. nov. spec.

Fuctificatio resupinata vel distincta reflexa, 0.4-0.6 mm crassa, supra infuscata; tramate distincto evoluto, 0.4-0.5 mm crasso; hymenium leve, cremeum vel lutescens, leviter rimosum; systema hyphale monomiticum; hyphis cum fibulis, rectis, levibus, laxe intertextis, luteobrunneis, circiter 4 μ m latis; hyphis subhymenialibus irregularibus, hyalinis; gloeohyphae et gloecystidia numerosa, amyloides; basidia anguste clavata, 20-25 x 7 μ m, 4 sterigmatibus; sporis subglobosis vel ellipsoidibus, echinulatis, 4 x 3-3.25 μ m, valde amyloidibus.

HOLOTYPE: Africa. Ghana. Ashanti Region, Bobiri Forest Reserve, ab. 30 km E of Kumasi. 18.-22. April 1974. L. Ryvar den 12875 (0).

Fruitbody resupinate to distinctly reflexed, 0.4-0.6 mm thick, pileus deep yellow to pale brown velutinate, azonate, trama well developed, pale yellowish brown, about 0.4-0.5 mm thick, hymenium cream-yellowish to strawcoloured with a light purple-brown tint, slightly cracked. Hyphal system monomitic, tramal hyphae loosely interwoven, pale yellowish brown, 3-5 μ m wide, smooth and with clamps. Subhymenial hyphae more irregular with the colour pale yellow or hyaline, mostly collapsed. Oleiferous hyphae abundant, 5-6 μ m wide, distinctly amyloid and with clamps at long intervals. Gloecystidia numerous, arising from the tramal layer, fusiform, with the apex obtuse, now and then with schizopapilles, 90-120 x 7-9 μ m, sometimes up to 150-200 μ m or more, distinctly amyloid, but negative in benzaldehyde. Basidia narrowly clavate, 20-25 x 4 μ m, with four sterigmata. Spores subglobose to ellipsoid, echinulate, 4 x 3-3.25 μ m, strongly amyloid.

Remarks. Laxitextum lutescens seems to be well located in the genus and is similar to L. bicolor, but delimited by its deep yellowish colour, the amyloid gloecystidia and oleiferous hyphae and the shorter spores. The tramal hyphae of L. bicolor are besides more distinctly brown.

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NOTES ON *ZOOPHTHORA OCCIDENTALIS*

(THAXTER) BATKO

(ENTOMOPHTHORALES: ENTOMOPHTHORACEAE)¹

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INTRODUCTION

The species *Entomophthora occidentalis* (Thaxter) 1888 was described by Thaxter (1888) as a species of *Empusa*. He recorded this pathogen as frequently and commonly infecting aphids on *Betula populifolia* Marsh in Maine, U.S.A. Batko (1964b) renamed this species *Zoophthora occidentalis* (Thaxter) and then placed it in the subgenus *Zoophthora* Batko (1966b) which includes the group of species referred to as the "*Entomophthora sphaerosperma* group" (Hutchison 1965; Waterhouse 1975; Remaudière et al 1976; Zimmermann 1978) and recently defined as *Zoophthora* Batko *sensu stricto* (Remaudière and Hennebert 1980).

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Further notes on the occurrence of this species are scarce and data on its morphology constitute only abbreviated records of the original description. Since the systematics of species within the subgenus *Zoophthora* has not yet been thoroughly defined and because the biology and morphological characteristics of *Zoophthora phalloides* Batko (1966a) closely resemble those of *Z. occidentalis*, it is useful to present the results of our observations of these fungi in recent years.

MATERIALS AND METHODS

In August and September of 1977 in Orono, Maine, five entomophthoraceous species were discovered on aphids *Myzus persicae* Sulz. infesting potatoes: *Z. occidentalis*, *Erynia neoaphidis* (Remaudière) Batko, *Z. exitialis* (Hall and Dunn) Batko, *Conidiobolus thromboides* Drechsler (= *Entomophthora virulenta* Hall and Dunn), and *Entomophthora major* (Thaxter) Gustafsson. Of these, *Z. occidentalis* was predominant. This pathogen was also collected from *Aphis fabae* Scopoli on pigweed, *Chenopodium album* L. The strains of this species were examined in the laboratory with application of usual microscopic techniques and suitable methods of cultivation on egg yolk and Sabouraud's maltose agar (EYSMA) and potato-dextrose agar (PDA) in 16 hours photophase. Laboratory infections were also attempted using several aphid species and caterpillars of *Choristoneura fumiferana* Clem. with conidial showers of the fungus. On July 13, 1978, a morphologically similar strain was found in the Wielkopolski National Park, Poland, on an aphid of the tribe Macrosiphina from *Deschampsia caespitosa* (L.) P.B. in the forest undergrowth. These materials have been preserved as microscopic preparations in lactophenol with cotton blue (LPCB) and the strains from the USA also in living cultures on media. In the course of the studies on the morphology and systematic position of these strains, a comparison was made with the holotype of *Zoophthora phalloides*, and data on the paratypes of this species were obtained.

RESULTS

A. Morphology and Diagnostics. The infected aphids produced rhizoids (Fig. 1) from the abdominal part of their bodies as 1-3 compact bundles which firmly attached them to plant leaves. The tips of the hyphae forming these rhizoidal bundles are widened and deeply sinuate (Fig. 1a).

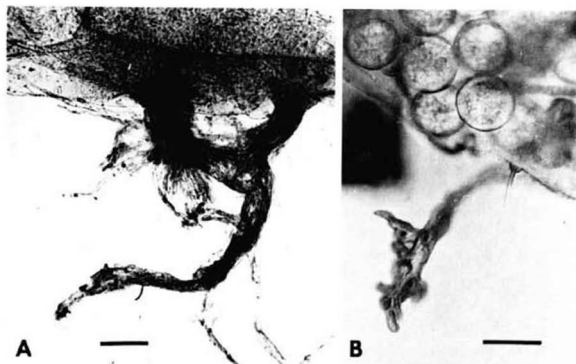


FIG. 1. Rhizoids of *Z. occidentalis* from hosts producing (a) conidia and (b) resting spores. Bar = (a) 20μ , (b) 10μ .

During the time of sporulation, the conidiophores covered the bodies with a compact mass of silver-white to white-cream mycelium; on the aphid from *D. caespitosa* the color of the mycelium was dirty lily. Pseudocystidia tapered with slightly blunt tips are scarce, and not a constant feature in the microstructure.

The conidia (Fig. 2) are elongated, with sides bowed slightly outward; conidia may be slightly curved. The conidium narrows slightly towards the papilla, and has a bluntly conical apex. In outline, the papilla is triangular, and often bears a small central apiculus or arch-wise and obtuse. The single nucleus is oval or elongate, and strains well in LPCB; the plasma appears somewhat hyaline with several granules. Dimensions of the conidia from aphids or from media differed remarkably: From aphids, they were found to be $25-26.5\mu \times 8-10\mu$ (av. $30 \times 9\mu$), L/D 2:3-3:5 (av. 3.0). From EYSMA conidia were $24-42\mu \times 7-10\mu$ (av. $31 \times 9\mu$); L/D 3:4-4:5 (av. 4.0). Secondary conidia of the first order are similar to the primary ones but a little shorter.

After the transfer of the conidia from EYSMA to PDA, capillispores are formed in great numbers, and a few secondary conidia of further orders usually appear; these

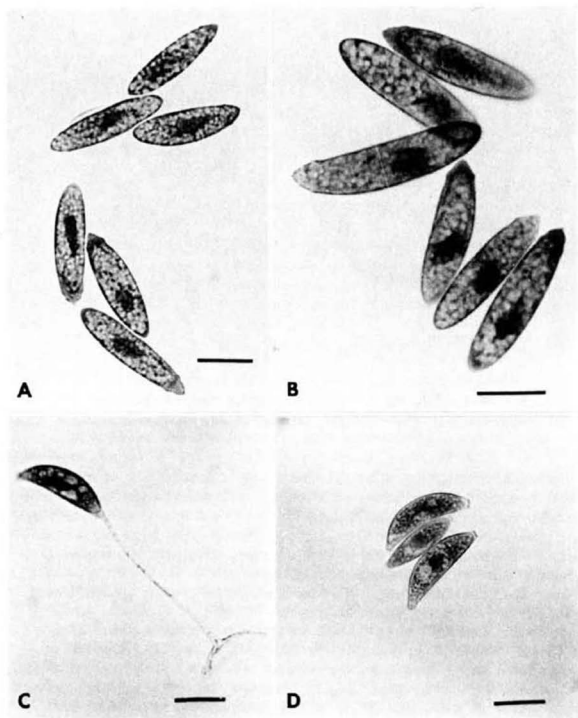


FIG. 2. Primary conidia of *Z. occidentalis* (a, b) are elongate and slightly curved. Primary conidia may produce secondary capillispores at the end of a fine tube (c); these secondary conidia are more strongly curved than primary conidia and lack a definite papilla (d). Bar = 10 μ .

latter conidia are short and considerably different from the primary, with L/D ratio less than 2. Capillisporos (Fig. 2c,d) are found relatively seldom on aphids. Capillisporos from aphids were found to be 20.2-25.1 X 7-9 μ , and, from media (EYSMA after transfer to PDA), 22-33.3 X 7-9 μ ; L/D 2:6-3:7 (av. 3:2).

Resting spores (Fig. 3) were formed in artificially infected aphids or on EYSMA at the temperature 25-27°C. They are smooth, globose, (20.5) 24-30 (32.5) μ diameter, with a light yellow wall 5-6 μ thick. On EYSMA at 25-27°C, resting spores were larger (20-41.5 μ) than in the insect. On the basis of many observations of early stages of spore development, it is concluded that resting spores are formed both as azygospores and zygozspores, even though the "classical" types of conjugation noted and illustrated by Thaxter (1888, p. 171-172) were not observed.

Misidentification of *Z. occidentalis* and *Z. phalloides* is possible because of the similarities of their published spore dimensions and, more importantly because of imprecise characterizations of the conidial apex in the description of these species. Thaxter (1888, p. 171) described the apex of *Z. occidentalis* as "tapering strongly" whereas Batko (1966a, p. 10-11) referred to *Z. phalloides* as "sharply pointed". On the basis of the analysis of differences described by Remaudière et al. (1976), studies which did not consider *Z. occidentalis*, the authors intended to provide a diagnosis for *Z. phalloides*. However, the comparison of the conidia of both species shows that the following alternatives would be more suitable:

- conidia cylindrical with parallel sides in outline, hemispherically convex papilla and broadly obtuse ends *Z. phalloides*
- conidia ellipsoidally spindle-shaped with the sides a little convex, papilla widely conical or slightly convex often with small sharp central apiculus, top part of the spore tapering with somewhat blunt apex *Z. occidentalis*

B. Biology and Occurance. *Zoophthora occidentalis* was the most frequent pathogen among entomophthoraceous fungi collected from the aphid *M. persicae* in Orono, Maine, causing 70-80% among the cases of mycosis; whereas the

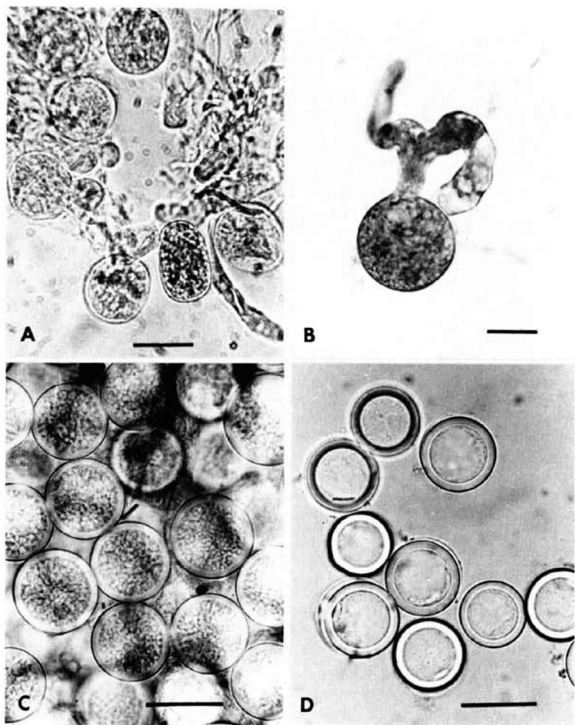


FIG. 3. Resting spores of *Z. occidentalis* are (a) thin walled in the early stages of formation and may be zygospores (b); the resting spore contents are granular at first (c) but contain an oil globule at maturity (d). Bar = (a,c,d) 20 μ , (b) 10 μ .

total mortality caused by entomogenous fungi was very high at the end of the growing season. On the media used in this investigation, the fungus grew and developed well, and produced all forms of spores. Under laboratory conditions, some species of potato aphids - viz., *M. persicae*, *Macrosiphum euphorbiae* Thom., *Aphis nasturtii* Kalt., and *Acyrtosiphon solani* Kalt. - were successfully infected with the spores produced on EYSMA. Attempts at similar infection against larvae of spruce budworm, *C. fumiferana*, failed.

The relative absence of *Z. occidentalis* from published reports probably derived from its misidentification as *Z. radicans* (Brefeld) Batko (= *Entomophthora sphaerosperma* Fres.). Even the investigations of Shands et al. (1958; 1962; 1963; 1972) on the entomogenous fungi of potato aphids in Maine, carried out between 1952-1972, did not report the presence of *Z. occidentalis*. Likewise, Gustafsson (1965) does not mention it from Sweden. Petch (1939; 1944) on the other hand reported it from two localities in Great Britain but contrary to his usual procedure offered no precise description. Batko (1962) listed it among Polish Entomophthoraceae from an aphid on nettle, *Urtica dioica* L., in the Bialowieza National Park, where, despite many repeated thorough searches, this species was found on few aphids only in one locality. Thus it may be interesting to determine the circumstances in which the increase of its frequency in natural conditions occurs, as well as to continue attempts to experimental infections in biological control of aphids.

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NEW CICADA PATHOGENS:

MASSOSPORA CICADETTAE FROM AUSTRALIA
AND *MASSOSPORA PAHARIAE* FROM AFGHANISTAN¹

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INTRODUCTION

The genus *Massospora* was described by Peck (1879) and emended by Soper (1974). Forbes (1888) and Thaxter (1888) working independently, correctly placed this genus in the Entomophthoraceae. Species of *Massospora* are pathogenic to cicadas and thus far are found only in adults. These fungi attack the reproductive organs of their host. As the fungus grows, the terminal abdominal segments of the host slough off revealing either conidia or resting spores. Both stages do not normally occur in the same individual. The infected cicadas remain alive and take an active part in transmission of conidia to other adult cicadas or in distribution of the resting spores (Soper et al. 1976a). Only one species, *Massospora levispora* Soper, has received extensive epizootiological study (Soper et al. 1976b). From this investigation and the observations of White et al. (1979) on

¹ Partial support for this project was provided by the United States/Australian Cooperative Science Program under travel authorization 2-AS-11 of NSF Contract INT 77-26877.

Massospora cicadina Peck, it is apparent that these pathogens play an important role in the population dynamics of their hosts.

Except for one report of *Massospora*, probably erroneously identified as *M. cicadina*, from the Japanese cicada *Platypleura kaempferi* F. (Kobayasi 1951), these fungi were known only in the Western Hemisphere. This paper describes two new *Massospora* species which indicates a much wider distribution by the addition of Afghanistan and Australian localities. This brings the total of known *Massospora* species to 13.

MASSOSPORA CICADETTAE Soper, sp. nov.

CONIDIA ochroleuca in massa, ellipsoideis, papilla indistincta, parietibus laevibus, $7.3-9.8 \times 20.7-28.1 \mu\text{m}$ ($9.0 \pm 0.8 \times 24.7 \pm 2.1 \mu\text{m}$, $\bar{x} \pm s$) binucleata vel interdum trinucleata vel uninucleata, nucleis nunquam bipolaribus autem ad partem latissimum conidii locatis. SPORAE PERDURES pallida vitellina in massa, reticulis profundis cameras distinctas formantes, cristis angustis laevibus vel raro minute papillatis, $26.8 - 38.7 \mu\text{m}$ in diam. ($35.2 \pm 3.2 \mu\text{m}$, $\bar{x} \pm s$).

CONIDIA pale brownish yellow in mass, ellipsoid, with an indistinct papilla, smooth-walled, $7.3 - 9.8 \times 20.7 - 28.1 \mu\text{m}$ ($9.0 \pm 0.8 \times 24.7 \pm 2.1 \mu\text{m}$, $x \pm s$), binucleate or occasionally tri- or uninucleate, nuclei located at the broadest part of the conidium, never bipolar. RESTING SPORES pale egg-yolk yellow in mass, with deep reticulations forming distinct chambers, the narrow ridges smooth or rarely minutely papillate, $26.8 - 38.7 \mu\text{m}$ in diam. ($35.2 \pm 3.2 \mu\text{m}$, $\bar{x} \pm s$). (Fig. 1)

Holotype: AUSTRALIA: New South Wales: 16 km east of Hay: Growing in the abdomen of an adult cicada, *Cicadetta murrayiensis* Distant, Det. M.S. Moulds. Conidial stage. 11 December 1978, *Coll. M.S. and B.J. Moulds*. CUP.

Paratypes: Same data as holotype. Resting spores present. CUP. AUSTRALIA: Queensland: Goomeri: near Kinbombi Falls. Growing in abdomen of *Cicadetta murrayiensis*, Det. M.S. Moulds. Resting spore stage present. 19 December 1976, *Coll. M.S. and B.J. Moulds*. CUP. Queensland: Aloomba: Growing in abdomen of *Cicadetta puer* (Walker), Det. K.J. Chandler. Conidial stage

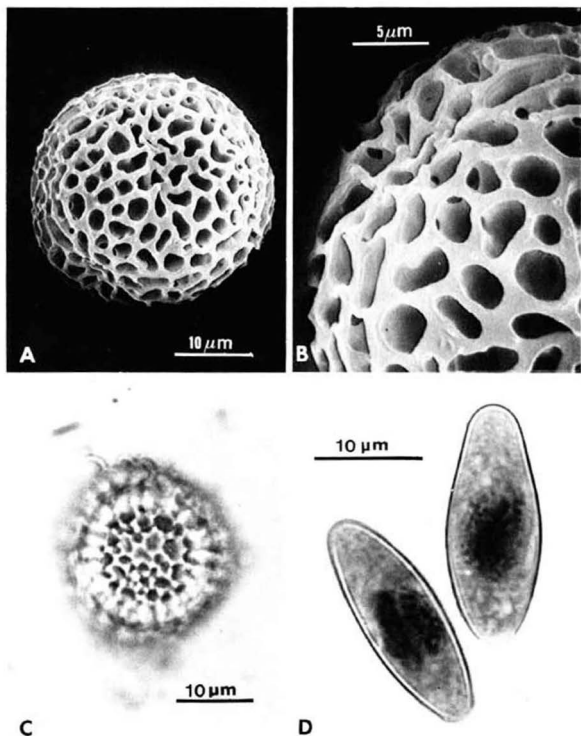


Figure 1. Spore stages of *M. cicadettae*: (A) and (B) scanning electron micrographs of resting spores; (C) resting spore as seen through light microscope; and (D) conidia.

present. 10 February 1973, Coll. K.J. Chandler. CUP. Tasmania: Davenport. Growing in the abdomen of *Cicadetta* sp., Conidia and resting spore stages present. No collection data. South Australia Museum.

Name: The species name of this *Massospora* is based on the generic name of its host, *Cicadetta*.

Host: *Cicadetta murrayiensis*, *Cicadetta puer*, and *Cicadetta* species (Cicadidae).

This species has the largest conidia yet described ($9.0 \times 24.7 \mu\text{m}$) for *Massospora*. Although *Massospora dorisiana* Soper measure $10.8 \times 21.8 \mu\text{m}$ thus approaching these dimensions, the spore surface is verruculose as opposed to smooth in *M. cicadettae*. The nuclei are distinctly bipolar in *M. dorisiana*, and although *M. cicadettae* is likewise generally binucleate, the nuclei are positioned centrally. The size and ornamentation of resting spores (*M. cicadettae*) are similar to the Chilean species *Massospora tettigatis* Soper which measures $36.3 \mu\text{m}$ in diameter. In the absence of conidial characteristics these species can be separated by the general lack of papillae on the reticulation of *M. cicadettae* resting spores as compared with numerous minute papillae found on *M. tettigatis* resting spores. The location (Australia) and host (*Cicadetta* sp.) are strong circumstantial support of *M. cicadettae* when only resting spores are present.

The host, *C. puer*, is a serious pest on sugar cane in northern Queensland, Australia. Although most cicadas have very long life cycles, viz., 4 to 17 years, *C. puer* is thought to complete its cycle in one year (K.J. Chandler personal communication). Potentially, *M. cicadettae* could have a correspondingly short life cycle, which would be the shortest known for any *Massospora* species. Intensive epizootiological studies on the periodical cicadas, *Magicicada* spp. indicate *M. cicadina* does not occur in the nymphal stages of the host nor in other genera of cicadas. This indicates *M. cicadina* must remain in the resting spore state for 17 years between the synchronized emergence of the adults (Soper et al. 1976a). A hiatus of 9 years between the occurrence of *M. levispora* resting spore production in an isolated population of *Okanagana rimosa* (Say) was given as evidence of a correspondingly long survival of *M. levispora* (Soper et al.

1976b). These observations strongly suggest that the life cycles of *Massospora* species are controlled in some way by those of their hosts rather than by timing mechanisms intrinsic to the fungi themselves.

MASSOSPORA PAHARIAE Soper, sp. nov.

CONIDIA ignotae. SPORAE PERDURES cinnamomeae in massa, reticulis profundis irregularibus indistinctae, cameras indistinctas formantes, cristis late separatis angustis papillatis, 32.8 - 44.7 μm in diam. ($39.5 \pm 2.6 \mu\text{m}$, $\bar{x} \pm s$).

CONIDIA unknown. RESTING SPORES dark red-brown in mass, with deep, irregular to indistinct reticulations, ridges widely separated, narrow, bearing numerous minute papillae, 32.8 - 44.7 μm in diam. ($39.5 \pm 2.6 \mu\text{m}$, $\bar{x} \pm s$). (Fig. 2)

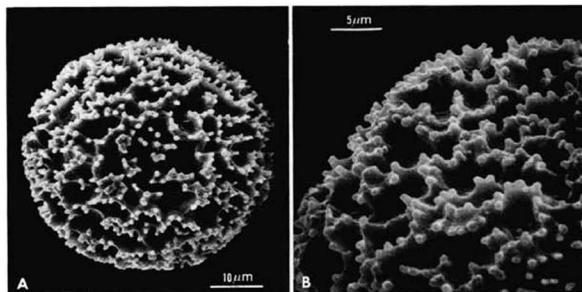


Figure 2. Scanning electron micrographs of *M. pahariae* resting spores, (A) and (B) note the many papillae on the reticulations.

Holotype: AFGHANISTAN: Paghman: Growing in the abdomen of an adult cicada, *Paharia casyapae* (Distant), collected on "shade and fruit trees", Det. R. Froeschner. Resting spores present. 3 July 1963, Coll. E.R. Millet AF 4 63, 23-14756. CUP.

Name: The species name of this *Massospora* is based on the generic name of its host, *Paharia*.

Host: Homoptera: *Paharia casyapae* (Cicadidae).

The resting spores of this species are similar to *Massospora levispora* Soper in their ornamentation (Soper, 1963, 1974). They can be differentiated by their average larger size 39.5 μm vs. 34.0 μm and fewer reticulations. In mass *M. levispora* resting spores are yellowish brown as opposed to the cinnamon brown of *M. pahariae*.

SYNOPTIC KEY TO SPECIES

The following key is a revised version of the original (Soper 1974). To identify a species the key can be entered at any point. As additional characters are examined, a unique combination will be found for each species. If only the resting spore is present, it will be necessary to utilize scanning electron microscopy. Complete descriptions of species 1-11 are given by Soper (1974). Species numbers are underlined when the characteristic is unique.

- | | |
|---|---|
| 1. <i>Massospora cicadina</i>
Peck | 5. <i>Massospora ocybetes</i>
Soper |
| 2. <i>Massospora spinosa</i>
Cifera, Machado & Vital | 6. <i>Massospora tettigatis</i>
Soper ² |
| 3. <i>Massospora levispora</i>
Soper | 7. <i>Massospora carinetae</i>
Soper ² |
| 4. <i>Massospora dorisianae</i>
Soper ² | 8. <i>Massospora diminuta</i>
Soper |

² Several orthographic errors were made in the original descriptions with respect to Latin endings. These have since been corrected in the Index of Fungi (1975. 4:312). The corrected spellings are used here.

- | | |
|---|---|
| 9. <i>Massospora platypediae</i>
Soper ² | 12. <i>Massospora cicadettae</i>
Soper |
| 10. <i>Massospora diceroproctae</i>
Soper ² | 13. <i>Massospora pahariae</i>
Soper |
| 11. <i>Massospora fidicinae</i>
Soper ² | |

CONIDIAL CHARACTERS

- | | |
|--|---|
| 1-1 <i>Conidial shape</i> | 1-3 <i>Conidial wall ornamentation</i> |
| a. ellipsoidal 2, 3, 4, 5, 9,
10, <u>12</u> | a. absent <u>3</u> , <u>6</u> , <u>9</u> , <u>12</u> |
| b. fusiform 10 | b. verrucose <u>1</u> , <u>2</u> , <u>7</u> |
| c. globose 7 | c. verruculose <u>4</u> , <u>5</u> , <u>10</u> , <u>11</u> |
| d. navicular 4 | 1-4 <i>Arrangement of nuclei</i> |
| e. obovate 4, 10 | a. bipolar <u>2</u> , <u>4</u> , <u>9</u> , <u>10</u> |
| f. ovoid <u>1</u> , 2, 3, 5, <u>6</u> , 11 | b. random <u>1</u> , <u>3</u> , <u>6</u> , <u>7</u> , <u>12</u> |
| g. subglobose 11, 7 | 1-5 <i>Conidial length</i> |
| 1-2 <i>Number of nuclei</i> | a. less than 8 μm 9 |
| a. one 3, <u>11</u> , 12 | b. 8 to 10 μm 2, 3, 5, 6,
7, 9 |
| b. two <u>1</u> , <u>2</u> , 3, <u>4</u> , 6, <u>7</u> , <u>9</u> ,
<u>10</u> , <u>12</u> | c. 10 to 15 μm 1, 2, 3, 4,
5, 6, 7, 8, 9, 10, <u>11</u> |
| c. three 3, 6, 12 | d. 15 to 20 μm 1, 2, 3, 4,
5, 6, 10 |
| d. four or more 6 | e. more than 20 μm 2, 3, 4,
10, <u>12</u> |

RESTING SPORE CHARACTERS

- | | |
|---|---|
| 2-1 <i>Ornamentation of resting spore reticulum</i> | 2-3 <i>Resting spore diameter</i> |
| a. papillae absent <u>5</u> | a. less than 25 μm 8 |
| b. papillae minute <u>6</u> | b. 25 to 30 μm 3, 5, 6, 8,
12 |
| c. papillae rounded <u>1</u> , 2, <u>3</u> ,
<u>12</u> , <u>13</u> | c. 30 to 45 μm 1, <u>2</u> , 3, 5,
6, <u>11</u> , <u>12</u> , <u>13</u> |
| d. papillae truncate 2, <u>8</u> , <u>11</u> | d. more than 45 μm 1 |
| 2-2 <i>Ridges of reticulum</i> | |
| a. broad forming small chambers <u>6</u> | |
| b. irregular forming indistinct chambers <u>3</u> , <u>13</u> | |
| c. narrow forming distinct chambers <u>1</u> , <u>2</u> , <u>5</u> , <u>8</u> , <u>11</u> , <u>12</u> | |

HOST

3-1 *Genus of cicada attacked*

- | | |
|----------------------------------|--------------------------------|
| a. <i>Carineta</i> <u>7</u> | g. <i>Magiccicada</i> <u>1</u> |
| b. <i>Cicada?</i> <u>8</u> | h. <i>Okanagana</i> <u>3</u> |
| c. <i>Cicadetta</i> <u>12</u> | i. <i>Paharia</i> <u>13</u> |
| d. <i>Diceroprocta</i> <u>10</u> | j. <i>Platypedia</i> <u>9</u> |
| e. <i>Dorisiana</i> <u>4, 5</u> | k. <i>Quesada</i> <u>2</u> |
| f. <i>Fidicina</i> <u>11</u> | l. <i>Tettigates</i> <u>6</u> |

DISTRIBUTION

4-1 *Collection localities*

- | | |
|--------------------------|----------------------------------|
| Afghanistan <u>13</u> | Honduras <u>11</u> |
| Argentina <u>5, 7</u> | Mexico <u>2, 9, 11</u> |
| Australia <u>12</u> | United States <u>1, 3, 9, 10</u> |
| Brazil <u>2, 4, 7, 8</u> | Venezuela <u>2</u> |

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LEOTIACEAE III. NOTES ON SELECTED TEMPERATE SPECIES
REFERRED TO *HELOTIUM* AND *HYMENOSCYPHUS*

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SUMMARY

Fourteen temperate species of *Helotium* and *Hymenoscyphus* were studied and their taxonomic placement discussed. The accepted species are redescribed and illustrated. Of the fourteen, six were shown to be good species of *Hymenoscyphus*, four were demonstrated to be taxonomic synonyms of other species of *Hymenoscyphus*, one additional species is a probable synonym, and three were shown to be members of the Sclerotiniaceae.

As was recently pointed out by Carpenter (1981) and by Dumont (1980, 1981), the identification of the majority of the neotropical species of Inoperculate Discomycetes is difficult and, at times, frustrating due to the lack of comprehensive and updated monographs and floristic studies. Dumont (1980) pointed out, in his study of *Helotium rufo-corneum*, that even the most common species are still difficult to name correctly because of the lack of adequate literature descriptions and because of existing, confusing synonymies.

We have been attempting to work out the names of some of the more commonly encountered species of Inoperculate Discomycetes which we have collected (Carpenter & Dumont, 1978; Carpenter, 1981; Dumont & Carpenter, 1981; Dumont, 1980; and Haines, 1980) from neotropical regions. Dumont & Carpenter (1981) reported the difficulty they encountered while attempting to identify their own field collections of the *Helotium-Hymenoscyphus* complex and *Helotium*-like organisms from Colombia and adjacent regions and pointed out the need to reinvestigate all of the species of that genus described from the neotropics. Dumont (1981) presented a preliminary summary of all of the species of *Helotium-Hymenoscyphus* reported from the neotropics. In that study many species were actually excluded from the genus, placed into synonymy, or redistributed to genera in the Sclerotiniaceae.

While attempting to work out some of the complicated synonymies, such as found in *Helotium rufo-corneum*, it became necessary to study the literature and specimens of several spe-

cies of temperate members of *Helotium-Hymenoscyphus*. The purpose in the paper is to present some of the information recently uncovered while studying the types of several temperate species described originally as *Helotium* or placed in *Helotium*, in hopes that this will add to the growing information on the genus *Hymenoscyphus*.

As I become more familiar with and study additional species of *Helotium-Hymenoscyphus*, I am beginning to see the emergence of two distinct groupings of species in the genus. One I refer to as the *Hymenoscyphus caudatus* group (which includes the type species *H. fructigenus*) and the second the *Hymenoscyphus epiphyllus* group. In the first group, the apothecia are stipitate, and the sterile tissue (outer ectal excipulum) from the base of the stipe to the margin is composed of a well developed and well defined *textura porrecta* to *textura prismatica*. In the *H. epiphyllus* group, the apothecia are generally turbinate to substipitate, and the outermost tissue of the stipe and frequently the lower portion of the receptacle adjoining the stipe is composed of a well to poorly defined *textura globulosa* to *textura angularis*. It is probable that these two distinct structural differences are worthy of recognition; either a separate genus for the *H. epiphyllus* group, or an infrageneric rank might be applicable. Since the majority of the species of *Helotium-Hymenoscyphus* still remain to be examined and redescribed, it would be premature to make additional modifications at this point.

The accompanying taxonomic key includes not only the species studied in the present work, but incorporates all of the species which I have previously studied (Dumont & Carpenter, 1981; and Dumont, 1980) and accept as valid species of *Hymenoscyphus*. The key should not be thought of as definitive, but one to which additional species should be added after they are studied. The methods used are the same as those reported by Dumont (1971).

Key to species of Hymenoscyphus studied

1. Apothecia sessile, substipitate to turbinate; stipe (or substipe) and base of receptacle of apothecium to the outside composed of *textura angularis* or *globulosa*.....(*Hymenoscyphus epiphyllus* group)2.
2. Ascospores 28-42x3-5µm, fusoid, anterior end pointed, posterior end attenuated.....*H. dearnessii* p. 62.
2. Ascospores less than 28µm long.....3.
3. Ascospores (7-)8-10(-12)x2.5-3.5(-4.5)µm, trapezoidal, obovoid, generally equilateral, if inequilateral then not flattened on one surface.....*H. immitabilis* p. 73.
3. Ascospores more than 12µm long.....4.
4. Apothecia occurring on wood, strongly umbilicate; ascospores (13-)15-18(-20)x(3-)4-5µm.....*H. umbilicatus*
(See Dumont, 1980)
4. Apothecia occurring on leaves, turbinate, not umbilicate; ascospores 15-18x3.5-5µm.....*H. epiphyllus* p. 66.
(See Dumont, 1980)
1. Apothecia stipitate; stipe (or substipe, if present) composed of *textura porrecta* to *textura prismatica* (if any globose cells present in stipe or receptacle, use first choice).....

-(*Hymenoscyphus caudatus* group)5.
5. Ascospores equilateral or if inequilateral then not flattened, generally trapezoidal to obovoid to obpyriform.....6.
6. Apothecia characteristically tiny, less than 1mm in diam, discoid; ascospores (10-)11-14(-15)x(3-)4(-5) μ m on leaves; temperate.....*H. translucens* p. 80.
6. Apothecia characteristically large, greater than 2mm in diam, irregular in outline, umbilicate; ascospores (4-)5-7(-9)x1.5-2(-2.5) μ m.....*H. leucopsis*
(See Dumont, 1980)
5. Ascospores strongly inequilateral, a high proportion flattened..7.
7. Ascospores with a large "nuclear staining area" visible in phloxine, cotton blue and analine blue dyes, rarely 1-septate, (22-)26-30(-35)x4-5(-6) μ m, ascus apex papillate; tropical.....*H. sclerogenus*
(See Dumont & Carpenter, 1981)
7. Ascospores without nuclear staining area.....8.
8. Ascospores regularly septate (more than 50% in any amount).....9.
9. Ascospores 1-septate, (17-)18-22(-24)x4-5(-6) μ m; apothecia without hairs at the base of the stipe.....*H. musicola*
(See Dumont, 1980)
9. Ascospores 3-septate, (20-)24-30(-35)x4-5(-6) μ m; apothecia with hairs at the base of the stipe.....*H. lasiopodius*
(See Dumont & Carpenter, 1981)
8. Ascospores aseptate.....10.
10. Ascospores (16-)18-22(-26)x2-3 μ m, with a basal cilium.....*H. scutulus*
(See Dumont & Carpenter, 1981)
10. Ascospores without a basal cilium.....11.
11. Ascospores hooked apically.....12.
12. Ascospores (16-)18-23(-30)x3-3.5(-4.5) μ m, strongly and obviously hooked, tapering gradually to the base.....*H. serotinus*
(See Dumont & Carpenter, 1981)
12. Ascospores (14-)16-23(-26)x4-5(-6) μ m, if hooked, only slightly and a few per mount, abruptly pointed at basal end...*H. caudatus*
(See Dumont & Carpenter, 1981)
11. Ascospores not hooked apically.....13.
13. Ascospores (11-)12-15(-16)x(2.5-)3-4 μ m, with internal, oily-resinous material.....*H. cereus*
(See Dumont, 1980)
13. Ascospores without internal resinous, oily material.....14.
14. Ascospores frequently curved and then indented, with anterior end pointed; apothecial margin with a gelatinous matrix present; ascospores (10-)12-16(-18)x3-4(-5) μ m...*H. erraticus* p. 66.
14. Ascospores if curved, then not indented, with anterior end rounded; apothecial margin lacking a gelatinous matrix; ascospores (14-)16-23(-26)x4-5

(-6) μm*H. caudatus*
(See Dumont & Carpenter, 1981)

1. *Helotium albopunctatum* Peck, Annual Rep. New York State Mus. 31: 47. 1879.

\equiv *Hymenoscyphus albopunctatus* (Peck) Kuntze, Revis. gen. Pl. 3(3): 485. 1898.

NOTES. *Helotium albopunctatum* was described from New York by Peck (1879) as occurring on leaves, and was fully re-described and illustrated by White (1943), who accepted its placement in *Helotium*. It was transferred to *Hymenoscyphus* by Kuntze (1898) and accepted in that genus by both Dennis (1964) and Arendholz (1979). After having studied the type deposited in NYS, I conclude that the species is the same as *Hymenoscyphus caudatus*, and I place it into synonymy and adopt the name *Hymenoscyphus caudatus* which has priority. This species fits well within my concept of *H. caudatus* as presented in Dumont & Carpenter (1981).

Holotype: U.S.A., New York Adirondack Mts., C. H. Peck (NYS, NY).

2. *Helotium caudatum* (Karsten) Velenovský, Monogr. Discom. Bohem. 1: 206. 1934.

\equiv *Peziza caudata* Karsten, Fungi fenn. exs. 547. 1866.

\equiv *Hymenoscyphus caudatus* (Karsten) Dennis, Persoonia 3: 76. 1964.

NOTES. *Helotium caudatum* was originally described by Karsten from a foliicolous collection made in Finland; it is a very common temperate and neotropical species. Dennis (1964), Arendholz (1979), and Dumont & Carpenter (1981) accept the species in *Hymenoscyphus*, a decision with which I concur. For a full description, illustrations, and discussion, see Dumont & Carpenter (1981).

3. *Helotium conocarpi* Seaver & Waterston, Mycologia 34: 517. 1942.

NOTES. *Helotium conocarpi* was originally described from Bermuda on leaves. It was accepted in *Helotium* by White (1943), but neither Dennis (1964) nor Arendholz (1979) treated the species. Dumont (1976) has shown the species to be a member of the genus *Moellerodiscus* (Sclerotiniaceae). See Dumont (1976) for full description, illustrations and discussion.

4. *Helotium dearnessii* (Ellis & Everhart) White, Mycologia 34: 167. 1942. FIGS. 1, 2.

\equiv *Phialea dearnessii* Ellis & Everhart, Proc. Acad. Nat. Sci. Philadelphia 1893: 146. 1894.

\equiv *Hymenoscyphus dearnessii* (Ellis & Everhart) Kuntze, Revis. gen. Pl. 3(3): 485. 1898.

Apothecial morphology — Apothecia minute, scattered or found in small groups, sessile, to ca 1mm in diam, ca 1mm high, when fresh disc drying concave, rehydrating convex. Hymenium when fresh at first subolivaceous,

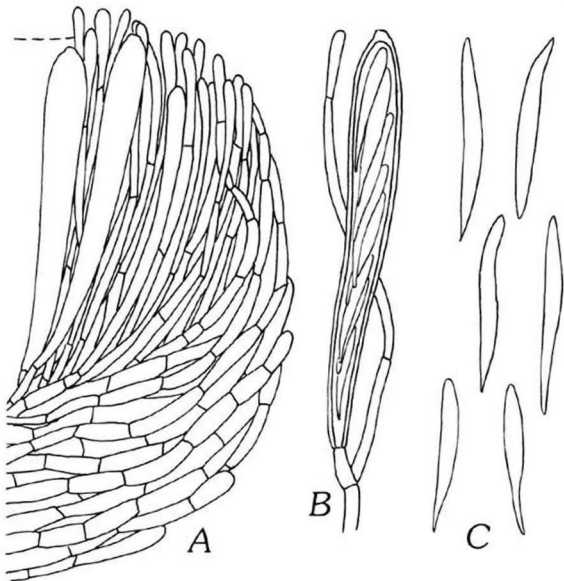


FIG. 1. *Hymenoscyphus dearnessii*, holotype ex NY, freehand drawings, x 1,000. A. Median longitudinal section of an apothecium through margin. B. An ascus and a paraphysis. C. 6 ascospores drawn after discharge from ascus.

becoming yellow, drying flesh-colored to ochraceous, rehydrating lighter and yellowish; receptacle generally concolorous with hymenium when fresh, dry and rehydrated; stipe coloration difficult to detect owing to small size, but appearing dark brown.

Apothecial anatomy — Asci 8-spored, $90-110 \times 9-12 \mu\text{m}$, croziers reported by White to be absent, presence or absence uncertain owing to poor staining of the asci, but possibly present, long-cylindric to clavate, tapering towards the base and there generally not becoming expanded to form a small foot, walls ca $1 \mu\text{m}$ thick, becoming enlarged at the subpapillate to papillate apex and there $3-4 \mu\text{m}$ thick; pore J+, the pore walls staining intensely blue in Melzer's Reagent. Ascospores $28-42 \times 3-5 \mu\text{m}$, details obscured in all collections owing to failure of spores to absorb stain ade-

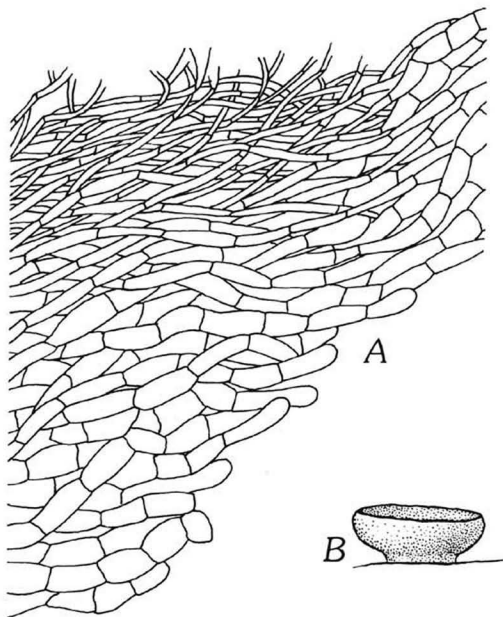


FIG. 2. *Hymenoscyphus dearnessii*, holotype ex NY, freehand drawings. A. Median longitudinal section of an apothecium at approximately juncture of stipe and receptacle, x 1,000. B. Habit sketch of an apothecium on the substrate, x ca 50.

quately, biseriate to irregularly uniseriate, hyaline, smooth, aseptate, elongate, fusoid, anterior end pointed, posterior end attenuated and drawn out to a fine point, a cilium reported by White not observed in these studies, in outline equilateral, frequently sigmoid and curved, guttules spherical to irregular in outline, irregularly dispersed in the individual ascospores and generally filling major portion of ascospores and separated by narrow bands of cytoplasm. Paraphyses equal to or slightly exceeding the asci, internally hyaline, occasionally branching towards the base of the asci, septate, filiform, becoming slightly expanded at the apex and there 2-4 μ m wide, walls thin, smooth and hyaline. Subhymenium absent, without a defined zone or differentiation of tissue beneath the asci. Medullary excipulum poorly developed, in the flanks consisting of narrow, parallel hyphae 2-3 μ m wide, in the center of the receptacle the

parallel hyphae becoming torn apart and interwoven, the individual hyphae hyaline with walls thickened, refractive and smooth. Ectal excipulum: inner ectal excipulum absent. Outer ectal excipulum non-gelatinized, refractive due to thickened walls of hyphae, to ca 40 μ m wide towards the margin and to ca 60 μ m towards the stipe, consisting of hyphae originating in the stipe, and at the juncture of stipe and receptacle extending almost perpendicularly to the surface of the apothecium, continuing towards the margin the individual hyphae extending to the surface at gradually lower angles until at the margin the hyphae parallel to each other and to the surface of the apothecium, the individual cells brick-shaped, (6-)8-12 (-20)x4-8 μ m, the walls thickened, hyaline and smooth, at the surface of the apothecium the apically free cells unmodified and remaining appressed to the surface. Outer covering layer absent. Hairs absent. Margin narrow, consisting of hyphae originating in the receptacle and extending parallel to the surface, as described for the receptacle, but less refractive. Stipe to the outside composed of a pigmented zone of irregularly arranged hyphae, in some areas the individual hyphae losing hyphal orientation, and appearing cellular in composition, the individual cells brick-shaped, globose or angular, lightly to intensely pigmented, the walls thickened, non-gelatinized and obviously roughened; inside the pigmented zone, the hyphae narrow, 2-3 μ m wide, extending parallel to each other and to the surface of the apothecium and continuing into the medullary excipulum of the receptacle, the individual cells hyaline or pigmented light brown, the walls slightly thickened, lightly pigmented or hyaline and generally smooth. Hairs absent.

Habitat: On old stems of *Steironema ciliatum*.

Etymology of the specific epithet: refers to the collector of the original and type specimen.

Holotype: Canada, London, dead stems of *Monarda*, May 1980, J. Dearness 1713 (FH). See additional notes below.

Illustrations: White, *Farlowia* 1: 615, figs. 31-34. 1944. Dennis, *Periconia* 3: 41, fig. 16. 1964.

NOTES. *Phialea dearnessii* was described by Ellis & Everhart from a collection made by J. Dearness (1713) in Canada. As White (1944) has pointed out the type collection was made in May 1890; and the host was originally described as *Monarda*, but Dearness apparently returned to the same locality in June and collected additional material of the new species and concluded that the host was not *Monarda*, but *Steironema ciliatum*. To add to the confusion, Dearness gave the new collection the same number as the original material, but fortunately they both do represent the same species.

Kuntze (1898) transferred *Phialea dearnessii* to *Hymenoscyphus*; White (1944) placed it in *Helotium*. Dennis (1964) accepted its placement in *Hymenoscyphus*, a decision with which I concur. The species appears to be most clearly related to other substipitate species in the genus, such as *H. epiphyllus*. In these species the stipe is composed of hyphae which do not form a well-structured *textura prismatica* or *porrecta*, as in species such as *Hymenoscyphus caudatus* and its related species, but the hyphae become disoriented with the individual cells losing hyphal orientation and also losing the layered effect. In *H. epiphyllus*, *H. immutibile* and *Helotium midlan-*

dense the outermost tissue of the stipe toward the receptacle is formed of *textura globulosa* to *textura angularis*. Further, in these species, the ectal excipulum towards the stipe is formed of hyphae originating in the interior of the stipe and extending nearly perpendicularly or at very high angles to the surface of the apothecium. From the stipe to the margin, the hyphae approach the surface at lower angles until at the margin the hyphae are parallel to the surface. In the *H. caudatus* group the ectal excipulum is formed of hyphae originating in the stipe and running parallel (or at a very low angle) to the surface from the stipe to the margin. *Hymenoscyphus dearnessii* is separated from these other species by the size of its ascospores.

White (1944), in his description of *H. dearnessii*, mentioned that the ascospores produced a delicate cilium 2.5-3.5 μm long. In the portion of the type which I studied, the ascospores only poorly absorbed the various stains utilized; and, for this reason, I was unable to observe certain features of the anatomy of the species. I did not observe the cilium described by White, but agree with him that the ascospores do become obviously attenuated.

5. *Helotium epiphyllum* (Persoon ex Persoon) Fr., Summa veg. Scand. p. 356. 1849.

= *Peziza epiphylla* Persoon, Ann. Bot. (Usteri) 11: 30. 1794.

= *Peziza epiphylla* Persoon, Mycol. Europ. 1: 295. 1822.

= *Hymenoscyphus epiphyllus* (Persoon ex Persoon) Rehm ex Kaufmann, Pap. Mich. Acad. Sci. 9: 177. 1929.

NOTES. *Helotium epiphyllum* is a very widely distributed species in temperate regions. As mentioned by Dumont (1981), the presumed type of *Hymenoscyphus epiphyllus* deposited at L contains only sclerotia of an unidentified fungus. Until the situation of the type of the species is resolved, I am following the concept of the species as set forth by Dennis (1956) and Arendholz (1979), both of whom illustrate the species. Dumont (1981) has fully described and illustrated a tropical variant of the species.

6. *Helotium erraticum* White, Farlowia 1: 606. 1944. FIG. 3.

Apothecial morphology — Apothecia scattered or rarely gregarious, generally produced from leaf blades, small, stipitate, 0.5-1.0mm in diam, to ca 1.0mm high, when fresh cupulate in youth, disc flat with age. Hymenium when fresh white to off-white, drying bright yellow, dull yellow, ochraceous, or rarely reddish; rehydrating translucent, pallid to yellow; margin generally concolorous with margin when fresh, dry and rehydrated; receptacle when fresh white, drying slightly lighter than hymenium or concolorous, yellow to occasionally reddish, rehydrating as hymenium or slightly lighter; stipe generally concolorous with the receptacle when fresh, dry or rehydrated.

Apothecial anatomy — Asci 8-spored, 85-110x8-11 μm , produced from small replicating croziers, broadly cylindrical to long cylindrical-clavate, tapering to the base and there becoming slightly expanded to form a small foot, wall to ca 1 μm thick, enlarged to 2-3 μm at the rounded to truncate apex;

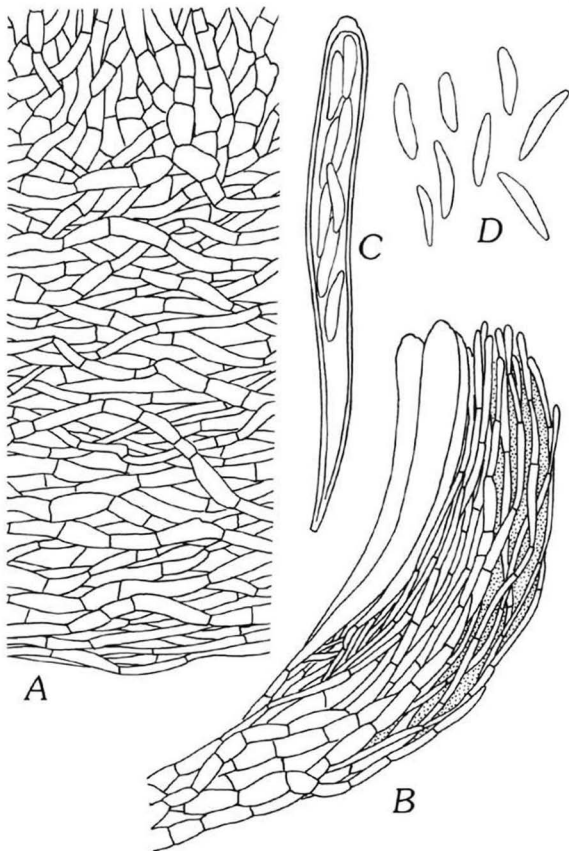


FIG. 3. *Hymenoscyphus erraticus*, freehand drawings, x 1,000. A. Median longitudinal section through an apothecium at approximately midpoint between stipe and margin. B. Median longitudinal section of an apothecium through margin. C. Ascus with 8 ascospores. D. 8 ascospores drawn after discharge from ascus.

pore J+ in Melzer's Reagent. Ascospores (10-)12-16(-18)x3-4(-5) μ m, biserial above and uniserial below, biserial throughout or less commonly obliquely uniserial, hyaline, smooth, aseptate or rarely 1-septate (and then apparently in poorly preserved material), oblong, obovoid, ends rounded to pointed, in outline inequilateral, flattened on one side and generally curved and then slipper-shaped, generally anterior end slightly broader than posterior end, guttules present, more or less equal, irregular in outline, bipolar, filling about half the spore at each end. Paraphyses equal to or rarely slightly exceeding the asci, rarely branched near the base of the asci, sparingly septate, internally hyaline, filiform, becoming slightly expanded at the apex and there 2-4 μ m wide, walls thin, hyaline and smooth. Subhymenium not well differentiated from the medullary excipulum, with a perpendicularly oriented zone to ca 40 μ m in the center staining more intensely than the remainder of the receptacle and composed of parallel, short, broad hyphae and croziers, the individual cells 2-4(-5) μ m wide, the walls thin, hyaline and smooth, these hyphae tightly compact and grading into the upper portion of the medullary excipulum. Medullary excipulum poorly to well developed, hyaline, obconical, non-refractive, consisting in the center of loosely interwoven, branched, septate hyphae, towards the flank and subhymenium becoming more lightly compact and parallel to subparallel, the individual hyphae 2-5 μ m broad, the walls thin, hyaline and smooth. Ectal excipulum not divided into recognizable layers, to the outside the hyphae more tightly compact and narrower than the hyphae in the center of the excipulum; in this region forming a well developed *textura porrecta* to *prismatica*, the individual hyphae hyaline, the walls thickened, hyaline and smooth; these hyphae grading toward and into the narrow hyphae of the medullary excipulum. Margin at its base constructed as the upper portion of the flanks, but soon becoming characteristically gelatinized, the spaces between the individual hyphae 1-2(-3) μ m, the apical cells unmodified. Stipe composed as the flanks, the individual hyphae larger than comparable cells in the flanks. Hairs absent from the margin, flanks, and stipe.

Habitat: On leaf blades of unidentified species, leaves of *Amelanchier* sp., *Ulmus* sp., *Hamelis* sp. and *Populus* sp., and last year's pods of *Robinia pseudo-acacia*.

Etymology of the specific epithet: relevance to the fungus uncertain.

Holotype: U.S.A., New York, Coy Glen, near Ithaca, on decaying leaves, 3 Oct 1938, H. H. Whetzel & W. L. White s.n. (White herb nos. 3421 and 3416, one collection apparently divided into two packets or two specimens collected on the same day with identical data.)

Paratype specimens: U.S.A., New York, Malloryville, W. of North Bog, on leaves of *Amelanchier* sp., *Ulmus* sp., and *Hamelis* sp., 18 Oct 1941, H. H. Whetzel & Niederhauser s.n. (FH ex CUP 29658); Newfield Gorge, Ithaca, last year's pods of *Robinia pseudo-acacia*, 26 Oct 1941, H. H. Whetzel & T. Sproston s.n. (FH ex CUP 29666); Canada, Quebec, Duchesnay, County Portneuf, on leaves of *Populus* sp., 25 Aug 1938, H. H. Whetzel & T. Sproston s.n. (FH ex CUP 27850).

Illustrations: White, Farlowia 1: 607, figs. 14-18. 1944. Arendholz, Morphologisch-taxonomische Untersuchungen an blattbewohnenden Ascomyceten aus der Ordnung der Helotiales, pl. 15, fig. 4. 1979.

NOTES. *Helotium erraticum* was described by White (1944) who indicated that the species was closely related to members of the *Helotium epiphyllum* group: *H. epiphyllum* *H. immuta-*

bile, *H. carpinicola*, and *H. midlandense*. These latter four share in common one feature which is absent in *H. erraticum*. In all four the ectal excipulum at the juncture of the receptacle and stipe is composed of globose cells to the outside, with hyphae progressing gradually from high angles to the surface in the lower portion to parallel at the margin. I have found no globose cells in any of the collections studied here of *H. erraticum*.

I conclude that *H. erraticum* is closely related to *Hymenoscyphus caudatus*. In *Helotium erraticum* the ascospores are generally curved and frequently indented or slipper-shaped, while in *Hymenoscyphus caudatus* the ascospores are inequilateral and generally not curved and when curved, not slipper-shaped. A high proportion of the ascospores in *Helotium erraticum* have ascospores with pointed apices, while this feature is absent in *Hymenoscyphus caudatus*. In *H. caudatus* the margin is composed of thin-walled, brick-shaped cells, while in *Helotium erraticum* the margin is unusual for a member of *Hymenoscyphus*. The margin is composed of narrow cells apparently imbedded in a gelatinous matrix; at times, however, the hyphae are tightly compact, and it is difficult to interpret the structure, and the walls merely appear thickened. In one collection (CUP 29666) the hyphae in the ectal excipulum are narrower than normal and very thick-walled, and the ascospores are 1-septate. The collection appears to be badly preserved, which may account for the abnormal formation of the sterile tissue and the septation in the ascospores. All other features are characteristic for *Helotium erraticum*. The shape and size is right for the species, as is the characteristic subhymenium. I conclude that this is merely a variant collection.

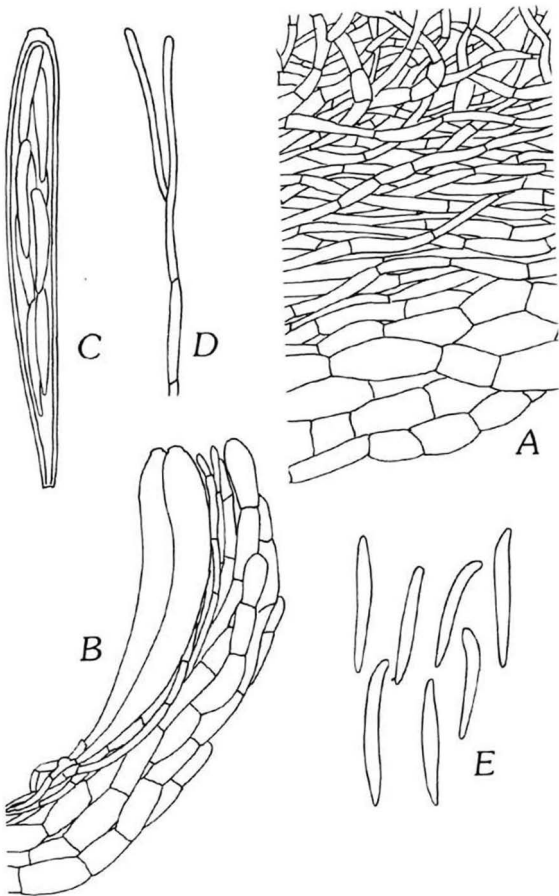
I am somewhat concerned about the placement of this species in *Hymenoscyphus*, since it produces a margin with at least some hyphae in a gelatinous matrix. This feature is more characteristic of the genus *Crociareas* than it is of *Hymenoscyphus*. However, the ectal excipulum is that of an *Hymenoscyphus*. Dennis (1964) did not treat the species, while Arendholz (1979) did place it in *Hymenoscyphus* as *H. erraticus*. I conclude that the species is perhaps intermediate between the two, but shows more affinities to *Hymenoscyphus* than to *Crociareas*. I will, thus maintain it in *Hymenoscyphus*.

7. *Helotium fastidiosum* Peck, Annual Rep. New York State Mus. 27: 107. 1875. FIG. 4.

Apothecial morphology — Apothecia scattered, stipitate, 1-1.5mm in diam, 1-1.5(-3)mm high, when fresh disc flat, drying flat to slightly cupulate, rehydrating convex. Hymenium when fresh off-white to pale yellow, drying darker, yellow, ochraceous to flesh-colored, rehydrating lighter, slightly translucent; receptacle when fresh generally slightly darker than hymenium and white to dark yellow, drying and rehydrating darker than hymenium; stipe narrow, cylindrical, when fresh above concolorous with the lower portion of the receptacle, becoming darker toward the base, drying darker, especially toward the base, or rehydrating lighter and pallid, darker below.

Apothecial anatomy — Asci 8-spored, (75-)85-100x9-12 μ m, produced from small croziers, long cylindric-clavate, tapering gradually to the base and there occasionally becoming slightly expanded to form a small foot, wall to ca 1 μ m thick, enlarged at the papillate apex and there 2-3 μ m thick; pore J+, visible as two, dark blue apical dots in Melzer's Reagent. Ascospores (23-)26-30(-35)x2.5-3.5(-4.5) μ m, obliquely uniseriate to irregularly arranged, hyaline, smooth, aseptate, subfusoid to clavate, apical end hooked and pointed, basal end pointed, in outline inequilateral, curved and rounded in the upper portion, flattened and straight below, anterior end broader than posterior, with a single row of spherical to irregular guttules filling majority of the spores. Paraphyses equal to or slightly exceeding the asci by 5 μ m, internally hyaline and devoid of pigmented contents, branching occasionally toward the base of the asci, septate, filiform, not expanded at the apex and there 2-3 μ m wide, walls thin, smooth and hyaline. Subhymenium not well differentiated from the medullary excipulum, poorly developed, with an indistinct hyaline zone beneath the asci with hyphae more tightly compact than the medullary excipulum below. Medullary excipulum poorly developed in the flanks and in the center of the apothecium, hyaline, non-refractive, consisting of septate, branched, vertically oriented to tightly interwoven hyphae 3-6 μ m wide, the walls thin, hyaline and smooth. Ectal excipulum: inner ectal excipulum well defined and well differentiated from the outer ectal excipulum and grading into the medullary excipulum, entire layer hyaline, non-refractive, to ca 10 μ m wide toward the margin and to 25 μ m toward the stipe, consisting of tightly compact, hyaline hyphae 2-4 μ m wide, with walls thin, non-refractive, hyaline and smooth. Outer ectal excipulum non-gelatinized, non-refractive or rarely appearing slightly refractive due to thickened walls, to ca 20 μ m broad toward the margin and to ca 30 μ m toward the stipe, consisting predominantly of a well defined *textura prismatica* with the individual hyphae extending parallel to the surface of the apothecium and without apically free hyphal tips; the individual cells toward the margin 8-15x4-8 μ m, 12-25x5-8 μ m toward the stipe, the walls thin or rarely slightly thickened, hyaline and smooth. Outer covering layer present or becoming detached and the apothecium then appearing naked, consisting of 1-2 layers 2-5 μ m broad, the individual hyphae extending parallel to the surface, overlapping, terminating before the margin, the apically free cells remaining appressed to the surface of the apothecium and unmodified, the individual cells hyaline or rarely light yellow-brown, the walls non-refractive, hyaline to light brown, and smooth. Hairs absent. Margin generally poorly developed, narrow above, broader below, entire zone hyaline or light brown, constructed similarly to the receptacle below, the individual cells approximately same size as upper receptacle or slightly smaller, apical cells unmodified. Stipe in the upper portion constructed similarly to the lower portion of the receptacle and apothecial flank, at approximately midpoint and to the outside 1-2 layers of narrow, hyaline or lightly pigmented hyphae (often difficult to detect) with thin, lightly to intensely pigmented and smooth or rarely roughened walls, to the inside a zone of *textura prismatica* with the individual cells hyaline to light brown, in the central core the hyphae narrow, parallel to slightly interwoven; hairs absent.

FIG. 4. *Hymenoscyphus fastidiosus*, holotype ex NYS, freehand drawings, x 1,000. A. Median longitudinal section of an apothecium at approximately midpoint between margin and stipe. B. Median longitudinal section of an apothecium through margin. C. An ascus with 8 ascospores. D. A branching paraphysis. E. 7 ascospores drawn after discharge from ascus.



Habitat: On leaves of *Alnus incana*, *Alnus* sp., and catkins of *Alnus*.

Etymology of the specific epithet: relevance to the fungus uncertain.

Holotype: U.S.A., New York, Forestburgh, on fallen petioles of *Alnus* leaves, Sept., C. H. Peck s.n. (NYS).

Additional specimens examined: U.S.A., New York, Adirondack Mts., on leaves of *Alnus* sp., date not given, C. H. Peck s.n. (NYS); Labrador Lake, near Tully, on old overwintered leaves of *Alnus incana*, 26 Aug 1935, H. H. Whetzel & W. L. White s.n. (CUP 24817 ex FH); Lloyd Preserve, McLean, on *Alnus incana*, leaves on ground, 6 Sept 1935, H. H. Whetzel, W. L. White & Rogers (White 2042 ex FH). Oregon, Tilly Jane Creek, Mt. Hood, on dead alder leaves and petioles, 26 Aug 1933, J. Kienholz K 137 (BPI ex FH). CANADA, Quebec, Duchesnay, County Portneuf, on petioles and lower midribs of *Alnus* leaves, 26 Aug 1938, H. H. Whetzel s.n. (FH); locality as previous collection, 24 Aug 1938, H. H. Whetzel s.n. (FH).

Illustrations: White, *Mycologia* 34: 158, fig. 2; 166, fig. 9. 1942. White, *Farlowia* 1: 153, fig. 7. 1943. Arendholz, *Morphologisch-taxonomische Untersuchungen an blattbewohnenden Ascomyceten aus der Ordnung der Helotiales*, pl. 15. fig. 2. 1979.

NOTES. *Helotium fastidiosum* was described by Peck from the U.S.A. as occurring on leaves of *Alnus*. The species does not produce a stroma and is surely referable to *Hymenoscyphus* as reported recently by Arendholz (1979) who also reported it for the first time from Europe. Structurally the sterile tissue of the apothecium is similar to the common species *Hymenoscyphus caudatus* and *Hymenoscyphus serotinus*, while the ascospore shape is like that of *H. serotinus*. Dumont & Carpenter (1981) pointed out that until recently *H. serotinus* was generally regarded as a species occurring on herbaceous stems. However, they have demonstrated from their studies of neotropical collections that the species is extremely variable, and occurs on leaves as well as on stems. They further reported the ascospores to be (16-)18-23(-30)x3-3.5(-5) μ m and indicated that the base of the stipe generally stains light pink in Melzer's Reagent, a reaction also observed in the material studied here. Dennis (1964) reported the ascospore measurements for *H. serotinus* to be 18-28x3-4 μ m, while Seaver reported them to be 4x22-24 μ m & Rehm (1893 in 1887-1896) found them to be 30-36x4-6 μ m. The ascospores studied in the present investigation of *Helotium fastidiosum* were (23-)26-30x2.5-3.5(-4.5) μ m and would thus appear to fall within the reported range for *Hymenoscyphus serotinus*, and is a probable synonym of *H. serotinus*. I have been unable to find a type of *H. serotinus* and am thus following the generally accepted concept as presented by Dennis (1956, 1964). A final decision on the tentative synonymy can be made only after type material is located or a neotype designated for *H. serotinus*.

8. *Helotium fraternum* Peck, Annual Rep. New York State Mus. 32: 47. 1879.

= *Hymenoscyphus fraternus* (Peck) Dennis, *Persoonia* 3: 76. 1964.

NOTES. *Helotium fraternum* was originally described from New York State as occurring on leaves, and was accepted by

White (1942). Dennis (1964) transferred the species to *Hymenoscyphus*; Arendholz (1979) did not treat the species. Dumont (1981) redescribed and illustrated the species. He found that the species formed a well-defined substratal stroma, produced an ectal excipulum with hyphae embedded in a gelatinous matrix and transferred the species to *Poculum* (Sclerotiniaceae). For a full description, illustrations and discussion, see Dumont (1981).

9. *Helotium immutabile* Fuckel, Jahrb. Nassauischen Vereins Naturk. 25-26: 338. 1871. FIG. 5.

= *Hymenoscyphus immutabilis* (Fuckel) Dennis, Persoonia 3: 76. 1964.

Apothecial morphology — Apothecia scattered, arising from leaf blades, short stipitate, ca 1.0-1.5mm in diam, to ca 0.5-0.75mm high, when fresh flat to slightly convex, drying same, rehydrating flat to slightly cupulate. Hymenium when fresh white, drying pale yellow, yellow-brown, or reddish orange, rehydrating slightly lighter and pallid; margin concolorous with hymenium; receptacle when fresh white, drying pale yellow-orange, rehydrating lighter and concolorous in the lower portion with the upper portion of the stipe; stipe in the upper portion similar to receptacle, coloration difficult to observe owing to small size.

Apothecial anatomy — Asci (70-)80-100(-105)x(6-)8-9 μ m, produced from small croziers, clavate to clavate-cylindric, gradually tapering toward the base and there becoming expanded to form a small foot; wall ca 1 μ m wide; pore J+. Ascospores (7-)8-10(-12)x2.5-3.5(-4.5) μ m, generally biseri-ate, but occasionally uniseriate, hyaline, smooth, obovoid to trapezoidal, anterior end round or rarely pointed, posterior generally slightly pointed, in outline more or less equilateral, if inequilateral only slightly so and not flattened on one side, walls slightly thickened, eguttulate or less commonly with polar guttulate areas (which are generally obscured and difficult to detect) or in youth biguttulate with guttules disappearing with age. Paraphyses equal to or slightly exceeding the asci, internally hyaline, branching toward the base of the asci, septate, filiform, becoming slightly expanded at the apex and there 2-3 μ m wide, walls thin, smooth and hyaline. Subhymenium not well differentiated from the medullary excipulum, consisting of parallel, vertically oriented hyphae grading into medullary excipulum. Medullary excipulum well developed, non-refractive, hyaline to pigmented light brown, consisting of septate, branched, parallel or loosely to tightly interwoven hyphae 2-4 μ m wide, the walls thin, non-refractive, hyaline or rarely pigmented light brown, and smooth. Ectal excipulum: inner ectal excipulum poorly defined and differentiated from medullary excipulum, entire layer non-refractive, hyaline to pigmented light brown, consisting of tightly compact, more or less parallel, hyaline hyphae 2-3 μ m wide, the walls thin, non-refractive, hyaline and smooth. Outer ectal excipulum non-refractive, entire layer hyaline to subhyaline, variable to ca 35 μ m broad toward the margin and the same toward the stipe, at the intersection of the stipe and the receptacle to the outside composed of thin-walled, globose cells comprising a well developed textura globulosa, to the inside the hyphae parallel, above the juncture the isodiametric cells giving way to hyphae originating in the stipe and extending almost perpendicularly or at very high angles to the surface, progressing toward the margin the hyphae extending at lower and lower angles to the surface until at the margin the hyphae parallel to the surface of the apothecium, the individual cells with walls thin to slight-

ly thickened, hyaline or rarely pigmented light brown and smooth. Outer covering layer as a distinct layer absent, but with an area to the outside formed of the apical cells of the hyphae extending at low to high angles to the surface and continuing to the surface, and then becoming somewhat expanded and frequently large and clavate, these cells hyaline to slightly yellow-brown with walls thin to slightly thickened and frequently roughened. Margin constructed similarly to the upper portion of the flank, with the individual cells smaller, the apical cells generally unmodified. Hairs absent on the receptacle and margin. Stipe in the upper portion constructed similarly to the lower portion of the receptacle, below to the outside the cells globose with parallel narrow hyphae to the inside; hairs absent.

Habitat: Leaves of *Populus tremula*, *P. nigra*, *Quercus pedunculata*, *Quercus* sp., *Ulmus* sp., *Robinia pseudo-acacia*.

Etymology of the specific epithet: relevance to the fungus uncertain.

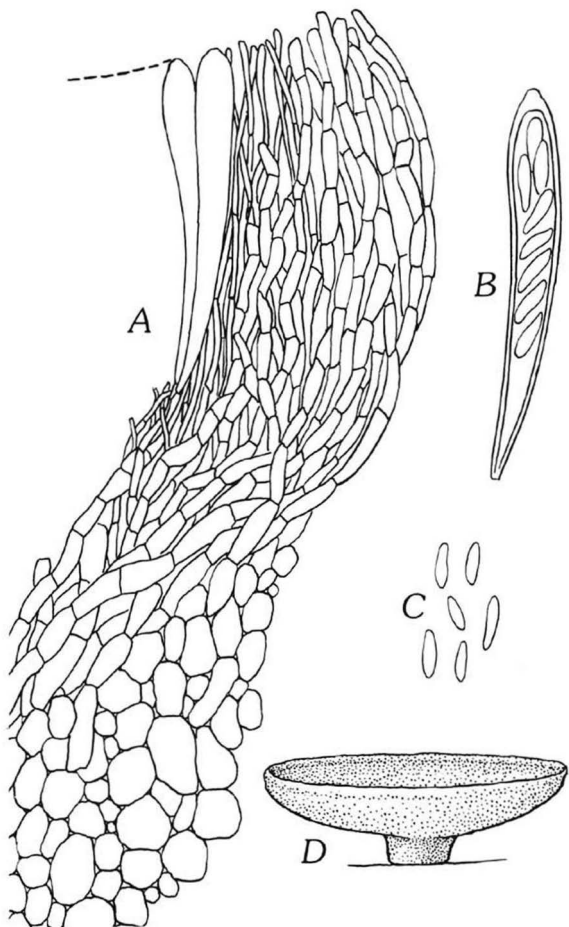
Holotype: Germany, Boss Pr. Ebenbach, ad *Populi tremulae* folia putrida, Fuckel (Fuckel, Fungi rhenani no. 2388, NY probably distributed in Herbarium Barbey-Boissier no. 1215.).

Illustrations: Dennis, Mycol. Pap. 62: 93, fig. 85. 1956. White, Farlowia 1: 143, fig. 3. 1943. Arendholz, Morphologisch-taxonomische Untersuchungen an blattbewohenden Ascomyceten aus der Ordnung der Helotiales, pl. 11, figs. 2, 3. 1979.

NOTES. According to White (1943) and Arendholz (1979), *Helotium immutabile* is a widely distributed species in Europe. Dennis (1964) transferred the species to *Hymenoscyphus*, a decision with which I agree, and it appears most closely related to *Helotium epiphyllum*, since both have similar structure of the sterile tissue of the apothecium; the outermost layers of the stipe and base of the receptacle are composed of globose cells, and the receptacle is composed of hyphae originating at the stipe and at the base extending almost perpendicularly to the surface. Further, from the stipe to the margin the hyphae extend at progressively lower angles until toward the margin the hyphae are parallel to the surface.

I have studied few collections of *H. immutabile*; but based on the shape of the ascospores, it appears distinct from *H. epiphyllum*. There is overlap in the measurements, *H. epiphyllum* 12-24x3-5 μ m (combined measurements according to Dennis, 1956, White, 1943 and Arendholz, 1979) and in *H. immutabile* they are said to be 10-13x4-5 μ m. It would appear that ascospore size is of little value in separating these two species. I have attempted to locate the type specimen of *Peziza epiphylla* in the Persoon herbarium at L, but no apothecia remain on the leaf, and all that is present in the type are several apparently unrelated sclerotia, probably non-Sclerotiniaceae. Thus, a final decision on the correct placement of *H.*

FIG. 5. *Hymenoscyphus immutabilis*, Fuckel, Fungi rhenani 2388 ex NY, free-hand drawings. A. Median longitudinal section through an apothecium showing lower portion of receptacle and margin, x 1,000. B. An ascus with 8 ascospores. C. 6 ascospores drawn after discharge from ascus, x 1,000. D. Habit sketch of an apothecium on the substrate, x ca 50.



epiphyllum and its relationship with *H. immutabile* can be made only after a neotype specimen is designated for *Peziza epiphylla*, a chore I am not now prepared to do.

10. *Helotium linderi* White, Farlowia 1: 154. 1943.

NOTES. White described *Helotium linderi* from collections from the U.S.A. Dennis (1964) did not treat the species, while Arendholz (1979) placed *H. linderi* into synonymy with *Hymenoscyphus caudatus*, a decision with which I concur.

Holotype: U.S.A., Tenn., Chimney Trail, Great Smokey Mts. Nat. Park, 3200 ft., on fallen leaves, 18 Aug 1939, D. H. Linder (FH).

11. *Helotium midlandense* White, Farlowia 1: 605. 1944.

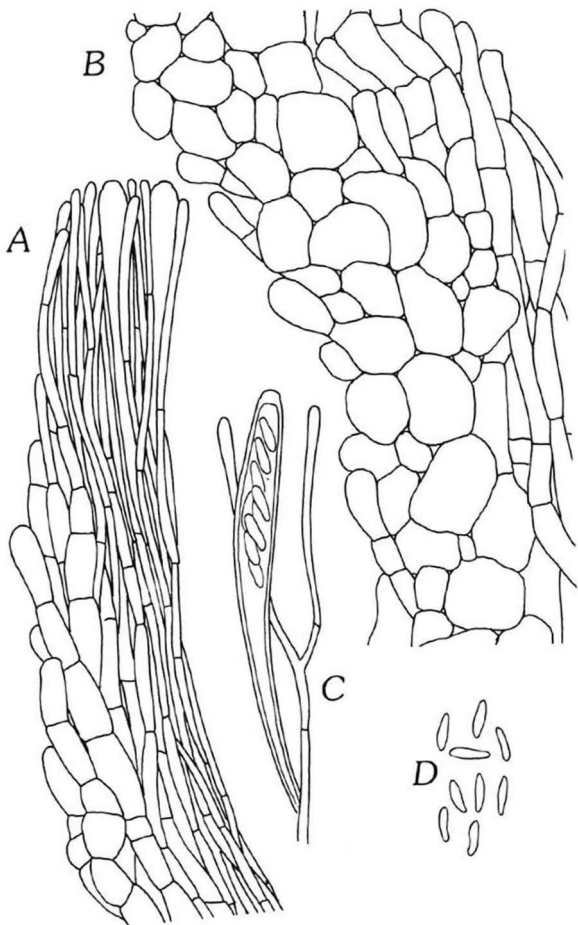
FIG. 6.

Stroma — Substratal, visible on the fruits only as blackened areas of the substrate; in section the blackened areas composed of an irregularly formed rind composed of irregular to epidermoid cells in face view, also visible in cross section of the base of the stipe of the apothecium; not known in culture.

Apothecial morphology — Apothecia solitary, gregarious or occasionally arising in clusters of 3-7, stipitate, unknown in fresh condition, when dry ca 0.5mm in diam and ca 0.5mm high, disc drying flat to slightly cupulate, rehydrating flat and expanded. Hymenium drying shades of yellow to flesh-colored, rehydrating glassy-translucent; margin drying lighter than hymenium, rehydrating concolorous with hymenium; receptacle drying concolorous with the margin, lighter than hymenium, rehydrating glassy-translucent and concolorous with hymenium; stipe concolorous with receptacle in dry and rehydrated conditions, cylindrical, short, generally less than 0.5 mm high and wide, occasionally subpapillate in appearance.

Apothecial anatomy — Asci 8-spored, (40-)52-58(-65)x(5-)6-7 μ m, produced from small, replicating croziers, cylindrical to slightly clavate, gradually tapering toward the base and there becoming slightly expanded to form a small foot, wall to ca 1 μ m thick, slightly enlarged at the apex and there to ca 2 μ m thick; pore apparently J- in Melzer's Reagent. Ascospores (5-)6-8(-9)x2-3 μ m, obliquely uniseriate, irregularly biseriate or biseriate throughout, hyaline, smooth (a few devoid of pigment possibly lightly punctate), aseptate, obovoid to obpyriform, ends rounded or rarely slightly pointed, in outline generally equilateral, anterior end obviously broader than posterior end, occasionally slightly indented below median point of spores; guttules generally absent or rarely with one tiny (ca 1 μ m or less), transversely positioned at the narrowest portion of the spore and appearing as a septum. Paraphyses equal to or slightly exceeding the asci, internally hyaline, branching toward the base of the asci, septate, cylindrical, generally not becoming expanded at the apex and there 2-3 μ m wide, walls thin, smooth, and hyaline. Subhymenium well developed, well

FIG. 6. *Helotium midlandense*, Stevens 59 ex NY, freehand drawings, 1,000. A. Median longitudinal section of an apothecium through margin. B. Median longitudinal section of an apothecium through the juncture of receptacle and stipe. C. An ascus with 8 ascospores and a paraphysis. D. 9 ascospores drawn after discharge from ascus.



differentiated from the medullary excipulum below, hyaline, to ca 50 μ m broad in the center, narrower toward the margin, consisting of parallel to slightly interwoven, vertically interwoven hyphae, the individual hyphae hyaline, 1.5-3 μ m wide, the walls thin, hyaline and smooth. Medullary excipulum poorly developed, obconical, non-refractive, hyaline, consisting of septate, branched, loosely to tightly interwoven (to parallel in the flank) hyphae 2-3(-4) μ m wide, the walls thin, hyaline and smooth. Ectal excipulum: inner ectal excipulum well to poorly defined, well-differentiated from the outer ectal excipulum and grading into the medullary excipulum, entire layer non-refractive, hyaline, 15-20 μ m broad toward the margin and approximately the same toward the stipe, consisting of parallel to slightly interwoven, hyaline hyphae 2-3 μ m wide, the walls thin, non-refractive, hyaline and smooth. Outer ectal excipulum non-gelatinized, non-refractive, entire layer hyaline, toward the stipe ca 50 μ m broad, consisting of large, globose, angular to irregular cells forming a well defined *textura globulosa* to *angularis*, the individual cells (5-)8-12(-20) μ m wide, the walls thin to slightly thickened, hyaline and smooth. Outer covering layer and hairs absent; progressing towards the margin the outer ectal excipulum becoming narrower, and grading from a *textura angularis* and *globulosa* to *textura prismatica*, the individual cells smaller and forming more and more regular rectangularly shaped cells until at the margin the layer composed of a well defined *textura prismatica*, lacking an outer covering layer. Margin simple, the brick-shaped cells if continuing into margin only a short distance and the remainder simple and composed of the narrow cells of the inner ectal excipulum and the adjoining paraphyses, hairs also absent in the margin. Stipe constructed similarly to the lower portion of the receptacle, to the outside a zone to ca 70 μ m broad composed of globose to angular cells and grading into narrow, parallel hyphae in the interior of the stipe; the individual cells with walls thin to slightly thickened, hyaline and smooth or rarely roughened; at the base rind cells visible; hairs absent.

Habitat: Old pods of *Gleditsia triacanthos* L. and reported by White (1944) to occur also on petioles and midveins of *Quercus* sp.

Etymology of the specific epithet: refers to the part of the country where the fungus was originally collected.

Holotype (not examined): Iowa, Homestead, pods of *Gleditsia triacanthos*, 26 Sept 1931, G. W. Martin 5186 (FH).

Illustrations: White, Farlowia 1: 607, figs. 10-13. 1944.

Specimen examined: Kansas, Lawrence, on pods (? of *Gleditsia* sp.), 1890, W. C. Stevens 59 (paratype ex NY).

NOTES. When White (1944) described *Helotium midlandense*, he noted that the stipe to the outside was composed of globose to angular cells, and he illustrated the ectal excipulum as formed of brick-shaped cells in the lower portion approaching the stipe. I have examined a paratype collection and conclude that the majority of the ectal excipulum is composed of a well defined *textura globulosa* to *angularis*, but that toward the margin the tissue becomes more organized into a well defined *textura prismatica*, and not toward the stipe as illustrated by White. Further, I have found a well developed rind on the substrate in association with the apothecia and in section have observed rind cells at the base of the stipe. I have also noted that the spores are relatively small for the *Scler-*

otiniaceae and are obovoid to obpyriform. Since the species produces a substratal stroma, it should be referred to the Sclerotiniaceae. Because (i) the ectal excipulum is composed of a *textura globulosa*, (ii) small pyriform ascospores are present, and (iii) it occurs on fruits, the species is probably best referred to *Ciboria*. At present, I am uncertain of species concepts in *Ciboria* and of the separation between *Ciboria* and *Moellerodiscus*; I will not make a new combination in *Ciboria*, but will defer action until the species are better known.

I have observed in several apothecia a rather high proportion of misshapen and malformed ascospores, some of which produced an apparent septum. I cannot satisfactorily explain the presence of the irregularly formed ascospores, nor the occasional production of a single septum.

My observations are then different from White's, who did not report the presence of a stroma and referred it to the Leotiaceae (*Helotium*). Neither Dennis (1964) nor Arendholz (1979) treat the species.

12. *Helotium phyllogenon* Rehm, *Hedwigia* 24: 14. 1885.

≡ *Hymenoscyphus phyllogenon* (Rehm) Kuntze, *Revis. gen. Pl.* 3(3): 485. 1898.

NOTES. *Helotium phyllogenon* was originally described from Hungary on leaves of "poplar." Kuntze (1898) transferred the species to *Hymenoscyphus*, and the placement there was accepted by Dennis (1964) and by Arendholz (1979). I find the type collection to be indistinguishable from *H. caudatus*, following the concept of Dumont & Carpenter (1981). They gave the ascospore measurements of *H. caudatus* to be (14-)16-23(-26)x4-5 (-6)µm, whereas in the original description of *H. phyllogenon* the ascospore measurements were 12-15x3.5µm. White (1943) gave them as 11-14x3.8-5µm, Dennis (1956) 14-16x4-6µm, Arendholz (1979) 11-15x3.5-4.5µm. I find the apothecial structure of the type of *H. phyllogenon* indistinguishable from *H. caudatus*, and conclude that they represent the same species, with the ascospore measurements of the type of *H. phyllogenon* falling at the lower limits of *H. caudatus* and perhaps extending the limits somewhat lower than previously reported by Dumont & Carpenter (1981).

Holotype: Hungary: bei Ungerisch-Altenburg (Ungarn), an faulen Pallel-Blättern, Oct 1883, Linhart (ex FH, Rehm, *Ascomyceten* 768).

13. *Helotium phyllophilum* (Desmazières) Fries, *Summa veg. Scand.* p. 356. 1849.

≡ *Peziza phyllophila* Desmazières, *Ann., Sci. Nat. Bot., Sér. 2*, 17: 98. 1842; *Pl. Crypt (du Nord) France* éd. 1. no. 1159. 1842; éd. 2. Sér. 1. no. 659. 1842.

≡ *Hymenoscyphus phyllophilus* (Desmazières) Kuntze, *Revis. gen. Pl.* 3(3): 485. 1898.

NOTES. *Peziza phyllophila* was originally described as occurring on leaves of *Acer* and *Fagus* from France and was trans-

ferred to *Helotium* by Fries and to *Hymenoscyphus* by Kuntze (1898). Its placement in *Hymenoscyphus* was accepted by both Dennis (1964) and Arendholz (1979). I find the sterile tissue of the apothecium of the lectotype specimen of the species to be indistinguishable from that of *H. caudatus*. The apothecia of *H. phyllophilus* were originally thought to be smaller than those of *H. caudatus*, but as Dumont & Carpenter have pointed out, the apothecia of tropical collections are frequently less than 1mm in diameter and those of the type of *H. phyllophilus* would then fall within the range of *H. caudatus*.

In the original description the measurements of the ascospores were not given, but they were reported to be 1-septate. Dennis (1956) reported the ascospores to be 14-16x3.5-4µm, White (1943) gave them as 11.5-15x3.2-4µm, and Rehm (1893, in 1887-1896) had them 10-15x3-3.5µm. Rehm further placed *Helotium phyllophilum* and *H. phyllogenon* into synonymy. The only difference which I consider noteworthy between *Hymenoscyphus caudatus* and *H. phyllophilus* is that in the latter all of the ascospores in the type collection are 1-septate, whereas, as pointed out by Dumont & Carpenter (1981), in *H. caudatus* they are only rarely 1-septate. However, I have noted in other clearly marked species, such as *H. rufocorneus* (see Dumont, 1981) where for some unexplained reason all of the ascospores in a single collection become 1-septate. I do not consider this then to represent the single criterion upon which to separate two species, *H. caudatus* and *H. phyllophilus*. Rather, I prefer to consider them as probable synonyms.

There is some question as to where the name "*Peziza phyllophila*" was actually first published and as to what the type specimen actually is. Desmazières (1842) presented a formal description of the species and a description also appeared on the labels of two specimens cited by Desmazières; "Pl. Crypt. éd. 1, no. 1159; éd. 2, no. 659." As nearly as can be determined (Sayre, 1969), fascicles containing these two specimens were issued simultaneously. I have been unable to determine if the article appeared before the exsiccati specimens.

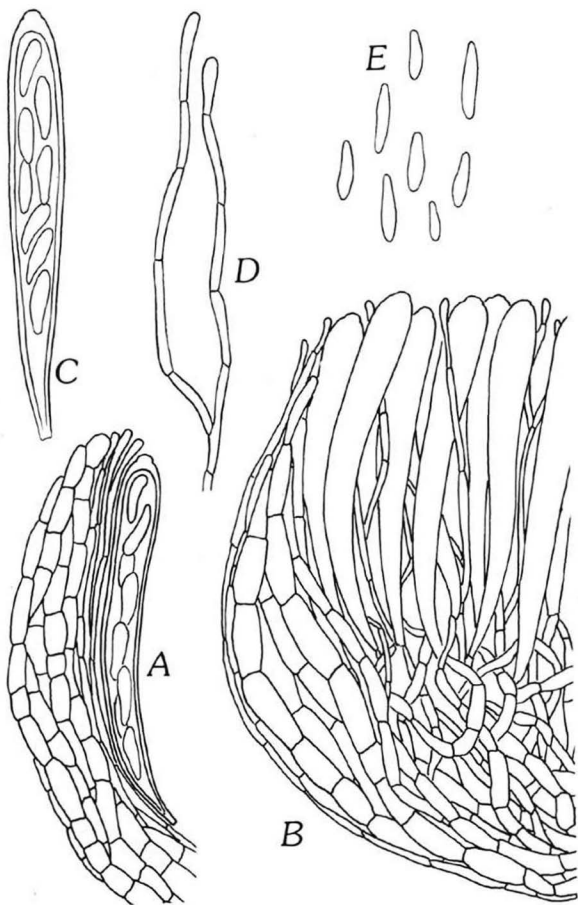
In this regard, White (1943) selected no. 1159 as the type, but indicated that he examined two specimens, one in the bound set of the exsiccata at FH and the other from the Curtis collections. I select the portion deposited in the bound exsiccata as the lectotype specimen for *Peziza phyllophila*.

Lectotype specimen: Plantes Cryptogames de France, éd. 1, no. 1159 (ex FH, bound set).

14. *Helotium translucens* White, Farlowia 1: 149. 1943.

FIG. 7.

FIG. 7. *Hymenoscyphus translucens*, holotype ex FH, freehand drawings, x 1,000. A. Median longitudinal section of an apothecium through margin. B. Median longitudinal section of an apothecium through margin. C. An ascus with 8 ascospores. D. A branching paraphysis. E. 8 ascospores drawn after discharge from ascus.



= *Hymenoscyphus translucens* (White) Arendholz, Morph.-tax. Untersuch. blattbe. Ascomyceten Helotiales p. 78. 1979.

Apothecial morphology — Apothecia tiny, solitary to occasionally gregarious, arising from leaf blades, especially veins of various sizes, with a minute stipe, to ca 0.5mm in diam and to ca 0.4mm high, when fresh disc flat, drying and rehydrating same. Hymenium pallid to hyaline when fresh, drying off-white to pale ochraceous, rehydrating lighter, pale flesh-colored to off-white; margin concolorous with hymenium when fresh, dry and rehydrated; receptacle generally concolorous with the hymenium when fresh, dry and rehydrated; stipe, difficult to observe coloration owing to minute size.

Apothecial anatomy — Asci 8-spored, 80-95x9-11 μ m, probably produced from tiny croziers, long cylindric-clavate, gradually tapering toward the base and there not expanded to form a small foot, wall ca 1 μ m wide, slightly enlarged at the rounded to subtruncate apex and there 2(-3) μ m thick; pore J+. Ascospores (10-)11-14(-15)x(3-)4(-5) μ m, uniseriate to biseriate throughout, hyaline, smooth, aseptate or very rarely 1-septate, trapezoidal, obovoid, ends rounded or less commonly slightly pointed, in outline generally equilateral, if inequilateral only slightly so and then not flattened on one side, anterior end broader than posterior end, frequently appearing eguttulate or with two polar guttulate areas generally obscure but granular and oily in composition, walls slightly thickened and occasionally to ca 1 μ m wide. Paraphyses equal to or slightly exceeding the asci, internally hyaline, branching at the base of the asci and toward the middle of the asci, filiform, becoming slightly expanded at the apex and there (2-)3-4 μ m wide, walls thin, smooth and hyaline. Subhymenium not well differentiated from the medullary excipulum, but with a tendency of the hyphae at the base of the asci to be slightly more compact and narrower than the hyphae in the center of the receptacle. Medullary excipulum poorly developed, obconical, non-refractive, hyaline, consisting of septate, branched, more or less parallel to slightly interwoven hyphae 2-4 (-5) μ m wide, the walls thin, non-refractive, hyaline and smooth. Ectal excipulum: inner ectal excipulum poorly defined and differentiated from medullary excipulum. Outer ectal excipulum non-refractive, non-gelatinized, entire layer hyaline to subhyaline, ca 15-20 μ m wide toward the margin and to ca 20-25 μ m toward the stipe, consisting predominantly of *textura prismatica* with the individual hyphae extending parallel to or at low to high angles to the surface of the apothecium and without apically free hyphal tips; the individual cells frequently collapsing, becoming disoriented owing to drying process and tissue then appearing to be composed of narrow hyphae; individual cells toward the margin 5-12(-15)x3-5 μ m and 12-22x3-7 μ m toward the stipe, the walls thin to slightly thickened and ca 1 μ m broad, hyaline to light brown and smooth. Outer covering layer present, but difficult to detect, 1-2 hyphal layers and 1-3 μ m broad, the individual hyphae extending parallel to the surface of the apothecium, overlapping, terminating before the margin, the individual cells subhyaline, the walls non-refractive, thin, pigmented light brown to intense brown, smooth to frequently slightly roughened. Hairs absent. Margin poorly developed, narrow above, broader below, entire zone hyaline to light brown, constructed similarly to the flank below, the individual cells smaller. Stipe in the upper portion constructed similarly to the lower portion of the flanks, not becoming noticeably interwoven in the center, toward the base the individual hyphae becoming disrupted, and frequently losing hyphal orientation, but generally oriented at a high angle to the surface. Hairs absent.

Habitat: Leaves of *Acer* sp., *Quercus* sp. and *Fagus silvatica* (fide Arendholz, 1979).

Etymology of the specific epithet: refers to the color of the apothecium in its fresh condition.

Holotype: U.S.A., Mass., swamp at east side of Great Blue Hill, near Milton, on decaying leaves of *Acer* sp., 11 Nov 1941, D. H. Linder, E. V. Seeler & W. L. White s.n. (FH).

Paratype: U.S.A., Mass., Stony Brook Reservation, West Tuxbury, fallen leaves of *Quercus* sp., 11 Nov 1941, W. L. White s.n. (FH).

Illustrations: White, Farlowia 1: 153, fig. 6. 1943. Arendholz, Morphologisch-taxonomische Untersuchungen an blattbewohenden Ascomyceten aus der Helotiales, pl. 10, figs. 3, 5. 1979.

NOTES. In his description of *Helotium translucens*, White (1943) concluded that this new species was closely related to *Helotium caudatum*, and I am in agreement with this. *Helotium caudatum* is distinguished from *H. translucens* in that the apothecia of *H. translucens* are smaller than those of *H. caudatum* and the ascospores in *H. caudatum* are predominantly inequilateral and frequently hooked apically; in contrast the spores of *H. translucens* are trapezoidal in shape and generally equilateral. The shape of the ascospores in *H. translucens* is suggestive of those of *H. immutabile*. The structure of the apothecia in these two species is fundamentally different. See discussion under *H. immutabile* for a comparison of apothecial structure.

The species was not treated by Dennis (1956, 1964), but I accept the placement of *H. translucens* in *Hymenoscyphus* by Arendholz (1979).

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MYCOTAXON

Vol. XIII, No. 1, pp. 85-104

April-June 1981

LICHENES EXSICCATI
DISTRIBUTED BY THE UNIVERSITY OF COLORADO
MUSEUM, BOULDER
FASCICLES 1-15, NOS. 1-600, 1961-1979

WILLIAM A. WEBER

Univ. of Colorado Museum,
Campus Box 218, Boulder, CO 80309

SUMMARY

Revision of identifications and nomenclature for the first 600 numbers of LICH. EXSICC. COLO is provided, and the following new combinations are proposed: Cladina galapagosensis, C. polia, Heterodermia barbifera, H. circinalis, H. stellata and H. verrucifera. Validation is provided for the publication of three new species distributed in the exsiccati: Lecanora pseudopinguis, L. texana and Psora cerebriformis.

Lichenes Exsiccati, distributed by the University of Colorado Museum (standard abbreviation LICH. EXS. COLO) began in 1961 with the distribution of Fascicle 1 (1-40). The latest fascicle was Fascicle 15 (561-600), distributed in 1979. Some of the early fascicles were distributed in 100 sets, but the current ones are reduced to 60 sets. The exsiccati are offered in exchange to most of the active lichen herbaria of the world. A brief notice was published in *Taxon* 13:31. 1964, wherein the institutions receiving the complete series were listed. The present paper provides an alphabetical arrangement of the numbers thus far distributed, with updated nomenclature and corrections of identifications up to the present moment. Additional corrections will be welcomed.

In the following list, the first name given is that under which the number was distributed. When that name is enclosed by brackets, this indicates that the name has been changed, and the current name is preceded by an (=) sign. If the identification has been changed, no (=) sign precedes the second name.

TAXON	PROVENANCE	NO.
<u>Acarospora badiofusca</u> (Nyl.) Th. Fr.	Colorado	1
[<u>A. flava</u> (Bell.) Trev.] = <u>A. chlorophana</u> (Wg.) Ach.	Colorado	2
<u>A. nitida</u> H. Magn.	Colorado	522
<u>A. schleicheri</u> (Ach.) Mass.	Arizona	416
[<u>Alectoria chalybeiformis</u> (L.) S. Gray] = <u>Bryoria chalybeiformis</u> (L.) Brodo & Hawksw.	Norway	466
[<u>A. fremontii</u> Tuck.] <u>Bryoria tortuosa</u> (Merr.) Brodo & Hawksw.	California	418
[<u>A. glabra</u> Mot.] <u>Bryoria lanestris</u> (L.) Brodo & Hawksw.	Wyoming	431
<u>A. lata</u> Tayl.	California	417
<u>A. nigricans</u> (Ach.) Nyl.	Tasmania	241
<u>A. ochroleuca</u> (Hoffm.) Mass.	Canada	220
<u>A. poeltii</u> Bystrek	N. Guinea	346
[<u>A. pubescens</u> (L.) R. H. Howe] = <u>Pseudephebe pubescens</u> (L.) Choisy	Canada	219
	Norway	468
<u>A. sarmentosa</u> (Ach.) Ach.	California	132
<u>A. smithii</u> DuRoi	N. Guinea	345
[<u>A. tenuis</u> E. Dahl] <u>Bryoria chalybeiformis</u> (L.) Brodo & Hawksw.	Colorado	125
[<u>Anaptychia barbifera</u> (Nyl.) Trevis.] = <u>Heterodermia barbifera</u> comb. nov.	N. Guinea	379
[<u>A. circinalis</u> (A. Zahlbr.) W. A. Web.] = <u>Heterodermia circinalis</u> comb. nov.	Galapagos	504
[<u>A. diademata</u> (Tayl.) Kurok.] = <u>Heterodermia diademata</u> (Tayl.) Awasthi	Arizona	49
	Mexico	47
	Mexico	77
<u>A. erinacea</u> (Ach.) Trevis.		
[<u>A. leucomela</u> (L.) Mass. f. <u>verrucifera</u> Kurok.] = <u>Heterodermia verrucifera</u> comb. nov.	Galapagos	509
[<u>A. multiciliata</u> Kurok.] = <u>Heterodermia multiciliata</u> Follm. & Redon	Chile	489
[<u>Anaptychia obscurata</u> (Nyl.) Vain.] = <u>Heterodermia obscurata</u> (Nyl.) Trevis.	Australia	257
[<u>A. stellata</u> (Vain.) Kurok.] = <u>Heterodermia stellata</u> (Vain.) W. A. Web., comb. nov.	Galapagos	508
<u>Anthracotheceum ochraceoflavum</u> (Nyl.) Muell.-Arg.	Galapagos	145
<u>Anzia angustata</u> (Pers.) Muell.-Arg.	Australia	281
<u>A. gregoriana</u> Muell.-Arg.	N. Guinea	377
<u>A. hypoleuca</u> Muell.-Arg.	N. Guinea	378
<u>A. wilsonii</u> Raes.	Australia	311
<u>Arthonia glebosa</u> Tuck.	Colorado	53

<u>A. impolita</u> (Ehrh.) Borr. <u>ex</u> Hook. & Sowerby	California	192
	California	477
<u>A. platyspilea</u> Nyl.	Galapagos	122
<u>A. rubella</u> (Fee) Nyl.	Texas	442
<u>Arthopyrenia halodytes</u> (Nyl.) Arn.	Canada	515
[<u>Arthothelium spilomatoides</u> (Nyl.) A. Zahlbr.]		
<u>A. galapagoense</u> Huneck & Follmann TYPE COLL.	Galapagos	113
<u>Aspicilia alpina</u> (Sommerf.) Arn.	Montana	524
<u>A. desertorum</u> (Kremp.) Mereschk.	Colorado	586
[<u>A. hispida</u> Mereschk.] <u>A. desertorum</u> (Kremp.) Mereschk.	Colorado	144
[<u>A. mutabilis</u> (Ach.) Koerb.] = <u>Pachyospora mutabilis</u> (Ach.) Mass.	N. Mexico	157
<u>A. quartzitica</u> W. A. Web. TYPE COLL.	Colorado	364
<u>A. transbaicalica</u> Oxner	USSR	595
<u>A. sp. indet.</u> (fruticose modification)	California	178
<u>Bacidia albescens</u> (Arn.) Zwackh.	California	182
<u>B. herrei</u> A. Zahlbr.	Oregon	419
<u>B. millegrana</u> (Tayl.) A. Zahlbr.	Galapagos	121
<u>Baeomyces absolutus</u> Hepp in Zollinger	N. Guinea	332
	N. Guinea	334
	N. Guinea	335
<u>B. trachypus</u> Nyl.	N. Guinea	333
<u>B. weberi</u> J. W. Thomson TYPE COLL.	N. Guinea	333
<u>Belonia americana</u> Fink <u>ex</u> Hedrick	Texas	355
<u>Bryoria tortuosa</u> (Merr.) Brodo & D. Hawksw.	Canada	579
<u>Buellia curtisii</u> (Tuck.) W. A. Web.	Louisiana	541
<u>B. flavoareolata</u> (Nyl.) Muell.-Arg.	Chile	546
<u>B. galapagona</u> W. A. Web. TYPE COLL.	Galapagos	344
<u>B. glaziouana</u> (Kremp.) Muell.-Arg.	Galapagos	110
<u>B. novomexicana</u> B. de Lesd.	Arizona	408
<u>B. oidalea</u> (Tuck.) Tuck.	California	88
<u>B. punctata</u> (Hoffm.) Mass.	Colorado	208
[<u>B. spuria</u> (Schaer.) Anzi] Incorrect, possibly an undescribed taxon.	Mexico	98
<u>B. cf. taltalensis</u> Dodge	Chile	548
<u>B. triphragmioides</u> Anzi	Colorado	587
<u>B. sp. indet.</u>	Peru	581
<u>B. sp. indet.</u>	Peru	582
<u>B. zahlbruckneri</u> Steiner	Arizona	100
<u>Calicium abietinum</u> Pers.	Idaho	215
[<u>C. hemisphaericum</u> Howard] = <u>C. adequatum</u> Nyl.	Montana	222
<u>Caloplaca amabilis</u> A. Zahlbr.	Arizona	413
<u>C. aurantiaca</u> (Lightf.) Th. Fr.	Arizona	403
[<u>C. bracteata</u> (Hoffm.) Mass.] <u>Fulgensia desertorum</u> (Tomin) Poelt, cited specimen!	Colorado	134
[<u>C. chrysothalma</u> Degel.] Determination doubtful.	N. Mexico	161
<u>C. cinnamomea</u> (Th. Fr.) Oliv.	Colorado	197

<u>C. cladodes</u> (Tuck.) A. Zahlbr. <u>in Engler</u>	Colorado	160
<u>C. epithallina</u> Lynge	Colorado	573
<u>C. fernandeziana</u> (A. Zahlbr.) Follm. & Redon	Chile	547
<u>C. lamprocheila</u> (DC.) Flagey	Colorado	64
<u>C. microphyllina</u> (Tuck.) Hasse	Arizona	412
<u>C. modesta</u> (A. Zahlbr.) Fink	Arizona	414
<u>C. pollinii</u> (Mass.) Jatta	Minnesota	535
<u>C. stanfordensis</u> H. Magn.	California	193
<u>C. subnitida</u> (Malme) A. Zahlbr.	Texas	446
<u>C. trachyphylla</u> (Tuck.) A. Zahlbr.	Colorado	282
[<u>Caloplacopsis submexicana</u> (B. de Lesd.) B. de Lesd.]		
= <u>Candelina submexicana</u> (B. de Lesd.) Poelt	Arizona	405
<u>Candelariella deflexa</u> (Nyl.) A. Zahlbr.	Colorado	398
<u>C. rosulans</u> (Muell.-Arg.) A. Zahlbr.	Colorado	202
	Colorado	283
<u>C. spraguei</u> (Tuck.) A. Zahlbr.	Colorado	3
<u>Catillaria griffithii</u> (Sm.) Malme	California	179
<u>C. sp. indet.</u>	Chile	553
<u>Cavernularia lophyrea</u> (Ach.) Degel.	Canada	517
<u>Cetraria canadensis</u> (Raes.) Raes.	Idaho	223
<u>C. cucullata</u> (Bell.) Ach.	Alaska	42
<u>C. delisei</u> (Bory ex Schaer.) Th. Fr.	Norway	467
<u>C. ericetorum</u> Opiz	Colorado	217
<u>C. fendleri</u> (Nyl.) Tuck.	Arizona	411
[<u>C. fendleri f. coralligera</u> W. A. Web. TYPE COLL.]		
= <u>C. coralligera</u> (W. A. Web.) Hale	Arizona	99
[<u>C. glauca</u> (L.) Ach.] = <u>Platismafia glauca</u> (L.)		
W. Culb. & C. Culb.	Colorado	4
<u>C. idahoensis</u> Essl.	Idaho	400
<u>C. islandica</u> (L.) Ach.	Colorado	46
	Australia	274
	Colorado	203
<u>C. pinastri</u> (Scop.) S. Gray		
[<u>C. richardsonii</u> Hook. <u>in Richards.</u>] = <u>Masonhalea richardsonii</u> (Hook. <u>in Richards.</u>) Kaernefelt	Alaska	54
<u>C. scutata</u> (Wulf.) Poetsch	Tasmania	423
<u>C. tilesii</u> Ach.	Alaska	212
<u>C. sp. indet.</u>	N. Guinea	483
<u>Cetrelia chicitae</u> (W. Culb.) W. Culb. & C. Culb.	N. Guinea	315
<u>Chaenotheca brunneola</u> (Ach.) Muell.-Arg.	Idaho	214
<u>Chiodecton effusum</u> Fee	Galapagos	503
<u>C. farinaceum</u> Fee	Galapagos	498
<u>C. sp. indet.</u>	Chile	551
<u>C. sp. indet.</u>	Chile	557
<u>Chondropsis semiviridis</u> (F. Muell. <u>ex Nyl.</u>) Nyl. <u>ex</u>		
Cromb.	Australia	225
	Australia	570

<u>Cladia aggregata</u> (Sw.) Nyl.	Nepal	148
	Australia	238
	Galapagos	497
<u>C. ferdinandii</u> (Muell.-Arg.) R. Filson	Australia	571
<u>C. retipora</u> (Labill.) Nyl.	Australia	254
[<u>C. retipora</u> (Labill.) Nyl.] <u>C. corallaizon</u> R. Filson	Australia	237
	Australia	255
<u>C. schizopora</u> (Nyl.) Nyl.	Australia	266
<u>C. sullivanii</u> (Muell.-Arg.) W. Martin	Tasmania	227
[<u>Cladonia alpestris</u> (L.) Rabenh.] = <u>Cladina stellaris</u> (Opiz) Pouzar & Vezda	Alaska	44
[<u>Cladonia cariosa</u> (Ach.) Spreng.] <u>C. robbinsii</u> Evans	Kansas	63
<u>C. ceratophylla</u> (Sw.) Spreng.	Galapagos	502
<u>C. chlorophaea</u> (Flk.) Spreng.	Galapagos	139
<u>C. coccifera</u> (L.) Willd.	N. Guinea	314
<u>C. corymbescens</u> Nyl.	N. Guinea	312
<u>C. ecmocyna</u> Leighton	Colorado	531
<u>C. foliacea</u> (Huds.) Willd.	Denmark	593
[<u>C. galapagosensis</u> Ahti] = <u>Cladina galapagosensis</u> comb. nov.	Galapagos	105
<u>C. leporina</u> Fr.	Alabama	285
<u>C. multiformis</u> Merrill	Colorado	5
	N. Mexico	433
<u>C. pertriosa</u> Kremp.	Australia	267
<u>C. cf. pityrea</u> (Flk.) Fr.	Australia	360
[<u>C. polia</u> R. Sant.] = <u>Cladina polia</u> (R. Sant.) comb. nov.	Galapagos	106
<u>C. scabriuscula</u> (Del. in Duby) Nyl.	Australia	359
	Chile	534
	N. Guinea	342
<u>C. solitaria</u> H. Magn.		
[<u>C. squamosa</u> (Scop.) Hoffm.] <u>C. dilleniana</u> Flk. fide J. W. Thomson	N. Guinea	313
[<u>C. subtenuis</u> (des Abb.) Evans] = <u>Cladina subtenuis</u> (des Abb.) Hale & W. Oulb.	Kansas	52
<u>C. sulphurina</u> (Michx.) Fr.	Montana	527
<u>C. verticillata</u> (Hoffm.) Schaer.	Australia	250
<u>C. vulcanica</u> Zoll. in Hasskarl	N. Guinea	320
[<u>C. xanthoclada</u> Muell.-Arg.] <u>C. capitellata</u> (Tayl.) Bab.	Australia	258
<u>C. zopfii</u> Vain.	Denmark	592
<u>Coelocaulon aculeatum</u> (Schreb.) Gyel.	Colorado	574
	Canada	166
[<u>C. australiense</u> nom. nud.] <u>Cetraria australiense</u> Kaernefelt TYPE COLL.	Australia	454
<u>Coenogonium implexum</u> Nyl.	Tasmania	306
<u>Collema cristatum</u> (L.) G. H. Web. in Wiggers	Colorado	340

<u>C. durietzii</u> Degel.	Australia	438
<u>C. leucocarpum</u> Hook. f. & Tayl.	Australia	253
<u>C. polycarpum</u> Hoffm.	Colorado	392
<u>Conotrema urceolatum</u> (Ach.) Tuck.	W. Virginia	464
[<u>Cora pavonia</u> (Sw.) Fr.] = <u>Dictyonema montanum</u> (Sw.) Mamm. ex Follm.	Galapagos	107
[<u>Cornicularia aculeata</u> (Schreb.) Ach.] = <u>Coelocaulon</u> <u>aculeatum</u> (Schreb.) Gyel.	Colorado	6
[<u>C. californica</u> (Tuck.) DuRietz] = <u>Coelocaulon cali-</u> <u>fornicum</u> (Tuck.) Howe f.	Oregon	130
<u>C. normoerica</u> (Gunn.) DuRietz	Oregon	128
[<u>Cypheliopsis bolanderi</u> (Tuck.) Vain.] = <u>Thelomma</u> <u>mamosum</u> (Hepp in Hartung) Mass. The collection is mixed, containing also <u>T. santessonii</u> L. Tibell	Mexico	75
<u>Cyphelium inquinans</u> (Sm.) Trev.	Colorado	204
<u>Cystocoleus ebeneus</u> (Dillwyn) Thwaites	Colorado	491
<u>Dactylina madreporiformis</u> (Ach.) Tuck.	Colorado	7
	Colorado	210
<u>Darbishirella gracillima</u> (Kremp.) A. Zahlbr.	Chile	554
<u>Dendrographa leucophaea</u> (Tuck.) Darb.	Mexico	83
	Mexico	493
<u>D. minor</u> Darb.	Calif.	177
<u>Dermatocarpon leptophyllum</u> (Ach.) Vain.	USSR	598
<u>D. miniatum</u> (L.) Mann	Arizona	8
<u>D. plumbeum</u> B. de Lesd.	Colorado	391
<u>D. reticulatum</u> H. Magn.	Oregon	209
<u>D. rivulorum</u> (Arn.) Dalla Torre & Sarnth.	Colorado	394
<u>D. tuckermanii</u> (Rav. ex Mont.) A. Zahlbr.	Arizona	95
<u>D. vellereum</u> Zschacke	USSR	596
<u>Dictyonema irpicinum</u> Mont.	N. Guinea	361
<u>D. sericeum</u> (Sw.) Berk.	N. Guinea	362
<u>Diploicea canescens</u> (Dicks.) Mass.	California	184
<u>Diploschistes ocellatus</u> (Vill.) Norm.	Mexico	578
[<u>D. scruposus</u> (Schreb.) Norm.] <u>D. cf. canadensis</u> Raes.	Colorado	9
<u>Dirina catalinariae</u> Hasse	Mexico	76
<u>D. herrei</u> A. Zahlbr.	Galapagos	367
<u>D. limitata</u> Nyl.	Chile	549
<u>Dolichocarpus chilensis</u> R. Sant.	Chile	560
<u>Endocarpon pulvinatum</u> Th. Fr.	Colorado	291
<u>E. pusillum</u> Hedw.	Colorado	10
	Colorado	429
<u>Enterographa atacamensis</u> Dodge	Peru	583
[<u>Enterographa atacamensis</u> Dodge] <u>Roccellina olivacea</u> Follm.	Chile	555
<u>Ephebe lanata</u> (L.) Vain.	Colorado	59
<u>Evernia prunastri</u> (L.) Ach.	California	187

<u>Everniastrum neocirrhatum</u> (Hale & V. Wirth) Hale	Mexico	487
<u>E. vexans</u> Hale	Galapagos	499
<u>Glypholecia scabra</u> (Pers.) Muell.-Arg.	USSR	600
<u>Graphis caesiella</u> Vain.	Galapagos	480
<u>G. striatula</u> (Ach.) Spreng.	Galapagos	142
<u>Gyrostomum scyphuliferum</u> (Ach.) Nyl.	Louisiana	542
<u>Haematomma babingtonii</u> Mass.	New Zealand	296
<u>H. subpuniceum</u> (Muell.-Arg.) B. de Lesd.	Texas	11
<u>Heppia lutosa</u> (Ach.) Nyl.	Iowa	297
<u>Heterodea muelleri</u> (Hampe) Nyl.	Australia	172
<u>Heterodermia leucomelos</u> (L.) Poelt	Mexico	577
[<u>Hubbsia lumbricoides</u> W. A. Web. TYPE COLL.]		
= <u>Reinkella californica</u> Raes.	Mexico	85
<u>Hydrothyria venosa</u> Russell	California	293
<u>Hypogymnia billardieri</u> (Mont.) R. Filson	Australia	382
<u>H. krogii</u> Ohlsson	N. Carolina	529
<u>H. lugubris</u> (Pers.) Krog	Australia	351
<u>H. mundata</u> (Nyl.) Rassadina	Australia	385
<u>H. oroarctica</u> Krog	Colorado	572
<u>H. physodes</u> (L.) Nyl.	Colorado	133
<u>Icmadophila ericetorum</u> (L.) A. Zahlbr.	Colorado	164
<u>Ingaderia pulcherrima</u> Darb.	Chile	558
<u>Koerberia biformis</u> Mass.	Arizona	12
<u>Lasallia pensylvanica</u> (Hoffm.) Llano	Colorado	13
<u>L. pustulata</u> (L.) Mereschk. <u>ssp. papulosa</u> (Ach.)		
W. A. Web.	Texas	14
	Colorado	15
<u>Lecanactis californica</u> Tuck.	California	194
<u>L. cf. myriadea</u> (Fee) A. Zahlbr.	Chile	550
<u>L. (subg. Bactrospora) sp. indet.</u>	California	456
[<u>Lecanora arizonica</u> (Tuck.) W. A. Web.] = <u>Omphalo-</u>		
<u>dium arizonicum</u> (Tuck.) Tuck.	N. Mexico	16
<u>L. badia</u> (Hoffm.) Ach.	Montana	523
[<u>L. bolanderi</u> Tuck.] <u>L. phryganitis</u> Tuck.	California	165
[<u>L. cancriformis</u> (Hoffm.) Vain.] <u>L. caesiorubella</u>		
Ach. <u>ssp. merrillii</u> Imsh. & Brodo	California	181
<u>L. carpinea</u> (L.) Vain.	California	183
<u>L. chlarotera</u> Nyl.	Colorado	200
<u>L. christoi</u> W. A. Web. TYPE COLL.	Colorado	458
<u>L. confusa</u> Almborn	Oregon	420
[<u>L. conizaea</u> (Ach.) Nyl.] Incorrect, but no alter-		
native identification available.	Galapagos	138
[<u>L. conizaeoides</u> Nyl. ex Cromb.] <u>L. confusa</u> Alb.	California	180
<u>L. coquimbensis</u> A. Zahlbr.	Chile	552
<u>L. frustulosa</u> (Dicks.) Ach.	Colorado	17
<u>L. garovaglii</u> (Koerb.) A. Zahlbr. (Disregard com-		
ments)	Colorado	118

[<u>L. cf. glabrata</u> (Ach.) Malme] Probably incorrect but no alternative name available.	Texas	436
[<u>L. lentigera</u> (G. Web.) Ach.] = <u>Squamarina lentigera</u> (G. Web.) Poelt	Colorado	18
[<u>L. marginalis</u> Hasse] = <u>Rhizoplaca marginalis</u> (Hasse) W. A. Web.	California	126
[<u>L. melanophthalma</u> Ram.] = <u>Rhizoplaca melanophthalma</u> (Ram.) Leuckert & Poelt	Colorado	368
	Wyoming	430
<u>L. mellea</u> W. A. Web. TYPE COLL.	California	462
[<u>L. peltata</u> (Ram.) Steud.] = <u>Rhizoplaca peltata</u> (Ram.) Leuckert & Poelt	Colorado	119
	Colorado	460
	California	475
<u>L. phryganitis</u> Tuck.	Mexico	87
<u>L. pinguis</u> Tuck.	Colorado	521
<u>L. polytropa</u> (Ehrh.) Rabenh.	Colorado	135
<u>L. pringlei</u> M. Lamb	Galapagos	500
<u>L. pseudopinguis</u> W. A. Web. sp. nov. TYPE COLL.		
[<u>L. rubina</u> (Vill.) Ach.] = <u>Rhizoplaca chrysoleuca</u> (Sm.) Choisy	Colorado	19
<u>L. saligna</u> (Schrad.) A. Zahlbr.	Colorado	395
<u>L. texana</u> W. A. Web., sp. nov. TYPE COLL.	Texas	451
[<u>L. thomsonii</u> H. Magn.] = <u>L. novomexicana</u> B. de Lesd. ex H. Magn.	Colorado	20
<u>L., stirps varia</u> (Ehrh.) Ach.	Colorado	563
<u>Lecidea aspidula</u> Kremp.	Australia	273
[<u>L. atrobrunnea</u> (Ram.) Schaer.] <u>L. leucothallina</u> Arn.	Colorado	155
<u>L. berengeriana</u> (Mass.) Th. Fr.	Colorado	21
	Oregon	428
[<u>L. cinnabarina</u> Sommerf.] <u>Protoblastenia russula</u> (Ach.) Raes.	Australia	259
[<u>Lecidea decipiens</u> Ach.] = <u>Psora decipiens</u> Hoffm. (Ignore the incorrect synonymy on the label)	Colorado	22
<u>L. dolodes</u> Nyl. ex Hasse	California	478
<u>L. elabens</u> Fr.	Australia	277
<u>L. elata</u> Schaer.	Colorado	481
[<u>L. friesii</u> Ach. in Liljebl.] = <u>Hypocenomyce friesii</u> (Ach. in Liljebl.) G. Schneider	Australia	272
[<u>L. icterica</u> (Mont.) Tayl.] = <u>Psora icterica</u> (Mont.) Muell.-Arg.	N. Mexico	435
<u>L. insularis</u> Nyl.	Montana	141
<u>L. leucothallina</u> Arn.	Montana	525
<u>L. limosa</u> Ach.	N. Guinea	567
[<u>L. luridella</u> Tuck.] = <u>Psora luridella</u> (Tuck.) Fink	Colorado	482
<u>L. lyngei</u> Degel.	Colorado	424

[<u>L. novomexicana</u> (B. de Lesd.) W. A. Web. ex R. Anderson] = <u>Psora nipponica</u> (A. Zahlbr.) G. Schneider	Idaho	224
<u>L. nylanderi</u> (Anzi) Th. Fr.	Minnesota	536
[<u>L. quadricolor</u> (Dicks.) Borr.] = <u>L. granulosa</u> (Hoffm.) Ach.	Arizona	103
[<u>L. rubiformis</u> (Wahlenb. ex Ach.) Ach.] <u>Psora tuckermanii</u> R. Anderson ex G. Schneider, invalid. no Latin description.	Colorado	23
[<u>L. rubiformis</u> (Wahlenb. ex Ach.) Ach.] <u>Psora cerebriformis</u> W. A. Web., sp. nov.	Colorado	24
[<u>L. rufonigra</u> (Tuck.) Nyl.] = <u>Psorula rufonigra</u> (Tuck.) G. Schneider	Colorado Texas	124 450
[<u>L. symmicta</u> (Ach.) Ach.] <u>Lecanora cadubriae</u> (Mass.) Hedlund	Colorado	81
[<u>L. texana</u> W. A. Web.] = <u>Xanthopsora texana</u> (W. A. Web.) G. Schneider & W. A. Web. TYPE OOLL.	Texas	448
[<u>L. wallrothii</u> Flk. ex Spreng.] = <u>Trapeliopsis wallrothii</u> (Flk. ex Spreng.) H. Hertel & G. Schneider	California	127
<u>Lecidella elaeochroma</u> (Ach.) Haszl.	Colorado	520
[<u>Leprocaulon albicans</u> (Th. Fr.) Nyl. ex Hue] = <u>Leprocaulon gracilescens</u> M. Lamb & Ward	Arizona Colorado	25 26
<u>L. arbuscula</u> (Nyl.) Nyl. ex Hue	N. Guinea	290
[<u>L. microscopicum</u> (Vill.) Gams ex D. Hawksw.] <u>L. tenellum</u> (Tuck.) Nyl.	Galapagos	496
<u>Leprocaulon tenellum</u> (Tuck.) Nyl.	Peru	584
<u>Leptogium brebissonii</u> Mont. in Webb	Australia	317
<u>L. burnetiae</u> Dodge var. <u>hirsutum</u> (Sierk) P. M. Joerg.	Colorado	486
<u>L. cyanescens</u> (Ach.) Koerb.	Galapagos	506
<u>L. denticulatum</u> Nyl.	Colorado	78
[<u>L. foveolatum</u> Nyl.] <u>L. punctulatum</u> Nyl.	Mexico Galapagos	56 505
[<u>L. furfuraceum</u> (Harm.) Sierk] <u>L. papillosum</u> (B. de Lesd.) Dodge	Arizona	96
<u>L. hypotrachynum</u> Muell-Arg.	Australia	388
<u>L. javanicum</u> (Mont. & v. d. Bosch) Mont.	N. Guinea	295
<u>L. lichenoides</u> (L.) A. Zahlbr.	Colorado Oregon	50 421
<u>L. millegranum</u> Sierk	Arizona	97
<u>L. moluccanum</u> (Pers. in Gaud.) Vain.	Arizona	159
<u>L. phyllocarpum</u> (Pers.) Mont.	N. Guinea	381
<u>L. rugosum</u> Sierk	Mexico	55
<u>Letharia vulpina</u> (L.) Hue	Arizona	102
<u>L. vulpina</u> (L.) Hue f. <u>californica</u> (Lev.) W. A. Web.	California	512
<u>Lichina confinis</u> (O. F. Muell.) C. A. Agardh	Nevada	120
	Tasmania	234

<u>L. tasmanica</u> Henssen	Tasmania	393
<u>Lobaria hallii</u> Tuck.	Oregon	401
<u>L. retigera</u> (Bory) Trevis.	India	149
[<u>L. stictaeformis</u> (Schaer.) Trevis.] <u>L. discolor</u> (Bory in Del.) Hue	N. Guinea	347
<u>Lopadium pezizoideum</u> (Ach.) Koerb.	Idaho	221
<u>Maronella laricina</u> Steiner ISOTYPES	Austria	588
<u>Mastodia tessellata</u> auct. (See Brodo, The Bryologist 79:396-398, 1977, for nomenclatural discussion)	Alaska	27
<u>Melanaria melanospora</u> (Nyl.) Erichsen	Chile	545
[<u>Menegazzia aeneofusca</u> (Muell.-Arg.) R. Sant.] <u>M. nothofagi</u> P. James ined.	Australia	232
<u>M. pertusa</u> (Schrank) Stein	N. Guinea	384
<u>Micarea denigrata</u> Fr.) Hedlund	Colorado	73
<u>Microthelia micula</u> Koerb.	Arizona	67
<u>Mycoglaena myrica</u> (Nyl.) R. C. Harris	Colorado	519
<u>Neophyllis melacarpa</u> (F. Wils.) F. Wils. G. Schneider maintains this over Yoshimura's placement in <u>Gymnoderma</u>	Australia	246
<u>Nephroma arcticum</u> (L.) Torss.	Alaska	43
<u>N. australe</u> A. Richard	Tasmania	569
<u>N. celluloseum</u> (Sm. ex Ach.) Ach.	Australia	239
<u>N. helveticum</u> Ach.	Colorado	28
<u>N. resupinatum</u> (L.) Ach. [<u>Nephromopsis californica</u> Gyel.] = <u>Cetraria orbata</u> (Nyl.) Fink	Alaska	170
<u>Neuropogon acromelanus</u> (Stirt.) M. Lamb	California	129
<u>N. sulphureus</u> (Koenig) Hellb.	Tasmania	247
<u>Normandina pulchella</u> (Borr.) Nyl.	Spitzbergen	591
<u>Ocellularia alba</u> (Fee) Muell.-Arg.	Australia	276
<u>Ochrolechia grimmiae</u> Lynge	Australia	352
<u>O. upsaliensis</u> (L.) Mass.	Spitzbergen	590
<u>Omphalaria kansana</u> Tuck. Determination doubtful.	Colorado	57
<u>Opegrapha saxicola</u> Ach.	Colorado	173
<u>Pannaria leucophaea</u> (Vahl) P. M. Joerg.	California	198
[<u>P. nigrata</u> Muell.-Arg.] Possibly <u>P. tavaresii</u> P. M. Joerg., cf. Opera Bot. 45:70. 1978.	USSR	599
<u>P. pezizoides</u> (G. Web.) Trevis.	Australia	386
<u>Parathelium</u> sp. indet.	Colorado	396
[<u>Parmelia bostrychodes</u> A. Zahlbr.] = <u>Hypotrachyna</u> <u>bostrychodes</u> (A. Zahlbr.) Hale	Chile	556
[<u>P. caperata</u> (L.) Ach.] = <u>Pseudoparmelia caperata</u> (L.) Hale	N. Guinea	376
[<u>P. chiricahuensis</u> R. Anderson & W. A. Web.] = <u>Neofuscellia chiricahuensis</u> (R. Anderson & W. A. Web.) Essl.	Australia	270
	Arizona	91
	Texas	444

[<u>P. conspersa</u> (Ach.) Ach.] <u>Xanthoparmelia chlorochroa</u> (Tuck.) Hale	Wyoming	29
[<u>P. dichotoma</u> Muell.-Arg.] = <u>Xanthoparmelia dichotoma</u> (Muell.-Arg.) Hale	Australia	322
[<u>P. dominicana</u> Vain.] = <u>Parmotrema dominicana</u> (Vain.) Hale	Galapagos	136
[<u>P. elegantula</u> (A. Zahlbr.) Szatala] = <u>Melanelia elegantula</u> (A. Zahlbr.) Essl.	Colorado	207
[<u>P. endosulphurea</u> (Hillm.) Hale] = <u>Parmotrema endosulphureum</u> (Hillm.) Hale	Dominica	336
[<u>P. exasperatula</u> Nyl.] = <u>Melanelia exasperatula</u> (Nyl.) Essl.	Colorado	154
[<u>P. furfuracea</u> (L.) Ach.] <u>Pseudevernia intensa</u> (Nyl.) Hale & W. Culb.	Texas	30
[<u>P. galbina</u> Ach.] = <u>Parmelina galbina</u> (Ach.) Hale	Australia	244
[<u>P. isidiotyla</u> Nyl.] <u>Neofuscelia loxodes</u> (Nyl.) Essl.	Colorado	153
[<u>P. novomexicana</u> Gyel.] = <u>Xanthoparmelia novomexicana</u> (Gyel.) Hale	Arizona	404
[<u>P. perlata</u> (Huds.) Ach.] = <u>Parmotrema perlatum</u> (Huds.) Hale	Australia	279
[<u>P. physodes</u> (L.) Ach.] = <u>Hypogymnia physodes</u> (L.) Nyl.	Colorado	68
<u>P. praesignis</u> Nyl.	N. Mexico	156
<u>P. pseudotenuirima</u> Gyel.	Australia	365
[<u>P. pulla</u> (Schreb.) Ach.] = <u>Neofuscelia pulla</u> (Ach.) Essl.	Australia	271
[<u>P. pulvinata</u> Fee] = <u>Hypotrachyna pulvinata</u> (Fee) Hale	Arizona	101
[<u>P. revoluta</u> Flk.] = <u>Hypotrachyna revoluta</u> (Flk.) Hale	Colorado	463
[<u>P. rutidota</u> Hook. f. & Tayl.] = <u>Pseudoparmelia rutidota</u> (Hook. f. & Tayl.) Hale	Australia	268
<u>P. saxatilis</u> (L.) Ach.	Colorado	538
[<u>P. saximontana</u> R. Anderson & W. A. Web.] TYPE COLL. = <u>Melanelia substygia</u> (R. Anderson & W. A. Web.) Essl.	Colorado	41
<u>P. signifera</u> Nyl.	Australia	251
<u>P. subalbicans</u> Stirt.	Australia	280
[<u>P. subcristata</u> Nyl.] = <u>Parmotrema subtinctorium</u> (Nyl.) Hale	Dominica	339
<u>P. subrudecta</u> Nyl.	Australia	349
[<u>P. substrigosa</u> Hale in W. A. Web.] = <u>Xanthoparmelia substrigosa</u> (Hale in W. A. Web.) Hale	Australia	338
[<u>P. subtinctoria</u> A. Zahlbr.] = <u>Parmotrema subtinctorium</u> (A. Zahlbr.) Hale	Australia	278
	Arizona	410

[<u>P. taractica</u> Kremp.] = <u>Xanthoparmelia taractica</u> (Kremp.) Hale	Arizona	409
[<u>P. tinctorum</u> Nyl.] = <u>Parmotrema tinctorum</u> (Nyl.) Hale	Galapagos	114
[<u>P. ulcerosa</u> A. Zahlbr.] = <u>Xanthoparmelia ulcerosa</u> (A. Zahlbr.) Hale	Venezuela	469
[<u>P. ulophyllodes</u> (Vain.) Sav.] = <u>P. soledica</u> Nyl.	Colorado	213
[<u>P. weberi</u> Hale] = <u>Xanthoparmelia weberi</u> (Hale) Hale	Arizona	415
[<u>P. wyomingica</u> (Gyel.) Hale] = <u>Xanthoparmelia wyo-</u> <u>mingica</u> (Gyel.) Hale	Wyoming	465
<u>Parmeliopsis aleurites</u> (Ach.) Nyl.	Arizona	406
<u>P. placorodia</u> (Ach.) Nyl.	Colorado	31
	Arizona	65
<u>Peccania</u> sp. indet.	Mexico	576
<u>Peltigera aphthosa</u> (L.) Willd.	Colorado	32
<u>Peltigera dolichorhiza</u> (Nyl.) Nyl.	Australia	245
	N. Guinea	321
<u>P. horizontalis</u> (Huds.) Baumg.	Colorado	71
	N. Guinea	319
<u>P. horizontalis</u> (Huds.) Baumg. <u>f. zopfii</u> (Gyel.) J. W. Thomson	Colorado	485
<u>P. polydactyla</u> (Neck.) Hoffm.	Alaska	171
	Oregon	422
[<u>P. scabrosa</u> Th. Fr.] <u>P. sp. nov. Vitikainen ined.</u>	Colorado	484
<u>P. venosa</u> (L.) Willd.	Colorado	33
<u>Peltula cylindrica</u> Wetmore	Georgia	402
<u>P. obscurans</u> (Nyl.) Gyel. <u>var. deserticola</u> (A. Zahlbr.) Wetmore	Arizona	397
<u>Pertusaria californica</u> Dibben	California	473
<u>P. saximontana</u> Wetmore	Colorado	459
<u>P. xanthodes</u> Muell.-Arg.	Louisiana	539
<u>Pertusaria</u> (Subg. <u>Lecanorastrum</u>) sp. indet.	Tasmania	479
[<u>Phaeographina tridactna</u> W. A. Web. sp. nov. ined.] <u>P. isidiosa</u> (Vain.) A. Zahlbr., det. Nakanishi.	N. Guinea	390
<u>Phaeographis exaltata</u> (Mont. & v. d. Bosch) Muell.- Arg.	Mexico	74
<u>P. sp. indet.</u>	Galapagos	514
<u>Phylliscum demageonii</u> (Moug.) Nyl.	Colorado	45
<u>Physcia adscendens</u> (Th. Fr.) Oliv. em. Bitt.	Colorado	152
[<u>P. aegialita</u> (Ach.) Nyl.] <u>Dirinaria subconfluens</u> Awasthi, cited specimen.	Australia	242
<u>P. caesia</u> (Hoffm.) Hampe Identification incorrect, but no alternative available.	Australia	269
<u>P. callosa</u> Nyl.	Colorado	60
<u>P. duplicorticata</u> W. A. Web. & J. W. Thomson TYPE COLL.	Calif.	476

[<i>P. pulverulenta</i> (Schreb.) Hampe f. <i>coralloidea</i> Suza in Nadv.] <i>Physconia</i> sp. indet.	N. Mexico	158
[<i>P. setosa</i> (Ach.) Nyl.] <i>P. hispidula</i> (Ach.) Frey	Arizona	93
[<i>P. setosa</i> (Ach.) Nyl.] Identification incorrect.	Australia	252
<i>P. stellaris</i> (L.) Nyl.	Colorado	201
<i>P. tribacoides</i> Nyl.	Louisiana	437
<i>P. vitii</i> Nadv.	Germany	589
<i>Physconia detersa</i> (Nyl.) Poelt	Colorado	399
<i>P. muscigena</i> (Ach.) Poelt	Colorado	341
<i>Physma byrsinum</i> (Ach.) Muell.-Arg.	N. Guinea	363
<i>Pilophoron robustum</i> Th. Fr.	Alaska	34
<i>Placynthium nigrum</i> (Huds.) S. Gray	Ohio	457
<i>Platismatia stenophylla</i> (Tuck.) W. Culb. & C. Culb.	Canada	516
<i>Polychidium muscicola</i> (Sw.) S. Gray	Colorado	58
	Colorado	490
<i>Porina epiphylla</i> Fee	Cocos I.	143
	Australia	370
<i>P. rubentior</i> (Stirt.) Muell.-Arg.	Cocoa I.	143
<i>P. sp. indet.</i>	Chile	561
<i>Pseudocyphellaria argyracea</i> (Del.) Vain.	Australia	307
<i>P. australiensis</i> H. Magn.	Australia	318
[<i>P. durvillei</i> (Del.) Vain.] <i>P. hirta</i> (G. Forst.) D. Gall. & P. James	Tasmania	383
[<i>P. freycinetii</i> (Del.) Malme] Identification dubious.	Australia	389
	Tasmania	387
<i>P. gilva</i> (Ach.) Malme	N. Guinea	373
<i>P. glabra</i> (Hook. f. & Tayl.) Dodge	Tasmania	310
<i>P. impressa</i> (Hook. f. & Tayl.) Vain.	Tasmania	308
<i>P. neglecta</i> (Muell.-Arg.) H. Magn.	Australia	372
[<i>P. orygmæa</i> (Ach.) Malme] <i>P. hirta</i> (G. Forst.) D. Gall. & P. James	Tasmania	264
<i>P. sp. indet.</i>	Australia	565
<i>P. rainierensis</i> Imshaug	Washington	123
<i>Psoroma soccatum</i> R. Br. in Cromb.	Australia	461
<i>P. sphinctrinum</i> (Mont.) Nyl.	Tasmania	256
<i>Psorotichia minuta</i> H. Magn.	Colorado	196
<i>P. sp. indet.</i>	Texas	449
<i>Pyrenotrichum splitgerberi</i> Mont.	Louisiana	286
<i>Pyrenula cerina</i> Eschw.	Galapagos	146
<i>P. nitidella</i> (Flk. ex Schaer.) Muell.-Arg. var. <i>maculata</i> R. C. Harris TYPE COLL.	Texas	443
<i>Pyxine pringlei</i> Imshaug	Galapagos	111
[<i>Ramalina ceruchis</i> (Ach.) DeNot.] = <i>Desmazieria</i> <i>ceruchis</i> (Ach.) Trevis.	Mexico	79
[<i>R. combeoides</i> Nyl.] = <i>Desmazieria combeoides</i> (Nyl.) Follm. & Huneck	Mexico	89
<i>R. complanata</i> (Sw. ex Ach.) Ach.	Galapagos	147

<u>R. farinacea</u> (L.) Ach.	Galapagos	168
	California	186
	Australia	348
<u>R. fraxinea</u> (L.) Ach. Identification doubtful.		
<u>R. geniculata</u> Hook. f. & Tayl. var. <u>compacta</u> Muell.-Arg.	Tasmania	350
[<u>R. homalea</u> Ach.] = <u>Desmazieria homalea</u> (Ach.) Mont.	Mexico	82
	California	185
<u>R. javanica</u> Nyl.	N. Guinea	343
<u>R. linearis</u> (Sw.) Ach.	Galapagos	167
<u>R. menziesii</u> Tayl., non Tuck.	Mexico	511
<u>R. pacifica</u> Asahina	Philippines	533
<u>R. peruviana</u> Ach.	Galapagos	510
<u>R. sinensis</u> Jatta	N. Mexico	434
<u>R. sorediantha</u> Nyl.	Galapagos	137
	Galapagos	169
<u>R. subleptocarpa</u> Rundel & Bowler	Mexico	564
<u>R. usnea</u> (L.) Howe f.	Panama	90
	Galapagos	115
[<u>Ramalodium succulentum</u> (R. Br.) Nyl.] <u>Arctomia</u> <u>fruticosa</u> Henssen & Weber. sp. nov. ined.	Tasmania	452
<u>Reinkella lirellina</u> Darb.	Peru	580
<u>R. parishii</u> Hasse	California	176
<u>Rhizocarpon macrosporum</u> Raes.	Colorado	518
<u>Rinodina applanata</u> H, Magn.	Louisiana	540
[<u>R. archaea</u> (Ach.) Vain. em. Malmé] Sheard questions the identification.	California	199
	California	206
<u>R. calculiformis</u> W. A. Web. TYPE COLL.	Mexico	94
<u>R. coloradiana</u> H. Magn.	Colorado	537
<u>R. marysvillensis</u> H. Magn.	California	474
[<u>R. oreina</u> (Ach.) Mass.] = <u>Dimelaena oreina</u> (Ach.) Norm.	Colorado	35
[<u>R. radiata</u> (Tuck.) Tuck.] = <u>Dimelaena radiata</u> (Tuck.) Hale & W. Culb.	California	189
<u>R. turfacea</u> (Wahlenb.) Koerb.	Colorado	292
<u>Roccella babingtonii</u> Mont.	Mexico	80
	Galapagos	109
<u>R. fimbriata</u> Darb.	Mexico	84
	Mexico	86
<u>R. fucoides</u> (Dicks.) Vain.	Italy	594
<u>R. portentosa</u> Darb.	Chile	492
[<u>R. portentosa</u> Darb.] <u>R. galapagoensis</u> Follm. TYPE COLL.	Galapagos	112
	Galapagos	117
<u>Roccellaria mollis</u> (Hampe) A. Zahlbr.	Chile	559
	Peru	585
<u>Roccellina condensata</u> Darb.	Chile	544

[<i>R. lutosa</i> Follm.] <i>Roccellina luteola</i> Follm.	Chile	543
<i>Sarcogyne clavus</i> (Ram.) Kremp.	Texas	445
<i>S. regularis</i> Koerb.	Texas	441
<i>Schismatomma cupressum</i> Herre	California	195
<i>Schistoporon tenue</i> Stirt.	Galapagos	495
<i>Schizopelte californica</i> Th. Fr.	California	175
<i>Siphula coriacea</i> Tayl. <u>ex</u> Nyl.	Australia	243
	Australia	275
<i>Siphula fragilis</i> (Hook. <u>f.</u> & Tayl.) J. Murray, <u>nom.</u> <u>nud.</u>	Tasmania	265
<i>S. sp. indet.?</i>	California	455
<i>Speerschneidera euploca</i> (Tuck.) Trevis.	Texas	447
<i>Sphaerophorus melanocarpus</i> (Sw.) DC.	Tasmania	289
<i>S. tener</i> Laur.	Tasmania	294
<i>Sporastatia testudinea</i> (Ach.) Mass.	Colorado	36
<i>Staurothele clopima</i> (Wahlenb. <u>ex</u> Ach.) Th. Fr.	California	427
<i>Stereocaulon glareosum</i> (Sav.) H. Magn.	Colorado	66
	Montana	528
<i>S. leprocauloides</i> M. Lamb TYPE COLL.	N. Guinea	299
<i>S. massartianum</i> Hue	N. Guinea	302
	Philippines	532
[<i>S. microcarpum</i> Muell.-Arg.] <i>S. weberi</i> M. Lamb, TYPE COLL.	Galapagos	494
<i>S. myriocarpum</i> Th. Fr.	N. Guinea	304
[<i>Stereocaulon piliferum</i> Th. Fr.] misspelling for <i>S. piluliferum</i> Th. Fr.	India	150
<i>S. pseudomassartianum</i> M. Lamb <u>ex</u> Frey	N. Guinea	300
	N. Guinea	301
	N. Guinea	303
<i>S. ramulosum</i> (Sw.) Rausch <u>var. nudatum</u> (Muell.-Arg.) Muell.-Arg.	Australia	249
<i>S. rivulorum</i> H. Magn.	Colorado	530
<i>S. staufferi</i> M. Lamb <u>ex</u> Frey	N. Guinea	305
<i>S. virgatum</i> Ach. <u>ex</u> Spreng.	Dominica	337
<i>Sticta byschiana</i> Mont. & v. d. Bosch <u>ex</u> Jungh.	N. Guinea	374
<i>S. filix</i> (Sw.) Nyl.	Australia	248
<i>S. [Pseudocyphellaria] cf. fragillima</i> Bab. <u>ex</u> Hook., <u>sensu</u> A. Zahlbr.	N. Guinea	375
<i>S. fuliginosa</i> (Dicks.) Ach.	Australia	240
<i>S. rubella</i> Hook. <u>f.</u> & Tayl.	Tasmania	309
<i>S. weigeli</i> Isert <u>ex</u> Ach.	Arizona	92
<i>Strigula elegans</i> (Fee) Muell.-Arg.	Louisiana	284
	Australia	371
<i>Teloschistes exilis</i> (Michx.) Vain.	Texas	356
<i>T. fasciculatus</i> Hillm.	Australia	286
<i>T. flavicans</i> (Sw.) Norm.	Galapagos	108
<i>T. velifer</i> F. Wils.	Australia	235

<u>T. villosus</u> (Ach.) Norm.	Mexico	191
<u>Thamnia vermicularis</u> (Sw.) Schaer.	Colorado	37
	Alaska	211
	Australia	226
<u>T. vermicularis</u> (Sw.) Ach. <u>ssp. solida</u> (Sato)		
W. A. Web.	N. Guinea	380
[<u>Thelidea corrugata</u> Hue] = <u>Knightsiella splachnirima</u>		
(Hook. f. & Tayl.) Gyel.	Tasmania	353
<u>Thelotrema diminutum</u> Hale, Cited specimen.	Australia	453
<u>Thrombium epigaeum</u> (Pers.) Wallr.	Colorado	426
<u>Thysanothecium hookeri</u> Berk. & Mont.	Australia	369
<u>T. hyalinum</u> (Tayl.) Nyl.	Australia	218
<u>Toninia bullata</u> (Mey. & Flot.) A. Zahlbr.	Australia	233
[<u>T. ruginosa</u> (Tuck.) Herre] <u>T. sp. nov. ined.</u>	Colorado	140
<u>T. tristis</u> Th. Fr.	Colorado	174
<u>Trypethelium grossum</u> Muell.-Arg.	N. Guinea	358
<u>T. mastoideum</u> (Ach.) Ach.	Louisiana	287
<u>T. tropicum</u> (Ach.) Muell.-Arg.	Louisiana	288
<u>Umbilicaria arctica</u> (Ach.) Nyl.	Norway	470
<u>U. coriacea</u> Tmschaug	Norway	471
<u>U. cylindrica</u> (L.) Del. <u>ex Duby</u>	Australia	228
<u>U. decussata</u> (Vill.) A. Zahlbr.	Australia	230
<u>U. deusta</u> (Huds.) Baumg.	Colorado	425
<u>U. havaasii</u> Llano	Norway	472
<u>U. hirsuta</u> (Sw.) Ach.	Colorado	69
<u>U. hyperborea</u> (Ach.) Hoffm.	Colorado	162
<u>U. krascheninnikovii</u> (Sav.) A. Zahlbr.	Colorado	216
<u>U. phaea</u> Tuck.	California	131
<u>U. proboscidea</u> (L.) Schrad.	Australia	229
<u>U. subglabra</u> (Nyl.) Harm.	Australia	231
<u>U. vellea</u> (L.) Ach.	Colorado	38
[<u>U. vellea</u> (L.) Ach.] <u>U. cinereorufescens</u> (Schaer.)		
Frey <u>sensu Llano</u>	Colorado	39
<u>Usnea angulata</u> Ach.	Mexico	51
<u>U. angulosa</u> (Muell.-Arg.) Mot.	Australia	331
<u>U. arbusculiformis</u> Mot.	Mexico	70
<u>U. capillacea</u> Mot.	Tasmania	261
	Tasmania	440
<u>U. cavernosa</u> Tuck. <u>in Agassiz</u>	Arizona	48
	Mexico	298
<u>U. cf. cladocarpa</u> Fee	Galapagos	507
<u>U. comosa</u> (Ach.) Roehl.	Arizona	104
[<u>U. diplotypus</u> Vain.] <u>U. herrei</u> Hale. <u>nom. nud.</u>	Kansas	61
	Texas	62
<u>U. aff. elongata</u> Mot.	Galapagos	501
<u>U. flexilis</u> Stirt.	Tasmania	263
	N. Guinea	568

<u>U. herrei</u> Hale, <u>nom. nud.</u>	Arizona	407
<u>U. igniaria</u> Mot.	Chile	562
<u>U. inermis</u> Mot.	Tasmania	262
<u>U. cf. merrillii</u> Mot.	Mexico	488
<u>U. microcarpoides</u> (Muell.-Arg.) Mot.	Australia	329
<u>U. mutabilis</u> Stirt.	Texas	357
<u>U. paradoxa</u> (A. Zahlbr.) Mot.	Galapagos	366
<u>U. poliotrix</u> Kremp.	Australia	326
	Australia	327
	Australia	328
[<u>U. rubiginea</u> (Michx.) Mass.] <u>U. rubicunda</u> Stirt.	Australia	260
<u>U. scabrata</u> Nyl. <u>ssp. nylanderiana</u> Mot.	USSR	597
<u>U. scabrida</u> Tayl.	Tasmania	323
	Tasmania	324
[<u>U. soreidiifera</u> auctt.] = <u>U. fulvoreagens</u> (Raes.) Raes. <u>cf. Brodo</u> in <u>The Bryologist</u> 79:406. 1976 [1977].	Colorado	72
	Colorado	205
	Wyoming	432
<u>U. spilota</u> Stirt.	Australia	566
<u>U. cf. strigosa</u> (Ach.) Eaton	Mexico	575
<u>U. torquescens</u> Stirt.	Australia	316
<u>U. torulosa</u> (Muell.-Arg.) A. Zahlbr.	Australia	325
	Tasmania	330
[<u>U. tristis</u> Mot.] <u>U. florida</u> (L.) Wigg., <u>fide</u> Asahina, J. <u>Jap. Bot.</u> 43:65. 1968.	N. Mexico	116
<u>U. xanthopoga</u> Nyl.	Tasmania	439
<u>Verrucaria laevata</u> Ach.	S. Dakota	40
<u>Xanthoparmelia wyomingica</u> (Gyel.) Hale	Colorado	513
<u>Xanthoria candelaria</u> (L.) Th. Fr.	Mexico	190
<u>X. fallax</u> (Hepp ex Arn.) Arn. <u>sens. latiss.</u>	Texas	354
<u>X. polycarpa</u> (Ehrh.) Rieb.	Colorado	151
<u>X. polycarpa</u> (Ehrh.) Rieb. Identification doubtful.	California	188
[<u>Xylographa spilomatica</u> (Anzi) Th. Fr.] = <u>X. vitiligo</u> (Ach.) Laundon	Colorado	163
<u>X. vitiligo</u> (Ach.) Laundon	Montana	526

NEW COMBINATIONS

Cladina galapagosensis (Ahti) W. A. Web. Cladonia galapagosensis
Ahti, Ann. Bot. Soc. 'Vanamo' 32(1):46. 1961.

Cladina polia (R. Sant.) W. A. Web. Cladonia polia (as "pohlia",
orth. error) R. Sant., Ark. Bot. 30A(10): 15. 1952.

Heterodermia barbifera (Nyl.) W. A. Web. Physcia barbifera Nyl.,
Syn. Lich. 1:416. 1860.

Heterodermia circinalis (A. Zahlbr.) W. A. Web. Anaptychia leuco-
melaena var. multifida f. circinalis A. Zahlbr., Beih. Bot. Centralbl.
19(2):84. 1905.

Heterodermia stellata (Vain.) W. A. Web. Anaptychia podocarpa var. *stellata* Vain., Acta Soc. Fl. Faun. Fenn. 7:131. 1890.

Heterodermia verrucifera (Kurokawa) W. A. Web. Anaptychia leucomelaena f. *verrucifera* Kurokawa, Nova Hedwigia, Beih. 6:72. 1962.

VALIDATIONS OF NEW SPECIES

LECANORA PSEUDOPINGUIS W. A. Web., sp. nov. Thallus saxicolous crustaceus effusus indeterminatus rimoso-areolatus, areolis minus quam $1\ \mu\text{m}$ diam ca. 0.5 mm crassis laevibus pallide citrinis vel albescentibus soredia et isidia desunt. Cortex $30\ \mu\text{m}$ crassus, granulis lutescentibus inspersis, medulla fulvescens, C+ aurantiaca, P-, K-, IKI-, UV+ rubro-violascens. Apothecia sessilia usque ad 1.5 mm diam margine crasso laevi vel undulato-crenulato disco plano livido pulverulento, epihymenio flavo-granuloso C+ aurantiaco hymenio $50\text{--}60\ \mu\text{m}$ crasso sporis 8/nae $7\text{--}9 \times 3\text{--}5\ \mu\text{m}$. Pycnidia cylindrica $200\ \mu\text{m}$ alta $\times 100\ \mu\text{m}$ lata margine prominenti elevato, conidiis arcuatis acicularis $15\text{--}20\ \mu\text{m}$ longis. Ad saxa vulcanica praecipue littoralis.

Thallus saxicolous, crustaceous, effuse, indeterminate, irregularly rimose-areolate with areoles less than 1 mm diam, ca. 0.5 mm thick, smooth, pale citrine to albescent, lacking soredia or isidia. Cortex $30\ \mu\text{m}$ thick, interspersed with yellowish granules; medulla discolored, C+ orange, P-, K-, IKI-, UV+ red-violet. Apothecia sessile, up to 1.5 mm diam, with thick smooth or crenulate-wavy margin, the disk plane, livid, pulverulent; epihymenium with yellow granules, C+ orange; hymenium $50\text{--}60\ \mu\text{m}$ high, spores 8/ascus, $7\text{--}9 \times 3\text{--}5\ \mu\text{m}$; pycnidia cylindric, $200\ \mu\text{m}$ deep $\times 10\ \mu\text{m}$ wide, with prominent raised rim resembling the apothecial margin; pycnoconidia arcuate, acicular, $15\text{--}20\ \mu\text{m}$ long. On volcanic rocks, primarily littoral.

HOLOTYPE. Ecuador. Galapagos Islands. Isla Santa Cruz, vicinity of Academy Bay, on exposed point along shore just east of Darwin Research Station, abundant on rocks just above high tide mark, 10 April 1976, Weber & Lanier, Lich. Exsicc. COLO No. 500 (COLO). Further distribution. Floreana I., Jervis I., Werman I., S. Plazas Islet, Champion I., and probably on every island of the archipelago. One population has been found inland from the seacoast but on a seaward exposure: Santiago I., on rocks at rim of easternmost crater in the highlands, 400 msm, 10 May 1971, L. H. Pike 2112 (COLO L-55204). This is a most unusual occurrence and may be correlated with atmospheric conditions (upwelling of fog) peculiar to the site. *L. pseudopinguis* is one of the dominant species coloring the rocks of the shoreline yellow on many of the islands.

Except for its effuse thallus the new species resembles the lobate littoral Californian *Lecanora pinguis* Tuck. Dr. Chicita Culberson, to whom I am indebted for analyses of the chemistry of *L. pseudopinguis* and

the next species, reported: "L. pseudopinguis contained a compound (Rf classes A4-5, B5, C4-5; spot color with 10% H₂SO₄ and heat is orange brown) in high proportion that is probably a xanthone pigment.... We have seen this same pigment before in a sample from Ahmadjian that was also from the Galapagos Islands but that had been identified as Buellia straminea. The proportions of the minor products of Ahmadjian's sample were different from the Lecanora, and the Lecanora also contained a trace of thiophanic acid and another unknown pigment (probably not a xanthone) that is common in lichens and that we have previously called "pigment SV-1".

LECANORA (Eulecanora) TEXANA W. A. Web., sp. nov. Prothallus destitutus. Thallus effusus indeterminatus laevis vel tartareus non-vel indistinctissime lobatus, usque ad 2 mm crassus, irregulariter rimoso-areolatus vel continuus, pallide flavus. Pseudocortex prosenchymaticus 30-40 μ m crassus ex hyphis erectis, strato algarum 30-50 μ m alto, continuo. Medulla densissima, alba, subtus fulvescens, K-, KC+ luteus. Apothecium immersum vel adnatum, usque ad 2 mm diam, margine vestigiali, excipulo proprio ex hyphis erectis parallelis, disco carneo, plano vel convexiusculo, epihymenio hyalino granuloso, granulis in K dissolvens, hymenio 50-60 μ m alto, subhymenio ex hyphis verticalibus, hypothecio hyalino subtus algifero, ascis IKI+ coeruleis, paraphysibus simplicibus, 2 m diam, gracilibus, non capitatis, sporis 8-nae anguste-ellipsoideis rectis vel plus minusve curvatis, 10-14 x 4-5 μ m. Pycnidium ostiolo aeruginoso loculo 200 μ m diam, pycnoconidiis Eulecanoroideis acicularis, arcte curvatis vel sigmoideis, 10-15 x 0.5 μ m. Ad saxa granitica, areas amplas rupium superficieram verticalibus tectus.

Prothallus lacking. Thallus effuse, indeterminate, smooth to tartareous, very distinctly if at all marginally lobate, up to 2 mm thick, irregularly rimose-areolate or continuous, pale yellow; pseudocortex prosenchymatous, 30-40 μ m thick, of erect hyphae; algal layer 30-50 m high, continuous; medulla very dense, white, becoming discolored brownish below, K-. KC+ yellow wash. Apothecia immersed to adnate, up to 2 mm diam, the margin vestigial or represented by a few small bead-like remnants; proper exciple of erect parallel hyphae; disk flesh-colored, plane to slightly convex; epihymenium hyaline, granulose, the granules dissolving in K; hymenium 50-60 μ m high, difficult to differentiate from subhymenium of vertical hyphae; hypothecium hyaline, with continuous or broken algal layer below; asci IKI+ blue; paraphyses simple, 2 μ m diam, slender, not capitate; spores 8/ascus, narrowly ellipsoid, straight or somewhat curved, 10-14 x 4-5 μ m. Pycnidia appearing as black dots on thallus surface, the ostiolar margin aeruginose, the loculus 200 μ m diam; pycnoconidia of Eulecanora type, acicular, curved in a semicircle, 10-15 x 0.5 μ m. On granite rocks, covering large areas of the vertical faces, rarely fruiting, but when fertile with clusters of apothecia in small plaques of the thallus.

HOLOTYPE. Texas, U. S. A. Gillespie Co. [not Llano as given on the label]: trail to "Balanced Rock", a granite dome rising above the surrounding plain, 4 mi n of Fredericksburg in the Texas "Hill Country"; abundant and dominant on vertical N-facing massive granite blocks, 29 April 1974, Weber, Lich. Exsicc. COLO No. 451 (COLO).

Lecanora texana is apparently closely related to *L. sulphurea* (Hoffm.) Ach., but the apothecium is pale from the beginning, never blackening. Dr. C. Culberson reported: "*L. texana* contained usnic acid, zeorin and a trace of atranorin. In addition there were two or three unidentified triterpenoids that are probably related to zeorin but which do not correspond to leucotylin or leucotylic acid (the only known compounds of this type in our collection). There is also a curious unidentified phenolic compound (R_f classes A3, B2-3, C5; spot color with 10% H_2SO_4 and heat is yellow). This may be a new lichen product."

PSORA CEREBRIFORMIS W. A. Web., *sp. nov.* Thallus terricola cretaceus tumulos hygrosopicos usque ad 5 cm diam formantes, squamulis dense aggregatis sed non imbricatis primo planis mox alte convexis rimosis marginibus revolutis. Cortex argillaceus pseudoparenchymaticus 50-60 m crassus, strato epinecroso crasso albo tectus. Stratum gonidialium continuum. Medulla alba solida crassa, coacto rhizo-hyphoso substrato affixo. Cortex et medulla K-, C-, KC-, P-, IKI-. Apothecia 1-2 mm diam atra marginalia convexa numerosa plerumque aggregata, excipulo ex hyphis parallelis pallide ferrugineo, ascis clavatis 30-40 m longis, sporis ellipsoideis (8)10-12(17) x 5-6 m. Epihymenium ferrugineum K+ vinosum C-, P-, IKI-.

Thallus terricolous, chalk-white, forming hygrosopic convex mounds up to 5 cm or more diam; squamules densely aggregated but not overlapping, up to 5 mm diam, at first plane but very soon highly convex, deeply cracked, with the margins rolled under. Cortex pale clay-colored, of pseudoparenchyma type, 50-60 m thick, covered by a thick white epinecral pruinose layer. Algal layer continuous. Medulla white, solid, thick, attached to the substrate by a rhizo-hyphal felt. Reactions K-, C-, KC-, P-, IKI-. Apothecia 1-2 mm diam, black, marginal, highly convex, abundant and often clustered; exciple of parallel hyphae, pale reddish-brown; asci clavate, 30-40 m long; spores ellipsoid, (8)10-12(17) x 5-6 m; epihymenium reddish-brown, K+ vinose, C-, P-, IKI-.

HOLOTYPE. U. S. A. Colorado. Montrose Co.: on gypsum knolls, floor of Paradox Valley 4 mi E of Bedrock, 1,500 msm, 30 May 1960, Shushan, Anderson & Weber, Lich. Exsicc. COLO No. 24 (COLO).

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CULTURAL STUDIES ON PORIA CINERASCENS,

P. RIVULOSA, AND P. SUBVERMISPORA

(Aphylliphorales, Basidiomycotina)

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ABSTRACT

The culture complex involving Poria cinerascens, P. rivulosa, and P. subvermispora is examined. Cultural descriptions are provided for each species. Although nearly identical, cultures of these three species can be separated by temperature studies, growth on cabbage extract agar, and microscopic characteristics.

INTRODUCTION

Sporophores of Poria cinerascens (Bres.) Sacc. et Syd., Poria rivulosa (Berk. et Curt.) Cke., and Poria subvermispora Pilát are distinct and easily identified by observing the hyphal systems and basidiospores (Lowe 1966). However, when grown in pure culture they are nearly indistinguishable. Nobles (1948) treated P. rivulosa (as P. albipellucida Baxter) and P. cinerascens together under one culture description. Later, Nobles (1965) and Stalpers (1978) distinguished these fungi on minor or variable characters.

Because of the difficulty in identifying these cultures, the importance of individual species of the complex in the decay of products, preservative-treated wood, and other wood forms cannot be assessed accurately. Duncan and Lombard (1965), in their summary report on Hymenomycetes associated with decay of wood products, report these fungi collectively as the Poria cinerascens complex. Thus, this study was undertaken to resolve the P. cinerascens culture complex.

MATERIALS AND METHODS

Cultures were grown at 25° C in the dark on 1.5% malt extract agar (MEA) in glass petri plates and were examined at weekly intervals.

^{1/} Maintained at Madison, Wis., in cooperation with the University of Wisconsin.

Cultures were grown also on 0.5% gallic acid agar (GAA), on 0.5% tannic acid agar (TAA) (Davidson *et al.*, 1938), and on cabbage extract agar (CAB) at 25° C and measured at 7 and 14 days.

Cabbage extract agar was made from fresh cabbage because growth of these fungi on commercial media was not satisfactory. CAB is prepared as follows: Place 500 gm chopped, fresh cabbage in 1000 ml distilled water, cover, and bring to a boil. Simmer about 15 hours (over several days if necessary) and add about 1,000 ml distilled water during this time. Filter through cheese cloth and bring final volume of cabbage extract to 1000 ml. Add 20 gm agar, 2.5 gm diastasic sodium phosphate, 5 gm sodium chloride, and 10 gm Difco Bacto-peptone to the 1000 ml of cabbage extract and autoclave.

Unless otherwise indicated, all cultures studied are of polysporous origin. Monosporous isolates were obtained from fruiting in culture. Key patterns of 2-week-old cultures follow the format of Davidson *et al.* (1942). Species codes of 6-week-old cultures follow the format of Nobles (1965). Cultures are on deposit at the Center for Forest Mycology Research.

Sporophores associated with the cultures were previously identified by Drs. J. L. Lowe, R. L. Gilbertson, or M. J. Larsen. Although the sporophores were not reexamined by the author, dikaryotic-monokaryotic matings were done with all cultures studied. Cultures listed under each species had successful matings with one or more haploids of that species.

CULTURE DESCRIPTIONS

Poria cinerascens (Bres.) Sacc. et Syd., Syll. Fung. 16:161. 1902.

Growth characters: Growth on MEA rapid, plates covered in 1 wk; mats white, appressed, thin, subfelty, with a network of thin, radial strands, sometimes slightly raised and downy or felty at 2 wk, may develop small white balls of mycelium at 6 wk; margin even, appressed; odor none; agar discoloration none; not fruiting by 6 wk. Oxidase reactions at 1 wk on GAA moderately strong, mat 37-51 mm diam; on TAA strong, mat 12-26 mm diam. Optimum temperature 36° C (Fig. 1). Growth on CAB 0-trace at 1 wk.

Microscopic characters: Hyphae of advancing zone 6-8 µm diam, thin-walled, simple septate, branched, becoming thick-walled and scattered by 2 wk. Hyphae of submerged mycelium 2-4 µm diam, thin-walled, nodose septate, branched. Hyphae of aerial mat (a) similar to submerged hyphae except occasionally encrusted with hyaline crystals; (b) "binding hyphae" very slender, 0.5 µm diam, aseptate, richly branched, hyaline, nonstaining in phloxine, rare to abundant at 2 wk; (c) fiber hyphae (Fig. 2) 4-6 µm diam, thick-walled, walls thinning toward apex, aseptate, hyaline nonstaining in phloxine, apparently lacking or numerous at 2 wk. Chlamydospores globose to limoniform, 8-13.5 x 8-13.5 µm, thin-walled, walls thickening in age, hyaline, terminal or intercalary, few to abundant in aerial and submerged mycelium. Bulbils sometimes present in submerged mycelium.

Key patterns: A-P-F-1-2-10; A-P-F-1-2-10-16; A-P-F-1-2-11; A-P-F-1-2-11-16; A-P-F-1-2-10-14.

Species code: 2.4.(7).(8).(22).34.36.38.42.55.59.

Monosporous cultures: Eight monosporous cultures of HHB 76 (4 of each mating type) examined were similar to the polysporous cultures except that they did not have clamp connections.

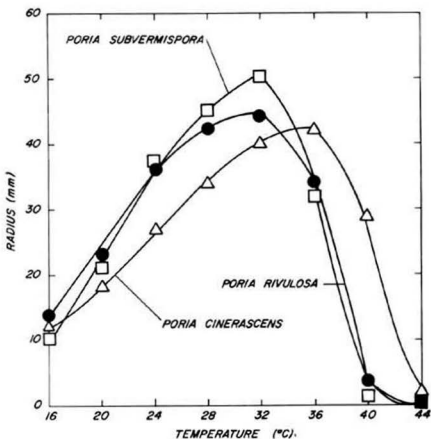
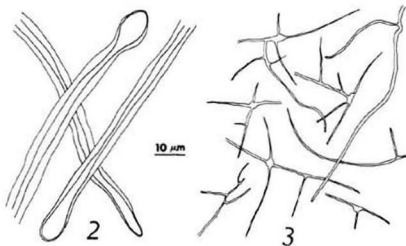


Fig. 1. Average radial growth of *Poria* species on MEA after 4 days at 8 temperatures. (M 149 135)



M 149 140

Fig. 2. Fiber hyphae from a culture of *Poria cinerascens* (FP 100506) at 2 wk.

Fig. 3. "Binding hyphae" from cultures of *Poria rivulosa* and *P. subvermispora* at 2 wk. (M 149 140)

Incompatibility system: Poria cinerascens has been reported by Nobles et al. (1957) to be heterothallic and bipolar. Pairing of 17 monosporous cultures of HHB 76 confirm their finding: $A_1 = 1,4,5,8,11,13,16,18$; $A_2 = 2,3,6,7,9,10,12,17$.

Cultural description: Nobles (1948); Singh (1966).

Cultures studied: U.S.A.: MARYLAND--FP 100506 on pine, Prince Georges County; SOUTH DAKOTA--FP 105939 on Pinus ponderosa Dougl. ex Laws. (ponderosa pine), Black Hills National Forest; TEXAS--HHB 76 on Pinus sp., San Jacinto County; WISCONSIN--FP 105349 on Pinus banksiana Lamb. (jack pine), Wood County.

Remarks: The development of fiber hyphae and optimum temperature of 36° C are diagnostic for P. cinerascens. Poria cinerascens can quickly be separated from P. rivulosa and P. subvermispora by growing the cultures on MEA at 40° C (Fig. 1). At 4 days P. cinerascens grows significantly more than the other species. Cultures of P. cinerascens will often deteriorate after growing several years on artificial media. Deteriorated cultures are slow growing, sodden, and do not develop fiber hyphae.

Poria cinerascens is widely distributed in the U.S.A. and is associated with a white rot of conifers.

Poria rivulosa (Berk. et Curt.) Cke., Grevillea 14:109. 1886.

Growth characters: Growth on MEA rapid, plates covered in 1 wk; mats white, azonate, at 1 wk appressed, thin, subfelty, at 2 wk slightly raised, thicker, felty, subfelty to felty; margin even, appressed; odor none; agar discoloration none; not fruiting by 6 wk. Oxidase reactions at 1 wk on GAA moderate, mat 64-90 mm diam; on TAA strong, mat 19-29 mm diam. Optimum temperature about 32° C (Fig. 1). Growth on CAB 20-48 mm diam at 1 wk.

Microscopic characters: Hyphae of advancing zone 6-7.5 mm diam thin-walled, simple septate, branched, by 2 wk often becoming thick-walled, rare to abundant, absent in older cultures. Hyphae of submerged mycelium 3-5 mm diam, thin-walled, nodose septate, branched. Hyphae of aerial mat (a) 1.5-5 mm diam, similar to submerged hyphae, sometimes encrusted with hyaline crystals; (b) "binding hyphae" (Fig. 3) very slender, 0.5 mm diam, aseptate, richly branched, hyaline, nonstaining in phloxine, usually present by 2 wk in clusters, difficult to separate. Chlamydo spores globose to limoniform, 8.5-11.5 x 8.5-11.5 mm, thin-walled, becoming thick-walled in age, hyaline, terminal or intercalary, abundant in aerial mat.

Key patterns: A-P-F-1-2-10-16; A-P-F-1-2-10; A-P-F-1-2-10-14-16; A-P-F-1-2-10-14.

Species code: 2.4.(7).(8).34.36.38.41.42.55.59.

Monosporous cultures: Eight monosporous cultures of FP 133035 (4 of each mating type) were similar to the polysporous cultures except that thick-walled advancing zone hyphae were present in 6-wk-old cultures, and they lacked clamp connections and "binding hyphae."

Incompatibility system: Nobles et al. (1957) reported that P. rivulosa is heterothallic and bipolar. Pairings of 17 monosporous cultures of FP 133035 confirm their result: $A_1 = 1,3,4,5,9,12,13,14,16,17$; $A_2 = 2,6,7,8,10,11,18$.

Cultural descriptions: Buckland (1946), Baxter (1937), and Nobles (1948) as Poria albidipellucida Baxter; Stalpers (1978) as P. lindbladii (Berk. et Br. ex Berk.) Cooke.

Cultures studied: U.S.A.: CALIFORNIA--Pirto 26 (rot) in Sequoiadendron giganteum (Lindl.) Bucholz (giant sequoia); JLL 10602 on Sequoia sempervirens (D. Don) Endl. (redwood), Del Norte County; FP 104207 (rot) in redwood, Wilson Creek; IDAHO--JLL 6987 on conifer, Kootenai County; OREGON--FP 133035 (rot) in Tsuga sp., Benton County, FP 133406 and FP 133696 on conifer, Lincoln County. CANADA: BRITISH COLUMBIA--DAOM 9757 on Pseudotsuga menziesii (Mirb.) Franco (Douglas-fir).

Remarks: Poria rivulosa is most similar culturally to P. subvermispora. They can be separated, however, by growing the cultures on CAB. Poria rivulosa will grow 20-40 mm in diameter at 1 wk but P. subvermispora only 0-trace.

Poria rivulosa is common in the northwest U.S.A. and British Columbia on conifers. It is associated with a white ring rot of living western redcedar, Thuja plicata Donn ex D. Don (Buckland 1946) and living redwood (Kimmey and Lightle 1955).

Poria subvermispora Pilát, Studia Bot. Cech. 3:2. 1940.

Growth characters: Growth on MEA rapid, plates covered in 1 wk; mats white, azonate, at 1 wk appressed and subfelty or raised and downy, at 2 wk appressed, subfelty to felty, or raised and woolly, sometimes tufted; margin even, appressed; odor none; agar slightly bleached in age; not fruiting by 6 wk. Oxidase reactions at 1 wk on GAA moderate, mat 69-90 mm diam; on TAA strong, mat 22-30 mm diam. Optimum temperature 32° C (Fig. 1). Growth on CAB 0-trace at 1 wk.

Microscopic characters: Hyphae of advancing zone 6-7 µm diam, thin-walled, simple septate, infrequently branched, by 2 wk becoming thick-walled, rare. Hyphae of submerged mycelium 2-4 µm diam, thin-walled or with slight wall thickening, nodose septate, branched. Hyphae of aerial mat (a) similar to submerged hyphae, occasionally encrusted with hyaline crystals; (b) "binding hyphae" very slender (Fig. 3), 0.5 µm diam, aseptate, richly branched, hyaline nonstaining in phloxine, usually present in clusters by 2 wk, difficult to separate; (c) fiber hyphae rare, 2 µm diam, thick-walled, aseptate, hyaline nonstaining in phloxine, infrequently branched, seen only in 6-wk-old cultures of FP 90031. Chlamydospores globose to limoniform, 11.5-13.5 (-16) x 7.5-13.5 (-16) µm, thin-walled, walls thickening in age, hyaline, terminal or intercalary, numerous in submerged and aerial mycelium.

Key patterns: A-P-F-1-2-10-16; A-P-F-1-2-10; A-P-F-1-2-10-14-16; A-P-F-1-2-10-14.

Species code: 2.4.(7).(8).34.36.38.41.(42).54.55.59.

Monosporous cultures: An examination of 8 monosporous cultures obtained from JLL 14807 (4 isolates of each mating type) reveals that they are similar to the polysporous isolates except that their mats tend to be appressed, develop delicate radial to reticulate patterns, and lack clamp connections.

Incompatibility system: Nobles *et al.* (1957), as P. notata Overh., and Domański (1969), as Fibuloporia subvermispora (Pilát) Domański, reported that P. subvermispora is bipolar. Pairings of 17 monosporous cultures of JLL 14807 confirm their results: $A_1 = 1, 3, 4, 5, 9, 12, 13, 15, 16, 18, 19$; $A_2 = 7, 8, 10, 11, 14, 17$.

Cultural descriptions: Baxter (1947) (as P. quercuum Baxter); Nobles (1948); Domański (1969); Stalpers (1978).

Cultures studied: U.S.A.: ARIZONA--JLL 8904 on ponderosa pine, Cochise County; COLORADO--JLL 6133 on conifer, Garfield County; JLL 6332 on lodgepole pine, Larimer County; FLORIDA--FP 104027 (sporophore tissue) on *Quercus laurifolia* Michx. (laurel oak), Duval County; MARYLAND--FP 90031 on oak and FP 105752 on hardwood board, Prince Georges County; NEW YORK--JLL 3292 on conifer, Essex County; JLL 15225 on hardwood, Ulster County; OREGON--ME 485 (rot) in Douglas-fir wood chips, Clatsop County. CANADA: Saskatchewan--JLL 14807 on *Populus tremuloides* Michx. (aspen).

Remarks: *Poria subvermispora* is widely distributed throughout continental U.S.A. It is associated with a white rot of conifers and hardwoods and has been isolated several times from wood chip piles in Washington and Oregon.

DISCUSSION

Cultures of *P. cinerascens*, *P. rivulosa*, and *P. subvermispora* can be identified by noting several critical characters. These are summarized in Table 1. Dikaryotic-monokaryotic pairings can be used also to identify cultures once monosporous cultures are obtained. Cultures of the three species will fruit on MEA and discharge spores if kept long enough. The shape and size of the basidiospores can also be used to differentiate between the species (Table 1).

Development of the slender "binding hyphae" found in the aerial mat of all three species has not been recorded previously. Their development in culture suggests their presence in the sporophore; however, this is not the case. *Poria rivulosa* and *P. subvermispora* have monomitic hyphal systems (Lowe 1966). However, *Poria cinerascens* sporophores have a trimitic hyphal system (Lowe 1966), and this is supported in culture.

Table 1.--Characteristics that differentiate cultures of *Poria cinerascens*, *P. rivulosa*, and *P. subvermispora*

	Optimum growth temperature	Growth at 40° C ^{1/}	Growth on CAB ^{2/}	Host	Basidio- spore size ^{3/}
	°C	mm	mm		µm
<i>Poria cinerascens</i>	36	30 - 36	0-trace	Conifers	4.0 - 7.0 x 1.5 - 2.0
<i>P. rivulosa</i>	32	1 - 7	20 - 48	Conifers	5.0 - 6.0 x 4.0 - 5.0
<i>P. subvermispora</i>	32	1 - 2	0-trace	Conifers and hardwoods	3.5 - 6.0 x 1.0 - 1.5

1/ Average radial growth after 4 days on malt extract agar.

2/ Average mat diameters on cabbage extract agar after 7 days at 25° C.

3/ Spore measurements taken from Lowe (1966).

Culturally, these three species belong to a unique group of fungi. The rapid growth, production of extracellular oxidases, simple septate hyphae of the advancing zone that later give rise to nodose septate hyphae, and the bipolar mating system are characters shared by all species of Group 54 (Nobles 1958). *Phlebia subserialis* (Bourd. et Galz.) Donk, *Phlebia subochracea* (Bres.) Erikss. et Ryv., and *Hyphodontia setulosa* (Berk. et Curt.) Maas G. are the most culturally similar to the *Poria* species discussed.

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A NEW SPECIES OF AMANITA

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ABSTRACT

Amanita marginata (fig. 1) is described as new from Tennessee. It is assigned to Section *Lepidella*, Subsection *Solitariae*.

Amanita marginata Jenkins, sp. nov.

Holotype: Tennessee, Loop Road, Cades Cove, Great Smoky Mountains National Park, 17. viii. 72, David T. Jenkins 580(DTJ).*

Pileus 110 mm mense diametro, convexus, margo leviter appendiculatus, non striatus; labrum distinctum sterileque subter, album convertens ad gilvum colorem; reliquae volvae ita verrucae adnatae factaeque in inaequalem formam, gilvae cum cacumenibus in vero colore, potius in circulos qui easdem medias partes habent. Lamellae stipatissimae, solutae. Stipes 90 x 20 mm, solidus, fibrillosus squameusque adversus basem; tegumen ad apicem partim, pendens, album, delicatum; bulbus ad basem in ovi formam factus, gilvus; maculans flammeus fuscusque adversus fundum; aliquando rimae longitudine; reliquae volvae ita circli squamarum adnatarum reflexarumque leviter, qui easdem medias partes habent. Sporae 5.46-6.25 x 7.0-10.15 μ m.

Fruit body medium to large, solitary. PILEUS: 110 mm diam, convex, margin non-striate, very slightly appendiculate, but with a distinct, sterile margin, up to 3 mm wide, pileipellis fairly easily separable, fibrillose between volval remnants, white to creamy-white, flesh white, up to 15 mm thick at center, tapering toward margin; universal veil remnants as fibrous, adnate, irregular-shaped warts, up to 7 mm wide at base, cream with tips cream to pale tannish-cream, more or less concentric ring arrangement, becoming decidedly smaller toward margin. LAMELLAE: very crowded, free, fairly broad, cream, edges smooth; lamellulae numerous, attenuate. STIPE: 90 x 20 mm, solid, white, fibrillose-scaly beneath annulus, becoming lacerate-scaly toward base; partial veil apical, creamy-white, pendant, large and thick, striate above, floccose below, double-edged, very delicate, soon falling off, ring of universal veil material approximately 10 mm below partial veil. Basal bulb ovoid, cream at apex, staining orangish-brown toward bottom, 65 x 45 mm, occasionally with longitudinal splits; universal veil remnants as concentric rings of adnate, slightly recurved scales, quite thick, on upper part of bulb. Faint chloride of lime smell.

PILEIPELLIS: filamentous hyphae densely interwoven, slightly gelatinized, hyaline, 2-7 μ m diam. PILEUS TRAMA: filamentous hyphae

* DTJ = The author's herbarium

undifferentiated, moderately branched, clamps rare, 3-8 μm diam; inflated cells abundant, mostly terminal, variform, mostly elongate, up to 162 x 31 μm . LAMELLA TRAMA: bilateral; filamentous hyphae undifferentiated, moderately branched, no clamps, 3-8 μm diam; inflated cells terminal, elongate. SUBHYMENIUM: hyphae ramose, clamps rare. BASIDIA: up to 48 x 3.9-8.6 μm , 4-sterigmate, thin walled, without clamps. UNIVERSAL VEIL: filamentous hyphae on pileus moderately abundant, sparsely to moderately branched, without clamps, 3-6 μm diam, irregularly disposed; inflated cells abundant, globose, subglobose, broadly elliptic, elliptic, clavate, cylindric, most being quite small, not exceeding 62 x 46 μm , usually short, terminal chains, irregularly disposed: tissue on basal bulb similar, but with a slightly higher percentage of filamentous hyphae. STIPE TRAMA: filamentous hyphae undifferentiated and abundant, sparsely branched, without clamps, 3-6 μm diam; the presence of gloeoplerous hyphae conspicuous; inflated cells terminal, clavate, longitudinally oriented, up to 312 x 25 μm . PARTIAL VEIL: upper surface mostly inflated cells, elliptic, ventricose, clavate, up to 125 x 63 μm , mostly terminal; filamentous hyphae sparse, sparsely branched, without clamps, 2-7 μm diam: lower surface similar to that above, but with many more smaller inflated cells, variform; filamentous hyphae also sparse, similar to above.



Fig. 1 *Amanita marginata* Jenkins 580(DTJ)

SPORES: 5.46-6.25 x 7.0-10.15 μ m, (\underline{E} - 1.25-1.62; \underline{E}^m - 1.51), elliptic to elongate, adaxially flattened, thin walled, hyaline, weakly amyloid, spore print white; contents guttulate; apiculus sublateral, short cylindrical.

Habitat and distribution: terrestrial, mixed coniferous and deciduous forest, Tennessee.

Collections examined: road cut, Loop Road, Cades Cove, Great Smoky Mountains National Park, Tennessee, 17. viii. 72, David T. Jenkins 580(DTJ).

Discussion: *Amanita marginata* is assigned to Section *Lepidella*, subsection *Solitariae*. This is due to the presence of such characters as amyloid spores, an appendiculate pileus margin, a white to cream pileus color, and variform volval elements.

As attempt has also been made to assign this organism to a particular stirps. Following the organization of Bas (1969) this organism would seem to belong to Stirps *Strobiliiformis*, based upon the absence of basidial clamps, white volval remnants with moderately abundant, interwoven elements, and the production of irregularly shaped volval warts on the pileus.

Although this organism is apparently related to the members of Stirps *Strobiliiformis*, only slight changes in a particular character would allow it to fit into other stirps. For example, if the basidia had clamps this organism could be placed into Stirps *Virgineoides* due to its small to medium, detersile volval warts and the irregularly disposed universal veil remnants, consisting of moderately abundant hyphae and inflated cells. If the elements of the volval remnants on the pileus had a more erect-parallel disposition, it would be more closely related to Stirps *Polypyraxis*, due primarily to its color and small to medium, detersile warts.

As the *Amanita* flora of the United States, and in particular the southeastern region, is further studied the variation and diversity becomes increasingly apparent. As can be seen with *Amanita marginata* it is frequently quite difficult to determine the specific relationship with other members of this section based upon the current classification system. Therefore, the establishment of new stirps within this section may be necessary in the future.

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CANDIDA PARATROPICALIS, A NEW SPECIES OF CANDIDA

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An unusual *Candida* isolated from clinical specimens has been found to be sufficiently different from existing taxa within this genus to warrant its establishment as a new species, *C. paratropicalis*.

MATERIALS AND METHODS

Twenty-nine isolates of the new *Candida* were recovered from specimens of human blood, sputa, urine, bronchial washings, throat swabs, a lung biopsy, and a decubitus ulcer. Their morphologic and physiologic characters were studied by procedures previously described (Baker, Salkin, Pincus, and D'Amato, 1981a, b).

TAXONOMIC PART

Division: Fungi imperfecti.
Form class: Blastomycetes.
Form order: Cryptococcales.
Form family: Cryptococcaceae.

Candida paratropicalis Baker, Salkin, Pincus et D'Amato, sp. nov. (Figure 1).

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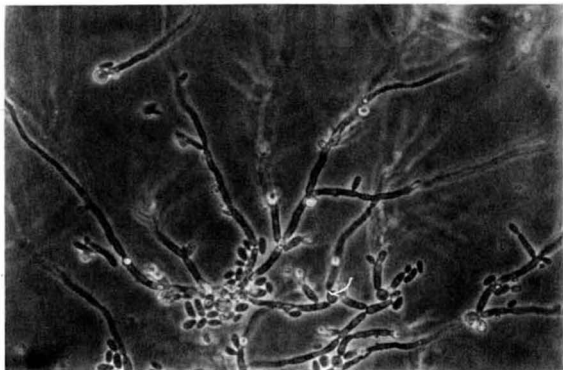


Fig. 1. *Candida paratropicalis* on cornmeal + 1% Tween 80 agar after 7 days incubation at 25°C.

Coloniae in extracti malti agaro cultae 2 mm diam usque sunt, ochroleucae farinulentae, lucidae vel sublucidae, planae subconvexae vel umbonatae, marginibus filamentosis patulis. Per dies 14 in glucoso-fermenti extracto cum aqua peptonica immersis sedimentum pelliculumque formantur, cellulis fertilibus ovoideis vel elongato-ovoides ($2.5\text{--}7.5 \times 3.0\text{--}12.5 \mu\text{m}$) cum quibus blasticonidia gemmantia numerosa collo angusto conjunguntur; pseudohyphae hyphaeque genuinae adsunt. In farinae Zeae agaro per dies 7 cultae fermenti cellulae ovoideae vel elongato-ovoides ($2.5\text{--}6.5 \times 3.5\text{--}14.0 \mu\text{m}$) cum pseudohyphis hyphisque genuinis formantur. Status sexualis ignotus. Notulis physiologicis (tab. 1) cum *C. tropicalis* (Cast.) Berkhout congruit, sed sacrosum melezitotum fermentare necnon L-arabinosum assimilare nequit simulque methyl-D-glucosidum sacrosum melezitotumque modo variabili assimilant.

Habitat: Man.

Holotype: 79MR8, isolated from human blood at autopsy, New York, New York.

The holotype slide preparations, as well as living cultures, have been deposited in the American Type Culture Collection, Rockville, Maryland (accession number ATCC 42678), and in the New York Botanical Garden Culture Collection, Bronx, New York.

The epithet *C. paratropicalis* was chosen to emphasize the resemblance of this species to *C. tropicalis*.

Colonies on malt-extract agar at 14 days are 2 mm in diameter; yellowish-white; pasty; glossy to dull glossy; and umbonate, flat, or slightly convex, with spreading filamentous margins. At 14 days in glucose -- yeast-extract -- peptone water a sediment and pellicle have formed; reproductive cells are ovoid to long-ovoid (2.5-7.5 x 3.0-12.5 μm) with multiple budding blastoconidia, each attached by a narrow neck; and pseudohyphae and true hyphae are present. When grown on cornmeal agar for 7 days, the cells are ovoid to long-ovoid (2.5-6.5 x 3.5-14.0 μm) with well-ramified pseudohyphae and true hyphae. The teleomorph is unknown. Physiologic characters (Table 1) are similar to those of *C. tropicalis* (Cast.) Berkhout except for (i) the inability of *C. paratropicalis* to ferment sucrose and melezitose and assimilate L-arabinose and (ii) its variable assimilation of methyl-D-glucoside, sucrose, and melezitose.

DISCUSSION

The morphology of *C. paratropicalis* (shape and size of blastoconidia, development of blastoconidia in sparse clusters at hyphal nodes, absence of chlamydo-spores and germ tubes) suggests a close similarity to *C. tropicalis* and its sucrose-negative variant (Ahearn, Meyer, Mitchell, Nicholson, and Ibrahim, 1977; Baker et al., 1981a). However, the new species' inability to ferment sucrose and melezitose clearly differentiates it from *C. tropicalis* (van Uden and Buckley, 1970). The utilization of methyl-D-glucoside, melezitose, and sorbose by many isolates of *C. paratropicalis* (24%, 52%, and 90% respectively; Baker et al., 1981a) and its inability to assimilate inulin distinguish this taxon from the sucrose-negative form of *C. tropicalis* (Ahearn et al., 1977).

Since isolates of *C. paratropicalis* frequently (41%) do not assimilate sucrose, cursory examination of such isolates could contribute to their misidentification as *C. stellatoidea*.

The lack of recognition of this new species despite its relative frequent recovery from diverse clinical specimens in several laboratories suggests that *C. paratropicalis* has been and probably still is being misidentified.

ACKNOWLEDGMENTS

The authors express their appreciation to Dr. Rupert Barneby for preparation of the Latin translation of the diagnosis, to Drs. G. A. Land and G. D. Roberts for supplying some of the cultures used in this study, and especially to Dr. Michael R. McGinnis for providing taxonomic assistance.

TABLE 1. FERMENTATION AND ASSIMILATION PATTERNS OF 29
C. paratropicalis ISOLATES^a

Substrate	Isolates positive (%)	
	Fermentation	Assimilation
Cellobiose	0	69
Dextrose	100	100
Galactose	100	100
Inulin	0	0
Lactose	0	0
Maltose	100	100
Melezitose	0	52
Melibiose	0	0
Raffinose	0	0
Sucrose	0	59
Trehalose	100	100
D-Arabinose		0
L-Arabinose		3
Citric acid		100
Dulcitol		0
Erythritol		0
Ethanol		100
Ethylamine hydrochloride		100
Glucitol		100
Glycerol		14
Inositol		0
2-Keto-gluconate		100
DL-Lactic acid		86
Mannitol		100
Methyl-D-glucoside		24
Potassium nitrate		0
Ribitol		100
Ribose		10
Rhamnose		0
Salicin		24
Sorbose		90
Succinic acid		100
Xylose		100

^aResults obtained with Wickerham broth technique at 3 weeks.

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NOTES ON CORTICIACEAE (BASIDIOMYCETES) VIII.

Two new species of *Tubulicrinis*.

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SUMMARY

Tubulicrinis cinctoides and *T. ovalisporus* are described from North Europe and Africa respectively. The first species is similar to *T. cinctus* G.H.Cunn. but delimited by its more globose spores and cystidia of divergent appearance. The African species is related to the *T. glebulosus*-group in its cystidial morphology, but well defined by having oval spores. The type of *T. cinctus* was studied.

TUBULICRINIS CINCTOIDES Hjortst. nov. spec. Fig. 1, a-c.

Species *Tubulicrinis cincto* affinis sed sporis oblique subglobosis (4.75-)5 x (3.75-)4-4.5 μ m; lycocystidiis indistincte capitatis, 60-70(-90) x 5-6 μ m.

Holotypus: Sweden. Östergötland. V. Tollstad par., Omberg, Storpissan state forest reserve, in herb-rich *Picea* forest, on decayed trunk of *Picea abies*. 1979-11-09. K.Hjortstam & T.Hallingbäck. Hjm 11348 (GB).

Paratypus: do. Hjm 11393 (priv.herb.).

Fruitbody resupinate, effuse, at first reticulate, then more or less continuous, thin, whitish, with the cystidia projecting, each with a globule of excreted matter easily seen under a lens (50-100 X).

Subiculum very sparse, composed of thin-walled, more or less uniform hyphae, about 2-2.5 μ m wide, with a clamp at each septum.

Lycocystidia faintly amyloid, mostly greyish in Melzer's reagent, cylindrical, up to 60-70(-90) μ m long and 5-6 μ m wide near the bi-, seldom tri-furcate base; the capillary lumen ending more or less abruptly; neck-width (3.5-)4-5 μ m, the apices of the cystidia slightly expanded but indistinctly capitate, usually not more than 6-7 μ m in diam.

Basidia small, 12(-15) x (4.5-) 5 (-6) μ m, with four sterigmata and a clamp at the base, not amyloid.

Spores obliquely subglobose, thin-walled, with neither amyloid nor cyanophilous reaction, (4.75-) 5 x (3.75-) 4-4.5

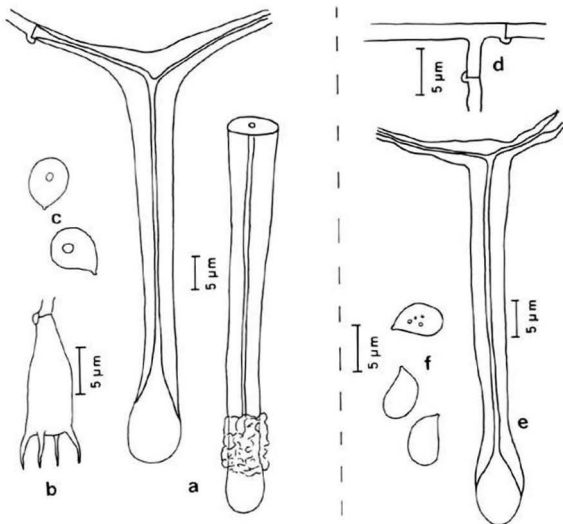


Fig.1. Tubulicrinis cinctoides a) cystidia b) basidium c) spores. - Coll. Hjm 11348 (holotypus). T. cinctus d) hypha e) cystidium f) spores. - Coll. Cunn. 17428 (holotypus). um, with a distinct apiculus.

Remarks. The new species T. cinctoides is undoubtedly very similar to T. cinctus G.H.Cunn. but seems to be well delimited by its more globose and slightly larger spores. In the type of the latter (as far as known the only material gathered) the spores are more or less ellipsoid and measure 4.2-4.5 (-5) x 2.75-3.25 (-3.5) um (in the original description 4-4.5 x 3-3.5 um), while most spores of T. cinctoides measure 4.75-5 x 3.75-4 um. The lycocystidia of the two species show a slight but clear difference. In T. cinctus the apical bulb is fairly distinct, starting somewhat abruptly from the neck (fig.1,e), and thick-walled in the basal part, while the cystidial-bulb of the new species is less marked and usually consists only of the thin-walled part of the apex (fig.1,a).

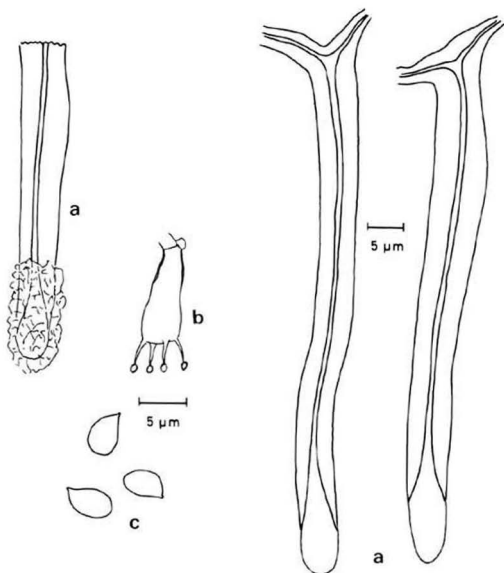


Fig.2. *Tubulicrinis ovalisporus* a) cystidia b) basidium c) spores. - Coll. Ryv. 11308 (holotypus).

TUBULICRINIS OVALISPORUS Hjortst. nov. spec. Fig. 2.

Species *Tubulicrinini* glebuloso affinis sed fructificatione subtiliter odontioide et sporis ovalibus $4.5 (-5) \times 2.5-2.75 (-3) \mu\text{m}$; lycystidiis cylindricis, pallide amyloidibus (fere griseis), circiter $100 \times 5-6 \mu\text{m}$, versus apicem leviter angustum.

Holotypus: Africa. Malawi. Southern Prov., Mulanje distr., Mulanje Mts., Lichenya Plateau, alt. 1800-2000 m.a.s.l., on coniferous wood, probably *Widdingtonia*. 1973-03-09/10. L. Ryvarden 11308 (0). Isotypus: in GB.

Paratypus: do. L. Ryvarden 11382 (0).

Fruitbody resupinate, effuse, thin, composed of very small aculei, about 15-20/mm, whitish or becoming yellowish brown, with numerous cystidia projecting above the hymenium and easily observed under a lens (50-100 X), each encrusted at the apex with brownish, excreted material.

Subiculum thin, mostly inconspicuous, composed of thin-walled, uniform hyphae with clamps, usually 2-3 μ m wide.

Lycystidia numerous, cylindrical, faintly amyloid, about 100 μ m long and 5-6 μ m wide near the bi-furcate base, tapering slight towards the apex and with a neck-width of 4-5 μ m, the capillary lumen ending gradually at the slightly tapering, but obtuse, apical part.

Basidia usually 13-15 x 4.5-5 μ m, with four sterigmata and with a basal clamp, not amyloid.

Spores ovate or ellipsoid, thin-walled, 4.5 (-5) x 2.5-2.75 (-3) μ m, with neither amyloid nor cyanophilous reaction.

Remarks. Owing to the cystidial morphology this new species seems to be closely related to I. angustus (Rog. & Weres.) Donk and I. glebulosus (Bres.) Donk but is distinguished by its ovate to ellipsoid spores. The finely odontoid fructification is somewhat unusual in other species of the genus, but recently Jülich described one species (I. corneri) which, according to the description, is quite close to I. hamatus (Jacks.) Donk but distinctly toothed, with fairly large aculei. In fresh material, similar growth structure can be demonstrated for example in I. inornatus (Jacks. & Rog.) Donk, where it is less pronounced than in I. ovalisporus.

Acknowledgement

The author is grateful to Carl Stenholm's foundation for supporting the investigation of the Storpissan nature reserve. In a subsequent paper the region of Omberg will be treated more thoroughly.

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NOTES ON
CORTICIACEAE (BASIDIOMYCETES) IX.Three new combinations in *Hypochniciellum*.

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SUMMARY

The genus *Hypochniciellum* with the type species *Leptosporomyces ovoideus* Jülich was described by Hjortstam and Ryvarden (1980) and characterized by pellicular to membranaceous fruitbodies and thick-walled, cyanophilous spores.

Three new combinations are proposed here: *H. cremeoisabellinum*, *H. molle*, and *H. subillaqueatum*. The generic circumscription is slightly emended to include also species with spores greyish but not distinctly blue in Melzer's reagent.

INTRODUCTION

When Eriksson and Ryvarden (1976) treated the genera *Leptosporomyces* and *Leucogyrophana* they discussed several similarities between *Leptosporomyces ovoideus* Jülich and *Leucogyrophana cremeoisabellina* (Litsch.) Parm., *L. mollis* (Fr.) Parm., and *L. subillaqueata* (Litsch.) Jülich. The authors also expressed the possibility of joining the species in a separate genus.

Later on Ginns (1978) circumscribed the genus *Leucogyrophana* and excluded the three species mentioned above without any suggestions as to their generic position. Consequently, the species need a place in the family Corticiaceae, and in my opinion *Hypochniciellum* seems to be suitable.

Without any doubt the thickness of the spore-wall is a character to consider, uniting the species. In spite of the fact that the spores of *H. ovoideum* are non-amyloid in comparison with the other species, which have a more or less greyish reaction, the species have many characters in common.

The genus *Amyloathelia* Hjortst. (1979) is in many respects not appropriate for the species here referred to *Hypochniciellum*. All species in that genus have strongly amyloid spores and thinner spore-walls.

Key to the species of *Hypochniciellum*

1. Cystidia present, basal hyphae with thickened walls, about 5-6 μ m wide..... H. molle
1. Cystidia absent, basal hyphae thin-walled up to 4-5 μ m wide..... 2
2. Spores 5-6 (-7) μ m long..... H. cremeo-isabellinum
2. Spores 3.5-4.5 (-5) μ m long..... 3
3. Spores unchanged in Melzer's reagent. Known only from deciduous wood..... H. ovoideum
3. Spores greyish in Melzer's reagent. Known only from coniferous wood..... H. subillaqueatum

HYPOCHNICIELLUM Hjortst. & Ryv., Mycotaxon 12(1): 176, 1980.

Type species: Leptosporomyces ovoideus Jülich

Emendation.

Fruitbody resupinate, effuse, pellicular to membranaceous, with a whitish, fairly well-developed subiculum, consisting of loosely interwoven hyphae; hymenium smooth or with small granules; margin inconspicuous or in some species fibrillose to rhizomorphic.

Hyphal system monomitic. Basal hyphae thin- or more or less thick-walled, subhymenial hyphae thin-walled, all hyphae with clamps.

Cystidia present or absent, somewhat hypha-like, with thin- or thickened walls.

Spores thick-walled, cyanophilous, in some species with walls greyish in Melzer's reagent.

HYPOCHNICIELLUM CREMEOISABELLINUM (Litsch.) Hjortst. nov. comb. Basionym: Corticium cremeoisabellinum Litsch., Ann. mycol. 39(2-3): 117, 1941.

Lectotypus: Sweden. Södermanland. Near Saltsjöbaden, on *Pinus sylvestris*. 1906-11-14. L. Romell (5). !

This is a little-known species, but during recent years a few collections have been found in Norway. One of these specimens (Hjm 10530), differs in having spores greyish in Melzer's reagent and in having slightly encrusted hyphae. It is possible that the species consists of more than one taxon but this cannot at present be satisfactorily solved.

Selected specimens studied: Sweden. Småland. Rumsquilla par., Norra Kvill Nat. Park, on coniferous wood. 1970-10. Gro Gulden s.n. (GB and O). Norway. Oppland. Dovre. Hjerkinholen, elev. 950 m, on *Pinus sylvestris*. 1979-08-25. Hjortstam 10497; do. Hjm 10530 (both in Hjm priv. herb.).

HYPOCHNICIELLUM MOLLE (Fr.) Hjortst. nov. comb.

Basionym: Thelephora mollis Fr., Syst. mycol. I, pag. 443, 1821.

Authentic material not located.

The modern interpretation of the species corresponds well with the protologue by Fries which reads as follows: effuse, carnosomembranacea, rubescenti-pallida, subtus tomentosa, papillis prominulis majusculis.

On well-developed specimens the small granules (which were observed by Fries) on the hymenium are easily distinguishable, at least under a lens (50 X), and together with the cystidia, spores, and wider basal hyphae with thickened walls, delimit the species well from others of the genus.

Selected specimens studied: Sweden. Västergötland. Bergstena par., Korpås, on decayed timber (coniferous wood). 1977-08-07. Hjortstam 8141. Norway. Akershus. Nannestad. Tømte farm, on coniferous wood. 1978-09-27. Hermansen & Hjortstam s.n. (both in Hjm priv. herb.). Hedmark. Grue. Meldalen, on *Picea abies*. 1974-10-12. L. Ryvarden 13352 (0).

HYPOCHNICIELLUM SUBILLAQUEATUM (Litsch.) Hjortst., nov. comb. Basionym: Corticium subillaqueatum Litsch., Ann. mycol. 39 (2-3): 128-129, 1941.

Lectotypus: Sweden. Stockholm. Lidingö, on *Pinus sylvestris*. 1910-05-29. L. Romell 2068 (S). !

For a good description and comprehensive discussion of the species see Eriksson and Ryvarden 1976.

Selected specimen studied: Sweden. Uppland. Ärentuna par., Störvreta, on decayed coniferous fencing. 1927-05-08. Seth Lundell No. 53 (GB).

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CLADOSPORIUM BANTIANUM AND ITS SYNONYM CLADOSPORIUM TRICHOIDES

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Caracas, Venezuela

SUMMARY

Examination of the type material for *Cladosporium bantianum* (Sacc.) Borelli and the living type culture of *C. trichoides* Emmons, as well as a number of other living cultures, has revealed that these two taxa are conspecific. Based upon priority, *C. bantianum* is the correct name for this taxon.

INTRODUCTION

During the past several years, there has been considerable controversy regarding *Cladosporium bantianum* (Saccardo) Borelli, 1960 and its relationship to *C. trichoides* Emmons, 1952. Borelli (4) studied the type material for *Torula bantiana* Saccardo, 1912, which was isolated from a human case of cerebral phaeohyphomycosis by Banti (1), and concluded that *T. bantiana* should be transferred to the genus *Cladosporium*, for which, he proposed the new combination *C. bantianum*. Borelli also considered *C. trichoides*, which was likewise isolated from a case of human cerebral phaeohyphomycosis (3), to be a synonym of *C. bantianum*. Emmons *et al.* (5) have disagreed with Borelli. They consider *C. trichoides* to be a distinct species and *Torula bantiana* a *nomen confusum*. This study was undertaken to reevaluate these two taxa and to

resolve the controversy surrounding *C. bantianum* and *C. trichoides*.

METHODS AND MATERIALS

Living cultures were studied on potato dextrose agar (PDA) after 2 weeks incubation at 25°C in the dark. Measurements were determined from camera lucida drawings of slide culture preparations.

Herbarium specimens examined. Exsiccatum labelled 499, *Torula bantiana* Sacc., Herbarium Mycol. Orto Botanico, Padova, Italy, P. A. Saccardo; 5 photomicrographs of *T. bantiana* prepared by P. A. Saccardo; DMD-38, slide culture preparation grown on PDA, isolated from sawdust mulch heap via spleen tissue of hamster, Williamsburg, VA, prepared by D. Dixon in 1977, received as *C. trichoides* (see NCMH 1145 for living culture); DMD-39, slide culture preparation grown on PDA, isolated from sawdust mulch heap via testes of hamster, Williamsburg, VA, prepared by D. Dixon in 1977, received as *C. trichoides* (see NCMH 1148 for living culture).

Living cultures of *C. bantianum* studied. NCMH 111 = CDC B-1937 = NIH 8595 = IP 509, from human brain abscess, sent to NIH by G. Segretain, Institut Pasteur, Paris; NCMH 112 = CDC B-1938 = NIH 8580, from human brain abscess, T. Collette, Sayre, PA.; NCMH 113 = CDC B-1940 = NIH 8579 = ATCC 10958 = CBS 173.52, type culture, from human brain abscess, C. Binford, Baltimore, MD; NCMH 114 = CDC B-1941 = NIH 8598, from soil, P. Klite *et al.*, Panama; NCMH 115 = CDC B-2003 = NIH 8504, from chronic abscess in human abdominal wall, J. Pereira, Washington, DC; NCMH 116 = NCMH 121 = CDC B-2283 = CDC B-1897, from brain abscess in a cat, S. Jang *et al.*, Davis, CA; NCMH 117 = CDC A-980 = NIH 8590, from human brain abscess, J. Barnola Duxans, Caracas, Venezuela; NCMH 121 = NCMH 116 = CDC B-1897, from brain abscess in a cat, M. Rinaldi, Davis, CA; NCMH 122 = CDC B-1898 = ATCC 22649, from human brain abscess, J. Bennett, FL; NCMH 474 = NCMH 767 = CDC B-2525, from brain abscess in a cat, Biberstein, Davis, CA; NCMH 767 = NCMH 474 = CDC B-2525, from brain abscess in a cat, S. Jang *et al.*, Davis, CA; NCMH 1145 = DMD-38, from sawdust mulch heap, D. Dixon, Williamsburg, VA; NCMH 1146 = DMD-41, from sawdust mulch heap, D. Dixon, Williamsburg, VA; NCMH 1147 = DMD-58, from stump of *Juniperus virginiana*, D. Dixon, Suffolk, VA; NCMH 1148 = DMD-39, from sawdust mulch heap, D. Dixon, Williams-

Table 1. Measurements of *Cladosporium bantianum* conidia in herbarium specimens

Specimen	Average size (μm)	Range (μm)
Exsiccatum labelled 499	4.6 X 8.4	3.1-7.3 X 5.6-13.0
DMD-38	3.1 X 6.9	2.2-3.9 X 5.7-9.2
DMD-39	3.4 X 7.5	2.7-4.1 X 5.1-14.0

Table 2. Measurements of *Cladosporium bantianum* conidia after 2 weeks on potato dextrose agar at 25°C

Isolate	Average size (μm)	Range (μm)
NCMH 111	2.5 X 6.4	1.8-3.2 X 4.5-10.0
NCMH 112	2.7 X 7.2	2.1-3.6 X 4.8-10.5
NCMH 113	3.0 X 6.3	2.3-4.0 X 4.7-11.0
NCMH 114	2.7 X 6.2	2.0-3.4 X 5.0-9.2
NCMH 115	2.8 X 7.2	2.0-3.8 X 5.0-13.8
NCMH 116	2.9 X 5.7	2.3-3.5 X 4.6-8.3
NCMH 117	2.7 X 6.4	2.0-3.5 X 4.6-10.6
NCMH 121	3.0 X 5.9	2.1-6.5 X 4.0-8.5
NCMH 122	2.7 X 5.5	2.1-3.3 X 4.0-8.0
NCMH 474	2.7 X 5.3	2.1-3.1 X 3.8-8.5
NCMH 767	3.1 X 6.5	2.1-4.0 X 4.1-12.0
NCMH 1145	2.9 X 5.8	2.5-3.5 X 4.1-9.0
NCMH 1146	3.0 X 5.3	2.3-3.7 X 3.7-9.0
NCMH 1147	2.9 X 6.3	2.1-3.5 X 5.0-8.5
NCMH 1148	3.1 X 6.2	2.0-4.4 X 4.0-11.8
NCMH 1151	3.1 X 5.9	2.5-4.1 X 3.8-9.0
NCMH 1152	2.8 X 6.6	2.1-3.2 X 4.8-11.0
NCMH 1168	2.6 X 5.5	2.1-3.1 X 3.8-11.5
NCMH 1181	2.8 X 5.8	2.1-3.7 X 4.2-8.5
NCMH 1182	3.0 X 5.6	2.5-3.5 X 4.0-7.6
NCMH 1186	3.2 X 8.0	2.5-4.0 X 5.0-14.6

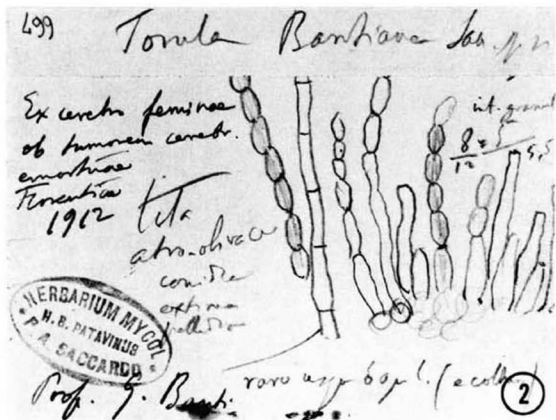
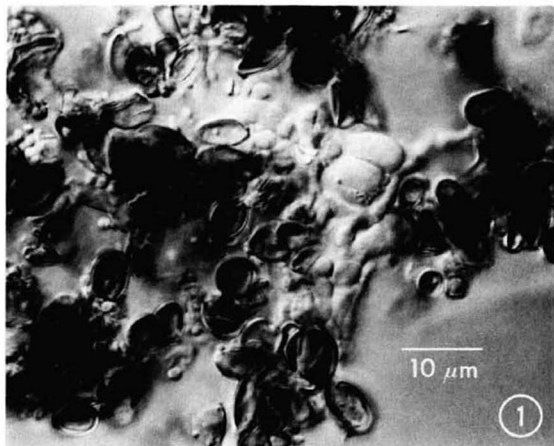
burg, VA; NCMH 1151 = ATCC 24928, from human brain abscess, P. F. Jurgensen; NCMH 1152 = ATCC 28255 = Crichlow 853, from human brain abscess, D. K. Crichlow; NCMH 1168 = CBS 328.65, from a dog, G. A. de Vries; NCMH 1181 = SM 1502, from human brain abscess, M. Hironaga, Otsu, Japan; NCMH 1182 = SM 1522, from human cutaneous abscess, M. Hironaga, Otsu, Japan; NCMH 1186 = FCM 7819, from human skin lesion, S. Amma, North Kerala, India.

DISCUSSION

Measurements of the conidia preserved in the holotype (Fig. 1) for *C. bantianum* are $4.6 \times 8.4 \mu\text{m}$ ($3.1\text{--}7.3 \times 5.6\text{--}13.0 \mu\text{m}$) (Table 1), which vary a little from the measurements included in the original description (6) for the species, and those measurements recorded by Saccardo in his notes (Fig. 2). Saccardo described the conidia as "...conidiis longe et sat persistenter catenulatis (in quaque catenula 5-10), concoloribus, oblongo-ellipsoideis, 8-11 X 5, levibus, intus granulosis, summis junioribus et pallidioribus." Even though Saccardo makes no statement concerning whether or not the chains were simple or branched, his photomicrograph of *C. bantianum* (Fig. 3) clearly shows that the chains were branched. Figure 3 also reveals that some of the chains consisted of more than ten conidia. The exsiccatum contains a few blastoconidia that have three hila, which confirms that the chains were branched.

Emmons *et al.* (5) have interpreted the original description of *C. bantianum* as follows: "Saccardo stated that *T. bantiana* produced conidia 8 to 11 X 5μ in unbranched chains which were only 5 to 10 cells in length". Their translation of the Latin diagnosis for *C. bantianum* is apparently in error since Saccardo considered only the conidiophores to be simple. He described them as "...conidiophoris ex mycelio hypostromatico brevi, celluloso ascendentibus, fasciculatis, cylindraceutis, simplicibus, subrectis, variae longitudinis, plerumque 15-30 μ

Figs. 1-2. *Cladosporium bantianum*. 1. Lactophenol preparation of exsiccatum labelled 499, *Torula bantiana* Sac., Herbarium Mycol. Orto Botanico, Padova, Italy, P. A. Saccardo, in differential interference contrast microscopy. 2. Notes prepared by P. A. Saccardo for *Torula bantiana* Sacc.

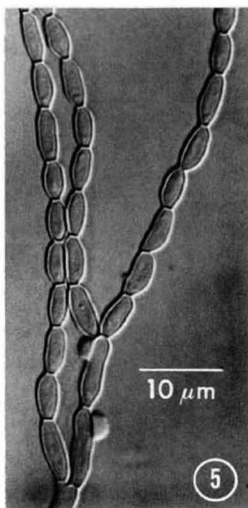
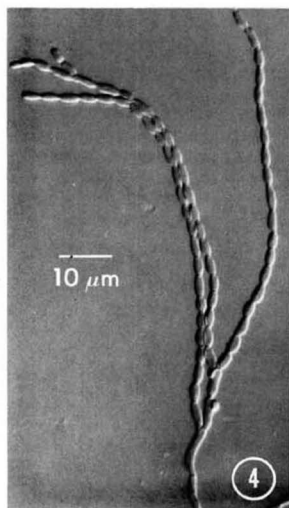


long, raro usque ad 60 μ (ex cultura)...". Owing to their incorrect translation of the nature of the conidial chains for *C. bantianum*, they stated: "If the name, *T. bantiana*, is based on both Saccardo's descriptions and Borelli's interpretation of a photomicrograph attributed to Saccardo, then the name, *T. bantiana*, apparently is based on two fungi and is a *nomen confusum*." Figure 3 is the photomicrograph published by Borelli in 1960 (4) that is referred to by Emmons *et al.* It clearly shows that the chains were branched and that only one fungus was involved.

It has been argued that *C. bantianum* and *C. trichoides* should be maintained as separate species because of differences reported in the number of conidia composing the chains (5). Emmons in Binford *et al.* (3) described the conidia of *C. trichoides*, in part, as "Conidiis ellipsoidis vel cylindraceis, 2-2.5 X 4-10 μ , continuis vel 1-septatis, fuscis, aequalibus, formatis in cateniis longis ramosisque." No mention was made by Emmons of the length of the conidial chains other than that they were long, but his figure 5 shows the longest chain to possess six conidia. Examination of NCMH 113 (Figs. 4-5), and additional cultures have revealed that the chains are much longer, often consisting of 35 or more conidia. Even though Saccardo stated that the conidial chains of *T. bantiana* consisted of 5-10 conidia, his photomicrograph (Fig. 3) shows that some of the chains were longer.

Emmons *et al.* (5) have observed that isolates of *C. bantianum* ... "have been remarkably uniform in morphology and size of spores, although a few strains have borne larger spores in primary cultures than those described by Emmons, reverting to spores of typical dimensions upon subculture." A comparison of DMD-38 and DMD-39 (Table 1) with NCMH 1145 and NCMH 1148, respectively (Table 2), supports the observation that the conidia tend to become smaller in size upon subculture. In a personal communication to Dr. Borelli, J. Bennett (National Institutes of Health, Bethesda, MD) studied a brain isolate that he

Figs. 3-5. *Cladosporium bantianum*. 3. Photomicrograph prepared by P. A. Saccardo of *Torula bantiana* showing its microscopic morphology. 4-5. Branched chains of blastoconidia (NCMH 113), potato dextrose agar after 2 weeks at 25°C, in differential interference contrast microscopy.



identified as *C. trichoides*. Bennett's isolate produced conidia that were approximately the same size as those described for *C. bantianum* by Saccardo. Unfortunately, Bennett's isolate could not be included in this study because it had died several years ago. After examining the living cultures listed in Table 2, it is obvious that the conidia of *C. bantianum* vary in size, ranging from 1.8-6.5 X 3.7-14.6 μ m. The size range seen in the exsiccatum is 3.1-7.3 X 5.6-13.0 μ m, which is compatible with the variation observed in the living isolates. This similarity in conidial size clearly supports the conclusion that *C. bantianum* and *C. trichoides* are conspecific, even though the conidia in the exsiccatum tend to be larger than those in the living cultures.

There is no doubt that Banti's isolate had the ability to grow at 37°C (6). In fact, Banti found that his fungus was pathogenic for rabbits (7), whose body temperature is approximately 40°C. Since there is no living culture of Banti's isolate, it is impossible to determine whether it could grow at 42-43°C. Borelli (4) demonstrated that isolates of *C. bantianum* can grow at this high temperature. The thermotolerance and neurotropism in man of the original isolates of *C. bantianum* and *C. trichoides* add additional support to the conspecificity of these two taxa.

Another controversial point is the reported color of the human brain lesions. According to Banti's description (1), the cerebral nodules were ... "of a deep brown color, giving one the impression of a melanotic sarcoma." According to Emmons *et al.* (5), the lesions caused by *C. trichoides* occur as gray abscesses. Judging by what is published in the literature, this aspect of the disease is variable. For example, in a Venezuelan case (2), the lesions were described as gray-blackish. In the necropsy protocol of Banti's case, which was reproduced by Stigliani (7), the lesions were described as a ... "dirty gray... from which a thick, mucous thread-like, grayish-greenish liquid came out...". The latter text was most likely dictated by Banti during the performance of the necropsy. Since the color of the lesions could have darkened within a few hours, this may have resulted in the different descriptions of the lesion color. The contention by Emmons *et al.* (5) that the pathological aspects of the two infections are significantly different enough to maintain *C. bantianum* and *C. trichoides* as separate taxa is unjustifiable.

It is apparent that *C. trichoides* and *C. bantianum* are conspecific. The arguments put forth by Emmons *et al.* for maintaining these as separate taxa are not compelling.

Cladosporium bantianum can be characterized as growing at temperatures up to 42-43°C and by producing long, sparsely branched chains of blastoconidia consisting of as many as 35 or more conidia. The conidia are smooth, 1-celled (rarely 2-celled), oval to oblong-ellipsoidal, 3.0 X 6.4µm (1.8-7.3 X 3.7-14.6µm), and pale brown. The conidia arise from hyphae or simple septate conidiophores that are pale brown in color. The nomenclature is:

Cladosporium bantianum (Sacc.) Borelli, Riv. Anat. Patol. Oncol. 17:618, 1960.

≡ *Torula bantiana* Sacc., Ann. Mycol. 10:320, 1912.
(basionym)

= *Cladosporium trichoides* Emmons in Am. J. Clin. Pathol. 22:540-541, 1952.

ACKNOWLEDGEMENTS

The authors wish to thank Dr. Donald P. Rogers for reviewing our manuscript, Professor Sergio Chiesa for loan of the herbarium material maintained in the Saccardoan Herbarium, and Dr. Dennis Dixon for loan of his slide culture preparations and living cultures.

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A PRELIMINARY DISCOMYCETE FLORA OF MACARONESIA: PART 2, HYALOSCYPHACEAE SUBF. ARACHNOPEZIZOIDEAE*

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"Much like a subtle spider which doth sit
In the middle of her web, which spreadeth wide."

Sir John Davies

THE IMMORTALITY OF THE SOUL

Order HELOTIALES

Suborder HYMENOSCYPHINEAE

Family HYALOSCYPHACEAE Nannf. 1932

Subfamily ARACHNOPEZIZOIDEAE Korf 1978

Tribe Arachnopezizeae Nannf. in Korf emend. Korf 1978

ONE KNOWN MACARONESIAN GENUS

ARACHNOPEZIZA Fuckel 1870 emend. Korf 1952

Key to the known Macaronesian species

- 1. Ascospores 5-7-septate at maturity. 3. *A. aurata*
- 1'. Ascospores 0-1-3-septate at maturity2
 - 2(1'). Ascospores 0-1(-2)-septate at maturity;
apothecia on husks of chestnut (Casta-
nea) burrs3
 - 2'(1'). Ascospores mostly 3-septate at maturity;
apothecia on wood, leaves, duff, acorns
.....4
- 3(2). Hairs thin-walled, tapering, usually 1-septate,

* This flora will appear in an irregular order, with Part 1, containing the introductory material, one of the last to be published. Reprints of individual parts will not be available. References will appear in the final part. For a geographical definition of *Macaronesia* and other special features of this flora, see the note on page 144.

not constricted at the septum.

- 3'(2). Hairs thicker-walled, multiseptate, constricted the septa.
- 4(2'). Apothecia 1-3 mm diam., ascospores 10.2-13.2(-16.1) x 2.2-3.7 μ m.
- 4'(2'). Apothecia less than 0.5 mm diam.5
- 5(4'). Apical cells of hairs without cyanophilic contents; ascospores (13.5-)18.3-22(-26) x 2.9-3.7 μ m; paraphysis apices variously deformed.
- 5'(4'). Apical cells of hairs with cyanophilic contents; ascospores (14.6-)16.1-25.7 x 2.2-4.4 μ m; paraphysis apices filiform, not deformed.

1. *A. aranea* f. *aranea*

2. *A. aranea* f. *monilipila*

4. *A. aurelia*

5. *A. obtusipila*

6. *A. zonulata*

1. *Arachnopeziza aranea* (De Not.) Boud., *Icones Mycol. Expl. Pl.*, sér. 3: 2. 1906 forma *aranea*.
- = *Lachnum aranea* (De Not.) Lindau, in Engler & Prantl, *Nat. Pflanzenfam.* I 1 (130): 203. 1896.
- = *Arachnoscypha aranea* (De Not.) Boud. ex Dennis, *Mycol. Pap.* 32: 87. 1949.

RECENT TAXONOMIC TREATMENTS: Dennis (1949, 1978), Korf (1952).

PREVIOUS MACARONESIAN RECORDS: None.

TYPE LOCALITY: Italy.



KNOWN MACARONESIAN DISTRIBUTION
MADEIRA.

Madeira. CUP-MM 1542.

CANARY ISLANDS.

La Palma. CUP-MM 870.

A. aranea f. *aranea*, 5 ascospores, CUP-MM 1542, x 1000.

SUBSTRATA: On cupules (burrs) of *Castanea sativa*.

Notes: I saw one 2-septate ascospore in my mounts (illustrated), but typically the spores are either continuous or 1-septate. Probably common wherever chestnut grows, but difficult to find among the hairs on the in-

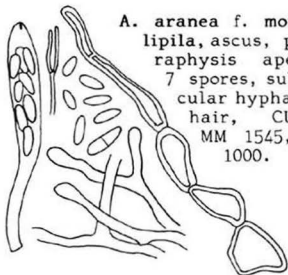
ner surface of the spiny husks. I cannot agree with Dennis, who separated this species in a distinct genus, *Arachnoscypha* Boud., of which it is the type species, but which Boudier himself later abandoned. In my earlier monograph (Korf, 1952), I missed Lindau's transfer to *Lachnum*, which should be added to the synonymy given there.

2. *Arachnopeziza aranea* (De Not.) Boud. f. **MONILIPILA**
Korf, f. nov.

PREVIOUS MACARONESIAN RECORDS: None.

A. aranea aranea
f. *aranea* pilis multiseptatis miniliformibus dif-
fert.

Differing from the type form in having multi-septate rather than 1-septate hairs, and by the constrictions at the septa, giving the moniliform aspect to the hairs.



A. aranea f. *moni-*
lipila, ascus, pa-
raphysis apex,
7 spores, subi-
cular hyphae,
hair, CUP-
MM 1545, x
1000.

HOLOTYPE: R.P. Korf, R. Fogel, G.L. Hennebert & L.M. Kohn, on cupules of *Castanea sativa*, *Castanea* grove below Pousada Vinháticos, Madeira, Portugal, 15.i.1977.

KNOWN MACARONESIAN DISTRIBUTION

MADERIA.

Madeira. CUP-MM 1545 (holotype).

SUBSTRATA: On spiny burrs of *Castanea sativa*.

Notes: The asci arise from repeating croziers, and the ectal excipular cells are quite large. The new form may represent merely a growth-stage of the species.

3. *Arachnopeziza aurata* Fuckel, Jahrb. Nassauischen
Vereins Naturk. 23-24: 304. 1870.

RECENT TAXONOMIC TREATMENTS: Dennis (1949, 1978), Korf (1952).

PREVIOUS MACARONESIAN RECORDS:

*Dennis & al. (1977).

TYPE LOCALITY: Germany.

KNOWN MACARONESIAN DISTRIBUTION

*AZORES.

*Terceira. *Dennis & al., 7.
iv.75 n.v.

CANARY ISLANDS.

Gomera. CUP-MM 1348, 1368(TFC).

Hierro. CUP-MM 1446(TFC), 1450.

La Palma. CUP-MM 898.

Tenerife. CUP-MM 223, 230, 246,
256, 298(TFC), 435(TFC),
508(TFC), 574, 1281.

A. aurata, 2
ascospores,
CUP-MM 1145,
x 1000.

SUBSTRATA: On wood and trunk of Erica arborea, on bark of Myrica faya, wood of Castanea sativa, and undetermined decorticated wood and bark.

Notes. This is surely the most common species of the genus in Macaronesia, as it tends also to be in North America and in Europe.

4. *Arachnopeziza aurelia* (Pers. : Fr.) Fuckel, Jahrb. Nassauischen Vereins Naturk. 23-24: 303. 1870.

RECENT TAXONOMIC TREATMENTS: Dennis (1949, 1978), Korf (1952).

PREVIOUS MACARONESIAN RECORDS:
None.

TYPE LOCALITY: France.

KNOWN MACARONESIAN
DISTRIBUTION

CANARY ISLANDS.

Hierro. CUP-MM 1459 (OSC,
TFC).

A. aurelia, 7 as-
cospores, CUP-MM
1459, x 1000.

SUBSTRATA: On fruits and husks of Castanea sativa and other vegetable debris.

Notes: In North America and Europe this is most common on cupules and leaves of Quercus.

5. *Arachnopeziza obtusipila* Grelet, Amateur Champignons 8(3): 45. 1922, emend. Korf, Mycologia 43: 213. 1951.

RECENT TAXONOMIC TREATMENTS:
Korf (1951, 1952).

PREVIOUS MACARONESIAN RECORDS:
*Baagøe & al. (1972) as "inoperculat discomycet."

TYPE LOCALITY: France.

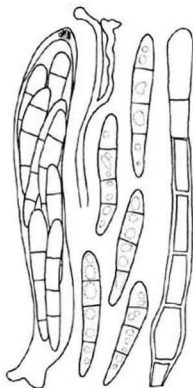
KNOWN MACARONESIAN
DISTRIBUTION

*MADEIRA.

*Madeira. *CUP-MM 2502(C).

SUBSTRATA: On decorticated wood
of Pinus pinaster.

Notes: I am indebted to the students of the University of Copenhagen who collected this (Baagøe & al., 1972) and for calling the specimen to my attention for identification.



A. obtusipila, ascus, paraphysis, 5 ascospores, hair, CUP-MM 2502, x 1000.

6. *Arachnopeziza zonulata* (Rolland) Boud., Hist. classific. discomyc. Europe p. 126. 1907.

RECENT TAXONOMIC TREATMENTS: Korf (1952).

PREVIOUS MACARONESIAN RECORDS: None.

TYPE LOCALITY: Corsica (France).

KNOWN MACARONESIAN DISTRIBUTION

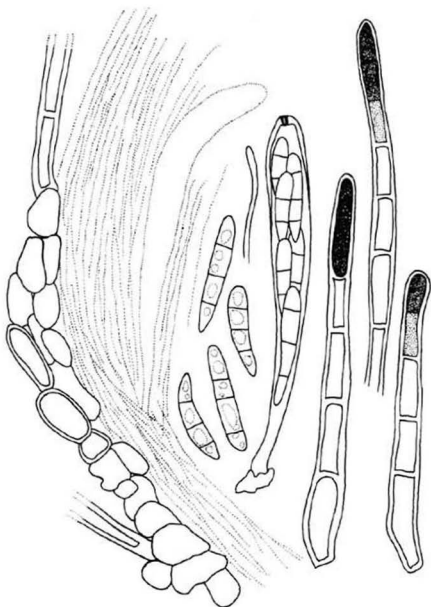
CANARY ISLANDS.

Tenerife. CUP-MM 1222.

SUBSTRATA: On wood of Pinus canariensis.

Notes: In the absence of type material, I (Korf, 1952) treated this as a doubtful species. If I am correct in interpreting this as Rolland's species, it is only criti-

cally distinct from *A. obtusipila*. Malençon and Bertault (1958) proposed varietal status for Grelet's species under *A. zonulata*, but their combination was not validly published. I have made 3 trips to Rolland's type locality near Corte, in Corsica, but did not find any species matching his description.



A. zonulata, section through the apothecial margin with 2 hair bases, 4 ascospores, paraphysis apex, ascus with J+ pore, 3 hairs in cotton-blue, CUP-MM 1222, x 1000.

Tribe Polydesmieae Korf 1978

ONE KNOWN MACARONESIAN GENUS

POLYDESMIA Boudier, Bull. Soc. Mycol. France 1: 113. 1885.

Key to the known Macaronesian species

1. Ascospores 1-3-septate at maturity, $9.0-13.5 \times 2.2-2.7 \mu\text{m}$. 1. *P. fructicola*
 1'. Ascospores 3-septate at maturity, $(13.9-22.0) \times 3.7-5.1 \mu\text{m}$. 2. *P. pruinosa*

1. *Polydesmia fructicola* Korf, Mycotaxon 7: 475. 1978.

RECENT TAXONOMIC

TREATMENTS: Korf (1978).

PREVIOUS MACARONESIAN RECORDS: *Korf (1978), †Beltrán Tejera (1980).

TYPE LOCALITY: Madeira.

KNOWN MACARONESIAN DISTRIBUTION

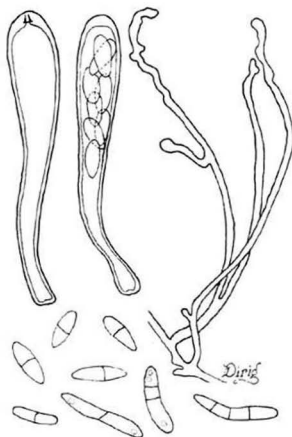
*MADEIRA.

*Madeira. *CUP-MM 1504 (holotype) (TFC), *1533, *1551, *1556, *1618, *1619, *2333, *2361, *2362, 2366, *2391.

*CANARY ISLANDS.

*Tenerife. *CUP-MM 1306.

SUBSTRATA: On peduncles, pods, twigs and wood of *Acacia* spp., bark and capsules of *Eucalyptus* spp., undetermined bark and branchlet.



P. fructicola, ascus with J+ pore, ascus with spores, paraphyses, 4 upper ascospores, CUP-MM 1504; 4 lower spores CUP-MM 2361; all $\times 1000$.

2. *Polydesmia pruinosa* (Jerd. in Berk. & Br.) Boudier,
Bull. Soc. Mycol. France 1: 113. 1885.

RECENT TAXONOMIC TREATMENTS:
Dennis (1978), Korf (1978).

PREVIOUS MACARONESIAN RE-
CORDS: *Korf (1978), †Beltran
Tejera (1980).

KNOWN MACARONESIAN
DISTRIBUTION

*MADEIRA.

*Madeira. *CUP-MM 1481(TFC),
*1627, *2277, *2345, 2406.

*CANARY ISLANDS.

*La Palma. *CUP-MM 639
(TFC), *709.

SUBSTRATA: On pyrenomycetes,
on pyrenomycetes immersed in
wood, on branch of Acacia sp.

SPECIAL FEATURES OF THIS FLORA

Since Part 1, the introductory material, is
to appear later, one reviewer of this part
confessed he did not know where Macarone-
sia is, and that he could not find it in
any of the atlases he consulted! Thus:

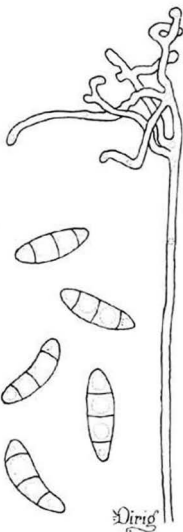
Macaronesia: the Atlantic island chain
comprising the archipelagos of the Azores,
Madeira, Salvage Islands, Canary Islands,
and Cape Verde Islands. Etymology: from
Greek, μάκαρος (fortunate) + νῆσος (is-
land): "Les Îles Fortunes," "The Fortunate
Islands." (Want to guess how often these
papers will be cited as **Macronesia** instead
of **Macaronesia**?)

Substrates follow O. Eriksson, A. Hansen
& P. Sunding. 1974. Flora of Macaronesia.
Check-list of vascular plants, 1974. Uni-
versity of Umeå, Sweden. iv. + 66 pp.

Distributional symbols: Papers with previous Macaronesian records citing spec-
imen data are indicated by one or more asterisks (* ** ***) and the cor-
responding specimens examined have been coded to those asterisks with their
CUP-MM numbers; where such specimens cannot be located or were not avail-
able on loan, the symbol "n.v." follows brief specimen data. Papers citing
taxa without giving specimen data, or merely repeating previous records,
are indicated by a dagger or daggers († † †).

Herbarium abbreviations are those of the Index Herbariorum, ed. 6, and up-
dates appearing periodically in the journal Taxon.

Without the generous financial assistance of the **National Science Foundation**
(Grant DEB75-23557) this preliminary flora would never have been undertaken.



P. pruinosa, 4 asco-
spores, paraphysis
apex, CUP-MM 1481,
x 1000.

MYCOTAXON

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April-June 1981

A PRELIMINARY DISCOMYCETE FLORA OF MACARONESIA: PART 3, HYALOSCYPHACEAE SUBF. TRICHOSCYPHELLOIDEAE*

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"A Hair perhaps divides the False and True."

Omar Khayyám [tr. Edward Fitzgerald]

RUBÁIYÁT, Stanza 19

Order HELOTIALES

Suborder HYMENOSCYPHINEAE

Family HYALOSCYPHACEAE Nannf. 1932

Subfamily TRICHOSCYPHELLOIDEAE Nannf. 1932

ONE KNOWN MACARONESIAN GENUS

LACHNELLULA Karsten 1884 emend. Dennis 1962

Key to the known Macaronesian species

1. Apothecia white, with a bright orange hymenium, on living trees and cut ends of stumps of Pittosporum spp. 1. *L. pittospori* subsp. *azorica*
- 1'. Apothecia greenish-glaucus or grey to nearly black 2
- 2(1'). Apothecia grey to black. 2. *L. pulveracea*
- 2'(1'). Apothecia greenish-glaucus, strongly plicate. 3. *L. viridi-glauca*

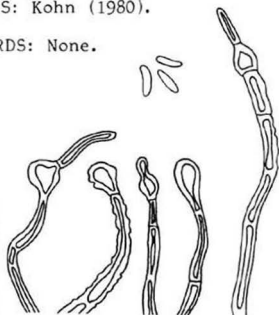
* The parts of this flora, under the editorship of Richard P. Korf, will appear in irregular order. Reprints of individual parts will not be available for distribution.

1. *Lachnellula pittospori* Kohn, Mycotaxon 12: 278. 1980, subsp. *AZORICA* Kohn, subsp. nov.

RECENT TAXONOMIC TREATMENTS: Kohn (1980).

PREVIOUS MACARONESIAN RECORDS: None.

Cum subspecie typica *Lachnellulae pittospori* omnia conveniens sed pilos erectos apice tumidos (in subspecie typica non praesentes) ad marginem praebens. Hi pili ex cellula apicali saepe pariunt processum filiformem aliquando tumescentum et cellulam alteram supra eam primam formantem. Huius subspeciei novae distributio jam connota ad Insulas Azoricas limitata est. Holotypus: CUP-MM 2140.



L. pittospori subsp. *azorica*, 5 marginal hairs, 3 ascospores, CUP-MM 2140, x 1000.

Agreeing with the description of *L. pittospori* subsp. *pittospori* in all respects except for the presence of erect hairs with swollen apices at the margin, not present in subsp. *pittospori*. These hairs often produce from the apical cell a filiform process, which in some cases swells to form a second swollen cell above the first. The distribution of the new subspecies is so far as known limited to the Azores Islands. The type subspecies was described from Bermuda.

HOLOTYPE: R.P. Korf, L.M. Kohn, N. Korf & A.Y. Rossman, on stump of *Pittosporum* sp., Ribeira da Cruz, Lombas north of Caveira, Flores, Azores, Portugal, 13. iv.1978. (CUP-MM 2140.)

KNOWN MACARONESIAN DISTRIBUTION

AZORES.

Flores. CUP-MM 2138 (TFC), 2140 (holotype).
Terceira. CUP-MM 2009 (TFC).

SUBSTRATA: On *Pittosporum* spp., (stumps, cut areas of living trees, and rotted wood of standing trees).

2. *Lachnellula pulveracea* (Alb. & Schw. : Fr.) Dennis, *Persoonia* 2: 184. 1962.

RECENT TAXONOMIC TREATMENTS: Dennis (1949), Höhnelt (1917).

PREVIOUS MACARONESIAN RECORDS: *Dennis & al. (1977).

TYPE LOCALITY: Germany.

KNOWN MACARONESIAN DISTRIBUTION

*AZORES.

***Terceira.** *CUP-MM 1681(K), 2011, 2382, *Dennis & al. 3.iv.75 n.v.

MADEIRA.

Madeira. CUP-MM 1560, 1621, 1623.

CANARY ISLANDS.

Gomera. CUP-MM 1343, 1354(TFC).

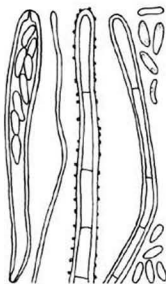
Hierro. CUP-MM 1424, 1444(TFC), 1477.

La Palma. CUP-MM 825.

Tenerife. CUP-MM 87, 248, 538, 1212(TFC).

SUBSTRATA: On twig of *Acacia* sp., of *Ilex perado* subsp. *platyphylla*, of *Ulex europaeus* and *Ulex* sp., on stem of *Rubus* sp., and on undetermined twigs, bark, wood, cut stumps, branches and roots.

Notes: This distinctive species, collected on various woody substrates, is characterized by apothecia that turn violet in KOH and reddish in iodine, as noted by Dennis (1949) and by Höhnelt (1917: No. 1020). As Höhnelt also noted, often the asci are immature. The apothecial structure is typically *Trichoscyphelloideus*, with a completely gelatinized long-celled ectal excipulum, unlike that in *Hyaloscyphoideae*. In lactic acid-



L. pulveracea, ascus, paraphysis, hair in KOH-phloxine-glycerine, hair in lactic-blue (CUP-MM 87); 5 upper spores (CUP-MM 1477); 5 lower spores (CUP-MM 1354); all x 1000.

cotton-blue the granulations on the hairs are no longer visible. Raitviir (1970) treated it as a doubtful or excluded species of the genus *Lachnellula*, and Dharne (1965) did not recognize it in the genus. Rehm's *Ascomyceten* No. 1580, issued as *Dasyscypha coerulescens* var. *dealbata* Rehm, was examined and found to be conspecific, confirming Dennis's (1949) synonymy.

3. *Lachnellula* VIRIDI-GLAUCA Kohn, sp. nov.

PREVIOUS MACARONESIAN RECORDS: None.

Apothecia gregaria vel fasciculata, viridi-glaucata, sessilata; pilis albis vestita, profunde cupulata, receptaculi parietibus penitus plicatis praesertim in apotheciis immaturis; receptaculum 0.25-0.50 mm in diam.; commutatio xanthochroica in 2% KOH eveniens. Excipulum medullare ex textura intricata hyalina formatum, cellulis 2-3 μ m latis. Excipulum ectale ex textura oblita formantum hyalina



L. viridi-glaucata, hair, ascus and paraphysis, CUP-MM 850, x 1000.

intertexta in gelatina copiosa flava contenta, cellulis 3-6 μ m latis, exterioribus pilos edentibus. Pili hyalini, erecti, septati, grosse granulati, 35-45 x 2 μ m. Asci clavati, 36-45 x 5-6 μ m, ex uncis iteratis enati, 8-spori, apice magnopere incrassato, pori canalis pariete J- cum/sine usu KOH antecedente. Ascospori ellipsoidei vel interdum allantoidi, 2-multi-guttulati, 6-7 x 2-2.5 μ m. Paraphyses filiformes, septatae, ramosae, ascos longitudine aequantes, 1 μ m latae, in gelatina contente. Holotypus: CUP-MM 850.

Apothecia gregarious to fasciculate, greenish-glaucous, sessile, clothed in white hairs, deeply cupulate, walls of receptacle deeply plicate, especially in young apothecia; receptacle 0.25-0.5 mm diam; xanthochroic reaction in 2% KOH. Medullary excipulum of hyaline textura intricata, cells 2-3 μ m wide. Ectal excipulum of hyaline, interwoven textura oblita bound in a copious, yellowish gel, cells 3-6 μ m wide, outer cells giving rise to hairs. Hairs hyaline, erect, septate, coarsely granulate, 35-45 x 2 μ m. Asci clavate, arising from repeating croziers, 8-spored, apex greatly thickened, pore channel wall J- with or without KOH pretreatment, 36-45 x 5-6 μ m. Ascospores ellipsoid or occasionally allantoid, 2-

multiguttulate, 6-7 x 2.0-2.5 μ m. Paraphyses filiform, septate, branched, same length as asci, 1 μ m wide, bound in gel.

HOLOTYPE: R.P. Korf, W.C. Denison, L.M. Kohn & M.A. Sherwood, on wood of ?Castanea sativa Mill., near mine entrance at Km mark 13, road between Buenavista and El Paso, La Palma, Canary Islands, Spain, 18.i.1976. (CUP-MM 850.)

KNOWN MACARONESIAN DISTRIBUTION

CANARY ISLANDS.

La Palma. CUP-MM 850 [holotype] (TFC, isotype).

SUBSTRATA: On wood (of ?Castanea sativa).

Notes: The greenish-glaucus color, the deeply plicate receptacle (giving young apothecia the appearance of a "bundt cake"), and the xanthochroic reaction in 2% KOH makes this species especially distinctive. Nevertheless, the ectal excipulum composed of textura oblita and the granulate hairs show Trichoscyphelloideus affinities. Note that young apothecia often have no asci and bear a copious gel layer above the hymenium.

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RESINOMYCENA GEN. NOV. (AGARICALES), AN ALLY OF HYDROPUS, MYCENA AND BAEOSPORA

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ABSTRACT

A new genus, *Resinomycena*, is proposed to accommodate *Agaricus rhododendri*, the type, *Mycena kalalochensis*, and three new species, *R. brunnescens*, *R. montana* and *R. acadensis*. At present the genus is known with certainty only from North America. The relationships to the genera *Baeospora*, *Hydropus* and *Mycena* are discussed.

Agaricus rhododendri Peck and *Mycena kalalochensis* A.H. Smith along with three previously undescribed taxa form a clearly defined taxon characterized by amyloid spores, pseudoamyloid tramal tissues and a turf-like pileal epicutis composed mainly of resin secreting oleocystidia.

In her monograph of the northeastern *Marasmius* species, Gilliam (1976), excluded *M. resinosus* Peck (= *A. rhododendri*) and suggested that a new genus might need to be proposed for it. A.H. Smith (1947) excluded both *Omphalia rhododendri* (Peck) Sacc. and *M. resinosus* from *Mycena* although at the same time describing *M. kalalochensis* as a new species of *Mycena*. These epithets were not dealt with in *The Agaricales in Modern Taxonomy*, (Singer 1975). In this last publication the species described here clearly key out to the *Tricholomataceae* tribus *Myceneae* and less clearly to the choice between *Hydropus* and *Mycena*. However, the delimitations of both genera exclude the group. Thus the recognition of a distinct genus is a viable option. Alternatively emendation of *Hydropus* or *Mycena*

from the concepts accepted in *The Agaricales in Modern Taxonomy* could be proposed. The emendation of *Basospora* by the inclusion of *Basospora pallida* Singer (1977), a species with close but not crowded lamellae and white carpophores, is cause for considering this last genus as a possible depository for *A. rhododendri* and allies. As detailed at the end of this paper, the erection of a new genus most clearly resolves the problems surrounding the placement of these species. Emendation of existing genera would make them too heterogeneous.

Resinomycena Redhead & Singer, gen. nov.

Habitus omphaliodeus vel marasmiodeus, superficeibus ± resinosis ex oleocystidiis (frequenter intermixtis elementis filamentosis vel dendroideis). Lamellis adnatis vel subdecurrentibus, confertis vel subdistantibus (haud confertissimis), ad aciem oleocystidiis numerosis instructis. Stipe aequalis, subcartilagineo, sicco subcorneo, ± resinaceo. Tramate debiliter vel sat manifeste pseudoamyloideo (partibus subgelatinascentibus exceptis); tramate hymenophorali regulari. Sporis hyalinis, amyloideis, levibus, acyanophilis.

Typus: Agaricus rhododendri Peck.

Basidiomes omphaloid or marasmioid, resinous, covered with oleocystidia intermixed with dendroid hyphae, white or dully pigmented. Lamellae adnate to slightly decurrent, close to subdistant but not crowded, with resinous edges formed by numerous oleocystidia. Stipe equal, cartilaginous but drying to a horny consistency, resinous, covered by scattered or clumped oleocystidia. Tramal tissues weakly to strongly pseudoamyloid where walls not slightly gelatinized, but frequently some inamyloid hyphae intermixed. Hymenophoral (lamellar) trama regular. Spores smooth, amyloid, hyaline, acyanophilic.

Type species: *Agaricus rhododendri* Peck.

A key to *Resinomycena* species

- A. From Europe *Marasmius rhododendri*
(see discussion)
- A. From eastern North America B
- A. From western North America D
- B. Pileus brown; hyphae incrustated with
 brown pigments *R. brunnescens*
- B. Pileus white or whitish; hyphae lacking
 pigmented incrustation C
- C. Lamellae close; spores small,
 5.4-8.5 X 2.4-4.1(-4.5)µm *R. rhododendri*
- C. Lamellae moderately spaced to subdistant;
 spores large, 9.4-12.8 X 4-5µm *R. acadensis*

- D. Pileocystidia mainly capitate, secondarily septate, mostly erect; on angiosperm litter *R. kalalochensis*
- D. Pileocystidia clavate but rarely capitate, rarely if at all secondarily septate and often collapsed to a repent condition; on coniferous litter *R. montana*

Resinomyces rhododendri (Peck) Redhead & Singer, *comb. nov.*

Figs. 1-5.

- ≡ *Agaricus rhododendri* Peck (1875: 94)
- ≡ *Omphalia rhododendri* (Peck) Saccardo (1887: 335)
- ≡ *Omphalopsis rhododendri* (Peck) Murrill (1916: 311)
- = *Marasmius decurrens* Peck (1872: 77) *nom. illeg. non. M. decurrens* Montag. (1854: 118)
- ≡ *Marasmius resinosus* Peck (1883: 181) *nom. nov. for M. decurrens* Pk.
- = *Marasmius resinosus* var. *niveus* Peck (1903: 38)
- ≡ *Marasmius resinosus* var. *candidissimus* Peck (1905: 40) *nom. nov. illeg. for M. resinosus* var. *niveus* Pk.

PILEUS: 4-15(-19) mm wide, convex becoming plano-convex to plane, usually depressed centrally, occasionally umbilicate, rarely subumbonate, white to yellow white, opaque to vaguely striate marginally becoming obscurely corrugated-striate on some and sometimes obscurely concentrically ridged, dry to tacky or slightly viscid, micaceous when dry; edges incurved at first, uneven with age; context tough-pliant, concolorous; odor and taste not distinctive.

LAMELLAE: adnate to subdecurrent or arcuate-decurrent, whitish, moderately narrow, close, often becoming forked in places or developing ladder-like anastomoses; edges crenulate, in some specimens beaded with resin. **STIPE:** 12-50 mm long, 0.5-1 mm wide, equal or slightly enlarged above, tough and pliant, drying to a corneous texture, whitish, varying from glutinous to tacky or dry and glistening from resinous cells in scattered beads, fistulose, with a silky radiating white basal disc or subiculum.

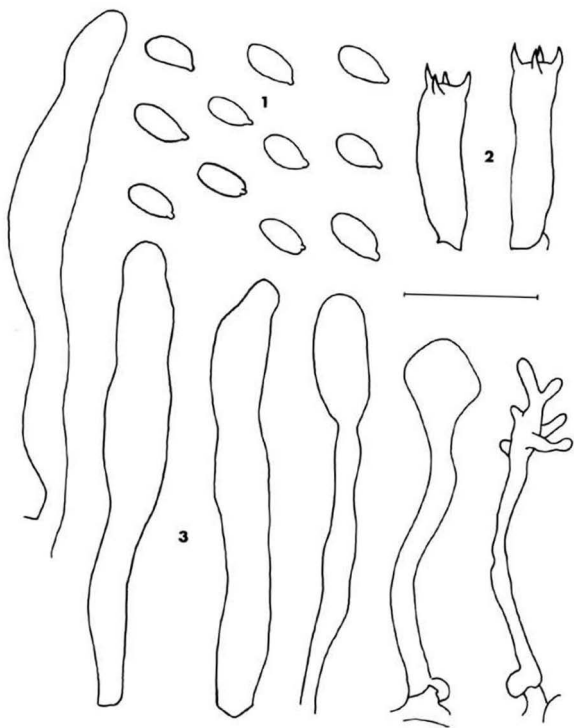
PILEAL EPICUTIS: a layer of polymorphic, suberect, later decumbent and tangled cystidia ranging from cylindrical narrowly clavate or capitate oleocystidia to narrow dendroid nonresinous forms with some intergradation, 40-60 X 3-9.5 μ m, hyaline, thin-walled, often filled or partially filled with vacuolate to homogeneous oily contents and covered with similar exudates which dry as slightly yellowish and hardened masses. **PILEAL TRAMA:** obscurely duplex, with slightly broader hyphae above, 5-15 μ m diam., than below, 3-10 μ m diam.; hyphae subparallel, clamped, smooth, walls thin to slightly thickened, faintly pseudoamyloid. **LAMELLAR TRAMA:** similar to pileal trama, varying to slightly interwoven, more definitely pseudoamyloid. **CHEILOCYSTIDIA:** abundant, forming a sterile edge, similar to the cylindrical to narrowly clavate or fusoid oleopileocystidia, 55-65 X 7-7.5 μ m. **PLEUROCYSTIDIA:** rare or absent, similar to the cheilocystidia. **BASIDIA:** 19-22 X 5-5.2 μ m, narrowly cylindrical to obscurely utriform, 4-spored, clamped. **BASIDIOSPORES:** 5.4-8.5 X 2.4-4.1(-4.5) μ m often being predominantly at one or the other end of the range, mostly ellipsoidal or broadly cylindrical, inequilateral in profile, prominently apiculate, smooth, hyaline, thin-walled, amyloid.

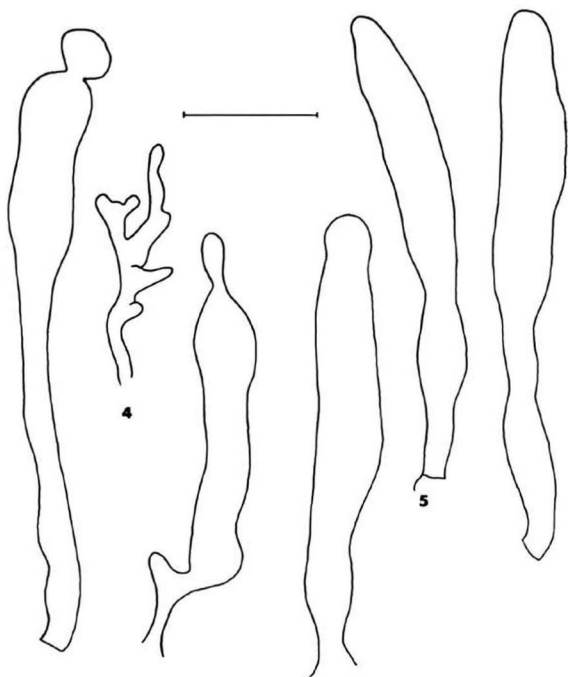
STIPE HYPHAE: parallel, 3-5 μ m diam. in the cortex, up to 10 μ m diam. in the medulla, mostly prominently pseudoamyloid in the medulla, walls uneven and slightly thickened. CAULOCYSTIDIA: similar to the pileocystidia, often clustered. BASAL MYCELIUM: interwoven, 2-3 μ m diam., smooth, inamyloid, clamped, hyaline, walls refractive and slightly thickened.

HABITAT, HABIT and SUBSTRATES: scattered to subcespitate on leaf litter, small twigs or bits of wood and woody fruits or husks from *Rhododendron*, *Quercus*, *Fagus*, *Castanea*, *Carya* and sometimes *Pinus* when mixed with *Quercus* in eastern hardwood forests.

COLLECTIONS EXAMINED: (only collectors initials cited after first full citation): CANADA: Ontario: London, Sept. 27, 1896, Aug. 18, 1897, Sept. 8, 1915, J. Dearness (DAOM); and probably London area, Sept. 11, 1903 and October 23, 1926, J.D. (DAOM); Toronto, July 12, 1932, H.S. Jackson (DAOM 50112, ex TRIC 3381). U.S.A.: Georgia: Rabun Co., Rabun Bald, Sept. 3, 1947, Walters 153 (MICH). Illinois: River Forest, bank of Desplaines R., July 1, 1902, E.T. & S.A. Harper 542 (F). Kentucky: Harlan, Sept. 4, 1916, C.H. Kauffman (MICH). Michigan: Lenawee Co., Cleveland L., Onsted State game area, Sept. 9, 1970, W. Patrick (M. Gilliam 939; MICH); Livingston Co., George Reserve, July 27, 1970, J. Williams (M.G. 834; MICH), Pinckney, Aug. 7, 1937, A.H. Smith 6823 (MICH). Oakland Co., Proud L., Oct. 10, 1970, M.G. 992 (MICH), Haven Hill, Aug. 8, 1972, A.H.S. 81375 (MICH); Washtenaw Co., Ann Arbor, July 25, 1912, C.H.K., Oct. 7, 1931, A.H.S., Aug. 3, 1935, A.H.S. 1699, Aug. 22, 1937, A.H.S. 7175, Aug. 11, 1960, A.H.S. 62873, July 11, 1970, C. Nimke (M.G. 693, 701), July 30, 1970, C. Nimke 49 (MICH), Gorman L., Aug. 20, 1972, A.H.S. 81620 (MICH), Halfmoon L., July 26, 1970, S.J. Mazzer 6186 (MICH), Manchester, July 4, 1935, A.H.S. 1470 (MICH), Sharon Hollow, Aug. 4, 1960, R.L. Shaffer 2513, Aug. 19, 1970, M. Gilliam 929, Sept. 16, 1970, M.G. 952 (MICH), Silver L. area, Aug. 22, 1960, R.L.S. 2646 (MICH), Waterloo Rec. area, Sept. 3, 1968, F. Hoseney 1094, July 10, 1970, F. Hoseney (M.G. 625), July 22, 1970, M.G. 786, July 2, 1971, F.H. 1800 (MICH), Winnewana, July 11, 1970, M. Gilliam 648 (MICH). Minnesota: Rice Co., Wheeling Twp., Nerstrand State Park, Aug. 29, 1965, M.G. Weaver 1243 (MICH). North Carolina: July 11, 1924, J.V. Couch 7380 (MICH), Swain Co., Flat Cr., Aug. 1, 1937, L.R. Hesler & A.J. Sharp (MICH). New York: Albany, rural cemetery, July-Aug. 1872, C.H. Peck [type of *M. decurrens*] (NYS); East Worcester, July, C.H.P. (NYS); Forestburgh, Sept. 1874, C.H.P. [type of *A. rhododendri*] (NYS); Genesee Co., Bergen Swamp, Sept. 4, 1972, H.S. Vishniac (MICH); Greenbush, Aug., C.H.P. (NYS); Ithaca, Aug. 11, 1904, C.H.K. (MICH), Aug. 1935, H.C. Beardlee Jr. 35074 (MICH); Port Jefferson, Aug. 6, C.H.P. [type of *M. resinosus* var. *niveus*] (NYS); Schuyler Co., Hector land rice area, Aug. 19, 1972, H.S.V. (MICH). Ohio: Lane, July 13, 1922, H.C.B. Jr. (MICH); Portage Co., West Branch State Park, July 8, 1972, M. Gilliam 1493 (MICH). Pennsylvania: Mt. Gretna, Sept. 5, 1926, C.H.K. (MICH); Media, June 29, 1940, P.M. Rea & Woodbury (MICH). Tennessee: Great Smoky Mts. Park, Cades Cove, Aug. 18, 1938, A.H.S. 10374, Indian Gap, Aug. 29, 1938, A.H.S. 10625, Aug. 7, 1942, L.R.H. 14476, Laurel falls trail, Aug. 8, 1938, A.H.S. 9909 (MICH).

Resinomycena rhododendri was first described by Peck (1872) as *Marasmius decurrens*. He apparently collected a similar fungus in





Figs. 1-5. *Resinomycena rhododendri* (DAOM, ex Dearness Oct. 23, 1926). 1, basidiospores. 2, basidia. 3, pileocystidia. 4, caulocystidia. 5, cheilocystidia.

Greenbush in 1869, #20, but lost the specimen before describing it (J. Haines, pers. comm.). Peck's validating description of *A. decurrens* indicated a dark taxon with a grayish or tawny pileus and a gray stipe. These colours are not apparent on the type of *A. decurrens*, microscopically or macroscopically, possibly indicating that more than one taxon was involved when the description was written. However, as the name is now lectotypified by Gilliam (l.c.), it applies to the pale form described in this paper. Peck (1883) not Saccardo (1887) as is often cited, later proposed the new name *M. resinus* for *M. decurrens* which was a later homonym of *M. decurrens* Mont. However, Peck (1875) had in the meantime unknowingly described it a second time as *Agaricus rhododendri* which is, therefore, the earliest valid and legitimate name. His variety *niveus* is herein treated as being con-varietal with the type variety as it is now typified. The proposal of the new name *M. resinus* var. *candidissimus* to replace *M. resinus* var. *niveus* is superfluous under present nomenclatural rules, but it should be kept in mind that Peck was following the Rochester Code.

Smith (1947) was the first to recognize that *Agaricus rhododendri* and *Marasmius resinus* were congeneric, although he reported the spores of *A. rhododendri* to be smaller, 3-4.5 X 2.5 μ m, than for *M. resinus*. Bigelow (1970) treated the two as a conspecific but offered no evidence. In our studies spores measuring 6.5-7.2 X 3-3.2 μ m were found on the type of *A. rhododendri* which in other respects is typical for *M. resinus*. Thus the two are treated as conspecific. Hesler's (1959) report of inamyloid spores for the type of *M. resinus* is evidently in error as we concur with Gilliam (1976) that amyloid spores are present.

Resinomycena rhododendri appears to be restricted to the eastern deciduous forest where it occurs on litter of the dominating hardwood trees. It is the largest of the *Resinomycena* species and the one most commonly collected.

Resinomycena kalalochensis (Smith) Redhead & Singer, *comb. nov.*

Figs. 6-10.

= *Mycena kalalochensis* A.H. Smith (1947: 99)

"Pileus 3-8 mm. broad, convex, remaining broadly convex, margin incurved at first, spreading in age, chalk white and appearing pruinose under a lens at first, glabrous and uneven in age but remaining chalky, slightly sulcate at maturity, not hygrophanous; flesh membranous and pliant (but not reviving), odor not distinctive, taste not recorded; lamellae adnate, broad, distant, 10-12 reach the stipe, two tiers of lamellulae, white over all, edges pruinose; stipe 3-7 mm. long less than 0.5 mm. thick, equal or the base flanged slightly, strigose, the remainder pruinose like the pileus, chalky white over all."
Smith (1947: 99-100).

PILEAL EPICUTIS: an erect to suberect tangled turf of polymorphic cystidia, 25-50 X 3-8 μ m, varying from cylindrical to narrowly clavate, usually capitate to subcapitate, often secondarily septate, scantily resinous oleocystidia to variously branched to

narrow dendroid and antler-like forms, which lack resinous exudates or contents, walls thin, hyaline, smooth, clamped basally. PILEAL TRAMA: somewhat duplex with slightly broader hyphae above, 5-15 μ m diam., than below, 5-10 μ m diam.; hyphae faintly pseudoamyloid initially but becoming more intense after several days in permanent Hoyer's-Melzer's reagent, clamped, smooth, thin-walled, subparallel. LAMELLAR TRAMA: hyphae similar to the pileus trama hyphae. CHEILOCYSTIDIA: abundant, forming a sterile edge, less variable than the pileocystidia, mostly narrowly cylindrical to slightly clavate, occasionally forked or branched, occasionally with an apical finger-like elongation, 31-48 X 5-6.5 μ m, scantily resinous. BASIDIA: 24-25 X 7.5-8.0 μ m, clavate to obscurely utriform, 4-spored, clamped. BASIDIOSPORES: 7.8-10.8 X 3.8-4.8 μ m, ellipsoid to obscurely fusoid or occasionally obovoid, inequilaterally flattened in profile, smooth, thin-walled, amyloid, hyaline, with a prominent apiculus. STIPE HYPHAE: parallel, 5-15 μ m diam., smooth, pseudoamyloid, hyaline, with the broadest hyphae in the medulla. CAULOCYSTIDIA: similar to the pileocystidia but only scantily resinous like the cheilocystidia. BASAL MYCELIUM: not studied.

HABITAT, HABIT and SUBSTRATES: Scattered on litter of *Rubus*, *Alnus rubra* Bong. and grasses in the coastal forest zone of western North America.

COLLECTIONS EXAMINED: CANADA: *British Columbia*: North Vancouver, Capilano Canyon, Oct. 8, 1973, S.A.R. #AS 9 (DAOM 166524), Vancouver, Univ. B.C., June 2, 1971, S.A.R. 342 (UBC). U.S.A.: *California*: Humboldt Co., Big Lagoon Park, Dec. 18 & 19, 1956, A.H.S. 56787, 56833, 56823, Prairie Cr. Park, Dec. 9, 1956, A.H.S. 56520 (MICH). *Oregon*: Seaside, Sept. 21, 1944, W. Gruber & A.H.S. 19020 (MICH). *Washington*: Jefferson Co., Kalaloch, April 30, 1939, A.H.S. 13035 [type] (MICH).

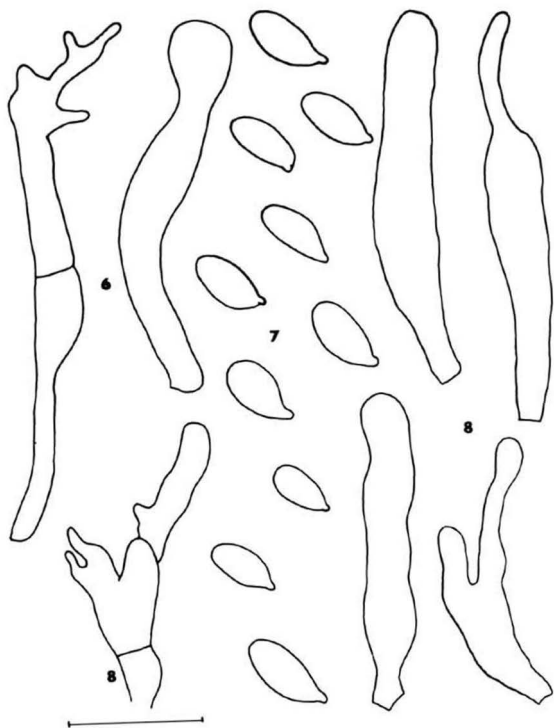
A.H. Smith (1947) reported slightly larger spores, 8-11 X 5-6 μ m, an inamyloid lamellar trama and a lack of incrustations on the pileocystidia. In our studies resinous exudates and dextrinoid tramal tissues were found on the type although the iodine reaction was not as prominent as that for the stipe. In addition, the resinous exudates tend to become clarified and dissolve in Melzer's reagent more so than in KOH 3% aqueous sol.

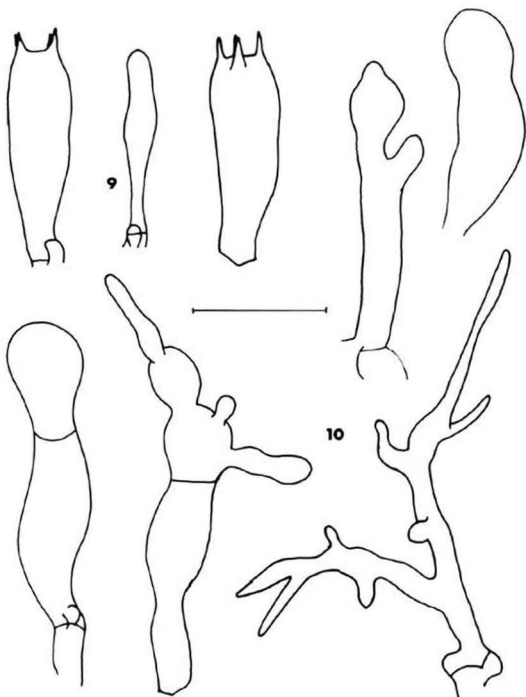
Resinomyceia kalalochensis differs from *R. rhododendri* by distant lamellae, smaller basidiomes, larger spores, and more scant resinous exudates, in addition to the differences in geographic range and to a lesser extent substrates.

Resinomyceia montana Redhead & Singer, sp. nov.

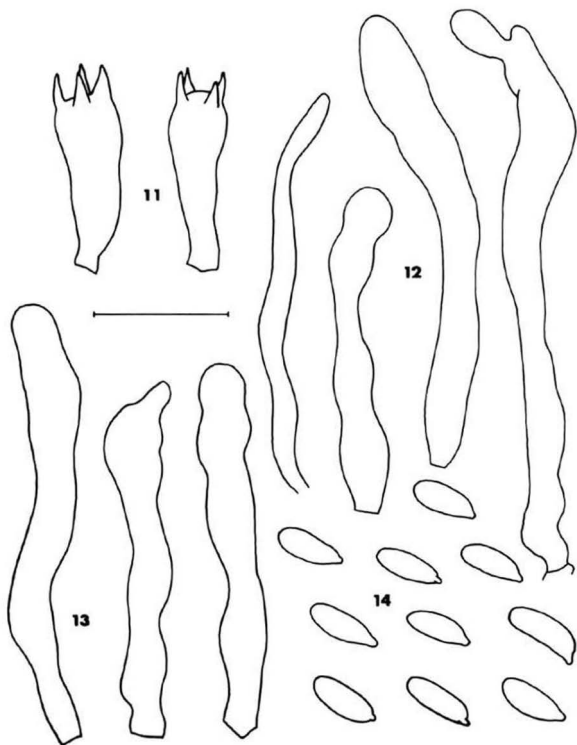
Figs. 11-14, 26.

Pileus 3.5-5.5 mm *latus*, *convexus vel depressus*, *candidus*. *Lamellae arcuatae, candidae*. *Stipes* 9-23 mm *long.*, 0.4-0.6 mm *crassus, candidus*. *Sporae* 8-9.3 X 2.8-4.5 μ m, *amyloideae, ellipsoideae*. *Pileocystidia* 45-67 X 2.5-7 μ m, *resinosa, clavata vel dendroidea ut nonresinosa*. *Cheilocystidia* 42-52 X 6-8.2 μ m, *resinosa, clavata vel strangulata*. *Tramate pseudoamyloideo*.





Figs. 6-10. *R. kalalochensis* (DAOM 166524). 6, caulocystidia. 7, basidiospores. 8, cheilocystidia. 9, basidia. 10, pileocystidia.



Figs. 11-14. *P. montana* (DAOM 178214). 11, basidia. 12, pileocystidia. 13, cheilocystidia. 14, basidiospores.

Holotypus: DAOM 178215. Donald Station, Rocky Mt. trench, B.C., Canada, Sept. 21, 1980, S.A. Redhead 3977.

PILEUS: 3.5-5.5 mm wide, convex and slightly depressed centrally, vaguely striate to chalky white, micaceous, obscurely corrugated-striate, developing greyish casts when dry; context white, pliant; odor not distinctive. **LAMELLAE**: arcuate decurrent, moderately spaced, white, with micaceous edges; lamellulae in 2 tiers. **STIPE**: 9-23 mm long, 0.4-0.6 mm wide, equal, cargiliginous or tougher, white, finely powdered overall.

PILEAL EPICUTIS: a suberect to tangled turf of polymorphic cystidia 45-67 X 2.5-7 μ m, varying from a majority of elongated clavate oleocystidia to filiform or dendroid nonresinous elements. **PILEAL TRAMA**: obscurely duplex, with slightly broader hyphae above, 3-11 μ m diam., than below 3-5 μ m diam.; hyphae subparallel, faintly pseudoamyloid at first, smooth, clamped, thin to slightly thick-walled, hyaline. **LAMELLAR TRAMA**: hyphae similar to the lower pileal trama hyphae. **CHEILOCYSTIDIA**: abundant, forming a sterile edge, 42-52 X 6-6.2 μ m, narrowly clavate to cylindrical and slightly strangulate, occasionally with a short apical elongation, with scant to abundant resin exudates. **BASIDIA**: 20-21 X 6.8-7.2 μ m, clavate subcapitate, 4-spored, clamped. **BASIDIOSPORES**: 8-9.3 X 2.8-4.5 μ m, cylindrical to narrowly ellipsoidal, slightly inequilateral in profile, amyloid, smooth, hyaline, with a prominent apiculus. **STIPE HYPHAE**: parallel, pseudoamyloid especially on the slightly broader medulla hyphae, 4-13 μ m diam., smooth, thin to slightly thick-walled, clamped. **CAULOCYSTIDIA**: scattered to abundant and often clustered, similar to the pileocystidia. **BASAL MYCELIUM**: not studied.

HABITAT, HABIT and SUBSTRATES: scattered to gregarious on coniferous needles, cone debris and small twig fragments in western Subalpine, Coastal and Columbian coniferous forests of North America.

COLLECTIONS EXAMINED: CANADA: *British Columbia*: North Vancouver, Mt. Seymour, 3500' alt., Sept. 17, 1973, S.A.R. #AJ 3 (DAOM 178218); Sept. 22, 1973, S.A.R. #AI 24 (DAOM 178214); Columbia R. valley in Rocky Mt. trench at Donald Stn. on Hwy. 1 crossing of river, Sept. 21, 1980, S.A.R. 3977 [type] (DAOM 178215). U.S.A.: *Washington*: Lower Elwha, July 3, 1939, A.H.S. 14732 (MICH).

Resinomycena montana has longer spores than *R. rhododendri*, occurs on coniferous debris and usually develops a greyish cast on drying. Geographically the two are widely separated. *Resinomycena kalalochensis* differs by its more capitate and shorter pileocystidia which are often secondarily septate and scarcely resinous. Also the latter has smaller basidiomes which have not been observed to develop a grey cast and it occurs on angiosperm debris.

Resinomycena acadensis Redhead & Singer sp. nov.

Figs. 20-25.

Pileus 2-10 mm latus, convexus demum depressus, candidus. *Lamellae* arcuatae, candidae. *Stipes* 10-40 (-70) mm long., 0.2-0.8 mm crassus, candidus. *Sporae* 9.4-12.8 X 4-5 μ m, amyloideae, ellipsoideae. *Pileocystidia* 30-58 X 3-9 μ m, resinosa, clavata vel nonresinosa ut

dendroidea. *Cheilocystidia* 38-40 X 6-8 μ m, aequae *pileocystidia*.
Tramate pseudoamyloideo.

Holotypus: DAOM 166073. Kouchibouguac Natl. Park, N.B., Canada
Sept. 24, 1977, J.E. & S.A. Redhead 2558.

PILEUS: 2-10 mm wide, convex becoming depressed to cyathiform, translucent-striate but readily becoming opaque on partial drying, white to whitish, occasionally tinted fawn on the disc, rugose with age, moist to dry; edges incurved initially; context white, pliant; odor and taste not distinctive. **LAMELLAE**: arcuate-decurrent, becoming more distinctly decurrent, white, subdistant to moderately spaced; edges crenulate and often beaded; lamellulae in 2 tiers. **STIPE**: 10-40 (-70) mm long, 0.2-0.8 mm wide, equal or tapering up, white or with ochreous to rosy buff tints basally, glutinous when wet, powdered when dry, often attached to the substrate by a small radiating silky white mycelial pad.

PILEAL EPICUTIS: a suberect turf and later repent tangle of polymorphic cystidia 30-58 X 3-9 μ m, varying from short to long clavate oleocystidial forms to narrow filiform to dendroid nonoleocystidial forms with some intergradation, thin-walled, smooth, hyaline, with copious to moderate resinous exudates. **PILEAL TRAMA**: obscurely duplex with the broadest forms in the upper half, hyphae subparallel, 5-11 μ m, smooth, hyaline, faintly pseudoamyloid at first, clamped. **LAMELLAR TRAMA**: hyphae mostly parallel, 3-5 μ m diam., pseudoamyloid. **CHEILOCYSTIDIA**: abundant forming a sterile edge, similar to the pileocystidia but rarely of the branched forms, mostly clavate, 38-40 X 6-8 μ m. **BASIDIA**: 20-32 X 7-8 μ m, (2-)4-spored, clavate, clamped. **BASIDIOSPORES**: 9.4-12.8 X 4-5 μ m, ellipsoidal to cylindrical or obscurely reniform to pip-shaped, inequilaterally flattened in profile, with a prominent apiculus hyaline, white in mass, amyloid. **STIPE HYPHAE**: 3-10 μ m diam., with narrower hyphae in the cortex than in the medulla, strongly pseudoamyloid, smooth, parallel, hyaline, clamped. **CAULOCYSTIDIA**: usually clustered, similar to the pileocystidia but less often with long pedicels and more often secondarily septate. **BASAL MYCELIUM**: slightly interwoven, agglutinated, 3-10 μ m diam., pseudoamyloid.

HABITAT, HABIT and SUBSTRATES: scattered on small broken pieces of woody angiosperm debris in predominantly Acadian coniferous forests of larch or spruce, fir and pine.

COLLECTIONS EXAMINED: CANADA: *New Brunswick*: Kouchibouguac Natl. Park, Aug. 11, 1977, R.L. Milikin (DAOM 169619), Sept. 24, 1977, J.E. & S.A.R. 2558 [type] (DAOM 166073), Oct. 9, 1978, S.A.R. 2753 (DAOM).

Resinomyцена acadiensis can be distinguished from *R. rhododendri* by the larger spores, more distant lamellae and the more flaccid flesh. *Resinomyцена kalalochensis* differs by its small stature, its more erect turf of capitate and scarcely resinous, secondarily septate pileocystidia and its geographic range. *Resinomyцена montana* differs by its smaller usually narrower spores, smaller basidia, a coniferous substrate and its geographic range.

The glutinous to viscid stipes are a result of copious resinous

exudates. These dry down to hardened masses in less humid situations creating a powdered appearance to the stipes. In herbarium specimens these masses become yellow with age.

Resinomyceia brunescens Redhead & Singer, sp. nov.

Figs. 15-19.

Pileus 6 mm *latus*, *planiconvexus* ut *depressus*, *rugulosus*, *brunneus*. *Lamellae* *adnatae* vel *subdecurrentes*, *tubalineae* vel *ochraceae*. *Stipe* 20 mm *long.*, 0.5 mm *crassus*, *concolorus*. *Sporae* 6.5-8 X 3.3-4µm, *amyloideae*, *ellipsoideae*. *Pileocystidia* 55-90 X 3-11µm, *resinosa*, *clavata* vel *strangulata*, vel *nonresinosa* ut *dendroidea*. *Cheilocystidia* 36-50 X 10-14µm, *resinosa*, *clavata* vel *capitata*, *raris nonresinosa* ut *dendroidea*. *Tramate pseudoamyloideo*.

Holotypus: DAOM 165884. Kouchibouguac Natl. Park, N.B., Canada, July 13, 1977, J.E. & S.A. Redhead 2336.

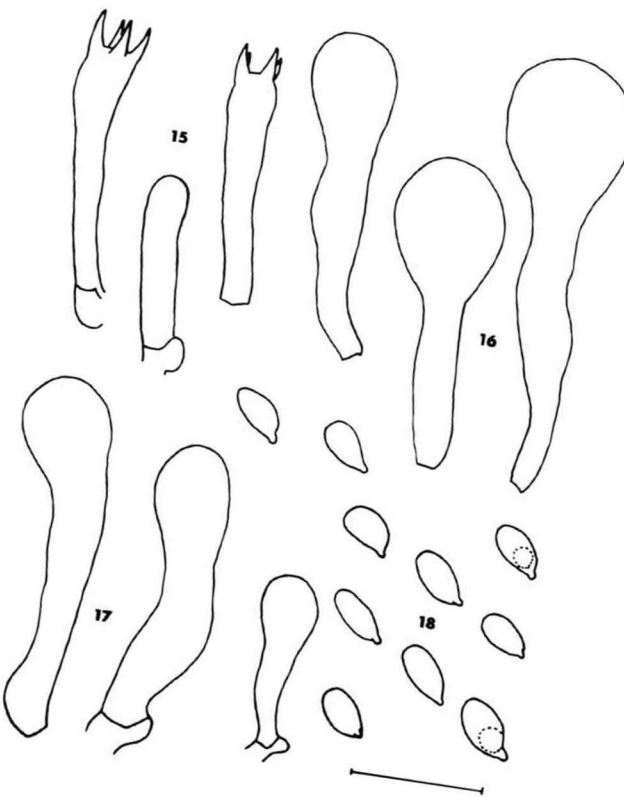
PILEUS: 6 mm wide, plano-convex and depressed centrally, rugose, tacky, scarcely translucent on edges, greyish sepia centrally, fulvous with ochreous to buff margins; context membranous, thin, concolorous above; odor not distinctive. **LAMELLAE:** adnate to subdecurrent, moderately spaced, buff to ochreous with whitish crenulate edges, sometimes forked; one tier of lamellulae. **STIPE:** 20 X 0.5 mm, concolorous with the pileus with the darkest pigments below, with a striated powdered appearance from dried resin, micaceous, attached to the substrate by a radiating white mycelial pad.

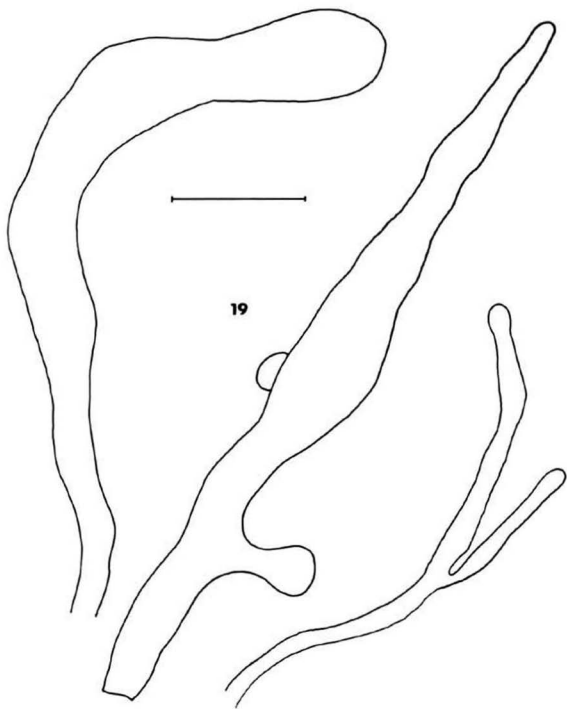
PILEAL EPICUTIS: a layer of tangled polymorphic cystidia 55-90 X 3-11µm, varying from clavate to fusoid often strangulate oleocystidia exuding copious resinous exudates to filiform often branched dendroid nonresinous forms. **PILEUS TRAMA:** duplex, with hyphae 10-15µm diam. subparallel, thin-walled, pseudoamyloid, yellowish brown and prominently incrustated with brownish resin-like materials in flat patches above, 5-10µm diam., hyaline, walls thin or pronounced below. **LAMELLAR TRAMA:** hyphae similar to those of the lower pileus trama. **CHEILOCYSTIDIA:** abundant, forming a sterile edge which on drying is a continuous hardened resinous line, 36-50 X 10-14µm, clavate and prominently capitate, hyaline, thin-walled. **BASIDIA:** 27-29 X 5-6µm, narrowly clavate (2-)4-spored, clamped. **BASIDIOSPORES:** 6.5-8 X 3.3-4µm, ellipsoid to obscurely obovoid or fusoid, hyaline, smooth, thin-walled, amyloid, with a prominent apiculus. **STIPE HYPHAE:** parallel, 3-5µm diam. in the cortex with pigments similar to the pileus tramal hyphae, 10-15µm diam. in the medulla, pseudoamyloid, clamped. **CAULOCYSTIDIA:** clustered, similar to the cheilocystidia but less swollen apically, occasionally forked, intermixed with a few dendroid filiform elements. **BASAL MYCELIUM:** not studied.

HABITAT, HABIT and SUBSTRATE: solitary on fallen leaf of *Rhododendron canadense* (L.) Torr. in a dense Acadian coniferous forest.

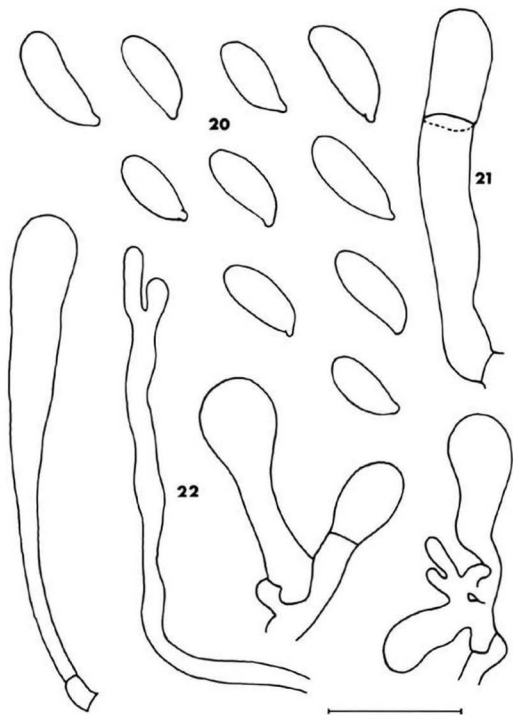
COLLECTION EXAMINED: CANADA: *New Brunswick:* Kouchibouguac Natl. Park, July 13, 1977, J.E. & S.A.R. 2336 [type] (DAOM 165884).

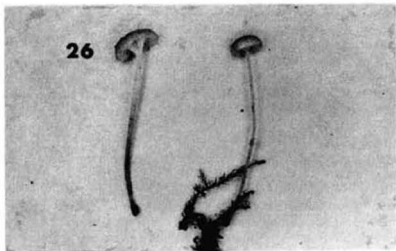
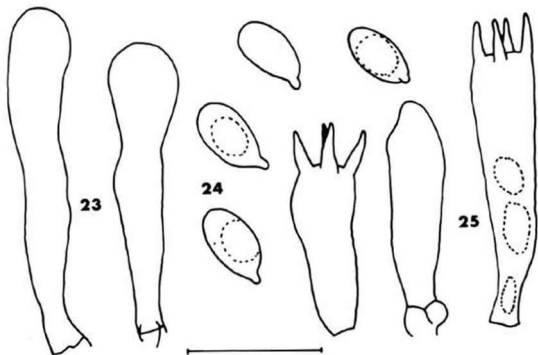
Resinomyceia brunescens is readily distinguished from all other species in the genus by its darkly pigmented pileus and stipe. The





Figs. 15-19. *R. brunnescens* (DAOM 165884). 15, basidia.
16, cheilocystidia. 17, caulocystidia. 18, basidiospores.
19, pileocystidia.





Figs. 20-25. *R. acadiensis* (all DAOM 166073 except 24, DAOM 169619).
 20, basidiospores. 21, caulocystidium. 22, pileocystidia.
 23, cheilocystidia. 24, basidiospores. 25, basidia.
 Scales = 15 μ m. 26, *R. montana* (DAOM 178214), carpophore, mag. X 2.

incrusted hyphae responsible for this darkened appearance are very distinctive microscopically compared to the nonincrusted hyphae in other *Resinomyceena* species. In addition, *R. brunnescens* is characterized by very prominently capitate cheilocystidia which are inflated more than in any of the other species.

As noted above, Peck's original description of *Marasmius decurrens* is of a brown fungus. The existence of a brown species of *Resinomyceena* in eastern North America suggests that Peck may have collected both *R. brunnescens* and *R. rhododendri*. However, this cannot be confirmed now.

GENERAL DISCUSSION

The three most probable generic depositories for *Resinomyceena* if the taxon is not recognized as a distinct genus are *Hydropus*, *Mycena* and *Baeospora*, there being a foregone conclusion that the species are excluded from the genus *Marasmius* because of amyloid spores. The genus *Baeospora* as it was defined in *The Agaricales in Modern Taxonomy* (Singer, 1975) represents a small cohesive genus characterized by a number of characters, but in particular, small amyloid spores, inamyloid tramal tissues and crowded lamellae. *Resinomyceena rhododendri* is the species most similar to a *Baeospora* in that it has close lamellae and small amyloid spores, however, the tramal tissues of the pileus and stipe are pseudoamyloid unlike any *Baeospora* species. Also the massed large oleocystidia mixed with dendroid elements on the pileus form a structure unlike any epicuticular formation found in *Baeospora*, including *B. pallida*. When all species of *Resinomyceena* are taken into account, species such as *R. acadensis* and *R. kalalochensis*, exhibit characters not found in *Baeospora* such as distant, slightly decurrent lamellae and large spores. As a result of these combinations of characters *Resinomyceena* cannot readily be incorporated into *Baeospora*.

The genus *Hydropus* as defined by Singer (1975) contains species having numerous pileo-, cheilo- and caulocystidia and in some species pseudocystidia with oily contents. The spores can be amyloid and the tramal tissues can be weakly pseudoamyloid. Again the abundant oleocystidia on the pilei, stipes and lamellae of *Resinomyceena* species, not found in *Hydropus*, in combination with intermixed dendroid elements form a tissue not found in *Hydropus* and clearly separate the genera.

The genus *Mycena* is treated as delimited earlier (Singer 1975). It includes species with amyloid spores, pseudoamyloid tramal tissues and oleocystidia as does *Hydropus*. The majority of *Mycena* species have an epicutis consisting of a repent layer of diverticulate narrow hyphae. Deviations occur in some sections of the genus where smooth repent filamentous hyphae replace the diverticulate elements. Those species which had smooth epicuticular hyphae bearing obtuse cystidia were removed to the genus *Hydropus*. The most aberrant type of epicutis for a species admitted to *Mycena* by Singer (1975) is found in *Mycena vorida* (Fr.) Kühner and its allies. These species have a hymeniform layer of sphaeropedunculate cells and, apparently missed by previous authors, in *M. vorida*, intermixed oleocystidia. Previously

(Singer 1975), it was indicated that these species were related to *Hydropus* and might need to be removed from *Mycena*. However, the strongly pseudoamyloid tramal tissues and a gelatinized stipe cortex weighed in favor of retaining the species in *Mycena*. Here again, the well developed layer of usually clavate oleocystidia and narrow dendroid elements found in *Resinomyceana* is not typical of any section of *Mycena* and represents a large divergence from a typical *Mycena* type of epicutis. Furthermore, the typical *Mycenae* have characteristically broad (broader than *Resinomyceana*) hypodermal and tramal hyphae; the hyphal cells often being short and voluminous. In a less restrictive delimitation of *Mycena* such as by Smith (1947) or Kühner (1938), *Resinomyceana* would be included.

From the above discussion it can be seen that *Baeospora*, *Hydropus* and *Mycena* are open to emendation to encompass *Resinomyceana* species but only by introducing a species group having an unusual combination of characters peripheral to the core of the genera involved. Instead the proposal to recognize a distinct genus seems more justified. *Resinomyceana* species are obviously closely related to each other. They are readily distinguished from *Baeospora*, *Hydropus* and *Mycena* by the abundance of resinous oleocystidia intermixed with dendrohyphidia. The segregation of *Resinomyceana* species maintains the integrity of the older genera, leaving them less heterogeneous and, therefore, more recognizable. There is no doubt that *Resinomyceana* is closely related to *Baeospora*, *Mycena* and *Hydropus*. Obviously one solution would be to merge all three genera and add *Resinomyceana*. Again there seems to be more merit in recognizing each of these genera including *Resinomyceana*.

Finally, *Marasmius rhododendri* Singer (1947, 1936), a species described from Spain in the subalpine zone of the Pyrenees on *Rhododendron ferrugineum* L. is probably congeneric. The type (F) now consists of stipes only. Spores obtained from the type were amyloid although weakly in some and up to 12.5µm long. The stipe hyphae are pseudoamyloid and in an outer layer more or less gelatinized (the stipe was described from fresh material as somewhat shiny and viscidulous). They bear cystidia with dried incrustations similar to that found on *Resinomyceana* species. In the original description (1936) yellowish incrustations were mentioned on the lamellar edges of vesiculose to ampulliform cheilocystidia ("echinulatis" possibly being an error in the 1947 description unless referring to dendroid elements).

The exact nature of these cystidia and the pileipellis could not be restudied in the damaged type collection and the identity of the species remains in doubt. The weakly developed collar is not found on the presently recognized *Resinomyceana* species. This species should be restudied from fresh materials.

ACKNOWLEDGEMENTS

We thank Drs. J. Haines (NYS), R.J. Bandoni (UBC), R.L. Shaffer, R. Fogel and A.H. Smith (MICH) for the use of herbarium materials and facilities. Drs. J. Ginns and J. Ammirati offered appreciated reviews and Mr. K.W. Spicer provided technical assistance. Parks Canada supplied collecting permits or facilities in New Brunswick and British Columbia and the Agriculture Canada research station, Kamloops,

provided transportation in 1980. Mrs. M. Meredith typed the final manuscript.

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SINOTERMITOMYCES, A NEW GENUS OF AMANITACEAE
FROM YUNNAN, CHINA

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SUMMARY

Detailed analysis has led the author to propose the termitophilous *Sinotermitomyces* as a new genus, including two new species, *S. cavus* Zang and *S. carnosus* Zang. Both were collected in Yunnan, China. Type specimens of the two new species are deposited in the Herbarium of the Kunming Institute of Botany, Academia Sinica (HKAS).

The new genus *Sinotermitomyces* consists of two new species growing on termite combs. These edible fungi are distributed in subtropical and tropical regions of southwestern Yunnan. In Yunnan, professional collectors gather both *Termitomyces* and *Sinotermitomyces* in the rainy season from May to September, and dry preservation with salt and vegetable oils are used for building up a year-round supply (Cheo, 1942, 1948; Batra, 1979).

The genus *Termitomyces* in China is commonly found southward of the Yangtze. The distribution of *Sinotermitomyces* is in forests of the southwestern part of Yunnan. Specimens were collected by the author under *Pinus khasya* Royle ex Cord., *Pinus yunnanensis* Fr., and *Castanopsis hystrix* (Hk. f. & Thoms.) A. DC.

SINOTERMITOMYCES Zang, gen. nov.

Pileus conicus vel campanulatus, subconvexus vel papillatus, siccus, glaber vel scabridus. Lamellae subliberae, liberae vel adnatae, albae. Stipes centralis, cavus, aequalis vel sub-fusoideus, carnosus, coriaceus vel fibrillosus. Annulus superus distinctus. Basidiosporae hyalinae; ellipticae vel obovatae, inamyloidea. Pleurocystidia ventricosa vel cylindrica, verrucosa vel tuberculosa. Cheilocystidia oblonga vel cylindrica, laeviuscula vel verrucosa. Fibulis nullis. Termiticola.

Typus generis: *Sinotermitomyces cavus* Zang.

Pileus conico-campanulate, subconvex and often with a prominent papillate apex, dry, never viscid, glabrous or scabrous. Gills free to almost free or adnate. Stipe central, often very long, variable in thickness, hollow, equal or subfusiform, fleshy or leathery or fibrillose. Annulus superus present, veil a pellicular veil that covers the gills in young specimens. Basidiospores from globose to ellipsoid, smooth, hyaline, inamyloid. Pleurocystidia present, cylindrical or some fusoid-ventricose, covered with wart-like outgrowths. Cheilocystidia oblong to cylindrical, smooth to warted. Clamp connections absent. The primordia develop in the holes of hypogeous termi-

taria.

This new genus can be distinguished from the genus *Termitomyces* Heim by the following characters: the small, subconvex pileus with prominent papillate apex; presence of veil remains on stipe and pileus; stipe leathery or fibrillose and hollow; pleurocystidia with sturdy, short processes (Heim, 1942; Singer, 1949).

1. *Sinotermitomyces* CAVUS Zang, sp. nov. (FIGS. 1-7)

Pileus 1-2.5 cm., conicus vel subconvexus, siccus, glaber, primo flavidus vel eburneus, demum fulvo-umbrinus vel rufo-brunneus. Lamellae subliberae vel adnatae, albiae vel eburneae, Stipes 20-30 cm. longus, 5-12 mm. crassus, aequalis, cavus, tubulosus, coriaceus vel fibrillosus, apice cicatricatus, sursam scabroso-furfuraceus, basim versus discoideus. Annulus superus distinctus. Basidiosporae 2.4-5 x 3.5-9 μ m, hyalinae, globosae, ovoideae, ellipsoideae, laeves, inamyloideae. Basidia clavata, 15-25 x 10-14 μ m, 4-sporigera. Pleurocystidia 12-20 x 25-45 μ m, cylindrica vel clavata, verrucosa vel tuberculosa. Cheilocystidia 10-15 x 20-30 μ m, oblonga vel cylindrica, tuberculosa. Fibulis nullis.

Hab. In sylvis praecipue *Pinetis yunnanensi* Fr. et *Pinetis khasyae* Royle ex Cord. nido Termitidarum (*Odontotermes*) erumpente.

Yunnan: Tengchung County, Tuan-Tian village, La-ba-qing. On nest of termites, in swampy grassland, under the pine woods above 2100 m. alt., 8. VIII. 1980. Li Xing-jiang 11. (Typus, HKAS 6533); Tengchung County, Pu-Chuan village, 17. VII. 1979. Ma Xi-xiong 1, 2. (HKAS 4612, 4613); Shweli County, Deng Ga, in pine woods, 13. VIII. 1980. Li Xing-jiang 22. (HKAS 6568); Mangshi. 10. VIII. 1980. Zang Mu 06545 (HKAS 6545).

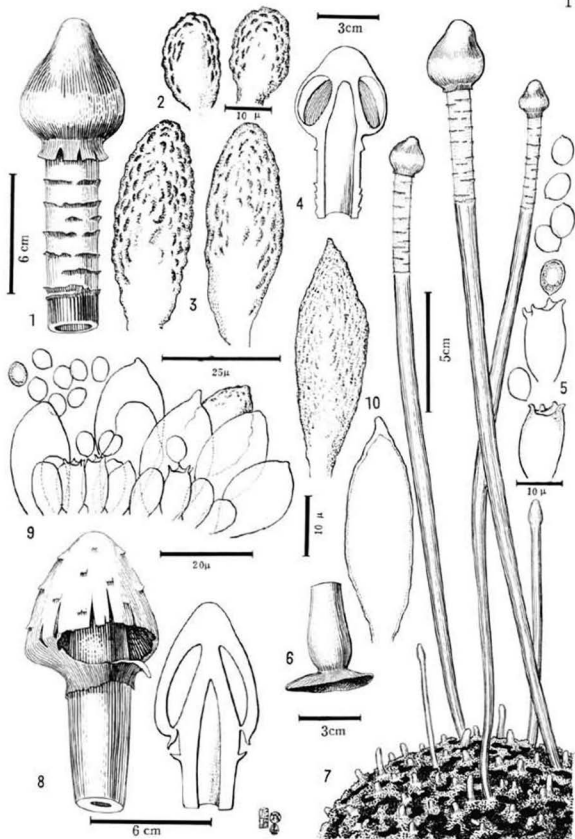
Pileus 1-2.5 cm broad, conical or subconvex, dry, glabrous, surface ivory white or pale yellow becoming yellowish-umber or reddish-brown towards the center, glabrous. Gills almost free or adnate, white to ivory white. Stipe 20-30 x 0.5-1.2 cm, equal, hollow, tube-like, leathery to fibrillose, upwards often covered with bran-like scales or scabrous and with a discoid base. Veil persistent as a thick, membranous leathery annulus, which is striate above. Basidiospores 2.4-5 x 3.5-9 μ m, hyaline, globose, ovoid to ellipsoid, smooth, inamyloid. Basidia 15-25 x 10-14 μ m, clavate, 4-spored. Pleurocystidia 12-20 x 25-45 μ m, cylindric to clavate, covered with irregular wart-like outgrowths. Cheilocystidia 10-15 x 20-30 μ m, oblong to cylindric, roughened. Clamp connections absent.

2. *Sinotermitomyces* CARNOSUS Zang, sp. nov. (FIGS. 8-10)

Pileus 4-6.5 cm., conicus vel campanulatus, demum plani-convexus, siccus, scabridus vel subtomentosus, primo albidus vel pallidus, demum cervinus vel brunneus. Lamellae subliberae vel adnatae, albiae. Stipes 15-20 cm. longus, 1-2.5 cm. crassus, subfusioideus, carnosus, cavus, concolorous, pseudorhizophorus. Annulus membranaceus, superus instructus. Basidiosporae 3-6 x 4.9-8 μ m, hyalinae, globosae, ovoideae, ellipsoideae, laeves, inamyloideae. Basidia 9.8-12 x 14-18 μ m, clavata, 4-sporigera. Pleurocystidia 10-18 x 24-48 μ m, lageniformia vel fusiformia, exasperata vel verrucosa. Cheilocystidia 12-18 x 18-34 μ m, rotunda, ovata, oblonga, laevicula vel exasperata. Fibulatae adsunt.

Hab. In sylvis praecipue *Castanopsis hystrix* (Hk. f. & Thoms.) A. DC. alt 1600-2100 m. nido Termitidarum erumpente.

Yunnan: Tsangyuan Autonomous County of Wa Nationality. On termites' nest. 30. VIII. 1980. Zang Mu 06752 (Typus, HKAS 6752).



FIGS. 1-10. *Sinotermatomyces* spp. 1-7, *S. cavus* Zang (HKAS 6533). 1, 4, 7, carpophores; 2, cheilocystidia; 3, pleurocystidia; 5, basidia and basidiospores; 6, the discoid base of stipe. 8-10, *S. carnosus* Zang (HKAS 6752). 8, carpophores; 9, cheilocystidia, basidia, and basidiospores; 10, pleurocystidia.

Pileus 4-6.5 cm broad, conical, campanulate to plano-convex, dry, scabrous or subtomentose, surface white or pale white to waxy yellow or brown. Gills almost free or adnate, white. Stipes 15-20 x 1-2.5 cm, subfusiform, hollow, more fleshy, uniform in colour, pseudorhiza present. Annulus superior, fragments adhering to stipe. Basidiospores 3-6 x 4.9-8 μ m, hyaline, globose, ovoid to ellipsoid, smooth, inamyloid. Basidia 9.8-12 x 14-18 μ m, clavate, 4-spored. Pleurocystidia 10-18 x 24-48 μ m, flask-shaped or fusiform, roughened with wart-like outgrowths. Cheilocystidia 12-18 x 18-34 μ m, almost round, broadly oblong, almost smooth to roughened. Clamp connections absent.

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MYCOTAXON

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TYPE STUDIES IN THE POLYPORACEAE 13.
SPECIES DESCRIBED BY J. H. LÉVEILLÉ.

by

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S U M M A R Y

Of the 145 polypores described by J. H. Lévèillé 12 are accepted, 83 are synonyms, 6 names are invalid, 5 are an uncertain status, most of them Ganoderma sp., while 39 types were not found. The names of many of the missing types are accepted as synonyms based on notes by previous mycologists. The combination Tyromyces dissectus is proposed. The latter species and Trichaptum perrottetii are described in detail.

The French mycologist J. H. Lévèillé was very industrious and described many fungi of which 145 are polypores. His basis was the collections in the Paris herbarium besides which he made a journey to Leiden and was allowed to take samples from the rich East-Asian collections of Jung-huhn and Zollinger. Most of his collections are today in the Paris Herbarium (PC), but many of the polypore types are also in the Leiden herbarium (L.). Isotypes are at Kew (K), Stockholm (S), the Farlow herbarium (FH) and the National Fungus Collection (BPI).

Lévèillé made very scanty labels, but often he fastened his specimens to the sheet with a red silky band attached both to the specimen and the sheet with a red lacquer. This made it possible to study both sides of the specimen without loosening it from the sheet.

This red lacquer has in many cases made it possible to trace some of his types when label and specimen have been separated, which was the case in many collections. Nevertheless, many of his types have been lost. However, both Lloyd (1912) and Bresadola (1916 & 1920) studied many of his specimens before they disappeared and it is possible to settle their identity. Lévèillé described polypores in a number of genera, and they are treated in the genus in which Lévèillé originally placed them. Within each genus

the species are placed alphabetically according to their specific epithets. After the name there is a reference to where it was published and an indication of the herbarium in which the lectotype and eventual isotypes can be found.

The label is then cited in inverted commas and, if the text indicates no type locality, this is added in brackets. If the type was found to be a synonym this is indicated by = followed by a correct citation of the name in question.

When the species has been accepted, it is cited in its proper genus with a reference to a recent description, or described in detail if no modern description seems to exist.

DAEDALEA Fr.

D. aulaxina 1844:197. The type has not been found (Java).
= Lenzites vespacea (Pers.) Ryv. teste Bresadola (1916:230).

D. flavida 1844:198 (PC).

"Java". = Lenzites acuta Berk.

D. fuliginosa 1844:199.

The type has not been found (Mauritius).

D. lurida 1844:198 (PC, isotype in S).

"Java" = Lenzites acuta Berk.

D. microzona 1846:142 (L).

"Zollinger no 2060" (Java) = Lenzites acuta Berk.

D. plumbea 1846:142.

The type has not been found (Noveboracum = New York, USA).

D. pruinosa 1844:198 (PC, isotype in S).

"Ins. pacifico". = Lenzites acuta Berk.

D. splendens 1844:197.

The type has not been found (Sumatra).

D. violacea 1846:142.

The type is apparently lost (Cuba).

FAVOLUS Fr.

F. fibrillosus 1844:201 (PC).

"Manille" (The Phillipines). = P. philippinensis Berk.

F. fissus 1844:201 (PC).

"Bresil, Rio de Janeiro, Mars 1836" = F. brasiliensis (Fr.) Fr.

F. granulosis 1863:286 (PC, isotype in K).

"Chachi alt. 2600 m, Sept. 1860 Lindig 2919" (Colombia).

= Panellus pusillus (Lév.) Burds. & Miller.

F. guadeloupensis 1846:144 (PC).

"Guadeloupe, M. l'Herminier".

The type does not represent a true polypore and has probably been fleshy and gelatinous when fresh. Today it is resinous and horny. Pileus glabrous and pustulate, dark brown, pores shallow and angular. 1-2 mm wide, dirty to dark brown. Context dark brown and very dense. Hyphal system monomitic with clamps, variable in diameter, in parts inflated, 2-8 μ m wide, but difficult to separate. Spores and basidia not seen. The species is unknown to me.

- F. junghuhnii 1844:202 (L, isotypes in PC & S).
 "Ad truncos, in Insula Bantan" (The Phillipines). = P. philippinensis Berk.
F. multiplex 1844:203 (PC, isotype in K & S).
 "Java". = F. spatulatus (Jungh.) Lév.
F. peltatus 1844:203 (L, isotype in PC).
 "Zollinger no. 707" (Java). = F. brasiliensis (Fr.) Fr.
F. tener 1844:202 (L).
 "Sumatra". = F. spatulatus (Jungh.) Lév.
F. tenuissimus 1844:202.
 Type not found (Mauritius).

GLOEOPORUS Mont.

- G. leptopilus 1844:194.
 Type has not been found (Surinam).
G. pusillus 1844:195.

I have not found the type, but the species has generally been accepted as a poroid agaric. Modern descriptions can be found in Singer (1945:224) who placed the species in Dictyopanus, and in Burdsall and Miller (1978:85) who transferred the species to Panellus.

HEXAGONIA Fr.

- H. blumei 1844:199 (PC).
 "Java, Blume". = H. tenuis (Hook.) Fr.
H. cingulata 1844:200.
 The type has not been located (Java).
H. cyclophora 1846:143 (PC).
 "Mahe" (Tahiti). = H. tenuis (Hook.) Fr.
H. dregeana 1846:143 (PC).
 "C.B.S. (Cape of Good Hope) Dreg. no. 17b." = H. tenuis (Hook.) Fr.
H. glabra 1846:143 (PC).
 "Bombay (India), Polydore Roux". = Lenzites acuta Berk.
H. molkenboeri 1844:260 (L).
 "Java, Junghuhn". = Lenzites vespacea (Pers.) Ryv. The synonymy was also noted by Bresadola (1916:231).
H. pulchella 1844:200 (L).
 "Java". = H. tenuis (Hook.) Fr.
H. tabacina 1854:17 (PC, isotype in K).
 "Coll. Zollinger" (Java). = Cyclomyces setiporus (Berk.) Pat.

LENZITES Fr.

- L. berkleyi 1846:122 (PC, isotypes in S & BPI).
 "Grand bassin Saint Jean (Louisiana, USA) Mougeot no 36".
 = L. betulina (Fr.) Fr.
 The type came from New York, but I have not found it. Bresadola (1920:66) had apparently examined it and indicated the synonymy given above. The cited specimen is selected as neotype until the lectotype reappears.

- L. ciliata 1844:181.
No type has been found. No type locality was given.
- L. guilleminiana 1846:122.
"Bresil meridional, Guillemin no. 1339" (PC). = Stipto-
phyllum erubescens (Berk.) Ryv.
- L. Junghuhnii 1844:180 (PC).
"ad truncos Java" Lèveillé's hand. "Pour moi identique
à Lenzites betulina L. juvenile" Bresadola's handwriting.
= L. betulina (Fr.) Fr.
- L. murina 1844:122 (PC, isotype in S).
"Java, Korthals". Selected as neotype. The type locality
is given as Sumatra, but no such collection has been
found. = Trametes menziesii (Berk.) Ryv.
- L. myriophylla 1863:282 (PC).
"Voyage de J. Triana 1851-57. Novo Granata, Colombia".
= L. elegans (Fr.) Pat.
- L. platyphylla 1844:179 (PC, isotypes in NY & BPI).
"Java, ad truncos". = Lenzites vespacea (Pers.) Ryv.
- L. platypoda 1844:180 (PC).
"Manille (The Phillipines)". = L. elegans (Fr.) Pat.
- L. tenius 1846:122 (PC).
"Guadeloupe". = L. elegans (Fr.) Pat.

POLYPORUS Fr.

- P. apalus 1848:124.
No type has been found. (Rentilly, France).
- P. abnormis 1844:186.
The type has not been found (Java). = P. sanguinaria Kl.
teste Lloyd (1910).
- P. aculeatus 1846:137 (PC, isotype in BPI).
"Zollinger no. 2055" (Java). = Trametes modesta (Fr.)
Ryv.
- P. albomarginatus 1844:191 (PC, isotypes in BPI & L).
No text on label, except the name written by Lèveillé.
(Java). This is an accepted species in Pyrofomes as P.
albomarginatus (Lév.) Ryv. For a description, see Ry-
varden & Johansen (1980:529).
- P. anisopilus 1844:191 (PC, isotypes in L & BPI).
"Java, Blume". = Coriolopsis sanguinaria (Kl.) Teng.
- P. appositus 1846:141 (PC, isotypes in L & BPI).
"Zoll. no. 19. Auf faulen Baumen in Wald bei Tjiboda
(Java) Mai 1842". This is an accepted species in Phelli-
nus as P. appositus (Lév.) Pat. For a description, see
Ryvarden & Johansen (1980:141).
- P. atypus 1844:184 (PC).
"Guadeloupe, M. L'Herminier". = Trametes modesta (Fr.) Ryv.
- P. auriculaeformis 1844:194 (L, isotypes in BPI and S).
"Java". = Phellinus senex (Nees & Mont.) Imaz.
- P. blumei 1844:185 (L, isotypes in PC and BPI).
"Java, Zoll. no. 11" = Trametes menziesii (Berk.) Ryv.
- P. bonplandensis 1846:301 (PC, isotype in BPI).
"Am equatoriale". = Trametes membranaceus (Fr.) Kreisel.
- P. botryoides 1846:128 (S, isotype in BPI).
"Frag. orig. ex." (in Bresadola's hand) (North America).
= Globifomes graveolens (Schw.) Murr.

- P. bracypus 1846:127 (PC, isotype in BPI).
"Guadeloupe, M. L'Herminier, Février 1843". = Trametes marianna (Pers.) Ryv.
- P. callimorphus 1846:133 (PC).
"Voyage de M. Perville N.O. de Madagascar 1841."
This is an accepted species in Phellinus as P. callimorphus (Lév.) Ryv. For a description, see Ryvarden & Johansen (1980:145).
- P. callochrous 1844:181.
The type has not been found, and no type locality was given.
- P. candicans 1863:285 (PC, isotype in K).
"Tequendama (Colombia), alt. 2600 m, Sept 1856, Lindig no. 2922". = Trichaptum biformis (Fr. in Kl.) Ryv.
- P. candidulus 1846:301 (PC, isotype in K).
"Caesarodunum" (?) = Trametes cervina (Schw.) Bres. as already noted by D. Reid on the sheet.
- P. chryseus 1846:301 (PC).
"Nouvelle Grenade (Colombia), M. Justin Goudot 1844."
This is an accepted species in Phellinus as P. chryseus (Lév.) Ryv. For a description, see Ryvarden & Johansen (1980:151).
- P. cinerascens 1844:184.
The type has not been located (Java).
- P. cineraceus 1846:139.
The type has not been seen by me, but Bresadola (1916:223, as "cinereus") reports the species to be a synonym of Trametes hirsuta (Fr.) Pil.
- P. cohaerens 1846:132 (L, isotype in PC, BPI, FH and S).
"Zollinger no. 13" (Java). = Corioloopsis asper (Jungh.) Teng. as already indicated on the sheet in herb. PC by Bresadola.
- P. confertus 1844:187 (PC, isotype in BPI).
"In truncos Java". = Corioloopsis asper (Jungh.) Teng.
- P. connexus 1846:135 (PC).
"Brasil, Rio de Janeiro. M. Gaudichaud 1831-33".
As already indicated by Bresadola (1920:68) the type collection is mixed. One specimen is Corioloopsis cape-rata (Berk.) Murr. while another is Phellinus gilvus (Schw.) Pat. The description is so vague that it may cover both specimens, and thus, the name has to be rejected.
- P. coriaceus 1846:137.
The type has not been found (Nelli Gherry, India). The name is invalid as a homonym of P. coriaceus Endl. 1830.
- P. corrugatus 1846, 136 (L).
"Java". Invalid name, being a homonym of P. corrugatus Pers. 1826. The type of the invalid name is a specimen of Corioloopsis asper (Jungh.) Teng.
- P. convolutus 1844:186 (PC, isotypes in L, S, NY and BPI).
"Java, Junghuhn". = Trametes menziesii (Berk.) Ryv.
- P. cyathiformis 1844:181.
The type has not been found (Haiti).
- P. demidoffii 1842:92 (S, isotypes in BPI and K).
"Frag. specimen orig. leg. Demidoff, comm. Patouillard".
This is an accepted species in Pyrofomes as P. demidoffii (Lév.) Kotl. & Pouz., for a description, see Ryvarden (1978:399).

P. dermatodes, see Trametes dermatodes.

P. dilitatus 1844:184 (L, isotype in S).

"Java, ex reliqui Junghuhnii". = Trametes menziesii (Berk.) Ryv.

P. disciformis 1844:193.

The type has not been found (Mauritius).

P. dissectus 1846:139 (PC).

"Herbier du Chile, donné par M. Gay".

This is a Tyromyces species, and the following combination is proposed: Tyromyces dissectus (Lév.) Ryv. comb. nov. Basionym: Polyporus dissectus Lév. Ann. Sci. Nat. Ser. 3 Vol. 5:139, 1846.

Fruitbody pileate, dimidiate with a contracted base, about 2.5 x 2.5 cm, base about 1 cm wide, 2-6 mm thick, fragile. Pileus applanate, margin fimbriate to distinctly split or lobed, surface white to dirty light brown, smooth, azonate, finely velutinate to almost glabrous in parts and weakly wrinkled radially. Pore surface light brown, pores 4-5 per mm, entire to slightly incised, tubes 2 mm deep, white. Context ochraceous, 1 mm thick.

Hyphal system monomitic, hyphae 2-5 μ m wide and with clamps, partly gelatinized in 2.5 % KOH, in Melzer with walls about 0.5 μ m thick, sparingly branched. Some hyphae have a dense yellowish protoplasm, distinct both in KOH and Melzer, 2-6 μ m wide and with scattered clamps, these hyphae may be interpreted as oleiferous or gloeopleurous. Cystidia and basidia not seen. Spores subcylindrical to oblong ellipsoid, hyaline, thin-walled and non-amyloid, 4-5 x 2-2.5 μ m. As long as only the type is known, the macroscopical characteristics have to be used with care.

This taxon may be the same as T. floriformis (Quel.) Bond. & Sing. Their microscopical characteristics are identical and the fruitbodies are similar. T. floriformis as collected in Europe is usually more pure white, but the darker colour of the type of T. dissectus is certainly due to bad preservation after it was collected in Chile and then sent to France.

P. dozyanus 1848:123.

The type has not been found (Java).

P. elatus 1846:129.

The type has not been found (Guadeloupe).

P. extensus 1846:129 (PC).

"Guadeloupe, L'Herminier". This is an accepted species in Phellinus as P. extensus (Lév.) Pat. For a description, see Ryvarden & Johansen (1980:158).

P. fastuosus 1844:190 (L, isotype in BPI).

"Singapore". The species is accepted in Phellinus as P. fastuosus (Lév.) Ryv. For a description, see Ryvarden & Johansen (1980:159).

P. fuscus 1854:17 (L, isotypes in PC and K).

"Insul. Flores prope Balie". = Coriolopsis asper (Jungh.) Teng.

P. fuscus 1846:137 (PC, isotype in S & L).

"Zollinger no. 1454" (Java). The type is a specimen of what Berkeley later described as P. setiporus, a species

which usually is placed in Cyclomyces. The type species of the latter is C. fuscus Fr. 1833 making it illegitimate to transfer Léveillé's name to Cyclomyces. However, should anyone want to place the taxon behind Léveillé's and Berkeley's names in a genus other than Cyclomyces, then Léveillé's name has priority.

P. gaudichaudii 1846b:178 (PC, isotypes in S & BPI).

"Singapore 1839". = Trametes menziesii (Berk.) Ryv.

P. gayanus 1846:127 (PC).

"Chile, M. Cl. Gay. 1839". = Trametes marianna (Pers.) Ryv.

P. gibberulosus 1846:139 (PC, isotypes in K & FH).

"Guyana, Batara Reidweg, Wegel no. 568". = Trametes villosa (Hook.) Ryv.

P. gossypinus 1848:124 (PC, isotype in BPI).

"Ad truncos in Voges (France)". = Tyromyces caesius (Fr.) Murr.

P. guadeloupensis 1846:134 (PC, isotypes in S & NY).

"Guadeloupe, M. L'Herminier". = Fomitopsis supina (Fr.) Ryv. as already indicated by Bresadola on the original label.

P. haskarlii 1844:190 (L, isotypes in PC & BPI).

"Java". = Phellinus pectinatus (Kl.) Quel. as indicated on the sheet by Bresadola.

P. hasseltii 1844:187.

The type has not been found (Java).

P. heteromorphus 1846:123.

"Guyana". = Amauroderma schomburgkii (Mont. & Berk.) Torr., as already indicated by Furtado (1968:268).

P. hymenius 1863:283 (PC, isotype in K).

"Cune M. 1100 Juillet 1860" (Colombia). = Corioloopsis brunneo-leuca (Berk.) Ryv.

The type is unfortunately sterile. I have not seen the species fertile from the neotropics, and the dextrinoid reaction found in the skeletal hyphae of many African specimens was not seen in Léveillé's specimen. It may be that two macroscopically similar species are involved. Fertile material from the neotropics may shed some light on the problem.

P. inquinatus 1846:140 (PC, isotypes in FH & BPI).

"Perrottet, Nelli Gherry, India orientalis". = Trichaptum biformis (Fr. in Kl.) Ryv.

P. kickxianus 1848:122.

The type has not been found (America).

P. korthalsii 1844:190 (L).

"Sumatra, Korthals". = Phellinus senex (Nees & Mont.) Imaz.

P. lenis 1848:123 (PC, isotype in BPI).

"America, ad truncos". = Corioloopsis polyzona (Pers.) Ryv.

P. lenziteus 1854:17 (L, isotype in PC).

"Zollinger 975 (Sumatra)". = Lenzites acuta Berk.

P. leucomelas 1846:140.

The type has not been found. Invalid name, non P. leucomelas Pers. 1825.

P. lindigii 1863:283 (PC, isotype in K).

"Teguendama (Colombia) Canoas alt 2600. Sept. 1860".

- = Trametes cfr. modesta (Fr.) Ryv. The collection is fertile with spores 6.5-8 x 3-3.5 um. These measurements are in accordance with those of Fidalgo & Fidalgo (1968:27). The spores of the paleotropical taxon called T. modesta and macroscopically identical with that of the neotropics, are seemingly smaller. However, a richly fertile specimen has never been found and no sporeprint has been made.
- P. longipes 1846:124 (PC; isotype in BPI).
"Guyane française". This is an accepted species and the type of Haddowia, Stey. in the Ganodermataceae. For a description, see Ryvarden & Johansen (1980:93) with further references.
- P. macropus 1848:122.
The type has not been found (Java).
- P. mangiferae 1846:130 (PC).
"Voayge de M. Perville. N. East de Mahe (Tahiti) 1841". The species belongs in Ganoderma and its final status has to be decided when the genus has been revised.
- P. manubriatus 1854:17 (L).
"Zollinger 974" (Sumatra). = Microporus scopulosus (Berk.) Ryv.
- P. marchionicus 1846:300.
The type has not been found (Marquesas Islands).
- P. mastoporus 1846b:176 (PC).
"Singapore, Février 1832". This is a Ganoderma species in the G. lucidum complex. Its status has to be settled when the genus is revised.
- P. megaloma 1846:128.
The type has not been found (New York, USA). In American literature it is commonly assumed that this is the same as Ganoderma applanatum, see Overholts (1953:99). From the description it is very probably so.
- P. melaneus 1846:131 (L, isotypes in S & BPI).
"Zollinger no. 2085 (Java)". = Coriolopsis caperata (Berk.) Murr.
- P. melanaleucus 1846:141.
The type has not been found (Réunion).
- P. microcyclus 1844:188 (L, isotypes in PC, K, NY & BPI).
"Java". = Cyclomyces tabacinus (Mont.) Pat.
- P. microloma 1844:183.
The type has not been found (Manilla, The Philippines). At Kew there is a specimen: "Phillipines 1908, Curran, cum typo comparato" ink, Bresadola's hand. This is a specimen of Microporus affinis (Nees & Blume ex Fr.) Kunt. The specimen mentioned above is selected as neotype until authentic material is found.
- P. moritzianus 1846:130 (L).
"Zollinger 360 (Java)". = Trametes marianna (Pers.) Ryv.
- P. murinus 1844:185 (L, isotypes in PC and FH).
"Java". = Trametes menziesii (Berk.) Ryv.
- P. nephelodes 1846:125.
The type has not been found (Paramaribo, Brazil). In the Stockholm herbarium there is a specimen: "182 P. nephelodes Lév. = Polyp. flabelliform. Kl. transiens Java v. Höhnel" Bresadola's hand. The specimen is Microporus affinis (Blume et Nees ex Fr.) Kunt. However, as the genus Microporus is not known from South America,

the specimen from Java cannot be selected as neotype of P. nephelodes.

P. nordmannii 1842:93.

The type has not been found (Korbek, USSR). Fries (1874:581) refers the species to Heterobasidion annosum (Fr.) Bref. but it is not stated whether he had seen the type or based his suggested synonymy on the description alone. The latter is rather vague, but could apply to a resupinate specimen of H. annosum.

P. notopus 1844:194 (PC).

"Java". = Trametes modesta (Fr.) Ryv.

P. ostreatus 1846:127 (PC, isotype in BPI).

"Bresil, Rio de Janeiro, M. Gaudichaud 1831-33". = Trametes cubensis (Mont.) Sacc.

P. pala 1844:183.

The type has not been found (Surinam).

P. pectunculus 1846:138 (PC).

"Indes orientalis, Nelli-Gherry (India)". = Trametes versicolor (Fr.) Pil.

P. perpusillus 1844:191 (PC).

"Am. Boreal". = Fomitopsis scutellata (Schw.) Bond. & Sing.

P. phaeus 1846:132.

The type has not been found (Sri Lanka).

P. placopus 1846:124 (S).

"Java, Junghuhn". This is a Ganoderma sp. in the Ganoderma lucidum complex. Its status has to be settled when the genus is revised.

P. platypilus 1844:192.

The type has not been located (Java).

P. plumbeus 1846:136 (PC, isotype in S).

"Guadeloupe, M. L'Herminier". = Rigidoporus fusco-lineatus (Pers.) Ryv.

P. rhodophaeus 1844:190 (S).

"Java". This is an accepted species in Fomitopsis as F. rhodophaeus (Lév.) Imaz. For a description, see Ryvarden & Johansen (1980:340).

P. rigidus 1844:189 (L).

"Java, Zollinger no. 732". = Nigroporus durus (Jungh.) Ryv.

P. rudis 1846:133 (PC).

"Guadeloupe, M. L'Herminier". Invalid name, non P. rudis Berk. 1839. The type of the invalid name is Fomitopsis supinus (Fr.) Ryv.

P. rugulosus 1844:189 (L).

"No. 57" (Java) (leg. Junghuhn?). = Coriolopsis sanguinaria (Kl.) Teng.

P. sclerodermus 1846:129 (PC).

"Guadeloupe, L'Herminier". = Fomes fasciatus (Fr.) Cooke.

P. scleropodius 1846:123 (BPI).

"Ile Bourbon (Réunion)". = Lignosus sacer (Fr.) Ryv.

P. sericellus 1846:125.

The type has not been located (Sri Lanka).

P. sideroides 1844:182 (PC, isotype in K).

"Sumatra". This is an accepted species in Coltricia as C. sideroides (Lév.) Teng., for a description, see Ryvarden & Johansen (1980:112). Lévillé named also Coll.

- Zollinger No. 90 as P. sideroides, but this collection is a specimen of Phylloporia chrysa (Berk.) Ryv. Duplicates of this collection have been seen in herb. L, S, FH, PC and BPI.
- P. sordidus 1844:192 (PC, isotype in S).
"Guadeloupe, L'Herminier". = Trametes modesta (Fr.) Ryv.
- P. splendens 1844:187 (PC, isotype in L).
"Java". = Trametes modesta (Fr.) Ryv.
- P. spurcus 1846:135 (PC, isotype in S).
"Guadeloupe, L'Herminier". = Phellinus gilvus (Schw.) Pat.
- P. stevenii 1842:91.
The type has not been seen. Lloyd (1915:285) indicates it to be Ganoderma applanatum based on the original description and plate. Donk (1974:67) agreed, and I subscribe to the conclusion.
- P. subflavus 1846:300 (PC).
"Nouvelle Orleans, Salle". = T. versicolor (Fr.) Pil.
- P. swartzianus 1846:132.
The type has not been found (Jamaica).
- P. tegularis 1846:131 (PC, isotype in FH).
"Zollinger 2054 (Java)". = Trametes scabrosa (Pers.) Cunnighg.
- P. tenax 1846:139.
The type has not been located (Tequendama, Colombia).
- P. tener 1846:139 (PC).
"Guadeloupe, M. L'Herminier". = Trametes villosa (Hook.) Ryv.
- P. tenuissimus 1844:188.
The type has not been found (Java, leg. Korthals).
- P. testaceus 1846:126 (PC).
"Brasilia meridionalis, Dupré, 1842". This is a Ganoderma sp. in the G. lucidum complex. Its status has to be settled when the genus has been revised.
- P. trachodes 1844:192.
The type has not been found (Java).
- P. tricolor 1846:134.
The type has not been found (Bahia, Brazil).
- P. trigonus 1846:134 (PC).
"Zollinger No. 2069 (Java)". = Coriolopsis sanguinaria (Kl.) Teng.
- P. tristis 1846:126 (PC, isotype in FH).
"Zollinger No. 2035 (Java)". Invalid name, non P. tristis Pers. 1825. The type of Lévillé's invalid species is a specimen of Nigroporus vinosus (Berk.) Murr.
- P. unguiformis 1846:138 (PC, isotype in FH).
"Zollinger No. 1437, Tjiboya (Java)". = Rigidoporus microporus (Fr.) Overeem.
- P. vulneratus 1844:188 (PC).
"Ad truncos, Java". = Coriolopsis sanguinaria (Kl.) Teng.
- P. zollingerianus 1846:131 (PC, isotypes in S and FH).
"Zollinger 1386 (Java)". = Trichaptum byssogenus (Jungh.) Ryv.

SISTOTREMA Fr.

S. ochroleucum 1846:145 (PC).

"Bombay (India), Polydore Roux" = Lenzites acuta Berk.

TRAMETES Fr.

T. acuta 1844:196.

The type has not been located (Sumatra). Specimens determined by Bresadola in herb. S and BPI are all Coriolorp-sis strumosa (Fr.) Ryv., and this was probably also Léveillé's concept.

T. crassa 1844:197 (K).

"Madagascar". = Hexagonia hirta (Fr.) Fr.

T. dermatodes 1844:196 (PC).

"Voyage de M. Gaudichaud sur la Bointé. 1836-37. Manille (The Phillipines)". = Oxyporus cervino-gilvus (Jungh.) Ryv. as already indicated by Reid (1953:135).

T. incana 1844:196 (BPI).

"Manille (The Phillipines)". This is an accepted species in Trametes, for a description, see Ryvar den & Johansen (1980:565).

T. perrottetii 1844:195 (PC).

"Guyane Francaise M. Poilleau" (printed on the label).

"Java, M. Perrottet", unknown handwriting. The name, however, is written by Léveillé.

There has been a mixing of labels or facts. The type represents a South American taxon which has never been found or reported from the paleotropics. The type is apparently the same specimen which was used by Montagne and Berkeley when they described Trametes trichomallus in 1849. The latter is the type of Trichaptum Murr.

Trichaptum perrottetii (Lév.) Ryv.

Norw. J. Bot. 19:237, 1972.

Fruitbody applanate, sessile, semicircular to elongated shelf-like, mostly broadly-attached, usually not decurrent on the substratum, 5-15 cm long, 3-7 cm wide and to 8 mm thick (tomentum not measured), tough and flexible. Pileus with a dense, strigose to villose or hirsute layer of forked hairs, dark brown, becoming darker towards the base and more greyish towards the margin, azonate or weakly-zonate, up to 10 mm thick at the base. Margin entire and sharp. Pore surface snuff brown, pores angular to round, first entire and thin-walled, 2-3 per mm, in older specimens often incised, coalescing and in parts sinuous to daedaleoid, in the latter case up to 2 mm wide and several mm long. Tubes deep brown, 2-5 mm deep. Context very thin, 0.1-0.4 mm, brown to dark ochraceous. Hyphal system dimittic, generative hyphae thin-walled, hyaline and with clamps, 2-4 μ m wide. Skeletal hyphae abundant, thick-walled to solid, mostly yellowish to light brown, 3-5 μ m wide. Cystidia abundant in the hymenium, clavate to ventricose with a tapering apex, smooth or with an apical crown of crystals, 10-18 μ m long, slightly projecting, basidia clavate 12-15 x 4-6 μ m and with 4 sterigmata. Spores subcylindrical to ob-

long ellipsoid, smooth, thin-walled and non-amyloid, 5-7 x 2-3(3.5) um. On deciduous wood.

Specimens have been examined from USA, Cuba, British Honduras, Guatemala, Costa Rica, Colombia, Brazil, French Guyana and Bolivia.

T. vittata 1844:196.

The type has not been found (Sumatra).

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The staff of the herbaria in Stockholm, Leiden, Paris, Kew, New York, Harvard and Beltsville are most kindly thanked for their cooperation during my type studies. The Norwegian agency for International Development has given financial support for which I am very grateful. R. Winter has suggested improvements in the English text.

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THE PUBLICATION DATE OF ARENDHOLZ'S THESIS ON LEAF-INHABITING HELOTIALES

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SUMMARY

July 7, 1979 is shown to be the date of effective publication of W.-R. Arendholz's doctorate thesis, "Morphologisch-taxonomische Untersuchungen an blattbewohnenden Ascomyceten aus der Ordnung der Helotiales," and the date of valid publication of the new taxa and new combinations proposed therein.

The status of publication of W.-R. Arendholz's Ph.D. thesis, entitled "Morphologisch-taxonomische Untersuchungen an blattbewohnenden Ascomyceten aus der Ordnung der Helotiales" was assumed to be "unpublished" by Korf and Kohn (1980). In order to protect the status of Arendholz's new names, they reported one as "an as yet unpublished new species referred to the genus" They had received in the mails from Arendholz a copy of his thesis (Arendholz, 1979), a soft-bound volume having the appearance of being some sort of photo-copy of the original typewritten thesis on deposit with the Universität Hamburg.

The purpose of this note is to call attention to the fact that over 180 copies of this thesis were actually distributed, and that in no way can the new names therein be ignored or treated as not validly published. On July 7, 1979, 139 copies were mailed by Arendholz to the

Staats- und Universitätsbibliothek, Hamburg, which subsequently distributed them to other libraries, and that date is held here to constitute the date of distribution constituting "effective publication" under the International Code of Botanical Nomenclature (Stafleu et al., 1978), Article 29.1. Twenty further copies were sent to the library of the Institut für Allgemeine Botanik der Universität Hamburg, and another 23 as gifts to scientists in different countries. Approximately 20 additional copies are still in Arendholz's possession out of a photo-offset lithography press run of some 200 copies.

While it seems unfortunate that the copies that were distributed carry no indication of the exact date of distribution, nor even that the volume constitutes a publication in the sense of the Code of Nomenclature, recent articles on effective publication under Articles 29-31 of the Code leave no doubt in our mind that the thesis was effectively published. The four articles in *Taxon* 29(4), the August 1980 issue, support our position [Brummit (1980); Hara and Eichler (1980); Nicholson (1980); Weresub and McNeil (1980)].

Since the thesis is printed by photo-offset lithography, it must apparently be accepted as printed matter (Art. 29): "... it may safely be assumed that matter produced by a conventional typewriter, consisting of one top copy plus one to several carbon copies, is not considered to be printed. On the other hand, copy originally produced on a typewriter and then reproduced by offset lithography (as for example the important taxonomic journal Mitteilungen der Botanischen Staatssammlung München) must certainly be accepted as printed matter" (Brummitt, 1980: 477-478).

The Code (Art. 29.1) requires that the printed matter be distributed "to the general public or at least to botanical institutions with libraries accessible to botanists generally." Neither the number of copies, nor the number of libraries (nor even that they be located in more than one country) is stipulated in the Code, "and presumably in an extreme case a printed work distributed to as few as two libraries should be taken as effectively published" (Brummitt, 1980: 479).

In the case in point, the requirements for effective publication would have been fulfilled by mere deposition of one copy of the printed thesis in the "Staats- und Universitätsbibliothek" and one in the library of the "In-

stitut für Allgemeine Botanik." As noted above, many more than two copies were distributed to several libraries and scientists worldwide. Nicholson's (1980) key similarly keys out this thesis as "effectively published."

Establishing an exact date of publication of any printed work is critical for nomenclatural purposes. For journals, the practice is to use the actual date of mailing of each issue as publication date [this is the procedure with MYCOTAXON, as it is also with MYCOLOGIA (C.T. Rogerson, Managing Editor, pers. comm.)]. Similarly, commercial books are published on the date they are "actually offered and available for purchase, not that on which they actually reach the public or libraries" (Brummitt, 1980: 480). Since the bulk of the copies of Arendholz's printed thesis were mailed on July 7, 1979 to the Staats- und Universitätsbibliothek, Hamburg, that date should now be accepted as the date of publication of the thesis for all nomenclatural purposes.

Various proposals by Hara and Eichler (1980) and by McNeil (in Weresub and McNeil, 1980) focus on the kind of problem presented by Arendholz's thesis with a view to eliminating any problems in the future, with worthwhile suggestions for requiring an indication of intention to offer material as "published" under the provisions of the Code, requirements for deposition in a specific number of (named) institutions, and indication of actual date of publication (distribution). Such proposals will be useful in the future, but in any case will not be retroactive, but apply from 1982 onward if they are adopted at the International Botanical Congress in Sydney later this year in the forms submitted.

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AN ALTERNATIVE VIEW OF CERTAIN TAXONOMIC CRITERIA USED IN THE ENTOMOPHTHORALES (ZYGOMYCETES)

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SUMMARY

Long-standing unresolved controversies about the various generic classifications of the entomopathogenic Entomophthorales require a review of the validity and the relative weights of the morphological and other taxonomic criteria used to construct a useful classification for this group. Major morphological criteria used in the Batko and Remaudière classifications are evaluated here in a survey of the taxonomic distributions of these characters and the significance of the irregularities in their distributions.

The number and nature of nuclei in the primary spores, the branching of sporophores, and the mode of discharge of primary spores are regarded as the major characters suitable for delimiting genera. The presence of rhizoids and (to a limited extent) the shapes of primary spores are significant secondary generic characters. The presence of cystidia, mode of resting spore formation, and presence of capillary secondary sporophores are not regarded as characters of generic importance. The conidial or sporangial nature of primary spores is discussed, as are difficulties in the interpretation and applications of the terms *zygospore* and *azygospore*.

The applications of the generic names used in the Batko and Remaudière classifications are reviewed in light of the findings reported here. The validity of both *Strongwellsea* and *Entomophaga* is upheld, and the sense of *Erynia* is restricted to those species having branched sporophores and uninucleate bitunicate primary spores which do not form secondary spores on capillary sporophores. A nomenclatural problem is noted which indicates the synonymy of *Zoophthora* and *Erynia* if these genera are not retained as separate as proposed in the Remaudière classification. *Tabanomyces* is regarded as a synonym of the nematophagous genus *Meristacrum*.

INTRODUCTION

A far-reaching revision of the entomopathogenic genera of the Entomophthorales (Zygomycetes) proposed by Batko (1964a-e, 1966; Batko and Weiser 1965) has spawned considerable controversy among the students of these fungi. Even though there is now wide agreement that it is unsatisfactory to continue classifying approximately 150 diverse species in the single genus *Entomophthora* Fresenius, there has been no agreement regarding the appropriateness of the criteria used by Batko to restrict *Entomophthora* and to establish segregate genera or of the circumscriptions of these genera.

King and Humber (1981) offer a basic reconsideration of the major morphological characters of this group and discuss their taxonomic significance. While King and Humber evaluate the basic strengths and weaknesses of the Batko classification, they consider that no definitive reworking of the genera would be appropriate without a final resolution of the long-standing problem of delimiting *Conidiobolus* Brefeld from *Entomophthora* sensu lato.

More recently, however, a major reworking of Batko's generic classification by Remaudière and Hennebert (1980) and Remaudière and Keller (1980) — hereafter referred to as the Remaudière classification — resulted in generic assignments for some species which led to no manifest increase in the biological homogeneity of the recognized genera. Many taxonomic decisions in the Remaudière classification, and particularly those regarding the validity and

importance of various characters used to construct classifications of these fungi, were presented without explanation and, in turn, have triggered further controversy and alternate views on classification (Ben-Ze'ev and Kenneth 1981a; Humber 1981a).

The time is appropriate to examine thoroughly those morphological and developmental characters which have been employed in the modern classifications proposed by Batko and by Remaudière and his colleagues. The values of these criteria are examined in the contexts of the available classification systems. Indeed, a sense of the relative values of their chosen criteria does emerge from a consideration of the major schemes of entomophthoralean genera and of the internal inconsistencies generated by the selection and ranking of the criteria accepted as valid in each of these generic schemes.

Despite its undisputed desirability, no general agreement regarding the generic classification of the Entomophthorales will be possible until the morphological and other criteria used in the past or being proposed now have been critically reviewed, evaluated, and agreed upon. This paper is the first in a series discussing the merits and weaknesses of the Batko, Remaudière, and other emergent classifications (*e.g.*, Ben-Ze'ev and Kenneth 1981a) in an attempt to find rational bases on which to build a generic scheme which (1) avoids the inconsistencies of the previous classifications of the Entomophthorales, (2) can accurately predict the natures of characters or aspects of the host-pathogen relationship, and (3) reflects what appear to be the evolutionary relationships among these fungi.

TAXONOMIC CRITERIA: ANALYSIS AND CONCLUSIONS

PRIMARY SPORES

Primary Spores: Conidia or Sporangia?

Throughout this paper I refer to *primary spores* or *spores* rather than to the more conventional *conidium* or the less common *sporangium* or *sporangiole*. *Conidium* and *sporangium* are terms referring to specific developmental patterns; their use should be for purposes of accuracy and must be backed by sufficient demonstration of the mode of

sporogenesis. The reference to any asexual propagules in the Zygomycetes as *conidia* has been questioned by the students of the Hyphomycetes and Coelomycetes, although there is no reason to believe *a priori* that conidia cannot be formed by these or any other lower fungi. If appropriate studies of the origins and fates of the various wall layers of spores (and sporophores) during sporogenesis and germination prove that, indeed, they are conidia (or sporangia), then these propagules should be referred to as such. Similar studies are also needed for the panoply of secondary spore types produced in the Entomophthorales.

Spores containing single nuclei and borne on branched (or simple, in *Strongwellsea* Batko & Weiser emend. Humber) sporophores have caused much interpretational difficulty. These spores have a layer which, in liquid, lifts away from the spore surface and balloons outward. This has been regarded variously as a gelatinous coat (e.g., *Entomophthora gloeospora* Vuillemin 1886) or, more frequently, as evidence for the sporangial nature of these fungi by regarding the detached layer as a sporangial wall enclosing a single sporangiospore (Thaxter 1888; Strong et al. 1960; Batko 1964b, 1974). This type of primary spore seems to be entirely restricted to the species of *Zoophthora* Batko, *Erynia* Nowakowski, and *Strongwellsea*. Kenneth (1977) claims that the outer wall of *Entomophthora turbinata* Kenneth [= *Neozygites turbinata* (Kenn.) Rem. & Keller, a generic assignment which remains debatable] lifts away but admits (personal communication) that this feature is doubtful and requires further investigation. If the spore of *E. turbinata* truly were bitunicate (with a separable outer wall layer), it would be the only instance of this character in a species with multinucleate spores.

Humber (1975; King and Humber 1981; Roberts and Humber 1981) notes ultrastructural evidence that the spore of *Strongwellsea magna* Humber, which has a separable outer wall layer, cannot be a monosporic sporangiole. The spore wall has no indigenous (sporangiospore) layer that is not shared with the sporophore itself. The separable layer is not a distinct (sporangial) wall, but represents only a separation of the outermost layer of the sporophore wall, all layers of which are continuous over the entire spore surface except for the papilla — which is covered only by an extension of the inner wall layer of the spore and sporophore (Humber 1975). This spore might be a true conidium, but this cannot be confirmed without also determining

the fates of the wall layers during germination (see Dykstra 1974).

The only evidence for the occurrence of sporangia in the Entomophthorales comes from several species of *Basidiobolus* Eidam, in which there are numerous instances when the cytoplasm inside a cell (usually but not always a spore) may be divided into several independent uninucleate units; *B. microsporus* Benjamin (1962) provides the most striking examples of this behavior.

I use the terms *unitunicate* and *bitunicate* introduced by Remaudière and Hennebert (1980) and Remaudière and Keller (1980) to refer to spores on which a separable outer wall layer is either absent or present, respectively. These terms are used almost exclusively within mycology to refer to the morphology (and function) of asci in the Ascomycotina, and constitute a primary character dividing the (bitunicate) Loculoascomycetes from all other (unitunicate) classes of this subdivision (Ainsworth et al. 1973). The use of these terms, however, should be generally accepted for the Entomophthorales since they (1) are convenient, (2) respect their etymological derivations, (3) should not be in any way confused with their applications for the ascomycetes, and (4) in the case of bitunicate spores, would help suppress the notion that such spores are sporangial in nature.

Morphology

The variety of shapes of primary spores in the Entomophthorales has been the source of much attention, particularly among those who have included nearly all entomopathogens of this group in the single genus *Entomophthora*. A series of artificial species groupings based on spore morphology have been proposed primarily to aid species identification (Lakon 1919; Hutchison 1963; Gustafsson 1965; MacLeod and Müller-Kögler 1970, 1973; MacLeod et al. 1976; Waterhouse 1975; Zimmermann 1978). None of these authors, however, intended their groupings to have taxonomic value. Remaudière and Keller (1980), however, chose the morphology of primary spores to be the main criterion for their generic classification.

Several genera do have characters of spore morphology which are unique to themselves. One of the strongest and most valid criticisms of the Batko classification, in fact,

was its separation of the species having campanulate spores (with relatively flat bases and a strongly apiculate apex) into *Entomophthora* sensu stricto and *Culicicola* Nieuwland depending upon the absence or presence of rhizoids. Remaudière and Keller (1980) rejected the significance of rhizoids for defining the genera, and redefined *Entomophthora* to include only species with this distinctive campanulate spore shape and associated mode of spore discharge (see MacLeod et al. 1976; Samson et al. 1979).

The spores of *Triplosporium* (Thax.) Batko (*nom. gen. conserv. prop.*, see Humber et al. 1981; = *Neozygites* Witlaczil) are variable in shape, but most often are round to slightly elongate and have a truncate rather than a conical or rounded papilla. As circumscribed by Batko (1964c), the spores of *Culicicola* species could be either campanulate (like those of *E. muscae* (Cohn) Fres.) or obpyriform with a smooth apex and rounded papilla; species with the former type were restored to *Entomophthora* s.str. while those with the latter type of spores were transferred to *Conidiobolus* by Remaudière and Keller (1980). King (1976b, 1977) regards *Conidiobolus* species to have round to pyriform primary spores, and notes the similarity of several entomopathogenic *Entomophthora* (sensu lato) species with *Conidiobolus*. Remaudière and Keller (1980) extended King's concept of *Conidiobolus* by transferring all entomopathogenic species with rounded to pyriform spores to that genus. The species of *Entomophaga* Batko have pyriform spores with a rounded apex and closely resemble the spores of *Conidiobolus* species. These two genera are unambiguously differentiated, however, by both the morphology and stain reactions of their nuclei (Humber 1981b).

The primary spores of *Zoophthora* Batko sensu Remaudière and Hennebert (1980) (= *Zoophthora* subg. *Zoophthora* Batko sensu Ben-Ze'ev and Kenneth 1981a) are rather cylindrical and usually taper apically to a blunt cone or a poorly defined point; just above the conical papilla, the spore flares to a very slight (but not always apparent) shoulder. The spores of *Erynia* Nowakowski species (see Remaudière and Hennebert 1980) display the greatest shape variation of any genus in the Remaudière classification. They range from elongate fusiform with a marked curvature and a shallow rounded papilla to somewhat cylindrical to ovoid or fusiform spores with rounded apices and papillae canted away from the spore axis. The spores of *Strongwellsea* species fit well within the range of morphological

variation found in *Erynia*, a fact which seems to have motivated Remaudière and Keller (1980) to merge this genus into *Erynia*. The spores of *Empusa caroliniana* Thaxter, which Remaudière and Hennebert (1980) transferred to *Erynia*, are obovoid to pyriform, a shape corresponding more with those of *Erynia* spores than with any other genus.

Karyology: Number and Type of Nuclei

Early studies noting the taxonomic importance of entomophthoralean nuclei have been largely ignored (Vuillemin 1895; Cavara 1899a-b), so that no surpassing importance was attached to the number of nuclei in primary spores until the proposition of Batko's classification (1964a-d, 1966; Batko and Weiser 1965), which necessarily drew upon the observations of earlier workers regarding this cytological detail.

One of the most significant characters used by Batko (1964b) to establish *Zoophthora* was that of uninucleate primary spores, a character which is readily detectable and seems to be correlated well with the other characters of this genus. *Strongwellsea* Batko & Weiser (1965) was also described as having uninucleate spores but was clearly distinguished from *Zoophthora* by its simple sporophores and unique habit. *Erynia* Nowak. was split from *Zoophthora* by Remaudière and Hennebert (1980) and is circumscribed so as to allow multinucleate primary spores even though *E. caroliniana* is the only such species included in *Erynia* by Remaudière and his colleagues. As discussed at length below, the morphological characters of this species do not appear to belong in this genus, and its probable misclassification is clearly indicated by its status as the only species with multinucleate spores in this large grouping of species with uninucleate spores.

As is the case with branched sporophores (see below), there is a very low but natural incidence of abnormally developed primary spores. The spores of *Zoophthora*, *Erynia*, and *Strongwellsea* receive very nearly the entire cytoplasmic contents of the terminal cell of the sporophore. In species with branched sporophores there are, occasionally, larger than usual volumes of cytoplasm containing two (or three) nuclei which are isolated in a branch by its basal septum. The spore subsequently formed from this branch will be both markedly larger than normal and contain the extra nucleus or nuclei. Such exceptionally rare

spores are clearly aberrant and cannot be used to invalidate the significance of uninucleate spores. Similarly, they do not justify the inclusion in *Erynia* (all of whose other species have uninucleate primary spores) of *E. caroliniana* (whose spores are always multinucleate and whose other major morphological characters do not match with the other species of *Erynia*).

Three genera are noted by Batko (1964a-b, 1974) to have plurinucleate spores. The spores of several *Entomophthora* species (sensu Remaudière and Keller 1980) contain 4-6 nuclei; those of *E. culicis* (Braun) Fres. are characteristically binucleate (or occasionally trinucleate) while those of *E. muscae* (Cohn) Fres. contain 5-8 nuclei. *Triplosporium* (= *Neozygites*) was described by Batko (1964b) to have quadrinucleate spores. Of the eight species included in *Neozygites* by Remaudière and Keller (1980), seven have 4-nucleate spores; *Entomophthora turbinata* Kenneth (1977) has spores with 5-7 or more nuclei. This deviation from the quadrinucleate condition of all other species in this genus indicates that close scrutiny must be given to the generic assignment of this relatively little studied fungus. On first examination, the only marked similarities of *E. turbinata* with the other species of *Triplosporium* are its simple sporophores and ovoid resting spores with a jet black epispore. Batko (1974) incorrectly indicated that the primary spores of *Massospora* Peck are binucleate; in fact, they vary from 1-6 (or more) nuclei, but most commonly contain 2-3 nuclei (Soper 1974, 1981).

The number of nuclei in spores of *Culicicola* species (as originally described by Batko 1964c-d) varied from pluri- to multinucleate, but really represent a bimodal distribution with either a few nuclei or many in the primary spores. This bimodal variation perfectly parallels the dispersal of *Culicicola* species for other reasons to either *Entomophthora* s.str. (plurinucleate) or *Conidiobolus* (multinucleate).

The taxonomic importance of nuclear morphology has not been appreciated until now even though some authors such as Batko (1964a-b, 1966; Batko and Weiser 1965) have drawn attention to the staining of nuclei in lactophenol/cotton or aniline blue or other stains, or to their general size and appearance, but never realized their potential taxonomic value. The large nuclei of *Basidiobolus* species, for example, have been the subjects of numerous cytological studies.

Humber (1981b) finds that nuclei of *Conidiobolus* species (sensu King 1976b) have relatively small nuclei with a prominent central nucleolus but no obvious heterochromatin; these nuclei remain unstained or only very weakly differentiated in lactophenol/aniline blue, aceto-orcein, or a number of other nuclear stains. The species of *Entomophthora* and most of the entomopathogenic genera have large nuclei with prominent heterochromatin but no large central nucleolus; these nuclei may be differentiated in lactophenol/aniline blue and are strongly and rapidly differentiated in aceto-orcein and other nuclear stains. This karyological criterion has proven to be of inestimable value for diagnostic work, particularly for distinguishing species of *Conidiobolus* from *Entomophaga* (whose primary spores are nearly identical in shape) or other genera with readily differentiated nuclei, and in determining incidences of double infections in insects when one of the fungi is a species of *Conidiobolus*.

Taxonomic Significance of Primary Spores

Batko's recognition of the generic significance of the number of nuclei (one, a few, or many) in primary spores has been widely embraced. The Remaudière classification, however, necessarily disregarded this character in order to place the greatest generic value on the shapes of primary spores. As will be made more apparent below, this dependence upon spore shapes is misplaced and leads to what seem to be several misclassifications which offer no improvement over earlier artificial schemes based on spore shapes and proposed without taxonomic or phylogenetic significance.

Nuclear cytology affords the only consistent and readily observed criteria which adequately distinguish the species of *Conidiobolus* from *Entomophthora* and its segregates (Humber 1981b). These criteria also uphold the validity of *Entomophaga* Batko for *E. grylli* (Fres.) Batko and several other species. *Entomophthora obscura* Hall & Dunn and *E. thaxteriana* (Petch) Hall & Bell were assigned to *Entomophaga* (Batko 1964d) but are now regarded as synonyms (see below); nuclear cytology confirms the transfer of *Entomophthora obscura* to *Conidiobolus* by Remaudière and Keller (1980).

The unitunicate or bitunicate nature of primary spores appears to be perfectly correlated with the nuclear number. With the sole unconfirmed exception of *E. turbinata*, all

known bitunicate spores also contain a single large nucleus. Except for two species of *Massospora* in which uninucleate primary spores can occur (Soper 1974), all uninucleate spores are pluri- or multinucleate. It is possible that this distinction in wall structures of primary spores may assume a greater significance as more entomophthoralean fungi are found and described.

Among the characters related to primary spores in the Entomophthorales, then, generic value should be accorded to at least the number and nature of the nuclei in the spore. The shape and uni- or bitunicate nature of the primary spore should always be noted. However, while various states of these characters may coincide perfectly with the generic limits defined by other criteria, no reason is apparent at this time why either of them should be accorded generic value equal to nuclear number and morphology.

PRIMARY SPOROPHORES

Morphology and Branching

The branching of sporophores was one of the first criteria used to divide the entomopathogenic Entomophthorales into two genera (Brefeld 1873, 1877; Nowakowski 1883), and has continued to be one of the leading characters used by Batko (1964a-c, 1974; Batko and Weiser 1965) to construct a contemporary classification. The historical significance of sporophore branching as a character with generic importance demands careful review of both the reasons for and the effects of its de-emphasis by Remaudière and Keller (1980). It is necessary first to examine the systematic distribution of simple and branched sporophores and then to evaluate the exceptions to the normal state in each genus.

For any practical purpose, the sporophores are always simple in *Entomophthora* (although in *E. culicis* several sporophores may arise from each hyphal body, thus giving an impression of branching), *Massospora*, *Triplosporium*, and *Completozia* Lohde, a parasite of fern prothallia. *Tabanomyces* Couch et al. (1979; from tabanid fly larvae) and *Meristacrum* Drechsler (1940; from nematodes) produce an upright, unbranched sporogenous hypha which becomes septate and produces forcibly discharged spores on lateral (or terminal) papillae. *Ballocephala* Drechsler (1951; from tardi-

grades) also produces an upright, unbranched sporogenous hypha on which small lateral cells are formed from which the spores are budded and forcibly discharged. Sporophores in species of *Ancylistes* Pfitzer (Berdan 1938; from desmid algae) and *Basidiobolus* (Drechsler 1964) may occasionally branch; those of the 27 *Conidiobolus* species recognized by King (1976b, 1977) also may, on rare occasion, be bi- or trifurcate (King and Humber 1981). Among the species transferred to *Conidiobolus* by Remaudière and Keller (1980) the sporophores of *Empusa apiculata* Thaxter and *E. major* Thaxter may show some branching on rare occasion (see Thaxter 1888, Fig. 63), but are almost always simple. There is no significant branching in any *Entomophaga* species.

All species of *Zoophthora* s.str. (Remaudière and Hennebert 1980; Keller 1980; Ben-Ze'ev and Kenneth 1981a-b) have digitately branched sporophores with the possible exception of *Z. crassitunicata* Keller (1980) in which the sporophores appear in histological sections to be mostly simple.* All but three *Erynia* species (Remaudière and Hennebert 1980; Remaudière and Keller 1980; Kramer 1981) have digitately branched sporophores.

Within the bounds of the usual biological variation, it is not surprising that some species with simple sporophores occasionally have a small number of sporophores that show a low order of branching. Whatever branching occurs in these genera is usually basal rather than apical as in *Zoophthora* and *Erynia* (where the apical, digitate branching pattern usually results in the formation of a tightly interwoven hymenium). The infrequent occurrences of branched sporophores in genera characterized by simple sporophores or of occasional simple sporophores in genera characterized by branched sporophores in no way indicates that this character should be de-emphasized or discarded.

The sporophores of *Zoophthora* and *Erynia* (apart from the three exceptions discussed below) do exhibit a variable degree of branching. Unbranched sporophores might occa-

* Keller's (1980) characterization of the sporophores of *Z. crassitunicata* as mostly simple seems to be based entirely on histological sections and should, therefore, be accepted as provisional until the publication of micrographs of carefully dissected pieces of hymenium showing whole sporophores of this species.

sionally be found on specimens from this large group of species in which nearly all sporophores are digitately branched. Similarly, species such as *Z. crassitunicata* may exist in which (digitately?) branched sporophores are less numerous than simple ones, but the affinities of these species with *Zoophthora* or *Erynia* will always be apparent from the natures of those branched sporophores, their primary spores, and other associated characters.

The natural degree of plasticity in the branching of sporophores in species of *Zoophthora* and *Erynia* does not imply, however, any real probability of finding species such as *Z. radicans* (Bref.) Batko (= *Entomophthora sphaerosperma* Fres.) or *Erynia neoaphidis* Rem. & Henneb. (= *Entomophthora aphidis* Hoffman sensu Thaxter) which form obviously digitate sporophores to yield an exceptional specimen in which a majority (much less all) of the sporophores are simple. It is even less feasible that a strain of such a species exists which produces branched sporophores when infecting most hosts but which forms only simple ones on one or more other hosts.

The uniform occurrence of digitate sporophores in *Zoophthora* sensu Remaudière & Hennebert — including those (digitately?) branched sporophores of *Z. crassitunicata* — leaves only the two species of *Strongwellsea* and *E. caroliniana*, the three exceptional species placed in *Erynia* by Remaudière and his colleagues, to account for the de-emphasis by Remaudière and Keller (1980) of sporophore branching as a generic criterion.

It is important to examine why Remaudière and Keller (1980) supposed that a significant probability exists for finding branched sporophores in the two *Strongwellsea* species. It is these fungi which seem to form the basis for their de-emphasis of sporophore branching. The statement that branched sporophores are "not always present" in *Strongwellsea*, their primary justification for their opinion, seems to draw solely upon Strong et al. (1960) who note that the sporophores are "rarely branched" but who also admit the difficulty of tracing complete structures in their prepared slides. Batko and Weiser (1965) used some of these slides and correctly described the sporophores of *S. castrans* Batko & Weiser (1965) to be unbranched; they neither illustrated nor noted any exception to this simple state. Humber (1975, 1976) also found the sporophores of *S. magna* Humber to be *always* simple, and included this

characterization in the emended generic description of *Strongwellsea*. The uniformly unbranched nature of sporophores in this genus reflects the means by which additional sporogenous hyphae interpolate themselves into and thus continually expand the surface area of the fungal ball in the host fly's abdomen (Humber 1975).

It has already been noted that *E. caroliniana* (with multinucleate and unitunicate spores, simple sporophores, and lack of rhizoids or cystidia) shares no major characteristics with *Erynia* species except for spore shape; the simple sporophores of this species do not constitute any significant exception to the common state of digitately branched sporophores in *Erynia*. However, some minor degree of branching of *E. caroliniana* sporophores may occur *inside* the host body, but the emergent sporophores are uniformly unbranched on the surface of affected craneflies (Thaxter 1888). Both Giard (1888, as *E. arrenoctona* Giard) and Keller (1978) also find the sporophores of *E. caroliniana* to be simple.

Several aspects of sporophore morphology — the presence and pattern of branching and the overall shape of the sporogenous cell or sporophore apex — should be considered to be important, but generic significance has usually been placed only on whether a sporophore is simple or branched. The sporophores of all *Entomophthora* species (in the strict sense) are markedly swollen and clavate below the spore, but this is the only genus in which this character is uniformly present. There is a less consistent tendency to produce clavate sporophores in species of *Zoophthora* or *Erynia*. In most other species and genera of the order, the apical portion of the sporophore is relatively cylindrical with only a slight tendency to become clavate.

Srinivasan et al. (1964) proposed to separate *Conidiobolus* from *Entomophthora* on the basis of sporophore shape. They suggested that *Conidiobolus* species have micronemous (hypha-like, indeterminate) sporophores while those of *Entomophthora* are macronemous (thicker than and distinctly differentiated from the vegetative hyphae, and of determinate length). This criterion was devised to separate *Conidiobolus* species from *Entomophthora muscae* (the type of its genus) and paid little attention to the many other species in *Entomophthora* sensu lato. In the more modern taxonomic systems considered here, the criterion proposed by Srinivasan et al. (1964) is applicable only to the few spe-

cies of *Entomophthora* s.str. (Remaudière and Keller 1980); it is also untenable, however, since *Conidiobolus adiaeretus* Drechsler (see King and Humber 1981, Fig. 4f) and some other *Conidiobolus* species have markedly differentiated (macronemous) sporophores.

Several genera can, in fact, be distinguished in part by the exact morphology of the sporophore. *Basidiobolus* species have a more or less prominent swelling of the sporophore immediately below the spore, and the sporophore is markedly narrowed at the point where the spore is attached. Species of *Meristacrum*, *Tabanomyces*, and *Ballocephala* form several spores on each unbranched erect sporogenous hypha. The mode of spore production in *Meristacrum* and *Tabanomyces* is identical, and these genera are here considered to be synonymous (see below). The spores of *Ballocephala* are produced on globose to elongate lateral cells produced sympodially at the growing apex of the sporogenous hypha (Drechsler 1951; Richardson 1970; Pohlád and Bernard 1978). Three species placed in *Conidiobolus* by Remaudière and Keller (1980) — *E. apiculata*, *E. major*, and *E. papillata* Thaxter — differ from all other species of that genus (in the sense of King 1976b, 1977) by having an extended and neck-like narrowing of the sporophore apex; this morphological difference suggests that all characters of these fungi should be compared carefully with those of *Conidiobolus* before accepting their placement in this particular genus.

Taxonomic Significance of Primary Sporophores

The presence of branched or simple sporophores was regarded as a primary generic character in the Batko classification but was de-emphasized (and effectively rejected) in the Remaudière classification, seemingly to justify the synonymy of *Strongwellsea* (in which the sporophores are always simple) with *Erynia* (in which the usual case is for sporophores to be digitately branched). This de-emphasis and the elevation of spore morphology to a primary generic character by Remaudière and Keller (1980) further confused the classification by placing *E. caroliniana* (with simple sporophores) in *Erynia* and *Entomophthora carpentieri* Giard (apparently with branched sporophores, see below) in *Conidiobolus*.

As has been advocated by Ben-Ze'ev and Kenneth (1981a) sporophore branching should be re-instituted as a prime

generic character. This restoration sweeps aside the apparent misclassifications of the Remaudière scheme by returning *Strongwellsea* to its rightful status as a separate genus (Humber 1975, 1976, 1981a), and leaving the placements of *E. caroliniana* and *E. carpentieri* undetermined and in need of further study.

For the entomopathogenic genera and *Conidiobolus*, the most taxonomically significant aspect of primary sporophores is only whether they are simple or branched. There does seem to be a significant difference, however, between a low level of basal branching (as in *Conidiobolus* and *E. caroliniana*) and the apically digitate branching restricted to the species of *Zoophthora* and *Erynia*. The means by which the primary spore or spores are produced on the sporophore and the morphology of the sporophore apex also assume taxonomic significance when considering the complete spectrum of entomophthoralean genera.

MODE OF FORCIBLE DISCHARGE OF PRIMARY SPORES

Three distinct mechanisms for the forcible discharge of primary spores are known from the Entomophthorales. Two of these mechanisms appear to be entirely restricted to single clearly defined genera while the predominant mechanism operates in nearly all the remaining genera. Primary spores are forcibly discharged in all generally recognized genera of this order except for *Massospora*, in which spores and hyphal bodies are passively dispersed from the disintegrating abdomens of the living host cicadas. All of the diverse types of secondary spores are also forcibly discharged except for those capillispores (see below) produced on long, narrow capillary sporophores.

All species of *Entomophthora* sensu stricto (Remaudière and Keller 1980) have campanulate and apiculate primary spores which are forcibly discharged on a stream of cytoplasm and vacuolar sap squirted from the sporophore as the wall layer(s) securing the primary spore to the sporophore break. The spore itself has a flat or slightly convex base rather than a distinct papilla, and appears to play no active role in the discharge process. Spores discharged in this manner are accompanied throughout their trajectories

by a voluminous drop of the protoplasm which serves to affix the spore to the surface on which it lands, and forms a characteristic radially striate corona around the primary spore upon drying.

The primary spores of all species of the non-entomogenous genus *Basidiobolus* are borne on sporophores with a pronounced swelling immediately below the primary spore. Discharge occurs with a circumscissile rupture of the lower portion of the swelling; the spore and swollen upper portion of the sporophore fly away as the hydrostatic pressure in the upper portion of the sporophore is released backwards like a small rocket (Ingold 1934). Ingold's rocket analogy is strengthened by the fact that it is often possible to obtain the firing of a "second stage" as the small conical projection of the *Basidiobolus* sporophore into the spore itself may evert during flight, thus giving a slight push against the sporophore fragment and assuring a somewhat longer trajectory for the spore than for the sporophore fragment from which the spore becomes detached.

With the exception of *Massospora*, all other genera of the Entomophthorales apparently have primary spores discharged by the eversion of the spore's papilla against the sporophore (Gallaud 1905; Couch 1939). This mechanism depends upon the establishment of high turgor pressures in both the sporophore and spore prior to the rounding off of the turgid cell (Ingold 1971). The sporophore wall breaks at the point of spore attachment, and the spore papilla everts rapidly from its original position pointed into the spore and pushes the spore away from the turgid sporophore. The shock of this eversion against the pressurized water column of the sporophore sometimes causes a rupture of the sporophore tip and exudation of a water droplet at the time of discharge (*e.g.*, Page and Humber 1973).

Despite the lack of detailed understandings of these discharge mechanisms and of the involvements of the various wall layers of the sporophores and spores, it is apparent that spore discharge is a major event in the life history of an entomophthoralean fungus requiring the expression of a considerable portion of the genome. In view of the significance of this process for the fungus and the restricted distributions of the three diverse mechanisms, it is appropriate to recognize the mode of forcible discharge of primary spores as having taxonomic value at least on the generic level.

MODE OF SECONDARY SPORULATION

The general ability of entomophthoralean fungi to produce one or more types of secondary spores from primary spores is one of the most unusual features of this group of fungi. Among other fungal taxa, the Sporobolomycetaceae (heterobasidiomycetous "mirror" yeasts), many ascomycetes whose ascospores may bud repeatedly in the ascus, and some other entomopathogens such as a few species of *Septobasidium* Patouillard (Couch 1938) and the hyphomycetous genera *Muogone* Thaxter (1920) and *Termitariopsis* Blackwell, Samson & Kimbrough (1980) may produce secondary spores. In none of these groups, however, has secondary sporulation assumed such a significant role as in the Entomophthorales. It is becoming increasingly apparent, for instance, that the secondary spore types are equally as infective as primary spores or that primary spores serve as dispersive units while the secondary spores produced from them may even be the major infective units (Carner 1976; Carner and Canerday 1968; Kramer 1980; Nemoto and Aoki 1975; Nemoto et al. 1979; Selhime and Muma 1966; Tsintsadze and Vartapetov 1976; Wilding 1970).

The morphological diversity (Batko 1974; King and Hummer 1981) and biological importance of these spore types suggests that they might have taxonomic value above the specific level. In fact, two types of secondary spore forms — microspores and capillispores — have formed the bases for taxonomic divisions at subgeneric or generic levels in the Entomophthorales.

Microspores are produced when the primary spores of some *Conidiobolus* and *Basidiobolus* species produce a few to several dozen small secondary spores which are forcibly discharged and act individually as asexual propagules. The formation of microspores by *C. coronatus* (Costantin) Batko was first used to distinguish the genus *Delacroixia* Sacc. & Sydow; Tyrrell and MacLeod (1972) proposed that *Delacroixia* should be regarded as a subgenus of *Conidiobolus* to accommodate all species of that genus capable of forming microspores.

Capillispores (despite a profusion of names which are applied to them) are passively detached spores produced singly atop a thin and elongate (capillary) sporophore. Batko (1966) proposed to divide *Zoopthora* among four sub-

genera with all species producing almond-shaped to elongate secondary spores on capillary sporophores to be restricted to the subgenus *Zoophthora* Batko. Remaudière and Hennebert (1980) restricted the genus *Zoophthora* to those species producing capillaries and capillispores and transferred all species remaining in the other three subgenera proposed by Batko (1966) to *Erynia* Nowakowski. In addition to their presence in all species of *Zoophthora* subg. *Zoophthora* sensu Ben-Ze'ev & Kenneth (1981a) or *Zoophthora* Batko sensu Remaudière & Hennebert (1980), capillispores are produced by all species of *Triplosporium* sensu Batko (but not *Neozygites* sensu Remaudière & Keller 1980), *Meristacrum* (including *Tabanomyces*; see below), and some species of *Conidiobolus* and *Basidiobolus*. Batko (1974) considers the presence of capillary sporophores to reflect an ancestral character of the Entomophthorales and rejects the possibility of their separate origins in these genera by evolutionary convergence.

If the Remaudière classification were perfectly even-handed in its adoption of a generic criterion based on capillispores, then *Entomophthora turbinata* Kenneth (1977), which produces no capillispores, should not have been included in *Neozygites* by Remaudière and Keller (1980), and both *Conidiobolus* and *Basidiobolus* should have been split. It is curious that Remaudière and his colleagues apparently to not accept the parallel notion of splitting *Conidiobolus* because of the presence of microspores in some species of this genus.

Ben-Ze'ev and Kenneth (1981a) correctly reject the generic emphasis placed on capillispores by Remaudière and Hennebert (1980), and propose to use microspores and capillispores as a character with subgeneric validity as did Batko (1966) and Tyrrell and MacLeod (1972) before them. However, it is neither advisable nor even possible to apply criteria based on specialized secondary spore forms to all genera in which they might occur since *Basidiobolus microsporus* forms both microspores and capillispores (Benjamin 1962).

RHIZOIDS

Historical Perspective and Taxonomic Distribution

Batko's (1964a-c, 1974) placement of generic value on the presence of rhizoids has been one of the most troublesome aspects of his classification of the entomopathogenic Entomophthorales. His adoption of this criterion narrowed his circumscription of *Entomophthora* s.str. and led him to disperse the few species now regarded to belong in this genus (Remaudière and Keller 1980) among two genera. This misplaced emphasis also led him to propose *Culicicola* and *Entomophaga*, separated by the respective presence or absence of rhizoids, genera whose validity and circumscriptions have remained suspect due to the general controversy over the validity of rhizoids and to Batko's inability to provide a definitive separation of *Conidiobolus* from *Entomophthora* sensu lato.

Humber et al. (1977) re-evaluated and rejected the significance of rhizoids described for *Entomophthora virulenta* Hall & Dunn (1957; = *Conidiobolus thomboides* Drechsler; see Latgé et al. 1980). The rhizoids described for this fungus are now regarded to have belonged to another entomophthoralean fungus affecting the same aphids but which produced no spores or whose spores were undetected.

Remaudière and Keller (1980) and King and Humber (1981) independently discuss those species which usually form rhizoids but occasionally fail to do so. Note that all these exceptions represent the absence of (or failure to find) rhizoids in species normally producing them; they never involve observations of rhizoids in species not known to form them. Remaudière and Keller (1980) believe that no weight should be put on the presence of rhizoids because their presence "in certain species" is not always constant. On the contrary, King and Humber (1981) regard the presence of rhizoids to be taxonomically significant while their absence is not a dependable criterion.* Ben-Ze'ev and Kenneth (1981a) regard the presence of rhizoids to be taxonomically significant at the subgeneric level. Brobyn and Wilding (1977), in the most thorough consideration of the develop-

* Editorial changes of the text unintentionally altered this opinion to state that the absence of rhizoids is not taxonomically significant.

ment of rhizoids and cystidia yet published, uphold the presence of rhizoids to be a taxonomically significant character.

It is necessary to consider the general situation to know if any confidence should be placed on the presence (or absence) of rhizoids as a taxonomic characters. Even though no absolute scale is available to quantify one's level of confidence, it is necessary to weigh how well a species is known: How many times has it been found? In how many sites? From how many hosts? Are rhizoids always present or always absent? An examination of the taxonomic distribution of rhizoids among the entomophthoralean species reveals that many have never been found with them present. In only a few species, particularly *Zoophthora radicans* (= *Entomophthora sphaerosperma*), is the presence of rhizoids usual but inconstant. The absence of rhizoids might be explained in some instances by improper handling of specimens, but it is also possible that their formation on some insects might be a function of that particular host (Remaudière and Keller 1980; King and Humber 1981).

GENERA WITHOUT RHIZOIDS. Rhizoids are completely absent or have not been found in several genera. These include *Triplosporium* (at least 7 species), *Entomophaga* (at least 4 species although the circumscription of this genus remains in dispute), *Tabanomyces* (1 species; = *Meristacrum*), *Massospora* (13 species), and *Strongwellsea* (2 species). None of the non-entomopathogenic genera — *Basidiobolus*, *Ancylistes*, *Completozia*, *Meristacrum*, and *Ballocephala* — produce rhizoids.

ENTOMOPHTHORA (SENSU STRICTO). Two of the six species of *Entomophthora* s.str. (Remaudière and Keller 1980) produce rhizoids. This is the least constant appearance of this structure in any entomophthoralean genus. It can be said with assurance that rhizoids do not occur in the type species, *E. muscae*, since this is the most frequently observed of all species in the order. Both *E. culicis* (Braun) Fres. and *E. planchoniana* Cornu produce abundant rhizoids on hosts bearing forcibly discharged primary spores, but these anchoring structures may not occur when resting spores are formed. Both species are cosmopolitan in distribution and affect a wide variety of dipterans and aphids (respectively). All of the remaining species — *E. weberi* Lakon ex Samson (a little known species from a neuropteran),

E. thripidum Samson & al. (1979; known only from thrips in Dutch greenhouses), and *E. erpata* (Dustan) Hall (which is known from several North American and European sites and hosts) produce no rhizoids and disperse their primary spores from living, mobile hosts. The same behavior occurs in the species of *Massospora* and *Strongwellsea*.

CONIDIOBOLUS. Among the 27 morphologically homogeneous species recognized by King (1976a-b, 1977), none is known to produce rhizoids; very few *Conidiobolus* species are currently known to be entomopathogenic. Remaudière and Keller (1980) transferred most entomophthoroid species with round to pyriform spores from *Entomophaga* (which was circumscribed to exclude species with rhizoids) and those species from *Culicicola* with round spores to *Conidiobolus*. All other *Culicicola* species have campanulate spores and were restored to *Entomophthora* by Remaudière and Keller (1980).

Among the 38 *Conidiobolus* species listed by Remaudière and Keller (1980), four produce rhizoids: *Empusa apiculata* Thax., *E. major* Thax., *E. papillata* Thax., and *Entomophthora carpentieri* Giard (1888). The first three species were placed in *Culicicola* by Batko (1964c-d) because of their rhizoids and multinucleate round spores borne on simple (or occasionally branched) sporophores. Their pronounced similarity to *Conidiobolus* species belies two obvious morphological differences from all other *Conidiobolus* species: the consistent formation of a few stout rhizoids ending in strongly differentiated holdfasts, and the nearly cylindrical, collar-like narrowing of the sporophore apex. Close study of these species will be required to verify if they actually belong in *Conidiobolus* as suspected by King (1976b) and affirmed by Remaudière and Keller (1980).

The only morphological character of *Entomophthora carpentieri* Giard (1888) fitting *Conidiobolus* is the round primary spore supposed by Turian (1957) to belong to this species. Several other characters of this fungus — the presence of rhizoids, uninucleate spores on branched sporophores (Turian 1957), and cystidia (Petch 1944, if Petch's belief that *E. coleopterorum* Petch and *E. carpentieri* are identical is correct) — correspond exactly with those of *Erynia* or *Zoophthora* in the Remaudière classification. *Entomophthora coleopterorum* (Petch 1944) has narrowly oval rather than round primary spores; the mode of rhizoidal attachment is identical with that of *E. carpentieri*, a spe-

cies which Giard (1888) characterized only the the distinctive nature of its rhizoidal attachment. In a situation resembling that described above for the "rhizoids" of *C. thromboides*, one cannot discard the possibility that Turian (1957) may have observed the round spores of a secondarily invasive *Conidiobolus* species on a beetle which was already infected by *E. carpentieri* which was able to form its very characteristic rhizoids but no spores. No cytological staining or photomicrographic evidence supported Turian's contention that these round spores were uninucleate; it is possible that he mistook a central oil droplet or vacuole for a nucleus. The discrepancies between the Giard, Petch, and Turian concepts of *E. carpentieri* can be reconciled only after examining any existing herbarium material of these collections. In any event, however, the entire habit of this species refers it to *Erynia* or *Zoopthora* rather than to *Conidiobolus*.

ZOOPHTHORA. The presence of rhizoids is strongly correlated with branched sporophores bearing uninucleate, bitunicate primary spores — the species of *Zoopthora* Batko or *Erynia* Nowakowski. All eight *Zoopthora* species recognized by Remaudière and Hennebert (1980) form rhizoids. By including four newly described species, *Z. lanceolata* Keller (1980), *Z. crassitunicata* Keller (1980), *Z. orientalis* Ben-Ze'ev & Kenneth (1981a), and *Z. petchii* Ben-Ze'ev & Kenneth (1981b), the total rises to at least 11 of 12 species known to produce rhizoids. Ben-Ze'ev and Kenneth (1981a) remain uncertain if rhizoids were present on the few specimens of *Z. orientalis* available to them.

ERYNIA. Among the species listed by Remaudière and Hennebert (1980) and Remaudière and Keller (1980), some 21 of 25 *Erynia* species (excluding those of *Strongwellsea*) produce rhizoids. Together with *Erynia delphacis* (Hori) Humber*, *E. delpiniana* (Cavara) Humber**, and *E. ithacensis* Kramer (1981), the total incidence of rhizoids rises to 22 of 28 species. The absence of rhizoids in the six *Erynia*

* *ERYNIA DELPHACIS* (Hori) Humber, comb. nov., basionym: *Entomophthora delphacis* Hori, 1906, Entomol. Mag. (Tokyo) 3: 81.

** *ERYNIA DELPINIANA* (Cavara) Humber, comb. nov., basionym: *Entomophthora delpiniana* Cavara, 1899, Nuov. Giorn. Bot. Ital. (N. Ser.) 6: 422.

species discussed below is not considered to overturn the value of rhizoids as a secondary generic character although *E. delphacis* and *E. delpiniana* constitute the most notable of these exceptions.

Thaxter (1888) suspected that rhizoids were present on *Empusa virescens* Thax., but could not be certain since he was not the original collector of the material which he later used to describe this species.

Erynia aquatica (Anderson & Anagnostakis) Humber* is known only from mosquito larvae and pupae floating on the surface of temporary pools in the northeastern United States. This floating habit of the stricken hosts maintains their position for favorable transmission of the fungus without any "need" for rhizoids. This species is not known well enough, however, to say whether rhizoids might not form, for example, on individuals trapped at the receding edges of rapidly drying pools.

Ben-Ze'ev and Kenneth (1979) report cystidia but no rhizoids from *Zoophthora erinacea* Ben-Ze'ev & Kenneth, but note the similarity of rhizoids and cystidia of species referable to *Zoophthora* subg. *Erynia* Batko (1966). They muse on the possible inter-relatedness of cystidia and rhizoids as did Thaxter (1888) and Gustafsson (1965). However, Brobyn and Wilding (1977) provide effective evidence that the functions and course of development of these structures are unrelated in species where they both occur together; it must be assumed, then, that rhizoids and cystidia are under completely separate genetic controls.

Empusa caroliniana, with its multinucleate unitunicate spores borne on simple sporophores and lack of cystidia or rhizoids, is once again shown to stand apart from *Erynia* in

* *ERYNIA AQUATICA* (Anderson & Ringo ex Anderson & Anagnostakis) Humber, comb. nov., basionym: *Entomophthora aquatica* Anderson & Ringo ex Anderson & Anagnostakis, 1980, Mycotaxon 10, 350, NON *Entomophthora aquatica* Anderson & Ringo, 1969, J. Invertebr. Pathol. 13, 386 (which was invalidly published without a designated type). The combination *Erynia aquatica* (Anderson & Ringo) Remaudière & Hennebert, 1980, Mycotaxon 11, 301, is also invalid according to Article 45 of the International Code of Botanical Nomenclature.

all major characters except for the shape of its primary spores. The correct generic assignment for this fungus remains problematic, although its spore karyology, sporophore morphology, and mode of spore discharge more closely resemble the species of *Entomophaga* than any other genus recognized by the Batko, Remaudière, or Ben-Ze'ev/Kenneth classifications.

Erynia delphiniana and *Erynia delphacis* may be true exceptions to the usually rhizoidal state of fungi in this genus. Cavara (1899b) is fastidious in his observations of the small, uninucleate, bitunicate primary spores, secondary spores like the primaries, digitately branched sporophores, and very large and occasionally branched cystidia of *E. delphiniana*. However, he never specifically mentions any hyphae affixing the affected flies to the damp piers of a bridge over a small stream. The flies were completely covered under a dense, dirty-white to straw-colored hymenium. This entire hymenial covering was easily broken away from the cadavers during attempts to detach the insects (Cavara 1899b). Except for the apparent absence of rhizoids all characters of this species match the characteristics used by describe *Zoophthora* subg. *Erynia* Batko (1966); all species in this subgenus have rhizoids and prominent cystidia. *Erynia delphiniana* is known only from Cavara's original collections, but it seems likely that examination of this material or any fresh collections would reveal the presence of rhizoids.

Erynia delphacis is a much more important exception to the presence of rhizoids than the other species considered above. This pathogen of leaf- and planthoppers in Asian rice paddies bears a strong morphological resemblance to *E. neoaphidis* Remaudière & Hennebert (= *Entomophthora aphidis* Hoffman sensu Thaxter) from aphids (Shimazu 1976, 1977; Remaudière and Hennebert 1980), but differs notably in the complete lack of rhizoids on any of its hosts, including two artificially infected aphid species, *Aphis gossypii* and *Macrosiphum akiebiae* (Shimazu 1977, and personal communication). *Erynia delphacis* produces cystidia both on its host and in culture. Even though Hori (1906) illustrates and describes spherical, yellow-white resting spores 24-28 μm in diameter and containing many fatty granules, no such spores have been found in subsequent collections of this species in Japan, the Philippines, or Indonesia, nor are they formed in culture.

Remaudière and Hennebert (1980) consider the differences of growth *in vitro* between *E. delphacis* and *E. neoaphidis* to be insignificant and regard *E. delphacis* as a *nomen dubium*. However, a comparison of a Japanese strain (Shimazu F32; our RS 134) and several Philippine isolates of *E. delphacis* (RS 458, RS 459, RS 461, RS 478, RS 479) from green leafhoppers, *Nephotettix virescens*, with numerous cultures of *E. neoaphidis* isolated by Insect Pathology Research Unit personnel indicate that *E. delphacis* grows substantially more rapidly on a wider variety of media (nutritionally simple or complex), and sporulates a great deal more prolifically over a longer time than any culture of *E. neoaphidis* we have yet observed. Until more extensive studies can be made comparing these two species, it appears to be preferable to recognize *E. delphacis* as a legitimate species of *Erynia*. It should also be noted that if later studies indicate the synonymy of *E. delphacis* and *E. neoaphidis*, the International Code of Botanical Nomenclature would require that *E. delphacis* (Hori) Humber must be the correct name for the fungus still generally known as *Entomophthora aphidis* Hoffm. (*sensu* Thaxter).

Much more pertinent information is needed about the morphology of rhizoids than the perfunctory characterization found in many descriptions that rhizoids are merely "present." As noted by King and Humber (1981), complete descriptive information about rhizoids should include their points of emergence, dimensions (particularly thickness), abundance (few or many?), relative spatial distribution (isolated, aggregated into one or more pseudorhizomorphs, or forming a veil?), and the morphology of any terminal holdfast apparatus. Thaxter (1888), Gustafsson (1965), Ben-Ze'ev and Uziel (1979), and Ben-Ze'ev and Kenneth (1981a-b) are exemplary in the clarity of their illustrations and characterizations of rhizoidal morphology. This descriptive information can be useful for detecting dual infections involving rhizoidal and non-rhizoidal species since Brobyn and Wilding (1977) affirm that the morphologies of rhizoids appear to be species specific.

At least three morphological classes of rhizoids can be recognized: Most rhizoidal species have either (1) numerous individual rhizoids which are relatively little differentiated from vegetative and may or may not have any terminal holdfast differentiation, (2) numerous hypha-like rhizoids aggregated into one or more columnar pseudorhizomorphs, or (3) relatively few, thick rhizoids with strongly

differentiated (usually discoid) terminal holdfasts. No comprehensive study of the development and morphology of rhizoids or of their taxonomic distribution has yet been undertaken. The thorough studies of Brobyn and Wilding (1977) were limited to a few species affecting aphids. In view of the observations presented here, however, such a comprehensive study would be exceptionally useful.

Rhizoids as a Taxonomic Character

The Batko classification adopted the widely accepted contention of Nowakowski (1883) that the presence of rhizoids is a major generic character. This criterion, superimposed on those of nuclear number in primary spores and the branching of sporophores had effects which have been regarded as unacceptable and have prevented the general acceptance of the Batko classification (Remaudière and Keller 1980; King and Humber 1981). The Remaudière classification, in turn, rejected the significance of the presence of rhizoids. It seem certain, however, that the absence of rhizoids in many genera is significant while their presence in others is an important secondary character.

Regardless of the level (if any) at which one may accept the taxonomic importance of rhizoids, species presenting exceptions to the normal rhizoidal or nonrhizoidal condition of their genus should be examined carefully. The classifications of *Strongwellsea castrans*, *S. magna*, and *E. caroliniana* in *Erynia* and of *E. apiculata*, *E. major*, *E. papillata*, and *E. carpentieri* in *Conidiobolus* are important examples of possible misclassifications pointed out by the lack or possession of rhizoids.

CYSTIDIA

Cystidia (pseudocystidia, paraphyses, or pseudoparaphyses for other authors) are short to long, spear-like to columnar structures of determinate growth that project above the level of the sporogenous hymenium. Until recently, little has been known of their function, but Brobyn and Wilding (1977) convincingly demonstrated that the cystidia of *Erynia neoaphidis* are penetration organs which rupture the host cuticle and aid the subsequent emergence of the

sporophores. The emergence of cystidia before the sporophores also has been noted in *E. erinacea* (Kenneth 1977) and *E. ithacensis* (Kramer 1981). It seems likely that cystidia serve to perforate the host cuticle prior to hymenial formation in all species in which they occur; future investigators should be careful to note the order of emergence of cystidia and sporophores.

Remaudière and Keller (1980) indicate that, apart from *Erynia* and (less commonly) *Zoophthora*, the only other genus in which cystidia occur is *Conidiobolus*. Among the 38 species of *Conidiobolus* accepted by Remaudière and Keller, the only ones ever mentioned to form cystidia are *C. obscurus*, *C. thomboides* (= *E. virulenta*), and *E. carpentieri* (as *E. coleopterorum* Petch 1944); as was noted above, the latter species almost certainly belongs in *Erynia* or *Zoophthora* rather than *Conidiobolus*.

The description of *E. virulenta* (= *C. thomboides*; see Latgé et al. 1980) states that "cystidia occur rarely" (Hall and Dunn 1957). However, as explained above for the "rhizoids" of this species, no subsequent treatment of it reports finding either rhizoids or cystidia. The "cystidia" of *C. thomboides* might have been those of another entomophthoralean fungus which produced both rhizoids and cystidia (see Humber et al. 1977). Hall and Dunn (1957) might also have interpreted vegetative or sporogenous hyphae whose indeterminate growth occasionally far exceeds the general level of the hymenium to be cystidia.

Remaudière et al. (1979) report the formation of long cystidia by *C. obscurus* on aphid cadavers placed in a humid environment. I have observed similar growth of long hyphae above the hymenium of aphids affected by *C. thomboides* but always find these structures to be undifferentiated vegetative hyphae which should not be regarded as cystidia by the definition given above.

The listing of *Lamia* (= *Entomophthora*) *culicis* in a chart by Nowakowski (1883) as having cystidia present and rhizoids absent was corrected by Batko (1964c) as being a typographical transposition not corresponding with the Polish text which correctly notes this species to have rhizoids but no cystidia.

The genus most closely related to *Zoophthora* and *Erynia* must be *Strongwellsea*, with its uninucleate and bi-

tunicate primary spores borne on simple rather than branched sporophores (Batko and Weiser 1965; Humber 1976). Remaudière and Keller (1980) consider the lack of rhizoids and cystidia to be insignificant criteria and use this information to explain their submergence of *Strongwellsea* into *Erynia* (see Humber 1981a). In fact, if cystidia function only for cuticular penetration, it is not surprising that *Strongwellsea* species form no cystidia. These fungi produce no hymenial structures outside the host's body. The abdominal hole (the only cuticular rupture caused by the fungus) begins with a point perforation of the pleuron and enlarges progressively. This point initiation of the rupture appears to be involved with the hypertrophic reaction of the host cuticle which leads, in turn, to the extensive lining of the lower portion of the fungal ball by the host's cuticle. It seems clear that some mechanism completely distinct from that of cystidial penetration is involved in the formation by *Strongwellsea* of an abdominal hole (see Humber 1975, 1976).

In view of the above discussion, it is probably correct to say that the only species in which cystidia (as determinate organs for cuticular penetration) occur are those with uninucleate, bitunicate primary spores borne on branched sporophores and classifiable in *Erynia* or (less commonly) *Zoophthora* sensu Remaudière & Hennebert.

Unlike for rhizoids, the presence or absence of cystidia has never been used to separate genera although the Batko classification recognized their value as a supporting character strongly correlated with the occurrence of uninucleate, bitunicate primary spores and branched sporophores. As with the other characters discussed here, exceptions to the usual state of the character in a genus should indicate that a close inspection of the correspondences of all characters of the suspect species with those of other species in the genus is in order. The presence of cystidia in *E. coleopterorum* (Petch 1944), a presumptive synonym of *Conidiobolus carpentieri* (Giard) Remaudière & Keller, offers a significant example of this principle.

RESTING SPORES

Zygosporos or Azygosporos:A Morphological/Cytological Dilemma

I applaud the de-emphasis by Remaudière and Keller (1980) of the distinction between zygosporos and azygosporos. They note correctly that resting spores are not known (or are not formed) in a series of species, and that the mode of their formation has not been observed in a large proportion of species that do produce resting spores. Both types of development may occur in closely related species or even within a single species (MacLeod and Müller-Kögler 1973; Humber 1976; Remaudière and Keller 1980; King and Humber 1981).

This developmental distinction has been used only twice as a taxonomic character of generic importance. The peculiar mode of conjugation of (amoeboid?) hyphal bodies or gametangia of *Empusa fresenii* Nowakowski 1883 (= *Neozygites aphidis* Witlaczil 1885) and the resultant jet-black, ovoid zygosporos formed at the point of conjugation prompted Thaxter (1888) to set this and related species apart in *Empusa* subg. *Triplosporium*. This taxon was later raised to generic status by Batko (1964b), who was apparently unaware of the nomenclatural priority of the generic name *Neozygites* Witlaczil (see Humber et al. 1981). King (1976b) notes that the double-walled resting spores formed by the species of *Conidiobolus* are anisogamous zygosporos. Other types of conidioboloid resting spores such as chlamydo-sporos and villose spores are distinguished from zygosporos (or azygosporos) by their single rather than double wall structures.

Couch (1939) notes that the zygosporos of *Conidiobolus* species are formed in the larger of two gametangia (and are almost always produced in the axis of the parental hypha) while those of *Entomophthora* (sensu lato) are produced in lateral buds from the conjugating hyphae or hyphal bodies. Very few species of *Entomophthora* s.l. produce their zygosporos in the hyphal axis: *E. virulenta* (= *C. thromboides*), *E. obscura* Hall & Dunn [= *E. ignobilis* Hall & Dunn = *E. thaxteriana* (Petch) Hall & Bell; see below], and *Empusa dipterigena* Thaxter. The former two species are now recognized as species of *Conidiobolus* by Remaudière and Keller (1980). With the apparent exception of *E. dipterigena*,

whose zygosporogenesis is illustrated by Gustafsson (1965), the resting spores of other species of *Entomophthora* or its segregated are budded off from the parental hypha or hyphal body.

Zygosporos, the bilayered, thick-walled spores derived from a conjugation of two gametangia, have been a well established phenomenon in the Entomophthorales since their first recognition (Brefeld 1873; Nowakowski 1877, 1882, 1883; Thaxter (1888). Nearly every subsequent major discussion of these fungi has referred to zygosporos as sexual spores and to azygosporos as asexual spores (Schröter 1897; MacLeod 1956, 1963; Gustafsson 1965; Waterhouse 1973).

In general morphological terms, zygosporogenesis may be called a sexual process, in which case azygosporogenesis must be regarded as asexual. However, in their strict sense, the terms *zygosporos* and *azygosporos* refer only to the mode of development of a thick-walled spore in the Zygomycotina (Fitzpatrick 1930; Bessey 1950; Ainsworth 1961; Snell and Dick 1971; Alexopoulos and Mims 1979).

There is a hazard in thinking of zygosporos as sexual and azygosporos as asexual: One can be misled easily into performing an act of semantic sleight-of-hand by then expecting that the *genetic* events of a sexual life cycle, karyogamy and meiosis, necessarily occur in a "sexual" zygosporos but cannot occur in an "asexual" azygosporos. A related terminological confusion can occur when one refers to the resting spore as the sexual spore in the entomophthoralean life cycle as opposed to the forcibly discharged primary (or secondary) spores which are, in every sense, asexual spores.

Nuclear events in entomophthoralean resting spores — reductions in nuclear number, karyogamy, and meiosis — remain completely uninvestigated in all but a few species. This gap in our knowledge reflects a lack of fundamental studies on the life cycles in this group.

Some cytological studies have noted the progressive reductions of nuclear numbers to two or even a single nucleus in fully mature resting spores (Vuillemin 1900a-b; Krenner 1961; Latgé 1976; Couch et al. 1979; D. McCabe and B. Martinell, unpublished). Other investigators using different species have found no obvious change in the nuclear number from the time of formation to the fully

mature, dormant state (Olive 1906; Riddle 1906; Goldstein 1929; Humber 1975). If the Entomophthorales follow patterns of karyogamy and meiosis similar to those in other Zygomycetes (Cutter 1942a-b), karyogamy should occur in the resting spore followed by meiosis in the spore just before or during its germination. No conclusive cytological evidence of nuclear fusions or of meiotic division has yet been found in entomophthoralean fungi despite the strong circumstantial evidence for their occurrences (*e.g.*, Couch et al. 1979). It must be hoped that synaptonemal complexes, a widely recognized ultrastructural marker for chromosomal synapsis, will be demonstrated at the time and site of the presumptive meiosis in one or more members of the Entomophthorales.

Data regarding changes in nuclear numbers presented by Latgé (1976) support the possibility that karyogamy and meiosis occur in the resting spores of *C. thombooides*, and that the vegetative nuclei of this species must be haploid. Albeit, Latgé feels that his data are also compatible with gametangial meioses of diploid nuclei such as occurs in the Oomycetes (see Alexopoulos and Mims 1979). Little ambiguity about the interpretation of the life cycle remains, however, in view of the results of electrophoretic analyses of isozyme patterns of several strains of *C. thombooides* and *Z. radicans* (May et al. 1979) and of *E. muscae*, *Massospora cicadina* Peck, and *Entomophaga grylli* (Fres.) Batko (Soper, May, and Martinell, in preparation). These results suggest that the vegetative nuclei of these species are haploid, although heterokaryosis may occur in some species since multiple alleles of some loci may be present in a mycelium.

There is no evidence yet of heterothallism within the Entomophthorales. All zygosporogenesis appears to be homothallic and leaves open at least two possible interpretations for the life cycles of these fungi: Those species whose zygospores or azygospores at some point contain only a single nucleus probably have autogamous (sexual) life cycles. Those species whose zygospores or azygospores undergo no reduction in nuclear numbers may have abandoned a sexual life cycle in favor of an apogamous (asexual) life which preserves the particular (successful) combination of characters in that species. No evidence now exists that parasexuality occurs in any species of the Entomophthorales.

The distinction between zygospores and azygospores cannot be retained as a character of taxonomic significance although the presence and nature of zygospores is a useful ancillary character in at least *Triplosporium* and *Conidiobolus*. The developmental origins of resting spores can be difficult to interpret correctly even if suitable material is available. Nuclear events in resting spores are of far greater importance for the fungus than whether conjugations precede sporogenesis. Karyogamy and meiosis, the critical genetic events in a sexual life cycle, apparently may occur in either zygospores or azygospores; similarly, other species may have apogamous life cycles regardless of whether their resting spores are zygospores or azygospores.

Indeed, the full details of entomophthoralean life cycles and their breeding strategies may yet prove to be useful taxonomic characters, but not enough is currently known of these processes to draw meaningful conclusions. There is no doubt, however, that this information is of absolutely vital importance for the practical development and manipulation of these fungi for use in the microbial control of their insect hosts.

Resting Spores: Problems and Interpretation of Taxonomic Traps

It appears that the generic events of sexual reproduction might occur in either zygo- or azygospores, and that even species producing "sexual" zygospores might be apogamous. Several further caveats and an example serve to reinforce the rejection of any taxonomic value which might be attached to the distinction between zygospores and azygospores (Remaudière and Keller 1980).

In most instances, authors simply state their observation that the resting spores of a given species are zygospores or azygospores and neither illustrate nor discuss the evidence supporting this conclusion. However, Sawyer (1931) notes the great difficulty in determining if conjugations occur in *Z. radicans*. Humber (1975) also notes difficulties with the resting spores of *Strongwellsea* species due to problems of interpreting the morphology of some hyphal bodies and to the evanescence of the emptied hyphal wall remnants which provide the only temporary evidence for the mode of formation of mature resting spores;

determining the "pedigree" of these resting spores is dependent upon seeing them during a relatively narrow period during their development. Consequently, it seems unwise to place full confidence in any characterization of resting spores as zygo- or azygospores unless convincing documentation is also presented.

The difficulty in correctly applying these developmental designations casts doubt on whether any taxonomic decisions should be based solely on this distinction (even though it may have some utility for characterizing, but not separating, some genera as noted above). The practical effects of differing viewpoints of the value of this distinction can be seen in the following example:

In studies on the identities of aphid-pathogenic species of the Entomophthorales, Humber (1978) noted that *Entomophthora ignobilis* Hall & Dunn should be used as the nomenclaturally valid name for *Entomophthora thaxteriana* (Petch) Hall & Bell, a species to which Hall and Bell (1963) considered *E. ignobilis* to be identical. *Empusa thaxteriana* Petch (1938) was proposed for a species misidentified by Thaxter (1888), and which was characterized as having azygospores. The resting spores of *E. ignobilis* are described by Hall and Dunn (1957) as zygo-spores. Humber (1978) considered the reported differences in the origins of resting spores to be so ambiguous as to be insignificant, an opinion which is reinforced here. A later study by Remaudière et al. (1979) used morphological and biochemical data to demonstrate the synonymy of *E. thaxteriana* and *E. obscura* Hall & Dunn (1957) [= *Conidiobolus obscurus* (Hall & Dunn) Rem. & Keller], which was described as producing azygospores.

In fact, the differences between *E. thaxteriana*, *E. ignobilis*, and *E. obscura* had always been troublesome for diagnosticians. The only major described difference between *E. ignobilis* and *E. obscura* is that of zygo-spores versus azygospores, a difference considered here, by Humber (1978), and by Remaudière and Keller (1980) to be insignificant. A more consistent and simple approach to this taxonomic problem would have been to accept *E. obscura* to be the correct name by virtue of page priority over *E. ignobilis* rather than to regard the latter species as a *nomen dubium* merely because of the described difference in the origins of the resting spores in these two species (Remaudière et al. 1979).

If there is to be even-handed application of the accepted taxonomic criteria and acknowledgement of the net effects of rejecting other criteria which have been applied in the past, it is now necessary to recognize the correct synonymy of *Conidiobolus obscurus* to be the following:

Conidiobolus obscurus (Hall & Dunn) Remaudière & Keller 1980, Mycotaxon 11, 331.

≡ *Entomophthora obscura* Hall & Dunn 1957, Hilgardia 27, 162.

≡ *Entomophaga obscura* (Hall & Dunn) Batko 1964, Bull. Acad. Polon. Sci., Sér. Sci. Biol. 12, 404.

= *Entomophthora ignobilis* Hall & Dunn 1957, Hilgardia 27, 162.

= *Entomophthora planchoniana* Cornu sensu Thaxter 1888, Mem. Boston Soc. Nat. Hist. 4, 165, NON sensu Petch 1938, Trans. Brit. Mycol. Soc. 21, 34.

≡ *Empusa thaxteriana* Petch 1938, Trans. Brit. Mycol. Soc. 21, 34. NAME INVALID WITHOUT LATIN DIAGNOSIS.

[≡ *Entomophthora thaxteriana* (Petch) Hall & Bell 1963, J. Insect Pathol. 5, 186.]

[≡ *Entomophaga thaxteriana* (Petch) Batko 1964 Bull. Acad. Polon., Sér. Sci. Biol. 12, 404.]

CHOOSING A CORRECT CLASSIFICATION:

TRIAL BY ANOMALY

One of the most severe tests for a classification system is to see how well it can handle the least typical species from the group being classified. Within the Entomophthorales, several seemingly anomalous generic assignments from both the Batko and Remaudière classifications are discussed here. However, in terms of testing the real capacities of these or any other classifications of these fungi, confirming the correlations of characters, or suggesting where primary taxonomic weights should be placed,

the most important fungus has not yet been considered. This is an undescribed species found by Bałazy (1978) in Poland which was, obviously, not known at the time of the proposition of the Batko classification, and was not considered by the Remaudière classification.

This fungus affixes gnats (Diptera: Tendipedidae) by rhizoids to wet, decayed wood just above a stream surface in a deciduous forest in the Wielkopolski National Park, Poland. Its primary spores are slender and curved, 45-75 (82) x 8-10.5 μm , strongly tapered apically and with a flattened, conical papilla. These characters almost exactly match those of *Erynia conica* (Thax.) Rem. & Henn., which has slender curved spores (of a size identical to Bałazy's fungus) with bitunicate walls and single nuclei, borne on branched sporophores, and with cystidia and rhizoids present. Bałazy's fungus is, however, clearly different from *E. conica* since its primary spores have unitunicate walls, contain 4-10 nuclei, and are borne on simple sporophores; cystidia are absent.

The Remaudière classification, with its primary emphasis on spore morphology, would place this species in *Erynia* even though that assignment would be impossible if one accepts the emended circumscription advocated here (excluding *Strongwellsea* and *E. caroliniana* from the genus by limiting it to species with uninucleate, bitunicate spores borne on branched sporophores and not producing capillisporos). The Bałazy fungus might be regarded as a species of *Culicicola* except that Batko (1974) apparently rejected *Culicicola* as a heterogeneous mixture of species now regarded to belong to *Entomophthora* s.str. or *Conidiobolus* (Remaudière and Keller 1980; King and Humber 1981). There is no basis for including the Bałazy fungus in *Conidiobolus* according to the circumscriptions of this genus either stated or implied by Batko (1974), Remaudière and Keller (1980), here, or in Humber (1981b). It might be assumed that the Polish fungus could be allied to *E. apiculata*, *E. major*, and *E. papillata* because of their multinucleate unitunicate spores, except that the nuclear morphology of the latter species corresponds to that of *Conidiobolus* species while that of the Polish fungus resembles *Entomophthora* and all of its segregates rather than *Conidiobolus* (see Humber 1981b). The generic system outlined by Ben-Ze'ev and Kenneth (1981a) would include this species in *Entomophthora* sensu stricto, but the mode of spore discharge in the Bałazy fungus is by the eversion of the papilla.

If, as suggested here, one rejects spore morphology as the primary generic criterion and accepts the restriction of *Erynia* noted above, and places generic value upon the karyology of primary spores, the morphology of the sporophore, and the mode of spore discharge, the only remaining sensible disposition for Bałazy's fungus is in a new and (for now) monotypic genus allied to but differing from *Erynia*. It seems reasonable to assume that other entomophthoralean species may yet be found with characters like those of the one characterized by Bałazy (1978).

It is imperative that this Polish fungus be fully characterized and more widely publicized. This fungus seems certain to provoke much useful discussion of the taxonomic criteria which should be accepted for entomophthoralean fungi, and of how much weight should be accorded to each of these criteria.

MODERN VIEWS OF ENTOMOPATHOGENIC GENERA OF THE ENTOMOPHTHORALES

This study has examined the major morphological criteria used in entomophthoralean taxonomy, particularly as they are applied in the generic classification schemes proposed by Batko and by Remaudière and his colleagues. It is apparent from all of the above considerations that the application of only three characters provides an effective separation of species into morphologically and biologically homogeneous genera. These characters are the following:

1. The number and nature of nuclei in the primary spores.
2. The morphology of the sporophores (and the mode of sporogenesis).
3. The mode of discharge of primary spores.

Neither the morphology of primary spores, types of secondary spore formed, presence or absence of rhizoids, presence or absence of cystidia, nor the mode of formation of resting spores was found here to be important for delimiting genera in the Entomophthorales. Each of these

characters, however, may have utility as secondary or supporting characters which do not in themselves determine the limits of genera. Some of these characters are appropriate for the circumscription of subgeneric taxa (Ben-Ze'ev and Kenneth 1981a).

It is appropriate to summarize the difference among the Batko and Remaudière classifications and that proposed preliminarily by Ben-Ze'ev and Kenneth (1981a), and to note those adjustments which seem to be necessary to circumscribe these genera still more accurately according to the criteria accepted here.

ENTOMOPHTHORA Fresenius. Batko's overestimation of the importance of rhizoids caused him to split a small and natural group of species (MacLeod et al. 1976; Samson et al. 1979) with simple sporophores, campanulate apiculate primary spores containing a relatively small number of nuclei, and a characteristic mechanism of spore discharge among two genera. The application of the mechanism of spore discharge as a generic criterion confirms the narrow definition of *Entomophthora* proposed by Remaudière and Keller (1980) despite the de-emphasis here of the value of spore morphology. The exceptionally high degree of morphological and biological homogeneity of this group of species would be manifestly lowered if the mode of spore discharge did not exclude both *Empusa caroliniana* Thaxter and the species found by Bałazy (1978) and discussed above. Both of these species have simple sporophores and plurinucleate primary spores but are discharged by papillar eversion rather than by the "fungal cannon" mechanism of *E. muscae*.

MASSOSPORA Peck. This venerable genus which affects gregarious cicadas (Homoptera: Cicadidae) was not treated in the Batko classification until his phylogenetic treatment (Batko 1974) since *Massospora* represents the only genus of the entomogenous Entomophthorales whose validity and circumscription (see Soper 1974, 1981) has never been questioned. The species of this genus have plurinucleate primary spores which are not forcibly discharged from the simple sporophores lining small cavities in the abdomen of affected insects.

TRIPLOSPORIUM (Thaxter) Batko, nom. gen. conserv. prop. Batko (1964b) accepted Thaxter's (1888) belief that *E. fresenii* and similar species should be accorded generic status. This small and natural group of species is distin-

guished primarily by its unique mode of zygosporogenesis and ovoid zygospores with black (or very dark) episporangia. Batko (1964b) circumscribed the genus to include species whose primary spores are quadrinucleate, borne on simple sporophores, and capable of producing capillisporangia as one type of secondary spore. Remaudière and Keller (1980) accept Batko's circumscription of this genus but they (1) reject the requirement for the primary spores to be quadrinucleate in order to justify the (contentious) inclusion of *E. turbinata* Kenneth (1977), and (2) replace *Triplosporium* (Thaxter) Batko with the older generic name *Neozygites* Witlaczil (1885). Humber et al. (1981) offer several reasons why *Triplosporium* should be conserved against the older and nomenclaturally correct name *Neozygites*; this nomenclatural issue will be resolved at the 13th International Botanical Congress during the summer of 1981.

ENTOMOPHAGA Batko. This genus, based on *Entomophthora grylli* Fres., was proposed to include species with multinucleate spores and simple sporophores but without forming rhizoids. Its validity has been questioned because of its linkage to the unacceptable *Culicicola* through Batko's emphasis on rhizoids and also because no satisfactory criterion has been proposed to distinguish unambiguously between any of these species and the morphologically similar *Conidiobolus*. The lack of this criterion seemingly prompted Remaudière and Keller (1980) to emphasize spore morphology by transferring all species of *Entomophthora* and those of *Culicicola* with round rather than campanulate primary spores to *Conidiobolus*. However, the criterion of nuclear morphology noted above and discussed fully by Humber (1981b) readily delimits *Conidiobolus* from *Entomophthora* sensu stricto and its segregate genera, and confirms the validity of *Entomophthora* for species with multinucleate unitunicate primary spores containing large, readily stainable nuclei, and discharged by papillar eversion from simple sporophores.

ZOOPHTHORA Batko and *ERYNIA* Nowakowski. Batko (1964b) described *Zoophtora*, with *Entomophthora radicans* Brefeld (= *Entomophthora sphaerosperma* Fresenius) as its type, for all species having uninucleate, bitunicate spores borne on branched sporophores and producing rhizoids. This genus has been accepted as a natural and valid grouping. Four subgenera were proposed by Batko (1966) based on the morphology of sporophores, rhizoids, and cystidia, and the type of secondary spores produced. Remaudière and Hennebert (1980) limited *Zoophtora* to species capable of form-

ing anadhesive secondary spores atop capillary sporophores (see King and Humber 1981), with all other *Zoophthora* species being transferred to *Erynia* Nowakowski (1881). The Remaudière classification modified the sense of Batko's grouping to accommodate species with simple sporophores and even multinucleate unitunicate primary spores so long as the spore shape was similar to that in other *Erynia* species.

For purposes of discussing the Remaudière classification in this paper, it has been necessary to accept this generic separation on at least a provisional basis. Considerable objection can be raised, as noted above, about the use of capillary sporophores as a character of generic significance. This issue has been treated in part by Ben-Ze'ev and Kenneth (1981a) and will be discussed more fully by these authors in a subsequent paper.

A nomenclatural problem reminiscent of the simultaneous use of *Empusa* Cohn and *Entomophthora* Fresenius as taxonomically distinct genera has been raised by Remaudière and Hennebert's use of both *Zoophthora* Batko and *Erynia* Nowakowski. In 1881, Nowakowski proposed the genus *Erynia* to include *Entomophthora ovispora* Nowakowski (1877) and *Entomophthora curvispora* Nowakowski (1877), but later rejected this generic name in favor of Brefeld's (1877) usage of *Entomophthora* as taxonomically distinct from *Empusa* (Nowakowski 1882, 1883). At the time of its description, *Zoophthora* Batko (1964b) included only its type species, *Entomophthora radicans* Brefeld. However, the inclusion in *Zoophthora* of *E. ovispora* and *E. curvispora* (Batko 1964d) automatically required the adoption of the earlier name. Batko (1966) was incorrect in believing that Nowakowski's later disuse of *Erynia* removed that name from consideration in matters of nomenclatural priority.

The ultimate resolution of this nomenclatural problem with *Erynia* and *Zoophthora* depends on the outcome of the debate to establish a consensus opinion of whether the Batko or Remaudière classification, or some modification of one of them shall be accepted for the species with bitunicate, uninucleate spores on branched sporophores. If, on the one hand, two genera are recognized with one based on *Entomophthora ovispora*, the type species chosen for *Erynia* by Remaudière and Hennebert (1980), and the other on *Entomophthora radicans*, there is no nomenclatural problem to resolve. If, on the other hand, common practice rejects

any generic separation and recognizes only a single genus which includes both *E. ovispora* and *E. radicans*, then either *Erynia* would have to be adopted as the correct generic name or *Zoophthora* might be proposed for conservation against *Erynia*.

It is unfortunate that such a confusing nomenclatural issue should arise once more in the Entomophthorales, but the three possible resolutions are unambiguous. My personal opinion should be clear from this paper: I do not believe that the species originally classified by Batko in *Zoophthora* should be split between two genera separated by characters such as spore morphology and the formation of capillary sporophores. As much as it might prevent some amount of further confusion in the literature if *Zoophthora* Batko were conserved against *Erynia* Nowakowski, I do not believe that such a proposal would be accepted for incorporation in the lists of *nomina conservanda* in the International Code of Botanical Nomenclature. The great preponderance of species proposed as or transferred to *Zoophthora* have already been transferred to *Erynia* by Remaudière and Hennebert (1980) or here; the relative number of species required to be transferred as a consequence of conservation of a younger name against an older and nomenclaturally correct one appears to be one of the major concerns of the nomenclatural committees which decide these matters. The new combinations in *Erynia* proposed here are fully justified and nomenclaturally correct whether one accepts the taxonomy of Batko (and his followers) or Remaudière and Hennebert for classifying the species in question.

STRONGWELLSEA Batko & Weiser emend. Humber. Humber (1976) provided an emended generic description and validation of this genus whose spores are uninucleate and bitunicate but are borne on simple rather than branched sporophores, and are discharged by papillar eversion. Remaudière and Keller (1980) provided no effective rationale for rejecting the importance of sporophore morphology or other diverse supporting data (Humber 1976) in order to synonymize *Strongwellsea* with *Erynia*. There can be little doubt from the criteria considered here to be significant at the generic level that *Strongwellsea* must be recognized as a separate genus. A more extensive support for this opinion is presented by Humber (1981a).

CULICICOLA Nieuwland. This genus is nomenclaturally superfluous since its type species, *Entomophthora culicis* (Braun) Fres., belongs in *Entomophthora* sensu stricto (Remaudière and Keller 1980; King and Humber 1981). The species included in this genus by Batko (1964c-d) are now dispersed to *Entomophthora* s.str. or *Conidiobolus* Brefeld by Remaudière and Keller, although some question remains whether the conidioboloid species producing rhizoids — *E. apiculata*, *E. major*, and *E. papillata* (which was not classified by Batko, 1964d, but clearly belonged in this group) — should be in *Conidiobolus* or in a closely allied but different genus. Batko's inclusion in *Culicicola* of *E. virulenta* (= *C. thromboides*; see Latgé et al. 1980) was based on the erroneous description of this species as producing rhizoids (Humber et al. 1977).

Batko regarded this as the most tentative of his proposed segregate genera, and he later apparently rejected *Culicicola* in his extended justification of his taxonomic ideas and discussion of phylogeny in the Entomophthorales (Batko 1974). *Culicicola* has been the most objectionable of his genera for other students of these fungi.

CONIDIOBOLUS Brefeld. Even though Batko and Weiser (1965) note that resting spores of *Conidiobolus* are not budded off to the side as in all other genera considered in the Batko classification, they again cited the widely used but untenable "criterion" which supposes *Conidiobolus* species to be basically saprobes while those of the other genera treated were entomopathogenic. Remaudière and Keller (1980) regard the lack of a suitable criterion to delimit *Conidiobolus* from morphologically similar species put in *Culicicola* and *Entomophaga* by Batko (1964b-d), with round to pyriform, multinucleate primary spores and simple spores, to justify their inclusion in a broadly redefined *Conidiobolus*.

As indicated here, however, nuclear cytology does provide the criterion sought to delimit *Conidiobolus* species from those of *Entomophthora* and its segregates (Humber 1981b). Some of the species transferred to *Conidiobolus* from *Entomophthora* sensu lato truly are species of *Conidiobolus*; these include *E. virulenta* (= *C. thromboides*; Latgé et al. 1980), *C. obscurus*, and possibly *C. apiculatus*, *C. major*, and *C. papillatus* (although these last three species may belong in a separate genus allied to *Conidiobolus* due to marked differences in the structure of the sporophore

with the presence of rhizoids as a supporting character). Other species — e.g., *E. grylli*, *E. batkoi*, and *E. gigantea* Keller — have nuclei whose morphology indicates their closer affinities to *Entomophthora* than to *Conidiobolus* (Humber 1981b).

TABANOMYCES Couch, Andreeva, Laird, & Nolan. This little known genus (Couch et al. 1979) was described for the peculiar mode of germination and spore formation from entomophthoralean resting spores found in larvae of tabanid flies in the Soviet Union. The thick-walled, colorless, ovoid resting spores of the type species, *T. milkoi* (Dudka & Koval) Couch & al., were originally described as *Coelomomyces milkoi* Dudka & Koval. During the germination of the spores, the single nucleus undergoes two (meiotic?) divisions, and a short, thick, quadrinucleate sporophore forms. This hypha becomes septate, and each uninucleate cell produces and discharges a lateral primary spore which is, in turn, capable of producing a secondary spore on a short capillary sporophore (Couch et al. 1979). These events and structures are essentially identical to those described for the germination of resting spores in the nematophagous genus *Meristacrum* Drechsler (Davidson and Barron 1973). The nuclei illustrated by Couch et al. (1979) are relatively small and have a prominent central nucleolus; this type of nucleus resembles that of *Conidiobolus* (Humber 1981b) and *Meristacrum* (Humber, unpublished).

Remaudière and Keller (1980) do not mention *Tabanomyces*, but this genus cannot be easily placed in any entomogenous genus included in the Batko or Remaudière classifications. The remarkable similarity of *Tabanomyces* and *Meristacrum* indicates that these genera are synonymous and that *T. milkoi* must be recognized as a heretofore unknown entomopathogenic species of *Meristacrum**.

TARICHIUM Cohn. This genus was created for those species of the Entomophthorales known only by their resting spores (see MacLeod and Müller-Kögler 1970). Neither the

* *MERISTACRUM MILKOI* (Dudka & Koval) Humber, comb. nov., basionym: *Coelomomyces milkoi* Dudka & Koval in Dudka, Koval & Andreeva, 1973, *Novitates Systematicae Plantarum non Vascularium* 10, 88-91. SYNONYM: *Tabanomyces milkoi* (Dudka & Koval) Couch, Andreeva, Laird & Nolan, 1979, Proc. Natl. Acad. Sci. (USA) 76, 2299-2302.

Batko nor Remaudière classifications fully accept this genus, but properly regard its members as having still undetermined affinities. It can be assumed that *Tarichium* species may be connected to a species producing primary spores and, thus, placed in the appropriate modern genera. A taxonomic system which accepts a narrow sense of *Entomophthora* and a number of other entomopathogenic segregate genera must also accept *Tarichium* as the correct provisional generic name for the species involved rather than the name *Entomophthora* (*Tarichium*) as was suggested (without any formal nomenclatural status as a subgenus of *Entomophthora*) by MacLeod and Müller-Kögler (1970).

Concluding Remarks on Generic Classifications

Despite its manifest shortcomings, the classification proposed by Batko (1964a-e, 1966; Batko and Weiser 1965) forms the basis for all other contemporary approaches to a generic classification for the Entomophthorales. This classification is discussed in depth (although not substantially amplified) in the first attempt to outline the course of evolution in the Entomophthorales (Batko 1974). The Remaudière classification attempted to eliminate the flaws which Remaudière and his colleagues perceived to prevent the general acceptance of the Batko classification. A generic arrangement more similar to that proposed by Batko than by Remaudière, Hennebert, and Keller emerges here in view of the acceptance of primary spore karyology, sporophore morphology, and the mode of primary spore discharge as the three characters best suited for delimiting genera. Ben-Ze'ev and Kenneth (1981a, 1981b, and other papers in preparation) also accept the karyology of primary spores and morphology of sporophores as characters with generic importance, but do not recognize the mechanism of spore discharge to be taxonomically significant.

No formal proposal of yet another new generic classification seems appropriate at this time. This study has sought only to evaluate the characters and criteria which have been or might be used to construct generic classifications of the Entomophthorales. I have extended the conclusions of this study regarding appropriate generic criteria to the existing classification schemes in order to show their net effects and to stimulate further discussion and debate.

Each attempt to propose a new generic classification for the Entomophthorales has suffered from the effects of one or more faultily circumscribed genera and from the less obvious effects of incompleteness. It is clear from the still incomplete catalogue of species described as or attributable to *Entomophthora* (sensu lato) (MacLeod 1963; MacLeod and Müller-Kögler 1970, 1973; MacLeod et al. 1976), that the available information about a distressing number of species is inadequate to allow a reliable classification in the narrowly defined genera comprising the contemporary classification schemes. I must also be noted that all of these latter-day efforts to reclassify the Entomophthorales have concentrated almost exclusively upon the entomopathogenic genera. Several morphologically diverse genera are known only as saprobes or pathogens of fern gametophytes, desmide algae, tardigrades, and nematodes; at least three species of this order are also known to affect humans and other vertebrates (King 1979). These fungi are no less valid members of the Entomophthorales than those species attacking insects; it is inappropriate to argue about the choice of criteria used to define genera or about the correctness of one or another classification of the Entomophthorales without taking these non-entomogenous genera into full account.

Despite these impediments to the proposition of a comprehensive generic scheme, however, I believe that most concerned parties now agree that following a period of further discussion of taxonomic criteria, it will be possible to propose a generally acceptable, realistic, and phylogenetically based classification for the Entomophthorales in which each genus will have a uniformly high degree of morphological and biological homogeneity. Despite the obvious difficulties in proposing such a comprehensive scheme, some comfort should be found in the thought that IF the generic scheme which is yet to emerge is accurate, then no further major adjustments of generic circumscriptions should be necessary. As new taxa are found and described, and as older taxa being held in abeyance become better known, they will be classified in the existing stable generic structure or seen to differ in enough significant characters to warrant the erection of new genera.

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LES HYMENOCHAETE A ELEMENTS HYMENIENS PINNATIFIDES

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SUMMARY

Hymenochaete acanthophysata nov.sp. with acanthophysate hyphae and *Hymenochaete hauerslevii* nov.sp. with pseudoacanthophysae are described. The author compares these African species with the only two known species with pinnatifid-tipped hyphidia in their hymenium: *H. digitata* Burt and *H. pinnatifida* Burt. The later has been found again: descriptions of a non stratose specimen from Africa and of a two-layered specimen from Guadalupe are given. An abstract in English including a detailed key to species is given in the end of the paper.

INTRODUCTION

Dans le cadre d'une étude des spécimens du genre *Hymenochaete* récoltés principalement en Afrique par J. BOIDIN d'une part et G. GILLES d'autre part, nous avons décrit précédemment une espèce, *H. spathulata*, remarquable par ses spinules à sommet spatulé (LEGER, 1980) ainsi que deux espèces, *H. separabilis* et *H. harpago* (LEGER, 1981) à spinules pourvues de diverticules.

Le présent travail a trait aux espèces dont l'hyménium possède des éléments stériles ornés de digitations. Jusqu'ici deux espèces présentent ce caractère: *H. pinnatifida* Burt et *H. digitata* Burt. *H. pinnatifida* a été retrouvé en République Centrafricaine, au Gabon et à la Guadeloupe. Une description complète est donnée qui fait apparaître notamment que cette espèce peut être bistratifiée. A titre comparatif, l'étude de *H. digitata* est reprise et complétée à partir du type de BURT. Enfin, deux espèces sont décrites : *H. acanthophysata* nov.sp. à hyphes acanthophysoïdes et *H. hauerslevii* nov.sp. à éléments hyméniens de type pseudoacanthophysés.

HYMENOCHAETE PINNATIFIDA Burt

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Basidiome résupiné, étalé, adhérent, mince, 60-150- (200) μm , en petites taches plus ou moins confluentes, brun jaunâtre (Munsell 10YR 5/4 ou encore Saccardo's umber selon Ridgway) puis brun rouillé (Munsell 5YR 4/6) quand il s'étale plus largement, à marge fibrilleuse appliquée, 0,5 mm, chamois (Munsell 10YR 7/8).

Cortex (Cx, fig.1) épais de 5 à 15 μm , sombre, formé d'un enchevêtrement serré d'hyphes brun-rouge foncé, plus ou moins cimentées et collapsées, x 2-2,5-(3) μm , à paroi épaisse.

Contexte (C, fig.1) assez lâche, 50 à 100 μm d'épaisseur, dimitique. Hyphes génératrices hyalines, x 1,5-2 μm , à paroi mince, ramifiées et cloisonnées. Hyphes squelettiques brun-jaunâtre, x 2-2,5 μm , à paroi épaisse, non cloisonnées, exceptionnellement ramifiées.

Spinules brun foncé, la plupart 15-30 x 3,5-5 μm , non émergentes; certaines, plus rares, mesurent 40-50 x 6-7 μm et émergent jusqu'à 20 μm . Toutes ces spinules naissent d'une couche sous-hyméniale sombre.

Hyménium (H, fig.1) de 15 à 20 μm ; rares basidioles et très rares basides mûres, 15 x 3 μm , à 4 stérigmates de 3 μm . Dendrophyses jaunes très abondantes, longues de 12 à 22 μm et de diamètre variant de 2 à 6 μm , à nombreuses ramifications très étroites (0,25 μm) en petites touffes terminales pour la plupart. Très rares hyphes paraphysoides grêles grimpant le long de certaines grandes spinules. Assez nombreuses masses cristallines volumineuses (souvent de 10-20 x 10 μm) à la base de l'hyménium.

Sous-hyménium (sH, fig.1) formant une zone sombre de 20 μm environ, d'aspect identique au cortex, constituée d'hyphes plus ou moins cimentées et collapsées, x 2-3 μm , à paroi épaisse.

Spores (sur sporée) cylindriques étroites, très légèrement déprimées, 3,8-4,5-(5) x 1,6-2 μm , hyalines, uninucléées (Giemsa), non amyloïdes, blanches en masse.

Description effectuée à partir de la récolte LY 5584, La Maboké, République Centrafricaine, 31 mars 1965, leg. J. Boidin (Herb. Boidin).

AUTRES RECOLTES :

LY 5931 sur *Carapa procera*, La Maboké, République Centrafricaine, 16 septembre 1967, leg. J. Boidin.

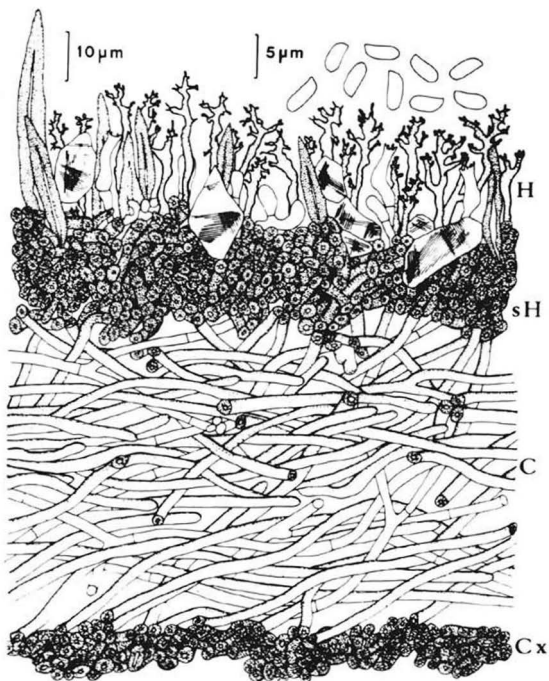


Figure 1 : *Hymenochaete pinnatifida* Burt
(LY 5584)

LY 6041 sur support indéterminé, Boubakiti, République Centrafricaine, 27 septembre 1967, leg. J. Boidin.

LY 9260 sur support indéterminé, Libreville, Gabon, 20 novembre 1978, leg. G. Gilles.

ESPECES DE REFERENCES EXAMINEES :

H. pinnatifida Burt, New Smyrna, Florida, January 1897, C.G. Lloyd 2139 (Type) et 2140. (FH).

H. pinnatifida Burt, *ad ligna emortua angiospermae in paroecia Aurelia*, Algiers, 17 septembre 1971, leg. M.T. Dunn, det. A.L. Welden (*Fungi Ludoviciani ex herbario Tulane* 7655). LY 6723.

LE CAS PARTICULIER DE LA RECOLTE LY 8091 :

Ce spécimen, trouvé par J. BOIDIN le 29 septembre 1976 à la Guadeloupe se présente comme un *H. pinnatifida* bistratifié (fig.2).

Tout se passe comme si un deuxième basidiome complet avec un cortex Cx2, un contexte C2, une zone sous-hyméniale sH2 et un hyménium H2 (pratiquement stérile) était né d'un basidiome précédent constitué (comme décrit pour LY 5584) d'un hyménium H1, d'un sous-hyménium sH1, d'un contexte C1 et d'un cortex basal Cx1. Dans le dessin de la figure 2, la base du champignon - c'est-à-dire le cortex Cx1 ainsi que les 4/5 du contexte C1 - n'est pas représentée. Le basidiome a une épaisseur totale de 150 à 230 μm dont 100 μm pour le contexte C1 (alors que le contexte C2 n'a qu'une quinzaine de μm d'épaisseur).

DISCUSSION :

Les spécimens africains présentent une très bonne concordance avec le type d' *H. pinnatifida* Burt examiné ainsi qu'avec la récolte américaine LY 6723 déterminée par A.L. WELDEN. Cependant, deux remarques s'imposent :

- L'assez grande variabilité de couleur des basidiomes, non seulement d'une récolte à l'autre mais aussi à l'intérieur d'une même récolte. Nous distinguons trois groupes selon la couleur: gris rosâtre (light cinnamon drab, vers 2,5 YR 6/2) pour LY 9260 et pour le centre de quelques taches de LY 6723; brun rougeâtre (reddish brown, vers 5YR 5/3 à 4/4 ou 4/6) pour LY 6041, LY 8091, LY 6723 *pro parte* et LY 5584 *pro parte*; brun jaunâtre (yellowish brown, vers 10YR 5/4) pour LY 5584 *pro parte* (petites taches très jeunes) et LY 5931. Cette couleur variable s'explique au moins en partie par le développement relatif de l'hyménium: ainsi, les taches de LY 6723 où l'hyménium est bien développé et fertile sont gris rosâtre alors que les taches brun rougeâtre ne présen-

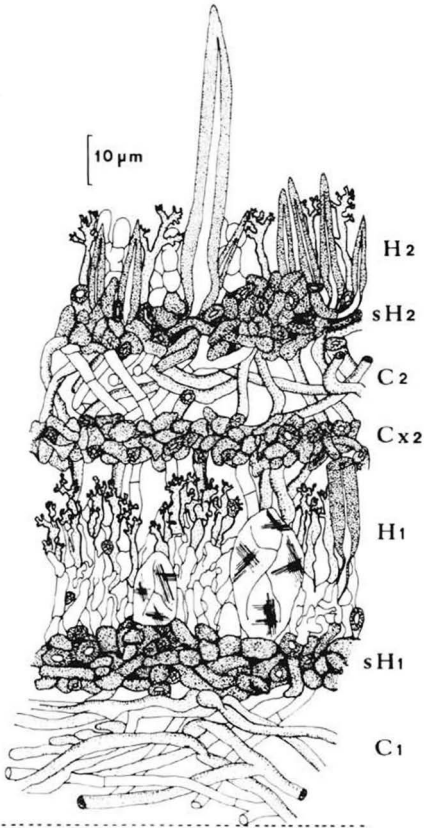


Figure 2 : partie supérieure d'*Hymenochaete pinnatifida* Burt (LY 8091)

-tent qu'un hyménium faiblement développé, pratiquement stérile et constitué presque exclusivement de dendrophyses. De même LY 9260, dont l'hyménium fertile est bien développé est uniformément gris rosâtre. La marge, chamois à rouillé, a presque toujours une couleur plus vive que le reste de l'hyménium.

- Concernant la stratification, l'examen de la récolte LY 8091 conduit incontestablement à *H. pinnatifida* Burt. Cette bistratification n'a jamais été signalée ni par E.A. BURT, ni ultérieurement lors d'autres récoltes.

En résumé, *H. pinnatifida* Burt est une espèce bien caractérisée par ses dendrophyses (à nombreuses et fines ramifications) issues d'une zone sous-hyméniale sombre, par un contexte dimitique assez lâche et un cortex basal sombre. La couleur de l'hyménium peut varier de façon sensible suivant l'état de maturité tandis que la marge est toujours de couleur plus vive.

L'aire de dispersion de *H. pinnatifida* s'étend non seulement à l'Afrique mais également à la Guadeloupe dont le spécimen montre un caractère jamais observé jusqu'ici pour cette espèce: une bistratification nette. Il serait intéressant de rechercher aux Etats-Unis notamment, une présence éventuelle de stratification lors de futures récoltes.

HYMENOCHAETE DIGITATA Burt

Ann. Missouri Bot. Gard. 5 : 347 . 1918

Basidiome résupiné, largement étalé, adhérent, épais de 700 à 800 μm , brun havane (Brussels brown à antique brown, 7,5YR 4/4 à 5/6), à surface granulée-tuberculée, à marge très mince, évanescence, concolore.

Cortex (Cx1, fig.4) d'une quinzaine de μm , formé d'un enchevêtrement serré et brun rouge sombre d'hyphes de 2,5-3 μm de diamètre, plus ou moins cimentées et collapsées, à paroi épaisse.

Ce cortex se prolonge par un feutrage assez lâche constituant le revêtement piléique (tomentum T), d'environ 40 μm d'épaisseur.

Contexte (fig. 3 et 4) stratifié, composé d'une alternance d'une quinzaine de couches sombres avec un nombre égal de couches plus claires. Les couches sombres (appelées Cx2 à Cx8 et dont seules Cx7 et Cx8 sont représentées sur la figure 3) ont une teinte brun rouge foncé et présentent une constitution identique à celle du cortex basal Cx1. Les couches plus claires ont une double constitution: d'une part, les

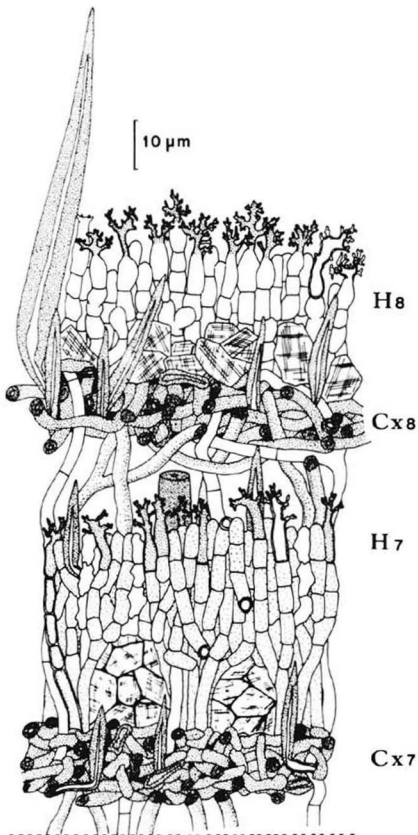


Figure 3 : partie supérieure d'
Hymenochaete digitata Burt (Type)

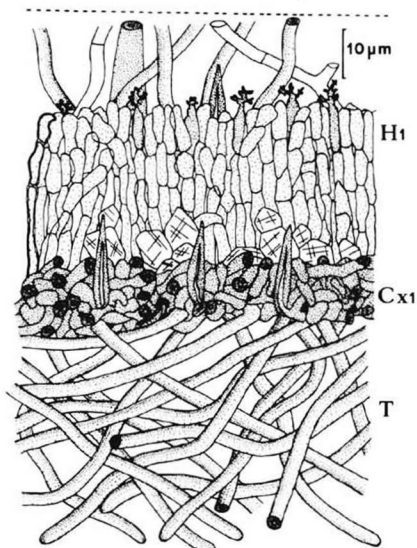


Figure 4: base d' *Hymenochaete digitata* Burt (Type)

hyméniums antérieurs (nommés H1 à H7 et dont seuls H1, H7, H8 sont représentés sur les figures 3 et 4) sont bruns, formés d'une dense palissade d'hyphes de 2-3 μm de diamètre, à paroi épaissie, à articles relativement courts (7 à 10 μm), très collapsées, auxquelles se mêlent des dendrophyses à paroi épaissie, entièrement jaunâtres. D'autre part, au dessus de chaque hyménium, des zones très claires sont formées d'un enchevêtrement très lâche d'hyphes cloisonnées et ramifiées, de 2-3 μm de diamètre, à paroi mince et d'hyphes non cloisonnées, 2-3 μm de diamètre, à paroi épaissie (dimitisme). A la base de chaque hyménium, de nombreuses masses cristallines plus ou moins volumineuses.

Spinules nées pour la plupart des couches très sombres; elles sont de deux types: très nombreuses petites

spinules 20-30 x 2-3 μm , non émergentes et grandes spinules 50-75 x 6-7 μm , moins abondantes et émergentes jusqu'à 40 μm .

Hyménium (H8, fig.3) de 25 μm environ; quelques basidiosoles mais basides extrêmement rares (non vues par BURT), 13 x 2,5-3 μm , à 4 petits stérigmates de 1,5 μm . L'essentiel est constitué par des sortes de pseudoacanthophyses nombreuses: ce sont des éléments hauts de 12 à 17 μm , à base renflée (3,5-4,5 μm) hyaline et à paroi mince, tandis que l'extrémité rétrécie, jaunâtre et à paroi épaisse porte de nombreuses ramifications en tous sens. Certaines sont cependant déjà entièrement jaunâtres et à paroi épaissie comme celles des hyméniums antérieurs.

Spores non observées.

Description d'après le type de *H. digitata* Burt, El Boquete, Chiriqui, Panama, W.R. Maxon, 5559. (FH).

DISCUSSION :

L'étude du type de BURT a permis d'apporter quelques précisions sur cette espèce qui n'a pas fait l'objet, depuis le travail originel, d'étude détaillée. Le point le plus intéressant, outre la présence non encore signalée d'un tomentum et la constitution précise des différentes couches de stratification, est sans doute l'aspect détaillé des pseudoacanthophyses qui n'avaient été que superficiellement décrites par BURT.

HYMENOCHAETE ACANTHOPHYSATA Léger nov.sp.

Basidioma resupinatum, late jacens, adherens, tenue, obscure brunneum, magis in medio propter pruina griseolum; ambitu ex rubiginoso fulvo, margine angustissimo, albo; trama bistrata, dimitica, hyphis genetricibus hyalinis, tunica tenui, x 2-3 μm , ramosis fibulatisque, ex hyphis skeletticis e flavis brunneis, tunica crassa, x 2-3 μm , neque ramosis neque septatis constante; cortice nullo; spinulis brunneis, amplis, 40-70 x 7-8,5 μm , usque ad 35 μm emergentibus; hyphis paraphysoideis gracilibus, x 1-1,5 μm , quasdam spinulas circumdantibus; basidiis rarissimis, 12 x 3 μm , 4 sterigmatis 3 μm longis; permultis hyphis acanthophysoideis, tunica crassa, x 2-2,5 μm , non vel vix emergentibus; passim crystallis inter hymenium praesentibus; sporis cylindratis, leviter depressis, 4-4,5-(5) x 1,5-1,8 μm , hyalinis, uninucleatis, haud amyloideis, in massa albis.

Holotypus : Lyon (LY), leg. J. Boidin n° LY 7741. In substrato incognito, Makokou, Gabon.

Ethymologie : espèce ainsi nommée du fait qu'elle possède de nombreuses hyphes acanthophysoides (acanthophyses définies par A. PILAT en 1926).

DESCRIPTION DU TYPE :

Basidiome résupiné, largement étalé, adhérent, mince, 125 à 200 μm , brun sombre (5YR 4/2, *fuliginus* de Saccardo) puis plus grisâtre au centre par la pruine (5YR 5/2, benzo brown R. à 2,5YR 5/2, cinnamon drab R.). Bordure fauve rouillée (7,5YR 5 à 6/6, vers antique brown R.). Marge très étroite, fibrilleuse, très appliquée, blanche.

Contexte (fig.5) bistratifié, dimitique. Hyphes génératrices hyalines, à paroi mince, 2-3 μm de diamètre, ramifiées et cloisonnées. Hyphes squelettiques brun jaunâtre, à paroi épaisse, 2-3 μm de diamètre, ni ramifiées ni cloisonnées.

A la base du champignon (pas de cortex), le contexte est enchevêtré peu sombre (30-50 μm) puis passe à une zone sous-hyméniale plus obscure (8-15 μm) où les hyphes sont plus fermement enchevêtrées et plus ou moins cimentées. Enfin, les hyphes se redressent pour donner l'hyménium supérieur.

Spinules brunes, massives, à extrémité peu pointue mais assez souvent mucronée, 40-70 x 7-8,5 μm , émergentes jusqu'à 35 μm . Des hyphes paraphysoides grêles (1-1,5 μm) peuvent les entourer.

Hyménium : l'ancien, collapsé, comprend des basidioles fripées, longues de 6-10 μm , des spinules et des éléments acanthophysoides brun jaunâtre. Ces éléments naissent de la zone sous-hyméniale sombre. L'hyménium supérieur est formé de basidioles de 6 à 10 (15) μm de haut, de spinules, de très rares basides 12 x 3 μm , à 4 stérigmates de 3 μm et surtout d'abondantes hyphes acanthophysoides à paroi épaissie, de 2 à 2,5 μm de diamètre, non ou peu émergentes (10 μm). Ces sortes d'acanthophyses à paroi mince lorsqu'elles sont jeunes, naissent d'hyphes squelettiques ou génératrices. Localement, des masses cristallines se rencontrent au niveau de l'hyménium.

Spores (sur sporée) cylindriques, légèrement déprimées, 4-4,5-(5) x 1,5-1,8 μm , hyalines, uninucléées (Giemsa), non amyloïdes, blanches en masse.

Sur support indéterminé, Makokou, Gabon, 5 mai 1976, leg. J. Boidin. Holotype : LY 7741 (Herb. Boidin).

AUTRES RECOLTES :

Côte d'Ivoire : LY 7053, sur support indéterminé, Abidjan, 5 novembre 1972, leg. G. Gilles.

Gabon : Makokou (leg. J. Boidin), sur support indéterminé : LY 7734, 4 mai 1976; LY 7750 et LY 7755, 5 mai 1976; sur

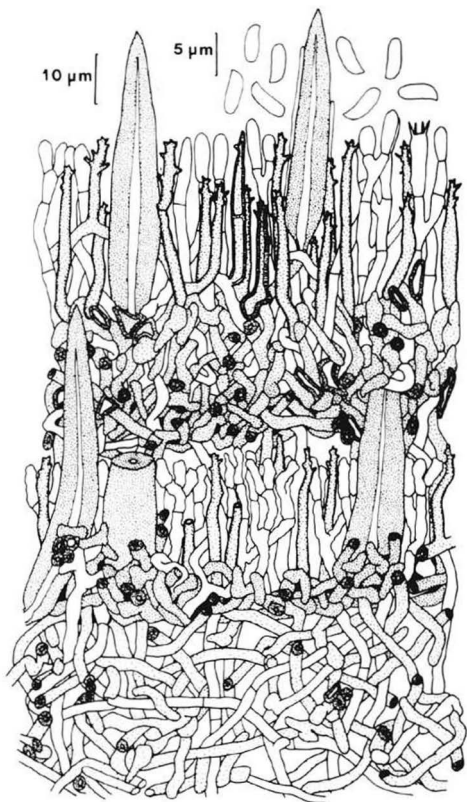


Figure 5 : *Hymenochaete acanthophysata*

Klainedoxa gabonensis : LY 7759, 7 mai 1976; sur *Ancistrophyllum* sp. : LY 7830, 13 mai 1976.

Gabon : Libreville (leg. G. Gilles), sur support indéterminé: LY 9114, 12 janvier 1979; LY 9151, 4 février 1979; LY 9198, 4 mars 1979; LY 9215, 12 mars 1979; LY 9258, 26 novembre 1978; LY 9261, 10 novembre 1978; LY 9273, 8 avril 1979; LY 9298, 22 avril 1979; LY 9310, 28 avril 1979 et LY 9417, 4 juin 1979.

DISCUSSION :

Ces nombreuses récoltes permettent de mieux cerner les caractères de *H. acanthophysata*, à travers les variations de la couleur de l'hyménium et surtout de la stratification ou non des échantillons :

Couleur de l'hyménium: l'abondance des hyphes acanthophysoides, lorsque l'hyménium n'est que très peu ou pas fertile, provoque un net assombrissement de la couleur. Ainsi, LY 7053, LY 7830 et LY 9417 par exemple qui sont à peu près stériles ont une couleur brun rougeâtre foncé (5YR 4/3 à 3/3 et même 2,5YR 2/4 pour LY 9298); cette couleur est renforcée par un aspect velouté souvent très marqué. Lorsque l'hyménium est bien développé par contre, la couleur est plus pâle: soit plus grisâtre (2,5YR 5/2 à 4/2, naval brown R. pour LY 9261 par exemple) soit plus jaunâtre (7,5YR 5 à 6/4, wood brown R. pour LY 7755). Une même récolte montre assez souvent une telle variation de couleur: par exemple, LY 7750 est tantôt chocolat (5YR 3/4) et velouté, tantôt gris rosâtre (5YR 5/2, benzo brown R.) et aride. De façon similaire, LY 7755 (ainsi que 7759) offre un très bel exemple: le centre est en effet gris légèrement rosé (5YR 6/2) alors que le reste de la surface est bai ferrugineux (5YR 4/6).

Stratification: sur un total de 17 récoltes, seules 5 présentent une stratification: LY 7734, LY 9258 et LY 7741 (type) sont bistratifiés. LY 9273 a quelques strates mais seul LY 7750 (dont l'épaisseur atteint 700 μm) montre 6 à 7 hyméniums successifs. Tous les autres spécimens (épais de 30 à 100 μm) n'offrent aucune stratification. Le basidiome a dans ce cas une structure plus simple: à la base, un mince contexte assez sombre (15-20 μm) formé d'hyphes enchevêtrées en tous sens donne naissance à un contexte vertical dont les hyphes se terminent presque toutes en éléments acanthophysoides; ceux-ci, comme les spinules, sont répartis dans toute l'épaisseur du basidiome. Qu'il s'agisse de spécimens stratifiés ou non, la plupart des récoltes sont presque ou totalement stériles: plusieurs sporées n'ont pu être obtenues et les autres n'ont que peu de spores.

En conclusion, *H. acanthophysata* nov.sp. est une espèce

stratifiée ou non, dont le basidiome étalé-adhérent est de couleur brun foncé, parfois velouté, à l'état substérile et plus grisâtre et aride lorsque l'hyménium est normalement constitué (les basides étant de toute façon très dispersées). La présence de nombreuses hyphes acanthophysoïdes est sans conteste le point le plus remarquable avec les spinules trapues et massives.

HYMENOCHAETE HAUERSLEVII Léger nov.sp.

Basidioma resupinatum, primum tenuissimum, e luteo ochraceo, dein paulo crassius, cinnamomeum, frustulosum, laxe ad subiculum e badio ferruginosum adnatum; ambitu indeterminato, concolore; trama laxe intermixta, ex hyphis brunneis, tunica incrassata, x 4-5-(6) μm , septatis ramosisque constante, atque ex hyphis hyalinis, tunica tenui, x 2-3 μm , septatis ramosisque; cortice nullo; crystallis permultis ad et inter hyphas; spinulis brunneis, acutis, longis angustisque, 90-170-(190) x (4)-6-8 μm , usque ad 100 μm emergentibus; hyphis paraphysoides gracilibus, x 2 μm , spinulas scandentibus; basidiis saepius paulum constrictis, 12-17-(19) x 4-5 μm , 4 sterigmatis 5 μm longis; pseudoacanthophysibus tunica tenui vel leviter incrassata, non vel vix emergentibus, 10-13-(20) x 3-5 μm , paucis apicalibus diverticulis 2-3 μm longis praeditis; sporis cylindratis (6)-6,5-7,5-(8) x 2,8-3-(3,2) μm , apiculo saepe manifesto, hyalinis, haud amyloideis.

Holotypus : Copenhague (C), leg. K. Bjørnekaer n° Afr.22, in substrato incognito, in monte Kenya, Kenya, 31-III-1963.

Ethymologie: espèce dédiée au Dr. K. HAUERSLEV qui nous a signalé et communiqué le spécimen.

DESCRIPTION DU TYPE :

Basidiome résupiné, étalé, d'abord très mince (60 μm), ocre jaune (10YR 6/6, brownish yellow) puis un peu plus épais (100-110 μm), cannelle (7,5YR 6/7, cinnamon), frustuleux, laissant apparaître un feutrage bai ferrugineux (5YR 4/6, reddish brown). Marge indéterminée, concolore.

Contexte (fig.6) lâchement enchevêtré d'hyphes brunes à paroi épaisse, 4-5-(6) μm de diamètre, cloisonnées et ramifiées et d'hyphes hyalines à paroi mince, de 2 à 3 μm de diamètre, également ramifiées et cloisonnées. Nombreux cristaux sur et entre les hyphes. Pas de cortex basal.

Spinules brunes aigües, étroites et très longues, 90-170-(190) x (4)-6-8 μm , naissant à différentes hauteurs, émergentes jusqu'à 100 μm . Des hyphes hyalines et grêles de

2 μm de diamètre grimpent souvent le long de ces spinules.

Hyménium formé de basidioles, de basides assez souvent un peu constrictées, 12-17-(19) x 4-5 μm à 4 stérigmates de 5 μm et de pseudoacanthophyses à paroi mince ou légèrement épaissie, 10-13-(20) x 3-5 μm , avec quelques diverticules terminaux (4 à 6 le plus souvent) de 2 à 3 μm de long. Basides et pseudoacanthophyses sont abondantes dans la partie jeune, ocre jaune, du basidiome et beaucoup plus rares dans la partie âgée, cannelle.

Spores (observées sur les coupes car sporée non obtenue) cylindriques, (6)-6,5-7,5-(8) x 2,8-3-(3,2) μm , à apicule souvent bien marqué, hyalines, non amyloïdes.

Sur support indéterminé, Mt Kenya Forest, Kenya, 31 mars 1963, leg. K. Bjørnekaer. Holotype: Afr.22 (C); petit isotype adressé par K. HAUERSLEV à J. BOIDIN en 1973 (LY 7280).

DISCUSSION :

Hymenochaete hauerslevii nov.sp. est un champignon parfaitement caractérisé par ses pseudoacanthophyses qui rappellent celles de *Stereum australe* Lloyd (voir J. BOIDIN, 1960, p. 67). Aucune espèce de *Hymenochaete* possédant de tels éléments hyméniens n'a été décrite précédemment. Un autre caractère intéressant est la grande ressemblance de cette espèce nouvelle avec *H. cinnamomea* (Pers.) Bres. : tout d'abord la couleur cannelle de la surface mais surtout la structure lorsque *H. cinnamomea* est jeune et à l'état unistratifié, c'est-à-dire le contexte lâche, les spinules longues, étroites et très émergentes; même la taille des spores rapproche les deux espèces. Cependant, aucune des récoltes européennes de *H. cinnamomea* unistratifié que nous avons pu examiner ne présente de pseudoacanthophyses, ce qui permet de distinguer aisément *H. hauerslevii*. Il serait intéressant de rechercher si des spécimens avec pseudoacanthophyses ne se cachent pas parmi les récoltes américaines unistratifiées de *H. cinnamomea*.

ABSTRACT

In this paper, species of *Hymenochaete* were studied which possess hymenial hyphidia with "pinnatifid tips" according to BURT's terminology. To the known species with such a feature, *H. pinnatifida* Burt and *H. digitata* Burt, two species are added : *H. acanthophysata* nov.sp. and *H. hauerslevii* nov.sp.. Although these species seem to form a group of their own in our study, it would certainly be excessive to consider them as a natural grouping. We give below a diagnostic key which summarizes the main characters of each species :

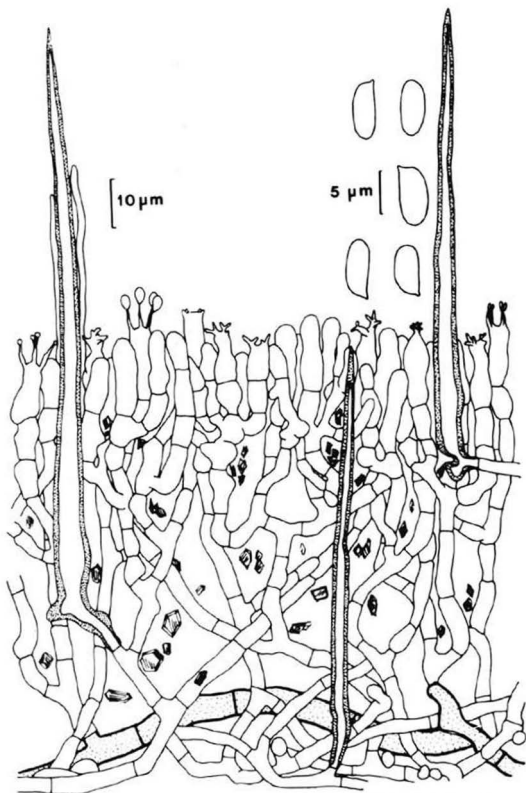


Figure 6: *Hymenochaete hauerslevii*

- 1- Cortex present (Cuticle *sensu* REEVES and WELDEN, 1967)2
- 1- Cortex absent3
- 2- No tomentum (abhymenial hairs); dark subhymenium of tightly interwoven hyphae, context not stratose (but a two-layered stage observed in LY 8091 from Guadalupe) and dendrophyses. West Indies, Gabon, Central African Republic and Guadalupe *H. pinnatifida* Burt.
- 2- Tomentum present; no dark subhymenium; context stratose of several hymenial layers and particular pseudoacanthophyses with dendrophyse-like apicis. Panama
..... *H. digitata* Burt.
- 3- Context not stratose of loosely interwoven hyphae; slender setae 90-170-(190) x (4)-6-8 μm and pseudoacanthophyses hyaline; spores (6)-6,5-7,5-(8) x 2,8-3-(3,2) μm . Kenya *H. haverslevii* Léger nov. sp.
- 3- Context stratose (but found frequently in first-stratum stage); setae somewhat massive, 40-70 x 7-8,5 μm , and very numerous acanthophysate hyphae; spores 4-4,5-(5) x 1,5-1,8 μm . Ivory Coast and Gabon
..... *H. acanthophysata* Léger nov. sp.

ACKNOWLEDGEMENTS

Thanks are due to H. ROMAGNESI for the translation of Latin diagnoses and to Dr. K. HAVERSLEV for sending the specimen from Mt Kenya. We are also very grateful to Dr. J. ERIKSSON and Dr. A.L. WELDEN for critical review and to the Curator of the Farlow Herbarium for the loan of specimens.

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CHLORIDIUM AND SOME OTHER DEMATIACEOUS HYPHOMYCETES
GROWING ON DECAYING WOOD. CORRECTIONS AND ADDITIONS

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Pruhonice near Praha.

Soon after publication of our paper under this title
(in Stud. Mycol. 13. 1976) our attention was drawn to some
homonymies (Index of Fungi 4: 469. 1978) and incomplete-
nesses which we wish to correct here.

CHAETOSPHAERIA FUSISPORA W.Gams & Hol.-Jech., l.c. p. 45
is a homonym of
Chaetosphaeria fusispora (Kawamura) Hino in Bull. Miyazaki
Coll. Agric. For. 4: 191. 1932 (Syn. *Miyoshia fusispora*
Kawamura; *Miyoshiella fusispora* (Kawamura) Kawamura);
a possibly congeneric fungus with much wider ascospores,
26-43 x 6-10 μ m - and
Chaetosphaeria fusispora P. Larsen in Dansk bot. Ark. 14(7):
7. 1952; which probably belongs to *Chaetosphaerella*
E. Müll. & C. Booth.

We therefore rename our fungus
Chaetosphaeria fusiformis W.Gams & Hol.-Jech. nom. nov.
replaced synonym: *Chaetosphaeria fusispora* W.Gams &
Hol.-Jech in Stud. Mycol. 13: 45. 1976.

The genus *CYLINDROTRICHUM* (l.c., p. 48) was incompletely
treated.

Species published before our paper include:
Cylindrotrichum proliferum Matsushima, Icones Microfungorum
a Matsushima lectorum: 47. 1975,
Cylindrotrichum triseptatum Matsushima, l.c.: 48. 1975 and
Cylindrotrichum triseptatum M.B.Ellis, More dematiaceous
Hyphomycetes: 470. 1976 (a homonym) = *C. ellisii* Morgan-
Jones in Mycotaxon 5: 490. 1977.

Since our publication four further species were described:
Cylindrotrichum oblongisporum Morgan-Jones in Mycotaxon
5: 487. 1977, which should be compared with our

C. zignoëllae,

Cylindrotrichum gori Lunghini in *Micol. ital.* 8: 25. 1979,
distinct from the previously known species by smaller,

often slightly curved conidia, 5.7-9.9 x 1.9-3.0 μm ,

Cylindrotrichum helisciforme Marvanová in *Trans. Br. mycol.*

Soc. 73: 368. 1979, with tricuspidate apical cells of the
conidia, and

Cylindrotrichum curvatum Morgan-Jones in *Mycotaxon* 12:

250. 1980, with curved conidia, 10-14 x 2.5-4.0 μm .

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TRICHOPHYTON RAUBITSCHKII, SP. NOV.

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A urease-positive Trichophyton isolated from skin scrapings has been found to be sufficiently different from existing taxa within this genus to warrant its establishment as a new species, T. raubitschekii.

MATERIALS AND METHODS

TEST ORGANISMS - Forty-one isolates of a urease-positive Trichophyton recovered from skin scrapings and 10 isolates each of T. rubrum and T. mentagrophytes recently obtained from clinical specimens were used in this study. The purity of all isolates was established by single spore isolation and other methods (2,5,8). The identification of T. rubrum and T. mentagrophytes isolates was established by standard methods (6,9) before they were selected for this investigation. All isolates were maintained on peptone-dextrose agar (Sabouraud dextrose agar: 4% crude glucose, 1% Neopeptone, 2% agar) at 26-28°C. Seven-day-old cultures on this medium served as the source of inocula for all morphologic and physiologic tests.

MORPHOLOGY - The gross and microscopic morphology of each isolate was observed after 2 and 4 weeks' incubation on peptone-dextrose agar at 26-28°C.

GROWTH AND PIGMENT FORMATION - A pinpoint inoculum from a 7-day-old culture was transferred to the center of

each agar slant. All slants were incubated at 26-28°C. The extent of growth and pigmentation on bromocresol purple-casein dextrose agar (5) and corn meal-dextrose agar (3) were assessed 7 days after inoculation. The extent of growth on lactose agar (4) was noted at weekly intervals for 3 weeks.

UREA HYDROLYSIS TEST - Ability to hydrolyze urea within 7 days was determined by the urea broth method of Kane and Fisher (7).

HAIR PERFORATION TEST - Following the technique described by Ajello and Georg (1), autoclaved human hair was examined for wedge-shaped perforations at weekly intervals for 4 weeks.

MATING REACTIONS - Mating tests with strains of *Arthroderma simii* and *A. benhamiae* were performed according to the method of Stockdale (10) on the oatmeal-salts agar described by Weitzman and Silva-Hutner (11).

RESULTS AND DISCUSSION

Division: Fungi imperfecti
Form class: Hyphomycetes
Form order: Moniliales
Form family: Moniliaceae

Trichophyton raubitschekii Kane, Salkin, Weitzman et Smitka, sp. nov. Colonia in agaro peptone-dextrose, postquam 14 dies apud 26°C lapsi sunt, 28 mm in diam. fit; plana, tenuis, lenis usque ad granosum in textu, cum centro fulvo sublatoque. Pars aversa erat sanguinea (#3 in charta colorum Raynor Mycology RMCC). Margo coloniae erat distinctus cum angusta fascia sine colore.

Post 28 diebus, colonia erat 53 mm in diam.; plana, tenuis, lenis, cum centro sublato exquo sulci radiantes ad peripheriam pertinuerant. Tres zonae colorum notatae sunt. Centrum erat submurinum, (RMCC #117) quod ad zonam mediam, colore grisea-sepiaceam (RMCC #106) pertinuit terminatumque ad aream peripheriam, lavendulo-griseam (RMCC #125). Margo erat distinctus cum fascia sine colore 3 mm in latitudinem. Pigmentum in parte aversa sanguineum (RMCC #3) remansit.

Sporulatio in agaro peptone-dextrose erat crebra et in primis et in subsequentibus subculturis. Macroconidia erant longa et angusta, 45-51 x 4.8-6.3 um, cum parietibus teribus, fines obtusi, et ex 5-9 cellulis composita. Microconidia erant in forma variabilia (clavata, subsphaeralia, globosa), 4.8-6.4 x 3.2-4.8 um, aut sessilia aut in brevibus conidiophoribus, a latere ferebantur. Holotypus OMH 1094, teleomorph ignotum.

Habitat: Man
Holotype: OMH 1094, isolated from human skin scrapings, Toronto, Ontario, Canada.

Living cultures of the holotype have been deposited in the American Type Culture Collection, Rockville, Maryland (accession number ATCC 42631); the University of Alberta Mold Herbarium, Edmonton, Alberta, Canada (accession number UAMH 4314); and the Medical Mycology Laboratory Culture Collection, Ontario Ministry of Health, Toronto, Ontario, Canada (accession number OMH 1094). The epithet T. raubitschekii was chosen to honor the late Dr. F. Raubitschek, dermatologist, medical mycologist, and teacher.

The colony on peptone-dextrose agar after 2 weeks at 26°C was 28 mm in diameter, flat, thin, velvety to granular in texture, with an elevated, buff-colored center (Figure 1A). The reverse was blood-red, #3 on the Raynor mycology color chart (RMCC: Commonwealth Mycological Institute, Kew, Surrey, England). The colony margin was distinct, with a narrow colorless band.

After 4 weeks the colony was 53 mm in diameter, flat, thin, velvety, with a raised center from which grooves radiated to the edge (Figure 1B). Three color zones were noted: a pale mouse-gray center (RMCC #117), a grayish sepia (RMCC #106) intermediate zone, and a lavender-gray (RMCC #125) peripheral area. The margin was distinct, with a 3-mm colorless band. The reverse remained blood-red (RMCC #3).

Sporulation on peptone dextrose agar was abundant in both primary and subsequent subcultures. Macroconidia were long and narrow, 46-51 x 4.8-6.3 μ m, with smooth, parallel-sided walls, blunt ends, and five to nine cells (Figure 2). Microconidia were variable in shape (clavate, subspherical, or globose), 4.8-6.4 x 3.2-4.8 μ m, and borne laterally, either sessile or on short conidiophores (Figure 3).

Teleomorph unknown.

Pronounced urease activity was observed within 7 days of incubation at 26-28°C. Growth on bromcresol purple-casein dextrose agar was restricted, and a reddish pigment developed in the center of the colony. Brown reverse pigmentation was observed on cornmeal dextrose agar. Growth on lactose agar was restricted after 3 weeks' incubation.

Mating of 12 isolates of T. raubitschekii with strains of A. simii mating type (A) resulted in the formation of either pseudogymnothecia or gymnothecia with immature, irregular asci. In contrast, no reaction occurred when the same isolates were paired with tester strains of A. benhamiae or mating type (a) of A. simii.

The key morphologic and physiologic features of T. rubrum, T. raubitschekii, and T. mentagrophytes are compared in Table 1. Trichophyton raubitschekii is similar to T. mentagrophytes in urease activity but differs in its inability to perforate hair *in vitro*, its restricted growth on bromcresol purple-casein dextrose agar, and its lack of sexual reaction with A. benhamiae. While resembling T. rubrum in its reddish pigmentation on glucose-peptone agar and sexual reaction with A. simii (A),

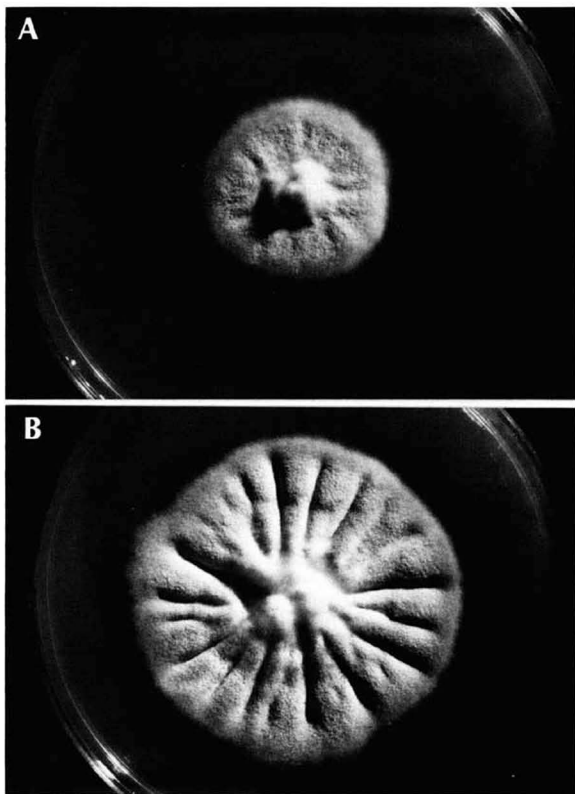


Figure 1: Culture of *T. raubitschekii* on peptone-dextrose agar at 26-28°C after (A) 2 weeks and (B) 4 weeks' incubation.

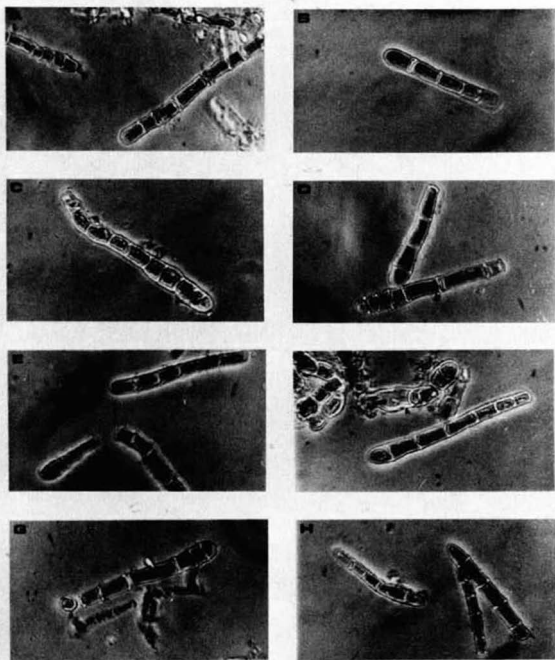


Figure 2: Macroconidia (A-H) on peptone-dextrose agar. Phase contrast, x 400.

T. raubitschekii differs from *T. rubrum* in four important characteristics: it is urease-positive; its growth is restricted rather than spreading on lactose agar; its pigmentation on casein dextrose agar is brown rather than red; and it produces abundant macroconidia.

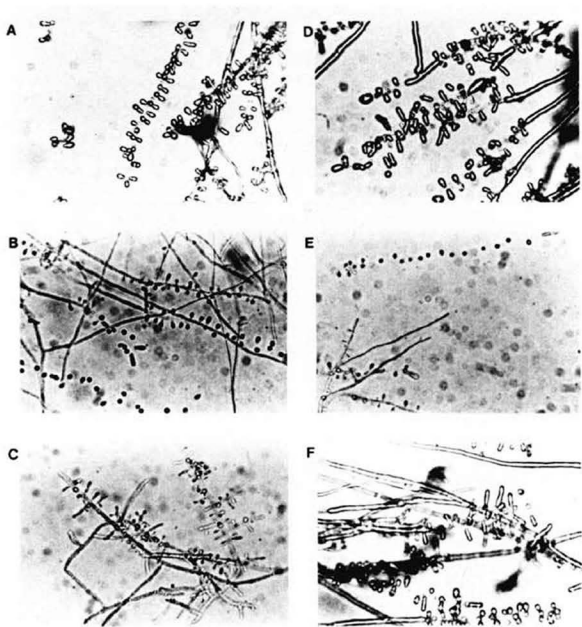


Figure 3. Microconidia (A-F) on peptone-dextrose agar, x 400.

Table 1. Comparison of morphologic and physiologic characters of T. rubrum, T. raubitschekii, and T. mentagrophytes

Character	<u>T. rubrum</u> (10) ^a	<u>T. raubitschekii</u> (41)	<u>T. mentagrophytes</u> (10)
Urease activity	Negative	Positive	Positive
Hair perforation	Negative	Negative	Positive
Pigment on cornmeal dextrose agar	Red	Brown	-
Growth on lactose agar	Spreading	Restricted	ND ^b
Growth on bromcresol purple-casein dextrose agar	Restricted	Restricted	Spreading
Macroconidia	Rare, pencil-shaped, 4-6 x 15-30 um	Abundant, pencil-shaped, blunt-ended, 4.8-6.3 x 46-51 um	Occasional, clavate, 6-8 x 20-50 um
Microconidia	Clavate, 2-3 x 3-5 um	Variable, 4.8-6.4 x 3.2-4.8 um	Variable, 3.6-4.2 x 2.2-2.8 um
Colony texture	Cottony, velvety	Velvety, granular	Cottony, velvety, granular, powdery

^aNumber of isolates used in this study.

^bND = not done in this study.

ACKNOWLEDGMENT

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CYRENELLA ELEGANS GEN. ET SP. NOV., A DIKARYOTIC ANAMORPH

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ABSTRACT

Cyrenella elegans Gochenaure is described and illustrated from sand around the base of the mushroom, *Laccaria trullisata* (Ellis) Singer. Its bright orange colony bears hyphae with clamp connections, small, obovate, dikaryotic conidia with 3 or 4 apical appendages and a single basal one, and clusters of globose chlamydospores. Monokaryotic blastospores may be produced. The organism is oxidative, mesophilic and acidophilic, utilizes nitrate, and requires the vitamins p-aminobenzoate and thiamine.

INTRODUCTION

Several *Rhodotorula*-like isolates were obtained from dilution plates prepared with sand scraped from the submerged stipe of *Laccaria trullisata*. One of these colonies during subculture was found to produce hyphae with clamp connections and small appendaged conidia. A new form genus is introduced for this isolate as no fungus with similar characteristics appears to have been described in the literature.

The holotype, a desiccated agar culture (No. 785), is deposited in The New York Botanical Garden Herbarium (NYBG), Bronx, N. Y. Living representative strains (isotypes) are deposited in The American Type Culture Collection (ATCC), Rockville, Md. and The National Culture Collection (NCC), Commonwealth Mycological Institute, Kew, England.

MATERIALS AND METHODS

Morphological descriptions are based on a strain that developed from a single conidium obtained from the original isolate. This dikaryon and a cloned blastospore strain derived from it were cultured on 2% malt extract agar with 2% glucose (MEA) at 20°C and examined microscopically using brightfield and phase optics. Measurements of microscopic structures are the mean of 50 determinations + the standard deviation. Colors refer to plates in the Methuen Handbook of Colour (10).

Nutritional studies using the dikaryon and blastospore cultures employed a basal medium composed of Bacto-Yeast Nitrogen Base without Amino Acids and Ammonium

Sulfate (Difco Laboratories, Chicago, Il.) and solidified with Bacto-Purified Agar. This was supplemented with glucose (1%) in tests for utilization of various nitrogen sources and with Bacto-Vitamin Free Casamino Acids in tests for utilization of various carbon sources. All basal media and test compounds were filter sterilized except compounds 17 through 23 (Table II) which were heat sterilized at 121°C. Vitamin requirements were determined for cultures grown on Bacto-Vitamin Free Yeast Base solidified with BBL-Ion Agar No. 3 (Baltimore Biological Laboratories, Cockeysville, Md.) and supplemented with 0.2% Bacto-Vitamin Free Casamino Acids and the appropriate vitamins. Vitamin-depleted inoculum was used.

Environmental studies, using the blastospore stage only, were done in liquid culture employing a medium composed of Bacto-Casamino Acids (0.5%) and glucose (1%). Growth was measured spectrophotometrically at a wave length of 450 nm. Additional temperature data were collected using the dikaryon grown on MEA.

Chlamydospore germination was studied using the technique employed by Newell and Fell (12) for *Rhodosporidium* teleospores. Chlamydospores produced in malt extract broth were collected by centrifugation, washed several times, resuspended in sterile distilled water and stored for up to 5 months at room temperature and 7°C. Aliquots were removed at various times and streaked on Bacto-V8 Juice Agar and MEA, incubated at 12°C and 25°C, and examined periodically over 4 days.

Nuclear number was determined using Furtado's Toluidine Blue (5). Cells were grown on MEA for 2 to 5 days at 30°C and then stained in wet mounts or smeared on cover glasses and pretreated for 5 min with 1N HCl at 60°C before being stained.

TAXONOMY

CYRENELLA Gochenaur, forma-gen. nov., nomen anamorphosis, Deuteromycotina, hyphomycetes pertains.

Mycelium ex hyphis hyalinis, ramosis, leviter tunicatae compositum zygo-desmatibus praeditae. Conidiophora hyalina, macronemata, solitariae vel congregatae, erecta, irregulariter ramosa. Cellulae conidiogenaе monoblasticae, apicalia et lateralia, globosae, ovoideae vel cylindricae vel mixtae, vacuae post dehiscentem conidiorum. Conidia hyalina, obovata, et cum brachiis apicalibus radiantibus, 2-5 et uniappendici basilari. Chlamydosporae et blastosporae sint productae.

Species typica: *Cyrenella elegans* Gochenaur

ETYMOLOGY: From the Greek, *Cyrene*, a mythical water sprite, in reference to the probable habitat of the fungus and the resemblance of its conidia to those of the aquatic hyphomycetes.

CYRENELLA ELEGANS Gochenaur, forma-sp. nov. Figs. 1-2.

Coloniae in agarо maltoso post dies decem ad 20°C 4-5mm diam. attingentes, humectae, calendulinae,

compactae, in centrum acervulatum, marginis planis, hyphis aereis raris. Hyphae immersae leves, zygoesmatibus praeditae, ad $2\mu\text{m}$ crassae. Conidiophora hyalina, macronemata, solitaria vel congregata, erecta, irregulariter ramosa, ad $100\mu\text{m}$ longa, $3-4\mu\text{m}$ latae. Cellulae conidiogenae monoblasticae, apicalia et lateralia, ovoideae vel cylindricae vel mixtae, vacuae post dehiscentem conidiorum, $5-12\mu\text{m}$ longae, $3-4\mu\text{m}$ latae. Conidia hyalina, unicellularia, binucleolata, $10 \times 4\mu\text{m}$, cum brachiis apicalibus radiantibus $3-4$, $13-20\mu\text{m}$ longis, $1\mu\text{m}$ latis ad basem et $0.5\mu\text{m}$ latis ad apicem, et uniaappendici eccentrica basilari, $7-11 \times 0.8\mu\text{m}$. Chlamydosporae abundae, solitariae et catenatae, intercalares et terminales, frequenter gregariae, hyalinae vel melleae pallidulae, binucleolatae, globosae vel subglobosae leves et crassitunicatae, $5.5-6.5\mu\text{m}$ latae. Status sexualis ignotis.

In arena, sub *Laccaria trullisata*, Hempstead Lake State Park, Long Island, New York, October, 1978. Holotypus cultura pura siccata (numero 785), NYBG Herb.

ETYMOLOGY: From the Latin *elegans*=graceful, describing the appendages of the conidia.

DEVELOPMENT

Germination of the conidia of *C. elegans* occurs within 24 hr on MEA at 20°C . One to 4 germ tubes arise apically between the appendages and/or from the base, but rarely from the lateral walls. Clamped septa and branches develop where the germ tubes emerge from the conidium. As the colony ages, conidia may germinate in place producing apically, laterally or from the base, one or more conidiogenous cells that develop conidia (Fig. 2C). Cytoplasm flows into the conidiogenous cell leaving the original conidium empty. This process may continue through several cycles, so that short chains of empty conidia are common. As a result of this proliferation, the color of the central area of the colony is diluted and much lighter than the periphery (Fig. 2A). Growth is slow. Colonies attain a diameter of 3 mm in 7 da and approximately 24 mm in 35 da. They are Persian to chrome orange (6A 7-8) with a raised, rugulose central region consisting of pulvinate masses of conidiophores and conidia and a narrow plane margin. With age, a broad, radiately wrinkled periphery composed of mycelium and chlamydospores develops (Fig. 2A). It is somewhat fibrous in appearance with small mounds of sporulating cells scattered over the surface or coalesced to form larger irregular or wedge-shaped patches. The surface is moist. Aerial hyphae are rare or absent; when present they are usually aggregated into tufts. The *mycelium* is hyaline and consists of sinuous and straight hyphae of uniform width, $2.0 \pm 0.3\mu\text{m}$ diam, with smooth, slightly thickened walls. Septa with clamp connections (Fig. 2B) first occur approximately $35 \pm 11\mu\text{m}$ from the tip and then at intervals of approximately $50\mu\text{m}$. The cells are binucleate. A dolipore apparatus was not revealed in preparations stained with ammoniacal Congo-Red (11).

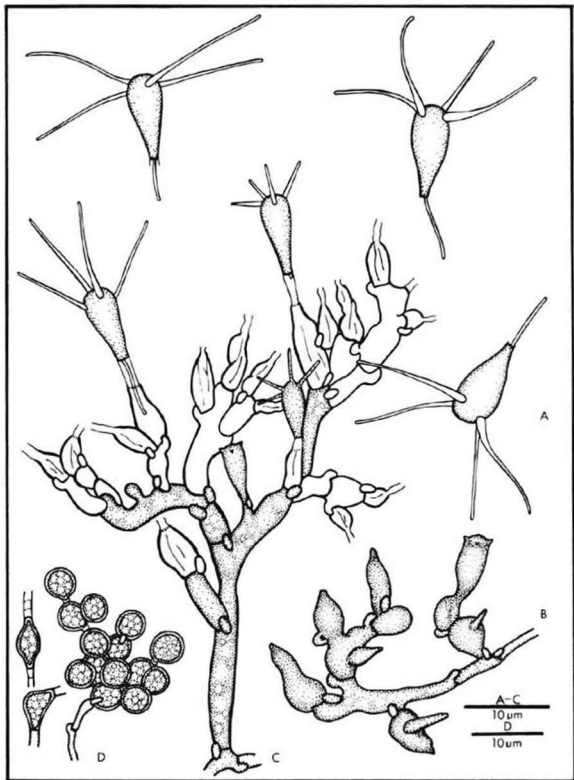


Fig. 1. *Cyrenella elegans*.

(A) Conidia. (B) Stages in the development of conidiogenous cells. (C) Mature conidiophore. (D) Single and clustered chlamydospores. Unstippled areas are devoid of cytoplasm.

Adventitious simple septa are common in older hyphae and coralloid or haustorial branches similar to those shown by Olive (13) for *Filobasidium* may occur.

Conidiophore development begins within 2-3 da. The conidiophores arise as erect, branched hyphae indistinguishable at first from the vegetative cells. They are hyaline, of variable length and produce thumb-shaped terminal and lateral branches that stain intensely with basic dyes (Fig. 1B). These branches are separated from the conidiophore early in development by clamp connections. They also may be produced individually along the vegetative hyphae (Fig. 2B) or less commonly from a clamp. They may differentiate directly into conidiogenous cells or may branch once or twice, each branch becoming a conidiogenous cell separated from its parent cell by a clamped septum. The irregularly branched conidiophore at maturity (Fig. 1C) is generally less than 100 μ m tall, 3-4 μ m wide at the base and is best observed using phase contrast optics since its upper half consists of cells devoid of cytoplasm and somewhat collapsed. Conidiophores are often aggregated into sporodochial-like clusters that arise from a basal mass of intertwined hyphae and chlamydospores. The conidiophores produce single or clusters of 2-3 conidiogenous cells laterally and terminally along their branches. The *conidiogenous cells* are phialide-like, binucleate, have a thick-walled body and a thinner-walled neck that is approximately 1-4 μ m long and 1-2 μ m wide. They vary in form from globose to ampulliform to cylindrical and measure 2-4 μ m wide, 5-12 μ m long including the neck. Each produces a single conidium.

Mature conidia appear within 4-5 da. *Conidia* (Fig. 1A) are blastic, orange in mass but hyaline by transmitted light, binucleate (Fig. 2C), obovate, thin-walled, 10 x 4 μ m (9.7+1.2 x 4.2+0.5 μ m) and bear 3-4 long, narrow, divergent, flexible, apical appendages, 16+2.4 μ m long, 1 μ m wide at the base and tapering to 0.5 μ m wide at the apex, and an eccentric basal appendage of uniform width, 8.9+2.1 x 0.8 μ m. At 20°C, approximately 33% of the conidia examined had 3 apical appendages; 66% had 4. At temperatures above 25°C, about 15% of the conidia had one, 2 and 5 arms.

Chlamydospores (Figs. 1D, 2D) appear after 3 da. They are single and catenulate and arise terminally, intercalary, or less commonly from clamps. They typically form in large compact clusters behind the margin of the colony and just above or below the agar surface. They are subglobose to globose, 6+0.4 μ m diameter, binucleate, hyaline to pale amber, thick-walled and filled with guttulae. Single ones are often oval and/or unsymmetrical and borne between several adventitious septa that form as the cytoplasm retracts into the developing chlamydospore.

Chlamydospores germinate within 3 da at 20°C (Fig. 2E). A single germ tube that branches and develops clamps where it exits from the spore is produced and a typical colony with conidia develops within 5 da. Chlamydospores remain viable after storage for up to 5 mo at room temperature in distilled water. Those held at 7°C exhibited a progressive

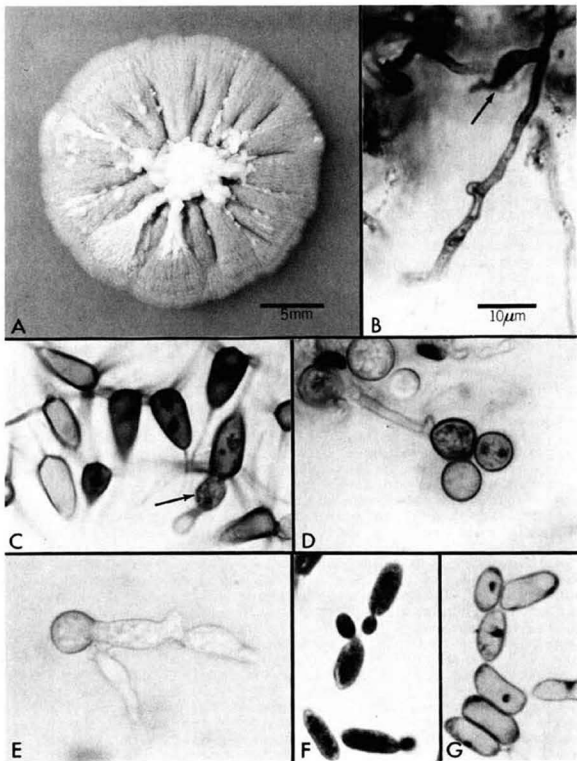


Fig. 2. Cyrenella elegans.

(A) Colony on malt extract agar after 35 da. at 20°C.
 (B) Vegetative hypha with a clamp connection. Arrow designates a young conidiogenous cell.
 (C) Binucleate conidia. Arrow designates a young conidiogenous cell and secondary conidium produced during in situ germination. Three empty conidia appear on the left.

decrease in the percent germinating after 45 da storage and showed no germination after 5 mos. Karyogamy, probasidia and soredia were not induced by these conditions, nor do these spores resemble, in either coloration or mode of formation, the teleospores produced by members of the genus *Rhodosporidium* (2, 3, and comparison in culture with *Rh. dacryoidum* Fell, Hunter, Tallman ATCC 24502 & ATCC 24503). In view of the above, it is considered likely that these structures are true chlamydospores and not the teleomorphic state of *C. elegans*.

Dedikaryotization occurs spontaneously, but infrequently, as the culture ages; a uninucleate yeast phase results. It can be detected readily by streaking material onto an agar medium containing raffinose, a sugar utilized less well by the filamentous stage. Origin of the blastospores is unknown. Subjecting the dikaryon to flooding with water or anaerobic or microaerobic conditions while growing it on media with various C/N ratios and low or abundant in nutrients failed to consistently induce the yeast phase.

The yeast colonies, like the dikaryon, grow slowly, reaching a diameter of 8-10 mm on MEA at 20°C after 30 da. They are slightly elevated when viewed in cross section and show faint radial lines. Their surface is semi-glossy, deep orange to Persian orange (6A8-7A8) in color with a slightly deeper carrot red reverse (7B7). The yeast cells (Fig. 2F-G) are unicellular, monokaryotic, non-encapsulated, elliptical to cylindrical with rounded ends, less commonly obclavate, $9.0 \pm 1.8 \times 3.2 \pm 0.8 \mu\text{m}$, and reproduce by terminal and subterminal budding. Fermentation is absent. In ME broth, most cells are single and only rarely are short chains of cells observed. No pellicle is produced.

Evidence suggests that the nuclei in the dikaryon are identical. A dikaryophase is not reestablished when yeast cells from randomly selected colonies are mixed on cornmeal agar. However, it develops spontaneously in every cloned yeast colony that is over a few days old. Cells in the center of a colony at the agar surface become binucleate. It is uncertain whether these occur following conjugation between two cells or by failure to bud during mitosis. Observation of empty single yeast cells with an attached binucleate conidium suggests that the latter is the case. A sparse mycelium always develops under every yeast colony but its further expansion is limited unless the overlying cells are scraped away. All cells on the surface of a colony remain monokaryotic so that the yeast phase can be maintained indefinitely in culture but always contaminated eventually with the dikaryon.

(D) Binucleate chlamydospores. (E) Germinating chlamydospore. (F) Budding blastospores. (G) Uninucleate blastospores. Material in photographs B-G was stained with toluidine blue. Cells in photograph G were pretreated with 1N HCl for 5 min at 60°C.

NUTRITION

C. elegans is an acidophilic, mesophilic, obligate aerobe. Growth occurs over a wide range of H-ion concentrations. It is best between pH 3.5-6.0 and absent below pH 2.5 and above pH 7.4. Conidia do not germinate at 5°C and 37°C. Pigmentation is more intense and colonial and cellular development best between 18-23°C. Growth is fastest at 30°C; absent above 34°C.

Table I. Growth of *Cyrenella elegans* on various nitrogen sources^{a)}

Compound	Growth	Compound	Growth
1. Potassium nitrate	+	6. Glutamic acid	+
2. Potassium nitrite	-	7. Asparagine	+
3. Ammonium sulfate	+	8. Casamino acids	++
4. Urea	+	9. Gelatin	++
5. Aspartic acid	+	10. Sodium Caseinate	++

a) Cultures were incubated for 5 da at 28°C.

++ Growth equal to growth on casamino acids

+ Growth weak but better than growth on a medium lacking a nitrogen source

- No growth

C. elegans utilized all nitrogen sources tested with the exception of potassium nitrite (Table I). Growth and development are best on a mixture of amino acids. The proteins, gelatin and sodium caseinate, are utilized but diffusible hydrolases were not detected when colonies on these substrates were flooded with acidified HgCl₂. A variety of compounds can serve as the carbon-energy source (Table II). Growth and development are best on glucose or fructose. Krebs Cycle acids (Nos. 37-38) are assimilated but not compounds with ester linkages (Nos. 23-24). Among the polysaccharides (Nos. 17-22), only the fructosan inulin and the starches (Nos. 18-19) were weakly degraded. Diffusible exoenzymes were not detected when colonies growing on the latter compounds were flooded with an IKI solution. Patterns of carbon and nitrogen assimilation by the blastosporic and dikaryotic stages are identical. The exogenous vitamins, p-aminobenzoate (PABA) and thiamine, are required for growth of both the mycelial and yeast phases. High concentrations of folic acid can be substituted for PABA but not a mixture of amino acids and purines.

DISCUSSION

Cyrenella elegans is a member of the Basidiomycotina because it produces mycelium with clamp connections but the absence of a telemorphic state makes its phylogenetic position in the subdivision uncertain and any attempt at ordinal placement a purely speculative venture.

Table II. Growth of *Cyrenella elegans* on various carbon-energy sources^{a)}

Compound	Growth	Compound	Growth
1. Xylose	++	22. Chitin	-
2. L-Arabinose	++	23. DNA	-
3. Ribose	-	24. Triglyceride	-
4. Glucose	++	25. Fucose	-
5. Fructose	++	26. L-Rhamnose	++
6. Mannose	++	27. Glucosamine	-
7. Galactose	-	28. Ethanol	-
8. L-Sorbose	-	29. Glycerol	++
9. Trehalose	+	30. Erythritol	-
10. Sucrose	++	31. Sorbitol	++
11. Maltose	++	32. Mannitol	++
12. Melibiose	-	33. Inositol	-
13. Cellobiose	+	34. Galactitol	-
14. Lactose	-	35. Acetate	-
15. Melezitose	++	36. DL-Lactate	-
16. Raffinose	+	37. Succinate	++
17. Inulin	+	38. Citrate	+
18. Amylose	+	39. Arbutin	+
19. Amylopectin	+	40. Salicin	+
20. Pectin	-	41. Vanillin	-
21. Cellulose	-	42. Casamino acids	-

a) All compounds were D-isomers except where noted. Cultures on compounds 17-24 were incubated for 14 da at 28°C, all others for 5 da at 28°C.

++ Growth equal to or better than growth on glucose

+ Growth weak but better than growth on a medium lacking a carbon source

- Growth equal to or less than growth on a medium lacking a carbon source

The blastosporic stage could be readily accommodated in the genus *Rhodotorula* (14) if the dikaryophase were lost. It resembles *R. aurantiaca* (Saito) Lodder and *R. lactosa* Hasegawa in producing cylindrical cells, in colony color, in utilizing nitrate, cellobiose and maltose and in requiring PABA and thiamine. It differs from the former in being galactose negative and L-rhamnose and raffinose positive and from the latter in being unable to utilize melibiose and lactose. The requirement for PABA is confined to a very few wild-type fungi, certain species of *Rhodotorula*, *Blastocladia pringsheimii* Reinsch and single strains of *Saccharomyces cerevisiae* Hansen (4). Because this requirement is so uncommon, there appears to be strong selection pressure against such fungi in nature. Its simultaneous occurrence in *Rhodotorula* and *Cyrenella* may signify a much closer relationship between these genera than is evident simply on the basis of similarities in morphology and nutritional patterns.

C. elegans was obtained from sand along the shore of

a fresh water lake in an area that is regularly submerged each spring for several months. Even though it came from a terrestrial habitat, *C. elegans* has several characteristics in common with organisms isolated from aquatic environments. Its thin-walled, tetra-radiate-like conidia bear appendages in a pattern reminiscent of that evolved by many fresh and a few salt-water fungi. They are remarkably similar to the basidiospores of the marine fungus *Nia* in size and shape but not development (9) and superficially resemble the conidia of aquatic hyphomycetes belonging to the genera *Acaulopage* Drechsler, *Clavatospora* S. Nilsson and *Clavariana* Nawawi. Diffusible exoenzymes are a characteristic feature of terrestrial fungi (1, 8) but would be a distinct disadvantage for an aquatic organism since such proteins and their digestion products would be lost into the water. *C. elegans* produces no diffusible hydrolases when utilizing starch, gelatin and casein. Since its hydrolases must be retained in protective association with the cell wall, it would be well adapted for life in dilute aqueous environments. Finally, *C. elegans*' requirement for PABA occurs less frequently among terrestrial fungi than among aquatic ones. In *R. aurantiaca*, for example, only strains of marine origin demonstrate an absolute need for PABA (14). These features suggest that *C. elegans* is not a terrestrial fungus but is an aquatic or possibly an amphibious one, whose propagules were trapped in the sand as the lake water receded, persisting there until isolated several months later.

If *Cyrenella* is an aquatic genus, it is an extremely uncommon one. Dr. R. Bandoni of the University of British Columbia (personal communication) reports that *C. elegans* appears to be identical to a fungus he collected from a driftwood chip on the shore of the Iowa River about 25 years ago. Other workers who have examined the conidia report they do not remember seeing spores of exactly this type in foam samples nor have illustrations of these spores appeared in the many works describing propagules from this environment (6, 7). With its slow growth rate in culture, superficial resemblance to *Rhodotorula* and tiny conidia, it is understandable why a *Cyrenella* could be overlooked.

ACKNOWLEDGMENTS

I wish to thank the following individuals for their aid: Dr. R. Bandoni for studying cultures of *Cyrenella* and offering many helpful suggestions as well as kindly reviewing the manuscript; Drs. W. Bridge Cooke, J. Leland Crane, C. T. Ingold, R. H. Petersen, and Shun-ichi Udagawa for examining slides of the fungus; Dr. C. T. Rogerson for assistance with the taxonomy; Dr. A. H. Brenowitz for taking the photomicrographs; Prof. L. Ascher, Language Dept. for correcting the Latin diagnosis; Ms. Julie Anderson, a former undergraduate in my laboratory who studied *Cyrenella* with me as part of her honors thesis; and Mrs. W. Jaworski for typing the camera-ready copy.

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REVUE DES LIVRES

par

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GENERA OF HYPHOMYCETES, par J.W. CARMICHAEL, W.B. KENDRICK, I.L. CONNERS and Lynne SIGLER, 390 p., 129 pl., 8°, reliure spirale, 1980, The University of Alberta Press, Edmonton, Alberta, Canada. Prix \$ 21.-.

Cet ouvrage est une seconde édition entièrement revue et complétée de la première présentation par les premiers auteurs d'une compilation illustrée et critique de la littérature originale des genres de Hyphomycètes dans *The Fungi* de Ainsworth, Sparrow and Sussman, Tome IV A, en 1973. Les auteurs ont recensé cette fois plus de 2000 noms de genre, contre 1541 en 1973. Les 61 planches de figures d'alors sont devenues 129 aujourd'hui. Après une introduction sur la méthode et la classification des types conidiogénétiques, on y trouvera une liste documentée des noms de genres, avec date, référence, type morphologique, hôte, espèce type, statut nomenclatural et synonymie, avec références. Cette liste distingue les genres acceptés par les auteurs des noms rejetés ou douteux. Viennent ensuite les illustrations, soit reproduites des descriptions originales, soit dessinées à partir de publications récentes ou encore du type. Les 853 figures représentent pour la plupart les espèces types des genres et sont identifiées à l'espèce. Le grossissement n'est pas donné mais n'est pas indispensable à ce niveau. Suivent encore 5 index: un index des genres par type conidiogénétique, un index des connections entre téléomorphes (stades sexués) et anamorphes (stades conidiens), une liste bibliographique, une liste des abréviations et une liste des figures. Les auteurs ont suivi la nomenclature illégale basée sur Persoon, *Synopsis Fungorum*, de 1801 comme point de départ, contrairement à l'art. 13, mais combien souhaitée par beaucoup. Le livre remplace les guides des genres publiés jusqu'à présent. Il est fort bien imprimé. Sa reliure en spirale s'inspire de celle du Barnett, *Illustrated genera of Imperfect Fungi*, mais on pourra le regretter. Vu son évidente utilité et son prix assez modique, le livre aura sans doute une large diffusion.

Les auteurs l'ont dédié à la mémoire de *Luella K. Woresub*, mycologue du Biosystematics Research Institute d'Ottawa, tant appréciée de tous.

CHAMPIGNONS D'EUROPE, par M. SVRCEK et J. KUBICKA, adapté en français avec coll. de G.L. HENNEBERT et al., 296 p., 448 ph. col., 8°, cartonné, 1980. Ed. Elsevier-Sequoia, B-1940 St Stevens-Woluwe, Belgique.

Ce guide des champignons est remarquable par le grand nombre d'espèces décrites - 448 - chacune convenablement illustrée d'une photo en couleurs dans le milieu naturel. Les descriptions sont précises

et donnent les dimensions sporales. Une introduction illustrée retrace la structure des champignons et leur classification, donne les conseils au récolteur et le documente abondamment sur la toxicité des champignons. Offrant un éventail exceptionnellement large des champignons d'Europe, tant de montagne que de plaine, ce guide surpasse un grand nombre d'autres moins complets.

ECOLOGY AND DISTRIBUTION OF FUNGI, Scripta Mycologica n° 9, 144 p., 12°, broché, 1980, Academie des Sciences d'Estonie S.S.R., Tartu Estonie, Prix. Rbl. 1,10.

Ce fascicule est intitulé en fonction des deux premiers articles qu'il comprend : *The composition and seasonal dynamics of the fungal cover on mineral soils and Trophic groups of Estonian Agarics*, par K. Kalamees. Dans le premier travail, l'auteur relève les flores d'agariques dans divers types de forêts et de prairies jusqu'à la tourbière et le marais et note sa composition et sa richesse spécifique à chaque écotype végétal. Dans un second article, il regroupe les Agaricales en 13 groupes trophiques, les mycorrhizogènes, les saprophytes de divers types de substrat et les parasites étant les principaux d'entre eux. Le fascicule se termine par une révision du genre *Lasiobelonium* par A. Raitviir, où l'auteur combine diverses espèces et en décrit cinq nouvelles. Texte en anglais.

THE CHEMOSYSTEMATICS OF THE LICHEN GENUS PERTUSARIA IN NORTH AMERICA NORTH OF MEXICO. par Martyn J. DIBBEN, Milwaukee Public Museum, Publications in Biology and Geology N). 5, iv + 162 p., 136 fig., 4°, broché, 1980. Milwaukee Public Museum, Milwaukee, WI. 53233, USA. Prix \$ 22.50 + postage.

L'examen de pas moins de 6800 specimens, incluant entre autres des matériaux historiques de quelques 40 herbiers reconnus, a été nécessaire à cette revue des 66 espèces North Américaines des lichens du genre *Pertusaria*. 14 espèces sont nouvelles. 33 espèces sont endémiques à l'Amérique du Nord. Du point de vue de leur distribution écologique, 60 % s'nt tempérées, 35 % arctoboréales et 5 % tropicales. 40 acides organiques sont recensés chez les *Pertusaria*. L'auteur a appliqué l'analyse chromatographie sur silicagel à tous les échantillons, selon la méthode de Culberson. Sa taxonomie des espèces est avant tout basée sur la morphologie du thalle et de la fructification apothéciale. Sur cette base elle distingue deux sous-genres, *P. sbg. Pertusaria* et *P. sbg. Pionospora*, l'un à fructification amplicariale ou pertusariale, l'autre à fructification lécanorale ou sorédiale. L'analyse chimique s'avère une confirmation précieuse dans la distinction des espèces. Confrontées entre elles, les deux méthodes permettent de réduire le poids donné aux variations et de ne plus subdiviser l'espèce en une série de variétés ou de formes. La différenciation chimique jointe à la ségrégation géographique a permis cependant de confirmer des espèces proches comme distinctes. L'espèce type *Pertusaria pertusa*, eurasiatique, n'est pas reconnue en Amérique du Nord, celle qui lui a été confondue est décrite comme *P. consocians*. L'auteur a le souci d'une nomenclature correcte. Elle indique le nom du genre comme *nomen conservandum*, notant qu'il est illégitime (Art. 63), et réfère sans plus de détail - à regret - à un article futur. Ses synonymies précisent le statut nomenclatural et sont justifiées par des commentaires. Les descriptions des espèces d'une liste détaillée des spécimens examinés et d'une distribution géographique illustrée. La

technique macrophotographique appliquée laisse à désirer. Cette monographie à la fois nouvelle, précise et didactique mérite l'attention.

A PRELIMINARY POLYPORE FLORA OF EAST AFRICA, par Leif RYVARDEN & Inger JOHANSEN, 64° p., 212 fig., 8°, couverture papier, 1980, Fungiflora Ed. P.O.Box 95, Blindern, Oslo 3, Norway. Prix Nkr 200

Durant les 10 ans passés, alors qu'il publiait à intervalle régulier les volumes des flores des Polyporaceae et des Corticiaceae d'Europe du Nord (revus dans Myco*axon), le premier auteur a élaboré cette flore des Polypores d'Afrique orientale par diverses explorations de récolte et l'étude des grands herbiers européens. La flore couvre 5 familles, les Corticiaceae, Ganodermataceae, Hymenochaetaceae, Polyporaceae et Tremellaceae, 75 genres dont 4 nouveaux (*Antrodiella*, *Echinoporia*, *Navi-sporus* et *Pseudopiptoporus*) et 337 espèces dont 3 nouvelles. Des clés réellement dichotomiques des genres et espèces sont proposées et, évidemment (et pour quoi?) limitées aux seuls genres et espèces traitées. Les espèces sont décrites avec une attention particulière aux détails anatomiques et microscopiques, lesquels sont dessinés dans la plupart des espèces. L'habitat et la distribution géographique sont indiquées, mais sans liste des spécimens étudiés (le livre se veut être une flore et non une monographie). Imprimé en caractères dactylographiques, le texte est sans relief, peu attirant. Les nombreux espaces blancs donnent au livre une épaisseur de 35 mm qui fera rapidement céder la faible couverture de papier. Le livre est dédié à la mémoire de M.A. DONK.

SAPROPHYTIC MICROFUNGI FROM TAIWAN. PART 1 HYPHOMYCETES, by Takashi MATSUSHIMA, Matsushima Mycological Memoirs No 1, 82 p., 46 fig., 8°, paperback, May 1980. Publication Matsushima Fungus Collection, 23-19-601 Mikageyamate-2-Chome, Higashinada-ku, Kobe, Japan 658. Price: free of charge through official request.

This is the first of a series of papers enumerating the microfungi collected in Taiwan during 1976 to 1978, by the author when he was guest as visiting senior mycologist at the Plant Pathology Division, Plant Protection Center, Taiwan. As in previous publications, the authors reports his profuse collections and cultures of the Hyphomycetes with the same wonderful ability and efficiency. His descriptions or comments on species are just what is needed, but his full page drawings speak more than words. Not less than 250 Hyphomycetes including 21 new species and 2 new genera (*Acumispora* and *Cheiropolyschema*) are recorded, documented and 46 illustrated on the 82 pages of this issue. This is the start of a new mycological series entitled *Matsushima Mycological Memoirs*. We are sure of the success which this series will receive through the mycological world. This series is distributed by the author's newly established institution, a very well standing fungus herbarium and culture collection, the *Matsushima Fungus Collection*.

BIBLIOGRAPHIA BOTANICA CECOSLOVACA 1973-1974, par A. Neuhäuslová-Novotná et D. Guthová-Jarkovská, 564 p., broché, 14x20 cm, Botanický Ústav ČSAV, Průhonice u Prahy, 1978.

Ce répertoire bibliographique donne toute la littérature botanique publiée en Tchécoslovaquie durant la période citée, classée par matière et par auteur.

UNTERSUCHUNGEN ZUR KONSERVIERUNG DE FRUCHTKÖRPER DES SPEISEPILZES *PLEUROTUS OSTREATUS* UND DER PARTIELLEN AUTOLYSE VON PILZSELLWÄNDEN, par Helga SCHMITZ, Bibliotheca Mycologica vol. 77, 98 p. 8°, dos papier, 1980, J. Cramer ed.

Dans l'étude du mécanisme enzymatique de l'attendrissement observé lors de la conservation de carpophores dans des solutions acidulées acétique, citrique et lactique (pH 4) l'auteur met en évidence l'action de R-glucanase et chitinase spécifiques du pied et du chapeau du fruit, tandis que les S-glucanase et chitobiase du mycelium sont sans action. Thèse réalisée sous la direction du Pr. G. Eger-Hummel, Marburg.

MUSHROOMS OF IDAHO AND THE PACIFIC NORTHWEST. DISCOMYCETES, par Edmund E. TYLUTKI, 133 p., 27 fig., 80 phot., 13x21cm, 1979, University Press of Idaho, University Station Box 3367, Moscow, Idaho 83843, USA.

Ce livre est écrit pour les amateurs "enthousiastes" qui veulent dépasser les frontières des basidiomycètes et s'aventurer à récolter les morilles, helvelles, truffes et autres grandes Pezizales, Tubérales et Geoglossacées. Chaque espèce décrite est illustrée d'une photo blanc-noir. Une clé "de terrain, purement macroscopique" est proposée: mais elle est dichotomique et couvre 5 pages bien pleines... qu'il faut lire à chaque récolte? Une clé synoptique serait plus didactique et plus rapide. Cette remarque dépasse le cadre de ce livre qui est bien écrit et agréable. Pas moins de 122 espèces y sont classées et reconnaissables par les clés.

CBS COURSE OF MYCOLOGY, par W. GAMS, H.A. VEN DER AA, A.J. VANDER PLAATS-NITERINK, R.A. SAMSON, J.A. STALPERS, 2e édition, 110 p., 75 fig., in 8°, dos papier, CBS, Baarn, Nederland. 1980.

Le texte de ce cours (1e édition revue dans Mycotaxon 3:558, 1976) est revu et complété de la littérature récente, sous à peu près le même volume. Les auteurs y développent les traits généraux de la morphologie des champignons, dans chaque groupe taxonomique, étayé de nombreuses figures et références. Le fascicule ne se veut pas un guide d'identification (voir von Arx, Champignons sporulant en culture pure, 3e éd. en prép.) mais une bonne introduction au règne des champignons.

A REVISION OF *CHRYSOSPORIUM* AND ALLIED GENERA, par C.A.N VAN OORSCHOT, Studies in Mycology, n° 20, 89 p., 36 fig., 3 pl., in 8°, dos papier, 1980, CBS, Baarn, Nederland. Prix Hfl 25.-.

Cette monographie donne les descriptions illustrées et les clés de *Chrysosporium* (22 espèces), *Myceliophthora* (8 espèces), *Emmonsia* (2 var.) *Zymonema* (1 esp.) *Trichosporiella* (2 esp.) et *Geomyces* (3 var.). La conidiogénèse blastique et thalique de ces champignons est décrite, ainsi que les relations avec les stades téléomorphiques appartenant aux Gymnoascaceae, Onygenaceae, Ascosphaeraceae et Sordariaceae. Cette revision est la première depuis le travail de Charmichael de 1962.

EESTI SEENTE KOONDNIMESTIK (LIST OF ESTONIAN FUNGI, WITH HOST INDEX AND BIBLIOGRAPHY) par J. JÄRVA et E. PARMASTO, 331 p., Din, dos papier, 1980, Institute of Zoology and Botany, Academy of Sciences of the Estonian S. S. R, Tartu, Estonia, Prix: Rbl. 2,10.-

Cette liste des champignons de l'Estonie est le résultat de 15 ans de travail des mycologues de l'Académie de Tartu pour le dépouillement de 1200 publications citées en annexe. Chaque espèce est citée avec hôtes et références.

BIOLOGY IN THE BLUE RIDGE, Fifty Years of the Highlands Biological Station, 1927-1977, by Ralph M. SARGENT, 156 p., ill., 8°, paperback, 1977, Highlands Biological Station, P.O. Drawer 850, Highlands N.C. 28741.

The then named Highlands Museum and Biological Laboratory, initiated in 1927 by a few determined people eager to preserve local human and natural history, has been, from the early years, associated with mycology. The Highlands station, in 1934, received the Mycological Society of America for its summer foray. Dr. William Chambers Coker, who presided the institution from 1933 to 1943, and subsequently as Honorary President, has been, with Dr L.R. Hesler, Vice-President, the promoters of the study of the Appalachian fungi. Coker published on basidiomycetes, particularly *Hydnium* and the Boletaceae (1939-1951). Hesler continued his *Notes on Southern Appalachian Fungi* in *Mycologia* from 1936 to 1955, and collaborated with Alexander H. Smith on *Hygrophorus* and *Lactarius* (1932-1962). Dr. L.S. Olive (1943-1953) published on Tremellales and in 1965 his book *Poisonous, Edible and Hallucinogenic Mushrooms*. Dr. Ronald H. Petersen, more recently investigated the clavarioid fungi and the aquatic hyphomycetes (1962-1963). Surely, the Blue Ridge Laboratory contributed much in 50 years to the American mycology. The Sargent's book, retracing, year after year, the multiple ways and reviving the unique spirit of the Highlands Biological Station will please the present and the future scientific people.

PHYTOPHTHORA CINNAMOMI AND THE DISEASE IT CAUSES, by G. A. ZENTMYER, Monograph No 10, 96 p., 39 figs., 4°, paperback, 1980, The American Phytopathological Society, 3340 Pilot Knob Road, St Paul MN 55121. Price US \$ 8.-.

How is it possible to cover such widespread and largely investigated fungus pathogen like *Phytophthora cinnamomi* in 95 pages. The author did, and on the base of 600 literature references. The pathogen is of importance for the decay of many tropical and temperate crops and trees. 900 hosts have been listed. The monograph details the many aspects of the biology, ecology, physiology and genetics of the pathogen, all data necessary in the search to an effective control. The text is dense and illustrated of many original documents. George A. Zentmyer of the University of California, Department of Plant Pathology is devoting himself on the study of that pathogen for already 20 years and published with his collaborators more than 100 papers on the subject. His mastery in that field will help many people engaged in research through the world.

A MANUAL OF ASSESSMENT KEYS FOR PLANT DISEASE, by Clive JAMES, 44 fiches illustrées, 10x18 cm., 1980, The American Phytopathological Society, St Paul. Prix US \$ 10.0.

Ces fiches illustrent les clés d'évaluation des dégâts par *Rhynchosporium*, *Puccinia*, *Septoria*, *Drechslera* sur céréales, *Phoma*, *Leptotrochila*, *Stemphylium* sur luzerne et trèfle, *Phytophthora*, *Streptomyces*, *Rhizocto-*

nia sur pomme de terre et *Xanthomonas* sur haricot. Elles aideront les phytopathologistes à effectuer des évaluations comparables.

LABORATORY HANDBOOK OF MEDICAL MYCOLOGY by Michael R. McGINNIS, 662 p., ill., 8°, hard cover, 1980. Academic Press, N.Y.

"Owing to the increased importance of fungi in medicine, there is a pressing need to discuss important topics such as laboratory safety and emergency procedures, quality control and modern concepts,... to assist microbiologists in safely isolating and accurately identifying fungi of medical importance.... The key to understanding any field of science necessitates a thorough understanding of its language..." (the author). Thus, the author devotes its first chapter to the morphological terms to be used in modern medical mycology for the description and the classification of the fungi, while chapter 4 (233 p.) and chapter 5 (63 p.) describe the systems of classification, detail the methods of study and comment the diagnoses of genera of, respectively the Ascomycetes, Coelomycetes, Hyphomycetes and Zygomycetes (sic) in the former and the yeasts in the later. Chapter 9 (50 p.) adds a synopsis of the mycoses, describing and illustrating symptoms, etiology and therapeutics of each disease. The other chapters of the book are descriptive of techniques: laboratory safety, handling clinical specimens, bioassay procedures, susceptibility testing, quality control, equipment maintenance, culture collection and culture media. Appendices provides with glossary, list of synonyms and list of fungal records on man.

The book is no doubt interesting and useful, not only for the techniques described, but as an effort of providing medical mycologists with the developments in modern descriptive mycology, especially in the field of the Hyphomycetes, after the First Kananaskis Conference of 1969.

Non obstant such evident value, the book presents several major defects that myself at least, I do regret. In the general disposition of the contents, the book might have been divided advantageously into two parts, one part, taxonomic, including chapters 1, 4 and 5 (the techniques excluded), chapter 9 and the appendices, the other part, technical, grouping chapters 2 and 3 with chapters 6 to 10, and the culture methods from chapters 4 and 5. That grouping might have avoided regrettable duplications between chapter 1 and chapter 4 in text and illustration.

The illustration is, beside a few line drawings, essentially made of black and white microphotographs of fungi, 11 x 8.5 cm (half page) in size, many of them being of good quality. Unfortunately, the illustration is defective in several ways. The magnification is not given. The numbering of the photographs, composed of the chapter number and the figure number within the chapter is complex and inconvenient to the reader. Furthermore from a count of 219 numbered figures, 52 are composed of 2 or 3 separate full-size photographs, not mounted in plate, but printed on distinct pages most often and provided with a full legend under fig. -A which is said "continued" under fig. -B and fig. -C. This constitutes another inconvenience. The procedure is in most case unjustified. Either the photographs composing a figure show the same fungus at low and high magnification, the one at low magnification being generally of no use or no value; or the photographs represent different fungi and could have been numbered and legended separately. At the same, the duplication of the same or similar pictures of the same fungus, illustrating here a term and further a generic concept might have been avoided. For instance, a picture of *Drechslera* sp. fig. 1.1 is the inverted photograph fig. 4.46 of the same. Why *Trichothecium roseum* needs to be shown at different focusing in figures 4.105A and 4.105B on p. 303 and 304? Why *Syncephalastrum ramosum* must

appear in 5 figures (1.9A, 1.9B, 4.120A, 4.120B, 4.121) and on distinct pages? Unacceptable is the fact that *Exophiala spinifera* is not only figured three times in figs. 1.27, 4.13 and 4.50 but also duplicated in fig. 4.49A, identical to fig. 4.13, under the name *Exophiala jeanselmei*. It will be also mentioned that amongst the 161 figs. illustrating chapters 1 and 4, 50 figs. are of unidentified species of filamentous fungi. Why, in such a handbook designed to be informative as much as exemplative, the author choose to illustrate unidentified isolates rather than properly identified fungal species of medical importance which are available in pure culture in official culture collections? Why finally the photographs are supplemented of such poor and inaccurate line drawings?

Other defects are in the text. The author decided not to cite the authors of the names of the fungi in the text, but did cite them, in Appendix C, only for the correct names of which he listed a synonym but not for many other names used in the text. The reader will have to consult other sources to distinguish for instance *Blastomyces dermatitidis* (conidial *Ajellomyces dermatitidis*) from *Wangiella* (*Hormiscium*, *Phialophora*, *Exophiala*) *dermatitidis*.

At the exception of the references inserted in the comments following the generic descriptions of the fungi, references to the literature are totally absent from the general text, like are the references to published sources for the 77 culture media cited. The reader need to be informed on the source of every statement, term, definition, classification, key, technique or formula other than the author's personal ones.

Dealing with terminology in chapter 1, the author restrict the application of the term *chlamydospore* to those terminal inflated thick-walled cells which are able to germinate and be reproductive and proposes the use of the term *vesicle* for the not germinating terminal chlamydospores as well for the intercallary ones. This proposition should be disrecommended, the term *vesicle* being currently designating the ampulliform tip of the *Aspergillus* conidiophore stipe. Further the author illustrates the term *favic chandelier*, in use in medical mycology, by a photograph of the appressorial hyphae in *Botrytis cinerea*. As soon as 1870, J. KLEIN introduced the term *appressorium* to designate that type of hypha differentiated for the attachment of the fungus. There is no reason for abandoning that term.

A final remark is needed to inform the author's statement "flagellate cells are produced only by the chytrids" on page 1 to 2.

Published by Academic Press, this handbook takes benefit of a high quality typographic printing and binding, as it will surely be in the hands of many mycologists in medical centers and laboratories.

FUNGAL BIOTECHNOLOGY, by J.E. SMITH, D.R. BERRY & B. KRISTIANSEN, x + 308 p., ill., 8°, hard cover, 1980. The British Mycological Society, Symposium series n° 3. Academic Press, 111 Fifth Avenue, New York, NY10003. Price \$ 32.50.

This book is the proceedings of a joint symposium of the British Mycological Society and the Society of Chemical Industry, held in Glasgow in September 1978. The recent developments of the fungal biotechnology has incited more research on the uses of fungi in the industrial processes. The book is a selection of topics related to the inoculation the growth and the exploitation of the fungi through the operation of liquid and solid state fermenters. The parameters of fungal growth, from the preparation of the inoculum, the genetical hazards, the rheological conditions to the reactions to the type of fermenter and the nature of the developed particles, are largely investigated and discussed for their

interference in the design of the fermenters. The production of edible fungi is also considered. The potentialities of fungal biotechnology are reviewed and the achievements in different regions of the world also described. Although the repeated orthographic errors in the names of the fungi, this book remains a very interesting account to applied mycology.

FUNGAL SAPROPHYTISM by Harry J. HUDSON, *Studies in Biology* n° 32, 2d ed., 76 p., ill., paper back, 1980. Edward Arnold Publ., 41 Bedford Square, London WC1B 3DQ. Price £ 2.25.

It is good to realize the author's ability to reach so many goals through these modest 70 pages: enhancing curiosity and interest for the fungi, considering many aspects of the fungal ecology, guiding the student on the field and in the laboratory and demonstrating the practical interest of the fungi in industry and human life. Some titles: wood decay, blue stain, basidiomycetes and microfungi in litter, chitinolytic fungi, lignine decomposition, pyrophilous fungi, coprophilous fungi, moisture requirements, osmophilous fungi, storage fungi, xerophytes, aquatic fungi, water pollution, thermophilous fungi, composts, fermented foods, antibiotics, vitamins. An exciting introduction to mycology, through ecology.

FUNGI, MAN AND HIS ENVIRONMENT, by R.C. COOKE, 144 p., ill., paper back, 1977. Longman Group Ltd, London. Price £ 3.5.

"I have attempted to write a book about fungi rather than a text-book on fungi". The author indeed realizes a similar approach to that of Houston, but with many more information and considering also the parasitic fungi, appealing the awareness of people on the danger and value of the fungi in the present and future world. From the pests and mycoses to their control, from the degradation of organic matter to the synthesis of protein and the cultivation of mushrooms for foods and drugs, the author clearly demonstrates the multiple, both destructive and elaborative potentialities of the fungi, in which, finally, man is concerned. With science and philosophy, he suscites the desire of reading more on fungi.

DIE BLÄTTERPILZE (AGARICACEAE) DEUTSCHLANDS UND DER ANGRENZENDEN LÄNDER BESONDERS OESTERREICHS UND DER SCHWEIZ, par Adalbert RICKEN xxiv + 482 p., 122 pl. col., in 8°, relié, Leipzig 1915. Réimprimé par Dr. Massimo CANDUSSO, Viale Europa 5, 21047 Saronno (Varese) Italia. Disponible dans l'allemand original (Lit. 98,000.- net) ou traduit en italien par Dr. Lazzari (Lit. 82,000.- net).

La réimpression de l'oeuvre mycologique originale et unique de Adalbert RICKEN (1851-1921) est un événement heureux. Publiée et distribuée par livraisons successives de 32 pages et 8 planches colorées de 1910 à 1915, l'oeuvre de RICKEN fut très bien accueillie par le monde mycologique de l'époque d'après guerre. En effet l'oeuvre est le bilan de quarante année de récoltes mycologiques (1870-1910) en Allemagne et régions limitrophes et portant sur plus de 1500 espèces. L'ouvrage présente dans un ordre systématique la description de 1482 espèces d'acariens sur la base d'observations précises tant microscopiques que macroscopiques, avec mention de l'habitat et, pour 662 espèces, une illustration

en couleurs reproduisant les peintures à l'huile que réalisa l'auteur sur le frais. La nomenclature est traditionnelle mais étayée des dates de publication. La dernière livraison comporta outre l'introduction et les index alphabétiques trois clés dichotomiques des genres, fort bien construites et basées sur les caractères les plus évidents.

La réimpression entreprise par l'Imprimeur mycologue, Dr. Massimo CANDUSSO est particulièrement réussie. Le texte est reproduit de manière régulière sur du papier de qualité. Les planches en couleurs, éditées sur un carton plus fort comme les originales, reproduisent les tons les plus stables des différents exemplaires originaux comparés. La reliure de type ancien est préservée dans une gaine. La qualité de cette réimpression mérite les encouragements du plus grands nombre de mycologues.

ICONOGRAPHIA MYCOLOGICA par Giacomo BRESADOLA, avec J.B. Traverso, L. Fenaroli, G. Catoni & J.B. Trener editeurs, 26 vols. 156 p., 1250 pl. col., Milan 1927-1933. Réimpression par Dr. Massimo CANDUSSO, Gruppo Micologico G. Bresadola, Viale Europa 5, 21047 Saronno (Varese) Italia, in 5 vols., 1250 pl. col., relié, Sept. 1981-Déc. 1982. Prix souscription Lit. 100.000 before September 1981, Lit. 100.000 au reçu des vols. 1, 2 and 3, le solde Lit. 60-85.000 au reçu du 4e vol.

L'examen de la planche proposée par l'imprimeur CANDUSSO au souscripteur semble présenter toutes les garanties de qualité que l'on peut espérer, voire exiger, d'une telle réimpression. L'ouvrage de Bresadola est une oeuvre majeure dans la connaissance des champignons. La réimpression ne sera réalisée que si la souscription est suffisante. On ne peut que recommander aux mycologues et aux institutions de science mycologique l'acquisition d'un ouvrage que les générations futures de mycologues à travers le monde ne pourront ne pas connaître ni ne pas consulter.

Also received:

LABORATORY GUIDE FOR IDENTIFICATION OF PLANT PATHOGENIC BACTERIA par N.W. SCHAAD, 72 p. 4 pl. col., mimeographed, 4° in folder, 1980. The American Phytopathological Society, St Paul, USA. Prix US \$10.-

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