

Studying Genome Heterogeneity within the Arbuscular Mycorrhizal Fungal Cytoplasm

Eva Boon, Sébastien Halary, Eric Bapteste, Mohamed Hijri

To cite this version:

Eva Boon, Sébastien Halary, Eric Bapteste, Mohamed Hijri. Studying Genome Heterogeneity within the Arbuscular Mycorrhizal Fungal Cytoplasm. Genome Biology and Evolution, 2015, 7 (2), pp.505- 521. $10.1093/gbe/evv002$. hal-01224215

HAL Id: hal-01224215 <https://hal.sorbonne-universite.fr/hal-01224215v1>

Submitted on 4 Nov 2015

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

[Distributed under a Creative Commons Attribution 4.0 International License](http://creativecommons.org/licenses/by/4.0/)

Studying Genome Heterogeneity within the Arbuscular Mycorrhizal Fungal Cytoplasm

Eva Boon^{1,‡}, Sébastien Halary¹, Eric Bapteste^{2,3,}*^{,†}, and Mohamed Hijri^{1,}*^{,†}

¹Département de Sciences Biologiques, Institut de Recherche en Biologie Végétale, Université de Montréal, Quebec, Canada

²CNRS, UMR7138, Institut de Biologie Paris-Seine, Paris, France

³Sorbonne Universités, UPMC Univ Paris 06, Institut de Biologie Paris-Seine (IBPS), Paris, France

*Corresponding author: E-mail: eric.bapteste@snv.jussieu.fr; mohamed.hijri@umontreal.ca.

z Present address: Department of Biology, Dalhousie University, Halifax, NS, Canada

[†]These authors contributed equally to this work.

Accepted: January 5, 2015

Data deposition: All raw pyrosequencing data (amplicon data for the single copy markers and WGS for the evolutionary network analyses) were deposited at the NCBI Trace Archive under Bioproject number PRJNA174749. The single copy marker allele alignments are provided in the [supplementary material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online.

Abstract

Although heterokaryons have been reported in nature, multicellular organisms are generally assumed genetically homogeneous. Here, we investigate the case of arbuscular mycorrhizal fungi (AMF) that form symbiosis with plant roots. The growth advantages they confer to their hosts are of great potential benefit to sustainable agricultural practices. However, measuring genetic diversity for these coenocytes is a major challenge: Within the same cytoplasm, AMF contain thousands of nuclei and show extremely high levels of genetic variation for some loci. The extent and physical location of polymorphism within and between AMF genomes is unclear. We used two complementary strategies to estimate genetic diversity in AMF, investigating polymorphism both on a genome scale and in putative single copy loci. First, we used data from whole-genome pyrosequencing of four AMF isolates to describe genetic diversity, based on a conservative network-based clustering approach. AMF isolates showed marked differences in genome-wide diversity patterns in comparison to a panel of control fungal genomes. This clustering approach further allowed us to provide conservative estimates of Rhizophagus spp. genomes sizes. Second, we designed new putative single copy genomic markers, which we investigated by massive parallel amplicon sequencing for two Rhizophagus irregularis and one Rhizophagus sp. isolates. Most loci showed high polymorphism, with up to 103 alleles per marker. This polymorphism could be distributed within or between nuclei. However, we argue that the Rhizophagus isolates under study might be heterokaryotic, at least for the putative single copy markers we studied. Considering that genetic information is the main resource for identification of AMF, we suggest that special attention is warranted for the study of these ecologically important organisms.

Key words: genome evolution, network analysis, genome heterogeneity, arbuscular mycorrhizal fungi, symbiosis, next generation sequencing.

Introduction

The multicellular individual is a functionally integrated assemblage of cells that share the same evolutionary fate, and can also be referred to as an organism. Even though there is no consensus on how to define "the individual," many definitions depend on genome homogeneity, that is, all cells in an individual are expected to contain the same nuclear genome (Santelices 1999). The popularity of this criterion is based on the assumption that intraorganismal genetic heterogeneity (IGH) leads to conflict within the organism and thus stands in the way of its survival. IGH can indeed be detrimental to the multicellular organism (Biesecker and Spinner 2013). However, recent reviews on IGH in nonmodel systems question the ubiquity of the genetically homogeneous organism and multiple occurrences of heterokaryosis have

© The Author(s) 2015. Published by Oxford University Press on behalf of the Society for Molecular Biology and Evolution.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License [\(http://creativecommons.org/licenses/by/4.0/\)](http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

SMBE

been reported (Santelices 1999; Pineda-Krch and Lehtila 2004; Pepper and Herron 2008; Folse and Roughgarden 2010).

A group of organisms that undoubtedly evokes questions about the defining criteria of individuality are the rootinhabiting arbuscular mycorrhizal fungi (AMF), which form their own phylum, the Glomeromycota (Schussler et al. 2001). AMF improve nutrient uptake in their host plants, and buffer the plant against abiotic and biotic stresses (Van Der Heijden and Sanders 2002). These fungi significantly increase plant growth rates, although benefits vary depending on the composition of the AMF and plant community (Van der Heijden et al. 1998). AMF are of great potential interest to agriculture, yet advances in understanding the genetics and biology of these organisms have been slow (Sanders and Croll 2010).

As obligate symbionts with a long generation time, AMF are challenging study organisms. Limited data are available for only a subset of taxa for which axenic cultures or pot cultures have been established from single spores (so-called "isolates") (Tisserant et al. 2012, 2013).

A major criticism against the possibility of heterokaryosis in AMF is that polymorphism could also be structured within nuclei, as duplicated genes (Hosny et al. 1999; Rosendahl and Stukenbrock 2004). A population genetic study of the highly polymorphic PLS gene has suggested that the observed genetic diversity (13 PLS alleles) occurred within each nucleus (Pawlowska and Taylor 2004). However, this evidence was debated: The copy number of the PLS marker used to demonstrate homokaryosis in AMF was found to be lower than its intraisolate allelic diversity (Hijri and Sanders 2005), and other hypotheses fit the data equally well (Bever and Wang 2005). The evidence against heterokaryosis brought forward by Stukenbrock and Rosendahl (2005a) is based on singlestrand conformation polymorphism and does not offer the resolution to distinguish between nucleotide differences within variants. Furthermore, for one of the markers used in this study, the Large Subunit rDNA, more alleles were recovered within the same isolate than the estimated gene copies per nucleus (Boon et al. 2010), indicating genetic differentiation between genomes for at least this locus. Recent publications of the Rhizophagus irregularis genome (Tisserant et al. 2013) and single nucleus sequencing (Lin et al. 2014) report evidence in favor of homokaryosis, but it is unclear whether the approach adopted in these studies is sufficient to provide a definite answer to the debate.

In contrast, several recent studies are in support of the heterokaryosis hypothesis. First, there is evidence for withinisolate sequence polymorphism in R. irregularis DAOM 197198 (synonym Glomus irregulare) and Glomus etunicatum (synonym Claroideoglomus etunicatum) transcripts (Boon et al. 2010; Tisserant et al. 2012). Second, the possibility of segregation of genetic variation between parent and offspring has been demonstrated for R. irregularis (Angelard et al. 2010) and G. etunicatum (Boon et al. 2013). Patterns of genetic segregation between parent and clonal offspring indicate that different fractions of genetic variation are passed on to different spores. Moreover, this variation appears to make a difference to the phenotype of the offspring isolate (Angelard and Sanders 2011). Third, within-isolate heterokaryosis has been demonstrated for several loci (review in Boon et al. 2010). Fourth, several AMF taxa seem at no part of their life cycle reduced to a single nucleus (Jany and Pawlowska 2010; Marleau et al. 2011; Ehinger et al. 2012). This latter observation offers a proximate, mechanistic explanation for high levels of genetic polymorphism in AMF isolates. This peculiar genomic organization might be the result of the absence of a bottleneck of genetic variation at any point in the AMF life cycle, which sets AMF apart from filamentous fungi, which are heterokaryotic only in a part of their reproductive cycle.

High levels of genetic variation within the AMF cytoplasm lead to conceptual as well as practical challenges to studying the real extent of IGH in AMF. An expanded array of methods is required to study genome structure and organization of AMF genetic diversity. We propose that relatively cost-effective and easily applicable methods inspired by metagenomics can be used within the cytoplasm of AMF isolates to provide estimates of genetic diversity in an organism with potentially genetically differentiated genomes. We adopted two complementary approaches to study the organization of genome diversity in AMF, focusing on polymorphism both at a genomewide scale and in single copy loci.

First, we estimated the genome-wide distribution of sequence differentiation. For this, we used a method that clusters data from whole-genome shotgun pyrosequencing runs of two R. irregularis isolates and one Rhizophagus sp. isolate, together also referred to as Rhizophagus. Clustering of reads was performed using sequence similarity networks (Yona et al. 2000; Medini et al. 2006; Halary et al. 2011; Alvarez-Ponce et al. 2013; Misner et al. 2013) (fig. 1). Then, we measured average percentage identity between sequences within clusters of overlapping homologous reads (henceforth referred to as "PID"), following Halary et al. (2009, 2013) (fig. 2). We also estimated clustering coefficients, which are measures of the connectivity of the clusters (Misner et al. 2013). This network analysis allowed us to compare Rhizophagus reads clusters to simulated wholegenome shotgun pyrosequencing runs of fully sequenced fungal genomes with a range of genome sizes. Distributions of PID and of clustering coefficients obtained for Rhizophagus data were compared with the corresponding distributions obtained from these controls, so significant deviations in Rhizophagus with respect to simulated data could be statistically assessed.

Second, we studied genome differentiation in detail by targeting polymorphic loci inferred to be present in single copy in two R. irregularis isolates and one Rhizophagus sp. isolate,

FIG. 1.—A graphic representation of the evolutionary network workflow.

which are together again referred to as Rhizophagus. Single copy markers have previously been reported in AMF (Stukenbrock and Rosendahl 2005b), although only polymorphism of intron sequences was assessed and copy number for these markers has never been published. To develop our markers, we screened open reading frames (ORFs) in 16 fully sequenced fungal genomes. As genetic variation between sequences that are present in single copy in the genome necessarily represents genetic variation between different nuclei within the same hyphal system, this method allowed us to infer intergenomic sequence variation for specific loci.

The two approaches combined attempt to address the question of the extent and physical partitioning of genome differentiation in Rhizophagus. This study represents the first genome-scale approach to tackle this question in a multigenomic organism. We find evidence for genome differentiation within the Rhizophagus cytoplasm, both genome-wide and on the scale of a single locus. The proposed genome differentiation has important implications for Rhizophagus identification using genetic information, and raises questions as to how these possibly differentiated genomes function as an integrated "individual."

SMBE

GENOME BIOLOGY AND EVOLUTION

SMBE

FIG. 2.—Description of the two variables employed to describe similarity clusters. Sequence alignments and their corresponding networks are shown. (A) The PID is the percentage of identical positions on the shortest sequence of an aligned reads pair. The PID per cluster is the average PID for all aligned reads pairs. (B)–(D) represent three scenarios of maximum, intermediate, and minimum connectivity, respectively. The clustering coefficient is the number of aligned reads pairs in the cluster, divided by the maximum number of possible pairs.

Materials and Methods

WGS Pyrosequencing of Rhizophagus Isolates

Approximately 1 million sterile spores of R. irregularis (synonym, G. irregulare) isolate DAOM 197198, formulated as commercial inoculant Mycorhise ASP, were provided by Premier Tech Biotechnologies (Rivière-Du-Loup, QC, Canada) in a liquid suspension of 4,000 spores/ml. This suspension was filtered on a sterile plastic 35-um sieve. Spores were checked for root contamination under a binocular microscope and root fragments were removed with forceps. The fungal material of isolates R. irregularis DAOM 234179 and Rhizophagus sp. DAOM 229456 (previously identified as Glomus diaphanum; Y. Dalpé personal communication) was obtained from in vitro cultures with Agrobacterium rhizogenens transformed carrot roots. An AMF isolate is a culture that was originally grown from a single spore. Spores and hyphae were freshly harvested by dissolving the Gellan-Gum matrix in which cultures were grown in a solution containing 0.0083 N sodium citrate and 0.0017 N citric acid, then gently crushed in a 1.5-ml microtube using a sterilized pestle. DNA was extracted using DNeasy Plant Mini kit (Qiagen), according to manufacturer's instructions. The purified DNA was then sent to the Genome Quebec Innovation Centre (McGill University, Montréal) for pyrosequencing using the GS FLX Titanium whole-genome shotgun kit (Roche 454 Life Science), employing a full run for each DNA sample.

Choice of Control Genomes and Pyrosequencing **Simulations**

To provide an internal control to interpret our sequence similarity network analysis (see below), we chose a wide range of fungal genomes with genome sizes from 15 to 150 Mb in order to cover same order of magnitude as the predicted R. irregularis genome sizes (Martin et al. 2008; Sedzielewska et al. 2011). To approach the 15-Mb genome size estimate, we chose genomes of Candida albicans (strains wo1 and sc5314) and Candida dubliniensis (abbreviated as wo1, sc and dub, respectively). These Candida genomes harbor GC contents (from 33.25% to 33.87%) close to R. irregularis (28%) (Tisserant et al. 2013). For the 150-Mb estimate, our simulations were based on the Puccinia triticina, Puccinia graminis, and Tuber melanosporum genomes (abbreviated as tri, gra, and tub, respectively), with GC content from 43.35% to 46.34%. Genomes were downloaded from National Center for Biotechnology Information (NCBI), or directly from the sequencing centre or genome consortium. Information regarding these genomes is summarized in [supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [table S1,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online. Pyrosequencing simulated data sets from these genomes, similar to our Rhizophagus WGS in terms of number of reads, length distribution, and technical bias, were performed with Metasim v0.9.1 (Richter et al. 2008). Simulation details are provided in [supplementary table S2,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online.

Sequence Similarity Network Analysis

As there is as yet no genome sequence available for most AMF isolates, except isolate DAOM 197198 (Tisserant et al. 2013), we used a method that allows us to describe the topology of variation in Rhizophagus without the need for detailed knowledge of genome content. By analogy, this approach can be likened to a restriction enzyme analysis such as restriction fragment length polymorphism (RFLP), where patterns of DNA variation are studied without knowledge of the actual genetic code. In RFLP, the same restriction enzymes are used for all DNA fragments under investigation, it becomes possible to study the relative behavior of the fragments that are cut by these enzymes. In a similar fashion, each control genome was "cut" using the sequencing parameters from actual Rhizophagus pyrosequencing runs to simulate exactly the same pyrosequencing run from an already published fungal genome. Subsequently, as in Misner et al. (2013), we used sequence similarity networks to cluster the real reads on the one hand, and simulated reads on the other hands, to compare the topological characteristics of these clusters (fig. 1).

All sequences sharing at least 25% identity and 75 identical nucleotides, with a BLAT e value cutoff of 1e-20, were grouped together following Halary et al. (2013). The resulting clusters are described by two variables. The first variable is the percentage of identical positions on the shortest sequence of an aligned reads pair (PID) (Misner et al. 2013) (fig. 2), which yields a highly conservative average percentage identity between sequence pairs in a cluster. The second is the clustering coefficient, which corresponds to the number of connected reads pairs in the cluster, divided by the maximum number of possible connections (also used in Misner et al. [2013]). This last variable quantifies similarities between sequences within a cluster of reads. The closer the clustering coefficient is to zero, the less connected, hence the more variable are the sequences in a cluster. The distributions of PID and clustering coefficient values from Rhizophagus and corresponding simulated reads networks were compared using a two-tailed Kolmogorov– Smirnov (KS) test.

Annotating Singletons from the Sequence Similarity **Network**

To determine whether the observed differences in singleton numbers were due to differences in the functional contents of the data sets, we annotated all singletons from our sequence similarity network. FragGeneScan (Rho et al. 2010) was first used to predict and translate ORFs. The resulting protein sequences were then aligned against the Uniref90 database (Suzek et al. 2007) using BLAT (Kent 2002). UniProt90 numbers from the functional annotation were translated into KEGG Orthology (KO) numbers using the ID mapping tools on the UniProt website [\(www.uniprot.org](www.uniprot.org) , last accessed April 2013). KO numbers were mapped to KEGG pathways using the KEGG Mapper web server [\(www.genome.jp/kegg,](www.genome.jp/kegg) last

accessed April 2013). KO numbers from all (simulated) pyrosequencing runs were compared with the run with the largest number of annotated genes as distributions of annotations, which was the tub genome simulated under the parameters of the Rhizophagus sp. DAOM 229456 pyrosequencing run (which also yielded the highest number of reads). Significant differences in the proportion of functionally annotatable singletons between runs were tested with KS tests in R [\(www.r](www.r-project.org)[project.org](www.r-project.org), last accessed April 2013).

De Novo Identification of Repetitive DNA in Rhizophagus Runs and Assembled Genomes

To evaluate the repeats content of genomic data of Rhizophagus and control genomes, RepeatScout (Price et al. 2005) was used to generate a de novo repeats library, with default parameters and the minimum element length to report set at 50 bp. Rhizophagus input data consisted of the pyrosequencing reads previously described. No read sets were available for the control genomes, so we used contigs, ultracontigs or scaffolds depending on availability, with preference for the highest assembly level. Choosing the highest assembly level will yield the least repeats, and is thus a conservative estimate relative to the Rhizophagus data, which was only available in reads. We estimated the total number of interspersed repeats (including processed pseudogenes, retrotranscripts, Short Interspersed Elements (SINES), DNA transposons, retrovirus retrotransposons, nonretrovirus retrotransposons, Long Interspersed Elements (LINES)), simple repeats (SR), and low complexity (LC) regions in the data sets with RepeatMasker Open-3.0 (Price et al. 2005) ([http://www.](http://www.repeatmasker.org) [repeatmasker.org\)](http://www.repeatmasker.org). The original RepeatScout library was used as a query. All parameters were set to default, except "cross_ match" as the search engine and the "slow" option, in order to obtain an increase of 0–5% in sensitivity with respect to the default parameters.

Estimating Rhizophagus spp. Genome Size

Assuming that Rhizophagus is not genetically heterogeneous and given that the Rhizophagus genome cannot be shorter than the sum of the DNA segments of its nuclear DNA that do not overlap, it becomes possible to calculate the minimum length of the Rhizophagus genome. To do so, we added the total length of each contig, assembled using Roche Newbler 454 assembly software, to singleton lengths (which are reads that could not be assigned to a cluster). When we assume that nuclei can be genetically different within an isolate, we can estimate the length of Rhizophagus "pangenome," understood as the entire collection of nuclear DNA in an isolate.

Development of Single Copy Markers

Proteins from 16 fully sequenced fungal genomes available at the time of analysis were investigated using a pipeline of custom-made perl scripts, to find all ORFs that were present

only once in all genomes. We used the genomes of Ascomycota Ashbya gossypii, Aspergillus fumigatus, Aspergillus nidulans, Aspergillus niger, Aspergillus oryzae, Candida glabrata, Debaryomyces hansenii, Kluyveromyces lactis, Magnaporthe grisea, Neurospora crassa, Saccharomyces cerevisiae, Schizosaccharomyces pombe, and Yarrowia lipolytica; of Basidiomycota Cryptococcus neoformans and Laccaria bicolor; and of the Microsporidia Encephalitozoon cuniculi. ORFs from A. gossypii were BLASTed against the nr database (BLASTP, threshold < 1e-5, max 5,000 hits) to retrieve homologs. Each gene was aligned with all its homologs using MUSCLE (Edgar 2004). Ambiguously aligned regions were excluded using GBlocks (Talavera and Castresana 2007) and double-checked with HoT (Landan and Graur 2007), retaining positions that were identically aligned in the reverse and forward alignments. Unambiguously aligned positions were used to reconstruct maximum-likelihood trees (applying the WAG + Gamma 4 categories model of nucleotide substitution, empirical character frequencies, estimated invariant proportion), using PHYML. These trees were scanned to define gene families in which fungi 1) were monophyletic and 2) were found in a canonical position with respect to other taxa (where "canonical" follows the phylogeny published in James et al. 2006). Reads from the Rhizophagus spp. runs were then aligned against these likely vertically inherited, highly conserved single copy genes. Alignment quality for selected markers was visually evaluated, applying unambiguity of alignment, sequence length, and conservation as criteria to design polymerase chain reaction (PCR) primers for 36 candidate single copy markers in Rhizophagus. Amplification of PCR primers was tested in the laboratory. Primer details for the retained markers are provided in [supplementary table S3,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online.

Relative Quantification of Marker Copy Number

To verify whether putative single copy markers were present in single copy in the Rhizophagus spp. genomes, we performed reverse transcription (RT)-PCR with the (monomorphic) single copy marker Rad15 as standard (Hijri and Sanders 2004; Corradi et al. 2007). Quantification of marker copy number was performed with SYBR green fluorescence for all the markers for all Rhizophagus isolates, and validated for a subset of the markers (40S-riboprot and Ef-tu) with TaqMan assays (Life Technologies, Canada) on R. irregularis DAOM 197198. The marker RLi-ABC was not validated with TaqMan assays because no sufficiently conserved region could be detected to design the probe on. Conserved fragments of marker regions were targeted using forward and reverse primer pairs and probes described in [supplementary table S3,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online. TaqMan probes and primers were designed using Primer Express 3.0 software (Life Technologies) and synthesized by Life Technologies. In each quantitative PCR (qPCR) experiment, we deployed the same amount of DNA for the amplification of marker fragments as for the amplification of a gene that is strongly suspected to occur in single copy in the genome of R. irregularis DAOM 197198, Rad15 (Hijri and Sanders 2004; Corradi et al. 2007).

Total DNA was extracted from spores and hyphae using DNeasy Plant Mini Kit (Qiagen, Canada). DNA was quantified using Quant-iT PicoGreen (Life Technologies). Two-fold serial dilutions of R. irregularis DAOM 197198 DNA (ranging from 21 to 0.65 ng) were performed in parallel for all samples including the reference gene Rad15. qPCR was performed in three replicates, with six dilutions per replicate using iTaq Universal Probes Supermix (BioRad, Canada) for TaqMan experiments and Maxima SYBR green qPCR Mix (Fermentas, Canada) for SYBR Green experiments. qPCR reactions were performed in a 20 µl volume in ViiA7 Thermalcycler (Life Technologies). The cycle threshold (Ct values) was then plotted against the log of the DNA concentration and relative copy number was established for each sample of target DNA using the Rad15 DNA regression line as a standard.

Amplicon Pyrosequencing of Single Copy Markers

To explore sequence polymorphism between alleles of our single copy markers within and between Rhizophagus spp. isolates, we performed pyrosequencing on five selected markers. Potential single copy marker sequences were amplified using DreamTaq DNA polymerase (Fermentas) using primers listed in [supplementary table S3,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online, with suitable adapter, key, and MID sequences added. DNA from the strains R. irregularis DAOM 197198, R. irregularis DAOM 234328, and Rhizophagussp. DAOM 229456 was extracted as previously described (Boon et al. 2010). The reaction was performed in 50 µl, containing 1 ng DNA, 1.25 U Taq polymerase (Fermentas), 0.2 mM dNTPs, 0.4 μ M of each primer, and the PCR buffer. PCR was carried out for 40 cycles, that is, 94 °C for 30 s, Ta for 30 s (see [supplementary table S3,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online, for annealing temperature per primer pair) and 72 °C for 1 min, which were preceded by an initial 3-min denaturation at 95 °C and followed by a 10-min hold at 72 °C, on a Mastercycler EPgradient S (Eppendorf). The PCR product was checked on an electrophoresis gel to ensure successful amplification of the gene, and then purified using a MinElute PCR Purification Kit (Qiagen) according to manufacturer's instructions. These purified samples were pooled by molecular weight and sent to the Genome Quebec Innovation Centre (McGill University) for pyrosequencing using the GS FLX Titanium emPCR kit (Roche 454 Life Science) with lib-L chemistry in one-eighth run.

Analysis of Pyrosequencing Results for Single Copy **Markers**

All analyses were performed using Mothur v. 1.22 (Schloss et al. 2009), unless specified otherwise. Low-quality reads

were eliminated according to previously published guidelines (Huse et al. 2007; Schloss et al. 2011); eliminated reads included those that 1) did not perfectly match the adaptor and primer sequences, 2) had ambiguous bases, 3) had a quality score below an average of 35 in a window of 50 bp, and 4) contained homopolymer lengths greater than 8 bp. Reads that passed quality control and differed by just 1 bp were preclustered following Huse et al. (2010). Chimeric molecules that could have formed during the PCR (Wang GCY and Wang Y 1997) or pyrosequencing (Haas et al. 2011) steps were defined as reads that did not match a database of previously obtained (Sanger sequenced) sequences with less than 90% bootstrap support. Chimeric sequences were detected and removed using the program Chimeraslayer (Haas et al. 2011). To assess whether sampling was representative of the actual diversity, we performed rarefaction analyses for every marker. We calculated the total number of alleles and the Chao1 diversity index, which measures the minimum richness in a sample (Chao et al. 2005). Only alleles that occurred more than once in the data set were considered.

Sequences were translated into amino acids by comparing them with homologous loci from the 16 fungal genomes that were originally used to find single copy markers in Rhizophagus spp. genomes. Recombination rate was calculated in DnaSP 5.10.01 (Librado and Rozas 2009) with the Rm estimator (Hudson and Kaplan 1985).

To assess whether our clustering of the data into alleles was the most appropriate approach to minimize the influence of sequencing error on our conclusions, we also tested three alternative clustering or denoising strategies, that is, preclustering by 2 bp differences instead of one, denoising of the sequences through SeqNoise as implemented in Mothur v1.28, and denoising of the flow files through AmpliconNoise (Quince et al. 2011). As error rate varies per run, we estimated per run error rate based on sequencing errors from Roche's internal homopolymer controls, which were kindly provided by the Genome Quebec Innovation Centre (McGill University).

Validation of Polymorphism

To investigate the possibility that single copy marker polymorphism was due to the sequencing methodology, we cloned and Sanger sequenced the TaqMan qPCR products of the 40S-riboprot, Ef-tu, and ARP markers. An amount of 4 μ l qPCR product was ligated into pGEM-T easy vector (Promega, Canada) and transformed into Escherichia coli competent DH5alfa cells. Bacterial colonies were randomly picked and PCR-screened with universal T7 and SP6 vector primers. Bacterial colonies that showed a PCR product with the expected size were considered as positive clones and were sent for sequencing to the Genome Quebec Innovation Centre (McGill, Canada).

Data Deposition

All raw pyrosequencing data (amplicon data for the single copy markers and Whole Genome Sequencing (WGS) for the evolutionary network analyses) were deposited in the NCBI Trace Archive under Bioproject number PRJNA174749. The single copy marker allele alignments are provided in the [supplementary material,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online.

Results

Genome-Wide Diversity Patterns through Simulations and Clustering

We estimated 1) the PID distribution and 2) clustering coefficients (where a high clustering coefficient indicates high overlap between sequences), following Halary et al. (2009, 2013) and Misner et al. (2013), see Materials and Methods and figures 1 and 2.We plotted the frequency distribution of average PID per cluster from each (real and simulated) pyrosequencing run (figs. 3 and 4). PID distributions from C. albicans sc5314, C. albicans wo1, and C. dubliniensis (from now on referred to as small control genomes) were significantly different from both Rhizophagus and tri, gra and tub (from now on referred to as large control genomes) [\(supplementary table S4](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). PID distributions from Rhizophagus pyrosequencing runs were not significantly different from those based on simulations from large control genomes. However, Rhizophagus PID distributions showed much elevated numbers of reads between 95% and 100% PID in comparison to the large control genomes. We also plotted clustering coefficients against average % PID per cluster (fig. 4). Except in the case of R. irregularis DAOM 234179 run 2, each analysis showed a higher % PID for the same clustering coefficient in Rhizophagus reads clusters, meaning that there is more sequence overlap in these data sets than in controls.

An interesting exception was the case of the gra simulated data set, which contained the largest amount of data (1,078,190 reads, table 1) (fig. 3d). The PID distribution was significantly different from the other large control genomes and Rhizophagus (KS test, $D = 0.3168$ [comparison with Rhizophagus sp. and tub] and 0.3267 [comparison with tri], P values 7.91E-05 and 4.15E-05, respectively, see also [supple](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1)[mentary table S4,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). The difference between the distributions for this Rhizophagus sp. run and for gra, the smallest of the large control genomes genome (88 Mb, [supplementary table S1,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online) reveals an important property of our approach. It shows that sequencing coverage plays an important role in the resolution of our network analyses. When the sampling depth of the real and simulated pyrosequencing runs approaches the actual size of the sampled genome, the sequence similarity network approach starts differentiating small and large genomes. In the case of gra, the genome size is

GENOME BIOLOGY AND EVOLUTION

SMBE

88 Mb and the pyrosequencing depth is 1,078,190 reads with an average length of 336 bp ([supplementary tables S1](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) and [S2,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Thus, a mere 4-fold coverage seems sufficient to start inferring differences between Rhizophagus and some of the large control genomes.

Estimating Genome Size with Sequence Similarity **Networks**

Clustering reads from control genomes resulted in very few clusters with many reads on the smaller Candida control genomes. This result is expected and consistent with analyses in Misner et al. (2013). We propose that these few clusters correspond to supercontigs of the Candida genomes, covering very large segments of these genomes. These "superclusters" are formed more readily in Candida genomes, as their small size leads to a high coverage faster than in the larger genomes. Therefore, our clustering approach effectively gathered large segments of the smaller control genomes by identifying overlapping reads. In contrast, our pyrosequencing efforts did not lead to a similar clustering for the large control genomes and for Rhizophagus. As Candida and Rhizophagus genomes share similar GC contents, unlike large control genomes, the effect of the pyrosequencing data set sizes (and thus, coverages) is likely much greater on clustering differences than a GC bias.

Because no Rhizophagus genome was thus fully "assembled," we used the total length of the assembled contigs plus that of singletons as a conservative estimate of the Rhizophagus genome (or pangenome) size. The sum of all contigs and singletons for R. irregularis DAOM 234179 was 178 Mb (based on an assembly comprising both run 1 and run 2), 163.7 Mb for Rhizophagus sp. DAOM 229456, and 64.7 Mb for R. irregularis DAOM 197198. These sums could be considered minimum genome sizes only in the sense that they are a sum of all overlapping and nonoverlapping genetic variation that a single pyrosequencing run retrieves. In the light of the heterokaryosis hypothesis, these minimum genome sizes do not give us information on whether the variation is located within or between nuclei. In principles, we may have inferred a conservative size for the pangenome of an isolate, and genome size of individual nuclei within such isolate may still vary.

Annotating Singletons

We annotated all singletons from Rhizophagus runs and from the large control genome simulated runs, for which distributions of PID and clustering coefficient are most similar to our

FIG. 4.—Scatterplots depict average PID in a similarity cluster and clustering coefficient. For all Rhizophagus spp. and control genomes, (a) R. irregularis DAOM 234179 run 1, (b) R. irregularis DAOM 234179 run 2, (c) R. irregularis DAOM 197198, and (d) Rhizophagus sp. DAOM 229456. Colors correspond to the number of reads that are represented by the respective data-points (see legend; for simplicity, only one legend is depicted for all graphs). The shape of the data-point refers to the reads set: Circle, Rhizophagus (specific isolates are identified in the graph title); square, P. graminis; diamond, T. melanosporum; triangle, P. triticina.

Rhizophagus pyrosequencing runs. Initial numbers of singletons varied between 4,474 for the tub-based simulation under the parameters of R. irregularis DAOM 234179 run 2 and 496,891 for the tub-based simulation under the parameters of Rhizophagus sp. DAOM 229456. Singletons could be assigned to ORFs and annotated [\(supplementary fig. S1,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online) with comparable success between Rhizophagus and simulated runs (Wilcoxon Signed Rank Test, [supplementary table S5](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Thus, Rhizophagus singletons do not consist of less ORFs than singletons from the large control genomes. We compared the KEGG annotations of the singletons, and tested the significance of differences between annotation distributions with KS tests. There were no differences between annotated singletons from Rhizophagus and control runs: Only the tub- and gra-based simulations under the parameters of Rhizophagus sp. DAOM 229456 were different from the other (simulated) runs [\(supplementary table S6,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Thus, annotation content, based on KEGG hierarchies, between Rhizophagus singletons and the large control genomes did not change with genome used as a basis for the (simulated) run.

De Novo Identification of Repetitive DNA in Rhizophagus Runs and Assembled Genomes

We estimated GC content and the percentage of masked bases for pyrosequencing runs of our Rhizophagus and for the control genomes [\(supplementary table S7](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Rhizophagus runs showed a typical low GC percentage (Hosny et al. 1997; Tisserant et al. 2013). They also showed a higher percentage of masked bases, although isolate DAOM 197198 fell in the lower percentage of masked bases and was in this respect similar to tri. SR were higher for Rhizophagus than for the larger control genomes, but lower with respect to the small genome control data sets. Finally, Rhizophagus runs showed a slightly higher percentage of LC regions with respect to all control genomes. Rhizophagus isolate DAOM 197198 actually has less masked bases than two of the control genomes, gra and tub.

Allele Diversity Estimates within Rhizophagus spp. Isolates

Our second approach to study genome diversity in AMF is a detailed investigation of polymorphism within and between single copy markers. We developed and pyrosequenced five

Results from Clustering Analyses for Rhizophagus spp. and Control Genomes

NOTE.-wo1, Candida albicans wo1; sc, C. albicans sc5314; dub, C. dubliniensis.

novel markers, which all represent partial exons from proteincoding sequences. Between 90 and 1,123 sequences fulfilling stringent quality criteria were recovered per marker (table 2). All markers were polymorphic, yielding between 2 and 103 alleles. Rarefaction curves are reported in [supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [figure S2,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online. Preclustering at different levels had an effect on allele count, but denoising strategy did not [\(supplementary table S8,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). The closest database matches for these markers after three psi-BLAST iterations (pBLAST search of translated marker sequences against nr, cutoff E-25) are detailed in [supplementary table S9](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online. Markers were named after the closest psi-BLAST hit.

Testing for Copy Number and Polymorphism Validation

Copy number for the five markers was tested by qPCR. Markers Ef-tu, 40S-riboprot, and RLi-ABC showed similar linear regressions of Ct values than the reference gene Rad15 (fig. 5 and [supplementary fig. S3](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). This indicates that these three markers have similar copy number than Rad15, which is likely a single copy gene. A R. irregularis genome search confirmed the occurrence of one copy of Rad15 sequence. Surprisingly, the markers ARP and ACOB showed approximately 2-fold higher Ct values compared with Rad15. Thus, in both TaqMan (ARP) and SYBR Green (ACOB) assays, the markers ARP and ACOB seem to be present in less than one copy per genome (fig. 5 and [supplementary fig. S3](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online).

We validated the polymorphism observed in the amplicon pyrosequencing runs for the markers 40S-riboprot, Ef-tu, and ARP, by cloning and Sanger sequencing of the TaqMan qPCR products. In spite of the low sampling intensity yielded by cloning/sequencing, most abundant alleles for these markers

Table 2

Amplification and Genetic Diversity of Single Copy Markers

^aR. irregularis DAOM 197198; R. irregularis DAOM 234328; Rhizophagus sp. DAOM 229456.

^bNumber of alleles that occur more than only once in the data set.

c Chao1 index; lci, lower 95% confidence interval; hci, higher 95% confidence interval (Chao et al. 2005).

were recovered [\(supplementary table S10,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online).

large portion of the clusters that are characterized by a high average PID (>95%) come from conserved coding genome regions located in different nuclei.

Discussion

Excess of Strongly Similar Coding DNA Regions in Rhizophagus

PID distributions from the small control genomes were significantly different from both Rhizophagus and large control genome distributions (fig. 3 and [supplementary table S4,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Thus, our sequence similarity network approach suggests that Rhizophagus genomes behave as a large genome. We predicted Rhizophagus minimum genome sizes. They range from 64.7 Mb for R. irregularis DAOM 197198 to 163.7 Mb for Rhizophagus sp. DAOM 229456 and 178 Mb for R. irregularis DAOM 234179, which all fall in the size range of the larger genomes. The most recent genome size estimates for R. irregularis also fall in this order of magnitude (Sedzielewska et al. 2011; Tisserant et al. 2013). It is important to note that comparing Rhizophagus to fungi with large genomes does not provide any information on the location of the variation: Clusters of reads and singletons on which these size estimates are based could be located within or between different nuclei in the cytoplasm.

The PID distributions of Rhizophagus show more reads clusters with higher average sequence identities (PID) than simulated genomes (fig. 3). Furthermore, reads clusters from Rhizophagus genomes have higher average PID than clusters of reads from control fungi (table 1). This excess of "strongly similar" regions (PID $> 95\%$) in Rhizophagus isolates suggests that the genetic organization in Rhizophagus differs from that in the control fungi. How can we interpret the excess of strongly similar regions in Rhizophagus spp. isolates? The reads from clusters that are characterized by a high average PID can come from the same or from different nuclei within the same cytoplasm. They can represent conserved coding genome regions in Rhizophagus, or noncoding repetitive elements (NCRE), such as tandem repeats. We propose that a

Indeed, for any cluster of reads, a high clustering coefficient indicates that a similar sequence is repeated multiple times in the cluster, whereas a low clustering coefficient indicates a cluster that is more comparable to a contig (e.g., a succession of overlapping reads, see also fig. 4). For the same average PID per cluster, Rhizophagus clusters have a higher clustering coefficient than clusters from the control genomes (fig. 4). Thus, nuclei in Rhizophagus isolates contain more highly similar regions, and a greater redundancy than the control fungi.

To get an idea of cluster content, we looked at repeat content in clusters. We compared NCRE in clusters from Rhizophagus and control genomes. NCRE numbers in Rhizophagus were not substantially different between large control and Rhizophagus genomes ([supplementary table S7](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Our current estimate of 45% aligns more closely with findings from the Rhizophagus genome (Tisserant et al. 2013). NCRE numbers are also high in the large control genomes [\(supplementary table S1](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online), so the peak Rhizophagus isolates show at greater than 95% PID cannot be explained by NCRE. Moreover, there were no differences in the proportions of annotated singletons between Rhizophagus genomes and large control genomes, indicating that all these genomes have comparable amounts of coding material [\(supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [fig. S1](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) and [table S8,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online)

Thus, we find redundant clusters of highly similar (although not identical) coding sequences in the Rhizophagus pyrosequencing runs. However, with our current data, it is not possible to assess whether these redundant coding sequences occur within or between nuclei in the same cytoplasm. Therefore, we cannot differentiate between highly conserved multigene families within the same nucleus, which would fit the homokaryosis hypothesis, or repeated low copy genes that are partitioned between nuclei, which would fit the heterokaryosis hypothesis. To distinguish between these

Downloaded from <http://gbe.oxfordjournals.org/> at UPMC on November 4, 2015

Downloaded from http://gbe.oxfordjournals.org/ at UPMC on November 4, 2015

FIG. 5.—Results of real-time qPCR for single copy markers. Linear regressions of the cycle threshold (Ct values) and the Log concentration of R. irregularis DAOM 197198 genomic DNA (pg) that was used as template for the reaction using TaqMan assays. Ct values of the markers Ef-tu and 40S-riboprot were compared with Ct values of the marker Rad15.

scenarios, we have to look in detail at selected single copy markers.

Rhizophagus Genome Variation in Close-Up with Single Copy Markers

This study reports extensive polymorphism on protein-coding single copy markers in AMF. We retrieved between 2 and 103 alleles for each putative single copy marker (table 2). Differentiation is not homogenous between loci: Some markers yielded relatively few variants (marker ACOB), whereas others revealed over a hundred different alleles (e.g., marker Ef-tu, see fig. 6a for the allele distribution). Our observations concur with previous findings of genetic differentiation between loci in AMF nuclei within the same cytoplasm

(Kuhn et al. 2001; Hijri and Sanders 2005; Boon et al. 2010; Ehinger et al. 2012; Tisserant et al. 2012).

Importantly, the allele counts we report most likely underestimate the total allele diversity in the isolates used in this analysis, for three reasons. First, none of the approaches we used to estimate minimum allele diversity for the Rhizophagus isolates reached a plateau of diversity, as defined by 1) rarefaction analyses [\(supplementary fig. S2](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [Supplementary](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online) and 2) the lower confidence interval of the Chao1 diversity index (table 1). For four of five markers (with the exception of the marker ACOB), the rarefaction analyses and Chao1 diversity index both indicate a spectrum of rare alleles that remains to be sampled. Second, we applied stringent quality checks, and reads differing by only 1 bp were Downloaded from <http://gbe.oxfordjournals.org/> at UPMC on November 4, 2015

Downloaded from http://gbe.oxfordjournals.org/ at UPMC on November 4, 2015

FIG. 6.—Allele distributions for selected single copy markers (a) 40S-riboprot and (b) Ef-tu (only alleles for which $f > 1$ are depicted).

clustered together. This clustering artificially lowers variation in a data set by erring on the conservative side. Three, we only counted alleles that occurred more than once in the reads from the pyrosequencing runs.

Although our conservative approach increases the chance of lumping different alleles together, this approach is necessary to avoid the inflated diversity counts that are often associated with pyrosequencing technology (Huse et al. 2010; Schloss 2010; Haas et al. 2011; Schloss et al. 2011; Schloss and Westcott 2011). More stringent preclustering is not expected to yield more stringent results: The error rate of our specific amplicon pyrosequencing run was 0.6%, which is lower than the amount of variation we removed by preclustering. Incidentally, this rate is also lower than the mean rates reported for this sequencing technology (Gilles et al. 2011). Finally, more stringent preclustering would compromise our ability to separate signal from noise. To elaborate on this point, if we would cluster reads differing by 2 bp together, the maximum amount of differences within a cluster would be 4 bp. On the length of one of the longest alignments, that for 40S-riboprot (175 bp), 4 bp represents 2.3% of the total sequence. As this level is typically the level of variation we are interested in, it thus becomes difficult to start inferring patterns of variation from the data. On shorter alignments, this effect would be even more pronounced. To actually distinguish signal from noise, one would need to implement denoising algorithms based on sequence or flow (.sff) data such as SeqNoise or AmpliconNoise, respectively. Applying

Table 3

Stopcodons, Frameshift mutations and Recombination in Single Copy Markers

^aNumber of nucleotide positions used in analysis.

^bStopcodons and frameshift events were counted before being removed from the analysis

'Minimum number of recombination events (as implemented in Librado and Rozas [2009]).

*Significant at $\alpha = 0.05$.

these denoising algorithms, we mostly find the same number or more alleles in our data [\(supplementary table S8,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) [Supplementary Material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) online). Thus, we are confident that we have underestimated and not overestimated the number of alleles for each single copy marker.

Another caveat might be the use of Rad15 as a single copy reference marker. Rad15 showed exactly the same pattern in RT-PCR experiments as Rad32 (Corradi et al. 2007), which was shown to be genetically homogeneous and likely present in single copy in isolate DAOM 197198 by a dot blot hybridization assay (Hijri and Sanders 2004). Therefore, our reference gene was the best available choice for estimation of relative copy number. It should be noted that copy number estimations in AMF can only be considered as an average over all mycelia, as it is possible that copy number variations occur between R. irregularis isolates as demonstrated by Corradi et al. (2007) for rRNA genes. However, we attempted to negotiate this difficulty by special precautions in the prescreening phase, through 1) our bioinformatics approach, in which we have excluded all loci that show signs of occurring in multiple copies in the Rhizophagus genome data or in any other fungal genome; 2) excluding all candidate markers that showed major deletions or rearrangements in the sequence alignment, which could be chimerical due to in vivo or in vitro recombination; and 3) only focusing on the most conserved parts of the loci under investigation.

If we consider the possibility of heterokaryosis, how could the polymorphism observed between single copy markers in the same AMF isolate (i.e., in the same cytoplasm) be maintained? Fusion between hyphal systems, anastomosis, could play an important role in conserving polymorphism between hyphal networks in soil ecosystems (Croll et al. 2009; Marleau et al. 2011; Boon et al. 2013). Little is known about generation and loss of genetic variation within AMF isolates. The possible effects of genetic drift have been described in AMF isolates (Cárdenas-Flores et al. 2010; Angelard et al. 2010; Angelard and Sanders 2011; Ehinger et al. 2012; Boon et al. 2013; de la Providencia et al. 2013). Alternatively, differences between allele frequencies might be due to differential selection pressures on the loci themselves or on the adjacent genome regions of the single copy markers. Unfortunately, we cannot test this latter hypothesis on our data, as preclustering reads from pyrosequencing technology means that all allele sequences are consensus sequences. Thus, variation between sequences cannot be used to confidently infer deviations from neutrality. Deviations from neutrality between loci are possible if linkage equilibrium is interrupted by recombination, as we observed for three of five loci (table 3), and as has been previously been reported in AMF (Vandenkoornhuyse et al. 2001; Gandolfi et al. 2003; Croll and Sanders 2009). However, the investigation of recombination and linkage in AMF is precarious, as there are few databases of sufficiently long reads available for any AMF.

This latter point stresses the potential importance of a single copy marker approach, as databases of variation between these or other single copy markers could be implemented in a manner very similar to multilocus sequence typing (MLST) to answer questions of linkage and recombination (Maiden et al. 1998). A major advantage of this strategy would be the possibility to study sequence polymorphism in AMF without assumptions on the inter- or intranuclear localization of this variation.

Probing the Rhizophagus Genome

We suggest that each Rhizophagus isolate harbors a population of differentiated genomes, based on observations from the two approaches presented in this study. First, we argue that differences in PID distributions between Rhizophagus and the large control genomes point to the presence of many similar but slightly differentiated sequences in the AMF cytoplasm (figs. 3 and 4).

Second, our observations of extensive genomic heterogeneity in Rhizophagus single copy markers indicate that this variation could be partitioned between and not within nuclei (and thus genomes) in the Rhizophagus cytoplasm. These observations agree with previous findings of genetic differentiation between specific loci in AMF nuclei (Kuhn et al. 2001; Hijri and Sanders 2005; Boon et al. 2010; Ehinger et al. 2012; Tisserant et al. 2012). Even though alleles from different loci found within AMF isolates have already been shown to be physically located between nuclei using FISH (Kuhn et al. 2001; Kuhn 2003), our reports of genetic heterogeneity between loci are the first to provide direct evidence of differentiation between genomes from single copy markers.

Some Speculation on the Rhizophagus Genome

The redundant clusters of slightly differentiated reads in our sequence similarity network analysis and high levels of

polymorphism for our putative single copy genes do not align with the homokaryosis hypothesis, even though some may still find the evidence provided here insufficient to support the heterokaryosis hypothesis. However the spatial organization of genetic variation in the AMF isolates under study, we can report with certainty on the unusually high polymorphism that we recovered. Even with a one-eighth pyrosequencing run we have not attained saturation of allele diversity in this study—a telltale of much variation we are still missing. Therefore, we suggest that each Rhizophagus isolate harbors a population of (at least partly) genetically differentiated genomes. If so, populations of nuclei within the AMF cytoplasm may act together to produce the Rhizophagus phenotype. Four observations support this interpretation. First, for several AMF it has been shown that they are at no point in their life cycle reduced to a single genome (Jany and Pawlowska 2010; Marleau et al. 2011; Boon et al. 2013). Second, Rhizophagus spores do not germinate below a certain number of nuclei per spore, which is roughly 65 nuclei for R. irregularis (Marleau et al. 2011). Third, for R. irregularis and G. etunicatum it was shown that genetic polymorphism is expressed in the transcriptome (Boon et al. 2010; Tisserant et al. 2012), which indicates that differentiation at the genome level could play a role in the functioning of Rhizophagus isolates. Finally, the high amounts of genetic variation in AMF isolates have been proposed to play a role in the ability of AMF to adapt to a wide range of host plants (Angelard et al. 2010).

Accordingly, we propose that this population of partly heterogeneous genomes in AMF is analogous to a pangenome, as there may not be one typical genome within an isolate, representative of all the other, but rather a population of partly differentiated genomes. The minimum size of the Rhizophagus pangenome would then be around 65 Mb for R. irregularis DAOM 197198, 178 Mb for R. irregularis DAOM 234179, and 163.7 Mb for Rhizophagus sp. DAOM 229456. These estimates are closer to the upper limit of the R. irregularis genome size that has been published to date, that is, 150 Mb (Martin et al. 2008; Sędzielewska et al. 2011; Tisserant et al. 2013), than to a previously published lower estimate of 15 Mb (Hijri and Sanders 2004).

The recognition of unprecedentedly high levels of genetic diversity within the Rhizophagus cytoplasm reported here and the possible organization of this genetic diversity into differentiated nuclei could lead to a careful consideration of the concept of the individual (Santelices 1999; Pineda-Krch and Lehtila 2004; Pepper and Herron 2008; Folse and Roughgarden 2010) in Rhizophagus, with important consequences for genetically based AMF studies in agriculture and ecology. A particularly exciting avenue is the role of anastomosis in the maintenance of this genetic variation, which could be tested by an MLST approach on multiple Rhizophagus isolates. Comparing interisolate single copy marker diversity profiles, such as those reported in this study, will allow AMF researchers to study linkage and

recombination between AMF isolates. Microbial ecology has already developed many metagenomics tools that can be used to study the evolution, function, and stability of a community whose components cannot be traced individually. We hope that this study will open similar avenues to the study of AMF.

Supplementary Material

[Supplementary material](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [figures S1–S3](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1), [tables S1–S10,](http://gbe.oxfordjournals.org/lookup/suppl/doi:10.1093/gbe/evv002/-/DC1) and references are available at Genome Biology and Evolution online [\(http://www.gbe.oxfordjournals.org/](http://www.gbe.oxfordjournals.org/)).

Acknowledgments

This work was supported by the Natural Sciences and Engineering Research Council of Canada (NSERC) to MH (grant number: 328098-2012) and an NSERC Vanier CGS to Ev.B and by the European Research Council FP7 (2007-2013 Grant Agreement No. 615274) to Er.B. The authors thank O. Deusch for providing the software HoT, P. Lopez for providing the phylogenetic pipeline to reconstruct the trees, E. Yergeau for help with AmpliconNoise, Y. Terrat for help with RepeatMasker, and A. Staffa for help with Roche internal controls. Mothur analyses were run on the supercomputer Cottos at the Université de Montréal, managed by Calcul Québec and Compute Canada. F.-J. Lapointe, B. Angers, F. Bouchard, and D. Walsh provided useful comments on the manuscript.

Literature Cited

- Alvarez-Ponce D, Lopez P, Bapteste E, McInerneya JO. 2013. Gene similarity networks provide tools for understanding eukaryote origins and evolution. Proc Natl Acad Sci U S A. 110:E1594–E1603.
- Angelard C, Colard A, Niculita-Hirzel H, Croll D, Sanders IR. 2010. Segregation in a mycorrhizal fungus alters rice growth and symbiosis-specific gene transcription. Curr Biol. 20:1216–1221.
- Angelard C, Sanders IR. 2011. Effect of segregation and genetic exchange on arbuscular mycorrhizal fungi in colonization of roots. New Phytol. 189:652–657.
- Bever JD, Wang M. 2005. Arbuscular mycorrhizal fungi—hyphal fusion and multigenomic structure. Nature 433:E3–E4.
- Biesecker LG, Spinner NB. 2013. A genomic view of mosaicism and human disease. Nat Rev Genet. 14(5):307–320.
- Boon E, Zimmerman E, Lang BF, Hijri M. 2010. Intra-isolate genome variation in arbuscular mycorrhizal fungi persists in the transcriptome. J Evol Biol. 23:1519–1527.
- Boon E, Zimmerman E, St-Arnaud M, Hijri M. 2013. Allelic differences among sister spores suggest genetic drift in an arbuscular mycorrhizal fungus. PLoS One 8:e83301.
- Cárdenas-Flores A, Draye X, Bivort C, Cranenbrouck S, Declerck S. 2010. Impact of multispores in vitro subcultivation of Glomus sp. MUCL 43194 (DAOM 197198) on vegetative compatibility and genetic diversity detected by AFLP. Mycorrhiza 20:415–425.
- Chao A, Chazdon RL, Colwell RK, Shen T-J. 2005. A new statistical approach for assessing similarity of species composition with incidence and abundance data. Ecol Lett. 8:148–159.
- Corradi N, et al. 2007. Gene copy number polymorphisms in an arbuscular mycorrhizal fungal population. Appl Environ Microbiol. 73:366–369.

- Croll D, et al. 2009. Nonself vegetative fusion and genetic exchange in the arbuscular mycorrhizal fungus Glomus intraradices. New Phytol. 181: 924–937.
- Croll D, Sanders I. 2009. Recombination in Glomus intraradices, a supposed ancient asexual arbuscular mycorrhizal fungus. BMC Evol Biol. 9:13.
- de la Providencia IE, Nadimi M, Beaudet D, Morales GR, Hijri M. 2013. Detection of a transient mitochondrial DNA heteroplasmy in the progeny of crossed genetically divergent isolates of arbuscular mycorrhizal fungi. New Phytol. 200:211–221.
- Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res. 32: 1792–1797.
- Ehinger MO, Croll D, Koch AM, Sanders IR. 2012. Significant genetic and phenotypic changes arising from clonal growth of a single spore of an arbuscular mycorrhizal fungus over multiple generations. New Phytol. 196:853–861.
- Folse HJ, Roughgarden J. 2010. What is an individual organism? A multilevel selection perspective. Q Rev Biol. 85:447–472.
- Gandolfi A, Sanders IR, Rossi V, Menozzi P. 2003. Evidence of recombination in putative ancient asexuals. Mol Biol Evol. 20: 754–761.
- Gilles A, et al. 2011. Accuracy and quality assessment of 454 GS-FLX Titanium pyrosequencing. BMC Genomics 12:245.
- Haas B, et al. 2011. Chimeric 16S rRNA sequence formation and detection in Sanger and 454-pyrosequenced PCR amplicons. Genome Res. 21: 494–504.
- Halary S, et al. 2011. Conserved meiotic machinery in Glomus spp., a putatively ancient asexual fungal lineage. Genome Biol Evol. 3: 950–958.
- Halary S, Leigh JW, Cheaib B, Lopez P, Bapteste E. 2009. Network analyses structure genetic diversity in independent genetic worlds. Proc Natl Acad Sci U S A. 107:127–132.
- Halary S, McInerney J, Lopez P, Bapteste E. 2013. EGN: a wizard for construction of gene and genome similarity networks. BMC Evol Biol. 13: 146.
- Hijri M, Sanders IR. 2004. The arbuscular mycorrhizal fungus Glomus intraradices is haploid and has a small genome size in the lower limit of eukaryotes. Fungal Genet Biol. 41:253–261.
- Hijri M, Sanders IR. 2005. Low gene copy number shows that arbuscular mycorrhizal fungi inherit genetically different nuclei. Nature 433: 160–163.
- Hosny M, deBarros JPP, Gianinazzi-Pearson V, Dulieu H. 1997. Base composition of DNA from glomalean fungi: high amounts of methylated cytosine. Fungal Genet Biol. 22:103–111.
- Hosny M, Hijri M, Passerieux E, Dulieu H. 1999. rDNA units are highly polymorphic in Scutellospora castanea (Glomales, Zygomycetes). Gene 226:61–71.
- Hudson RR, Kaplan NL. 1985. Statistical properties of the number of recombination events in the history of a sample of DNA sequences. Genetics 111:147–164.
- Huse S, Huber J, Morrison H, Sogin M, Welch D. 2007. Accuracy and quality of massively parallel DNA pyrosequencing. Genome Biol. 8: R143.
- Huse SM, Welch DM, Morrison HG, Sogin ML. 2010. Ironing out the wrinkles in the rare biosphere through improved OTU clustering. Environ Microbiol. 12:1889–1898.
- James TY, et al. 2006. Reconstructing the early evolution of Fungi using a six-gene phylogeny. Nature 443:818–822.
- Jany J-L, Pawlowska TE. 2010. Multinucleate spores contribute to evolutionary longevity of asexual glomeromycota. Am Nat. 175: 424–435.
- Kent W. 2002. BLAT—the BLAST-like alignment tool. Genome Res. 12: 656–664.
- Kuhn G. 2003., Organisation of genetic variation in multinucleate arbuscular mycorrhizal fungi. Lausanne (Switzerland): University of Lausanne.
- Kuhn G, Hijri M, Sanders IR. 2001. Evidence for the evolution of multiple genomes in arbuscular mycorrhizal fungi. Nature 414:745–748.
- Landan G, Graur D. 2007. Heads or tails: a simple reliability check for multiple sequence alignments. Mol Biol Evol. 24:1380–1383.
- Librado P, Rozas J. 2009. DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. Bioinformatics 25:1451–1452.
- Lin K, et al. 2014. Single nucleus genome sequencing reveals high similarity among nuclei of an endomycorrhizal fungus. PLoS Genet. 10: e1004078.
- Maiden MCJ, et al. 1998. Multilocus sequence typing: a portable approach to the identification of clones within populations of pathogenic microorganisms. Proc Natl Acad Sci U S A. 95:3140–3145.
- Marleau J, Dalpe Y, St-Arnaud M, Hijri M. 2011. Spore development and nuclear inheritance in arbuscular mycorrhizal fungi. BMC Evol Biol. 11: 51.
- Martin F, et al. 2008. The long hard road to a completed Glomus intraradices genome. New Phytol. 180:747–750.
- Medini D, Covacci A, Donati C. 2006. Protein homology network families reveal step-wise diversification of Type III and Type IV secretion systems. PLoS Comput Biol. 2:e173.
- Misner I, et al. 2013. Sequence comparative analysis using networks: software for evaluating de novo transcript assembly from next-generation sequencing. Mol Biol Evol. 30:1975–1986.
- Pawlowska TE, Taylor JW. 2004. Organization of genetic variation in individuals of arbuscular mycorrhizal fungi. Nature 427:733–737.
- Pepper JW, Herron MD. 2008. Does biology need an organism concept? Biol Rev. 83:621–627.
- Pineda-Krch M, Lehtila K. 2004. Challenging the genetically homogeneous individual. J Evol Biol. 17:1192–1194.
- Price AL, Jones NC, Pevzner PA. 2005. De novo identification of repeat families in large genomes. Bioinformatics 21:i351–i358.
- Quince C, Lanzen A, Davenport R, Turnbaugh P. 2011. Removing noise from pyrosequenced amplicons. BMC Bioinformatics 12:38.
- Rho M, Tang H, Ye Y. 2010. FragGeneScan: predicting genes in short and error-prone reads. Nucleic Acids Res. 38:e191.
- Richter D, Ott F, Auch A, Schmid R, Huson D. 2008. MetaSim—a sequencing simulator for genomics and metagenomics. PLoS One 3:e3373.
- Rosendahl S, Stukenbrock EH. 2004. Community structure of arbuscular mycorrhizal fungi in undisturbed vegetation revealed by analyses of LSU rDNA sequences. Mol Ecol. 13:3179–3186.
- Sanders IR, Croll D. 2010. Arbuscular mycorrhiza: the challenge to understand the genetics of the fungal partner. Annu Rev Genet. 44: 271–292.
- Santelices B. 1999. How many kinds of individual are there? Trends Ecol Evol. 14:152–155.
- Schloss PD. 2010. The effects of alignment quality, distance calculation method, sequence filtering, and region on the analysis of 16S rRNA gene-based studies. PLoS Comput Biol. 6:e1000844.
- Schloss PD, et al. 2009. Introducing mothur: open-source, platformindependent, community-supported software for describing and comparing microbial communities. Appl Environ Microbiol. 75: 7537–7541.
- Schloss PD, Gevers D, Westcott SL. 2011. Reducing the effects of PCR amplification and sequencing artifacts on 16S rRNA-based studies. PLoS One 6:e27310.
- Schloss PD, Westcott SL. 2011. Assessing and improving methods used in operational taxonomic unit-based approaches for 16S rRNA gene sequence analysis. Appl Environ Microbiol. 77:3219–3226.
- Schussler A, Schwarzott D, Walker C. 2001. A new fungal phylum, the Glomeromycota: phylogeny and evolution. Mycol Res. 105: 1413–1421.

- Sedzielewska KA, et al. 2011. Estimation of the Glomus intraradices nuclear DNA content. New Phytol. 192:794–797.
- Stukenbrock EH, Rosendahl S. 2005a. Clonal diversity and population genetic structure of arbuscular mycorrhizal fungi (Glomus spp.) studied by multilocus genotyping of single spores. Mol Ecol. 14: 743–752.
- Stukenbrock EH, Rosendahl S. 2005b. Development and amplification of multiple co-dominant genetic markers from single spores of arbuscular mycorrhizal fungi by nested multiplex PCR. Fungal Genet Biol. 42: 73–80.
- Suzek BE, Huang H, McGarvey P, Mazumder R, Wu CH. 2007. UniRef: comprehensive and non-redundant UniProt reference clusters. Bioinformatics 23:1282–1288.
- Talavera G, Castresana J. 2007. Improvement of phylogenies after removing divergent and ambiguously aligned blocks from protein sequence alignments. Syst Biol. 56:564–577.
- Tisserant E, et al. 2012. The transcriptome of the arbuscular mycorrhizal fungus Glomus intraradices (DAOM 197198) reveals functional tradeoffs in an obligate symbiont. New Phytol. 193:755–769.
- Tisserant E, et al. 2013. Genome of an arbuscular mycorrhizal fungus provides insight into the oldest plant symbiosis. Proc Natl Acad Sci U S A. 110:20117–20122.
- Van der Heijden MGA, et al. 1998. Mycorrhizal fungal diversity determines plant biodiversity, ecosystem variability and productivity. Nature 396: 69–72.
- Van Der Heijden MGA, Sanders IR. 2002. Mycorrhizal ecology. Heidelberg (Germany): Springer-Verlag.
- Vandenkoornhuyse P, Leyval C, Bonnin I. 2001. High genetic diversity in arbuscular mycorrhizal fungi: evidence for recombination events. Heredity 87:243–253.
- Wang GCY, Wang Y. 1997. Frequency of formation of chimeric molecules is a consequence of PCR coamplification of 16S rRNA genes from mixed bacterial genomes. Appl Environ Microbiol. 63:4645–4650.
- Yona G, Linial N, Linial M. 2000. ProtoMap: automatic classification of protein sequences and hierarchy of protein families. Nucleic Acids Res. 28:49–55.

Associate editor: Bill Martin