Evidence of non-tandemly repeated rDNAs and their 1

intragenomic heterogeneity in Rhizophagus irregularis 2

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13 Abstract

14 Arbuscular mycorrhizal fungus (AMF) species are one of the most widespread symbionts of land 15 plants. Our substantially improved reference genome assembly of a model AMF, Rhizophagus irregularis 16 DAOM-181602 (total contigs = 210), facilitated discovery of repetitive elements with unusual 17 characteristics. R. irregularis has only ten or eleven copies of complete 45S rDNAs, whereas the general 18 eukaryotic genome has tens to thousands of rDNA copies. R. irregularis rDNAs are highly heterogeneous 19 and lack a tandem repeat structure. These findings provide evidence for the hypothesis that rDNA $\hat{20}$ heterogeneity depends on the lack of tandem repeat structures. RNA-Seq analysis confirmed that all $\overline{21}$ rDNA variants are actively transcribed. Observed rDNA/rRNA polymorphisms may modulate translation 22 23 by using different ribosomes depending on biotic and abiotic interactions. The non-tandem repeat structure and intragenomic heterogeneity of AMF rDNA/rRNA may facilitate adaptation to a various 24environmental condition including the broad host range.

26 Introduction

27 The arbuscular mycorrhizal fungus (AMF) is an ancient fungus with origins at least as old as 28 the early Devonian period^{1,2}. AMF colonizes plant roots and develops highly branched structures 29 called arbuscules in which soil nutrients (phosphate and nitrogen) are efficiently delivered to the 30 host plant³. AMF forms symbiotic networks with most land plant species^{4,5}, and the mycelial 31 network formed by various AMF species contributes to plant biodiversity and productivity within 32 the terrestrial ecosystem⁶. The distinctive features of AMF have made it an important model in 33 ecology and evolution^{7,8}; these features include coenocytic mycelia⁵, nutrition exchange with plant, 34 classification as an obligate biotroph⁹, signal crosstalk during mycorrhiza development^{9,10} and 35 extremely high symbiotic ability^{9,11}.

36 Recently, multiple genome projects have advanced the understanding of AMF species. 37 Genomic data have been provided for *Rhizophagus irregularis* DAOM-181602^{12,13,14}, *Gigaspora* 38 rosea¹², Rhizophagus clarus¹⁵, and other isolates of R. irregularis^{,14,16}. These studies revealed 39 potential host-dependent biological pathways¹⁷¹² and candidate genes for plant infection and sexual 40 reproduction ^{12,15,16}. However, fragmented genome sequences limit the ability to analyze repetitive 41 structures and to distinguish between orthologous and paralogous genes¹⁴. The first published 42 genome sequence of R. irregularis DAOM-181602 (JGI v1.0)¹⁷ contained 28,371 scaffolds and an 43 N50 index of 4.2 kb (Supplementary Table 1). The second sequence, by Lin et al. 2014 (Lin14)¹³, 44 contained 30,233 scaffolds with an N50 of 16.4 kb (Supplementary Table 1). Recently published 45 assemblies by Chen et al. 2018 (JGI v2.0)¹⁴ contained 1,123 scaffolds with an N50 of 336.4 kb 46 (Supplementary Table 1). The quality of genomic sequence data for other AMF species did not 47 surpass that of DAOM-181602^{12,15}. In contrast, many fungi that are not AMF species contain less 48 than several hundred scaffolds and N50 lengths over 1 Mb¹⁸. For example, a genomic sequence of an 49 asymbiotic fungus closely related to AMF, Rhizopus delemar (GCA000149305.1), was constructed 50 from 83 assemblies with an N50 of 3.1 Mb¹⁹. Thus, we here present an improved whole-genome 51 sequence of R. irregularis DAOM-181602 to facilitate examination of the genomics underlying 52 specific features of AMF species. Taking an advantage of the highly contiguous assembly with little 53 ambiguous regions, we focus on the investigation of the repetitive structures including transposable 54 elements, highly duplicated genes, and rDNA gene copies.

55 A general eukaryotic genome has tens to thousands of rDNA copies²⁰ (Supplementary Fig. 56 1a), and the sequences of the copies are identical or nearly identical. However, since Sanders et al. 57 (1995)²¹, many studies have indicated intracellular polymorphisms of rDNA (ITS) in various AMF 58 species²²⁻²⁴, and the sequencing of isolated nuclei from *Claroideoglomus etunicatum* and 59 Rhizophagus irregularis DAOM-181602 suggested sequence variation among the paralogous 60 rDNAs, i.e., intragenomic heterogeneity^{13,25}. This heterogeneity has potentially high impact of 61 studying AMF species, because the rDNA is a fundamental marker of the AMF phylogeny and 62 ecology^{8,26,27,28}, and studies have assumed that these rDNAs have no intragenomic sequence 63 variation²⁹. Hence, determining the variation degree could cause a reevaluation of the previous understanding of geographic distribution⁸, species identification²⁸, and evolutionary processes of 64 65 AMF. However, the degree of the variation among the 48S rDNA paralogs has been ambiguous 66 because previous studies by Sanger or Illumina sequencing were unable to distinguish each rDNA 67 paralog in a genome. Moreover, the number of rDNA genes in an AMF genome has never been 68 investigated.

The tandem repeat structure (TRS) of the rDNAs is also an attractive topic for evolutionary studies. General organisms require many rDNA copies to make a sufficient amount of rRNA for protein translation^{30,31}. However, in the evolutionary time-scale, multicopy genes reduce in number due to homologous recombination (Supplementary Fig. 1b)^{32,33} and single-strand annealing (Supplementary Fig. 1c)³⁴. To maintain the number of rDNAs, eukaryotes increase the number of copies by unequal sister chromatid recombination (USCR) using the rDNA TRS (Supplementary Fig. 1d,e)³². Because this rDNA replacement causes a bottleneck effect in the genome, almost all eukaryotes have homogenous rDNAs in their genomes²⁰. This process, termed "concerted evolution," is an essential system to maintaining eukaryotic protein translation by ribosomes³⁰. The heterogeneous rDNAs observed in AMF species implies the collapse of their concerted evolution, and suggest the unique maintenance system of rDNA copy number.

80 In this study, we built an improved reference genome assembly of *Rhizophagus irregularis* 81 DAOM-181602, which allowed us to discover repetitive elements with unusual characteristics of the

82 AMF genome. We identified unusually small number of rDNA genes in the *R. irregularis* genome.

83 We also found that the rDNA copies are highly heterogeneous and lack a tandem repeat structure.

84

86 **Results**

A highly contiguous and complete reference genome of DAOM-181602 generated by PacBio-based *de novo* assembly

89 We primarily used single-molecule, real-time (SMRT) sequencing technology for sequencing 90 and assembling the *R. irregularis* genome. We generated a 76-fold whole-genome shotgun sequence 91 (11.7 Gb in total) (Supplementary Table 2) from genome DNA isolated from a spore suspension of a 92 commercial strain of R. irregularis DAOM-181602 using the PacBio SMRT sequencing platform. A 93 total of 766,159 reads were generated with an average length of 13.1 kb and an N50 length of 18.8 94 kb (Supplementary Table 2). We assembled these PacBio reads using the HGAP3 program³⁵ (149.9 95 Mb composed of 219 contigs). To detect erroneous base calls, we generated 423 Mb of 101 bases-96 paired-end Illumina whole-genome sequence data (Supplementary Table 2) and aligned them to the 97 HGAP3 assembly. Through variant calling, we corrected 3,032 single base call errors and 10,841 98 small indels in the HGAP3 assembly. Nine contigs were almost identical to carrot DNA sequences 99 deposited in the public database (Supplementary Table 3), and these were removed as contaminants 100 derived from a host plant used by the manufacturer. We evaluated the completeness of the final 101 assembly using CEGMA³⁶; of the 248 core eukaryotic genes, 244 genes (98.4%) were completely 102 assembled (Table 1 and Supplementary Table 1). Consequently, we obtained a high-quality 103 reference genome assembly of *R. irregularis* DAOM-181602, which is referred to as RIR17.

104 Compared with previous assemblies^{13,14,17}, RIR17 represents a decrease in assembly 105 fragmentation (1,123 to 210) and an improvement in contiguity using the N50 contig length as a 106 metric (Table 1 and Supplementary Table 1). The total size of the assembly was 9-59 Mb greater 107 than that of previous versions, reaching 97.24% coverage of the whole genome $(154 \text{ Mb})^{17}$ (Table 1 108 and Supplementary Table 1). The new assembly contains no ambiguous bases (N-bases), whereas 109 previous assemblies had 30,115-6,925,426 N-bases (Table 1 and Supplementary Table 1). 110 Approximately 1-7 Mb of sequences from previous assemblies were not contained in RIR17, and 111 JGI v2.0 has one more conserved gene family than RIR17 (Supplementary Table 1), indicating that 112 a few genomic parts remain to be uncovered by our improvement with continuous sequences. On the 113 other hand, RIR17 was aligned with 95-99.2% of previous assemblies (Supplementary Table 4), 114 suggesting that RIR17 covers the majority of the previously sequenced areas with high sequence 115 contiguity. Moreover, RIR17 contained 8-47 Mb of regions unassigned in previous genomes 116 (Supplementary Table 4). These regions are newly revealed by our improvement.

117 RIR17 contains a greater extent of repetitive regions than JGI_v2.0. The RepeatModeler³⁷ and 118 RepeatMasker³⁷ pipeline identified 64.4 Mb (43.03%) of RIR17 as repetitive regions 119 (Supplementary Table 5). These regions total 18.9 Mb more than those of JGI_v2.0 (Supplementary 120 Table 5). Previous fosmid sequences predicted that DAOM-181602 contains ~55 Mb of repetitive 121 regions¹⁷, suggesting that RIR17 covers the majority of the repetitive regions of DAOM-181602.

122 We confirmed a unique repeat profile in the AMF genome. The majority of the interspersed 123 repeats (62.83%) could not be categorized with known repeat classes (Supplementary Table 5), 124 indicating that the AMF genome accumulated novel classes of interspersed repeats. Moreover, 125 DAOM-181602 lacks short interspersed nuclear elements (SINEs), which are abundant in closely 126 related fungi (Supplementary Table 5). Several types of SINEs proliferate using transposases on 127 long interspersed nuclear elements (LINEs)³⁸. Although the AMF has 23 LINEs containing the 128 transposase gene (Supplementary Table 6), SINEs have never been found in previous genomes^{13,14,17} 129 or RIR17. DAOM-181602 may have a system to suppress the invasion and proliferation of SINEs 130 (e.g., a high number of very active Argonaute proteins, as predicted by Tisserant et al.¹⁷).

132 New gene annotation for DAOM-181602 details gene family expansion 133 in AMF

134 Using the RIR17 assembly together with strand-specific RNA-Seq data ("Rir RNA SS" in 135 Supplementary Table 2), we built a set of 41,572 gene models (Supplementary Tables 6 and 7). Of 136 the genes predicted, 27,859 (67.0%) had either RNA-Seq expression support, homology support or 137 protein motif support (Supplementary Tables 6 and 7). The gene models having any support were 138 submitted to the DDBJ as standard genes and were used in downstream analyses. The models having 139 no support were assigned as "PROVISIONAL" gene models (Supplementary Tables 6 and 7). Using 140 Orthofinder with previous genomic gene sets indicated that our gene models cover the majority of 141 previously provided genes (Supplementary Fig. 2). Although new models showed more coverage of 142 "Benchmarking Universal Single-Copy Orthologs" (BUSCOs)³⁹ (Supplementary Table 7) than 143 JGI v1.0 and Lin14, their gene completeness was slightly lower than that of JGI v2.0 (9 BUSCO 144 families overlooked, Supplementary Table 7), indicating the advantage of using the JGI annotation 145 pipeline to discuss the gene variety in DAOM-181602 (Chen et al 2018¹⁴). However, we considered 146 our model set suitable for the analysis of the repetitive region and highly paralogous genes because 147 our model is based on highly continuous assemblies, and the number of genes on repetitive regions 148 was increased to 2,349-12,559 genes from the number in JGI 2.0 (Supplementary Table 8).

149 R. irregularis has one of the largest numbers of genes in fungi (Supplementary Fig. 3). Our 150 ortholog analyses indicates that the gene number inflation was caused by lineage-specific expansions 151 of gene families and not by whole-genome duplications. An Orthofinder analysis of nine fungal 152 genomes and two animal data sets (Supplementary Table 9) showed that many of the single-copy 153 genes in other fungi were also single copies in RIR17 (216/239 families, Supplementary Table 10), 154 negating the possibility of whole-genome duplication in R. irregularis. The large number of 155 "species-specific single-copy" (SSSC) genes in DAOM-181602 (10,354 genes, Fig. 1a, 156 Supplementary Table 10) suggests that the AMF genes inflated by new gene constructions through 157 gene fusion and mutation accumulation. Moreover, several common gene families in Opisthokonta 158 also contributed to gene inflation; the R. irregularis lineage had 92 rapidly expanded (RE) families, 159 containing 8,952 genes (Fig. 1a, d-e, Supplementary Tables 10 and 11), suggesting that R. 160 irregularis has also acquired many genes by the duplication of particular gene families.

161 The motif annotation indicates that inflated genes may contribute to signaling pathways of 162 AMF species. Our Pfam search annotated 1,620 SSSC genes and 6,755 RE genes (Fig. 1b, f, 163 Supplementary Tables 6 and 10). The most frequently observed motif was "Protein tyrosine kinase; 164 PF07714" (Fig. 1b, f, Supplementary Table 12), which is often found in signaling proteins in 165 multicellular organisms⁴⁰, consistently with previous genome studies¹⁴. Other signal-related motifs 166 (e.g., Sell repeat and BED zinc finger) were also found in the inflated genes (Fig. 1b, f, 167 Supplementary Table. 12). AMF has developed a unique signal pathway for symbiosis (e.g., 168 establishments of symbiosis with pathways using sisI and $strigolactones^{41}$). This inflation of 169 signaling-related genes may have led the development of a complex signaling pathway in AMF.

170 We then investigated the contribution of the transposable elements (TEs) to gene inflation 171 based on the overlapping of highly paralogous genes and the TEs. Previous studies hypothesized that 172 the gene inflation in *R. irregularis* relates with the expansion of TEs¹⁴. Our analysis showed that in 173 several RE families (e.g., OG0000090 and OG000020), over 90% of the genes were located with 174 TEs (Fig. 1d,e, Supplementary Table 13), suggesting that TEs accelerated the gene expansion in 175 these families. However, some of the families had no correspondence with TEs (e.g., "OG0000025", 176 and "OG0000058" in Fig. 1c, e). In SSSC genes, TEs were slightly more frequently found in SSSCs 177 with motifs than in all gene sets but were less frequently found in SSSCs without motifs (Fig. 1c). 178 This detailed analysis supports the contribution of TEs to gene inflation in several gene families but 179 also clarified that several families show TE-independent expansion. Although more genome data for

AMF species and sister groups are required to reveal the gene expansion process and its contribution
 to AM symbiosis, our data provide a fundamental dataset to reveal the evolution of gene redundancy
 in AMF species.

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185 Losing conserved fungal genes

186 Previous AMF studies suggested the loss of several categories of genes by symbiosis with 187 plant^{12,13,17}. Our RIR17 genome assembly confirmed the loss of genes involved in the degradation of 188 plant cell walls such as cellobiohydrolases (GH6 and GH7 in the CAZy database), polysaccharide 189 lyases (PL1 and PL4), proteins with cellulose-binding motif 1 (CBM1), and lytic polysaccharide 190 monooxygenases (Supplementary Tables 6 and 14) and nutritional biosynthetic genes, including 191 fatty acid synthase (FAS) and the thiamine biosynthetic pathway (Supplementary Table 15). Given 192 that fatty acids and thiamine are essential nutrients for fungi42,43, R. irregularis should take up those 193 essential nutrients from a host plant without digestion of the plant cell wall. Several recent papers 194 have described the transport of lipids from plants to AMF⁴⁴⁻⁴⁶.

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196 *R. irregularis* has an exceptionally low rDNA copy number among

197 eukaryotes

198 The general eukaryotic genome has tens to thousands of rDNA copies²⁰ (Supplementary Fig. 199 1a). However, the RIR17 genome assembly contained only ten copies of the complete 45S rDNA 200 cluster, which was composed of 18S rRNA, ITS1, 5.8S rRNA, ITS2, and 28S rDNAs (Fig. 2a, 201 Supplementary Table 16). To confirm that no rDNA clusters were overlooked, we also estimated the 202 rDNA copy number based on the read depth of coverage. Mapping the Illumina reads of the genomic 203 sequences ("Rir DNA PE180") onto the selected reference sequences indicated that the coverage 204 depth of the consensus rDNA was 8-11 times deeper than the average coverage depth of the single-205 copy genes (Fig. 2b, Supplementary Table 17), the number of rDNA copies is approximately 10, and 206 the RIR17 assembly covers almost all of the rDNA copies. This AMF rDNA copy number is the 207 lowest among eukaryotes⁴⁷ other than pneumonia-causing *Pneumocystis* (one rDNA)⁴⁸ and malaria-208 causing Plasmodium (seven rDNAs)49.

209 This low copy number suggests a unique ribosome synthesis system in AMF. The rDNA copy 210 number has relevance for the efficiency of translation because multiple rDNAs are required to 211 synthesize sufficient rRNA. For instance, an experimental decrease in rDNA copy number in yeast 212 (approximately 150 rDNAs in wild type) resulted in no isolated strain having <20 copies, which is considered the minimum number to allow yeast growth³⁰. The doubling time of yeast with 20 rDNA 213 214 copies (TAK300) was 20% longer than that of the wild type³⁰. In DAOM-181602, successive 215 cultivation in an infected state with a plant has been widely observed, suggesting that this 216 exceptionally small rDNA copy number is enough to support growth. The multinucleate feature of 217 AMF would increase the rDNA copy number per cell and thereby perhaps supply enough rRNA to 218 support growth. A similar trend in rDNA reduction is observed in the organellar DNA (e.g., 219 mitochondria and plastids)⁵⁰. Revealing the details of translation in AMF requires a future tracking 220 study of the rRNA production and degradation process in AMF. Elucidation of the mechanism to 221 produce mass rRNAs from a few rDNAs may contribute the understanding of not only AMF 222 evolution but also other polynuclear cells (e.g., striated muscle and ulvophyceaen green algae) and 223 symbiont-derived organelles.

224

225 *R. irregularis* rDNAs are heterogeneous and completely lack a TRS

226 Interestingly, none of the RIR17 rDNAs form a TRS, in contrast to most eukaryotic rDNAs, 227 which comprise tens to hundreds of tandemly repeated units²⁰. Most of the rDNA clusters in RIR17 228 were placed on different contigs; a single copy of rDNA was found in "unitig 311", " 312", " 35", 229 " 356", " 4", and " 52", and two copies were found in "unitig 39" and " 62" (Fig. 2b, 230 Supplementary Table 16). In the cases where two rDNA clusters were found, the two copies resided 231 apart from each other and did not form a tandem repeat; the distances between the clusters were over 232 70 kb (76,187 bases in unitig 62 and 89,986 bases in unitig 39, Fig. 2b, Supplementary Table 16), 233 the internal regions contained 31 and 42 protein-coding genes, respectively, and the two clusters 234 were located on reverse strands from each other (Fig. 2a, Supplementary Table 16). Since all rDNA 235 copies are located over 28 kb away from the edge of each contig (Fig. 2a, Supplementary Table 16), 236 the lack of TRSs is unlikely to be an artifact derived from an assembly problem often caused by 237 highly repetitive sequences.

238 The lack of tandem rDNA structure was also supported by mapping our PacBio reads to 239 RIR17 and searching for rDNA on JGI v2.0 assemblies. BWA-MEM⁵¹ mapping showed multiple 240 PacBio reads across the 5' non-coding region, 48S rDNA and 3' non-coding regions of each rDNA 241 contig (Fig. 3a, Supplementary Fig. 4). Because our PacBio analysis directly sequenced the DNA 242 molecules in AMF, this syntenic structure is not due to chimeric fragments from DNA amplification 243 but reflects the natural sequence. The 5' and 3' non-coding regions of each rDNA have sequences 244 that are not similar other than the highly similar 5' regions on c62-1 and c62-2 (Fig. 3b and 245 Supplementary Fig. 4), negating the possibility of mapping confusion due to sequence similarity. We 246 reproducibly obtained the PacBio reads passing the rDNA regions from our three PacBio datasets, 247 which had been constructed from different spore suspensions. Furthermore, our rDNA searching by 248 RNAmmer detected a non-tandem 48S rDNA region from three JGI v2.0 scaffolds (Fig. 3c, 249 Supplementary Fig. 5 and Supplementary Table 18). Although the seven rDNAs cannot be 250 reconstructed from JGI v2.0, two partial rDNA sequences on JGI v2.0 had corresponding down- or 251 upstream sequences that matched our RIR17 rDNAs (Supplementary Fig. 5, and Supplementary 252 Table 18), indicating that our assembly around the rDNA genes is consistent with previous 253 assemblies.

254 We then examined polymorphism among the 45S rDNA clusters on RIR17. rDNA 255 heterogeneity has been reported in various AMF species, including DAOM-181602^{13,17,25,29}. 256 However, the distribution and degree of the variation among the rDNA paralogs were unclear. 257 Pairwise comparisons of the ten rDNA copies detected 27.3 indels and 106.1 sequence variants with 258 98.18% identity on average (Supplementary Tables 19 and 20), whereas the sequences of rDNA 259 clusters at c311-1 and c52-1 were identical. Polymorphisms were distributed unevenly throughout 260 the rDNA; percent identities were 99.91% in 18S rDNA, 97.93% in 28S rDNA, 96.65% in 5.8S 261 rDNA, 93.45% in ITS1, and 90.28% in ITS2 (Fig. 4, Supplementary Tables 19 and 20). The number 262 of polymorphic sites in R. irregularis rDNAs reached 4.07 positions per 100 bases, much higher 263 than in other fungi, which have polymorphic sites at 0.04-1.97 positions per 100 bases (Table 2). 264 The rDNA polymorphisms observed in RIR17 covered most of the polymorphisms previously 265 reported in this species (Fig. 5), providing incentive to review the molecular ecology of AMF. The 266 degree of intragenomic variation was not high enough to disrupt species-level identification but was 267 sufficient to cause erroneous identification of R. irregularis strains (Fig. 5). These findings pose a 268 caution that previous studies on geographic distribution⁸, species identification²⁸, and evolutionary 269 processes of AMF assuming rDNA homogeneity require reevaluation considering the high-level 270 intra-genomic heterogeneity of rDNA sequences in AMFs.

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272 A model for the relaxation of rDNA homogeneity in *R. irregularis*

The revealed non-tandem structure of AMF rDNA led to a model for the mechanism responsible for its intragenomic heterogeneity. Pawlowska and Taylor (2004) predicted that rDNA

275 heterogeneity is caused by relaxation of "concerted rDNA evolution" in Glomerales including 276 Rhizophagus²⁵. However, details of the "relaxation" have been unclear. Here, we propose a 277 hypothetical mechanism: the loss of TRSs precludes the presence of DNA conformations associated 278 with rDNA amplification and the maintenance of its homogeneity. The standard model of "concerted 279 rDNA evolution" needs two or more tandemly repeated rDNA segments because the rDNA 280 duplicates using tandemly repeated rDNAs as binding sites and templates for replication 281 (Supplementary Fig. 1c)⁵². Although non-tandem rDNAs are rare in eukaryotes, this trend of heterogeneity in non-tandem rDNAs has been detected by laboratory systems as well as in wild 282 283 organisms; Arabidopsis thaliana has one pseudogenic rDNA (lacking 270 bases of an important 284 helix as rRNA) besides the main tandem repeat rDNA arrays^{53,54}, and the lack of rDNA tandem repeats in malaria-causing *Plasmodium* parasites^{49,55} indicates intragenomic rDNA polymorphisms. 285 286 These observations support our hypothesis that rDNA heterogeneity in AMF is related to their lack 287 of TRSs. AMF species may not amplify their rDNA by the general eukaryotic rDNA amplification 288 system (USCR), which may increase their rDNA heterogeneity.

289 On the other hand, our phylogenetic analysis suggests that AMF has a system to maintain 290 weak similarity among the paralogs without TRSs. Previously observed rDNA heterogeneity in 291 Glomerales suggests that concerted evolution was relaxed before the diversification of *Rhizophagus* 292 species^{25,29}. When the observed ten rDNAs duplicated before speciation and evolved independently, 293 each of the duplicated genes formed a clade with orthologs in other species. However, we found no 294 orthologous rDNA genes from other Rhizophagus species (Fig. 5). Our tree suggests that the 295 observed rDNAs in R. irregularis either expanded or were assimilated after speciation. One 296 hypothetical mechanism that would cause this similarity is homologous recombination via 297 "synthesis-dependent strand annealing" (Supplementary Fig. 6)⁵⁶. This conserved system to repair 298 double-strand breaks (DSBs) results in non-crossover recombination and gene conversion wherein 299 nonreciprocal genetic transfer occurs between two homologous sequences (Supplementary Fig. 6). 300 Decreases in divergence by gene conversion are widely observed in duplicated genes. RIR17 301 showed that two rDNA pairs on the same contigs (c39-1 and c39-2, c62-1 and c62-2) had higher 302 similarity than other paralogs (Fig. 4c). This similarity may be caused by the high gene conversion 303 rate between these loci.

304 Our model raises a new question about the mechanism that maintains the number of rDNAs 305 without gene duplication by USCR. Even if rDNA lacks TRSs, crossover recombination and single-306 strand annealing delete paralogous genes. Observed inverted repeat structures between rDNAs in 307 proximity may contribute to inhibiting "single-strand annealing" between them and prevent copy 308 number reduction. Plastidial rDNAs of land plants also make inverted repeat structures and conserve 309 two rDNA copies on their plastidial DNA. Another probable system is the suppression of crossover 310 recombination by the limitations of meiosis. When Holliday junctions dissociate without crossover, 311 DSBs are repaired without gene number reduction. The majority of these crossovers arise during 312 meiosis in eukaryotes⁵⁶, and sexual reproduction had never observed in AMF. AMF species may 313 keep their rDNA copy number by asexual spore-making and the rarity of their meiotic cell division.

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315 **RNA-level impact and probable biological significances of non-**

316 tandemly repeated rDNAs

317 To confirm the transcriptional activity of each rDNA, we conducted total-RNA-Seq (RNA 318 sequencing without poly-A tail selection. See material & method section.). Illumina sequencing of a 319 modified library for rRNA sequencing ("Rir RNA rRNA" in Supplementary Table 2) produced 320 18,889,290 reads (read length = 100-301 bases) from DAOM-181602. We mapped the reads to all 321 gene models from RIR17 (43,675 protein-encoding isoforms and ten 48S rDNA paralogs) and 322 estimated the expression levels of each gene by eXpress software⁵⁷. All rDNA paralogs had over 323 5,000 FPKMs (Fragments Per Kilobase of exon per Million mapped fragments) (Table 3), and 324 multiple reads were matched to the specific region of each paralog, indicating that the ten rDNA

copies are transcriptionally active. In general, eukaryotes silence a part of the rDNA copies⁵⁸, and
some eukaryotes change the transcribed rRNA sequences by "RNA editing"⁵⁹. These editing and
silencing processes were not detected in the AMF, and the rRNA were as polymorphic as the rDNA.
These results show that DAOM-181602 has multiple types of ribosomes, each containing different
rRNAs. Additionally, we detected highly duplicated ribosomal protein genes (e.g., ribosomal protein
S17/S11) (Supplementary Tables 6 and 21) and tRNA genes, indicating unknown amino acid
isotypes, which may also account for the heterogeneity of ribosomes (Supplementary Table 22).

332 The evolutionary significance of the of non-tandemly repeated heterogeneous rDNAs is 333 unclear. One of the probable factors is a reduction in the need to maintain numerous rDNAs in a 334 genome. As described in the above sections, the AMF rDNA copy number suggests a system that 335 efficiently produces rRNA from a few rDNAs, and the inverted repeats structure of rDNAs and 336 asexual spore reproduction will also reduce the deletion rate of rDNAs. AMF may thus no longer 337 need to rapidly amplify rDNA copies using TRSs, and the slowed replacement rate of rDNA may 338 then cause the heterogeneity as a side effect. Another possibly significant effect is the enhancement 339 of phenotypic plasticity by ribosomal heterogeneity (Fig. 6). Recent studies have started to reveal 340 that various eukaryotes (e.g., yeast, mice, and Arabidopsis) produce heterogeneous ribosomes and 341 subsequently alter phenotypes via proteins translated by particular ribosomes⁶⁰. Accelerated 342 accumulation of AMF rDNA mutations by the lack of TRSs may lead to functional variety in 343 produced ribosomes and increases in the rate of adaptation by different translation activities within 344 the same species. Although the functional effects of observed rDNA mutations remain to be 345 determined, the middle area of our 28S rDNA (4,450-4,500 bases on c62-1) had a higher mutation 346 rate than ITS regions (Fig. 4a). Because the ITS regions (encoding non-functional RNA) vary under 347 neutral mutation rates, the accumulated variants in the middle-28S region may have functional 348 effects favored by natural selection (via diversifying selection). This region is thus a useful target for 349 the future functional analyses of AMF rRNA.

350 AMF species are similar to the malaria parasite in that they both have heterogeneous non-351 tandem rDNAs and infect distantly related host species⁴⁹. In the malaria parasite, changes in the 352 ribosome properties depend on the host (human or mosquito), which is likely able to alter the rate of 353 translation, either globally or of specific messenger RNAs, thereby changing the rate of cell growth 354 or altering patterns of cell development⁴⁹. The relationship between the diversity of host organisms 355 and rDNA polymorphisms will be an important area for further research. The phenotypic plasticity 356 caused by heterogeneous translation machinery may allow adaptation for various host species having 357 slightly different symbiotic systems. Previous studies have proposed that the heterokaryosity in 358 AMF species drives variable genetic combinations of mycelia in the absence of sexual 359 recombination⁶¹. Recent genomic studies, furthermore, discovered signatures of sexual reproduction 360 within the dikaryon-like stage 16,62 . Our hypothesis does not exclude current theories for the genetic 361 and phenotypic plasticity of AMF species (heterokaryosis and sexual reproduction) but proposes a 362 multilayered diversification mechanism leading to their widespread distribution.

364 Materials and Methods 365 PacBio-based assembling

366 **DNA preparation**

367 The DNA sample for the PacBio and Illumina sequencing was extracted from a commercial 368 strain of R. irregularis DAOM-181602 (MYCORISE® ASP, Premier Tech Biotechnologies, 369 Canada). The DNA extraction followed the method of Fulton et al., 1995⁶³ with some modifications 370 described below. Purchased spore suspensions (including approximately 1,000,000 spores) were 371 centrifuged (4500 rpm, 20 min), and washed three times with distilled water. Precipitated spores 372 were frozen with liquid nitrogen, ground with pestle, and dispersed in extraction buffer (100 mM 373 Tris-HCl pH 8.0, 20 mM EDTA, 0.75% sarkosyl, 0.1% PVP, 0.75% cetyl trimethylammonium 374 bromide (CTAB), 0.13 M sorbitol, 0.75 M NaCl, and 0.1 mg/ml proteinase K). After incubation at 375 37 °C for 10 min, the aqueous phase was centrifuged (15000 rpm, 4 min), and the pellet was 376 discarded. An equal volume of phenol/chloroform (1:1, vol/vol) was added, and the sample was 377 gently mixed and centrifuged (15000 rpm, 2 min). The aqueous phase was collected, and an equal 378 volume of chloroform was added to the sample, which was then mixed and centrifuged (15000 rpm, 379 2 min). The aqueous phase was collected again, and 1/10 vol of sodium acetate and 0.7 vol of 380 isopropanol were added. The sample was then mixed and centrifuged (12000 rpm, 20 min). The 381 resulting pellet was washed twice with 70% EtOH and eluted with TE buffer. Extracted DNA was 382 purified with Genomic-tip (Qiagen, Germany) following the manufacturer's instructions.

383 PacBio sequencing

Long-read sequences were generated with a PacBio RS II sequencer (Pacific Biosciences, Menlo Park, CA, USA) using a DNA/Polymerase Binding Kit P6 v2 (Pacific Biosciences) and a DNA Sequencing Reagent Kit 4.0 (Pacific Biosciences). The library was prepared according to the 20-kb Template Preparation Using BluePippin[™] Size-Selection System (Sage Science, MA, USA). A total sequence of 11.7 Gb in 955,841 reads (76× coverage of the genome, assuming a genome size of 154 Mb) was obtained from 29 SMRT cells (Supplementary Table 2). The N50 length of the raw reads was 13,107 bases.

391 PacBio-based genome assembly

392 The R. irregularis genome was assembled using the RS HGAP Assembly.3 protocol for 393 assembly and Quiver for genome polishing in SMRT Analysis v2.3.0 (Pacific Biosciences). The 394 procedure consisted of three parts, involving (1) generation of preassembled reads with improved 395 consensus accuracy; (2) assembly of the genome through overlap consensus accuracy using Celera 396 Assembler; and (3) one round of genome polishing with Quiver. For HGAP, the following 397 parameters were used: PreAssembler Filter v1 (minimum subread length = 500 bases, minimum 398 polymerase read quality = 0.80, minimum polymerase read length = 100 bases); PreAssembler v2 399 (minimum seed length = 6,000 bases, number of seed read chunks = 6, alignment candidates per 400 chunk = 10, total alignment candidates = 24, minimum coverage for correction = 6, and BLASR 401 options = 'noSplitSubreads, minReadLength = 200, maxScore = 1,000, and maxLCPLength = 16'); 402 AssembleUnitig v1 (genome size = 150 Mb, target coverage = 25, overlapper error rate = 0.06, 403 overlapper min length = 40 bases and overlapper k-mer = 14); and BLASR v1 mapping of reads for 404 genome polishing with Quiver (maximum divergence = 30, minimum anchor size = 12). Assembly 405 polishing with PacBio reads was carried out with Quiver using only unambiguously mapped reads. 406 The statistics of the PacBio-only assembly set and previously sequenced data (Lin14, JGI v1.0, 407 JGI v2.0) were evaluated using QUAST ver. 4.3⁶⁴. The percentage of genome coverage was 408 estimated assuming the genome size to be 154 Mb based on Tisserant et al¹⁷.

409 Error correction with HiSeq data and identification of host plant contamination

410 After polishing using Illumina data, we eliminated the sequences derived from contaminated 411 DNAs during the sample preparation. BLASTn search of the polished assemblies against the 412 "refseq genomic" database detected nine assemblies showing similarity with sequences from carrot 413 (BLAST ver. 2.2.31+, query coverage per subject >95%, percentages of identical matches >90%, bit 414 score > 1000) (Supplementary Table 2), which might be used as a host plant by the manufacturer for 415 the cultivation of *R. irregularis* samples. After elimination of the nine contaminated contigs, we 416 submitted the assemblies to the DDBJ as whole-genome shotgun sequence data (RIR17) of R. 417 irregularis DAOM-181602 (BDIQ01).

418 Genomic alignment with previous genome assemblies

The quality of our genome assembly was evaluated by alignment with previously available *R*. *irregularis* DAOM-181602 genome assemblies. A one-by-one genome alignment was constructed by MUMmer ver. 4.0.0beta2⁶⁵ between RIR17, JGI_v2.0, Lin14, and JGI_v1.0 assemblies. Each genome set was aligned by the nucmer function in MUMmer, and the statistics of the alignments were extracted by the dnadiff wrapper with the default setting.

424

425 Gene prediction and annotation

426 *De novo* repeat motifs were identified using RepeatModeler ver. 1.0.8, which combines 427 RECON and RepeatScout programs³⁷. Based on the identified motif, the repetitive region in the 428 assemblies was masked with RepeatMasker ver. $4.0.5^{37}$. We used the default parameters for the 429 identification and the masking.

430 For the gene models constructed from RIR17 assemblies, standard RNA-Seq data were 431 obtained from R. irregularis spores and hyphae. The RNA was extracted with an RNeasy Plant Mini 432 kit (Qiagen) after incubation of the purchased spores (MYCORISE® ASP) in a minimum nutrient 433 medium for one day. An Illumina RNA-Seq library was constructed with a TruSeq Stranded mRNA 434 Library prep kit (Illumina). The library was sequenced (101 bases from each end) on a HiSeq 1500 435 platform (Illumina). A total of 16,122,964 raw reads (3.2 Gb) were obtained from the library 436 (Supplementary Table 2). After filtering low-quality and adapter sequences, RNA-Seq data were 437 mapped to RIR17 assemblies with TopHat ver. 2.1.1⁶⁶ with the default setting.

438 Then, the RIR17 assemblies were processed through the RNA-Seq-based gene model 439 construction pipeline using AUGUSTUS ver. 3.2.1 software⁶⁷. We constructed R. irregularis-440 specific probabilistic models of the gene structure based on 495 manually constructed gene models 441 from the longest "unitig 392" sequence in RIR17. Manual gene models were made with ab initio 442 AUGUSTUS analysis based on probabilistic models for *Rhizopus oryzae* and by manual refinement 443 using the homology data with already-known genes and mapped RNA-Seq data. Then, with the 444 trained probabilistic models and the intron-hints data from the mapped RNA-Seq read, 37,639 445 optimal gene models were constructed using the AUGUSTUS pipeline. We then confirmed whether 446 the AUGUSTUS pipeline overlooked the called genes in previous genome studies. We mapped all 447 transcript sequences obtained from previous gene modeling on Lin14 and JGI v1.0 against our 448 RIR17 genomic sequences with Exonerate⁶⁸ (ver. 2.2.0, option --model est2genome --bestn 1), 449 resulting in the recruitment of 3,933 overlooked genes. The completeness of the constructed gene 450 model was evaluated with BUSCO ver. 2.039. The BUSCO analysis used the "Fungi odb9" gene set 451 (http://buscodev.ezlab.org/datasets/fungiodb9.tar.gz) as a benchmark and employed the "-m 452 proteins" option to analyze the preconstructed protein data without the *ab initio* gene modeling step.

The confidences of the obtained 41,572 gene models were estimated based on 1) RNA-Seq expression support, 2) homology evidence, and 3) protein motif evidence. For the calculation of 455 gene expression levels, we mapped our "Rir_RNA_SS" data and 32 RNA-Seq data submitted to the 456 sequence read archive (SRA) database (24 data sets from DRP002784 and 8 data sets from 457 DRP003319) and calculated the gene expression levels (FPKM) using FeatureCounts⁶⁹ with the 458 default setting (Supplementary Table 6). Homology with previously known genes was determined 459 by BLAST searches against the orthoDB (odb9) (Supplementary Tables 6 and 21). The protein motif 460 was searched using Pfam analysis in InterProScan ver. 5.23-62.0⁷⁰ (Supplementary Table 6).

461 Constructed gene models were annotated by several *in-silico* searches. Gene functions were 462 predicted based on BLASTp (Database = nr, RefSeq and UniProt), and Pfam in InterProScan 463 (Supplementary Table 6). We manually selected the descriptive nomenclatures from those four 464 searches and submitted to the DDBJ. Orthologous relationships were classified with Orthofinder 465 (ver. 1.1.2)⁷¹, and rapidity expanded/contracted families were analyzed with CAFE (ver. 4.1)⁷² from Orthofinder results. Phylogenetic trees for the CAFE analysis were constructed with IQ-tree (ver. 466 467 $(1.6.1)^{73}$ for maximum likelihood (ML) analysis and r8s (v1.81) for a conversion for an ultrametric 468 tree. An ML tree was made from 159 single-copy genes from the Orthofinder results (Supplementary 469 Table 6) and was converted to an ultrametric tree based on the divergence times of AMF-470 Mortierellales (460 Myr)²⁷ and Deuterostomia-Protostomia (550 Myr)⁷⁴. Overlapping genes with 471 TEs were extracted from AUGUSTUS and RepeatMasker results using bedtools (ver. 2.26.0, 472 "bedtools intersect" with -wa option)⁷⁵.

473 The MACG (missing ascomycete core gene) orthologs were sought using BLAST with the "-474 evalue 0.0001" option, and the reference sequences for the MACG search were selected from protein 475 data from an S288C reference in the Saccharomyces genome data base (SGD) (Supplementary Table 476 15). Genes involved in the degradation of plant cell walls were sought by BLAST with the same 477 settings as the MACG search, and the reference sequences were selected from Aspergillus niger 478 CBS 513.88 data in GenBank based on CAZY classification (Supplementary Table 15). Other gene 479 annotations based on the CAZy database were performed with the dbCAN HMMs 6.0 web service⁷⁶ 480 (Supplementary Table 6).

481

482 Detection of Ribosomal DNA and intragenomic

483 polymorphisms

Ribosomal DNA regions were detected by RNAmmer ver. 1.2⁷⁷ from whole RIR17
assemblies and were manually refined based on the MAFFT v7.294b⁷⁸ alignment to the 48S rRNA
in *Saccharomyces cerevisiae* S288C. The genomic positions of rDNAs were visualized with Python
ver. 3.4.0 (BasicChromosome ver. 1.68, and GenomeDiagram ver. 0.2 modules) (Fig. 2a).

488 The number of rDNA paralogs in the genome was estimated by mean depth of coverage. We 489 masked repetitive regions (based on RepeatModeler analysis) and all rDNA regions on RIR17 490 except one rDNA copy (c62-1). Then, trimmed R1 Illumina reads from "Rir DNA PE180" library 491 were mapped to the repeat-masked RIR17 using bowtie2 ver. 2.2.9⁷⁹. The coverage depth of the 492 rDNA region and 243 single-copy BUSCOs were obtained using bedtools ("bedtools coverage" 493 command with -d option), and the statistics of each region were calculated and visualized by R 494 software ver. 3.4.2 with the ggplot2 library (Fig. 2b). To prevent copy number estimation from depth 495 fluctuation due to the intragenomic heterogeneity, we confirmed the coverage depth using the 496 consensus sequences of all ten rDNA paralogs; the joined Illumina reads (from "Rir DNA PE180" 497 library) were mapped back to a consensus rDNA sequences and ten single-copy BUSCO genes from 498 RIR17, and the depth of coverages was then counted by bedtools (genomeCoverageBed) 499 (Supplementary Table 17).

500 The syntenic structure around rDNA genes was confirmed by the mapping of PacBio raw 501 reads and comparison with JGI v2.0 assemblies. All of the "filtered-subreads" from SMART 502 Analysis software were mapped to RIR17 assemblies by BWA-MEM (ver. 0.7.15-r1140) with the "-503 x pacbio" option. Mapped reads were visualized with Integrative Genomics Viewer (ver. 2.4), and 504 the reads covering the rDNA regions were selected by eye. Alignment between JGI v2.0 and RIR17 505 was done by a combination of MUMmer, LASTz (ver. 1.04.00), and AliTV⁸⁰ (ver. 1.0.4) software. 506 JGI v2.0 scaffolds having regions corresponding with RIR17 sequences were selected by the 507 nucmer and delta-filter (with -1 option) functions in MUMmer. Then, we extracted the JGI v2.0 508 scaffolds corresponding to RIR17 contigs with rDNAs ("unitig 311", " 312", " 35", " 356", " 4", 509 and "_52"). Selected scaffolds were aligned to the corresponding RIR17 contigs by ality.pl scripts 510 (with "alignment: program: lastz" and "--ambiguous=n" settings) and alitv-filter.pl (with "--min-link-511 identity 80" and "--min-link-length 10000" option) in the AliTV package and visualized with the 512 AliTV web service (http://alitvteam.github.io/AliTV/d3/AliTV.html).

The difference among the rDNA paralogs was calculated from the aligned sequences by MAFFT ver. 7.309 (options: --localpair, --op 5, --ep 3, --maxiterate 1000) using the pairwise comparison with CLC Main Workbench 7.8.1 (Qiagen). The mutation type was called by eye from the alignment, and we chose the c62-1 paralog as a reference sequence for mutation calling (Fig. 4a). Phylogenetic trees (Figs. 4c and 5) were constructed from the MAFFT alignment by the neighborjoining method with MEGA⁸¹ ver. 7.0.21 under the maximum composite likelihood model and were tested for robustness by bootstrapping (500 pseudoreplicates).

521 Heterogeneity of translation machineries

522 The expression levels of the rDNA paralogs were examined with modified Illumina 523 sequencing of R. irregularis spores and hyphae. Total RNA was extracted with an RNeasy Plant 524 Mini kit (Qiagen) after the incubation of the MYCORISE® spores in a minimum nutrient medium 525 for seven days. An Illumina RNA-Seq library was constructed with a TruSeq Stranded mRNA 526 Library prep kit (Illumina). To skip the poly-A tailing selection step in the library construction, we 527 started from the "fragmentation step" of the standard manufacturer's instructions. The library was 528 sequenced (301 bases from each end) on a MiSeq platform (Illumina). A total of 16,122,964 raw 529 reads (3.2 Gb) were obtained from the library (Supplementary Table 2). After filtering low-quality 530 and adapter sequences, RNA-seq data were mapped to the RIR17 assembly with TopHat with the 531 default settings. Fragments Per Kilobase of exon per Million mapped fragments (FPKMs) for each 532 gene were calculated with eXpress ver. 1.5.1 with the "--no-bias-correct" option. Transfer RNAs 533 were identified with tRNAscan-SE⁸² ver. 1.3.1.

534

535 Data Availability

Raw reads, genome assemblies, and annotations were deposited at INSDC under the
accessions as follows; Sequence read archive: DRA004849, DRA004878, DRA004889,
DRA004835, DRA005204, and DRA006039; Whole genome assembly: BDIQ01000001BDIQ01000210; Annotations: GBC10881-GBC54553. All the other data generated or analyzed
during this study are included in this published article and its Supplementary information.

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544 Author contributions

545 T.M., S.S., M.K., conceived of and designed the experiments; T.M., Y.K., H.K., N.T., K.Y., and 546 T.B. performed the experiments; T.M., N.O., S.S., and T.B. analyzed the data, T.M., Y.K., H.K.,

547 K.Y., T.B., S.S., and M.K. wrote the manuscript.

548

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555

556 **Competing Interests**

⁵⁵⁷ The authors declare no conflicts of interest associated with this manuscript.

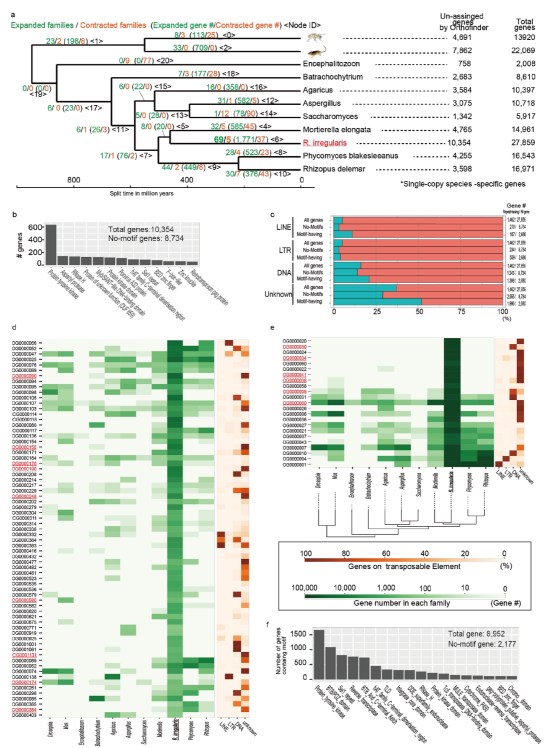


Fig. 1 Gene inflation in R. irregularis

a. Rapidly expanded/contracted ortholog groups based on CAFE analysis. Total gene number of analyzed species and unassigned genes by Orthofinder analysis (species-specific single-copy genes) are described on the right side of the tree. b. The number of *R. irregularis*-specific single-copy genes having protein motifs. Minor motifs (<50 genes) were omitted from the Fig. (raw-data; Supplementary Table 12). c. The proportion of genes among the SSSC genes having each repeat element. d. Sixty-nine rapidly expanded orthologous groups (OGs). Green heat map shows the number of genes in each OG. Orange heat map indicates the proportion of genes with each repeat element. The OGs containing the "protein tyrosine kinase" domain are marked in red. e. Rapidly expanded OGs based on z-score analysis. The colors have the same meaning as in 1d. f. The number of rapidly expanded ortholog genes having protein motifs. Minor motifs (<100 genes) are omitted from the Fig. (raw-data; Supplementary Table 12).

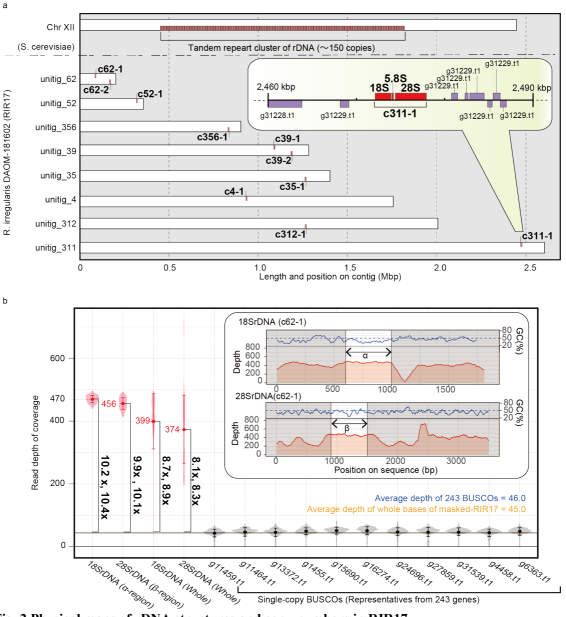


Fig. 2 Physical maps of rDNA structures and copy numbers in RIR17

a. Distribution of *R. irregularis* rDNA units in the genome. Each 48S rDNA unit is represented as a red box. For comparison, rDNA clusters on *Saccharomyces cerevisiae* chromosome XII are shown^{83,84}. Inset is a magnified view of a 48S rDNA unit (c311-1) with nearby protein-encoding genes (purple boxes). Genes encoded by the plus-strand genome are depicted on the top side, and those encoded by the minus strand are shown on the bottom side. **b. Copy number of rDNA in DAOM-181602 based on the read depth of coverage**. Averages of the "read depth of coverage" are represented as dots and with italic labels. Error bars and violin plots show standard deviations and normalized coverage distribution. The depths of rDNA regions are marked in red. For comparison, the data from representative single-copy BUSCO genes on RIR17 are shown in black. The mean depth of means from 243 BUSCOs is marked with a horizontal blue line, and the mean depth of all RIR17 bases is marked with an orange line. The changes in the depth of rDNA regions are in vertical bold labels and square brackets. rDNA regions adapted for the copy number estimation (α - and β -regions) are marked in the inset with the depth of coverage and the GC content of each sequence position.

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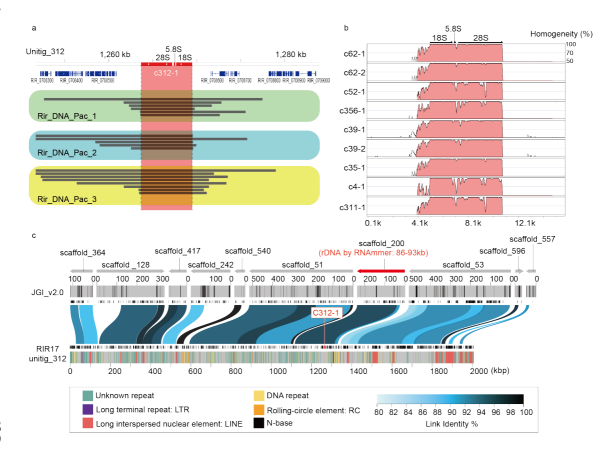


Fig. 3 Evidence for the lacking of tandem repeat structures of rDNA

a. Mapped PacBio read for the rDNA regions on unitig_312 contig in RIR17. The top bar and tick marks indicate sequence positions on the contig. The rDNA region (c312-1) is indicated in red. Blue boxes show the predicted protein-coding genes. The mapped read was indicated black bar, and reads from different DNA samples and libraries (Supplementary Table 2) boxed with green, light-blue and yellow colors, in each. Mapped reads for the other rDNA regions were summarized in Supplementary Fig. 4. b. Sequence similarity of c312-1 rDNAs with other rDNA regions on RIR17. The 5 kb upstream and downstream sequences of each rDNA region are separated from each contig. Alignment and similarity were calculated with mVISTA⁸⁵. Red color shows the sequence regions with similarity over the threshold (>70% similarity for 100b). c. Positions and identities of JGI_v2.0 scaffold aligned against unitig_312 contigs of RIR17. The top area indicates aligned scaffolds and their strands. A scaffold containing the predicted rDNA gene is marked in red. The positions of N-base on JGI_v2.0 are marked with black bars in the next line. Predicted protein-coding genes from Chen et al¹⁴. are indicated with the next black boxes. Aligned positions and their similarity are marked with blue or black bands on the next line. The area below the black boxes show the predicted genes in the present study. Repetitive regions are marked with colored lines on the bottom band. Types of repetitive elements and the legend of similarity coloration are indicated in the bottom box.

 $\begin{array}{c} 588\\ 5590\\ 591\\ 592\\ 593\\ 595\\ 596\\ 597\\ 598\\ 5990\\ 601\\ 602\\ 603 \end{array}$

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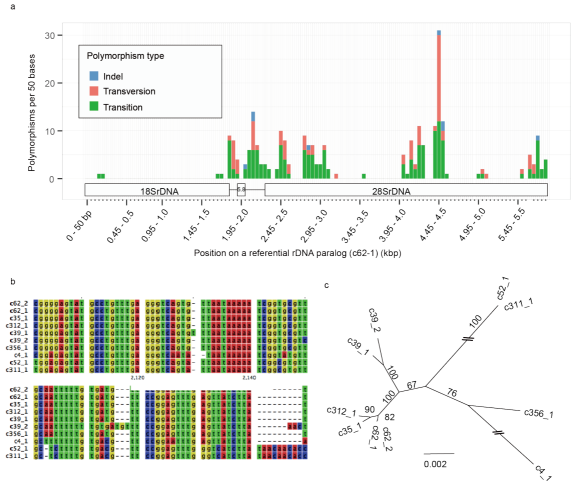
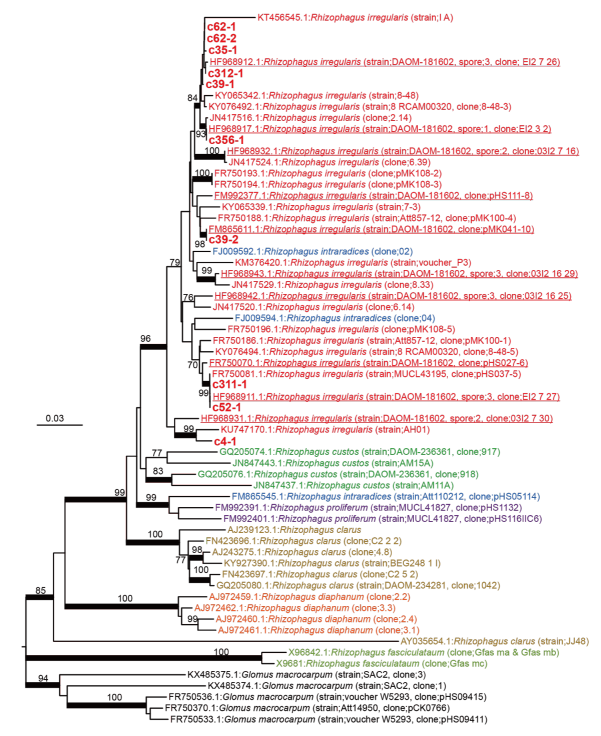


Fig. 4 Polymorphisms of 48S rDNA paralogs in RIR17

a. The distribution of rDNA sequence variants within the 48S rDNA of RIR17. The position and types of
polymorphisms were called based on the paralog c62-1. b. Alignment of a heterogeneous region among the 48S
rDNA paralogs. Partial sequences of MAFFT-aligned 48S rDNAs (corresponding to 2,049-2,136 bases positions on
c62-1). c. Neighbor-joining tree for phylogenetic relationships among the ten rDNA paralogs based on 5,847
aligned positions. Bootstrap values are described at each node.



620

621 Fig. 5 NJ tree based on 586 positions of 48S rDNA.

Partial 18S, ITS1, 5.8S, ITS2 and partial 28S rDNAs were used. The ten rDNA paralogs from RIR17 and 58 *Rhizophagus* sequences from the DDBJ were chosen as operational taxonomic units (OTUs). The 58 *Rhizophagus* sequences were selected from 329 OTUs in the DDBJ (209 OTUs for DAOM-181602, 57 OTUs for other *R. irregularis* strains, and 63 OTUs for other *Rhizophagus* species) using CD-Hit clustering (-c 0.98 -n 5). Five *Glomus* sequences were used as outgroup OTUs. Red underlined OTUs are sequences from *R. irregularis* DAOM-181602, and other red OTUs are data from other strains of *R. irregularis*. Nodes supported by over 80 bootstrap values are marked by a bold line. All *R. irregularis* OTUs made a single clade with *Rhizophagus intraradices* that is a morphologically non-distinct sister group of *R. irregularis*.

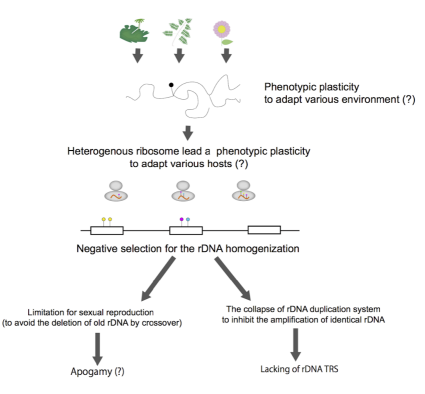


Fig. 6 Hypothetical model for the evolution of unique rDNAs/rRNAs in AMF

Evolutionary model for the lack of TRSs in AMF and its sequence heterogeneity. The various environmental conditions (e.g., various host species) may lead to the evolution of phenotypic plasticity via multiple types of ribosomes in AMF. If the rDNA is exposed to disruptive selection, rDNA duplication by TRSs and USCR may be nonadaptive because the duplication of particular rDNA types reduces the variety of rDNA types. Sexual reproduction, combined with crossover recombination, may also be limited to inhibit the reduction of mutated rDNA.

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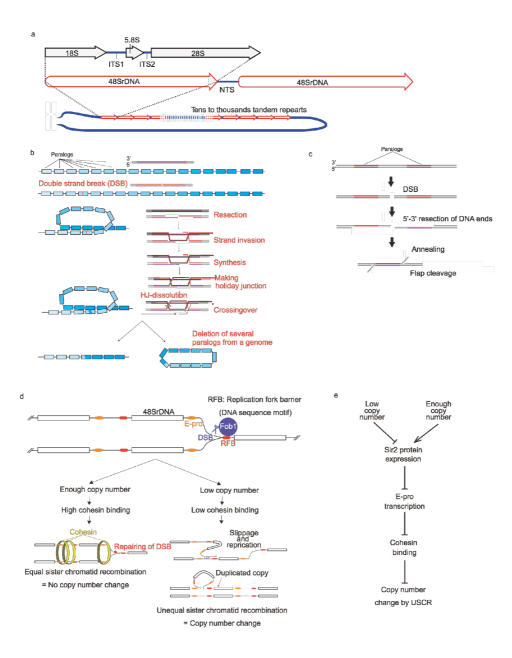
640 Table 1 Assembly statistics of <i>R. irregularis</i> genome

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	RIR17
Accession number	BDIQ01000000
Predicted genome size by flow cytometry	154 Mb
Total length of contigs (% of genome)	149,750,837 bases (97%)
# contigs	210
# N bases	0
Longest contig (bp)	5,727,599
Contig N50 (bases)	2,308,146
L50	23
GC %	27.9%
CEGMA completeness for genome contigs	98.4%
# of predicted genes	41,572
BUSCO completeness for gene models (DB; fungi_odb9)	94.1% (273/290)
Complete single copy	83.8% (243/290)
Complete duplicated	10.3% (30/290)
Fragmented	3.8% (11/290)
Missing	2.1% (6/290)

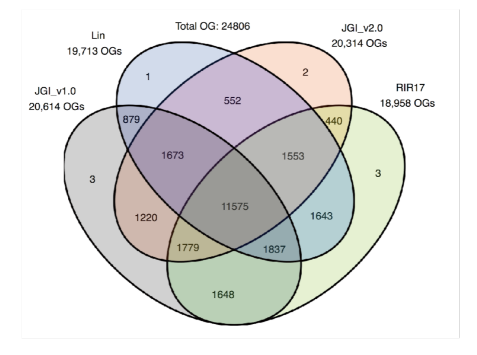
Species	# polymorphic sites	Repeat unit length (bp)	# of units in genome	# of polymorphic sites/100 bases
Rhizophagus irregularis	238	5847	10	4.07
Rhizophagus irregularis ¹³	38	1563	-	2.43
Ashbya gossypii ³⁰	3	8147	50	0.04
Saccharomyces paradoxus ³⁰	13	9103	180	0.14
Saccharomyces cerevisiae ³⁰	4	9081	150	0.04
Aspergillus nidulans ³⁰	11	7651	45	0.14
Cryptococcus neoformans ³⁰	37	8082	55	0.46
Phoma exigua var. exigua ⁸⁶	27	1672	-	1.61
Mycosphaerella punctiformis ⁸⁶	26	1669	-	1.56
Teratosphaeria microspora ⁸⁶	16	1671	-	0.96
Davidiella tassiana ⁸⁶	33	1672	-	1.97

645 <u>Table 2 Numbers of intragenomic polymorphic sites in fungal rDNAs</u>



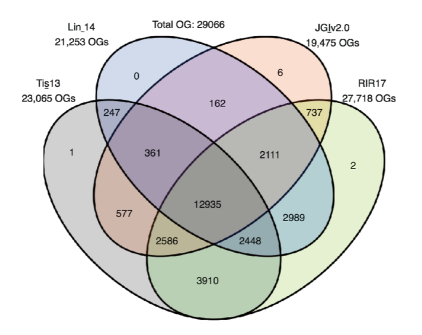
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Supplementary Fig. 1 General concerted evolution of eukaryotic rDNA a. A general structure of eukaryotic rDNA clusters²⁰. b. Deletion of homologous genes by crossover recombination^{33,56}. c. Deletion of homologous genes by single-strand annealing³⁴. d. A model of the rDNA number 656 657 maintenance system³². e. Copy number-controlling pathway in yeast³².



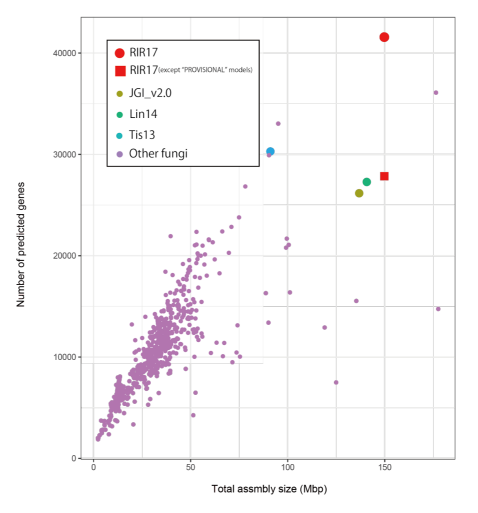
Gene models without "Provisional" models

All gene medels including "Provisional" models



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Supplementary Fig. 2 Cross comparison of *R. irregularis* DAOM-18160219 orthologous genes
 from three genomic studies and our RIR17 assemblies.

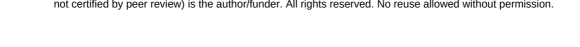


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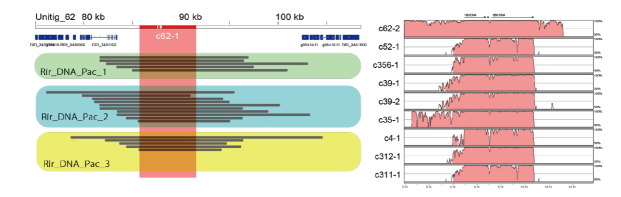
664 Supplementary Fig. 3 Total assembly size and predicted gene number in fungi.

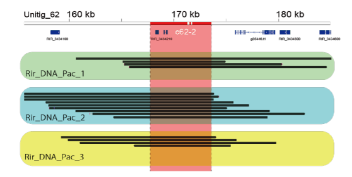
665 *Rhizophagus irregularis* genomes (RIR17 (this work) and two previously assembled genomes, JGI_v1.0 and Lin14) and 768 genomes registered in GenBank. The fungal assembly statistics were obtained from the registered information in GenBank (ftp://ftp.ncbi.nlm.nih.gov/genomes/ASSEMBLY_REPORTS/All/).

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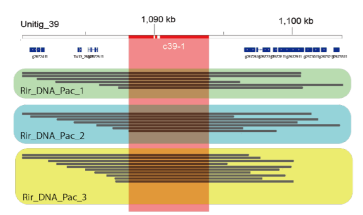


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	Rir_DNA_Pac_	g18257.H			,900	RR_SHEE	c52-1 c356-1 c39-1 c39-2	Philip Philip Ando Ando	<u> </u>	
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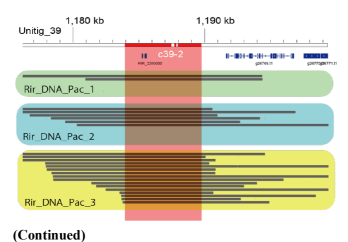


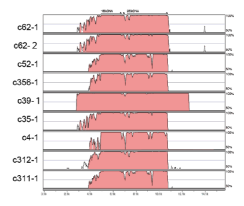


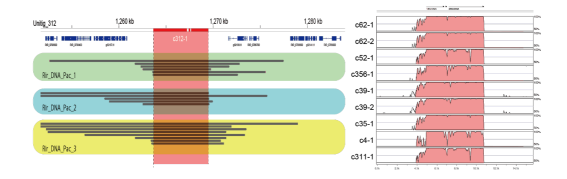
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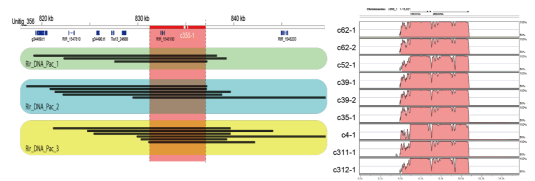


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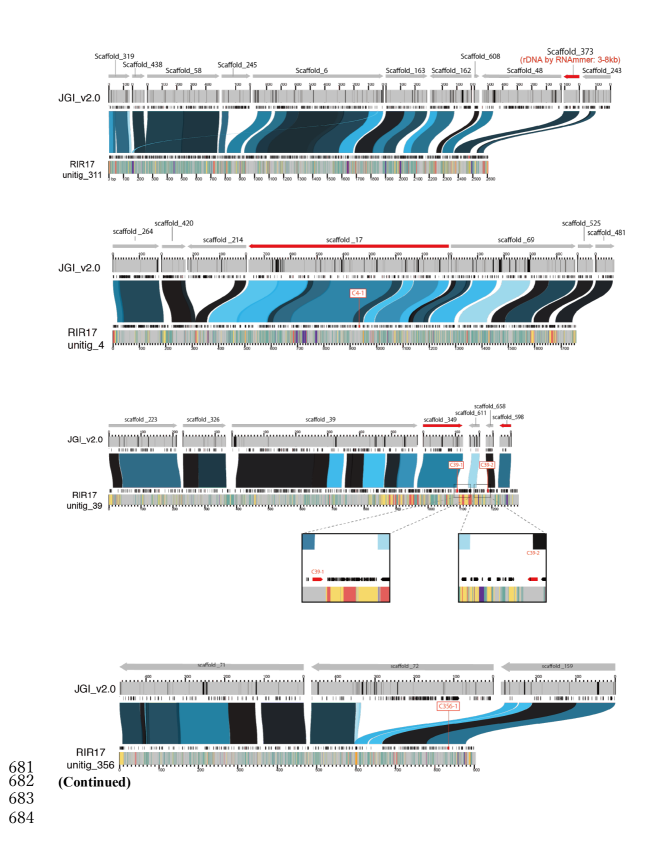


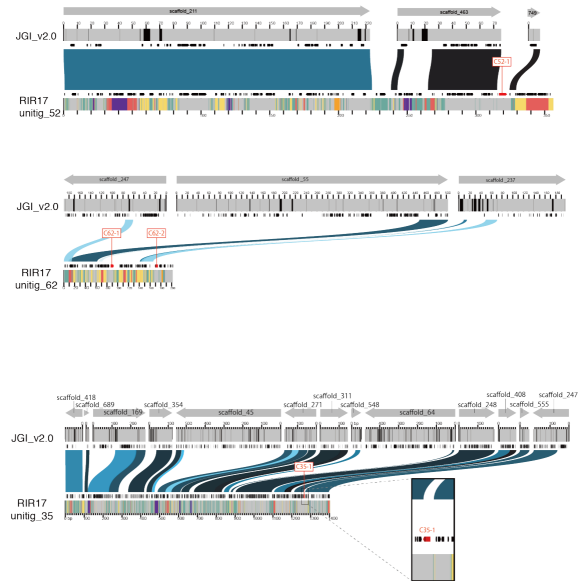


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Supplementary Fig. 4 Mapped PacBio reads of rDNA regions of RIR17 contigs and the sequence similarity of rDNAs with other rDNA regions in RIR17 678

679 The colors have the same meanings as in Fig. 3a-b.





- Figure S5 Positions and identities of JGI_v2.0 scaffold aligned against RIR17 contigs with
 rDNAs.
- 688 The colors have the same meanings as in Figure 3c.
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