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Glomeromycota: three new genera and glomoid species reorganized

Fritz Oehl^{1*}, Gladstone Alves da Silva², Bruno Tomio Goto³ & Ewald Sieverding^{4*}

 ¹Federal Research Institute Agroscope Reckenholz-Tänikon ART, Ecological Farming Systems, Reckenholzstrasse 191, CH-8046 Zürich, Switzerland
 ²Departamento de Micologia, CCB, Universidade Federal de Pernambuco, Av. Prof. Nelson Chaves s/n, Cidade Universitária, 50670-420, Recife, PE, Brazil
 ³Departamento de Botânica, Ecologia e Zoologia, CB, Universidade Federal do Rio Grande do Norte, Campus Universitário, 59072-970, Natal, RN, Brazil
 ⁴Institute for Plant Production and Agroecology in the Tropics and Subtropics, University of Hohenheim, Garbenstrasse 13, D-70599 Stuttgart, Germany
 ^{*}CORRESPONDENCE TO: fritz.oehl@art.admin.ch & sieverdinge@aol.com

ABSTRACT —Species in the orders *Glomerales* and *Diversisporales* (*Glomeromycetes*) with glomoid spore formation are reorganized based on combined ribosomal sequence and morphological analyses. Within the *Glomerales* two genera in the *Glomeraceae* (*Septoglomus, Simiglomus*) and one genus in the *Claroideoglomeraceae* (*Viscospora*) are proposed as new. *Paraglomerales* species (thus far monogeneric) also form glomoid spores that may all germinate directly through the spore wall instead through subtending hyphae as in *Glomerales*.

KEY WORDS - Glomus group A, Glomus group B, molecular phylogeny, rDNA

Introduction

Molecular biological studies have dramatically improved the understanding of phylogenetic relationships across a broad swathe of organisms. Arbuscular mycorrhizal (AM) fungi are no exception, as molecular analyses have expanded our knowledge of this group (e.g. Simon et al. 1992, Redecker et al. 2000a, b, Morton & Redecker 2001, Schwarzott et al. 2001, Schüßler et al. 2001, Souza et al. 2005, Palenzuela et al. 2008, 2010). This has led to taxonomic re-organization and the description of several new genera and families (e.g. Morton & Redecker 2001, Walker et al. 2007, Oehl et al. 2008). Schüßler et al. (2001) established the phylum *Glomeromycota* with four orders (*Archaeosporales, Diversisporales, Glomerales, Paraglomerales*) within one class (*Glomeromycetes*) described earlier by Cavalier-Smith (1998).

Within the Glomeromycota, species with glomoid spore formation represent the largest group. These species had previously been included within the Glomeraceae (Pirozynski & Dalpé 1989) of the suborder Glomerineae of the Glomerales (Morton & Benny 1990). However, after this large group of Glomus species was found to be polyphyletic (e.g. Redecker et al. 2000b, Schwarzott et al. 2001), several glomoid spore-forming species were separated from the Glomerales sensu Morton & Benny. Based on DNA sequence analyses, Gl. occultum was declared type species of the new genus Paraglomus (Paraglomeraceae; Morton & Redecker 2001) in the new order Paraglomerales (Schüßler et al. 2001). Glomus spurcum was moved as type species to a new genus Diversispora (Walker & Schüßler 2004) in the Diversisporaceae (Diversisporales) (Schüßler et al. 2001). More recently Oehl & Sieverding (2004) established Pacispora (Pacisporaceae, Walker & Schüßler 2004) with Glomus scintillans S.L. Rose & Trappe (Rose & Trappe 1980) as type species for all glomoid species possessing two spore walls and specific germination characteristics. Moreover, some glomoid species were recently transferred into the Ambisporaceae of the Archaeosporales (e.g. Gl. callosum, Gl. fecundisporum, Gl. leptotichum; Walker et al. 2007, Walker 2008). However, the remaining large group of Glomus spp. is considered particularly difficult to differentiate morphologically (Morton & Redecker 2001, Schwarzott et al. 2001, Walker & Schüßler 2004).

Phylogenetic reconstructions within AM fungi over the last 10 years indicate that this large group of Glomus species likely originated from a single ancestor, from which (according to Schwarzott et al. 2001) descend the Glomus groups Aa (related to Gl. mosseae and Gl. geosporum), Ab (related to Gl. intraradices and Gl. irregulare), Ac (related to Glomus sp. W3347, now Gl. hoi; see Daniell et al. 2001, Öpik et al. 2006), and B (related to Gl. etunicatum). Schüßler et al. (2001) also showed that the clades of groups Aa, Ab and B divide further. Adapting and slightly modifying the numbering system of Schwarzott et al. (2001), we differentiate the clades into group Aa1 (lead sp. Gl. mosseae), Aa2 (lead sp. Gl. hoi), Aa3 (lead sp. Gl. constrictum), Ab1 (lead spp. Gl. microcarpum, *Gl. macrocarpum*), Ab2 (lead spp. *Gl. intraradices*, *Gl. irregulare*, *Gl. aggregatum*), Ab3 (lead spp. Gl. sinuosum, Gl. coremioides), B1 (lead sp. Gl. etunicatum), and B2 (lead sp. Gl. viscosum). In the present study, "lead species" are those with typical morphological characters and not necessarily type species. Under "Results & discussion" we present the supporting phylogenetic clades and detail the major morphological spore characteristics for each of our groups - Aa1, Aa2, Aa3, Ab1, Ab2, Ab3, B1, B2. In those cases where morphological and molecular data are congruent, species of the same group and number can be classified together at the genus level (e.g., group Aa1 represents one genus, while Aa2 represents another genus), while the major groups A and B are suggested to be family levels. In the case of clades Ab1, Ab2, Ab3, however, the molecular database remains under-represented with regard to the known species and the results are contradictory, as some sequences are derived from species that have been only poorly characterized. Although the morphological data indicate a division of Ab into at least three subgroups (genera), we currently lack sufficient genetic data to support new genera. Thus, clades Ab1, Ab2, Ab3 will be considered in one single genus (group Ab) at this time.

Previous phylogenetic analyses indicated that several *Glomus* spp. (e.g. *Gl. versiforme*) belong to clade '*Glomus* group' 'C' (e.g. Schwarzott et al. 2001, Schüßler et al. 2001, Redecker et al. 2007), currently *Diversispora* (*Diversisporaceae*). Only six glomoid spore-forming species have so far been included in this genus (Walker & Schüßler 2004, Gamper et al. 2009, Schüßler & Walker 2010). The morphology of *Diversispora* spores differs, however, from spores of groups A and B. We call this group of species "group Ca" and will transfer them to the genus *Diversispora*.

Recently, Redecker et al. (2007) identified another *Glomus* group (including *Gl. fulvum* and *Gl. pulvinatum*) that was genetically related to group C but phylogenetically and morphologically clearly distinct from group Ca (*Diversispora*). The authors did not separate this group from the other species of group C (*Diversispora*), possibly due to lack of recognition of the morphologically distinct features of spores. Recently, Schüßler & Walker (2010) erected the genus *Redeckera* ("group Cb") with three species (*Gl. fulvum*, *Gl. megalocarpum*, and *Gl. pulvinatum*) within the *Diversisporaceae*.

When Schüßler et al. (2001) proposed a new phylum for the AM fungi, although they initiated a principal re-organization of species with the four orders, they did not propose families and genera that were clearly separated genetically from each other. For example, *Diversisporales* and *Diversisporaceae*, lacking either type genus or type species, were at that time invalidly described, while the same authors mentioned only that some Glomus groups in the Glomerales and Glomeraceae were clearly genetically different. This had the unfortunate result that other taxonomists or ecological mycorrhizologists continued to refer to Glomus group A, group B, and 'the like' as Schüßler et al. (2001) had called them. Even new taxa were being justified on the basis of small differences in partial rDNA sequences within such Glomus groups as A, B or C (e.g. Błaszkowski et al. 2004, 2006, 2008, Oehl et al. 2005). Furthermore, in a recent study (Gamper et al. 2009) species of Glomus, Diversispora and others, e.g. *Glomus versiforme* and *D. celata*, were shown to be phylogenetically strongly related, although no attempts were made to reorganize Gl. versiforme and related 'Glomus' spp. into Diversispora. In ecological studies, so-called 'environmental sequences' (DNA sequences from AM species as yet unrecorded in databases, often directly recovered from mycorrhizal root or soil analyses instead of from identified spores) are named and referred to as 'Glomus sp. uncultured' or *'Glomus* unknown,' not particularly helpful to someone who wishes to know whether the species so named represents *Diversispora*, *Glomus* group A or B sensu Schüßler et al., or a different group that might have completely different spore formation.

It is clear that an improved phylogeny of *Glomus* is urgently needed (see also Krüger et al. 2009, Stockinger et al. 2009). Currently, new or unknown species that are related to 'groups of genetically related fungi' but belong to clearly distinct phylogenetic groups continue to be called '*Glomus*' and thus remain incorrectly classified. One motivation for this paper was the feeling that because *Gl. versiforme* has been supported as belonging to *Diversispora*, it should officially be included within that genus. From that it followed that if group A could phylogenetically be separated from groups B and C (*Diversispora*), a new organization of the known and described species is both justified and necessary. A first attempt was tried only recently by Schüßler & Walker (2010); however, these authors based their conclusions almost exclusively on molecular analyses and did not correlate their phylogenetic results with the morphological characters of the different phylogenetic clades that our analyses indicate.

The objectives of the present study were therefore to: 1) identify and describe morphological differences or similarities of species in the various '*Glomus* groups,' and 2) reorganize the species into existing or new genera as accurately as possible according to currently available information. Thus, we aim here at re-organizing the AM fungal species with glomoid spore formation using existing genetic information combined with existing or new morphological spore characteristics within the *Glomerales* and *Diversisporales*.

Material & methods

Specimen analyses

We analyzed specimens representing 101 of a total 117 described *Glomus* and *Diversispora* spp. (TABLE 1). Holotype, isotype, paratype and ex-type materials were examined along with representatives from institutional herbaria — OSC, FH, Z+ZT, URM (Recife, Brazil), Embrapa Agrobiologia (Seropédica, Brazil), DCS-UFLA (Lavras, Brazil), International Culture Collection of Vesicular-Arbuscular Mycorrhizal Fungi (INVAM), Swiss collection of Arbuscular Mycorrhizal Fungi (SAF, Agroscope ART Zurich) — and private AM fungal collections curated by Sieverding, Oehl, Trappe, Błaszkowski, McGee, and Goto. The Hall & Abbott (1979) photographic slide collection was also reviewed. For this paper, all original species descriptions and published species emendations were also considered.

Older specimens (mounted on microscopic slides prior to 1990) were mostly mounted in lactophenol, while others were fixed with polyvinyl alcohol-lactic acid-glycerol (PVLG) or in a mixture of PVLG + Melzer's reagent, which post-1990 are the principal fixing media (Brundrett et al. 1994). Newly mounted spores and sporocarps from collections or from cultures were fixed using the latter two fixing media, or occasionally also in a mixture of 1:1 lactic acid to water, in Melzer's reagent, and in water.

When available, spores freshly isolated from soils or bait cultures were also mounted and analyzed. Spore wall terminology follows the nomenclature of Walker (1983) and Stürmer & Morton (1997). Analyses of the spore walls, the germination structures and all other mycorrhizal structures were performed using compound microscopes at 100– 1000×. Most photographs were taken with a digital camera (Olympus model DP70-CU) mounted on a Zeiss Axioplan compound microscope, although a few were taken at OSC, URM or INVAM during short-term visits. Legends and scales were inserted with Adobe Photoshop CS2 9.0.

All spore observations and all information on spore characteristics are based on spores extracted from soil, from trap cultures or from single or multiple spore-derived pure cultures. No information is provided from in-vitro-cultured materials.

Molecular and phylogenetic analyses

GENES ANALYZED: Partial sequences of β -tubulin and rRNA (SSU and LSU) genes were used to reconstruct, by independent analyses, the phylogeny of the *Glomeromycota*. Intron sequences from the β -tubulin gene were excluded and only exon regions were considered within analyses. ITS sequences were also used to clarify relationships among members of the *Diversisporales*.

SEQUENCE ALIGNMENT: The sequences (all obtained from the National Center for Biotechnology Information-NCBI) were aligned using the program ClustalX (Larkin et al. 2007) and edited with the BioEdit program (Hall 1999) to obtain a final alignment.

Species name	Type material (observer)	Non type material (observer)	
Diversispora spurca	Ex type at INVAM (Oehl)	Specimen from Bolivia & Oman (Oehl)	
Di. celata	Ex type, inclusive pure cultures (Oehl)	Specimen from Switzerland (Oehl)	
Glomus achrum	Type (Oehl), ex type (Goto)	*	
Gl. africanum	No access		
Gl. aggregatum	Type OSC #40254 (Oehl)	OSC #45840 (Oehl)	
Gl. albidum	Holotype OSC #40400 (Oehl)		
Gl. ambisporum	Holotype OSC #44289 (Oehl)	Specimen from Bolivia, Benin (Oehl)	
Gl. antarcticum	No access		
Gl. arborense	No access		
Gl. arenarium	Type at OSC (Oehl), ex type (Goto)	Specimen from Chile and UAE (Sieverding)	
Gl. atrouva	Ex type (Oehl, Goto)		
Gl. aurantium	Type (Oehl)	Specimen from Germany (Oehl)	
Gl. aureum	Holotype Z+ZT, isotypes OSC #; FB;	Specimen & pure cultures from	
	Pure culture (Oehl)	Central Europe (Oehl)	
Gl. australe	No access	(Trappe, McGee, Oehl)	
Gl. avelingiae	No access		
Gl. badium	Holotype Z+ZT, isotypes deposited at	Specimen from Central Europe	
	OSC (Sieverding, Oehl)	(Oehl, Sieverding)	
Gl. bagyarajii	No access		
Gl. bistratum	Type (Oehl), ex type (Goto)		
Gl. boreale	Photographic slide collection of Hall &		
Cl hotrwoides	Abbott (1979)	Trappe collection (Oebl)	
Gl. brohultii		Specimen from Benin (Oehl) and	
0	Ex type (Sieverding)	Costa Rica (Sieverding)	
Gl. caesaris	Holotype Z+ZT, isotypes OSC #: FB:	Specimen and pure cultures from	
	Pure culture (Oehl)	Germany (Oehl)	
	· · · ·		

TABLE 1. Type and non-type specimens from 115 *Glomus* and 2 *Diversispora* species analyzed for reorganization of the *Glomerales*.

TABLE 1, continued		
Gl. caledonium	Type FH (Oehl)	Specimen and pure cultures from
Gl. canadense		Central Europe (Oehl) Thaxter collection, Trappe collection
Gl. candidum	No access	(Oelli)
Gl. canum	Ex type (Oehl, Goto)	
Gl. cerebriforme	Ex type (Oehl)	
Gl. citricola	No access	
Gl. claroideum	Type OSC #40252 (Oehl)	Specimen from Brazil, Europe (Oehl),
		cultures from Benin (Oehl)
Gl. clarum	Ex type specimen (Sieverding)	INVAM; specimen from Colombia
Cl. alamiata amuna	Err terme (Oahl))	(Sieverding)
Gl. constrictum	Paratype (Oell)) Paratypes OSC #38850 (Oehl	Specimen and pure cultures from
Gi. constructum	Sieverding)	Central Furope (Oehl)
Gl. convolutum	Paratype OSC #30986, ex isotype (Oehl)	Sentral Europe (Sent)
Gl. coremioides	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Specimen from Brasil (Goto); Trappe
		collection (Oehl)
Gl. coronatum	Ex type (Oehl)	Specimen from India, Mexico, Spain
		(Oehl)
Gl. corymbiforme	Type OSC # 53909 (Oehl)	
Gl. cuneatum	Ex type (Goto)	
Gl. delhiense	No access Fr trme (Ochl)	Specimen from Central Europe
Gi. utupnunum	Ex type (Oelli)	Bolivia (Oebl)
Gl dimorphicum	No access	Bolivia (Gelli)
Gl. deserticola	Holotype 44474, Paratype OSC #47834	Colombia (Sieverding)
	(Oehl)	
Gl. dolichosporum	No access	
Gl. drummondii	Type (Oehl), ex type (Goto)	
Gl. eburneum	Ex type at INVAM (Oehl)	Specimen from Bolivia & Oman
C1		(Oehl)
Gl. epigaeum	Holotype OSC #39475 (Oeni)	Spacimon from Ponin Polivia
Gi. etunicatum	Holotype OSC (Oelli)	Paraguay Europe Mexico (Oehl)
Gl. fasciculatum	Type sensu (Walker & Koske 1987)	Specimen from Central Europe
,	/1	(Oehl, Sieverding)
Gl. flavisporum		Trappe collection (Oehl)
Gl. formosanum		Specimen from UAE (Sieverding)
Gl. fragile		Trappe collection (Oehl)
Gl. fragilistratum	Ex type at INVAM (Oehl)	Specimen from Switzerland (Oehl)
Gl. fuegianum		Trappe collection (Oehl)
Gl. fulvum		Trappe collection (Oehl)
Gl. geosporum	Type OSC #29419 (Oehl)	Specimen and pure culture and from
Gl øibhosum	Type: Błaszkowski online pages	Specimen from UAE (Sieverding)
Gl. globiferum	Holotype OSC #46718 (Oehl)	Specimen at INVAM (Oehl)
Gl. glomerulatum	Isotype OSC #46674, ex type	Specimen from Brazil (Goto)
8	(Sieverding)	1
Gl. goaense	No access	
Gl. halonatum	Paratype Trappe 3594 (Oehl)	Specimen from Brazil (Goto)
Gl. heterosporum	Holotype 44288 (Oehl)	Specimen from Germany (Oehl)
Gl. hoi	Paratype Trappe 3058 (Oehl)	Specimen from Germany (Oehl),
Cl. I. J. J. J. J. J. J. Star		cultures from Benin (Oehl, IITA)
Gi. nyaérabaaénse		from Colombia (Sieverding)
Gl indicum	No access	nom Colonidia (Sieverunig)
Gl. insculptum	Type (Oehl)	
Gl. intraradices	/ A ` '	Specimen and cultures from Europe
		&, specimen from INVAM (Oehl)
Gl. invermaium	Isotype OSC #43941 (Oehl)	Specimen from UAE (Sieverding)
Gl. iranicum	No access	· · · · · · · · · · · · · · · · · · ·
Gl. irregulare	Type (Oehl)	Specimen from Switzerland (Oehl)

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TABLE 1, concluded.		
Gl. kerguelense		Specimen from UAE (Sieverding)
Gl. lacteum	Type OSC (Oehl)	1
Gl. lamellosum	Isotype OSC #50183 (Oehl)	Specimen from Poland, Germany
	/1	(Oehl, Sieverding)
Gl. liquidambaris		Specimen from Brazil (Oehl, Goto)
Gl. luteum	Type OSC, ex type INVAM (Oehl)	1
Gl macrocarbum	-//	INVAM: specimen from Central
		Furope Benin Mexico (Oehl)
Gl. magnicaule	Ex type, photographic slide collection	Europe, Denni, Mexico (Oem)
5118.	by Hall & Abbott (1979) (Oehl)	
Gl manihotis	Holotype OSC #41498 (Oehl)	
Gl megalocartum	Holotype OSC #28739 ex type (Oebl)	
Cl. melanochorum	Holotype 000 #20700, ex type (0em)	
Gi. meunosporum	Holotype OSC #, ex type (Oell)	Constitution for an Marrian (Oall)
Gi. microaggregaium	Folotype OSC #46/19 (Oeni)	Specimen from Mexico (Oeni)
Gl. microcarpum	Ex type (Oehl)	Specimen from Central Europe
		(Oehl)
Gl. minutum	Ex type (Oehl)	
Gl. monosporum	Type OSC (Oehl)	Specimen from Switzerland (Oehl)
Gl. mortonii	Holotype OSC #49460 (Oehl)	Specimen from Germany (Oehl,
	• *	Sieverding)
Gl. mosseae	Holotype FH (Oehl)	Specimen and pure cultures from
		Germany (Oehl)
Gl_multicaule	Holotype OSC #35386 (Oehl)	
Gl multiforum	Type (Oehl)	Specimen from Switzerland &India
Gi. mangoram	Type (Oem)	(Och)
Cl. nanoluman	Isotupe OSC #49585	(Oelli)
Cl. backweaula	1sotype 03C #49585	Specimen from India and Penin
Gi. puchycuuie		(Osh)
CL to all'i form	Torrest DDD (O.11)	(Oeni)
Gi. painaum	Type at PDD (Oeni)	
Gl. pansihalos	Holotype OSC (Oehl)	Specimen at INVAM (Oehl)
Gl. pellucidum	Ex type (Oehl, Goto)	
Gl. perpusillum	Type (Oehl), Ex type (Goto)	
Gl. proliferum	Ex type (Oehl)	
Gl. przelewicense	No access	Specimen and pure pot cultures from
1		Switzerland (Oehl, Sieverding)
Gl pubescens		OSC #30990 (Ochl)
Cl publicatum		Trappe collection (Oehl)
Cl. putulatum	Holotimo OSC #46721 (Oahl)	Placekowski collection (Ochl)
Gi. pustulatum	Holotype OSC #46721 (Oeni)	Transmission (Ochl)
Gl. radiatum		Trappe collection (Oeni)
Gl. rubiforme	Type OSC #30931, paratypes (Oehl)	Specimen from Switzerland (Oehl)
Gl. segmentatum	Type , ex type (Oehl)	
Gl. sinuosum		Specimen and cultures from Central
		Europe, India, Oman, Benin, Bolivia
		(Oehl, Sieverding)
Gl spinosum	Ex type (Sieverding)	(o enil, ore veraning)
Cl spinuliforum	Holotume 7+7T isotopes at OSC & EB	Specimen from Switzerland (Oehl)
Gi. spinniger um	(O the Circuit in a)	Specifien from Switzenand (Oeni)
CI taiwayaya	(Oehl, Sieverding)	Smaaiman from Banin (Oahl)
Gi. taiwanense		Specimen from Benin (Oeni),
		Thailand (Sieverding), Brasil (Goto)
Gl. tenebrosum		Specimen from Switzerland (Oehl)
Gl. tenerum		Specimen from Australia (Oehl)
Gl. tenue		Specimen from alpine areas (Oehl)
Gl. tortuosum	Type OSC #40'251 (Oehl)	OSC #32894
Gl. trimurales	Holotype OSC #49584 (Oehl)	
Gl. verruculosum	Ex type, INVAM (Oehl)	
Gl. versiforme	71-7-1-2000	Specimen at INVAM: specimen from
Gi. reisgornie		Central Europe (Oebl. Sieverding)
Cl mariculiforum		OSC #28727 (Oabl)
Gi. vesiculijerum	En tre a (Oahl)	Coc #20/2/ (Utill)
GL VISCOSUM	Ex type (Oeni)	specifien at INVAM
Gl. walkeri	Type (Oeni), ex type (Goto)	
Gl. warcupii	Ex type (Oehl)	
Gl. xanthium	Type (Oehl) ex type (Goto)	
Gl. zaozhangianum	No access	

PHYLOGENETIC ANALYSES: The main tree was constructed by a heuristic search using the maximum likelihood (ML) method with the Phylogenetic Analysis Using Parsimony (PAUP) program version 4 (Swofford 2003). Maximum parsimony (MP) and neighbor joining (NJ) analyses with 1000 bootstrap replications also were performed by PAUP. Bayesian (two runs over 1 x 10⁶ generations with a burnin value of 2500) and maximum likelihood (1000 bootstrap) analyses were executed, respectively, in MrBayes 3.1.2 (Ronquist & Huelsenbeck 2003) and PhyML (Guindon & Gascuel 2003), launched from Topali 2.5. The model of nucleotide substitution was estimated using Topali 2.5 (Milne et al. 2004). Sequences from *Neurospora crassa, Boletus edulis, Rhizophydium sphaerotheca* were used as outgroups for *Glomeromycota*, and *Glomus intraradices* and *G. xanthium* for *Diversisporales* (ITS analysis).





FIG. 1A–B. Phylogenetic reconstruction of the *Glomeromycota* obtained from partial SSU rDNA sequences (~1800 bp). Group C are *Diversisporaceae*. The neighbor-joining (NJ), maximum likelihood (ML) and bayesian analyses were performed with GTR+G+I substitution model. Sequences are labeled with their database accession numbers. Bootstrap values (in %) are from NJ, maximum parsimony (MP) and ML analyses (1000 bootstraps), respectively. The lasting numbers below each branch line denotes the credibility value from the bayesian analysis. Only topologies with bootstrap values of at least 50% are shown (Consistency Index = 0.4701; Retention Index = 0.8241). (FIG. 1A left on p. 82; FIG. 1B above)

Results & discussion

Relationships between general phylogenetics and morphological features

All four orders of the *Glomeromycota* have species that form glomoid spores. *Paraglomerales* and *Archaeosporales* have low phylogenetic relationship with *Glomerales* and *Diversisporales* (e.g. Redecker et al. 2007, Palenzuela et al. 2008; FIGS. 1–2, 4). Representatives of *Ambispora, Archaeospora, Intraspora,* and *Paraglomus* form extraradical mycelia and mycorrhizal structures that stain faintly to not at all in trypan blue (Spain & Miranda 1996, Spain 2003, Spain et al. 2006, Sieverding & Oehl 2006). (The reasons for this are unclear: vesicle formation is rarely reported or might have been misinterpreted in *Archaeospora, Intraspora* and *Paraglomus*.) This differs from the current glomeralean and diversisporalean species, which have fungal structures that stain blue to deep



blue with trypan blue. They are also reported to form vesicular arbuscular mycorrhiza (Schüßler et al. 2001), except for *Gigasporaceae, Scutellosporaceae, Racocetraceae* and *Dentiscutataceae*, which form arbuscular mycorrhiza but not intraradical vesicles, or at least vesicles have never been observed (Bentivenga & Morton 1995, Oehl et al. 2008). To date *Paraglomerales* species are monogeneric and form glomoid spores that may all germinate directly through the spore wall instead of through the subtending hyphae as in *Glomus* and *Diversispora* (Spain and Miranda 1996; Oehl pers. obs. on *P. occultum*).



FIG. 2. Phylogenetic reconstruction of the *Glomeromycota* obtained from partial LSU rDNA sequences (~600 bp). The neighbor-joining (NJ), maximum likelihood (ML) and bayesian analyses were performed with GTR+G+I substitution model. Sequences are labeled with their database accession numbers. Bootstrap values (in %) are from NJ, maximum parsimony (MP) and ML analyses (1000 bootstraps), respectively. The lasting numbers below each branch line denotes the credibility value from the bayesian analysis. Only topologies with bootstrap values of at least 50% are shown (Consistency Index = 0.4207; Retention Index = 0.7818). (FIG. 2A left on p. 84; FIG. 2B above.)

Phylogenetic reconstructions for glomeromycotan species based on analyses of the 18S, 28S, and ITS regions of the ribosomal gene, and of the β -tubulin gene

The major clades for groups Aa, Ab, B and C were monophyletically obtained in the phylogenetic trees constructed for the genome regions analyzed (FIGs. 1-4). Clade group Aa was divided into three subclades with Gl. mosseae, Gl. hoi, and Gl. constrictum as lead species of Aa1, Aa2 and Aa3, respectively. Further separation of clade Ab is not clear, although former Sclerocystis spp. (group Ab3) and *Gl. badium* (morphologically also in group Ab3) are monophyletic (FIG. 1). Clades Ab1 and Ab2, which differ consistently in glomoid spore morphology (see below) and contain the largest number of species, were not re-organized because the genetic database is not yet complete. Of the species analyzed, only Gl. aureum represents clade Ab1, which should have Gl. macrocarpum as type species according to Clements & Shear (1931). However, the 18S rDNA region (FIG. 1) does not provide sufficient support by itself for creating a new genus. The two lead species of group Ab2 are Gl. intraradices and Gl. irregulare, but this group also is not yet recognized at the genus level. The group B clade clearly stands apart from the others and splits into *Glomus* groups B1 and B2. Glomus viscosum is hereafter attributed as a single species to group B2, while Gl. etunicatum was recognized as the lead species for the group B1 (FIG. 1). The ITS regions from clade groups Ca and Cb were analyzed because other sequence data were either unavailable for group Cb species or (for the 18S) available only for the lead species, Gl. fulvum. ITS analyses showed the Ca and Cb clades well separated from each other (FIG. 3).

Morphology of glomoid, diversisporoid, and pacisporoid spores

All glomoid spores of the present *Glomeraceae* and *Diversisporaceae* have one single spore wall with several layers. In many species the structural wall layer is the inner layer, although other species possess additional innermost (semi-)flexible layers. Because these additional layers are generally thin (often <1µm), closely adhere to the structural layer, and can be separated from the latter only through hard pressure on the cover slides, they are not considered separate walls. The presence or absence of innermost flexible layers is not sufficient to distinguish morphologically between *Glomus* groups since the majority of groups have some species with and some species without innermost flexible layers. Such characters as mycelial/ mycorrhizal infection features, spore size, spore wall color, wall ornamentation, and layer numbers or their staining features could rarely be attributed to all species within a specific clade.

Spores of groups A and B have a structural layer that forms a continuum from attached hyphal to spore wall layers (FIGS. 5–17). We refer to such spore types as GLOMOID.

Because Ca and Cb group spores have a structural layer that appears to be inserted into the wall of the terminal element of the attached hypha, the



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FIG. 3. Phylogenetic reconstruction of the *Diversisporales* obtained from ITS sequences (~500 bp). The neighbor-joining (NJ), maximum likelihood (ML) and bayesian analyses were performed with GTR+G+I substitution model. Sequences are labeled with their database accession numbers. Bootstrap values (in %) are from NJ, maximum parsimony (MP) and ML analyses (1000 bootstraps), respectively. The lasting numbers below each branch line denotes the credibility value from the bayesian analysis. Only topologies with bootstrap values of at least 50% are shown (Consistency Index = 0.5539; Retention Index = 0.8092).

structural spore wall layers may not form a continuum with the hyphal wall layers (FIGS. 18–21). The inner wall layer appears to differentiate de novo, similar (but not equal) to inner walls in *Acaulosporaceae* or *Entrophosporaceae*

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FIG. 4. Phylogenetic reconstruction of the *Glomeromycota* obtained from partial β -tubulin sequences (~600 bp). The neighbor-joining (NJ), maximum likelihood (ML) and bayesian analyses were performed with GTR+G+I substitution model. Sequences are labeled with their database accession numbers. Bootstrap values (in %) are from NJ, maximum parsimony (MP) and ML analyses (1000 bootstraps), respectively. The lasting numbers below each branch line denotes the credibility value from the bayesian analysis. Only topologies with bootstrap values of at least 50% are shown. (Consistency Index = 0.3709; Retention Index = 0.6969).

(Oehl et al. 2006, Sieverding & Oehl 2006). In Ca the structural and (if present) inner flexible spore wall layers form a septum at the base or at a short distance from the base of the spore within the subtending hypha. In Cb the inner lamina of the structural wall layer always forms a septum at the spore base and the outer lamina may continue for $1-10 \mu m$ into the subtending hypha. We refer to this spore type as DIVERSISPOROID.

Pacispora spores superficially resemble glomoid spores, as the spore wall structural layer forms a continuum with the hyphal wall layers. However, *Pacispora* spores strongly differ from glomoid spores: i) in possessing an additional thick inner germinal wall that forms de novo and is clearly separate from the outer wall; ii) by species-specific germination structures resembling a germ shield; and iii) by germ tubes that arise from the 'shields' on the inner wall and penetrate directly the adjacent outer wall. We refer to this spore type as PACISPOROID.

Morphology of the subtending hypha, spore base, and spore pore closure

A combination of two major morphological features and one secondary feature are congruent with the major molecular phylogenetic groups of the glomoid, diversisporoid, and pacisporoid species (TABLE 2): i) the morphology of the spore base including the pore closure and position of the septum (FIGs. 5–23, 24–114); ii) the morphology of the subtending hypha (SH) (FIGS. 5–23, 24–114); and iii) secondarily, the prevailing formation and spore organization level within either sporocarps or spore clusters or singly in soil.

Spores of *Glomus* groups Aa(1,2,3) and Ab(1,2,3) all have sH that are concolorous with (or slightly lighter in color than) the structural spore wall, and the spore wall differentiation extends completely into the sH (TABLE 2, FIGS. 5–14, 24–89). In group B1 and B2 spores the structural spore wall differentiation also extends into the sH, but the sH colour changes abruptly to hyaline or white at the spore base, even when the spore wall layers are distinctly pigmented (TABLE 2, FIGS. 15–17, 90–98).

Group	Aai	Aa2	Aa3	Аві	Ab2
Lead species	Gl. mosseae	Gl. hoi	Gl. constrictum	Gl. macrocarpum Gl. microcarpum	Gl. intraradices Gl. irregulare
Recent or New genus	Funneli- formis	Simiglomus	Septoglomus	Glomus	Glomus
SH CHARACTE	RS				
Color	Same as spore wall	Same as spore wall	Same as spore wall	Same as spore wall	Same as spore wall
Shape	Funnel-shape to cylindric,	Cylindric/sl. funnel-	Cylindric to constricted,	Cylindric to slightly funnel-	Cylindric,
&	·	shape,	rarely sl. funnel-shape,	shape,	
thickness	thick walled	thick walled	+/- thick walled	thick walled	thick walled
Pore closure	Conspicuous septum under / at spore base	Sev. conspic. septa distant to spore base	Septum under / at spore base	Introverted sw thickening; +/- bridging septum	Open or septum distant to spore base; introverted sw thickening rudimentary
Spore formation	Singly or in few-spored sporocarps	Singly or open spore clusters	Singly or open spore clusters	Compact, unorganized multi-spored sporocarps, clusters or singly	Loose or compacted, unorganized clusters; also singly

 TABLE 2. Morphology of subtending hyphae (SH) and spore formation in glomoid spored species in *Glomerales* and *Diversisporales*

TABLE 2, concluded.

Group	Авз	B1	B2	Са	Св
LEAD	Gl. coremioides	Gl. etunicatum	Gl.viscosum	Gl. versiforme	Gl. fulvum
Recent or New genus	Glomus	Claroideo- glomus	Viscospora	Diversispora	Redeckera
SH CHARACTE	RS				
Color	Same as spore wall	White to hyaline, abrupt color change at spore base if spore wall pigmented	White to hyaline	Hyaline distant to septum that is concolorous with structural SWL	Hyaline distant to septum that is concolorous with structural SWL
Shape; &	Cylindric, funnel-, or bill-shaped,;	Conspicuously bill-shaped;	Cylindric;	Generally fragile, cylindric distant to septum, or constricted:	Generally fragile and inflated ;
Thickness	thick-walled	+/- thick-walled	thick-walled	thin-walled	thin-walled
Pore closure	Introverted sw thickening	Septum at spore base	Constricted wall or open	Septum at or beneath spore base, or open	Broad septum at spore base
Spore formation	In compact, organized sporocarps, around hyphal center; initially also singly	Singly	Singly or in loose clusters	In large, compact sporocarps, or cluster aggregates, or singly	In large, compact sporocarps/ clusters, open aggregates, or singly

In contrast, the spore wall differentiation generally does not extend into the SH in group Ca and Cb spores (TABLE 2, FIGS. 14–19, 99–114), as lamination and differentiation of the structural spore wall layer either does not extend into the SH or extends only a short distance from the base. Within these Ca and Cb, therefore, the SH are usually quite fragile (FIGS. 99–107) and hyaline to white, distal to the pore closure at the spore base (*Gl. spurcum* = *Di. spurca*) or in the SH (*Gl. eburneum*). The hyaline to white color of the SH behind the septum changes conspicuously when spores are pigmented. Cb species have broad SH and broad septa at the spore bases (FIGS. 108–114). Their SH are inflated (*Gl. pulvinatum*) or (sometimes) very short within compact sporocarps (e.g. *Gl. fulvum* and *Gl. canadense*).

Type of spore formation

Glomus group spore formation is intercalary or terminal on SH, both for spores formed either singly in soil (e.g. *Gl. arborense, Gl. etunicatum*) or in loose/dense spore clusters or sporocarps (e.g. *Gl. fasciculatum, Gl. pallidum, Gl. glomerulatum*).

FIGS 5–17. Examples of spore bases (with 1–3 spore wall layers; swL1–3) and subtending hyphae (sH) of species assigned to phylogenetic clades Aa1, Aa2, Aa3, Ab1, Ab2, Ab3, B1 and B2; structural spore wall layers are continuous with the sH wall. 5–14. sH of A clade spores are concolorous with or slightly lighter than the spore wall. 5–8. Clade Aa species never have an introverted wall thickening at the base; spore pores are generally closed by a conspicuous septum (sp) arising



from the structural layer and/or the (semi-)flexible layer beneath. **5–6**. Aa1. **5**. Lead species *Glomus mosseae*. **6**. *Gl. geosporum*. **7**. Aa3: lead species *Gl. constrictum*. **8**. Aa2: so far single lead species *Gl. hoi*. **9**. Ab2: without introverted wall thickening; spore base pore regularly open; lead species *Gl. intraradices*. **10–13**. Ab1: typical *Glomus* species with an introverted wall thickening at the spore base; sometimes a short bridging septum arising from the structural wall layer or innermost flexible lamina closes the spore base pore. **10**. Lead species *Gl. microcarpum*. **11**. *Gl. aureum*. **12**. Lead species *Gl. macrocarpum*. **13**. *Gl. pellucidum*. **14**. Ab3; also with a slight introverted wall thickening but pores generally closed by a short septum arising from the structural layer and/or the (semi-)flexible layers beneath. In some clade Ab3 species the spore head sometimes is also thickened towards the sporocarp surface; lead species *Gl. coremioides* and *Gl. sinuosum*. **15–17**. B1: spore SH bill-shaped and hyaline to white, even when the spore wall is pigmented. **15–16**. lead species *Gl. etunicatum*. **17**. *Gl. claroideum*.



FIGS 18–23. Clades Ca (*Diversispora*) and Cb (*Redeckera*) spore bases (with 1–3 spore wall layers; swL1–3) and subtending hyphae (sH): structural spore wall layer seemingly extends only a very short distance into the subtending hyphae, soon becoming hyaline to white and undetectable; often only the adherent outer spore wall layer(s) appears continuous with the mycelial hyphal wall to connect the two neighboring spores in compact sporocarps. The structural layer often appears inserted into the 'swollen hyphal tips'. **18–22**. Ca: spore SH generally white and cylindrical at the spore base, soon fragile. Septum (sp) arises from the structural spore wall layer and either closes the pore immediately or at some distance from the spore base (**18–21**) or the spore pore appears open (**22**). **23**. Cb: spore SH generally inflated a very short distance from the spore base, spores are 'endospore-like,' with a broad, thick septum formed by the structural layer.

Aa1 species generally form spores singly (e.g. *Gl. geosporum, Gl. caledonium*) or occasionally in sporocarps containing just a few (2–20) spores (e.g. *Gl. mosseae, Gl. monosporum, Gl. coronatum*). Aa2 and Aa3 species form spores preferably singly in soil or roots, or in very loose spore clusters (*Gl. hoi, Gl. constrictum, Gl. deserticola, Gl. xanthium*). Ab1 and Ab2 species generally form spores in multi-spored loose/dense variably sized spore clusters. During initial spore formation they may also form spores singly, although some isolates appear never to form clusters in single species cultures on certain hosts. Ab3 species form spores strictly in sporocarps but may initially form single spores on their mycelia (*Gl. sinuosum*).

B1 species generally form spores either singly in soil (e.g. *Gl. etunicatum*, *Gl. claroideum*, *Gl. walkeri*) or in variably sized loose spore clusters (*Gl. viscosum*). Sporocarp formation is extremely rare, if ever. The solitary B2 species — *Gl. viscosum* — forms spores in clusters of highly variable size. Ca species form spores singly in soils (e.g. *Gl. spurcum* = *Di. spurca*; *Gl. eburneum*) or in small to large and irregular shaped compacted clusters with numerous spores (e.g. *Gl. versiforme*, *Gl. epigaeum*) depending on culture and site conditions. Finally, Cb species generally form spores in large, compact sporocarps (e.g. *Gl. fulvum*, *Gl. fragile*) that are among the largest known within the *Glomeromycota* (especially *Gl. megalocarpum*). As the formation of a peridium around the sporocarps or single spores does occur in groups Aa1, Ab1, Ab3, and Cb, it is not a consistent morphological indicator at the family or genus level. Peridium formation has not yet been observed for B1, B2, and Ca species.

Organization level of sporocarps

Except for group Ab3, spores of other Glomus groups have (with some exceptions) limited to no organization in sporocarps or compacted spore clusters. As spores in sporocarps appear to be randomly distributed, SH are variably long (e.g. Gl. mosseae, Gl. coronatum, Gl. macrocarpum, Gl. ambisporum, Gl. versiforme, Gl. fulvum, Gl. pulvinatum). In contrast, Ab3 sporocarps are highly organized with spores forming around a central hyphal plexus. Variation in SH length is much lower (e.g. Gl. sinuosum, Gl. rubiforme). Sporocarp size variability of Ab3 species is low, largely dependent on the diameter of the central hyphal plexus and spore size in individual species. In unorganized sporocarps of the other 'Glomus groups' sporocarp size is much more variable due to a lack of symmetry, greater SH length variability, and greater variability of numbers of spores formed per sporocarp. According to our analyses, species that form dense, highly compact sporocarps (e.g., Gl. radiatum, Gl. vesiculiferum, Gl. segmentatum, Gl. convolutum, Gl. fuegianum) do not form spores around a central hyphal plexus arising from a single hypha but sporocarps appear to have several hyphal attachments. They thus probably belong to group Ab1 and so are not attributed to group Ab3.

Taxonomic revision

Glomeraceae Piroz. & Dalpé emend. Oehl, G.A. Silva & Sieverd. (*Glomus* group A) EMENDED DESCRIPTION: Spores formed terminally on or intercalary in hyphae, in soil and sometimes in roots, either singly, in spore clusters, or in multiple-spored loose to compact sporocarps when compact spores are randomly distributed or organized around a central plexus of hyphae. Compact sporocarps with or without peridium. Spores with one mono-to-multiple layered wall. Wall of the sH conspicuously continuous and concolorous with the spore wall,

or slightly lighter in color than the spore wall; SH funnel-shaped, cylindrical or constricted; forming typical vesicular-arbuscular mycorrhiza, with mycorrhizal structures that stain blue to dark blue in trypan blue.

TYPE GENUS: <i>Glomus</i> Tul. & C. Tul.	
OTHER GENERA: Funneliformis C. Walker & A. Schüssler, Septoglomus Sieverd.	
et al., <i>Simiglomus</i> Sieverd. et al.	

Glomus Tul. & C. Tul. emend. Oehl, G.A. Silva & Sieverd. FIGS. 25–74 (Glomus groups Ab1, Ab2 & Ab3)

EMENDED DESCRIPTION: Spores formed singly within soil or sometimes roots, in disorganized, multiple-spored loose spore clusters or in compact sporocarps; compact sporocarps without or with peridium, spores are either not organized in sporocarp, or organized around a central hyphal plexus. Spores with a mono-to-multiple layered wall. Wall of the SH conspicuously continuous and concolorous with the spore wall, or slightly lighter in color than the spore wall. Spore pore closure often by introverted wall thickening, sometimes supported by a short bridging septum, rarely open. Forming typical vesicular-arbuscular mycorrhiza, with mycorrhizal structures that stain blue to dark blue in trypan blue.

Type species: Glomus macrocarpum Tul. & C. Tul.

- *Glomus achrum* Błaszk., D. Redecker, Koegel, Schützek, Oehl & Kovács, Botany 87: 262. 2009.
- Glomus aggregatum N.C. Schenck & G.S. Sm., Mycologia 74: 80. 1982.

Glomus ambisporum G.S. Sm. & N.C. Schenck, Mycologia 77: 566. 1985.

Glomus antarcticum Cabello, Mycotaxon 51: 124. 1994

Glomus arborense McGee, Trans. Br. Mycol. Soc. 87: 123. 1986.

Glomus atrouva McGee & Pattinson, Austral. Syst. Bot. 15: 115. 2002.

Glomus aureum Oehl & Sieverd., J. Appl. Bot. 77: 111. 2003.

Glomus australe (Berk.) S.M. Berch, Can. J. Bot. 61: 2611. 1983. = *Endogone australis* Berk., Fungi Flora Tasman. 2: 282. 1859.

- Glomus badium Oehl, D. Redecker & Sieverd., J. Appl. Bot. Food Qual. 79: 39. 2005.
 = Funneliformis badius (Oehl, D. Redecker & Sieverd.) C. Walker & A. Schüssler, The *Glomeromycota* – a species list: 13. 2010.
- Glomus bagyarajii V.S. Mehrotra, Philipp. J. Sci. 126: 235. 1997.
- *Glomus bistratum* Błaszk., D. Redecker, Koegel, Symanczik, Oehl & Kovács, Botany 87: 267. 2009.
- *Glomus boreale* (Thaxt.) Trappe & Gerd., Mycol. Mem. 5: 58. 1974. = *Endogone borealis* Thaxt., Proc. Am. Acad. Arts Sci. 57: 318. 1922

- Glomus botryoides F.M. Rothwell & Victor, Mycotaxon 20: 163. 1984.
- Glomus brohultii Sieverd. & R.A. Herrera, J. Appl. Bot. 77: 37. 2003

Glomus canum McGee, Austral. Syst. Bot. 15: 116. 2002.

- Glomus cerebriforme McGee, Trans. Br. Mycol. Soc. 87: 123. 1986.
- Glomus citricola D.Z. Tang & M. Zang, Acta Bot. Yunn. 6: 301. 1984.
- Glomus clarum T.H. Nicolson & N.C. Schenck, Mycologia 71: 182. 1979.
 = Rhizophagus clarus (T.H. Nicolson & N.C. Schenck) C. Walker & A. Schüssler, The Glomeromycota a species list: 19. 2010.
- *Glomus clavisporum* (Trappe) R.T. Almeida & N.C. Schenck, Mycologia 82: 710. 1990.
 - = *Sclerocystis clavispora* Trappe, Mycotaxon 6: 359. 1977.
 - = Sclerocystis microcarpus S.H. Iqbal & Perveen, Trans. Mycol. Soc. Japan 21: 58. 1980.
- Glomus convolutum Gerd. & Trappe, Mycol. Mem. 5: 42. 1974.
- *Glomus coremioides* (Berk. & Broome) D. Redecker & J.B. Morton, Mycologia 92: 284. 2000.
 - = Sclerocystis coremioides Berk. & Broome, J. Linn. Soc., Bot. 14: 137. 1873.
 - = Xenomyces ochraceus Ces., Atti Accad. Sci. Fis. 8: 26. 1879.
 - = Ackermannia coccogena Pat., Bull. Soc. Mycol. Fr. 18: 182. 1902.
 - = Sphaerocreas coccogena (Pat.) Höhn., Sitzungsber. K. Akad. Wiss. Wien, Math.-Naturwiss. Kl., Abt. 1, 118: 401. 1909.
 - = Sclerocystis coccogena (Pat.) Höhn., Sitzungsber. K. Akad. Wiss. Wien, Math.-Naturwiss. Kl., Abt. 1, 119: 399. 1910.
 - = Ackermannia dussii Pat., Bull. Soc. Mycol. Fr. 18: 181. 1902.
 - Sphaerocreas dussii (Pat.) Höhn., Sitzungsber. K. Akad. Wiss. Wien, Math.-Naturwiss. Kl., Abt. 1, 118: 401. 1909.
 - = Sclerocystis dussii (Pat.) Höhn., Sitzungsber. K. Akad. Wiss. Wien, Math.-Naturwiss. Kl., Abt. 1, 119: 399. 1910.
- Glomus corymbiforme Błaszk., Mycologia. 87: 732. 1995.

Glomus cuneatum McGee & A. Cooper, Austral. Syst. Bot. 15: 117. 2002.

Glomus custos C. Cano & Dalpé, Mycotaxon 109: 502. 2009.

- = Rhizophagus custos (C.Cano & Dalpé) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.
- *Glomus delhiense* Mukerji, Bhattacharjee & J.P. Tewari, Trans. Br. Mycol. Soc 81: 643. 1983.
- Glomus diaphanum J.B. Morton & C. Walker, Mycotaxon 21: 433. 1984.
 - = Rhizophagus diaphanus (J.B. Morton & C. Walker) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.

Glomus dolichosporum M.Q. Zhang & You S. Wang, Mycosystema 16: 241. 1997.

Glomus fasciculatum (Thaxt.) Gerd. & Trappe, Mycol. Mem. 5: 51. 1974.

- = Endogone fasciculata Thaxt., Proc. Am. Acad. Arts & Sci. 57: 308. 1922.
- = Rhizophagus fasciculatus (Thaxt.) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.
- = Endogone arenacea Thaxt., Proc. Am. Acad. Arts & Sci. 57: 317. 1922
- = Rhizophagites butleri Rosend., Bull. Torrey Bot. Club 70: 131. 1943.

Glomus flavisporum (M. Lange & E.M. Lund) Trappe & Gerd., Mycol. Mem. 5: 58. 1974.

= Endogone flavispora M. Lange & E.M. Lund, Friesia 5: 93. 1954.

Glomus formosanum C.G. Wu & Z.C. Chen, Taiwania 31: 71. 1986.

Glomus fuegianum (Speg.) Trappe & Gerd., Mycol. Mem. 5: 58. 1974. *= Endogone fuegiana* Speg., Anal. Soc. Cient. Argent. 24:125. 1887.

Glomus globiferum Koske & C. Walker, Mycotaxon 26: 133. 1986.

Glomus glomerulatum Sieverd., Mycotaxon 29: 74. 1987.

Glomus goaense Khade, Mycorrhiza News 20(4): 21. 2010.

Glomus heterosporum G.S. Sm. & N.C. Schenck, Mycologia 77: 567. 1985.

Glomus hyderabadense Swarupa, Kunwar, G.S. Prasad & Manohar., Mycotaxon 89: 247. 2004.

Glomus indicum Błaszk., Wubet & Harikumar, Botany 88: 134. 2010.

Glomus intraradices N.C. Schenck & G.S. Sm., Mycologia 74: 78. 1982.
 = Rhizophagus intraradices (N.C. Schenck & G.S. Sm.) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.

Glomus invermaium I.R. Hall, Trans. Br. Mycol. Soc. 68: 345. 1977.

FIGS 24–46. *Glomus* clade Ab1 forms spores in unorganized clusters/sporocarps or (sometimes) singly in soil; subtending hyphae (SH) are concolorous with (or slightly lighter than) the spore wall, normally with introverted, species-specific basal wall thickenings; SH often cylindrical (very short if formed in compact sporocarps). Species-specific SH wall thickening may continue over rather long distances ($\leq 200-500 \ \mu m$) even in sporocarps. Spores with 1–4 wall layers (SWL1–SWL4); pore at spore base pores often closed by introverted wall thickenings or with an additional septum (sp) that arises from the structural layer, from an adherent innermost (semi-)flexible layer, or from both layers. Individual layers might be species-specifically ornamented (ORN). 24–25. *Gl. microcarpum.* 26. *Gl. macrocarpum.* 27. *Gl. spinuliferum.* 28. *Gl. magnicaule.* 29. *Gl. multicaule.* 30. *Gl. heterosporum.* 31. *Gl. aureum.* 32. *Gl. invermaium.* 33. *Gl. glomerulatum.* 34. *Gl. ambisporum.* 35. *Gl. warcupii.* 36. *Gl. melanosporum.* 37. *Gl. atrouva.* 38. *Gl. botryoides.* 39. *Gl. spinosum.* 40. *Gl. mortonii.* 41. *Gl. australe.* 42. *Gl. fuegianum.* 43. *Gl. tenerum.* 44. *Gl. pellucidum.* 45. *Gl. segmentatum.* 46. *Gl. convolutum:* spore peridium was removed by strong pressure on the cover slide.





FIGS 47–58. *Glomus* clade Ab2 form spores in disorganized loose/dense clusters/sporocarps (sometimes singly in soil); subtending hyphae (SH) concolorous with (or slightly lighter than) the spore wall. SH generally openly cylindric; introverted wall thickenings rudimentary, but usually not existent; spore base pore diagnostic: open or (rather rarely) with a septum arising from innermost adherent (semi-)flexible layer. Spores with 1–4 wall layers (SWL1–SWL4). 47. *Gl. intraradices.* 48. *Gl. aggregatum.* 49. *Gl. vesiculiferum.* 50. *Gl. globiferum.* 51. *Gl. tortuosum.* 52. *Gl. corymbiforme.* 53. *Gl. manihotis.* 56. *Gl. fasciculatum.* 57. *Gl. diaphanum.* 58. *Gl. pallidum.*

Glomus iranicum Błaszk., Kovács & Balázs, Mycologia 102: 1457. 2010.

- = Rhizophagus iranicus (Błaszk., Kovács & Balázs) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.
- Glomus irregulare Błaszk., Wubet, Renker & Buscot, Mycotaxon 106: 252. 2008.
 - *Rhizophagus irregularis* (Błaszk., Wubet, Renker & Buscot) C. Walker
 & A. Schüssler, The *Glomeromycota* species list: 19. 2010.
- Glomus liquidambaris (C.G. Wu & Z.C. Chen) Y.J. Yao, Kew Bull. 50: 306. 1995.
 - = *Sclerocystis liquidambaris* C.G. Wu & Z.C. Chen, Trans. Mycol. Soc. Rep. China, 2: 74. 1987.
 - = Sclerocystis cunninghamia H.T. Hu, Quart. J. Chinese For. 21: 52. 1988.

Glomus macrocarpum Tul & C. Tul., Giorn. Bot. Ital., Anno 1, 2(7–8): 63. 1845. *Endogone macrocarpa* (Tul. & C. Tul.) Tul. & C. Tul., Fungi Hypog.: 182. 1851.

- = Endogone guttulata E. Fisch., Ber. Schweiz. Bot. Ges. 32: 13. 1923.
- = Endogone nuda Petch., Ann. R. Bot. Gdns Peradeniya 9: 322. 1925.
- = Endogone pampaloniana Bacc., Nuovo Giorn. Bot. Ital., n.s. 10: 90. 1903.
- = Paurocotylis fulva var. zelandica Cooke, Grevillea 8: 59. 1879.

Glomus magnicaule I.R. Hall, Trans. Br. Mycol. Soc. 68: 345. 1977.

Glomus manihotis R.H. Howeler, Sieverd. & N.C. Schenck, Mycologia 76: 695. 1984.

Rhizophagus manihotis (R.H. Howeler, Sieverd. & N.C. Schenck) C. Walker
 & A. Schüssler, The *Glomeromycota* – a species list: 19. 2010.

Glomus melanosporum Gerd. & Trappe, Mycol. Mem. 5: 46. 1974.

Glomus microaggregatum Koske, Gemma & P.D. Olexia, Mycotaxon 26: 125. 1986.

Glomus microcarpum Tul. & C. Tul., Giorn. Bot. Ital., Anno 1, 2(7–8): 63. 1845. *= Endogone microcarpa* (Tul. & C. Tul.) Tul. & C. Tul., Fungi Hypog.: 182. 1851.

Glomus minutum Błaszk., Tadych & Madej, Mycologia 76: 189. 2000.

Glomus mortonii Bentiv. & Hetrick, Mycotaxon 42: 10. 1991.

Glomus multicaule Gerd. & B.K. Bakshi, Trans. Br. Mycol. Soc. 66: 340. 1976.

Glomus nanolumen Koske & Gemma, Mycologia 81: 935. 1990.

Glomus pachycaule (C.G. Wu & Z.C. Chen) Sieverd. & Oehl, comb. nov. MycoBank MB 519622

= Sclerocystis pachycaulis C.G. Wu & Z.C. Chen, Taiwania 31: 74. 1986.

Glomus pallidum I.R. Hall, Trans. Br. Mycol. Soc. 68: 343. 1977.

Glomus pansihalos S.M. Berch & Koske, Mycologia 78: 832. 1986.

Glomus pellucidum McGee & Pattinson, Austral. Syst. Bot. 15: 120. 2002.

Glomus perpusillum Błaszk. & Kovács, Mycologia 101: 249. 2009.

Glomus proliferum Dalpé & Declerck, Mycologia 92: 1180. 2000.

= Rhizophagus prolifer (Dalpé & Declerck) C. Walker & A. Schüssler, The Glomeromycota – a species list: 19. 2010.

Glomus pubescens (Sacc. & Ellis) Trappe & Gerd., Mycol. Mem. 5: 57. 1974.

- = Sphaerocreas pubescens Sacc. & Ellis, Michelia 2: 582. 1882.
- = Stigmatella pubescens (Sacc. & Ellis) Sacc., Syll. Fung. 4: 680. 1886.
- = Sclerocystis pubescens (Sacc. & Ellis) Höhn., Sitzungsber. K. Akad. Wiss. Wien, Math.-Naturwiss. Kl., Abt. 1, 119: 399. 1910.

= Endogone pubescens (Sacc. & Ellis) Zycha, Krypt.-Fl. Brandenburg 6a: 214. 1935.

Glomus radiatum (Thaxt.) Trappe & Gerd., Mycol. Mem. 5: 46. 1974.

= Endogone radiata Thaxt., Proc. Am. Acad. Arts & Sci. 57: 316. 1922.

⁼ *Endogone neglecta* Rodway, Proc. Roy. Soc. Tasmania 1917: 107. 1918.



FIGS 59–74. *Glomus* clade Ab3, with species virtually all belonging to the former genus *Sclerocystis*; form spores in highly organized, compact sporocarps, generally with > 50 spores per sporocarp; subtending hyphae (sh) generally short, bill-shaped or cylindrical, and concolorous with (or slightly lighter than) the spore wall. Spores species-specifically with (1–)2 wall layers (SWL1–SWL2) and wall thickening at spore base, but pore generally closed by a septum that regularly arises from the structural wall layer. **59–60**. *Gl. coremioides*. **61–63**. *Gl. clavisporum*. **64–65**. *Gl. badium*. **66–67**. *Gl. taiwanense*. **68–70**. *Gl. sinuosum*. **71–72**. *Gl. pachycaule*. **73–74**. *Gl. rubiforme*.

Glomus rubiforme (Gerd. & Trappe) R.T. Almeida & N.C. Schenck, Mycologia 82:

- 709. 1990.
- = Sclerocystis rubiformis Gerd. & Trappe, Mycol. Mem. 5: 60. 1974.
- = Sclerocystis indica Bhattacharjee & Mukerji, Acta Bot. Indica 8: 99. 1980.

Glomus segmentatum Trappe, Spooner & Ivory, Trans. Br. Mycol. Soc. 73: 362. 1979.

Glomus sinuosum (Gerd. & B.K. Bakshi) R.T. Almeida & N.C. Schenck, Mycologia 82: 710. 1990

= Sclerocystis sinuosa Gerd. & B.K. Bakshi, Trans. Br. Mycol. Soc. 66: 343. 1976.

= Sclerocystis pakistanica S.H. Iqbal & Perveen, Trans. Mycol. Soc. Japan 21: 59. 1980.

Glomus spinosum H.T. Hu, Mycotaxon 83: 160. 2002.

Glomus spinuliferum Sieverd. & Oehl, Mycotaxon 86: 158. 2003.

Glomus taiwanense (C.G. Wu & Z.C. Chen) R.T. Almeida & N.C. Schenck ex Y.J. Yao, Kew Bull. 50: 306. 1995.

= Sclerocystis taiwanensis C.G. Wu & Z.C. Chen, Trans. Mycol. Soc. Rep. China 2: 78. 1987.

- *Glomus tenebrosum* (Thaxt.) S.M. Berch. Can. J. Bot. 60: 2615. 1983. = *Endogone tenebrosa* Thaxt., Proc. Am. Acad. Arts Sci. 57: 314. 1922.
- *Glomus tenue* (Greenall) I.R. Hall, Trans. Br. Mycol. Soc. 68: 350. 1977. = *Rhizophagus tenuis* Greenall, New Zealand J. Bot. 1: 398. 1963.
- Glomus tortuosum N.C. Schenck & G.S. Sm., Mycologia 74: 83. 1982.

Glomus vesiculiferum (Thaxt.) Gerd. & Trappe, Mycol. Mem. 5: 49. 1974.

- = Endogone vesiculifera Thaxt., Proc. Am. Acad. Arts Sci. 57: 309. 1922.
- = Funneliformis vesiculifer (Thaxt.) C. Walker & A. Schüssler, The Glomeromycota – a species list: 14. 2010.

Glomus warcupii McGee, Trans. Br. Mycol. Soc. 87: 125. 1986.

Glomus zaozhuangianum F.Y. Wang & R.J. Liu, Mycosystema 21: 522. 2002.

Funneliformis C. Walker & A. Schüssler, emend. Oehl, G.A. Silva & Sieverd. (Glomus group Aa1)
FIGS

FIGS. 75-84

EMENDED DESCRIPTION: Spores formed within soil or rarely roots, singly or sometimes in sporocarps with a few to several spores per sporocarp only; the conspicuous sH is concolorous with spore wall color (or slightly lighter in color), sH is species-specific and generally funnel-shaped to cylindrical. Wall differentiation and pigmentation may continue over long distances from the spore base (often > $50-250 \mu m$), then mycelium may become hyaline. Pore regularly closed by a conspicuous septum that species-specifically arises from the structural wall layer, from an additional adherent innermost, (semi-)flexible lamina, or from both but not by introverted wall thickening, which is lacking. Forming typical vesicular-arbuscular mycorrhiza, with mycorrhizal structures that stain blue to dark blue in trypan blue.

GENDER: *Funneliformis* is a compound Latin adjective with a masculine/ feminine termination. Although under ICBN [Vienna Code] Article 62.3, the authors had the prerogative to choose either of these genders, their actual choice

of neuter gender was not an available option. We therefore choose to treat this genus as masculine (Article 62.3) and have made the appropriate corrections to all adjectival epithets.

- TYPE SPECIES: *Funneliformis mosseae* (T.H. Nicolson & Gerd.) C. Walker & A. Schüssler
- *Funneliformis caesaris* (Sieverd. & Oehl) Oehl, G.A. Silva & Sieverd., comb. nov. MYCOBANK MB 518445
 - = Glomus caesaris Sieverd. & Oehl, Mycotaxon 84: 381. 2002.
- *Funneliformis caledonius* (T.H. Nicolson & Gerd.) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 13. 2010.
 - = Endogone macrocarpa var. caledonia T.H. Nicolson & Gerd., Mycologia 60: 318. 1968.
 - = Glomus caledonium (T.H. Nicolson & Gerd.) Trappe & Gerd., Mycol. Mem. 5: 56. 1974.
- Funneliformis coronatus (Giovann.) C. Walker & A. Schüssler, The Glomeromycota

- a species list: 13. 2010.

- = *Glomus coronatum* Giovann., Can. J. Bot. 69: 162. 1990.
- *Funneliformis dimorphicus* (Boyetchko & J.P. Tewari) Oehl, G.A. Silva & Sieverd., comb. nov.

MycoBank MB 518447

- = Glomus dimorphicum Boyetchko & J.P. Tewari, Can J. Bot. 64: 90. 1986.
- *Funneliformis fragilistratus* (Skou & I. Jakobsen) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 13. 2010.
 - = Glomus fragilistratum Skou & I. Jakobsen, Mycotaxon 36: 276. 1989.
- *Funneliformis geosporus* (T.H. Nicolson & Gerd.) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 14. 2010.
 - = Endogone macrocarpa var. geospora T.H. Nicolson & Gerd. Mycologia 60: 318. 1968.
 - = Glomus macrocarpum var. geosporum (T.H. Nicolson & Gerd.) Gerd. & Trappe, Mycol. Mem. 5: 55. 1974.
 - = Glomus geosporum (T.H. Nicolson & Gerd.) C. Walker, Mycotaxon 15: 56. 1982.
- Funneliformis halonatus (S.L. Rose & Trappe) Oehl, G.A. Silva & Sieverd., comb. nov. MYCOBANK MB 518453
 - = Glomus halonatum S.L. Rose & Trappe, Mycotaxon 10: 413. 1980.
- *Funneliformis kerguelensis* (Dalpé & Strullu) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518454
 - = *Glomus kerguelense* Dalpé & Strullu, Mycotaxon 84: 53. 2002.
- *Funneliformis monosporus* (Gerd. & Trappe) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518456
 - = *Glomus monosporum* Gerd. & Trappe, Mycol. Mem. 5: 41. 1974.
- *Funneliformis mosseae* (T.H. Nicolson & Gerd.) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 13. 2010.
 - = Endogone mosseae T.H. Nicolson & Gerd., Mycologia 60: 314. 1968.
 - = Glomus mosseae (T.H. Nicolson & Gerd.) Gerd. & Trappe, Mycol. Mem. 5: 40. 1974.



FIGS 75–84. *Funneliformis* species (clade Aa1) generally form big (150–400 μm) spores, singly and (species-specifically) also in 'small' sporocarps with only a few to several spores per sporocarp; subtending hyphae (SH) are concolorous with (or slightly lighter than) the spore wall without introverted wall thickening at the spore base and are species-specifically cylindrical, funnel-shaped, or (rarely) slightly constricted. SH wall thickening may extend over long distances (> 100–300 μm). Spores with 1–5 wall layers (swL1–swL5); pore regularly closed by a conspicuous septum (sp) that species-specifically arises from the structural layer, from an adherent innermost, (semi-)flexible layer, or from both layers. 75. *Fu. mosseae.* 76. *Fu. monosporus.* 77. *Fu. fragilistratus.* 78. *Fu. coronatus.* 79. *Fu. multiforus.* 80. *Fu. geosporus.* 81. *Fu. verruculosus.* 82. *Fu. kerguelensis.* 83. *Fu. caledonius.* 84. *Fu. caesaris.*

Funneliformis multiforus (Tadych & Błaszk.) Oehl, G.A. Silva & Sieverd., comb. nov.

MycoBank MB 518459

= Glomus multiforum Tadych & Błaszk., Mycologia 89: 805. 1997.

Funneliformis verruculosus (Błaszk.) C. Walker & A. Schüssler, The Glomeromycota

- a species list: 14. 2010.

= Glomus verruculosum Błaszk., Mycologia 89: 809. 1997.

Simiglomus Sieverd., G.A. Silva & Oehl, gen. nov.

МусоВанк МВ 518435

FIGS. 85-86 (Glomus group Aa2)

Sporae singulariter vel in congregatos apertos cum sporis pauciaribus efformatae; tunica sporarum cum tunica hypharum coniuncta; porum sporarum apertum; hyphae cylindricae vel infundiles pori sporarum aperti, pauciores septa in distanica ex tunicae hyphalis in hypha; mycorrhizas vesicular-arbusculares formans caeruleas colorantes cum 'trypan blue'.

ETYMOLOGY: from the Latin: *simi(laris)* = similar; *glomus* = cluster; referring to the relation with *Glomus*, to which species of the new genus previously belonged.

Key Characters: Spores formed singly or in very loose, small clusters. Spores with a mono-to-multiple layered spore wall. Wall of the sH conspicuously continuous and concolorous with the spore wall, or slightly lighter in color than the spore wall. SH are funnel-shaped to cylindrical. Wall at spore attachment not with introverted wall thickening. Pore at spore base open but several septa in hyphae in some distance from spore base can separate spore contents from mycelia contents. Walls of sH thickened over very long distances from the spore base (up to > 1000 μ m). Forming typical vesicular-arbuscular mycorrhiza, with mycorrhizal structures that stain blue to dark blue in trypan blue.

TYPE SPECIES: Simiglomus hoi (S.M. Berch & Trappe) G.A. Silva et al.

Simiglomus hoi (S.M. Berch & Trappe) G.A. Silva, Oehl & Sieverd., comb. nov.

MycoBank MB 518461

= *Glomus hoi* S.M. Berch & Trappe, Mycologia 77: 654. 1985.



FIGS 85–86. *Simiglomus hoi* (clade Aa2). Spores formed singly or in very loose small clusters; the subtending hyphae (SH) are concolorous with (or slightly lighter than) the spore wall without introverted wall thickening at the spore base and generally cylindrical to slightly funnel-shaped or slightly constricted. SH wall thickening may extend over very long distances. The spore pore is generally closed by conspicuously recurved septa at varying distances from the spore base. Often several recurved septa (sp) visible that arise from the structural layer in the subtending hyphae.



FIGS 87–89. *Septoglomus* (clade Aa3) species form spores singly in soil or in very loose small clusters; subtending hyphae (sH) are concolorous with (or slightly lighter than) the spore wall without introverted wall thickening at the spore base and generally cylindrical to constricted. sH wall thickening may extend over long distances. Spores with 1–3 wall layers (swL1–swL3); if closed, pore often with one to several septa (sp) that species-specifically arise from the structural layer in the subtending hyphae. 87. *Se. constrictum.* 88. *Se. deserticola.* 89. *Se. xanthium.*

Septoglomus Sieverd., G.A. Silva & Oehl, gen. nov.

FIGS. 87–89 (Glomus group Aa3)

МусоВанк МВ 518436

Sporae singulariter vel in congregatos apertos efformatae; tunica sporarum cum tunica hypharum coniuncta; hyphae cylindricae vel infundiles pori sporarum occlusi septo. Mycorrhizas vesicular-arbusculares formans caeruleas colorantes cum 'trypan blue'.

ETYMOLOGY: from the Latin: *septum* = septum; *glomus* = cluster; referring to the relation with *Glomus*, to which species of the new genus previously belonged.

KEY CHARACTERS: Spores formed singly or in very loose, small clusters. Spores with a mono-to-multiple layered spore wall. Wall of the SH conspicuously continuous and concolorous with the spore wall, or slightly lighter in color than the spore wall. SH are cylindrical to constricted or slightly funnel shaped at spore base. Pore at spore base or in some distance from spore based closed by a septum. Forming typical vesicular-arbuscular mycorrhiza, with mycorrhizal structures that stain blue to dark blue in trypan blue.

TYPE SPECIES: Septoglomus constrictum (Trappe) Sieverd. et al.

Septoglomus africanum (Błaszk. & Kovács) Sieverd., G.A. Silva & Oehl, comb. nov. MycoBank MB 519466

- = Glomus africanum Błaszk. & Kovács, Mycologia 102: 1452. 2010.
- = Funneliformis africanus (Błaszk. & Kovács) C. Walker & A.
 - Schüssler, The Glomeromycota a species list: 13. 2010.

Septoglomus constrictum (Trappe) Sieverd., G.A. Silva & Oehl, comb. nov.

МусоВанк МВ 518462

- = *Glomus constrictum* Trappe, Mycotaxon 6: 361. 1977.
- = Funneliformis constrictus (Trappe) C. Walker & A. Schüssler, The Glomeromycota – a species list: 14. 2010.

- Septoglomus deserticola (Trappe, Bloss & J.A. Menge) G.A. Silva, Oehl & Sieverd., comb. nov.
 - MyCoBank MB 518463
 - = Glomus deserticola Trappe, Bloss & J.A. Menge, Mycotaxon 20: 123. 1984.
- Septoglomus xanthium (Błaszk., Blanke, Renker & Buscot) G.A. Silva, Oehl & Sieverd., comb. nov.

МусоВанк МВ 518464

- = Glomus xanthium Błaszk., Blanke, Renker & Buscot, Mycotaxon 90: 459. 2004.
- = Funneliformis xanthius (Błaszk.) C. Walker & A. Schüssler, The Glomeromycota – a species list: 14. 2010.

Claroideoglomeraceae C. Walker & A. Schüssler, emend. Oehl, G.A. Silva & Sieverd. (Glomus group B)

EMENDED DESCRIPTION: Spores formed singly or in clusters with low spore numbers, extremely rarely in sporocarps; in soils or rarely in roots; SH hyaline to white, rarely subhyaline, often conspicuously bill-shaped. Spores with 1–4 wall layers (SWL1–SWL4); pore closure at spore base often with a septum that species-specifically may arise from the structural layer, from an adherent inner, (semi-)flexible layer, or from both layers.

TYPE GENUS: *Claroideoglomus* C. Walker & A. Schüssler Other GENUS: *Viscospora* Sieverd. et al.

Claroideoglomus C. Walker & A. Schüssler, emend. Oehl, G.A. Silva & Sieverd. (Glomus group B1) FIGS. 90–98

EMENDED DESCRIPTION: Spores generally formed singly in soil or rarely in roots; SH hyaline to white, rarely subhyaline, conspicuously bill-shaped. Spores with 1–4 wall layers (SWL1–SWL4); pore closure at spore base often with a septum that species-specifically may arise from the structural layer, from an adherent innermost, (semi-)flexible layer, or from both layers.

TYPE SPECIES: *Claroideoglomus claroideum* (N.C. Schenck & G.S. Sm.) C. Walker & A. Schüssler

Claroideoglomus candidum (Furrazola, Kaonongbua & Bever), Oehl, G.A. Silva & Sieverd., **comb. nov.**

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МусоВанк МВ 519459
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- = Glomus candidum Furrazola, Kaonongbua & Bever, Mycotaxon 113: 103. 2010.
- *Claroideoglomus claroideum* (N.C. Schenck & G.S. Sm.) C. Walker & A. Schüssler, The *Glomeromycota* – a species list: 21. 2010.
 - = *Glomus claroideum* N.C. Schenck & G.S. Sm., Mycologia 74:

84. 1982 (emend. Walker & Vestberg 1998).

- *Claroideoglomus drummondii* (Błaszk. & Renker) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 22. 2010.
 - = Glomus drummondii Błaszk. & Renker, Mycol. Res. 110: 559. 2006.



FIGS 90–98. *Claroideoglomus* (clade B1) species generally form spores singly in soils or (rarely) in roots; rarely also in very loose spore clusters; subtending hyphae (SH) conspicuously bill-shaped but hyaline to white, rarely subhyaline. Spores with 1–4 wall layers (SWL1–SWL4); pore closure at spore base generally with a septum (sp) that species-specifically may arise from structural wall layer, an adherent inner (semi-)flexible layer, or both layers. **90–91**. *Cl. etunicatum*. **92**. *Cl. lamellosum*. **93–94**. *Cl. claroideum*. **95–96**. *Cl. luteum*. **97**. *Cl. drummondii*. **98**. *Cl. walkeri*.

- *Claroideoglomus etunicatum* (W.N. Becker & Gerd.) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 22. 2010.
 - = Glomus etunicatum W.N. Becker & Gerd., Mycotaxon 6: 29. 1977.
- Claroideoglomus lamellosum (Dalpé, Koske & Tews) C. Walker & A. Schüssler, The Glomeromycota a species list: 22. 2010.
 = Glomus lamellosum Dalpé, Koske & Tews, Mycotaxon 43: 289. 1992.
- Claroideoglomus luteum (L.J. Kenn., J.C. Stutz & J.B. Morton) C. Walker & A. Schüssler, The Glomeromycota a species list: 22. 2010.
 = Glomus luteum L.J. Kenn., J.C. Stutz & J.B. Morton, Mycologia 91: 1090. 1999.
- *Claroideoglomus walkeri* (Błaszk. & Renker) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 22. 2010.
 - = Glomus walkeri Błaszk. & Renker. Mycol. Res. 110: 563. 2006.

Viscospora Sieverd., Oehl & G.A. Silva, gen. nov.

МусоВанк МВ 518439

Sporae in congregatos apertos efformatae; tunica sporarum cum tunica hypharum coniuncta; hyphae hyalinae vel albae; stratum exterior hyphae et sporarum cum materia mucilagina; porum sporarum non occlusum vel occlusum hyphae crassae. Mycorrhizas vesicular-arbusculares formans caeruleas colorantes cum 'trypan blue'.

ETYMOLOGY: from the Latin: *viscosus* = sticky; *spora* = spore; referring to the adhesive nature of the spore surface of the type species of the genus.

KEY CHARACTERS: Spores generally formed in loose clusters; SH hyaline to white, rarely subhyaline, often thick-walled. Spores with 1–4 wall layers (SwL1–SwL4); outer wall layer exuding a mucigel-like substance. Pore closure at spore base often open, or semi-closed by wall thickening.

TYPE SPECIES: Viscospora viscosa (T.H Nicolson) Sieverd. et al.

Viscospora viscosa (T.H Nicolson) Sieverd., Oehl & G.A. Silva, comb. nov.

MycoBank MB 518471

= Glomus viscosum T.H. Nicolson, Mycol. Res. 99: 1502. 1995.

Diversisporaceae C. Walker & A. Schüssler, emend. Oehl, G.A. Silva & Sieverd. (Glomus group C)

EMENDED DESCRIPTION: Spore formation either diversisporoid or otosporoid. Diversisporoid spores either formed singularly, in clusters or in large sporocarps with high spores numbers, not organized; in pigmented spores SH conspicuously change color, becoming hyaline to white behind the septum, i.e. immediately or in a minimal distance to this septum; SH generally straight, cylindrical, sometimes species-specifically constricted or inflated. Spores with 1–3 wall layers (SWL1–SWL3); pore closure often with a septum that may species-specifically arise from innermost wall lamina or overlaying laminate layer, or from both; pore of hyphal attachment rarely (species-specifically) open. Otosporoid spores with two multiple-layered walls, spore attachment generally closed by a septum.

TYPE GENUS: Diversispora C. Walker & A. Schüssler

OTHER GENERA: Otospora Oehl et al., Redeckera C. Walker & A. Schüssler

Diversispora C. Walker & A. Schüssler, emend. G.A. Silva, Oehl & Sieverd. (Glomus group Ca) FIGS. 99–107

EMENDED DESCRIPTION: Spores either formed singularly, in small open clusters or in large multi-spored clusters or sporocarps where spores are not organized. In pigmented spores SH conspicuously change color, becoming hyaline to white behind the septum; SH generally straight, cylindrical, sometimes species-specifically constricted, often hyphal attachment looks like inserted in spore wall. Spores with 1–3 wall layers (SWL1–SWL3); pore closure often with a



FIGS 99–107. *Diversispora* (clade Ca) species form spores singly or in unorganized clusters or sporocarps; in pigmented spores subtending hyphae (SH) with a conspicuous color change distant to the first septum at or near the spore base; SH generally cylindrical, species-specifically sometimes constricted. Spores species-specifically with 1–3 wall layers (SwL1–SwL3); pore closure at spore base often with a septum (sp) that species-specifically may arise from a structural wall layer, an adherent inner (semi-)flexible layer, or both layers; rarely (species-specifically) pore open. 99. *Di. spurca.* 100. *Di. eburnea.* 101. *Di. insculpta.* 102. *Di. arenaria.* 103–104. *Di. aurantia.* 105. *Di. epigaea.* 106. *Di. tenera.* 107. *Di. pustulata.*

septum that may species-specifically arise from the innermost wall layer; rarely (species-specifically) pore of attachment open.

TYPE SPECIES: Diversispora spurca (C.M. Peiff. et al.) C. Walker & A. Schüssler

Diversispora arenaria (Błaszk., Tadych & Madej) Oehl, G.A. Silva & Sieverd., comb. nov.

МусоВанк МВ 518472

= Glomus arenarium Błaszk., Tadych & Madej, Acta Soc. Bot. Pol. 70: 97. 2001.

- *Diversispora aurantia* (Błaszk., Blanke, Renker & Buscot) C. Walker & A. Schüssler, – a species list: 43. 2010.
 - = Glomus aurantium Błaszk., Blanke, Renker & Buscot, Mycotaxon 90: 450. 2004.

Diversispora celata C. Walker, Gamper & A. Schuessler, New Phytol. 182: 497. 2009.

- *Diversispora eburnea* (L.J. Kenn., J.C. Stutz & J.B. Morton) C. Walker & A. Schüssler, The *Glomeromycota* – a species list: 43. 2010.
 - = Glomus eburneum L.J. Kenn., J.C. Stutz & J.B. Morton, Mycologia 91: 1084. 1999.
- *Diversispora epigaea* (B.A. Daniels & Trappe) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 43. 2010.
 - = Glomus epigaeum B.A. Daniels & Trappe, Can. J. Bot. 57: 540. 1979.
- Diversispora gibbosa (Błaszk.) Błaszk. & Kovács, comb. nov. МусоВанк MB 518475 = Glomus gibbosum Błaszk., Mycologia 89: 339. 1997.
- Diversispora insculpta (Błaszk.) Oehl, G.A. Silva & Sieverd., comb. nov.
 - MycoBank MB 519546
 - = Glomus insculptum Błaszk., Mycotaxon 89: 227. 2004.
- Diversispora przelewicensis (Błaszk.) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518477
 - = Glomus przelewicense Błaszk., Bull. Pol. Acad. Sci., Biol. Sci. 36: 272. 1988.
- *Diversispora pustulata* (Koske, Friese, C. Walker & Dalpé) Oehl, G.A. Silva & Sieverd., comb. nov.
 - МусоВанк МВ 518478
 - = Glomus pustulatum Koske, Friese, C. Walker & Dalpé, Mycotaxon 26: 143. 1986.
- Diversispora spurca (C.M. Pfeiff., C. Walker & Bloss) C. Walker & A. Schüssler, Mycol. Res. 108: 982. 2004.
 - = Glomus spurcum C.M. Pfeiff., C. Walker & Bloss, Mycotaxon 59: 374. 1996.
- Diversispora tenera (P.A. Tandy) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518479
 - = *Glomus tenerum* P.A. Tandy, Austral. J. Bot. 23: 864. 1975.
- *Diversispora trimurales* (Koske & Halvorson) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 43. 2010.
 - = Glomus trimurales Koske & Halvorson, Mycologia 81: 930. 1990.
- Diversispora versiformis (P. Karst.) Oehl, G.A. Silva & Sieverd., comb. nov.
 - MycoBank MB 518481
 - = Endogone versiformis P. Karst., Hedwigia 23: 39. 1884.
 - = Glomus versiforme (P. Karst.) S.M. Berch, Can. J. Bot. 61: 2614. 1983.
- Redeckera C. Walker & A. Schüssler, emend. Oehl, G.A. Silva & Sieverd. (Glomus group Cb) FIGS. 108–114

EMENDED DESCRIPTION: Spore formation disorganized in large and compact sporocarps, containing hundreds to thousands of spores per sporocarp; spores



FIGS 108–114. *Redeckera* (clade Cb) with spore formation in unorganized but large and compact sporocarps (usually with hundreds to thousands spores per sporocarp); spores species-specifically with 2(–3) wall layers (swL1–swL3); subtending hyphae (sH) generally broad at spore base and with a conspicuous thick, broad septum (sp) arising from the inner lamina of the generally bi-laminated spore wall layer (swL2); structural swL2 generally extend over very short (2–10 µm) distances into sH; swL1 fragile and usually inflating where swL2 becomes invisible in the sH. **108**. *Re. fulva.* **109**. *Re. megalocarpa.* **110**. *Re. pulvinata.* **111–112**. *Re. canadensis.* **113–114**. *Re. fragilis.*

with 2 to rarely 3 wall layers (swL1–swL3); sH generally broad at spore base and with a conspicuous, thick and broad septum that arises from the inner lamina (swL2) of the generally bi-laminated, structural wall layer; structural swL2 generally continue over very short distances (2–10 μ m) into sH; swL1 fragile, usually inflating in a short distance to the spore base where swL2 becomes invisible in the sH.

GENDER: *Redeckera* is a Latin feminine noun, formed according to ICBN [Vienna Code] Recommendation 60B.1(b). Because the feminine gender is clearly apparent (and reinforced by the implication of Recommendation 20A.1(i)), Article 62.3 does not apply, and the authors did not have the prerogative to choose any other gender. We therefore correct all adjectival epithets to feminine.

TYPE SPECIES: Redeckera megalocarpa (D. Redecker) C. Walker & A. Schüssler

Redeckera avelingiae (R.C. Sinclair) Oehl, G.A. Silva & Sieverd., comb. nov.

MycoBank MB 518482

= Glomus avelingiae R.C. Sinclair, Mycotaxon 74: 338. 2000.

Redeckera canadensis (Thaxt.) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518483

= Endogone canadensis Thaxt., Proc. Am. Acad. Arts Sci. 57: 317. 1922.

= Glomus canadense (Thaxt.) Trappe & Gerd., Mycol. Mem. 5: 59. 1974.

- Redeckera fragilis (Berk. & Broome) Oehl, G.A. Silva & Sieverd., comb. nov. MycoBank MB 518484
 - = Paurocotylis fragilis Berk. & Broome, J. Linn. Soc. Bot. 14: 137. 1873.
 - = Glomus fragile (Berk. & Broome) Trappe & Gerd., Mycol. Mem. 5: 59. 1974.
- *Redeckera fulva* (Berk. & Broome) C. Walker & A. Schüssler, The *Glomeromycota* a species list: 44. 2010.
 - = Paurocotylis fulva Berk. & Broome, J. Linn. Soc. Bot. 14: 137. 1873.
 - = Glomus fulvum (Berk. & Broome) Trappe & Gerd., Mycol. Mem. 5: 59. 1974.
- *Redeckera megalocarpa* (D. Redecker) C. Walker & A. Schüssler, The *Glomeromycota*: 44. 2010.
 - = Glomus megalocarpum D. Redecker, Mycol. Progress 6: 38. 2007.
- Redeckera pulvinata (Henn.) C. Walker & A. Schüssler, The Glomeromycota: 44. 2010.
 - = Endogone pulvinata Henn., Hedwigia 36: 212. 1897.
 - = Glomus pulvinatum (Henn.) Trappe & Gerd., Mycol. Mem. 5: 59. 1974.

Otospora Oehl, Palenz. & N. Ferrol

TYPE SPECIES: Otospora bareae Palenz. et al.

Otospora bareae Palenz., N. Ferrol & Oehl, Mycologia 100: 298. 2008.

Paraglomeraceae J.B. Morton & D. Redecker

Paraglomus J.B. Morton & D. Redecker

TYPE SPECIES: Paraglomus occultum (C. Walker) J.B. Morton & D. Redecker

Paraglomus albidum (C. Walker & L.H. Rhodes) Oehl, G.A. Silva & Sieverd., comb. nov.

MycoBank MB 518488

= *Glomus albidum* C. Walker & L.H. Rhodes, Mycotaxon 12: 509. 1981.

Paraglomus brasilianum (Spain & J. Miranda) J.B. Morton & D. Redecker, Mycologia 93: 190. 2001.

= *Glomus brasilianum* Spain & J. Miranda, Mycotaxon 60: 139. 1996.

Paraglomus laccatum (Błaszk.) Renker, Błaszk. & Buscot, Nova Hedwigia 84: 400. 2007.

= Glomus laccatum Błaszk., Bull. Pol. Acad. Sci., Biol. Sci. 36: 271. 1988.

Paraglomus lacteum (S.L. Rose & Trappe) Oehl, G.A. Silva & Sieverd. comb. nov. MycoBank MB 518489

= Glomus lacteum S.L. Rose & Trappe, Mycotaxon 10: 415. 1980.

Paraglomus occultum (C. Walker) J.B. Morton & D. Redecker, Mycol. Res. 93:190. 2001.

= *Glomus occultum* C. Walker, Mycotaxon 15: 50. 1982.

Taxonomic problems

We were unsure whether we had attributed *Gl. dolichosporum* and *Gl. citricola* to the correct genus, as the available descriptions were either short or in Chinese, images were missing or relatively poor, or the type specimen was not accessible. Moreover, in several species descriptions (some even very recent), the focus on the spore base and sH features was minimal to absent. For such cases, we decided to retain these species within *Glomus*, pending future revision. Placement of *Gl. pubescens* was similarly difficult: Already transferred to four different genera other than its basionym genus, this particular species may not even represent an AM fungus (e.g. Trappe, pers. comm.). Of all clades discussed here and fungal genera known, this species appears to fit best in *Glomus* group Ab1.

During our spore analyses, it was apparent that several genetic groups (clades) possess species with spores so similar that either the species are difficult to differentiate morphologically or may be conspecific. Examples include Aa3-Se. deserticola, Se. xanthium; Ab2-Gl. clarum, Gl. manihotis, Gl. zaozhuangianum; and Ca-Di. versiformis, Di. przelewicensis, Di. epigaea. As it was beyond the scope of this study to clarify all vagaries and possible taxonomic uncertainties at the species level, we have placed them into genera based on morphological characteristics, especially as many type specimens were unavailable (see TABLE 1). In some cases confusion may have resulted from misidentification or incorrect re-naming prior to genetic analysis. For example, the Gl. clarum isolate BR147B (supplied by Bioplanta) that grouped with Gl. manihotis in the initial phylogenetic study by Schüßler et al. (2001) was an ex-type of Gl. manihotis when sent in 1987 to Bioplanta (Sieverding pers. com.) but which some time later was relabeled as *Gl. clarum* by someone else. Not surprisingly, this particular 'Gl. clarum' isolate grouped with other Gl. manihotis ex-type material in the study by Schüßler et al. (2001). Consequently, genetic information provided for some species as well as the conclusion that other species are conspecific may be based on erroneous information.

Until recently, both germination and germination structures were regarded as important features to distinguish between AM fungal genera and families (e.g., *Pacisporaceae* vs. *Glomeraceae* by Oehl & Sieverding 2004, Walker & Schüßler 2004; among *Gigasporaceae*, *Scutellosporaceae*, *Racocetraceae*, and *Dentiscutataceae* by Oehl et al. 2008, 2011a; and *Archaeospora* vs. *Ambispora* by Spain et al. 2006). There are clear indications that *Paraglomus* species germinate directly through the spore wall as observed for *Pa. occultum*, *Pa. brasilianum* and *Pa. lacteum* (Spain & Miranda 1996, Rose & Trappe 1980, Oehl pers. obs.) in contrast to all other species forming single-walled glomoid or diversisporoid spores where a germ tube emerges apparently through the sH (e.g. *Glomus*,

Diversispora). There are uncertainties surrounding germ tube formation in some *Glomus* spp., such as *Gl. pallidum*, which Hall (1977) reported as germinating directly through the spore wall, although his description and photos suggest instead either intercalary spore formation or two hyphal attachments on the same spore. Intercalary spore formation (= two attachments) is a frequent feature for many *Glomus* sporocarpic (Ab) species.

Molecular difficulties

The major problem encountered during the current study revolved around the limited molecular database, especially for species of the former *Glomus* groups Ab1, Ab3, and Cb. Moreover, only a limited number of *Glomeromycota* species in the public databases have been sequenced for all currently available markers (LSU rDNA, SSU rDNA, ITS region, β -tubulin). Some individual clades were supported in just one dataset. Another major difficulty was that either many cultures of glomoid or diversisporoid spore-forming species are lost or yet to be cultured successfully in single species cultures or the species were not culturable. This renders analyses difficult or impossible. During the study, DNA extraction was attempted from several species using field spores or sporocarps but they largely failed (e.g., *Gl. invermaium, Gl. taiwanense*). These obstacles have left the database incomplete and slowed progress in complete molecular AM fungal identification.

Discussion

In this paper, we have revised the taxonomic organization of AM fungal species with glomoid spore formation, based largely on newly available molecular data retrieved from the literature and public databases. This organization is also based upon morphological spore characteristics, which were either already well established or newly established during this study. Phylogenetic trees support all genera proposed in this paper and some genera proposed by Walker & Schüßer (2010), with bootstrap support > 80% (except Septoglomus in the beta-tubulin phylogeny) in at least one analysis (NJ, ML, Bayesian or MP). However, Rhizophagus (Ab2) and Sclerocystis (Ab3) were not supported by our phylogenetic analyses and thus, the resurrection of these genera Walker & Schüßer (2010) cannot be accepted. The genera proposed or confirmed in the present study are shown as clusters in the trees: Glomus (Ab1, Ab2, Ab3), Funneliformis (Aa1), Simiglomus (Aa2), Septoglomus (Aa3), Claroideoglomus (B1), Viscospora (B2), Diversispora (Ca) and Redeckera (Cb). In TABLE 3, we summarize the new taxonomic organization of AM fungal species within the Glomeromycota.

There is no doubt that glomoid and diversisporoid spores in *Glomeraceae*, *Claroideoglomeraceae* and *Diversisporaceae* can clearly be differentiated by molecular phylogeny (FIGS. 1–4), as shown earlier (Schüßler et al. 2001, Silva

Class	Order	Family	Genus
Glomeromycetes	Glomerales	Glomeraceae	Glomus
			Funneliformis
			Simiglomus
			Septoglomus
		Claroideoglomeraceae	Claroideoglomus
			Viscospora
	Diversisporales	Diversisporaceae	Diversispora
			Redeckera
			Otospora
		Entrophosporaceae	Entrophospora
		Acaulosporaceae	Acaulospora
			Kuklospora
		Pacisporaceae	Pacispora
	Gigasporales	Gigasporaceae	Gigaspora
		Scutellosporaceae	Scutellospora
		-	Orbispora
		Racocetraceae	Racocetra
			Cetraspora
		Dentiscutataceae	Dentiscutata
			Fuscutata
			Quatunica
Archaeosporomycetes	Archaeosporales	Archaeosporaceae	Archaeospora
			Intraspora
		Ambisporaceae	Ambispora
		Geosiphonaceae	Geosiphon
Paraglomeromycetes	Paraglomerales	Paraglomeraceae	Paraglomus

TABLE 3. New classification of the Glomeromycota

The taxa Archaeosporomycetes, Paraglomeromycetes, Gigasporales and Orbispora are presented in this volume (Oehl et al. 2011a, 2011b).

et al. 2006, Stockinger et al. 2010). *Glomeraceae* and *Claroideoglomeraceae* differ morphologically in the hyphal attachments at the spore base and in spore clustering/aggregation or sporococarp formation. The phylogenetic trees also show *Glomeraceae* and *Claroideoglomeraceae* as separate clades with high bootstrap values in SSU, LSU, and beta-tubulin analyses.

Differences in hyphal connections at the spore base are also basic morphological indicators for other related but clearly genetically separate families of the *Glomeromycota* (e.g., *Acaulosporaceae*, *Pacisporaceae*, *Scutellosporaceae*). It has been argued earlier that unequivocal assignment of glomoid and diversisporoid spore-forming species to the correct major phylogenetic clade through morphological spore analyses alone would be impossible (e.g. Schüßler et al. 2001, Morton & Redecker 2001, Walker et al. 2007), primarily because glomoid and diversisporoid spores have but few differentiating morphological characters (which may yet hold true). Features other than hyphal connections (such as spore size and color) overlap between many glomoid and diversisporoid species, and intraspecific variation might be considerable. Spore aggregation and cluster/sporocarp formation in glomoid and diversisporoid species are not well investigated and have thus far received only limited attention. However, sporocarp formation and organization may be stable characteristics, as found in the former genus Sclerocycstis, which at some point may need to be raised again to genus level within the Glomeraceae. Other species of Glomeraceae are also always formed in sporocarps (e.g. Gl. glomerulatum, Gl. formosanum) but as yet there are insufficient molecular data to support their placement within their own genus. Other glomoid spores form in clusters or sporocarps occurring in relatively small to sometimes huge spore numbers (Gl. macrocarpum, Gl. microcarpum, Gl. aureum) as well as form singly in soil or in roots, so that spore aggregation alone cannot be used to diagnose the taxonomic order. We acknowledge that the large genus Glomus currently has morphologically heterogeneous spores, and recent findings by Błaszkowski et al. (2009a, b) show that several clades with small-spored species also belong to this group. To date, there are insufficient sequence sets available for more detailed analyses, although new genera could result from such analyses. This applies also for Gl. intraradices versus Gl. macrocarpum and Gl. microcarpum and Glomus Ab1 species in general. Recent research has also shown that genetic information generated years ago may actually be derived from another species, Gl. irregulare (Stockinger et al. 2009). Based on morphology alone, species such as Gl. intraradices should be excluded from Glomus.

Within *Diversispora* (respective former *Glomus* group C, our clade Ca) and *Redeckera* (our clade Cb), there has been no previous attempt to explain how these species differ morphologically from the former *Glomeraceae* groups A and B (Schüßler et al. 2001, Walker & Schüßler 2004). While morphological differentiation between *Diversispora* and *Redeckera* is possible, the morphology of *Diversispora* spores that differentiates them from spores of some *Claroideoglomus* spp. may be difficult to recognize, even for taxonomic specialists.

To conclude, our paper reports on the congruence of morphological groups (genera and family) with molecular phylogeny that has shown the need for a substantial reorganization of taxa in *Glomeromycota*.

Unsolved obstacles in the Glomeraceae

With approximately 70 species, the revised *Glomus* remains morphologically heterogeneous and the largest genus in the *Glomeromycota*. Species such as

Gl. intraradices, Gl. aggregatum, and *Gl. proliferum* are atypical in the genus regarding spore base closure by wall thickening and/or septum. These are normally quite stable features and not highly variable within species: a septum not readily seen at the spore base is either species specific or indicates the spore development stage. Mature spores have septa more frequently formed by the innermost lamina of the inner spore wall layer. Morphologically speaking, species like *Gl. intraradices, Gl. aggregatum,* and *Gl. proliferum* should be included in a separate genus, but the absence of sufficient genetic information (in particular of the other *Glomus* spp.) does not support this hypothesis. Additionally, we include several small-spored, cluster-forming species within the revised *Glomus* (e.g. *Gl. perpusillum, Gl. indicum, G. iranicum*) whose spores quickly decompose, suggesting that some earlier may have erroneously been designated 'non-sporulating AM fungi' (Błaszkowski et al. 2009b). Verification is still needed to establish whether there are indeed species of 'non-sporulating' fungi within the *Glomus* clade.

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Literature cited

Bentivenga SP, Morton JB. 1995. A monograph of the genus *Gigaspora*, incorporating developmental patterns of morphological characters. Mycologia 87(5): 719–731.

Błaszkowski J, Adamska I, Czerniawska B. 2004. Glomus insculptum, a new arbuscular mycorrhizal species from Poland. Mycotaxon 89: 225–234.

- Błaszkowski J, Renker C, Buscot F. 2006. Glomus drummondii and G. walkeri, two new species of arbuscular mycorrhizal fungi (Glomeromycota). Mycol. Res. 110: 555–566. doi:10.1016/ j.mycres.2006.02.006
- Błaszkowski J, Czerniawska B, Wubet T, Schäfer T, Buscot F, Renker C. 2008. Glomus irregulare, a new arbuscular mycorrhizal fungus in the Glomeromycota. Mycotaxon 106:247–267.
- Błaszkowski J, Ryszka P, Oehl F, Koegel S, Wiemken A, Kovács GM, Redecker D. 2009a. Glomus achrum and G. bistratum, two new species of arbuscular mycorrhizal fungi (Glomeromycota). Botany 87: 260–271. doi:10.1139/B08-138
- Błaszkowski J, Kovács GM, Balázs TK. 2009b. Glomus perpusillum, a new arbuscular mycorrhizal fungus. Mycologia 101(2): 247–255. doi: 10.3852/08-087
- Cavalier-Smith T. 1998. A revised six-kingdom system of life. Biol. Reviews 73: 203-266.
- Clements FE, Shear CL. 1931. The genera of fungi. Hafner Publishing Co., New York, USA.
- Daniell TJ, Husband R, Fitter AH, Young JPW. 2001. Molecular diversity of arbuscular mycorrhizal fungi colonising arable crops. FEMS Microbiol. Ecol. 36: 203–209.
- Gamper HA, Walker C, Schüßler A. 2009. Diversispora celata sp. nov.: molecular ecology and phylotaxonomy of an inconspicuous arbuscular mycorrhizal fungus. New Phytol. 182(2): 495-506. doi: 10.1111/j.1469-8137.2008.02750.x
- Guindon S, Gascuel O. 2003. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. System. Biol. 52(5): 696-704. doi: 10.1080/10635150390235520
- Hall IR. 1977. Species and mycorrhizal infections of New Zealand *Endogonaceae*. Trans. Br. Mycol. Soc. 68: 341–356.
- Hall IR, Abbott LK. 1979. Photographic slide collection illustrating features of the *Endogonaceae*. 3rd edition. Invery Agricultural Research Centre and Soil Science Department, University of Western Australia, p. 1–27, plus 400 color transparencies.
- Hall TA. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucl. Acids Symp. Ser. 41: 95–98.
- Krüger M, Stockinger H, Krüger C, Schüßler A. 2009. DNA-based species-level detection of arbuscular mycorrhizal fungi: one PCR primer set for all AMF. New Phytol. 183(1): 212–223. doi: 10.1111/j.1469-8137.2009.02835.x
- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, Thompson JD, Gibson TJ, Higgins DG. 2007. Clustal W and Clustal X version 2.0. Bioinformatics 23: 2947–2948. doi: 10.1093/bioinformatics/btm404
- Morton JB, Benny GL. 1990. Revised classification of arbuscular mycorrhizal fungi (*Zygomycetes*): a new order, *Glomales*, two new suborders, *Glomineae* and *Gigasporinae*, and two families, *Acaulosporaceae* and *Gigasporaceae*, with an emendation of *Glomaceae*. Mycotaxon 37: 471–491.
- Morton JB, Redecker D. 2001. Two new families of *Glomales, Archaeosporaceae* and *Paraglomaceae*, with two new genera *Archaeospora* and *Paraglomus*, based on concordant molecular and morphological characters. Mycologia 93(1): 181–195.
- Oehl F, Sieverding E. 2004. *Pacispora*, a new vesicular-arbuscular mycorrhizal fungal genus in the *Glomeromycetes*. J. Appl. Bot. Food Qual. 78(1): 72–82.
- Oehl F, Redecker D, Sieverding E. 2005. *Glomus badium*, a new sporocarpic arbuscular mycorrhizal fungal species from European grasslands of higher soil pH. J. Appl. Bot. Food Qual. 79(1): 38–43.
- Oehl F, Sýkorová Z, Redecker D, Wiemken A, Sieverding E. 2006. *Acaulospora alpina*, a new arbuscular mycorrhizal fungal species characteristic for high mountainous and alpine regions of the Swiss Alps. Mycologia 98(2): 286–294.

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- Oehl F, Souza FA de, Sieverding E. 2008. Revision of *Scutellospora* and description of five new genera and three new families in the arbuscular mycorrhiza-forming *Glomeromycetes*. Mycotaxon 106: 311–360.
- Oehl F, Silva DKA, Maia LC, Ferreira NM, da Silva GA. 2011a. Orbispora gen. nov., ancestral in the *Scutellosporaceae* of the *Glomeromycetes*. Mycotaxon 116: 161–169. doi: 10.5248/116.161
- Oehl F, Silva GA, Goto BT, Maia LC, Sieverding E. 2011b. *Glomeromycota*: two new classes and a new order. Mycotaxon 116: 365–379. doi: 10.5248/116.365
- Öpik MM, Moora M, Liira J, Zobel M. 2006. Composition of root-colonizing arbuscular mycorrhizal fungal communities in different ecosystems around the globe. J. Ecol. 94(4), 778–790. doi: 10.1111/j.1365-2745.2006.01136.x
- Palenzuela J, Ferrol N, Boller T, Azcón-Aquilar C, Oehl F. 2008. Otospora bareai, a new fungal species in the Glomeromycetes from a dolomitic shrub-land in the National Park of Sierra de Baza (Granada, Spain). Mycologia 100(2): 282–291.
- Palenzuela J, Barea JM, Ferrol N, Azcón-Aquilar C, Oehl F. 2010. Entrophospora nevadensis, a new arbuscular mycorrhizal fungus from the Sierra Nevada National Park (southeastern Spain). Mycologia 102(3): 624–632. doi: 10.3852/09-145
- Pirozynski KA, Dalpé Y. 1989. Geological history of the *Glomaceae* with particular reference to mycorrhizal symbiosis. Symbiosis 7(1): 1–36.
- Redecker D, Morton JB, Bruns TD. 2000a. Ancestral lineages of arbuscular mycorrhizal fungi (*Glomales*). Mol. Phylogenet. Evol. 14(2): 276–284.
- Redecker D, Morton JB, Bruns TD. 2000b. Molecular phylogeny of the arbuscular mycorrhizal fungi *Glomus sinuosum* and *Sclerocystis coremioides*. Mycologia 92(2): 282–285.
- Redecker D, Raab P, Oehl F, Camacho FJ, Courtecuisse E. 2007. A novel clade of sporocarp-forming glomeromycotan fungi in the *Diversisporales* lineage. Mycol. Progress 6(1): 35–44. doi: 10.1007/ s11557-007-0524-2
- Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19(12): 1572–1574. doi: 10.1093/bioinformatics/btg180.
- Rose SL, Trappe JM. 1980. Three new endomycorrhizal *Glomus* spp. associated with actinorrhizal shrubs. Mycotaxon 10: 413–420.
- Schwarzott D, Walker C, Schüßler A. 2001: Glomus, the largest genus of the arbuscular mycorrhizal fungi (Glomales), is non-monophyletic. Mol. Phylogenet. Evol. 21: 190–197.
- Schüßler A, Walker C. 2010. The *Glomeromycota*. A species list with new families and new genera. Gloucester, UK. 56 p.
- Schüßler A, Schwarzott D, Walker C. 2001. A new fungal phylum, the *Glomeromycota*: phylogeny and evolution. Mycol. Res. 105: 1413–1421.
- Sieverding E, Oehl F. 2006. Revision of *Entrophospora* and description of *Kuklospora* and *Intraspora*, two new genera in the arbuscular mycorrhizal *Glomeromycetes*. J. Appl. Bot. Food Qual. 80(1): 69–81.
- Silva GA da, Lumini E, Maia LC, Bonfante P, Bianciotto V. 2006. Phylogenetic analysis of *Glomeromycota* by partial LSU rDNA sequences. Mycorrhiza 16(3): 183–189. doi: 10.1007/ s00572-005-0030-9
- Simon L, Lalonde M, Bruns TD. 1992. Specific amplification of 18S fungal ribosomal genes from vesicular-arbuscular endomycorrhizal fungi colonizing roots. Appl. Environ. Microbiol. 58(1): 291–295.
- Souza FA de, Declerck S, Smit E, Kowalchuk GA. 2005. Morphological, ontogenetic and molecular characterization of *Scutellospora reticulata (Glomeromycota)*. Mycol. Res. 109: 697-706. doi: 10.1017/S09532756205002546

- Spain JL. 2003. Emendation of Archaeospora and of its type species, Archaeospora trappei Mvcotaxon 117: 109–112.
- Spain JL, Miranda JC. 1996. Glomus brasilianum: an ornamented species in the Glomaceae. Mycotaxon 60: 137–142.
- Spain JL, Sieverding E, Oehl F. 2006. Appendicispora, a new genus in the arbuscular mycorrhizalforming *Glomeromycetes*, with a discussion of the genus *Archaeospora*. Mycotaxon 97: 163–182.
- Stockinger H, Walker C, Schüßler A. 2009. 'Glomus intraradices DAOM197198', a model fungus in arbuscular mycorrhiza research, is not Glomus intraradices. New Phytol. 183(4): 1176-1187. doi: 10.1111/j.1469-8137.2009.02874.x
- Stockinger H, Krüger M, Schüßler A. 2010. DNA barcoding of arbuscular mycorrhizal fungi. New Phytol. 187(2): 461–474. doi: 10.1111/j.1469-8137.2010.03262.x
- Stürmer SL, Morton JB. 1997. Developmental patterns defining morphological characters in spores of four species in *Glomus*. Mycologia 89(1): 72–81.
- Swofford DL. 2003. PAUP*. Phylogenetic Analysis Using Parsimony (* and Other Methods), Version 4. Sinauer Associates, Sunderland, Massachusetts.
- Walker C. 1983. Taxonomic concepts in the *Endogonaceae*: spore wall characteristics in species descriptions. Mycotaxon 18: 443–455.
- Walker C. 2008. Ambispora and Ambisporaceae resurrected. Mycol. Res. 112: 297-298.
- Walker C, Schüßler A. 2004. Nomenclatural clarifications and new taxa in *Glomeromycota*. Mycol. Res. 108: 981–982.
- Walker C, Vestberg M. 1998. Synonymy amongst the arbuscular mycorrhizal fungi: Glomus claroideum, G. maculosum, G. multisubstensum and G. fistulosum. Ann. Bot. 82(5): 601–624.
- Walker C, Vestberg M, Demircik F, Stockinger H, Saito M, Sawaki H, Nishmura I, Schüßler A. 2007. Molecular phylogeny and new taxa in the Archaeosporales (Glomeromycota): Ambispora fennica gen. sp. nov., Ambisporaceae fam. nov., and emendation of Archaeospora and Archaeosporaceae. Mycol. Res. 111: 137–153. doi: 10.1016/j.mycres.2006.11.008