A NEW WOOD-DECAYING SEQUESTRATE FUNGAL GENUS RELATED TO Armillaria (Physalacriaceae, Agaricales, Basidiomycota) \(^1\)

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- **Premise of the study:** Sequestrate basidiomycete fungi (e.g. “gasteromycetes”) have foregone ballistospory and evolved alternative, often elaborate mechanisms of basidiospore dispersal with highly altered basidiomorphology. Sequestrate fungi have independently evolved in numerous Agaricomycete lineages, confounding taxonomic arrangements of these fungi for decades. Understanding the multiple origins and taxonomic affinities of sequestrate fungi provides insight into the evolutionary forces that can drastically alter basidiomorphology. In the neotropical rainforests of the Guiana Shield, we encountered a remarkable sequestrate fungus fruiting directly on decaying hardwood roots. The fungus’ singular combination of traits include a wood-decaying habit; black, verrucose peridium; reduced stipe; and gelatinized basidiospore mass.

- **Methods:** **Guyanagaster necrorhiza** gen. et sp. nov. is described. Macro- and micromorphological characters were assessed and compared to most similar taxa. To determine the phylogenetic affinities of the fungus, DNA sequence data were obtained for the 18S, ITS, and 28S rDNA, RBP2, and EF1\(\alpha\) regions and subjected to single- and multi-gene analyses. DNA sequences from fungal vegetative organs growing on decaying woody roots confirmed the wood-inhabiting lifestyle of **Guyanagaster**.

- **Key results:** **Guyanagaster** is morphologically unique among sequestrate fungi worldwide. Phylogenetic evidence places **Guyanagaster** in close relation to the wood-decaying mushroom genus *Armillaria* in the Physalacriaceae (Agaricales, Agaricomycetes, Basidiomycota).

- **Conclusions:** **Guyanagaster** represents an independently evolved sequestrate form within the Physalacriaceae. Although molecular data confirm that **Guyanagaster** is closely related to *Armillaria*, the unusual features of this fungus suggest a case of radically divergent morphological evolution.

**Key words:** Agaricomycetes; basidiomycetes; fungal taxonomy; gasteromycete syndrome; Guyana; neotropics; rDNA systematics; sequestrate fungi; statismosporic fungi.

Within the Basidiomycota, “gasteromycetes” comprise a diverse, artificial assemblage of puffballs, earthstars, false earthstars, earthballs, bird’s nest and cannonball fungi, stinkhorns, se-cotoid agarics and boletes, and false truffles (Miller and Miller, 1988). The basidiospores of these fungi mature within their basidiomata, not on exposed hymenial surfaces (i.e., they are “sequestrate”), and are ultimately dispersed via mechanisms other than direct forceable discharge from basidia into the airstream. Sequestrate basidiomorpha morphologies adapted for dispersal via rain splash, animal mycophagy, forceable discharge of peridioles, physical weathering, or a bellows mechanism represent some of most dramatic evolutionary experiments in the Fungi (Ingold, 1965; Kendrick, 1992). The sequestrate Basidiomycota were traditionally treated as a cohesive taxonomic unit (e.g., class Gasteromycetes) with the assumption that ballistospory was lost rarely in the evolutionary history of fungi and that the advent of hymenial enclosure was rare (e.g., Coker and Couch, 1928). Alternatively, it had been suggested that the sequestrate state was ancestral in hymenomycetes before the evolution of ballistospory (e.g., Singer, 1971). Other early authors regarded sequestrate fungi as a polyphyletic group based on morphological and developmental evidence (e.g., Rejnders, 1963; Heim, 1971).

Molecular systematics studies of Agaricomycetes have since revealed that sequestrate fungi have independently evolved many times within the class and that new sequestrate taxa are being discovered in family or genus-level lineages previously thought to be entirely nonsequestrate (Bruns et al., 1989; Mueller and Pine, 1994; Hibbett et al., 1997; Miller et al., 2000; Miller and Aime, 2001; Peintner et al., 2001; Binder et al., 2006). Some sequestrate fungi represent recent, isolated evolutionary events that led to one or a few sequestrate species within a clade of nonsequestrate relatives (e.g., Kretzer and Bruns, 1997; Martin et al., 1999; Smith et al., 2006), whereas other sequestrate clades of earlier origin have speciated and radiated across the globe (e.g., Grubisha et al., 2002; Binder and Hibbett, 2006; Hosaka et al., 2006). Additionally, understanding of the multiple origins and taxonomic affinities of sequestrate fungi provides insight into the evolutionary forces that can...
drastically alter basidiof orm and function. (Thiers, 1984; Kretzer and Bruns, 1997; Reijnjders, 2000; Trappe and Claridge, 2005; Albee-Scott, 2007)

Traditional classifications of sequestrate basidiomycete fungi relied on overall basidioform as well as morphology of the peridium, basidiospores, and basidia (Miller and Miller, 1988; Trappe et al., 2009). Natural classifications aligning sequestrates with their nonsequestrate relatives have been problematic because many sequestrate taxa have reduced or unusual features that obscure phylogenetic inference. Sequestrate fungi often lack a stipe, their basidia may be atypically shaped and/or disappear at maturity, peridial structures may be poorly differentiated, and their basidiospores are usually highly ornamented in comparison to those of their closest nonsequestrate relatives (Trappe et al., 2009). In addition, many sequestrate fungi fruit below ground, they are infrequently collected and relatively understudied (Montecchi and Sarasini, 2001; Trappe et al., 2009).

Tropical sequestrate fungi are poorly known. For example, Mueller et al. (2007) estimated that ca. 30 species of hypogeous sequestrate taxa are currently described from the neotropics with ca. 200 species remaining unknown to science. Recent studies in the Guiana Shield region of northeastern South America have revealed a diverse assemblage of sequestrate fungi in forests dominated by ectomycorrhizal Dicymbe spp. (Fabaceae subfam. Caesalpinioideae). They are from divergent basidiomycete orders, including Agaricales, Boletales, Geastrales, Hysterangiales, and Phallales, as well as the ascomycetous Eurotiales (T. W. Henkel, M. C. Aime, M. E. Smith, S. L. Miller, University of Wisconsin; M. A. Castellano, Oregon State University; unpublished data; Miller et al., 2001; Aime and Henkel, 2008).

Here we describe Guyanagaster necrorhiza gen. et sp. nov. from the Pakaraima Mountains of Guyana, using morphological, molecular, and ecological data. This extraordinary new fungus is the only sequestrate taxon known with close affinities to the root parasitic, lamellate mushroom genus Armillaria (Fr.) Stude (Physalacriaceae, Agaricales, Agaricomycetes, Agaricomycotina, Basidiomycota), a genus of immense economic and ecological importance worldwide (Termsorhuizhen, 2000).

While these affinities are corroborated by the root-decaying habit of G. necrorhiza, its sequestrate form suggests a case of extreme divergence in fruiting morphology from Armillaria, the evolutionary pressures for which are currently unknown.

MATERIALS AND METHODS

Collecting and morphological analyses—Collecting expeditions were conducted during the rainy seasons of May–July of 2001–2003, 2008–2010 and December 2009 in the Upper Potaro River Basin in the west-central Pakaraima Mountains of Guyana. Fungi were collected within a 15-km radius of a previously established base camp (5°1’8.4”N, 59°54’4.0”W) in forests dominated by Dicymbe corymbosa Spruce ex Benth. and D. alstonii Sandw. (Henkel, 2003). Basidiomata of the black, truffle-like fungus were examined in the field for their fruiting habit and substratum relationships and described for their fresh morphological characteristics. Color was subjectively described and recorded according to Kornerup and Wanscher (1978) with corresponding color plates noted in parentheses (e.g., 3C4). Macroscopic spot tests followed those of Singer (1986). To elucidate the nutritional habit of the fungus, we collected 0.5-3 cm diameter small roots of D. corymbosa that were found in the vicinity of the basidiomata. We selected roots with attached hyphal cords, mycelial fans in the cambium, and white rot decay. Basidiomata, hyphal cords, and mycelial fans were field-dried with silica gel for molecular analyses (Miller et al., 2002). Additionally, mature basidiomata exhibiting arthropod excavation channels were collected and examined.

Micromorphological features of fresh basidiomata were examined in the field with an EPOI microscope and on dried specimens in the laboratory using an Olympus BX51 microscope with bright field and phase contrast optics. DNA samples were isolated by hand or by CTAB method (Gardes and Bruns, 1993) or with the Wizard Genomic DNA Purification Kit on field tissue samples stored in 600 μL of Nuclei Lysis Solution and extracted using the manufacturer’s protocols (Promega, Madison, Wisconsin, USA). PCR reactions were performed with published primers and according to standard protocols. For basidiomata, the ITS and the first ca. 1000 bp of the large subunit ribosomal (28S) DNA were amplified with several primer combinations. ITS: 28S-Gardes and Bruns/SIP/LR3 (Smith et al., 2007), LROR/LR5F (Tedersoo et al., 2008), and LSU4/BR/LR6 (Aime and Phillips-Mora, 2005). Approximately 1100 bp of the small subunit (18S) was amplified with primers SS3 and SS8 (White et al., 1990). The variable region between domains 6 and 7 of the RPB2 gene was amplified with primers RB2P-6F and RB2P-7R (http://faculty.washington.edu/mbhennell/; Matheny, 2005). DNA sequences of the Elongation Factor 1-alpha (EF1α) gene were amplified using primers EF1-αf and EF1-2218r (AFTOL website: http://www.aftol.org/primers.php; Rehner and Buckley, 2005). For an individual hyphal cord and mycelial fan, the ITS region was amplified to allow comparison with basidioma ITS sequences.

PCR products were visualized on 1.5% agarose gels stained with SYBR Green I (Molecular Probes, Eugene, Oregon, USA). Successful amplifications were cleaned with ExoSAP-IT (USB Corp., Cleveland, Ohio, USA). Bidirectional sequencing was performed with the described primers using the Big Dye Sequencing Kit v.3.1 (Applied Biosystems, Foster City, California, USA) on an ABI3730xl capillary sequencer (Applied Biosystems). Sequences were edited with the program Sequencer v.4.1 (Gene Codes, Ann Arbor, Michigan, USA).

Initial blastn (http://blast.ncbi.nlm.nih.gov/Blast.cgi) analyses of the 28S sequence data set indicated that our fungus had close affinities with Armillaria within the Physalacriaceae (Agaricales, Agaricomycetes, Basidiomycota), a genus of immense economic and ecological importance worldwide (Tormorshuizhen, 2000). While these affinities are corroborated by the root-decaying habit of G. necrorhiza, its sequestrate form suggests a case of extreme divergence in fruiting morphology from Armillaria, the evolutionary pressures for which are currently unknown.

Sequence alignments were conducted manually in the program Se-Al v2.0a11 (Andrew Rambaut, Department of Zoology, University of Oxford, UK; http://evolve.zoo.ox.ac.uk/). Ends were trimmed, and regions too variable to be reliably aligned were excluded from final analyses; adjustments were as follows: 28S: 93 of 95 total base pairs were included; ITS: 166 bp containing the 5.8S region were included; 18S: the taxon Xerula radicata contained a single 419 bp intron that was removed from the remaining 1734 bp; RPB2: 110 of 960 bp were removed; EF1α: 132 of 1130 bp were removed. Maximum parsimony (MP) analyses were conducted in PAUP* v4.0b10 (Swofford, 2003) as heuristic searches with 100 random addition replicates and TBR branch swapping. Support for the branching topologies was evaluated by bootstrap analysis derived from 1000 replicates with 10 random addition replicates each. Maximum likelihood (ML) analyses were conducted by the quartet puzzling method in PAUP* with 10000 puzzling steps; transition/transversion ratio = 2 (Strimmer and von Haeseler, 1996). For the multigene alignments, Lentinula lateritia, Gymnopus

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Fig. 1. Basidiomata of *Guyanagaster necrorhiza*. (A) Holotype; *Henkel 9065*. (B) Longitudinal section of basidioma showing immature gleba and well-developed peridium, columella, and stipe. (C) Mature gleba; note strong reddening of all internal tissues. Bar = 10 mm.
contrarius, and *Marasmius rotula* as additional members of the marasmioid clade were selected as outgroup taxa (Matheny et al., 2006); for single-gene data matrices, the non-Physalacriaceae taxa were selected as outgroups for rooting purposes. Bayesian analyses were conducted on the five-gene data set in the program Mr. Bayes v. 3.12 (http://mrbayes.csit.fsu.edu/index.php; Huelsenbeck and Ronquist, 2001) with the optimum models of DNA evolution determined using the program Mr. Modeltest2 v2.2 (http://www.abc.sel–nylander; Nylander, 2004). Posterior probability values were obtained by running one cold and three heated Markov chain Monte Carlo (MCMC) chains for 20 million generations, sampling trees every 1000 generations and discarding the initial 10% of trees as the burn in. The GTR+I+G model was selected for all three data sets, and in each case, the non-Physalacriaceae taxa were selected as outgroups for rooting purposes.

**TAXONOMY**

*Guyanagaster T. W. Henkel, Aime and M. E. Smith gen. nov. (Figs. 1–5)—* Basidiomata globosa usque ovata vel irregulater lobata, 22–52 × 11–34 mm, nigra, verrucosa; stipes concolor, laevis, 7–10 × 4–6 mm, ad radices lignosae affixus; gleba roseo-pallida usque lateritia, loculis gelatinosis; columella continua cum stipite, roseo-pallida usque rubro-aurantia; basidiosporae globosae, echinulatae, pedicellatae, avellinae, dextrinoidea, 15.5–18.5 × 15.6–18.8 µm.

*Guyanagaster necrorhiza T. W. Henkel, Aime and M. E. Smith sp. nov. (Figs. 1–5)—* Basidiomata gasteroida, globosa usque ovata vel irregulariter lobata, 22–52 × 11–35 mm, firma; peridium nigrum, verrucosum, verrucis obtusis, pyramidalibus, 0.3–0.5 mm crassis; endoperidium album, 0.8–1.5 mm crassum; stipes niger, laevis, 7–20 × 3–6 mm, ad radices lignosae affixus; gleba roseo-pallida usque carnea vel lateritia, loculis gelatinosis, cum albis hyphis separatis; columella continua cum stipite, roseo-pallida usque rubro-aurantia, evidenter distinctus gleba; basidiosporae globosae, echinulatae, pedicellatae, parietibus crassis, avellinae, dextrinoidea, 15.5–18.5 × 15.6–18.8 µm; basidia cylindrica, 2-sterigmata, genita hyphis loculorum immaturorum; fibulae carentes.

*Holotypus—* Henkel 9065 (BRG; isotype: HSU, FH, NY, DUKE)

Macromorphological features of basidiomata—Basidiomata gasteroid, subepigeous to erumpent, scattered or in linear troops, attached directly to woody roots of *Dicymbe* trees; 22–52 mm broad, 11–35 mm tall, globose to subglobose to ovoid and irregularly broadly lobate, dense, base with smooth, sterile concolorous stipe, 7–10 × 3–6 mm, attached directly to substratum (Figs. 1, 2). *Peridium* black during all stages of development, moist, tough, covered in blunt, 4–5 sided polygonal nondeciduous warts, these 0.75–1.5 mm broad, 0.3–0.5 mm tall, more acuminate with age, sharply differentiated from endoperidium; *endoperidium* white to light pink (7A1–7A2), tough, 0.8–1.5 mm thick, of matted hyphae transitioning evenly to sterile, white hyphal veins between the glebal locules; *gleba* composed of well-defined locules and intervening veins; locules globose to ovate to subangular, 0.5–4 mm broad, initially pinkish white (5A2–5A3), maturing in stages to flesh pink (6A3–6A4, to 6B0) and eventually a rich brick red (8B8–8C8–8D8) evenly throughout, unchanging; hyphae separating locules initially white, light pink (7A3) in older specimens, darkening slightly.
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at apex. With advanced age, entire basidioma permeated with invertebrate excavation channels; tissues between excavation channels remaining firm; without evidence of vertebrate consumption (Fig. 5). Odor mild, somewhat musty, clay-like in older specimens, taste mildly "chemical" or soapy, texture brittle when chewed. Macrochemical spot tests: KOH reddening endoperidium, columella, and gleba, darkening exoperidium; NH₄OH reaction lacking on all surfaces.

Vegetative features—Hyphal cords proliferating along exterior of long axis of individual Dicymbe roots from 0.5−3 cm diameter, white, 2−4 mm diameter, roundish to subflattened, tuberculate under lens, in cross section exterior rind undifferentiated, of loosely packed, white hyphae, with gelatinous, gray core 0.5−0.75 mm diameter, occasionally bridging adjacent roots; forming dense yellowish mycelial fans proliferating along root cambial region, from which feeder hyphae ingress via sapwood rays; wood decay reddish when incipient, advancing to a water-saturated, stringy white rot that eventually gelatinizes (Fig. 3).

Micromorphological features of basidiomata—Exoperidium composed of agglutinated masses of anticlinal hyphae forming conical to rounded warts, in mass 168−336 × 160−168 µm; individual hyphae of outer layer dark brown in H₂O, inamyloid, curved to tortuously curved, nongelatinous, 30−42 × 5−6 µm, swollen at septa, apically coalescing into protruding clumps, giving warts a hairy appearance under low magnification, hyphal walls rapidly staining deep red in KOH, with red pigment leaching into mounting medium; outer peridial hyphae subtended by a layer of subisodiametric cells, 14−22 × 12−22 µm, walls in H₂O yellow to light brown, 2−3 µm wide, intergrading below with hyphae of endoperidium (Fig. 4A). Endoperidium 700−760 µm thick, well-differentiated from exoperidium and glebal locules, composed centrally of highly interwoven, branching, anastomosing hyphae that are irregularly inflated, 2−5 µm wide, hyaline in water and KOH, inamyloid, thin-walled, moderately gelatinizing to yield an irregularly “honey-combed” appearance in mass; near extreme distal transition with exoperidium cells becoming more inflated and subanticlinal, toward the locules hyphae becoming thinner, more tightly interwoven. Sterile hyphae separating locules similar to endoperidial hyphae, parallel-interwoven, weakly gelatinous. Glebal locules in thin section globose to subglobose, 163−1020 × 255−1500 µm, tissues densely gelatinous such that locules are difficult to flatten on a microscope slide, preventing permeation of stains; locule tissue primarily of tightly packed basidiospores encased in a gel matrix, at maturity mostly lacking intact hyphae, when present hyphae 1−2 µm wide; well-organized hymenium lacking (Fig. 4B). Basidia apparently 2-sterigmate, difficult to observe even in microtome sections, more or less cylindrical and undifferentiated from subtending hyphae, occasionally inflated near apex and tapering evenly toward base, basal septum usually inevident, hyaline in water, thin-walled, originating as side branches from locular hyphae in immature locules; stigmatic difficult to observe, apparently long, arising at angles from sides of basidium apex. Basidiospores globose; in H₂O and KOH light tannish brown with faint pinkish overtones, in Melzer’s rich burgundy brown and therefore darkly dextrinoid, main body thick-walled, outer spore wall 1−2.4 µm wide, inner wall 0.75−1 µm wide, with regularly globose lumen, covered in entirety with straight to slightly recurved, narrow, cylindrical, apically acuminate spines, which are tightly packed and equidistant throughout, 1.7−2.2 µm tall, apical recurvations sometimes

Fig. 3. Vegetative development of Guyanagaster necrohiza on Dicymbe corymbosa. (A) Excavated roots with well-developed external hyphal cords. (B) Decaying woody roots with mycelia fans in cambium. (C) Wet, stringy, white rot. Bar = 10 mm.

with exposure; columella well-defined, base continuous with stipe, 9−22 × 7−15 mm, initially light pink (~7A3), orange-red (7B7–8B7) with age, dense, waterlogged, of longitudinally arranged hyphae under hand lens, sharply demarcated from gleba
unidirectional and giving the spore ornaments a “combed” appearance, spines apparently immersed in a gelatinous matrix; without spines (10.8–12.1−15.8 × 12−15.5(–16.1) μm (mean = 13.5 ± 0.3 × 13.5 ± 0.3 μm, Qr = 0.93–1.06, Qm = 1.0 ± 0.01); with spines included 15.5−18.5 × 15.6−18.8(–19.4) μm (mean = 17.0 ± 0.2 × 16.8 ± 0.3 μm, Qr = 0.95–1.05, Qm = 1.0 ± 0.01), (N = 20 basidiospores); in side view with robust, symmetrically positioned pedicel, 2.9−4.7 × 2.1−2.5 μm; immature basidiospores hyaline to light pink, wall thinner and spines longer relative to overall diameter, this situation reversing with maturity apparently due to progressive outward thickening of wall and resulting increase in basidiospore diameter, without attendant increase in spine length (Fig. 4C). **Columella tissue** composed of moderately interwoven, tightly packed, nongelatinous hyphae; hyphae thin-walled, hyaline in water and KOH, of variable widths, (2.5–)5−10(–18) μm. **Clamp connections** absent in all tissues.

**Habit, habitat, and distribution**—Rare, scattered to abundant between leaf litter and mineral soil attached to decaying woody roots of *Dicymbe corymbosa* and *D. altsonii* in tropical rainforest, known only from the Upper Potaro Basin of Guyana.

**Etymology**—Nekros = death, corpse; rhiza = root (Greek), referring to the root wood decaying, putatively parasitic habit of the fungus.

**Specimens examined**—Guyana, Region 8 Potaro-Siparuni—Pakaraima Mountains, Upper Potaro River Basin, elevation 710–750 m; vicinity of base camp, 29 June 2001, Henkel 8418 (BRG; HSU); vicinity of base camp, 11 June 2002, Aime 1956 (BRG, LSUM); vicinity of base camp, 4 July 2003, Aime 2312 (BRG, LSUM); 2.5 km southeast of base camp, *Dicymbe* plot 1, 27 July 2008, Henkel 8989 (BRG; HSU, FH); 2 km southwest of base camp, on roots of fallen *Dicymbe corymbosa* tree, 22 July 2009, Henkel 9065 (Holotype: BRG; isotype: HSU, FH, NY, DUKE), Henkel 9065R hyphal cord (HSU), Henkel 9065MF mycelial fan (HSU); 8 km southeast of base camp at Tadang Creek, on roots of standing dead *Dicymbe altsonii* tree, 24 December 2009, Henkel 9135 (BRG; HSU); 2 km southwest of base camp, on roots of fallen *Dicymbe corymbosa* tree, 17 May 2010, Aime 3950 (BRG; LSUM).

**MycoBank numbers**—MB 515527 (genus); MB 515528 (species).

**Commentary**—*Guyanagaster necrorhiza* is easily recognized in the field by its black, verrucose, subglobose to broadly lobate, firm basidiomata, which emerge slightly from the leaf litter in the vicinity of standing dead trees or recent treefalls. A longitudinal section reveals a firm, pinkish-flesh to brick red, loculate gleba, and distinct columella. Excavations of basidiomata reveal a black, smooth stipe directly attached to decaying woody roots, with associated white, exterior hyphal cords, mycelia fans in the root cambium, and wet, stringy,
Mature basidioma in the field showing arthropod excavation channels through the peridium, with associated frass (*altsonii* vaccinated by arthropods. The five-gene ML analysis (Fig. 6) and the *Armillaria* the closest related genus within the family is the lamellate anagaster maintain their firmness and moisture content, but are heavily excavated by arthropods.

Phylogenetic analyses fully support the placement of *Guyanagaster* within the Physalacriaceae in the Agaricales and that the closest related genus within the family is the lamellate *Armillaria* (Fig. 6). However, the exact relationship between *Armillaria* and *Guyanagaster* could not be definitively resolved. The five-gene analysis (Fig. 6) and the EF1α ML and MP analyses (Appendix S1, see Supplemental Data with the online version of this article) suggest that *Guyanagaster* represents a sister taxon to *Armillaria*, whereas the five-gene Bayesian and parsimony analyses (not shown) as well the 28S rDNA ML and PP analyses (online Appendix S2) suggest that *Guyanagaster* may be derived from within *Armillaria*, rendering *Armillaria* paraphyletic. With due consideration to these molecular results, we have chosen to erect *Guyanagaster* as a new genus for the following reasons: (1) the highly divergent basidioma morphology of *Guyanagaster* relative to all known species of *Armillaria* worldwide, including the sequestrate subhypogeous fruiting habit, strongly reduced stipe, gelatinized locules, bright-red glebal and black peridial colors, spinose basidiospores, inconspicuous basidia, and disorganized fertile tissues; (2) the radi-

cally different basidiospore dispersal in *Guyanagaster* relative to *Armillaria*; and (3) the differences in vegetative organs (e.g., the invasive hyphal cords of *Guyanagaster* are loosely organized, nonsclerotized, and white, whereas the rhizomorphs of all known *Armillaria* spp. are highly organized with a dense, melanized rind; Pegler, 2000). For basidiomata, hyphal cords, and mycelial fans, the ITS rDNA sequences were identical, confirming the root decay habit of *Guyanagaster*.

**DISCUSSION**

*Guyanagaster* is unique among sequestrate genera throughout the Agaricomycetes due to its combination of a black, verrucose peridium that leaches reddish pigments in KOH, short stipe and radiating columella, pink to brick-red gel-filled glebal locules, inconspicuous basidia, pinkish, spinose, strongly dextrinoid basidiospores with prominent pedicels, habit of fruiting directly from woody roots, and its phylogenetic affinities with *Armillaria*.

Morphologically similar sequestrate fungi—Among sequestrate genera of the Agaricales, *Guyanagaster* is similar to *Cribbea* A. H. Sm. and D. A. Reid (Physalacriaceae) in its basidiospore characteristics and wood decay habit. However, *Cribbea* species have basidia arranged in an obvious hymenium with robust cystidia and a hymeniform pileipellis (Lebel and Cratcheside, 2009). Our phylogenetic analyses of 28S sequence data show *Cribbea* as derived from within the *Xerula/Oudemansiella* lineage, and thus only distantly related to *Guyanagaster* (online Appendix S2; Lebel and Cratcheside, 2009). Species of *Hydnangium* Wallr. (*Hydnangiaceae*) have pediculate basidiospores with similar spore ornaments to those of *Guyanagaster*, but differ in having hyaline basidiospores, robust basidia, and smooth, usually pink or orange peridia; additionally, *Hydnangium* sensu stricto has originated within the lamellate genus *Laetaria* (Pegler and Young, 1979; Kropp and Mueller, 1999), which is now known to belong to the agaricoid lineage of Agaricales, rather than the marasmioid lineage to which Physalacriaceae belongs (Matheny et al., 2006). Species of *Stephanospora* (Berk.) Pat. (Stephanosporaceae) have spinulose basidiospores and putatively saprotrophic habits, but differ from *Guyanagaster* in their orange or yellow basidiomata that lack a stipe or columella and have nondextrinoid basidiospores with peri-appendicular coronas. In addition, *Stephanospora* is phylogenetically distant to *Guyanagaster* (Martin et al., 2004; Tedersoo et al., 2010).

Among sequestrate genera of the Boletales, *Guyanagaster* could potentially be confused with *Durianella* Desjardin, A. W. Wilson and Binder. The monotypic *Durianella echinulata* (Corner and Hawker) Desjardin, A. W. Wilson and Binder from tropical Asia has a verrucose peridium, gel-filled locules, spinose basidiospores, and a subhypogeous fruiting habit (Desjardin et al., 2008). However, *D. echinulata* has tissues that oxidize blue, a golden-yellow peridium that does not leach red pigment in KOH, inamyloid basidiospores lacking a pronounced pedicel, an ectomycorrhizal habit, and is a well-supported member of the Boletineae clade (Desjardin et al., 2008). Some species of the polyphyletic *Octaviania* Vittad. (Boletales) superficially resemble *Guyanagaster* in peridial and glebal structure and basidio-

spore ornamentation but lack a glebal columella and exhibit an oxidation reaction when cut or bruised (Montecchi and Sarasini, 2001; Trappe et al., 2009; Orihara et al., 2010).
Some *Sclerogaster* species (Geastrales) share features with *Guyanagaster*, including a pinkish or orangish gleba and ornamented basidiospores (Hosaka and Castellano, 2008). However, most *Sclerogaster* species have smooth, light-colored peridia, astipitate, acolumellate basidiomata, and basidiospores that are <10 µm diameter. The neotropical *Sclerogaster luteocarneus* (Bres.) Zeller and C. W. Dodge is similar to *Guyanagaster* in having spinose, pedicellate basidiospores and a short columella, but differs in its smooth, yellowish to brown peridium, yellowish gleba, and distinct, clavate basidia (Zeller and Dodge, 1935).

**Fig. 6.** Phylogram generated from maximum likelihood analysis of five gene regions (18S, ITS, and 28S rDNA, *RBP2*, and *EF1α*) from 24 taxa of Agaricales, showing a sister relationship of *Guyanagaster necrorhiza* with the genus *Armillaria* in the Physalaciaceae. *Lentinula lateritia*, *Gymnopus contrarius*, and *Marasmius rotula* were used as outgroup taxa. The phylogram was obtained from 10,000 quartet puzzling steps, with a transition/transversion ratio = 2. Support values shown above branches represent MP bootstrapping/ML quartet puzzling/Bayesian posterior probabilities. See Methods for analytical details.
mycelia fans, and the associated stringy, white rot suggest a similar wood decay habit, and possibly root parasitism, between Guyanagaster and Armillaria (Termorshuizen, 2000). While Guyanagaster may be able to colonize live roots of host trees by belowground proliferation of hyphal cords in a manner similar to many Armillaria species, more information is needed to ascertain its parasitic capacity. From the standpoint of basidioma morphology, there is little congruence macro- or microscopically between the sequestrate Guyanagaster and lamellate, agaricoid Armillaria, although some degree of basidiospore ornamentation has been noted in Armillaria spp. (Bennell et al., 1985). At the least, Guyanagaster is an entirely novel sequestrate taxon close to Armillaria within the largely nonsequestrate Physalacriaceae and as such represents a highly significant discovery.

Although the exact basidiospore dispersal mechanism of G. necrorhiza is currently unknown, field observations suggest that arthropods may be involved. First, many mature basidiomata were excavating. Third, basidiomata of G. necrorhiza were examined at specific collecting sites on different host tree species (TH 9065 and TH 9135; Fig. 5). Second, G. necrorhiza basidiomata appeared resistant to decay over a period of weeks and basidiospores remained immersed in a firm, gelatinized gleba except in portions where arthropods were excavating. Third, basidiomata of G. necrorhiza lack a powdery spore mass or strong odors, suggesting that neither wind or vertebrate mycophagy are likely dispersal vectors. Given the root decomposing, and possibly parasitic, habit of G. necrorhiza it is conceivable that basidiospores are successfully vectored into the belowground environment by arthropods. Lilkeslov and Bruns (2005) demonstrated that arthropods, including flies and beetles, were effective dispersers of basidiospores of the nonsequestrate, subterranean-fruiting ectomycorrhizal fungus Tomentella subbilacina. If similar mechanisms occur with G. necrorhiza, this would be the first recorded case combining such specialized dispersal and nutritional strategies in a fully sequestrate fungal species (J. M. Trappe, Oregon State University, personal communication).

LITERATURE CITED

KENDRICK, B. 1992. The fifth kingdom. Focus, Newburyport, Massachusetts, USA.


Appendix

**Taxon:** GenBank accession: 28S, 18S, ITS, RPB2, EF1α; Voucher specimen; Collection locale; Herbarium.

**Armillaria affinis** (Singer) T.J. Volk & Burds.; AF261356, —, —, —.
**Armillaria fumosa** Kile & Watling; DQ338552, —, —, —.
**Armillaria gemina** Bérubé & Dessur.; —, —, —, —.
**Armillaria heimi** Pegler; FJ618737, —, —, —.
**Armillaria hinnutae** Kile & Watling; DQ338555, —, —, —.
**Armillaria limonea** (G. Stev.) Pegler; FJ618733, —, —, —.
**Armillaria montagnei** (Singer) Herink; FJ711623, —, —, —.
**Armillaria nabsnona** T.J. Volk & Burds.; DQ338542, —, —, —.
**Armillaria novae-zealandiae** (G. Stev.) Herink; FJ618736, —, —, —.
**Armillaria padiellula** Kile & Watling; FJ618732, —, —, —.
**Armillaria psammophila** Kile & Watling; FJ618731, —, —, —.
**Armillaria tabescens** (Scop.) Emel; AF042593, DQ851372, AY123590, —, —.
**Armillaria villosa** (Romagn.) Herink; FJ618722, —, —, —.

**Armillaria sparrei** (Singer) Herink; FJ618750, —, —, —.
**Armillaria tabescens** (Scop.) Emel; AF042593, DQ851372, AY123590, —, —.

**Cyphellopsis anomala** (Pers.) Chamuris; DQ234541, AF518576, DQ205682, AY536283, —.
**Cyphellopsis digitalis** (Alb. & Schwein.) Fr.; AY635771, —, —, —.
**Cylindrobasidium torrendii** (Pers.) Chamuris; DQ234541, —, —, —.
**Cyphellopsis anomala** (Pers.) Donk; AF426955, AF426949, AY571034, —, —.
**Cyptotrama aspra** (Berk.) Redhead & Ginnis; AF261353, DQ440637, DQ973355, —.
**Cylindrobasidium torrendii** (Pers.) Chamuris; DQ234541, —, —, —.

**Cylindrobasidium torrendii** (Pers.) Chamuris; DQ234541, —, —, —.

**E. Horak & Ginns**; AF261353, —, —, —.
**Cyphella digitalis** (Alb. & Schwein.) Fr.; AY635771, —, —, —.
**Cyphella digitalis** (Alb. & Schwein.) Fr.; AY635771, —, —, —.

**Oudemansiella canarii** (Schrad.) Höhn.; AY207260, —, —, —.
**Oudemansiella mucida** (Schrad.) Höhn.; AY207260, —, —, —.
**Oudemansiella radicata** (Relhan) Singer; DQ717191, —, —, —.
**Oudemansiella radicata** (Relhan) Singer; DQ717191, —, —, —.
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**Oudemansiella radicata** (Relhan) Singer; DQ717191, —, —, —.
**Oudemansiella radicata** (Relhan) Singer; DQ717191, —, —, —.

**Porodisculus pendulus** (Schwägrichen{?}) Schwein.; AF261353, —, —, —.
**Schizophyllum commune** (Bull.) Maire; AF261353, —, —, —.

**Schizophyllum commune** (Bull.) Maire; AF261353, —, —, —.
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**Xerula furfuracea** (Bres.) Fr.; AY635771, —, —, —.
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