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1 **The virome of the arbuscular mycorrhizal fungus *Gigaspora margarita* reveals**
2 **the first report of DNA fragments corresponding to replicating non-retroviral**
3 **RNA viruses in Fungi**

4

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21 **Running title:** The virome of *Gigaspora margarita*

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26 **Originality-Significance statement:** this paper reports the first virome characterization of
27 *Gigaspora margarita*, an arbuscular mycorrhizal fungus already shown to harbor a complex
28 associated bacterial microbiota therefore adding a further layer of complexity to this organism;
29 furthermore, it reports for the first time the occurrence of cDNA fragments corresponding to
30 replicating non retroviral RNA viruses outside the phylum arthropoda, a discovery that, in those
31 organisms, has uncovered new anti-viral defence pathways.

32

33 **SUMMARY**

34 Arbuscular Mycorrhizal Fungi (AMF) are key components of the plant microbiota. AMF genetic
35 complexity is increased by the presence of endobacteria which live inside many species. A further
36 component of such complexity is the virome associated to AMF, whose knowledge is still very
37 limited. Here, by exploiting transcriptomic data we describe the virome of *Gigaspora margarita*. A
38 BLAST search for viral RNA-dependent RNA polymerases sequences allowed the identification of
39 four mitoviruses, one Ourmia-like narnavirus, one Giardia-like virus, and two sequences related to
40 *Fusarium* mycoviruses. Northern blot and RT-PCR confirmed the authenticity of all the sequences
41 with the exception of the *Fusarium*-related ones. All the mitoviruses are replicative and functional
42 since both positive strand and negative strand RNA are present. The abundance of the viral RNA
43 molecules was not regulated by the presence or absence of *Candidatus Glomeribacter*
44 *gigasporarum*, the endobacterium hosted by *G. margarita*, with the exception of the Ourmia-like
45 sequence which is absent in bacteria-cured spores. In addition, we report, for the first time, DNA
46 fragments corresponding to mitovirus sequences associated to the presence of viral RNA. These
47 sequences are integrated in neither the mitochondrial nor the nuclear genome, but likely exist as
48 extrachromosomal fragments.

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51 INTRODUCTION

52 The majority of land plants, including many crops and horticultural species, establish root
53 symbiotic interactions with a small group of soil fungi, the so called Arbuscular Mycorrhizal Fungi
54 (AMF), which belong to the Glomeromycotina (Spatafora *et al.*, 2016). As very ancient and
55 widespread in nature, AMF are key components of the plant microbiota. The benefits that AMF
56 provide to host plants, which include an improved mineral nutrition and an increased tolerance to
57 biotic and abiotic stresses, have raised the interest towards their exploitation as key components of
58 sustainable low input agricultural practices (Berruti *et al.*, 2016; Rodriguez and Sanders, 2015).

59 AMF obligate biotrophic nature and the lack of stable genetic transformation protocols have
60 made them a recalcitrant biological system to study. The recent publication of genomic data for
61 *Rhizophagus irregularis* (Tisserant *et al.*, 2013; Liu *et al.*, 2014; Ropars *et al.*, 2016) and gene
62 repertoires of other species such as *R. clarus* (Sędziewska Toro and Brachman, 2016), *Gigaspora*
63 *margarita* (Salvioli *et al.*, 2016) and *Gigaspora rosea* (Tang *et al.*, 2016) have shed light on the
64 biology and evolution of AMF (Kamel *et al.*, 2016). These findings have allowed to propose a new
65 phylogenetic classification (Spatafora *et al.*, 2016) and to infer the occurrence of a sexual cycle in
66 AMF (Corradi and Brachmann, 2017), for long considered clonal organisms. *R. irregularis*
67 possesses a large genome, compared to other fungi, with about 28,000 protein encoding genes and
68 rich in transposable elements (Tisserant *et al.*, 2013; Liu *et al.*, 2014). One striking feature is the
69 almost complete lack of glycoside hydrolases, a sign of the inability of the fungus to degrade plant
70 cell wall polysaccharides, possibly as a strategy to guarantee an intimate and compatible interaction.
71 The genome data have also been instrumental to describe a large repertoire of putative secreted
72 proteins to look for molecular tools exploited by the fungus to colonize the host plant (Kamel *et al.*,
73 2017). Very recently, new findings have shed light on their nature of these obligate biotrophs. The
74 lack of a fatty acid synthase gene in the genome of *R. irregularis* is mirrored by fatty acid
75 auxotrophy: fatty acids synthesized in the host plants are transferred to the fungus to sustain

76 mycorrhizal colonization (Jiang *et al.*, 2017; Bravo *et al.*, 2017; Luginbuehl *et al.*, 2017; Keymer *et*
77 *al.*, 2017).

78 The genetic complexity of AMF is also increased by the presence of endobacteria living
79 inside hyphae and spores of many AMF (Bonfante and Desirò, 2017). Two types of endobacteria
80 have been described in AMF: rod-shaped and Gram-negatives, associated to members of the
81 Gigasporaceae family (Ghignone *et al.*, 2012) and coccoid Mollicutes-related endobacteria,
82 distributed across different lineages of AM fungi (Naumann *et al.*, 2010; Desirò *et al.*, 2014; Torres
83 *et al.*, 2015). To date, only *Candidatus* Glomeribacter gigasporarum, the rod-shaped bacterium
84 hosted by *Gigaspora margarita* has been investigated in more detail. Thanks to the comparison
85 with a cured, endobacteria-free, strain (Lumini *et al.*, 2007) it has been demonstrated that the
86 endobacterium enhances fungal sporulation, bioenergetic capacity by priming mitochondrial
87 metabolic pathways and ability to detoxify reactive oxygen species possibly leading to an improved
88 ecological fitness of the fungus (Salvioli *et al.*, 2016; Vannini *et al.*, 2016).

89 All major fungal lineages were shown to possess additional genetic components among
90 which are viruses also called mycoviruses (Ghabrial *et al.*, 2015; Son *et al.*, 2015). In most cases the
91 genome of mycoviruses is an RNA molecule, mainly a double-stranded RNA (dsRNA) (Fauquet *et*
92 *al.*, 2005), containing at least a gene encoding an RNA-dependent RNA polymerase (RdRp), which
93 is necessary for replication (Ghabrial and Suzuki, 2009). Mycoviruses are transmitted horizontally
94 *via* hyphal anastomosis between vegetatively compatible individuals or transmitted vertically
95 through asexual and, less frequently, through sexual spores (Hillman *et al.*, 2004; Ghabrial *et al.*,
96 2015). External route of infection are apparently lacking; however, recent studies suggest that
97 mycophagous insects function as mycovirus transmission vectors in the case of ssDNA
98 mycoviruses (Liu *et al.*, 2016). The infection by mycoviruses is often asymptomatic, but in some
99 specific conditions alterations in the phenotype of the host fungus can be observed (van
100 Diepeningen *et al.*, 2006; Nerva *et al.*, 2017). In some cases higher order biological interactions are

101 also influenced; for example, the presence of a virus can lead to attenuation of virulence in a plant
102 pathogenic fungus (Choi and Nuss, 1992) or can provide the capability to enhance thermal tolerance
103 of the host plant to an endophytic fungus (Márquez *et al.*, 2007).

104 The knowledge of mycoviruses in mycorrhizal fungi is still very limited. In the
105 ectomycorrhizal fungus *Tuber aestivum*, three mycoviruses - a mitovirus (Stielow *et al.*, 2011b), a
106 totivirus (Stielow and Menzel, 2010) and an endornavirus (Stielow *et al.*, 2011a) - have been
107 described but their biological role was not characterized. Concerning AMF, Ikeda *et al.*, (2010)
108 demonstrated, for the first time, the presence of mycovirus-related dsRNAs, in the mycelium of
109 *Glomus sp.* (now named *Rhizophagus*) strain RF1. A 4.557 nucleotides segment, called GRF1V-M,
110 which could not be phylogenetically assigned to known genera of mycovirus, was characterized in
111 detail. The GRF1V-M encodes an RdRp and a structural protein. Remarkably, by subculturing
112 single spores, a fungal virus-free line was obtained: the absence of the virus led to the production of
113 a higher number of spores and to an enhanced stimulation of plant growth compared to the GRF1V-
114 M-positive line (Ikeda *et al.*, 2012). In a second work the same group has characterized by deep
115 sequencing of dsRNA in the AMF *Rhizophagus clarus* strain RF1, a sequence showing similarity to
116 RdRp of mitoviruses (Kitahara *et al.*, 2014). In this case a biological function has not been reported.

117 Here, by exploiting recently published transcriptomic data (Salvioli *et al.*, 2016) we describe
118 for the first time the components of the virome of the AMF *G. margarita*, which is phylogenetically
119 distantly related to *Rhizophagus* species. We provide *in silico* and experimental evidence of the
120 existence of a population of six viral sequences (4 mitoviruses, one Giardia-like and one Ourmia-
121 like virus). With one exception (the Ourmia-like virus), we found that the abundance of the viral
122 RNA molecules was not regulated by the presence of the endobacterium hosted by *G. margarita*.
123 Interestingly, we for the first time report cDNA fragments corresponding to mitovirus sequences
124 associated to the presence of their corresponding viral RNA: these sequences are integrated in
125 neither the mitochondrial nor the nuclear genome, but likely exist as extrachromosomal fragments.

126 RESULTS

127 Identification of viral sequences from *Gigaspora margarita* transcriptome

128 Contigs assembled from RNAseq data for the AMF *Gigaspora margarita* BEG34 (Salvioli
129 *et al.*, 2016) were searched for sequences annotated as RdRp; these are proteins performing
130 enzymatic activities typically encoded by RNA viral genomes, easily distinguishable from host
131 RdRp based on homology to other viral or host (fungal) RdRps. Eight sequences were retrieved and
132 a BLAST search allowed us to identify four mitoviruses, one ourmia-like virus, one Giardia-like
133 dsRNA virus, and two sequences related to Fusarium mycoviruses (Supporting information Table
134 S1 and Fig. S1).

135 The four putative mitoviral sequences and the Ourmia-like contig encode for a single ORF
136 for each genome, which, based on the presence of the conserved motifs GDD, can be annotated as
137 putative RdRps (Supporting information Fig. S1). The 5' UTRs present a length of 472 bp for
138 33086, 304 bp for 34036, 281 bp for 34770, 275 bp for 34875 and 268 bp for the Ourmia-like
139 contig. The ORFs encoding the RdRps end at 2854 bp for 33086, 3262 bp for 34036, 3311 bp for
140 34770, 3299 bp for 34875 and 3061 for the Ourmia-like virus. In the case of the Giardia-like, whose
141 genomic sequence predicts the presence of two ORFs, the first starts at nt 311 and ends at nt 1567
142 while the second one starts at nt 1611 and ends at nt 4679. Each ORF encodes a putative RdRp
143 protein of 794 aa for 33086, 986 aa for 34036, 1010 aa for 34770, 1008 aa for 34875 and 931 aa for
144 the Ourmia-like virus. The first ORF of the Giardia-like encodes a putative protein of 418 aa with
145 no similarity with proteins currently present in databases; the second ORF of 1022 aa shows
146 similarity to RdRps of another Giardia-like virus isolated in the AMF *Rhizophagus sp.* (Ikeda *et al.*,
147 2012) (Suppl. Table 1).

148 To confirm the authenticity (presence as biological entities, i.e. RNA molecules, and not as mere *in*
149 *silico* artifacts) of the sequences assembled from the transcriptome, RT-PCR assays were performed
150 using as template RNA from *G. margarita* germinating spores with specific primers (Supporting

151 information Table S2) designed to amplify a cDNA fragment in the size-range of about 500-700 bp.
152 A fragment of the expected size was obtained for each of the four mitoviruses and the Giardia-like
153 virus (Fig. 1). The sequencing of recombinant plasmids, after cloning the PCR fragments confirmed
154 the identity of the sequence of the PCR product with that predicted *in silico* (data not shown). More
155 primer pairs spanning different regions of the *in silico* predicted genomes were designed for the
156 remaining sequences (those that could not be confirmed in the first round of PCR, such as contig
157 23972, 39980 and 36178, the Fusarium micovirus and the Ourmia-like related sequences) and tested
158 again in RT-PCR assays. No amplified fragment was obtained for the 23972 and 39980 contigs, the
159 two Fusarium mycovirus-related sequences. Only for the Ourmia-like sequence (contig 36178) we
160 could confirm by RT-PCR and sequencing the presence of an RNA molecule corresponding to a
161 short portion (324 bp) identical to the virus RdRp predicted *in silico* (Supporting information Fig.
162 S2). Although the 36178 contig obtained *in silico* (Ourmia-like putative virus) encodes a long ORF,
163 the amplified segment correspond only to conserved RdRp motifs, whereas the rest of the sequence
164 does not show similarity to existing RdRps; several attempts to verify the presence of other cDNA
165 fragments corresponding to other regions of the Ourmia-like contig extending outside the conserved
166 GDD domain failed: we therefore hypothesized that the *in silico* assembled sequence may be a
167 chimera generated by an artifact of RNAseq assembly, and that only the conserved region we could
168 amplify is corresponding to a specific Ourmia-like RNA molecule.

169 To confirm the presence of an active RdRp encoded by the different identified sequences,
170 northern blot assays were carried out on *G. margarita* germinating spores using positive and
171 negative strand riboprobes. Northern blot allowed to detect genomic (+sense) RNA corresponding
172 to the four mitoviruses and the narnavirus-like sequence (Ourmia-like contig 36178) (Fig. 2-A-D),
173 but we could provide evidence for active replication only for the mitoviruses since both full length
174 positive strand and negative strand RNA were detected in RNA extracts; in addition, they seem
175 rather more abundant (Fig. 2 A-D) compared to the other virus tested. For the Ourmia-like sequence

176 only the positive sense-detecting probe gave a hybridization signal; no signal was obtained for the
177 negative sense-detecting probe probably due to the low abundance of the target sequence (Fig. 2-E),
178 close to detection limit of a northern blot assay.

179 No hybridization signal was detected for the Giardia-like sequence with both probes,
180 suggesting that the abundance of the RNA was below the threshold for northern detection in our
181 experiments (data not shown). The very low abundance of RNA corresponding to the Ourmia-like
182 virus and the Giardia-like virus was confirmed by qRT-PCR assays (see below).

183 **Phylogenetic placement of viral sequences**

184 We then further characterized our viral sequences with a phylogenetic analysis comparing
185 conserved regions of the RdRp, with those present in the databases. We first aligned the *G.*
186 *margarita* mitovirus and Ourmia-like virus to a dataset of RdRp that comprises the family
187 *Ourmiaviridae*, *Leviviridae*, *Narnaviridae* and a number of related viruses still not classified
188 taxonomically that resulted from NGS virome characterization (Shi *et al.*, 2016) The conserved
189 RdRp region among this wide dataset is limited to 109 amino acids surrounding the GDD catalytic
190 motif. This phylogenetic analysis allows to show that *G. margarita* putative mitoviruses are indeed
191 members of the *Mitovirus* genus, whereas the Ourmia-like fragment present in *G. margarita*
192 belongs to a clade of viruses still not classified taxonomically (Supporting information Fig. S3),
193 which we have previously proposed to form a new genus called Ourmia-like viruses (Turina *et al.*,
194 2017).

195 A further more defined phylogenetic analysis, that includes only mitoviruses, was carried
196 out on aligned sequences spanning 467 amino acids using a plant endogenized mitovirus as
197 outgroup (Fig. 3). This analysis showed that mitoviruses infecting *G. margarita* are
198 phylogenetically diverse, in particular the 33086 is distantly related to the other three sequences
199 which instead constitute a statistically well supported clade (34036, 34770 and 44875) that is
200 distinct from the other well supported mitovirus clades (Fig. 3). Indeed, contig 33086 shares the

201 highest RdRp protein identity score with *Botrytis cinerea* mitovirus (38.0%), while the best identity
202 value shared with the other *G. margarita* mitoviruses is 30.1% (33086 vs 34875). The other three
203 mitoviruses, instead, share higher percentage identity among each other, respectively from 32.0%
204 (34036 vs 34875) to 53.5% (34770 vs 34036) (Table 1 and Supporting information Table S3).
205 Interestingly, in all the 4 mitovirus sequences 100% of tryptophan (W) residues are encoded by the
206 TGG codon (Supporting information Fig. S1): therefore, a functional RpRd can be hypothetically
207 translated both in the cytosol and the mitochondria.

208 A phylogenetic tree for the Giardia-like viral sequence was also generated including the most
209 closely related members of dsRNA virus families (*Totiviridae* and *Partitiviridae*); as expected,
210 virus RdRPs belonging to the *Totiviridae* and *Partitiviridae* are grouped in statistically well
211 supported clades; the rest of the aligned sequences (including the *G. margarita* Giardia-like virus)
212 forms a clade with a relatively low statistical support value (39%) in bootstrap analysis (Fig. 4).

213 **Are the viral sequences endogenized into the *G. margarita* genome?**

214 A widespread endogenization of mitoviral sequences in plant genomes was recently reported
215 (Bruenn *et al.*, 2015) and this prompted us to look for possible genome endogenization events of the
216 viral sequences we identified in *G. margarita*.

217 We performed PCR assays on DNA extracted from *G. margarita* spores with the same specific
218 primers that were used to amplify the cDNA corresponding to the RNA (Supporting information
219 Table S1). In case of positive outcome, to confirm that the PCR products were indeed originated
220 from DNA, a control sample using DNase-treated genomic DNA was also analysed in parallel to
221 exclude any spurious activity of the DNA polymerase that might function as reverse transcriptase.
222 A PCR product of the expected size, which was absent in the DNase-treated sample, was obtained
223 for the four mitoviruses and for the Fusarium mycovirus 3-related 23972 sequence (Fig. 5). For
224 these five virus-like sequences a phenomena of genome endogenization can be therefore
225 hypothesized.

226 To further explore this issue we checked the sequence of the previously characterized *G. margarita*
227 mitochondrial genome (BEG34 strain, the same used in this study) (Pelin *et al.*, 2012): no viral
228 sequence was found within the complete mitochondrial DNA. We then tested the hypothesis that
229 these DNA sequences corresponding to viral RdRps could be endogenized into the nuclear genome.
230 A draft genome sequence of *G. margarita* (BEG 34 strain) is available (Ghignone, Venice, Salvioli,
231 Bonfante, unpublished). We mapped the genome sequencing reads on viral contigs by using BWA
232 software. The DNA sequence encoding the elongation factor, used as a positive control, was well
233 represented (590 paired end reads). We also had a confirmation of the genome endogenization for
234 the 23972 sequence (the Fusarivirus fragment) for which 164 paired end reads were found.
235 Surprisingly, coverage was zero for the 4 mitoviral sequences rejecting the hypothesis of a genomic
236 integration.

237 To investigate quantitatively the abundance of these viral DNA portions, we performed quantitative
238 PCR assays by using specific primers covering about 100-120 bp fragments of the four mitoviruses
239 (Supporting information Table S2) on three independent DNA preparations from 300 *G. margarita*
240 spores. As endogenous controls we used PCR primers targeting a portion of the ribosomal DNA
241 18S and of the elongation factor gene. Only the DNA corresponding to the mitovirus 33086
242 accumulated to a level statistically significantly different from the other three mitoviruses. The
243 amplicon corresponding to 33086 was the least abundant, while that of 34036 was the most
244 abundant but in all cases the amount of viral DNA was statistically significantly lower than that of
245 the DNA sequence encoding the elongation factor, which is supposed to be a single copy gene
246 (Table 2). Even with the more sensitive q-PCR assay we could not detect any DNA fragment
247 related to the Ourmia-like sequence (data not shown). We cloned and sequenced the DNA
248 fragments corresponding to the 34470 and 34875 mitoviruses, and confirmed that the sequence
249 fragments had the same sequence as the one assembled *in silico*, without gaps or any signs of
250 internal recombination (not shown).

251 To further clarify the nature of the DNA corresponding to the mitovirus sequences, we then
252 investigated whether i) DNA corresponding to the whole virus sequence could be amplified and ii)
253 the size of DNA amplified corresponded to genome fragments or to the complete viral sequence.
254 For this purpose a series of specific PCR primers was designed to span different portions of the
255 genome of the contig 34036 and to obtain amplicons of different size for the same mitoviral
256 sequence 34036 (Fig. 6): we choose the mitovirus 34036 because is the one with the highest DNA
257 titer. The primers were tested on two independent DNA extractions and on two independent cDNA
258 preparations from germinating spores.

259 We could amplify relatively abundant DNA corresponding to fragments spanning the 3' terminal
260 2400 bp, and only a much fainter band corresponding to the a fragment at the 5' of the genomic
261 sequence (Fig 6a). The consensus sequence obtained from a number of different clones amplified
262 from DNA corresponded exactly to that predicted from the *in silico* analysis, with only 1
263 synonymous nucleotide mutation (C to T) at position 1399 (Supporting information Fig. S4). When
264 we tested the maximum length of the amplified segment using different primer combination,
265 independently from the genome fragments where primers were designed, only amplicons below 700
266 bp in length could be amplified from DNA, while larger PCR products were obtained from cDNA
267 (Fig. 6b). This provides indirect evidence of the existence of small fragments of DNA
268 corresponding to the various regions of most of the viral genome.

269 **The mitoviral sequences are present in *G. margarita* B- (endobacteria free) but not in**
270 ***Gigaspora rosea* genomic DNA**

271 In order to investigate the presence of homologous viral sequences in other AMF, we performed
272 PCR experiments on genomic DNA from *Gigaspora rosea*, a phylogenetically related AMF which
273 does not host endobacteria, using the same *G. margarita* virome specific primers (Supporting
274 information Table S2). We also considered DNA of a *G. margarita* BEG34 isolate that has been
275 cured from the *Candidatus* Glomeribacter gigasporarum endosymbiotic bacteria (Lumini *et al.*,

276 2007) and has been called B- (endobacteria free). The quality of the DNA preparations was first
277 verified by PCR amplifications with AM-specific ribosomal primers AML1 and AML2 (data not
278 shown). A PCR product of the expected size was obtained from the B- isolate for all the analysed
279 viral sequences (4 mitoviruses and the Fusarium mycovirus 3-related sequence). The sequencing of
280 the PCR products confirmed the correspondence to the expected virus sequence (not shown). By
281 contrast, the *G. rosea* sample always gave negative results (Fig. 7). This suggests that these portions
282 of viral DNA are constantly present in *G. margarita* independently from the presence of
283 endobacteria, but are absent from the phylogenetically close *G. rosea*.

284 **Abundance of viral RNA molecules in B+ and B- germinating spores**

285 We then investigated whether the abundance of the viral RNA molecules was regulated by the
286 presence or absence of endobacteria. Quantitative RT-PCR reactions were set up on cDNA obtained
287 from B+ and B- germinating spores using the elongation factor as housekeeping gene. On average,
288 the mitoviral sequences led to a Ct (threshold cycle) between 11 and 14 (Ct =12 for 33086; Ct =14
289 for 34036; Ct = 11; 34470; Ct = 12 for 34875). For the Ourmia-like virus and the Giardia-like virus
290 sequences an average of 22 and 30 Ct was registered, respectively, confirming the lower abundance
291 compared to the mitoviral RNAs that was already noticed in the northern blot experiments.
292 Remarkably, no statistically significant different virus accumulation between B+ and B- spores was
293 observed for the mitoviral sequences and the Giardia-like virus (Fig. 8) but, surprisingly, the
294 Ourmia-like sequence seems to be present only in B+ spores: Ct values for this virus registered for
295 B- spores were below the detection threshold.

296

297 **DISCUSSION**

298 **A further layer of complexity in the organisms associated to *G. margarita***

299 The obligate biotrophism and the lack of stable genetic transformation protocols have made
300 AMF a complex biological system to study. Recent works, such as the description of the genome

301 sequence of *R. irregularis* (Tisserant *et al.*, 2013; Lin *et al.*, 2014), have increased our knowledge
302 on AMF genetic and genomic complexity, and shed light on the molecular determinants that play a
303 key role in the establishment of the symbiosis with the host plant. A further contribute to this
304 complexity is given by the presence of endobacteria (Bonfante and Desirò, 2017), whose biological
305 functions, with a single noticeable exception (Salvioli *et al.*, 2016), are still largely unknown.
306 A few studies recently revealed that a further level of complexity is given by the presence of
307 mycoviruses, that are widespread in all major taxa of fungi. Mycoviruses in AMF have possibly
308 evolved under unique selection pressures. In one case only it has been demonstrated that they are a
309 biologically active component of the symbiosis: a *Glomus (Rhizophagus)* strain free of the GRF1V-
310 M virus (a virus related to the Giardia-like virus described in this work) produced a higher amount
311 of spores and promoted plant growth more efficiently than the fungal line containing the virus
312 (Ikeda *et al.*, 2012). Similarly a fungal strain of *Mortierella elongata* cured from its endobacteria
313 produces higher spore numbers and grows better (Li *et al.*, 2017). Nevertheless, virus and
314 endobacterial association with AMF seem to be fairly common in nature, raising the possibility of
315 some ecological advantage difficult to measure in the laboratory experimental conditions, where
316 most of the competition aspects are not under scrutiny. Despite the above-mentioned major
317 acquisition, the knowledge of mycoviruses biological role in AMF is still limited.

318 Recently published RNAseq data of the AMF *G. margarita* (Salvioli *et al.*, 2016) allowed us
319 to identify through computational analysis, four putative mitoviral sequences, a putative Ourmia-
320 like mycovirus, a putative Giardia-like dsRNA virus and two sequences related to Fusarium
321 mycoviruses. The existence as RNA molecules of *in silico* predicted sequences was confirmed by
322 RT-PCR experiments, for the 4 mitoviral sequences, and the Ourmia-like and Giardia-like
323 sequences. We could not confirm the sequence related to Fusarium mycovirus 1 while that related
324 to Fusarium mycovirus 3 turned out to be corresponding to a stably endogenized viral fragment into
325 the genome of *G. margarita* likely not sufficiently transcribed to be detected by RT-PCR.

326 We could confirm the presence of the Ourmia-like virus RNA by RT-PCR assays only based on a
327 short sequence of RNA molecule corresponding to a small portion of the sequence (Supporting
328 information Fig. S2) which encodes for an RdRp conserved domain. We hypothesize that the whole
329 3.2 kb contig is probably a result of a misassembly. It is worth to note that the contig is rich in AT
330 stretches which often give many difficulties during the *in silico* assembly process. Nevertheless, the
331 small segment corresponding to the part amplified by RT-PCR is indeed part of a self replicating
332 RNA molecule, since no DNA corresponding to this fragment was detected, excluding
333 endogenization events. Moreover, we demonstrated that the four mitoviruses are replicative and
334 their RdRP is active since both positive and negative sense RNA strands were detected in northern
335 blot assays. Regarding the Ourmia-like fragment, we obtained a signal only from the positive probe
336 while no signal was detected neither by the positive nor by the negative probe for the Giardia-like,
337 probably due to low RNA abundance. The size of the Ourmia-like RNA detected with the antisense
338 probe (detecting + sense) is approximately 3 kb: therefore indeed an RNA molecule larger than the
339 small fragment we could amplify by RT-PCR exists.

340 Since many AMF are characterized by the presence of endobacteria which live inside hyphae and
341 spores, and it has been demonstrated that their presence may have an important role for the fungal
342 fitness (Lumini *et al.*, 2007; Salvioli *et al.*, 2016; Vannini *et al.*, 2016), we investigated whether the
343 presence or absence of endobacteria could affect the occurrence and abundance of viral RNA in *G.*
344 *margarita*. At least in germinating spores, the presence or absence of the endobacteria does not
345 affect the occurrence and the abundance of viral RNA molecules, with the exception of the Ourmia-
346 like virus that seems to be present only in B⁺ spores. This result opens the possibility that the
347 replication of this virus is dependent on endobacteria likely inside the endobacteria itself. It can be
348 also hypothesized that this sequence is associated to endobacteria; indeed a number of viruses
349 phylogenetically related to Ourmia-like viruses are known phages (family *Leviviridae*) (Dolja and
350 Koonin, 2012).

351 **Phylogenetic analysis of the *G. margarita* virome showed both evidence of virus-host co-**
352 **evolution and of horizontal gene transfer**

353 Since viral RNA-dependent RNA polymerases (RdRp) are markers of mycoviruses with an RNA
354 genome, we carried out a phylogenetic analysis considering the conserved catalytic domain of the
355 identified RdRps. This first analysis allowed us to associate the four mitoviruses to the genus
356 *Mitovirus* and the Ourmia-like fragment to a new clade called Ourmia-like group that was recently
357 shown to exist after a wide NGS (Next Generation Sequencing) analysis of invertebrate virome (Shi
358 *et al.*, 2016).

359 A further analysis allowed us to focus on the diversity of the four mitoviruses; in particular,
360 three of them cluster together (showing evidence of co-evolution with their host) while the 33086
361 sequence is separated and more closely related to mitoviruses from a phylogenetically distant host
362 (*Botrytis cinerea*), providing some indirect evidence of possible horizontal virus transmission.
363 Taxonomically, the criteria to establish a new mitovirus species requires the identity in the whole
364 aligned RdRp to be below 40% (Hillman and Esteban, 2011); when we first checked the best
365 sequences match retrieved by BLAST, none of them shared more than 40% identity, implying that
366 the four mitoviruses constitute new mitovirus species. When we performed a pairwise alignment
367 among the *G. margarita* mitovirus pairs using the Needleman-Wunsch Global Align Protein
368 Sequences software, inside the BLAST suite, none of the mitovirus pairs shared an identity
369 percentage over 40% (39% being the highest, between contig 34470 and 34875) (Table 1).
370 Therefore we propose that each of the mitovirus sequence we detected belongs to a new mitovirus
371 species, respectively called *Gigaspora margarita* mitovirus 1 (GmMV1), GmMV2, GmMV3 and
372 GmMV4.

373 The Giardia-like sequence also constitutes a new viral species called *G. margarita* Giardia-like
374 virus 1 (GmGLV1). Given that in the case of the Ourmia-like sequence only a small fragment was
375 confirmed by RT-PCR, we think that the data is overall too preliminary to establish a new virus

376 species. The phylogenetic inference of the Giardia-like sequence shows that RdRp belonging to
377 *Totiviridae* and *Partitiviridae* are grouped in a clade statistically supported while the other aligned
378 sequences, including Giardia-like, show low RdRp sequences identities (39%) and still do not form
379 a statistically well supported clade.

380 **The *G. margarita* mitoviruses display the same TGG codon frequency for tryptophan as the**
381 **mitochondrial genes**

382 The Mitovirus genus includes fungal viruses with small RNA genomes with a single ORF encoding
383 a RdRp; most mitoviruses replicate in their host's mitochondria (Cole *et al.*, 2000; Hillman and Cai,
384 2013; Wu *et al.*, 2016). For translation, mitoviruses relies on the endogenous mitochondrial
385 translational code which can use for tryptophan (Trp) the codons TGG (as in the cytosolic/nuclear
386 genetic code) and TGA (which, by contrast, serves as a stop codon in the standard cytosolic/nuclear
387 genetic code). Interestingly, in the four *G. margarita* mitoviral sequences all the Trp residues are
388 encoded by the TGG codon. This observation indicates that, apparently, functional RdRps from the
389 four *G. margarita* mitoviruses can be translated both in cytosol and mitochondria. The use of TGG,
390 compatible with both cytoplasmic and mitochondrial translation, is a feature of a few mitoviruses,
391 including RcmV1-RF1, the only mitovirus described so far in an AMF, that is *Rhizophagus clarus*
392 (Kitahara *et al.*, 2014). It has been hypothesized that this feature might be an advantage for
393 horizontal transmission among AMF (Kithara *et al.*, 2014). However, it has been recently observed
394 that host fungi whose mitoviruses have no or few TGA codons are distinctive in also having no or
395 few TGA codons in their core mitochondrial genes. For example, *G. margarita* mitochondrial DNA
396 has only 2% Trp encoded by TGA and similar low percentages occur in the mitochondrial genomes
397 of other AMF (Nibert, 2017). Thus, the exclusion of such codons in some mitoviruses appears to
398 reflect most fundamentally that TGA is a rare mitochondrial codon in their particular hosts.
399 Whether the four *G. margarita* mitoviruses are capable to replicate also in the cytosol is still an
400 open question and needs to be experimentally proved.

401 **First report of cDNA corresponding to replicating non retroviral RNA viruses present in**
402 **hosts other than diptera**

403 Since integration events of fungal mitovirus cDNA in the mitochondrial (and to a minor extent
404 nuclear) DNA of vascular plants was shown to be a fairly common event (Bruenn *et al.*, 2015), we
405 investigated whether genome integration occurred also for the viral sequences identified in *G.*
406 *margarita*. Our PCR assays revealed the presence of DNA templates corresponding to the four
407 mitoviruses and to Fusarium mycovirus 3-related sequence (23972), but when we checked the
408 published sequence of the mitochondrial genome of *G. margarita* we didn't find any viral sequence.
409 The presence of multiple viral sequences in the same individual organism is not uncommon for
410 mycoviruses (Nerva *et al.*, 2016; Kondo *et al.*, 2013; Xie and Gahbrial, 2012; Jiang *et al.*, 2013),
411 even in the case of closely related viruses as for the 4 *G. margarita* mitovirus species. It would be
412 interesting to assess the exact cellular niche for each mitovirus and whether they all infect each
413 mitochondrion (mixed infection) or whether one mitovirus species for each mitochondria exists,
414 contributing to a differentiation among the mitochondrial identities inside a single organism.

415 We had also the possibility to check for integration of those mitoviral DNA templates in the
416 genomic DNA, looking into reads from an ongoing sequencing project of the same *G. margarita*
417 isolate: we could map reads from the endogenized Fusarium-like virus fragment, but no evidence of
418 endogenized mitovirus sequence could be obtained. The nature of the amplified DNA for mitoviral
419 sequences remains therefore undefined. Our attempts to perform Southern blot assays failed in
420 detecting specific hybridization signals related to mitovirus endogenization (data not shown). Taken
421 as a whole, these data suggest that the mitoviral DNA sequences are not integrated in the genome
422 but they likely exist as extrachromosomal fragments of relatively small size as it can be deduced by
423 our different amplicon length amplification PCR experiments. This is the first time that viral DNA
424 has been detected in the presence of the corresponding homologous replicant RNA for mitoviruses.
425 The presence of genome endogenization in plants has been so far associated with the absence of

426 replicative capability of mitoviruses (Koonin *et al.*, 2014). To our knowledge this is also the first
427 time that cDNA corresponding to a non retroviral replicating RNA virus is detected outside the
428 phylum arthropoda. In fact a few reports have pointed to the existence of episomic cDNA of RNA
429 viruses in the order Diptera (Nag *et al.*, 2016; Goic *et al.*, 2013; Goic *et al.*, 2016). Recently, it was
430 shown that two species of *Aedes* mosquitoes infected with two arboviruses from distinct families
431 (dengue or chikungunya) generate, by endogenous reverse transcriptase activity, a viral-derived
432 DNA that is essential for mosquito survival and viral tolerance, being at the base of persistent viral
433 infections (Goic *et al.*, 2016); inhibition of cDNA synthesis results in higher mortality without
434 affecting the RNAi anti-viral system. In our system, we have shown that cDNA sequences
435 correspond exactly to viral genomic sequences without signs of re-arrangement as it is often the
436 case in the *Drosophila melanogaster* viral infection system (Goic *et al.*, 2013). A possible analogy
437 between the two system is that in mosquito it was shown that cDNA was associated to persistent
438 infection and our virus-fungal system also has the hallmarks of persistent infections (lack of external
439 infectivity of mitoviruses, widespread association between these viruses and AMF hosts, lack of
440 obvious detrimental effects on the host, presence of a relatively high virus titer); it is tempting to
441 speculate that the same anti-viral mechanism bringing to persistent infections is common to such
442 distantly related hosts. Furthermore, it would be interesting to identify the mechanisms by which
443 the mitoviral DNAs are generated in *G. margarita* and whether they may have an anti-viral
444 functional role as is the case of mosquito infecting virus *via* the piRNA pathway (reviewed in Olson
445 and Bonizzoni, 2017). Given the large amount of retrotransposon sequences present in the *G.*
446 *margarita* genome (Ghignone, Venice, Anselem, Salvioli, Bonfante, unpublished), we can envision
447 retro-transcriptase activity that can occasionally use as template RNA mitovirus sequences; the
448 presence of such retroviral activity in mitochondria is, to our knowledge, unknown and should be
449 further tested.

450 In conclusions, the augmented genome concept that scientists often use to describe human beings at
451 the organismal level, including the whole microbiota genomes, can be in part applied to the
452 fascinating biological system of AMF, where a fungus, its associated bacteria, and the associated
453 viruses seem to constitute an interacting superorganism; understanding the biological relevance of
454 each of these components and how they interact will be the challenging task to be pursued in the
455 near future.

456

457 **EXPERIMENTAL PROCEDURES**

458 **Biological material**

459 Spores of *Gigaspora margarita* Becker and Hall (BEG 34), the corresponding cured strain (without
460 the endobacterium *Candidatus Glomeribacter gigasporarum*; Lumini *et al.*, 2007) and *Gigaspora*
461 *rosea* (BEG 9) were propagated by using white clover (*Trifolium repens*) as trap plant. Clover
462 plants were inoculated with ca. 100 spores and after 2-3 months new spores were generated and
463 collected by the wet sieving technique. To generate germinating spores, *G. margarita* spore
464 suspensions were divided into aliquots of 100, surface sterilized twice for 10 minutes with 3%
465 chloramine-T and 0.03% streptomycin sulfate, rinsed several times with sterile distilled water and
466 then incubated in 1 ml of sterile distilled water for 5-7 days in the dark at 30°C. Germinated spores
467 were collected, immediately frozen in liquid nitrogen and stored at -80°C.

468 **Total nucleic acids extraction**

469 About 200-300 spores were crushed by a pestel using 1 ml of lysis CTAB buffer containing 2%
470 cetyl trimethylammonium bromide, 1% polyvinyl pyrrolidone, 100 mM Tris-HCl, 1.4 M NaCl, 20
471 mM EDTA. After incubation at 65°C for 15 min, samples were centrifuged at 10,000 rpm for 10
472 min. The supernatant was added with an equal volume of phenol-chlorophorm-isoamyl alcohol
473 (25:24:1; vol:vol:vol) and mixed. The upper aqueous phase of a centrifugation at 10,000 rpm for
474 10 min was transferred into a new tube to which an equal volume of chlorophorm was added. The

475 aqueous phase was then collected and precipitated with 2/3 vol of isopropanol. After an incubation
476 at 4°C for 2 hrs, samples were centrifugated at 14,000 for 30 min at 4°C. The pellet was washed
477 with 80% ethanol and resuspended into 50 µl of sterile water. To confirm that the PCR products
478 were indeed originated from DNA, a control sample was generated by a DNase treatment
479 performed on 1 µg of total nucleic acids for 1 h at 37°C using DNase (Ambion) according to
480 manufactures's instructions.

481 **PCR and cloning**

482 PCR reactions were set up following standards procedures. For sequencing the high fidelity Phusion
483 DNA polymerase (ThermoFisher) was used. Primers are shown in Supporting information Table
484 S2. PCR products were cloned into the TOPO Vector (Invitrogen) vector following manufactures's
485 instructions. Recombinant DNAs were extracted with QIAGEN plasmid Minikit and sequences
486 were obtained from the Sequencing Service, LMU Biozentrum, Großhaderner, Germany). Sequence
487 analyses were performed with CHROMAS LITE
488 (http://www.technelysium.com.au/chromas_lite.html).

489 **RNA extractions and RT-PCR**

490 Total RNA was extracted using the RNeasy Microarray Tissue Mini Kit (Qiagen, Hilden,
491 Germany), according to the manufacturer's instructions. The concentration and quality of the
492 nucleic acids were assessed with a Nanodrop1000 (Thermo Scientific, Wilmington, NC, USA).
493 Samples were treated with TURBO DNase (Ambion) according to the manufacturer's instructions.
494 The RNA samples were routinely checked for DNA contamination by means of RT-PCR analysis,
495 using primers Efgig2F 5'-TGAACCTCCAACCAGACCAACTG-3' and EfgigR 5'-
496 CGGTTTCAACACGACCTACAGGGAC-3' for *G. margarita* translation elongation factor
497 (Efgig, Salvioli *et al.*, 2014) and the One-Step RT-PCR kit (Qiagen). For single-strand cDNA
498 synthesis samples were denatured at 65°C for 5 min and then reverse-transcribed at 25°C for 10
499 min, 42°C for 50 min and 70° for 15 min in a final volume of 20 µl containing 10 µM random

500 hexamers, 0.5 mM dNTPs, 4 μ l 5X buffer, 2 μ l 0.1 M DTT, and 1 μ l Super-ScriptII (Invitrogen).
501 Quantitative RT-PCR (qRT-PCR) experiments were carried out in a final volume of 10 μ l
502 containing 5 μ l of iTaq™ Universal SYBR® Green Supermix (Bio-Rad), 0.2 μ l of 2 μ M specific
503 primers (Table 2), and about 20 ng of cDNA. Samples were run in the iCycler iQ apparatus (Bio-
504 Rad) using the following program: 3 min pre-incubation at 95 °C, followed by 40 cycles of 10 s at
505 95 °C, and 30 sec at 60 °C. Each amplification was followed by melting curve analysis (60 °C to 94
506 °C) with a heating rate of 0.5 °C every 15 s. All reactions were performed on at least three
507 biological and three technical replicates and only Ct values with a standard deviation that did not
508 exceed 0.3 were considered. The comparative threshold cycle method (Rasmussen, 2001) was used
509 to calculate relative expression levels using as a reference gene for transcript normalization the *G.*
510 *margarita* elongation factor (Efgig). Statistical analyses were performed through one-way analysis
511 of variance (one-way ANOVA) and Tukey's post hoc test, using a probability level of $p < 0.05$. All
512 statistical analyses were performed using the PAST statistical package (version 2.16; Hammer *et*
513 *al.*, 2001).

514 **Northern blot**

515 For northern blot analyses, total RNA from fungi was prepared using Total Spectrum RNA Reagent
516 (Sigma-Aldrich, Saint Louis, MO, USA) as suggested by the manufacturer. RNA samples were
517 separated in gel electrophoresis under denaturing conditions using glyoxal in HEPES-EDTA buffer
518 as detailed in Sambrook *et al.* (1989). Hybridization were performed using a radio-labeled RNA
519 probe prepared from EcoRI linearized purified plasmids containing cDNA fragments corresponding
520 to both orientation (sense and antisense probes) of each of the 6 viruses through T7 transcription
521 using the Maxiscript T7 kit reagents (Thermo Fisher Scientific Inc., Waltham, MA, USA), as
522 detailed before (Nerva *et al.*, 2017). In some cases, some fragments in specific orientations were
523 toxic to *E. coli* after transformation, and recombinant plasmid could not be obtained. The problem
524 was circumvented using as template for transcription, instead of a linearized plasmid, a PCR

525 fragment amplified with T7 promoter primer, and a fragment specific reverse primer directly
526 amplified from the ligation reaction.

527 **Phylogenetic analyses**

528 We identified groups of conserved sequences related to the different mycoviruses present in *G.*
529 *margarita* through BLAST searches of the databases (Accession numbers reported in Supporting
530 information Table S4) and the representative protein sequences identified were used for multiple
531 sequence alignments using MUSCLE (Edgar, 2004) implemented in the MEGA 6 (Tamura *et al.*
532 2013). Aligned sequences were used to infer phylogenetic trees using the Maximum Likelihood
533 method based on the Le_Gascuel_2008 model (Le and Gascuel 2008). The best amino acid
534 substitution model was calculated with MEGA 6. Statistical analysis was carried out through
535 bootstrap analysis with 1000 replicates. Further details of the phylogenetic analysis are included in
536 figure legends. Multiple aligned sequences were also used to calculate pairwise identity and
537 similarity percentages using MatGat (Campanella *et al.*, 2003).

538 **Bioinformatic analyses**

539 *G. margarita* BEG34 genomic DNA libraries (PE, MP-3kb, MP-8kb), currently used for an ongoing
540 genome assembly project, were first checked for quality with FASTQC (Andrews, 2010) and then
541 trimmed with TRIM GALORE! (Krueger, 2012). Cleaned PE reads were mapped onto *G.*
542 *margarita* transcript comp11141_c0_seq1 (GBYF01010162.1), coding for Translation Elongation
543 Factor EF-1 alpha, and onto putative viral sequences (33086, 34036, 34470, 34875) using BWA (Li
544 and Durbin, 2009). Mapping outputs were handled and analyzed with SAMTOOLS (Li *et al.*,
545 2009). Clues of endogenization of the putative *Fusarium graminearum* dsRNA mycovirus-3-like
546 sequence in the *G. margarita* genome were searched querying the assembly with the
547 comp23972_c0_seq1 (GBYF01024012) sequence using BLASTN 2.6.0+ (Zhang *et al.*, 2000), and
548 mapped reads were counted after BWA analysis.

549 The viral sequences have been submitted to GenBank under the following accession numbers:
550 33086: MG256173, 34036: MG256174, 34470: MG256175, 34875: MG256176 and 33452:
551 MG256177.

552

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560 **REFERENCES**

561

562 Andrews, S. (2010) FastQC: a quality control tool for high throughput sequence data. URL:
563 <http://www.bioinformatics.babraham.ac.uk/projects/fastqc>

564

565 Berruti, A., Lumini, E., Balestrini, R., and Bianciotto, V. (2015) Arbuscular mycorrhizal fungi as
566 natural biofertilizers: let's benefit from past successes. *Front Microbiol* 6: 1559.

567

568 Bonfante, P., and Desirò, A. (2017) Who lives in a fungus? The diversity, origins and functions of
569 fungal endobacteria living in Mucoromycota. *ISME J* 11: 1727-1735.

570

571 Bravo, M., Brands, V., Wewer, P., Dörmann, P. and Harrison, M.J. (2017) Arbuscular mycorrhiza-
572 specific enzymes FatM and RAM2 fine-tune lipid biosynthesis to promote development of
573 arbuscular mycorrhiza. *New Phytol* 214: 1631-1645.

574

575 Bruenn, J.A., Warner, B.E., Yerramsetty, P. (2015) Widespread mitovirus sequences in plant
576 genomes. *PeerJ*. 3:e876.

577

578 Campanella, J.J., Bitincka, L. and Smalley, J. (2003) MatGAT: An application that generates
579 similarity/identity matrices using protein or DNA sequences. *BMC Bioinform* 4.

580

581 Choi, G.H. and Nuss, D.L. (1992) A viral gene confers hypovirulence-associated traits to the
582 chestnut blight fungus. *EMBO J* 11: 473-477.

583

584 Cole, T.E., Hong, Y., Brasier, C.M. and Buck, K.W. (2000) Detection of an RNA-dependent RNA
585 polymerase in mitochondria from a mitovirus-infected isolate of the Dutch Elm disease fungus,
586 *Ophiostoma novo-ulmi*. *Virology* 268: 239-243.

587

588 Corradi, N., and Brachmann, A. (2016) Fungal mating in the most widespread plant symbiosis?
589 *Trends in Plant Sci* 22: 175-183.

590

591 Desirò, A., Salvioli, A., Ngonkeu, E.L., Mondo, S.J., Epis, S., Faccio, A. *et al.* (2014) Detection of
592 a novel intracellular microbiome hosted in arbuscular mycorrhizal fungi. *ISME J* 8: 257-270.

593

594 Dolja, V.V., and Koonin, E.V. (2012) Capsid-Less RNA Viruses. In: eLS. John Wiley & Sons Ltd,
595 Chichester. URL: <http://www.els.net>

596

597 Edgar, R.C. (2004) MUSCLE: multiple sequence alignment with high accuracy and high
598 throughput. *Nucleic Acids Res* 32(5): 1792-1797.

599

600 Fauquet, C.M., Mayo, M.A., Maniloff, J., Desselberger, U., and Ball, L.A. (2005) Virus taxonomy:
601 the eighth report of the International Committee on Taxonomy of Viruses. Amsterdam, NL:
602 Elsevier Academic Press.

603

604 Ghabrial, S.A., Castón, J.R., Jiang, D., Nibert, M.L., Suzuki N. (2015) 50-plus years of fungal
605 viruses. *Virology* 479-480: 356-68.

606

607 Ghabrial, S.A., and Suzuki, N. (2009) Viruses of plant pathogenic fungi. *Annu Rev Phytopathol* 47:
608 353-384

609

610 Ghignone, S., Salvioli, A., Anca, I., Lumini, E., Ortu, G., Petiti, L. *et al.* (2012) The genome of the
611 obligate endobacterium of an AM fungus reveals an interphylum network of nutritional interactions.
612 ISME J 6: 136-145.
613
614 Goic, B., Vodovar, N., Mondotte, J.A., Monot, C., Frangeul, L., Blanc, H. *et al.* (2013) RNA-
615 mediated interference and reverse transcription control the persistence of RNA viruses in the insect
616 model *Drosophila*. Nat Immunol 14: 396-403.
617
618 Goic, B., Stapleford, K. A., Frangeul, L., Doucet, A. J., Gausson, V., Blanc, H., *et al.* (2016) Virus-
619 derived DNA drives mosquito vector tolerance to arboviral infection. Nat Commun 7: 12410.
620
621 Jiang, Y., Wang, W., Xie, Q., Liu, N., Liu, L., Wang, D. *et al.* (2017) Plants transfer lipids to
622 sustain colonization by mutualistic mycorrhizal and parasitic fungi. Science 356: 1172-1175.
623
624 Jiang, D., Fu, Y., Li, G. and Ghabrial, S.A. (2013) Viruses of the Plant Pathogenic Fungus
625 *Sclerotinia sclerotiorum*. In: Ghabrial SA, editor. Adv Virus Res 86: 215-248.
626
627 Hillman, B.I. and Cai, G. (2013) The family Narnaviridae: simplest of RNA viruses. Adv Virus Res
628 86: 149-176.
629
630 Hillman, B.I, and Esteban, R. (2011) Narnaviridae, In: King AMQ, Adams MJ, Carstens EB,
631 Lefkowitz EJ. (Eds.), Virus taxonomy: classification and nomenclature of viruses: Ninth Report of
632 the International Committee on Taxonomy of Viruses. Elsevier, San Diego, 2011. pp.1055-1060.
633
634 Hillman, B.I., Supyani, S., Kondo, H. and Suzuki, N. (2004) A reovirus of the fungus
635 *Cryphonectria parasitica* that is infectious as particles and related to the coltivirus genus of animal
636 pathogens. J Virol 78: 892-898.
637
638 Ikeda, Y., Shimura, H., Kitahara, R., Masuta, C., and Ezawa, T. (2012). A novel virus-like double-
639 stranded RNA in an obligate biotroph arbuscular mycorrhizal fungus: a hidden player in
640 mycorrhizal symbiosis. Mol Plant-Microbe Interact 25: 1005-1012.
641
642 Kamel, L., Keller-Pearson, M., Roux, C., and Ané, J.-M. (2016) Biology and evolution of
643 arbuscular mycorrhizal symbiosis in the light of genomics. New Phytol 213: 531-536.
644
645 Keymer, A., Pimprikar, P., Wewer, V., Huber, C., Brands, M., Simone L Bucerius, S *et al.* (2017)
646 Lipid transfer from plants to arbuscular mycorrhiza fungi. eLife 6, e29107.
647
648 Kitahara, R., Ikeda, Y., Shimura, H., Masuta, C. and Ezawa T. (2014). A unique mitovirus from
649 Glomeromycota, the phylum of arbuscular mycorrhizal fungi. Arch Virol 159:2157-2160.
650
651 Kondo, H., Kanematsu, S. and Suzuki, N. (2013) Viruses of the White Root Rot Fungus, *Rosellinia*
652 *necatrix*. In: Ghabrial SA, editor. Adv Virus Res 86: 177-214.
653
654 Koonin, E.V., and Dolja, V.V. (2014) Virus world as an evolutionary network of viruses and
655 capsidless selfish elements. Microbiol Mol Biol Rev 78: 278-303.
656
657 Krueger, F. (2012) A wrapper tool around Cutadapt and FastQC to consistently apply quality and
658 adapter trimming to FastQ files, with some extra functionality for MspI-digested RRBS-type

659 (Reduced Representation Bisulfite-Seq) libraries. URL:
660 https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/
661
662 Le, S.Q. and Gascuel, O. (2008) An improved general amino acid replacement matrix. *Mol Biol*
663 *Evol* 25:1307-1320.
664
665 Liu, S., Xie, J., Cheng, J., Li, B., Chen, T., Fu, Y. *et al.* (2016) Fungal DNA virus infects a
666 mycophagous insect and utilizes it as a transmission vector. *Proc. Natl. Acad. Sci. USA* 113:
667 12803-12808.
668
669 Li, H., and Durbin, R. (2009) Fast and accurate short read alignment with Burrows-Wheeler
670 Transform. *Bioinformatics* 25: 1754-60.
671
672 Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N. *et al.* (2009) The Sequence
673 alignment/map (SAM) format and SAMtools. *Bioinformatics* 25: 2078-2079.
674
675 Li, Z., Yao, Q.M., Dearth, S.P., Entler, M.R., Castro Gonzalez, H.F., Uehling, J.K. *et al.* (2017)
676 Integrated proteomics and metabolomics suggests symbiotic metabolism and multimodal regulation
677 in a fungal-endobacterial system. *Env Microb* 19:1041-1053.
678
679 Lin, K., Limpens, E., Zhang, Z.H., Ivanov, S., Saunders, D.G.O., Mu, D.S. *et al.* (2014) Single
680 nucleus genome sequencing reveals high similarity among nuclei of an endomycorrhizal fungus.
681 *PLoS Genet* 10:e1004078.
682
683 Luginbuehl, L.H., Menard, G.N., Kurup, S., Van Erp, H., Radhakrishnan, G.V., Breakspear, A.,
684 Oldroyd, G.E.D. and Eastmond, P.J. (2017) Fatty acids in arbuscular mycorrhizal fungi are
685 synthesized by the host plant. *Science* 356: 1175-1178.
686
687 Lumini, E., Bianciotto, V., Jargeat, P., Novero, M., Salvioli, A., Faccio, A., Bécard, G. and
688 Bonfante, P. (2007) Presymbiotic growth and sporal morphology are affected in the arbuscular
689 mycorrhizal fungus *Gigaspora margarita* cured of its endobacteria. *Cellular Microbiology* 9: 1716-
690 1729.
691
692 Márquez, L.M., Redman, R.S., Rodriguez, R.J., and Roossinck, M.J. (2007) A virus in a fungus in a
693 plant: three-way symbiosis required for thermal tolerance. *Science* 315: 513-515.
694
695 Nag, D.K., Brecher, M. and Kramer, L.D. (2016) DNA forms of arboviral RNA genomes are
696 generated following infection in mosquito cell cultures. *Virology* 498: 164-171.
697
698 Naito, M., Morton, J.B., and Pawlowska, T.E. (2015) Minimal genomes of mycoplasma-related
699 endobacteria are plastic and contain host-derived genes for sustained life within Glomeromycota.
700 *Proc Natl Acad Sci USA* 112: 7791-7796.
701
702 Naumann, M., Schußler, A., Bonfante, P. (2010) The obligate endobacteria of arbuscular
703 mycorrhizal fungi are ancient heritable components related to the Mollicutes. *ISME J* 4: 862-871.
704
705 Nerva, L., Ciuffo, M., Vallino, M., Margaria, P., Varese, G.C., Gnani, G., Turina M. (2016)
706 Multiple approaches for the detection and characterization of viral and plasmid symbionts from a
707 collection of marine fungi. *Virus Res* 219: 22-38.
708

709 Nerva L, Silvestri A, Ciuffo M, Palmano S, Varese GC, Turina M. (2017) Transmission of
710 *Penicillium aurantiogriseum* partiti-like virus 1 to a new fungal host (*Cryphonectria parasitica*)
711 confers higher resistance to salinity and reveals adaptive genomic changes. *Env Microb* 19: 4480-
712 4492.

713

714 Nibert, M.L. (2017) Mitovirus UGA(Trp) codon usage parallels that of host mitochondria.
715 *Virology* 507: 96-100.

716

717 Olson, K.E. and Bonizzoni, M. (2017) Nonretroviral integrated RNA viruses in arthropod vectors:
718 an occasional event or something more? *Curr Opin Insect Sci* 22: 45-53.

719

720 Pelin, A., Pombert, J.F., Salvioli, A., Bonen, L., Bonfante, P., and Corradi, N. (2012) The
721 mitochondrial genome of the arbuscular mycorrhizal fungus *Gigaspora margarita* reveals two
722 unsuspected transsplicing events of group I introns. *New Phytol* 194: 836-845.

723

724 Petrzik K., Sarkisova, T., Starý, J., Koloniuk, I., Hrabáková, L. and Kubešová, O. (2016) Molecular
725 characterization of a new monopartite dsRNA mycovirus from mycorrhizal *Thelephora terrestris*
726 (Ehrh.) and its detection in soil oribatid mites (Acari: Oribatida). *Virology* 489: 12-19.

727

728 Rasmussen, R. (2001) Quantification on the LightCycler. In S Mener, C Wittwer, K Nakagawara,
729 eds, *Rapid Cycle Real-Time PCR: Methods and Applications*. Springer Press, Heidelberg, pp 21-34.

730

731 Rodriguez, A. and Sanders, I. R. (2015) The role of community and population ecology in applying
732 mycorrhizal fungi for improved food security. *ISME J* 9: 1053-1061.

733

734 Ropars, J., Toro, K.S., Noel, J, Pelin, A., Charron, P., Farinelli, L., *et al.* (2016) Evidence for the
735 sexual origin of heterokaryosis in arbuscular mycorrhizal fungi. *Nat Microbiol* 21: 16033.

736

737 Salvioli, A., Ghignone, S., Novero, M., Navazio, L., Venice, F., Bagnaresi, P. and Bonfante, P.
738 (2016) Symbiosis with an endobacterium increases the fitness of a mycorrhizal fungus, raising its
739 bioenergetic potential. *ISME J* 1-15.

740

741 Sambrook, J., Fritsch, E.F. and Maniatis T. (1987) *Molecular cloning: a laboratory manual*. - 2nd
742 ed. Cold Spring Harbor, N.Y. Cold Spring Harbor Laboratory Press.

743

744 Sędziewska Toro, K. and Brachmann, A. (2016) The effector candidate repertoire of the
745 arbuscular mycorrhizal fungus *Rhizophagus clarus*. *BMC Genomics* 17:101.

746

747 Shi, M., Lin, X.D., Tian, J.H., Chen, L.J., Chen, X., Li, C.X., Qin, X.C., Li, J., Cao, J.P., Eden, J.S.,
748 Buchmann, J., Wang, W. Xu, J., Holmes, E.C., and Zhang, Y.Z. (2016) Redefining the invertebrate
749 RNA virosphere. *Nature* 540: 539-543.

750

751 Son, M., Ju, J., and Kim, K-H. (2015) Five questions about mycoviruses. *PLoS Pathog* 11:
752 e1005172.

753

754 Spatafora, J.W., Chang, Y., Benny, G.L., Lazarus, K., Smith, M.E., Berbee, M.L. *et al.* (2016) A
755 phylum-level phylogenetic classification of zygomycete fungi based on genome-scale data.
756 *Mycologia* 108: 1028-1046.

757

758 Stielow, B., Klenk, H.P., Winter, S., and Menzel, W. (2011) A novel *Tuber aestivum* (Vittad.)
759 mitovirus. Arch Virol 156: 1107-1110.
760
761 Stielow, B., Klenk, H.P., Winter, S., and Menzel, W. (2011) Complete genome sequence of the first
762 endornavirus from the ascocarp of the ectomycorrhizal fungus *Tuber aestivum* Vittad. Arch Virol
763 156: 343-345.
764
765 Stielow, B., and Menzel, W. (2010) Complete nucleotide sequence of TaV1, a novel totivirus
766 isolated from a black truffle ascocarp (*Tuber aestivum* Vittad.). Arch Virol 155: 2075-2078.
767
768 Tamura, K., Stecher, G., Peterson, D., Filipiński, A., Kumar, S. (2013) MEGA6: Molecular
769 Evolutionary Genetics Analysis Version 6.0. Mol Biol Evol 30: 2725-2729.
770
771 Tang, N., San Clemente, H., Roy, S., Bécard, G., Zhao, B., and Roux, C. (2016) A survey of the
772 gene repertoire of *Gigaspora rosea* unravels conserved features among Glomeromycota for obligate
773 biotrophy. Front Microbiol 7:233.
774
775 Tisserant, E., Malbreil, M., Kuo, A., Kohler, A., Symeonidi, A., Balestrini, R., *et al.* (2013)
776 Genome of an arbuscular mycorrhizal fungus provides insight into the oldest plant symbiosis. Proc
777 Natl Acad Sci USA 110: 20117-20122.
778
779 Torres-Cortés, G., Ghignone, S., Bonfante, P., and Schüßler, A. (2015) Mosaic genome of
780 endobacteria in arbuscular mycorrhizal fungi: transkingdom gene transfer in an ancient
781 mycoplasma-fungus association. Proc Natl Acad Sci USA 112: 7785-7790.
782
783 Turina, M., Hillman, B.I., Izadpanah, K., Rastgou, M., Rosa, C., Consortium, I.R. (2017) ICTV
784 Virus Taxonomy Profile: Ourmiavirus. J Gen Virol 98: 129-130.
785
786 van Diepeningen, A.D., Debets, A.J., and Hoekstra, R.F. (2006) Dynamics of dsRNA mycoviruses
787 in black Aspergillus populations. Fungal Genet Biol 43: 446-452.
788
789 Vannini, C., Carpentieri, A., Salvioli, A., Novero, M., Marsoni, M., Testa, L. *et al.* (2016) An
790 interdomain network: the endobacterium of a mycorrhizal fungus promotes antioxidative responses
791 in both fungal and plant hosts. New Phytol 211: 265-275.
792
793 Wu, M., Deng, Y., Zhou, Z., He, G., Chen, W. and Li, G. (2016) Characterization of three
794 mycoviruses co-infecting the plant pathogenic fungus *Sclerotinia nivalis*. Virus Res 223: 28-38.
795
796 Xie, J., Ghabrial, S.A. (2012) Molecular characterizations of two mitoviruses co-infecting a
797 hyovirulent isolate of the plant pathogenic fungus *Sclerotinia sclerotiorum*. Virology 428: 77-85.
798
799 Yaegashi, H., Nakamura, H., Sawahata, T., Sasaki, A., Iwanami, Y., Ito, T., and Kanematsu, S.
800 (2013) Appearance of mycovirus-like double-stranded RNAs in the white root rot fungus,
801 *Rosellinia necatrix*, in an apple orchard. FEMS Microbiol Ecol 83: 49-62.
802
803 Young, J.P.W. (2015) Genome diversity in arbuscular mycorrhizal fungi. *Curr Opin Plant Biol* 23:
804 113-119.
805
806 Zhang, Z., Schwartz, S., Wagner, L., and Miller, W. (2000) A greedy algorithm for aligning DNA
807 sequences. J Comput Biol 7: 203-214.

808 **TABLES**

809 **Table 1.** Percentages similarity (*italics*) and identity (underlined) of pairwise alignments of the four *G. margarita*
 810 mitovirus sequences along the entire protein sequence. Identity and similarity among pairwise alignments were
 811 calculated using the Needleman-Wunsch Global Align Protein Sequence algorithm.

Mitovirus contigs	33086	34036	34470	34875
33086		<i>34%</i>	<i>32%</i>	<i>33%</i>
34036	<u>21%</u>		<i>38%</i>	<i>40%</i>
34470	<u>19%</u>	<u>23%</u>		<i>56%</i>
34875	<u>20%</u>	<u>24%</u>	<u>39%</u>	

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813 **Table 2.** Ct values (+/- standard deviation) registered by qPCR on *G. margarita* genomic DNA (3 independent samples)
 814 with primers for the 4 mitoviral sequences. 18S rDNA and *Tef* were used as endogenous genes corresponding to
 815 multiple or single copy genes, respectively. Different letters indicate statistically significant differences (ANOVA, p <
 816 0.05).

	33086	34036	34470	34875	Tef	18S rDNA
Ct	30.49	27.72	28.46	29.16	25.37	20.93 ⁸¹⁷
standard deviation	+/- 0.6647	+/-0.2121	+/-0.0141	+/-0.1273	+/-0.1697	+/-0.3111 ⁸¹⁸
Tukey test	d	b	bc	c	a	e ⁸¹⁹

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832 **FIGURE LEGENDS**

833 **Fig. 1.** Gel electrophoresis of RT-PCR products amplified from cDNA from *G. margarita*
834 germinating spores with the following primer pairs specific for the putative viral sequences:
835 33086genF1 and 33086genR1, 34036Fgen and 34036Rgen, 34770Fgen and 34770Rgen,
836 34875Fgen and 34875Rgen, 36178Fgen and 36178Rgen, 33452Fgen and 33452Rgen, 23972Fgen
837 and 23972Rgen, 39980Fgen and 39980Rgen. M: 100 bp ladder (Invitrogen); no cDNA sample (-).

838 **Fig. 2.** Northern blot analysis of *G. margarita*-associated viral sequences. Time of autoradiography
839 exposure (Exp) is indicated on top of each panel. MUT4330 (total RNA from an unrelated
840 filamentous fungus) was used as negative control in the hybridization experiment. Lower panels are
841 methylene blue stained membranes to show ribosomal RNA loading. gRNA= genomic RNA;
842 rRNA=ribosomal RNA.

843 **Fig. 3.** Molecular phylogenetic analysis by Maximum Likelihood method of Mitovirus sequences
844 (*G. margarita* sequences are indicated by arrows). The tree with the highest log likelihood (-
845 22958.4232) is shown. Initial tree(s) for the heuristic search were obtained by applying the
846 Neighbor-Joining method to a matrix of pairwise distances estimated using a JTT model. A discrete
847 Gamma distribution was used to model evolutionary rate differences among sites: 5 categories (+G,
848 parameter = 1.0199). The rate variation model allowed for some sites to be evolutionarily invariable
849 ([+I], 3.8500% sites). The tree is drawn to scale, with branch lengths measured in the number of
850 substitutions per site. All positions with less than 95% site coverage were eliminated. There were a
851 total of 467 positions in the final dataset.

852 **Fig. 4.** Molecular Phylogenetic analysis by Maximum Likelihood method of the Giardia-like viral
853 sequence in the context of some representative of the established family taxa *Totiviridae* and
854 *Partitiviridae*. The tree with the highest log likelihood (-11851.6550) is shown. The percentage of
855 trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for
856 the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms

857 to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with
858 superior log likelihood value. The rate variation model allowed for some sites to be evolutionarily
859 invariable ([+I], 3.5294% sites). The tree is drawn to scale, with branch lengths measured in the
860 number of substitutions per site. The analysis involved 29 amino acid sequences. All positions
861 containing gaps and missing data were eliminated. There were a total of 170 positions in the final
862 dataset. The star indicates the position of the *G. margarita* Giardia-like virus characterized in this
863 study.

864 **Fig. 5.** Gel electrophoresis of PCR products of *G. margarita* DNA untreated (1) or treated (2) with
865 DNase with primers specific for the viral sequences (the same primer pairs used in figure 1).
866 Samples 3 correspond to reactions with no DNA. M: 100 bp DNA marker (Invitrogen).

867 **Fig. 6.** Characters of the DNA corresponding to mitovirus sequences. A: Gel electrophoresis of
868 PCR products obtained from *G. margarita* cDNA (1), DNA extractions after DNase (2) and DNA
869 extractions (3); B: Gel electrophoresis of PCR products obtained from two *G. margarita*
870 independent genomic DNA extractions (1, 2), two independent cDNAs preparations (3, 4) or no
871 DNA (5) samples using the different primer pairs. The size of the amplicon in base pairs (bp) is
872 indicated in brackets. C: Scheme of the 34036 viral sequence showing the position of primers
873 (F=forward; R=reverse) indicated by arrows.

874 **Fig. 7.** Gel electrophoresis of PCR products from genomic DNA samples: *G. margarita* B+ (1) and
875 B- (2) harbouring endobacteria or endobacteria-cured, respectively; *G. rosea* (3); *G. margarita*
876 positive control (4); no DNA (5). The specific primer pair used for each virus is listed in the legend
877 of Figure 1.

878 **Fig. 8.** Quantitative RT-PCR on cDNA of B+ and B- (harbouring endobacteria or endobacteria
879 cured, respectively) germinating spores of *G. margarita* with primers specific for the 4 mitovirus
880 and the Giardia-like virus. *GmTef* was used as housekeeping gene and B- sample as reference. No

881 statistically significant difference was found between B+ and B- samples for any of the five viruses
882 displayed (ANOVA; $p < 0.05$).

883

884 SUPPORTING INFORMATION

885 **Fig. S1.** Nucleotide sequences of the full-length contigs assembled in silico corresponding to the
886 virus sequences in fasta format. RdRp start and stop codons are indicated in bold and cyan. When
887 relevant, the corresponding translation frame is given in order to outline the codons encoding for
888 tryptophan (W highlighted in yellow). Amino acid sequence of the RdRp are also then reported in
889 fasta format and the GDD conserved motif is indicated in pink.

890 **Fig. S2.** Gel electrophoresis of PCR products amplified from *G. margarita* genomic DNA (1); *G.*
891 *margarita* genomic DNA treated with DNase (2); *G. margarita* cDNA from germinating spores (3);
892 no DNA sample (4) with primers specific for the Ourmia-like sequence (36178 contig). M: 1 kb
893 ladder (Invitrogen).

894 **Fig. S3.** Molecular Phylogenetic analysis by Maximum Likelihood method of the four mitoviruses
895 and the Ourmia virus (positions in the tree pointed by red arrows). The evolutionary history was
896 inferred by using the Maximum Likelihood method based on the Le_Gascuel_2008 model (Le
897 and Gascuel, 2008). The tree with the highest log likelihood (-11687.6554) is shown. The
898 percentage of trees in which the associated taxa clustered together is shown next to the branches.
899 Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and
900 BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting
901 the topology with superior log likelihood value. A discrete Gamma distribution was used to model
902 evolutionary rate differences among sites (5 categories (+G, parameter = 1.0325)). The rate
903 variation model allowed for some sites to be evolutionarily invariable ([+I], 2.9115% sites). The
904 tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The
905 analysis involved 64 amino acid sequences. All positions containing gaps and missing data were

906 eliminated. There were a total of 109 positions in the final dataset. Evolutionary analyses were
907 conducted in MEGA6 (Tamura *et al.*, 2013).

908 **Fig. S4.** Schematic representation of the assembly of the clones used for deriving the 2kb consensus
909 sequence corresponding to 34036 mitovirus sequence amplified by PCR using as template DNA
910 preparations (compare with Fig. 6). In blue the amplified genomic region. Vertical green bars
911 correspond to single nucleotide mutations in specific position in at least one of the cloned
912 amplicons in respect to the consensus sequence. The only conserved mutation compared to the *in*
913 *silico* assembled sequence from transcriptome is a synonymous C to T change at position 1399.
914 Gray arrows represent the full-length viral contig sequence (derived from transcriptome, top arrow)
915 and all the different sequences derived from cloning the PCR amplification products displayed in
916 Fig. 6.

917

918 **Table S1:** BLAST searches of databases using as query the viral RdRp contigs present in *G.*
919 *margarita* transcriptome.

920 **Table S2.** List of primers used in this study.

921 **Table S3:** Percentage identity (upper right triangle) and similarity (lower left triangle) among
922 aligned RdRp from different mitoviruses.

923 **Table S4:** Accession numbers and virus names of proteins used in the phylogenetic analysis.