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A novel clade of sporocarp-forming species of glomeromycotan fungi in the *Diversisporales* lineage

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Abstract In the early times of taxonomy of arbuscular mycorrhizal fungi (Glomeromycota), exclusively sporocarpic species were described. Since then the focus has mainly shifted to species forming spores singly. For many of the sporocarpic species, no molecular data have been made available, and their phylogenetic position has remained unclear. We obtained small subunit ribosomal rDNA and internal transcribed spacer data from specimens of glomeromycotan sporocarps from tropical areas that were assigned to three morphospecies. The complete sequence of the 18S small rDNA subunit sequence, internal transcribed spacers (ITS) 1 and 2 and 5.8S rDNA subunit, was determined from a sporocarp of Glomus fulvum. Partial sequences of the small subunit and the other regions were obtained from Glomus pulvinatum and the newly described species Glomus megalocarpum. Molecular phylogenetic analyses placed all species analyzed as a monophyletic sister group to the Diversispora spurca/Glomus versiforme clade group ("Glomus group C") within the Diversis-

Taxonomic novelties: Glomus megalocarpum D. Redecker

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porales. The phylogenetic divergence from other known species suggests that this clade may constitute a new genus. These findings will have important consequences for taxon definition in the Diversisporales. They will facilitate identification of these fungi using rDNA sequences within colonized roots or the environment.

Introduction

Arbuscular mycorrhiza is a symbiosis formed by the vast majority of land plants. In this interaction, the plant partner provides carbohydrates to the fungal symbiont, which improves the uptake of mineral nutrients using its fine hyphal network. Approximately 200 morphospecies of arbuscular mycorrhizal fungi in ten genera have been described, mostly based on spore morphology. These fungi were previously placed in the family Endogonaceae (Gerdemann and Trappe 1974), later in the order Glomales of the Zygomycota (Morton and Benny 1990). Recently, a separate phylum Glomeromycota was established for them (Schüßler et al. 2001). As the spore morphology is rather simple in many species, molecular phylogenetic methods have helped to clarify the evolutionary relationships in the Glomeromycota (Redecker and Raab 2006). One result of these analyses was that the largest genus Glomus, with about 70 species, is polyphyletic (Redecker et al. 2000a; Schwarzott et al. 2001). Fungi falling within the previous morphological definition of Glomus have been shown to be in at least four different monophyletic lineages. In fact, some dimorphic species even form glomoid spores as well as other types of spores (Redecker et al. 2000a). Some of these lineages were separated in the new genera (Morton

and Redecker 2001), partly based on molecular characters. One remaining clade of *Glomus* comprises *Glomus* group A (e.g. *G. intraradices*, *G. mosseae*) and group B (e.g. *G. etunicatum*, *G. claroideum*), which are rather distantly related but still form a monophyletic group together (Schwarzott et al. 2001).

Another *Glomus* clade, referred to as *Glomus* group C, is more closely related to the Acaulosporaceae in rDNA phylogenies. It comprises the species *G. versiforme*, *G. spurcum*, and *G. aurantium*, as well as an isolate fitting the morphological definition of *G. etunicatum*, which is otherwise thought to belong to group B. *Glomus* group C was recently defined as a new family Diversisporaceae with *Diversispora spurca* as type species of the new genus (Walker and Schüßler 2004). Other species have not been formally renamed. Because of the scarcity of morphological characteristics on the light microscopic level that could aid separation from other *Glomus* species, signature sequences in the 18S ribosomal subunit were used for the definition of *Diversispora* as well as the corresponding family and order.

Numerous Glomus species form sporocarps, which mostly are relatively undifferentiated assemblages of spores. An exception are some of the species previously placed in the genus Sclerocystis. However, it was shown that Sclerocystis is an advanced clade within a group of well-defined nonsporocarpic Glomus group A species (Redecker et al. 2000b). Generally, the sporocarp-forming habit is not well correlated with phylogeny, and sporocarpic as well as nonsporocarpic representatives can be found in different phylogenetic groups. Very often, sporocarp formers are closely related to species forming spores singly in the soil, or a single species is known to form sporocarpic and nonsporocarpic spores (e.g. G. mosseae, G. coronatum). Early taxonomy of the Glomales focused very much on sporocarpic species, a trend that has not continued recently. The phylogenetic relationships of many of the sporocarpic species listed in the groundbreaking study of Gerdemann and Trappe (1974) have not been elucidated. Therefore, it is impossible to recognize these taxa if they are detected by molecular methods within colonized roots.

In this study, we sequenced regions of the ribosomal rDNA of some sporocarpic glomeromycotan species collected in tropical regions to elucidate their phylogenetic relationships and to allow the design of specific primers for molecular detection of these species.

Materials and methods

Specimen AC/Pohn99-001 was collected by Francisco J. Camacho in 1999 in a native forest at the ridge trail near the top of Sokehs Mountain on the pacific island of Pohnpei, Federated States of Micronesia (6.97°N 158.22°E). Other sporocarps were collected on the Caribbean islands of Guadeloupe and Martinique in August/September 2005 by Christophe Lécuru. Details were as follows: specimen CL/Mart05-35: Ste. Luce, Forêt départementalo-domaniale de Montravail, on forest floor; specimen CL/Mart05-049: Case pilote, Crête Jean Louis (Martinique), fruiting on dead wood; specimen CL/Guad05-51: Vieux-Fort (Guadeloupe), Ravine Blondeau, mesophile forest, on forest floor, close to *Boletellus* sp.; specimen CL/Mart05-111: Forêt départamentalo-domaniale de La Source Berry (Martinique), plantation of mahogany with indigenous species.

Slides to examine the spore morphology were obtained by scratching a few spores from the surface of crosssections and embedding them into Polyvinyl alcohol lactoglycerol (PVLG, Koske and Tessier 1983). Pictures of the whole sporocarps or parts of them were taken using a Hewlett Packard ScanJet 5470 flatbed scanner at maximum resolution (2,400 dpi). Microscopical pictures of the spores were taken with an Olympus DP70-CU CCD camera attached to a Zeiss Axioplan compound microscope equipped with differential interference contrast optics. For size measurements, 20 spores of each specimen were used, unless indicated otherwise. Colors were described as a percentage of CMYK values using the INVAM color chart (http://invam. caf.wvu.edu/otherinfo/articles/colorchart.htm).

DNA was extracted from 10 mg subsamples using a Qiagen DNeasy Plant Mini Kit (Qiagen, Hilden, Germany). Polymerase chain reaction (PCR) was conducted in two steps as a nested PCR using various primer combinations. For amplifying the complete 18S small subunit of specimen 1, universal primers were used (White et al. 1990). Preliminary experiments with universal primers had shown that the specimens belonged to the Diversisporales clade. Therefore, a primer was designed for specific amplification of the 3' end of the small subunit and the ITS of this lineage (GLOC1355: TCTTAGAGGGACTATTGGCATTT). It was used in the second step of the nested PCR together with ITS4i (Redecker et al. 2003), as second primer. In the first PCR step primers NS5/ITS4 (White et al. 1990) were used. Cycling parameters were as described by Hijri et al. (2006).

PCR products were ligated into pGEM-T (Promega, Wallisellen, Switzerland) and used to transform competent *Escherichia coli* DH5alpha. Clones were screened by blue/ white screening and colony PCR and sequenced in both directions using a Applied Biosystems Prism Dye Terminator 3.1 kit on an ABI 310 capillary sequencer (Applied Biosystems, Foster City, USA).

Sequence Navigator (Applied Biosystems) and Bio Edit (Hall 1999) were used to proofread and assemble the sequences. The near-complete sequence of the small subunit was assembled from sequenced fragments using variable regions for verifying that the fragments were from the same organism. To exclude chimaera formation, the fragments were subjected separately to phylogenetic analysis in PAUP* 4 (Swofford 2001). Moreover, the sequences were tested using Chimaera Check (Cole et al. 2003) and Bellerophon (Huber et al. 2004).

The small subunit sequence was included into a dataset comprising 39 selected taxa from the Glomeromycota with special focus on the Diversisporales. The alignment was done manually, and some regions were excluded from analysis because they could not be aligned unambiguously, resulting in 1,541 positions that were eventually analyzed. Phylogenetic trees were obtained under distance and maximum likelihood criteria using PAUP* 4. The neighbor joining and heuristic search algorithms were used, respectively. A gamma shape parameter of 0.5 was assumed for neighbor joining. Bayesian analyses were performed using MrBayes 3.1.1. (Ronquist and Huelsenbeck 2003) for Macintosh. Four chains were run over 9.2×10^6 generations under a general time-reversible model. Every 500th tree was sampled, and the first 10% of trees were discarded as a "burnin" before constructing a 50% consensus tree.

Partial small subunit sequences from the 3' end of the gene from more specimens were included into another dataset, comprising 31 taxa and 314 characters, containing sequences from spores of representative species of all glomeromycotan clades, as well as numerous partial 18S subunit sequences obtained from colonized roots. For Bayesian analysis, 5×10^6 generations were run, and the trees were processed as described above. For maximum likelihood analysis, the GTR+G+I model was used as determined using Modeltest 3.5 (Posada 2004).

ITS1/5.8S subunit/ITS2 sequences were placed in a dataset containing all available sequences from the *Diversispora/Glomus* group C clade. Some sequences from the public databases labeled *G. intraradices* were excluded because of probable misidentification. The 5.8S and ITS2 regions were used for phylogenetic analysis. The dataset comprised 22 taxa, and a total of 352 characters were used for the analysis. For Bayesian analysis, 5×10^6 generations were run, and the trees were processed as described above. The GTR+I model was used for maximum likelihood analyses.

DNA sequences were submitted to the European Molecular Biology Laboratory database under the accession numbers provided in Table 1. The alignments were deposited in TreeBase with the accession numbers M3072 and M3073.

Results

Morphological analyses

Specimen AC/Pohn99-001 from Micronesia (Fig. 1) had a size of $7 \times 7 \times 4$ mm. The peridium was smooth and had a

Table 1 Specimens studied and sequences obtained from them

Species	Specimen	Origin	Sequences
G. fulvum	AC/Pohn99-001	Pohnpei,	AM418543,
	CT 0.5 0.40	Micronesia	AM418544
	CL/Mart05-049	Martinique,	AM418545,
		Caribbean	AM418546
	CL/Mart05-111	Martinique,	AM418547,
		Caribbean	AM418548
G. pulvinatum	CL/Mart05-035	Martinique,	AM418549,
		Caribbean	AM418550
<i>G. megalocarpum</i> sp. nov	CL/Guad05-051	Guadeloupe,	AM418551,
		Caribbean	AM418552

tan to yellowish-brown color (20/60/100/10 on the INVAM color chart). One flat side of the sporocarp was not entirely covered by the peridium ("umbilicate sporocarp"). Spores were not embedded into hyphae. The spore mass was yellowish-brown (20/40/80/0).

Spores were oval to oblong to slightly claviform with a size of $80-138 \times 63-85 \ \mu m$ (average $126 \times 81 \ \mu m$). The spore content was yellowish-brown and had a dense, fibrous appearance. The spore wall was 2.5 to 5 μm thick and layered, the sublayers were not separating under pressure. The hyphal attachment was 7.5 to 10 μm broad, as well as the hyphae themselves. Spore contents were occluded by a prominent, bulging septum, apparently originating from the innermost wall layer(s) (Fig. 2d). These morphological features place the sporocarp into the species *Glomus fulvum* (Berk. & Broome) Trappe and Gerd.

Two specimens from the Caribbean (CL/Mart05-049 and CL/Mart05-111; Fig. 2a and b) were morphologically similar and also fit the description of *G. fulvum*. The sporocarps had diameters of 7 and 10 mm, respectively. However, their spores were generally shorter than in the Micronesian specimen (dimensions, $65-95 \times 75-95 \mu m$ and $80-95 \times 65-75 \mu m$, respectively). Two layers of the spore wall could be clearly distinguished in some spores. The spore wall was 2.5–4 μm thick, the layers were 1.2 to 2.4 μm wide.

The remaining two Caribbean specimens (CL/Mart05 05–35 and CL/Guad05–51) were less strongly pigmented. Sporocarp CL/Mart05 05–35 (Fig. 2c) had a diameter of 5 mm and fit the description of *Glomus pulvinatum* (P. Hennigs) Trappe and Gerd. The spore mass had a color of pale tan to ochre (0/20/40/0). Its spores had a diameter of 60–85 μ m differing from *G. fulvum* by their globose to subglobose shape. The hyphal attachment was equally broad (8–10 μ m) as in *G. fulvum*. The spore wall was 3 to 5 μ m thick with no apparent subdivision into layers. Fig. 1 Morphology of *G. fulvum* AC/Pohn99-001 from Micronesia. **a** Two halves of the sporocarp, the *top left* part shows the cutting plane, the *bottom right* the exterior surface. **b** Separated spores scraped from the sporocarp, showing the variation in spore shape. Note the dense spore contents. **c** Spore showing the hyphal attachment with the septum. **d** Close-up of the septum showing the wall layers



Specimen CL/Guad05-51 was a sporocarp considerably larger than the others (Fig. 3). It could not be assigned to a known morphospecies and is therefore described as *Glomus megalocarpum* sp. nov. None of the specimens mentioned showed a reaction with Melzer's reagent in the spores or attached hyphae.

Taxonomy

Glomus megalocarpum D. Redecker sp. nov.

Sporocarp firm, irregularly lobate, length approximately 3.8×2 cm, and with a brownish rind, whitish-velvety on the



Fig. 2 Spores of the sporocarp specimens from the Caribbean. a *G. fulvum*, specimen CL/ Mart05-049, b *G. fulvum*, specimen CL/Mart05-111, c *G. pulvinatum*, specimen CL/ Mart05-035, d *G. megalocarpum* sp. nov. CL/Guad05-51 **Fig. 3** Morphology of *G. megalocarpum* sp. nov. **a** Sporocarp CL/Guad05-51 in the field: approximate size along its long axis, 38 mm. **b** Spores in PVLG, note the triangular cavity separated by the septum (*asterisk*). **c** Dried slices of the sporocarp. **d** Spores in PVLG, showing variation in shape. *Scale bars* 50 μm in **b**, 1 cm in **c**, and 100 μm in **d**

outside. No smell perceptible. Composite spore mass in the interior pale tan to ochre (0/20/30/10 on the INVAM color chart). Spores obovate to pyriform or slightly irregular with dimensions of (56)65–95(115)×38–57(62) μ m (arithmetic mean, 81×47 μ m; *n*=40). Spore wall single-layered, hyaline, and 1.7–4 μ m thick. Hyphal attachments usually not persistent. Spores occluded by a prominent septum, 7–8 μ m broad, continuous with an inner lamina of the spore wall, separating a triangular cavity from the remainder of the spore lumen. Sporogenic hyphae thin-walled, usually not remaining attached. Gleba indistinct, spores tightly packed in irregular arrangement. Peridial hyphae rust-colored at higher magnification. Distinct by characteristic ribosomal DNA sequences: e.g., TTGAAGTTTGG GAGCGG in the 18S small subunit.

Sporocarpium firmum, irregulare, lobatum, 38×20 mm, interiore pallide fulva vel ochracea, peridio brunneo, superficie tomentosa, inolens. Sporae obovatae, piriformes, (56)65–95(115)×38–57(62) µm, tunicis hyalinis, 1.7–4 µm crassis, strato singulare. Hyphae sporophorae ephemerae, tunicis tenuibus. Gleba indistincta, sporis ordinatione densa et irregulare. Sporae septo occlusae, cavitatem

triangularem 7–8 µm latam segregante. Hyphae peridii ferrugineae. Sequentia typica acidi desoxyribonucleici ribosomatum: TTGAAGTTTGGGAGCGG.

Distribution and habitat	Known only from type collection
Mycorrhizal associations	Unknown
Etymology	Referring to the unusually large sporocarp
Collection examined	Guadeloupe, Vieux-Fort, Ravine Blondeau, mesophile forest, on forest floor, leg. C. Lécuru, 5.9. 2005. Holotype CL/Guad05-051 (LIP).

This species differs from other sporocarpic *Glomus* species by a number of characters, among them the large sporocarp size. *G. fulvum* and *G. pulvinatum* are distinct by the shape and color of their spores, as well as their rDNA sequences (see below). *G. canadense* (Thaxt.) Trappe and Gerd. was reported to have hyaline to pale yellowish spores with an oval to subglobose shape (Thaxter 1922), whereas



the spores of *G. megalocarpum* consistently have a higher length/width ratio. *G. fragile* (Berk. & Broome) Trappe and Gerd. also differs in spore shape and size and was reported to have a two-layered wall. *G. flavisporum* (M. Lange & Lund) Trappe and Gerd. has larger spores with thick, brown walls. *G. boreale* (Thaxt.) Trappe and Gerd. has red-brown spores (Gerdemann and Trappe 1974). Sporocarps of *G. cuneatum* McGee & A. Cooper contain dark-colored spores and differ in their unique sporocarp anatomy (McGee and Trappe 2002).

Fig. 4 Phylogenetic tree of the Diversisporales and outgroups based on Bayesian analysis of partial sequences (314 base pairs) of 18S small ribosomal subunit sequences. The tree was rooted with *Archaeospora/Paraglomus*. The first/second numbers on the nodes denote neighbor-joining bootstrap values from 1,000 replications or posterior probabilities from Bayesian analysis, respectively

Molecular phylogeny

A sequence comprising the complete 18S small subunit, ITS1, 5.8S subunit, and ITS2 regions was obtained from specimen AC/Pohn99-001 (*G. fulvum*, Micronesia). From the other specimens, a region of 315 base pairs at the 3' end of the small subunit, as well as ITS1, 5.8S subunit, and ITS2 were sequenced. Two clones spanning these regions were analyzed from each specimen to estimate the amount of intraspecific diversity.



Figure 4 shows a phylogenetic tree obtained by Bayesian analysis of partial 18S subunit sequences. The sequences from all sporocarp specimens form a monophyletic sister group to *D. spurca/G. versiforme* within the Diversisporales. The common clade as well as the sister group relationship received medium to high bootstrap support. The topology of the tree concerning these clades was verified using maximum likelihood analyses. In the analyses using the full-length small subunit sequence of *G. fulvum* AC/Pohn99-001 (results not shown), higher support (97% bootstrap support in distance analysis and 100% posterior probability in Bayesian analysis) could be obtained for the clade uniting *G. fulvum* and *D. spurca*/*G. versiforme*.

In the ITS tree (Fig. 5), the sporocarp specimens are clearly separated from the *Diversispora* clade, which is represented here by *G. versiforme* and *G. aurantium*. ITS sequences of *D. spurca* are not available in the databases. In some regions, the ITS are difficult to align between the two major clades in the tree. The branching order among the sporocarp specimens is almost identical to the small subunit tree. The different clones obtained from specimens *G. fulvum* CL/Mart05-049 and CL/Mart05-111 are extremely similar (1 to 6 positions different out of 314 in the 18S subunit



and 1 to 9 positions out of 351 in 5.8S subunit and ITS2) and cannot be separated conclusively. Their intermingled grouping in the tree proves that they belong to the same species. The sequence from *G. fulvum* AC/Pohn99-001 is closely associated with the latter two sequences, but shows some sequence differences. *G. megalocarpum* CL/Guad05-51 and *G. pulvinatum* CL/Mart05-35 are clearly separated from the *G. fulvum* sequences and from each other.

No root colonization could be detected by Trypan Blue staining 6 months after the inoculation of *Plantago media* with *G. fulvum*. Therefore, its mycorrhizal status remains to be elucidated.

Discussion

The morphological differences among the sporocarps are in concordance with the rDNA phylogenetic trees. The *G. fulvum* isolates have spores similar in shape, size, and color, whereas the other two can be easily distinguished.

G. fulvum belongs to a group of sporocarpic species of *Glomus* that were described at a time when taxonomy of that genus was very much focused on sporocarp formers (Thaxter 1922; Gerdemann and Trappe 1974). *G. fulvum* was originally described as *Paurocotylis fulva* by Berkeley and Broome (1873), and transferred to *Glomus* by Gerdemann and Trappe (1974). Thaxter (1922) and Gerdemann and Trappe (1974) also listed several similar, putatively related species (*G. canadense, G. pulvinatum*). In particular, Thaxter (1922) highlighted the septum as a distinctive characteristic of *G. fulvum* and relatives.

Interestingly, *G. fulvum* was reported to be common in tropical areas and to have a wide distribution (Sri Lanka,

Caribbean, Brazil, South Pacific; Gerdemann and Trappe 1974). The present specimens from Micronesia and the Caribbean fit well into that context. *G. pulvinatum* was reported from tropical America, Tasmania, and New Zealand (Gerdemann and Trappe 1974). Some other morphologically related species mentioned by these authors (*G. boreale, G. canadense, G, flavisporum*) were reported from temperate/boreal habitats.

In rDNA-based molecular phylogenies, *Diversisporal Glomus* group C forms a clade well separated from other *Glomus* groups (Schwarzott et al. 2001). However, it is not united by any obvious morphological characteristics on the light microscopic level that could aid distinguishing it from other *Glomus* species. Walker and Schüßler (2004) reported that *D. spurca* can be differentiated from *Glomus* species by a flexible inner wall. However, this is not undisputed (Kennedy et al. 1999), and there is no clear evidence that the other species in the Diversisporaceae share this putative morphological feature. No clear evidence for such an inner flexible wall could be found in the sporocarpic species analyzed here, even in spores with cracked walls.

All species in the *G. fulvum* group share the ability to form relatively large, compact sporocarps. The sporocarp of *G. megalocarpum* may be one of the largest glomeromy-cotan sporocarps reported so far. The spores of fungi of the *G. fulvum* clade show a prominent septum that closes a broad hyphal attachment. A collar-like remnant of the spore wall often protrudes beyond the septum. The spore content is often extremely dense, sometimes appearing stringy.

The prominent septum that occludes the mature spore appears as a potential uniting morphological feature between the *Diversispora* and *G. fulvum* clades. *G. versiforme* also has a rather prominent septa that look

Target taxon of signature	Diversisporales	Diversisporaceae	
Signature sequence Specimen	TYACCGGRAGGTRT	CTTTGGATTRGGGTTTAGGGRTC	
AC/Pohn99-001		ATTCG.	
CL/Mart05-049			
		ACTTCG.	
CL/Mart05-111			
		ATTCG.	
CL/Mart05-035	T		
	T	ATA.G.	
CL/Guad05-051		G.	
		G.	

Table 2 Signature sequences defined by Walker and Schüßler (2004), and their occurrence in the partial 18S subunit sequences presented in this study

Only the signatures located in the gene region obtained from all specimens are shown here.

similar to the ones in *G. fulvum*. In *G. spurcum*, it is apparently more difficult to see, in contrast to a putative relative *G. eburneum* (Kennedy et al. 1999). However, many other *Glomus* groups also feature a septum between hypha and spore. In many cases, it is not as obvious as in the *G. fulvum* group but the broad hyphal attachment of the latter may contribute to the visibility of the septum.

The sequence signatures used by Walker and Schüßler (2004) to define *Diversispora* and the corresponding higher taxa only partly match the new sequences presented here. In the 18S subunit sequence of AC/Pohn99-001, the first signature sequence GGTTTH characteristic for the order Diversisporales is present, as well as the second (TYACCG GRAGGTRT). However, the sequence signatures for the family Diversisporaceae show numerous changes (three changes out of 16 positions in the first signature, 5/20 in the second, and 1/14 in the third). The same is true for the genus-specific positions (4/23, 1/16, 6/23). From the two signatures located in the gene region sequenced from all specimens in this study, the order-specific signature is relatively well conserved in the taxa presented in this study, in contrast to the family-specific signature (Table 2).

These findings indicate that either these taxa have to be revised to comprise the sporocarpic species analyzed here or a new taxa have to be established. The *G. fulvum* group is phylogenetically clearly distinct from *Diversispora/ Glomus* group C, and it shows some uniting morphological features. Therefore, it may be preferable to erect a separate genus rather than include it into *Diversispora*, which has been highly problematic to define morphologically. However, we will await the analysis of more potential members of the clade before taking formal action and, if possible, additional gene sequences.

Mycorrhizal associations of *G. fulvum* are unknown, and it was reported to fruit on leaves and rotten wood (Gerdemann and Trappe 1974). We did not succeed in establishing it in mycorrhizal symbiosis but of course this does not prove that the fungus is not able to form mycorrhizas. It should be noted that ectomycorrhizal fungi fruiting on pieces of wood were mistaken for saprophytes for a long time until they were detected in the symbiosis by molecular methods (Gardes and Bruns 1996).

In the past few years, a large database of environmental glomeromycotan rDNA sequences that originate from molecular field studies of colonized roots has accumulated. Although the *G. fulvum* group should be detected by the primer pair NS31/AM1, which was used in a number of studies (Helgason et al. 1998), no close matches to *G. fulvum* etc. could be detected among those sequences by Basic Local Alignment Search Tool searches or phylogenetic analyses. Because of the sampling locations of the respective studies, most database sequences were found in temperate regions, but it should be noted that Glo36, the

closest match to the *G. fulvum* group detected in an extensive study of a tropical forest in Costa Rica (Husband et al. 2002), is much closer to *G. versiforme/spurcum* than to *G. fulvum*. Therefore, the nutritional mode of *G. fulvum* and relatives remains to be demonstrated. We used the new sequence data to design the primer GLOC1355, which will be refined to obtain a primer that can be applied to identify members of this clade within colonized roots. More environmental studies, in particular from the tropical areas where the sporocarps were found, are necessary to understand the ecological role of this glomeromycotan clade.

We demonstrated that a fascinating and spectacular group of species of glomeromycotan fungi that have been known for a long time constitutes a major, previously unknown monophyletic clade. These findings reiterate that glomeromycotan diversity is still far from being exhaustively characterized on the molecular level.

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