Abstract

Arbuscular mycorrhizal fungi (AMF) are part of the most widespread fungal-plant symbiosis. They colonize at least 80% of plant species, promote plant growth and plant diversity. These fungi are multinucleated and contain either one or two haploid nuclear genotypes (monokaryon and dikaryon) identified by the alleles at a putative mating-type locus. This taxon has been considered as an ancient asexual scandal because of the lack of observable sexual structures. Despite identification of a putative mating-type locus and functional activation of genes related to mating when two isolates co-exist, it remains unknown if the AMF life cycle involves a sexual or parasexual stage. We used publicly available genome sequences to test if Rhizophagus irregularis dikaryon genomes display signatures of sexual reproduction in the form of reciprocal recombination patterns, or if they display exclusively signatures of parasexual reproduction involving gene conversion. We used short-read and long-read sequence data to identify nucleus-specific alleles within dikaryons and then compared them to orthologous gene sequences from related monokaryon isolates displaying the same putative MAT-types as the dikaryon. We observed that the two nucleus-specific alleles of the dikaryon A5 are more related to the homolog sequences of monokaryon isolates displaying the same putative MAT-type than between each other. We also observed that these nucleus-specific alleles displayed reciprocal recombination signatures. These results confirm that dikaryon and monokaryon isolates displaying the same putative MAT-type are related in their life-cycle. These results suggest that a genetic exchange mechanism, involving reciprocal recombination in dikaryon genomes, allows AMF to generate genetic diversity.

Introduction

Arbuscular mycorrhizal fungi are plant endosymbionts, forming symbioses with most plant species, promoting plant growth [1], plant community diversity [2, 3] and affect how plants cope with biotic [4] or abiotic stresses [5]. As a consequence, they are widely used in
In a single agricultural field, the presence of at least 17 different genotypes of *Rhizophagus irregularis* displaying spatial genetic structure have been detected [7, 8]. Furthermore, these *R. irregularis* isolates display different levels of within isolate genetic diversity [9], which has been reported to produce differential effects on plant growth [10]. Understanding how genetic variability is generated in AMF, is important because it could be harnessed to generate genetic variants that could be beneficial for their plant hosts [11].

AMF are part of the Glomeromycotina subphylum [12], which fossil records date to at least ~400 Million years ago [13]. They are coenocytic (without septa separating otherwise adjacent compartments), their hyphae harbor hundreds of nuclei within the same cytoplasm [14] and no single-nucleus state has been recorded. The nuclei of these fungi have been reported as haploid [15–18]. This group of fungi has been previously considered as an ancient asexual scandal [19], due to low morphological diversification and the absence of observable sexual structures. However, different reports suggest that sexual reproduction could be possible in AMF since these fungi contain a complete meiosis machinery [20]. A putative mating-type determining locus (MAT) has been proposed [16], population genetic data suggests the existence of recombination in AMF populations [21] and activation of genes related to mating has been detected when different isolates of the same species co-exist in plant roots [22, 23].

Several *R. irregularis* isolates issued from the same geographic location have been reported to have one haploid nuclear genotype (monokaryon: isolates A1,B12 and C2) and two haploid nuclear genotypes (dikaryon: isolates A4, A5, C3 and G1) [16, 24–26], evidencing that monokaryon and dikaryon isolates co-exist in the same location. Single nucleotide polymorphisms (SNP) profiles from single-nucleus from dikaryon isolates (A4, A5 and SL1) cluster into two genetically different groups, that are associated with the identity of a putative MAT-locus [24]. This demonstrates that the presence of two copies of the putative MAT-locus is a reliable marker of the dikaryon state.

Like most fungi, AMF can undergo anastomosis, the fusion of hyphae. Through these connections, bi-directional flow of cytoplasm has been observed between genetically different AMF individuals [27]. Via anastomosis, the transfer of genetic material between vegetative compatible isolates (parasexuality) has been suggested as a mechanism of maintenance of genetic diversity in the absence of sexual recombination [28]. In fungi, hyphal fusion between different individuals leads to cell death, however, non-self vegetative compatibility has also been observed in AMF when different isolates form perfect hyphal fusions [29]. In the model species *Aspergillus nidulans* parasexuality involves fusion of two haploid nuclei, mitotic recombination and haploidization of the diploid nuclei by i.e. chromosome loss [30]. In consequence, a genomic signature of parasexuality is gene-conversion where there is a loss of heterozygosity. Although sexual reproduction can also produce signatures of gene-conversion, sexual reproduction also involves meiotic recombination and allele segregation on reciprocal products [31].

The existence and relevance of sexuality and/or parasexuality for the evolution of AMF remains unknown [32]. It has been hypothesized that monokaryon isolates could fuse to form dikaryon isolates during the AMF life-cycle [16]. However, it is still unclear whether the transition between a dikaryon and monokaryon life stages involves a sexual event involving meiotic recombination, or a parasexual event.

In the absence of stable transformation methods in AMF [33], analyses of continuous genomic sequences (haplotype analyses) could be an important resource to identify genomic signatures of recombination in dikaryon isolates. Nucleus-specific alleles from dikaryon isolates (identified through haplotype phasing) can then be compared to orthologous sequences of related monokaryon isolates (that share the same MAT-type) and could allow to identify
genomic signatures of a sexual event, involving meiotic recombination, or a parasexual event which results in the loss of heterozygosity or gene conversion.

AMF genomes display large gene duplication events [18], which make it difficult to distinguish nucleus-specific sequences (orthologs) from sequence duplications (paralogs) in dikaryon isolates. The analysis of nucleus-specific sequences could be used to better differentiate orthologs from paralogs. The identification of nucleotype specific sequences in dikaryon isolates could be made by: 1) Drop in coverage analyses: use of short-read sequences to identify genome-wide copy number variation. These analyses consist in obtaining the read depth, or coverage, after mapping the reads to a genome assembly and identify changes in coverage across the genome [34]. A drop in coverage could represent that the sequences are nucleotype specific. In AMF, a drop in coverage analysis was used to originally identify a putative MAT-locus [16]. 2) Analyzing sequences issued from long-reads sequencing platforms to identify haplotypes and, consequently, nucleotype-specific sequences in fungal dikaryon isolates [35].

Here, we demonstrated that AMF dikaryons display reciprocal recombination genomic signatures by analyzing nucleotype-specific sequences in dikaryon (A5) and monokaryon isolates (A1-C2). In this study we used publicly available data of bulk whole genome short-reads sequencing, single-nucleus short-read sequence data and long-read bulk sequence data to identify nucleotype-specific sequences in dikaryon isolates. We identified regions displaying drops in coverage in short-read whole genome sequence data. In these regions we detected the presence of genes that have two alleles in the dikaryon isolates and one allele in the respective monokaryons. We then confirmed independently, with short-read genome sequence data from single-nucleus, that in dikaryon isolates, different nuclei have different alleles and that they are not always associated to the nucleus genotype (putative MAT identity), evidencing a reciprocal recombination genomic signature. Finally, we validated the analysis, by evaluating long-read genome sequence assemblies and confirmed that A5 dikaryon display genomic signatures of reciprocal recombination.

**Materials and methods**

**Source data**

We used public-available sequence reads, genome assemblies and annotations of isolates A1, A4, A5, C2 of *R. irregularis* for this study, including data from bulk-isolate and single-nucleus sequencing (S1 Table). We used short-read whole genome assemblies from isolates A1, A4, A5 and C2 [16]. We used single-nucleus raw short-reads to generate genome assemblies from individual nuclei of isolates A1, A5 and C2 [24]. We also analyzed long-read genome assemblies of isolates A1, A5 and C2 [36]. The long-read genome assembly of the dikaryon isolate A5 is a phased assembly, where the contigs are divided in two parts, the primary assembly and the haplotig assembly. We downloaded the sequence reads from the sequence read archive (SRA) using the SRAtoolkit software with the prefetch and fastq-dump tools [37].

**Coverage analysis on bulk sequencing short-reads**

We first trimmed the bulk sequence reads using Trim Galore! [38] with the default parameters. We then used BWA [39] to index the reference genome assemblies and BWA mem -M [39] to map the reads to the reference whole-genome assemblies. We mapped the reads coming from a given isolate to the reference genome assembly of the same isolate (i.e. reads A1 mapped to reference A1). We then kept the reads that display a mapping quality of at least 30. We used the genomecov tool from bedtools [40] to calculate the coverage for each position. We then created a ready-to-use algorithm that detects genome-wide drop in coverage analysis in whole-genome data (S1 File). The algorithm divides the data in portions of 50kb. Then, with a
sliding window approach consisting of windows of 400bp and steps of 100bp, the algorithm searches for drops in coverage of 0.3–0.6 times lower than the median coverage of the entire genome and that with a minimum length of 1000bp (Please refer to S1 File for the algorithm specifications implemented in the R programming language). We then further filtered the regions exhibiting a drop in coverage by keeping only the regions that display an average of 1.25 coverage difference between the neighboring regions and the drop in coverage region.

**Gene detection in coverage drops**

We identified all the genes located within genomic regions that presented a drop of sequencing coverage. We used the `intersect` command from the BEDTools suite with the existing gene annotations corresponding to each *R. irregularis* isolate (GTF format) and their query regions with drops in coverage (BED format) to identify overlapping genes [40]. Genes in scaffolds smaller that 1kb were not considered for further analyses.

**de novo single-nucleus assemblies**

We trimmed the single-nucleus raw reads by using TrimGalore-0.6.0 [38] with default parameters. After trimming, we performed single-nucleus *de novo* assemblies with SPAdes v3.14 [41] with the following parameters: -k 21,33,55,77—sc—careful—cov-cut off auto. The resulted single-nucleus genome assemblies were used for further analysis. The length, number of contigs and N50 value of the *de novo* assemblies was evaluated with quast- 5.1.0rc1 with default parameters [42].

**Identification of genes in genome assemblies**

To identify the position of gene sequences on the different genome assemblies, we first extracted a query sequence. We then used the console NCBI+ blast suite [43] to blast the query against the desired target. In the case of the putative MAT-locus, we used the homeodomain genes HD2 and HD1-like as query (HD2:KT946661.1, HD1-like: KU597387 from isolate A1). For further downstream analyses, we extracted the sequences from the genome assemblies by using the blastdbcmd command from the NCBI+ suite. We used a reciprocal blast approach to identify the gene sequences corresponding between the whole genome sequence data and the single-nucleus data. We considered the best hits by evaluating the % identity, mismatches, e-value and bitscore. We used the reciprocal blast in the comparison between gene sequences in the short-reads bulk assemblies and the single nucleus assemblies. We do not expect that a high gene copy number plays a role at this step because we worked with highly filtered data. i.e. we are sure that the genes to compare display two copies in the dikaryon and 1 in the monokaryon. So, from the resulting blast output we keep only the best match of the sequence.

**Orthology inference**

We used Orthofinder 2.3.11 [44] to identify orthologs of genes found inside the drop in coverage regions within the same isolate. We also identified orthologs in the long-read assemblies and identified single copy orthologs in isolates A1, C2, the primary assembly of A5 and the haplotig assembly of A5. We used the orthogroups output from Orthofinder for the different analyses.

**Synteny plots**

We compared genomic regions by performing synteny plots computed with EasyFig2.2.3 [45]. We provide full Genbank files of each contig to compare genomic regions to each other. The
software executes a blast comparison between the regions to determine their homology. We used the default parameters for the blast. The figure result can be modified by using the tools on the image menu.

**Genetic distance between nucleus-specific haplotypes on short-reads data**

Coding sequences for the 12 confirmed nucleotype-specific genes were extracted using the Blast+ command line blastdbcmd tool [43]. The sequences were then aligned with MAFFT [46] using the—auto option. Then, the ape package [47] of R was used to calculate the pairwise distance between the 4 alleles (2 from A5, and 1 each from A1 and C2).

**Recombination detection in short-reads data**

We compared the sequences from drop in coverage regions from both nuclei specific haplotypes of isolate A5 and isolates A1 and C2 to detect if isolate A5 display recombination events between the two putative parental isolates. After identification of the syntenic region among the different isolates, we aligned the sequences with MAFFT [46] and evaluated whether the sequence of one of the nucleotype of isolate A5 was similar to A1 and the other similar to C2.

**Recombination detection in long-reads data**

We identified genomic regions in isolate A5 that display long phased sequences containing more than 10 genes. We then identified single-copy orthologs in the monokaryon isolates A1, C2 and on the phased assembly of isolate A5 (primary and haplotig assemblies). We then visualized the phylogenetic relation of each single-copy ortholog among the four genome assemblies by using the gene tree output Orthofinder [44]. We considered a sequence as recombined when within the same continuous phased sequence, we observed that an allele of the A5 primary assembly clusters with an allele of isolate A1, and then a contiguous gene of the same A5 primary assembly clusters with the allele of isolate C2. We did not compare phased sequences separated by unphased sequences because this could lead to artificial recombination signals.

**Phylogenetic analyses**

We used MEGA-X [48] for the different phylogenetic reconstructions shown in the study. We find the best DNA models describing the relation between the sequences. Finally, we used a maximum likelihood phylogeny reconstruction with 100 bootstraps to infer the phylogenetic relation among the samples. In several cases, we were not able to perform maximum likelihood phylogenies because of the low number of samples to compare, so UPGMA trees were done instead. Phylogenetic reconstructions of the different orthologous groups on Fig 4 were produced by the Orthofinder software.

**Results**

**Drop in coverage analysis on bulk short-reads genome assemblies reveals potential nucleotype-specific haplotypes**

Previously, a drop in coverage analysis of selected regions was used for the identification of a putative MAT-locus in *R. irregularis* [16]. To perform a similar analysis across the entire genome, we developed a script (S1 File) that allows us to identify genome-wide drop in coverage events (for the accessions of raw data and genome assemblies used in this study see S1 Table).

We identified drops in sequencing coverage in 4 different isolates of *R. irregularis* which are reported to be dikaryons (A4 and A5) and monokaryons (A1 and C2) (Fig 1, S2 Table). The
number of coverage drops and genes inside the drop in coverage regions were different between dikaryons (A4-A5) and monokaryons (A1-C2) (S3 Table). These results indicated that dikaryotic isolates displayed more heterozygous regions than the monokaryons, suggesting that the genes present within the regions showing a drop in coverage are potential candidates for highly divergent, nucleotype-specific alleles in dikaryons. Confirming the reliability of our approach, we detected the expected drop in coverage in the putative MAT-locus region in isolates A4 and A5 but not in isolates A1 and C2 (Fig 1D).

One cause for a drop in coverage could be copy number variation between the nucleotypes in a dikaryon. To test for this, we inferred orthologous gene families among the different isolates to identify if the genes present in the drop in coverage regions displayed more than one copy in their own genome. We used the gene annotation available for each isolate and inferred the orthology of all the genes present in each genome. Many orthologous groups had more than one copy within each isolate (A4: 20%, A5: 18%, A1: 17% and C2: 19%; Fig 1 and S1 Fig, S4 Table) consistent with the high reported incidence of paralogs in these fungi [18]. We further identified orthologous gene families of genes detected in drop in coverage in isolates A4 and A5 independently. Under the assumption of a monokaryon-dikaryon genome organization in *R. irregularis*, to avoid the confounding effect of duplications and reduce the complexity of the dataset, we retained only the orthologous groups that are present in the drop in coverage regions and that display two copies in the dikaryon isolates (A4, A5) and a single copy in the monokaryons (A1, C2) (Fig 2A, S5 Table). In the regions where a drop in coverage was detected, we identified 32 orthologous groups that are present with two copies in isolate A4 and only a single copy in isolates A1 and C2. We also identified 27 orthologous groups in isolate A5 that display two copies. Only two orthologous groups were common between the two isolates: namely, HD2 and HD1-like which are part of the putative MAT-locus in *R. irregularis* (Fig 2B). As reported in Ropars *et al.*, we observed that the two copies of the putative MAT-locus in the dikaryons were located in different contigs. One copy of HD2 and HD1-like genes were present in a long contig of the genome assembly, while the second copy was present in a much shorter contig (Fig 2C). We observed the same pattern for the other orthologous groups, where the second copy was always present in a second shorter contig (for several examples see Fig 2D).

To further sort paralogous genes from true orthologs, we performed a synteny analysis to compare the genomic location of the presumed orthologous genes among isolates. We identified that 12 out of 32 predicted orthologs in A4 and 16 out of 27 predicted orthologs in A5, were located in the same genomic location on the different isolates, suggesting that they should be considered as orthologs (S5 Table, for examples of inferred orthologs and paralogs see Fig 2, S2 Fig).

Hence, in the whole genome assemblies of dikaryon isolates, two divergent alleles were assembled into different scaffolds; one longer scaffold containing neighboring regions and a shorter scaffold without the neighboring regions. Given that the nuclei are haploid in the dikaryon isolates, two possibilities are consistent with this previous fact: The two copies could be present within the same or in different nuclei.

**Drop in coverage signatures represent nucleotype-specific sequences**

To confirm that genes found inside drop in coverage regions are nucleotype-specific, we used sequencing reads of individual nuclei of dikaryon isolates A4 and A5 [24] to produce *de novo* single-nucleus assemblies. The *de novo* assemblies were very fragmented and incomplete (Fig 3, S3A and S3D Fig, S6 Table) and their utilization was highly limited. This limitation resulted in the inability to identify some genes and some complete gene sequences. However, a reciprocal blast approach between the whole genome assembly and the single-nucleus assemblies...
allowed us to detect sequences in the single-nucleus assemblies corresponding to the genes detected in the whole genome assemblies.

We tested in the dikaryons if the genes identified in the drop in coverage regions were present in the form of different alleles in different single nuclei by using a reciprocal blast approach. We confirmed 9 orthologous genes to be nucleotype-specific in isolate A4 and 12 orthologous genes in isolate A5 (S7 Table). We found that the population of nuclei clustered in two groups that corresponded to the identity of the putative MAT-locus contained in each nucleus (Fig 3). This result confirms that nucleotype-specific alleles in dikaryon isolates can be identified based on genes found in drop in coverage regions and that are represented by a duplication within the genome assembly.

**Nucleotype-specific alleles from A5 share a more recent evolutionary origin with monokaryon isolates A1 and C2 than among them**

The origin of dikaryon isolates could be investigated through comparisons of monokaryon isolates that display the same alleles of the putative MAT-locus, to those found in the dikaryons.
Fig 2. Identification of orthologs of genes present in regions showing a drop in coverage. 

a, Orthologous groups that display two or more genes in dikaryons and only a single gene in monokaryons. This analysis was performed independently on isolate A4 and isolate A5. Orthologous groups where HD2 and HD1-like genes are present.
b, Venn diagram representing the number of shared orthologous groups within drop in coverage regions between isolates A4 and A5. Only two orthologous groups were shared between the isolates, they contain the putative MAT-locus genes HD2 and HD1-like.
c, Synteny plot between the two contigs containing the different alleles of the putative MAT-locus of isolates A4 and A5.
d, Synteny plot between the two contigs containing different alleles of other orthologous genes. Please note that the synteny figures are made from the public available annotations of each genome assembly.

Differences in size of open-reading frames (ORF) among isolates are due to differences in detection of ORF on each isolate and likely could be the result of the annotation process.

https://doi.org/10.1371/journal.pone.0270481.g002
To confirm the previous findings, for each previously defined nucleotype-specific gene, we compared the phylogenetic relationship of the two nucleotype-specific alleles in isolate A5 isolate and in isolates A1, C2 and A4. We observed that for several nucleotype-specific alleles, the allele from isolate A1 clustered with one of the alleles of isolate A5, but it was not always the case for isolate C2 (Fig 4A).

We then analyzed the genetic distance of each of the 12 nucleotype-specific genes independently between the two alleles of isolate A5 and the homologous allele in isolates A1 and C2. We observed that for several nucleotype-specific alleles, the allele from isolate A1 clustered with one of the alleles of isolate A5, but it was not always the case for isolate C2 (Fig 4A).

We then analyzed the genetic distance of each of the 12 nucleotype-specific genes independently between the two alleles of isolate A5 and the homologous allele in isolates A1 and C2. The mean nucleotide distance between the two A5 alleles was 0.147. The mean distance between A1 and C2 was 0.132. In contrast, the mean of the minimum distance between an allele of isolate A5 and isolate A1 was 0.013 and between A5 and C2 was 0.043 (Fig 4B). We did not observe any case where the two A5 alleles clustered together, instead we observed that for all 12 nucleotype-specific gene the two A5 alleles were more similar to the allele from isolate A1 or C2 (Fig 4C). The mean distances calculated between alleles in this study are much higher than average distances calculated on the whole genome between different isolates [50], reflecting our selection criteria for nucleotype-specific regions. As each of the A5 alleles was closer to A1 or C2, instead of the two A5 alleles being most similar, this indicates that the alleles of A5 share a more recent evolutionary origin with these monokaryons than the two alleles within A5.

Fig 3. Single-nucleus sequence data confirms that genes contained in regions where a drop in coverage was observed are nucleotype-specific. Phylogenetic reconstruction of single-nucleus for genes found in regions where a drop in coverage was detected in A4 and A5 isolates. The genes are named by their membership to the orthologous groups previously defined. Branch support consisting of 100 bootstraps is shown. When only sequences from three nuclei were included, we performed an UPGMA hierarchical clustering. a, data for nuclei from A4 isolate. b, data for nuclei from A5 isolate.

https://doi.org/10.1371/journal.pone.0270481.g003
Identification of reciprocal recombination between nucleotype-specific haplotypes in isolate A5 using single-nucleus data

Knowledge about nucleotype-specific alleles of dikaryon isolate A5 and their orthologs in isolates A1 and C2 allowed us to test whether parasexual or sexual genomic signatures could be identified in dikaryon isolate A5 (Fig 5A). We scanned the different nucleotype-specific genes for the detection of recombination events within the two haplotypes of isolate A5. Comparison of gene sequences issued from the short-read whole genome assemblies from isolates A1, C2 and the two haplotypes of A5 showed that each nucleotype-specific allele from isolate A5 was highly similar to either the allele of isolate A1 or C2, but we did not identify any recombination events on these sequences (S4 Fig).

To further assess the potential for clonal relationships between the two alleles within A5 and isolates A1 and C2, we compared the nucleotype-specific alleles on the single-nucleus assemblies from isolate A5 and their orthologs on the single-nucleus assemblies of isolates A1 and C2 (S8 Table). The difference with the previous analysis is that with the single-nucleus assemblies from isolate A5 and their orthologs, we used the minimum distance of the two alleles from A5, A1, and C2 to assess the potential for clonal relationships.
data, we are able to identify the identity of the A5 alleles by identifying the identity of the putative MAT-type on each nucleus. We found that for several nucleotype-specific genes (i.e. OG2995, OG3981 and OG4715), the alleles from isolates A5 (MAT-3 type) and A1 (MAT-3 type) clustered together (S5 Fig, Fig 5B). However, we found that for other nucleotype-specific genes (OG4925, OG4492 and OG4493), the alleles from isolate A5 (MAT-3 type) clustered with the alleles from isolate C2 which has a MAT-6 type (Fig 5, S5 Fig, Fig 5C). The alignments on these nucleotype-specific genes show that A5 nuclei with MAT-3 have similar, but not identical alleles as C2 nuclei (MAT-6 type). In the same way, A5 nuclei with MAT-6 type harbor alleles similar to those of A1 nuclei (MAT-3 type). These results demonstrate that the A5 isolate harbors two nuclei genotypes, where one genotype is highly similar to the genotype displayed on nuclei of isolate A1 and the other A5 genotype is similar to the genotype of isolate C2. However, the A5 nuclei that is similar to the A1 nuclei, contain regions that are more similar to the equivalent region of isolate C2. This happens as well, for the A5 nuclei that is similar to isolate C2, where some regions are more similar to the equivalent region of isolate A1, demonstrating a reciprocal recombination pattern. The presence of reciprocal recombinant nuclei in isolate A5, involving isolates sharing the same MAT-type, strongly suggest that isolate A5 results from a sexual and not parasexual event between isolates similar to A1 and C2.

**Confirmation of reciprocal recombination events in isolate A5 using long-read genome assemblies**

We compared single-copy orthologous genes among the A1, C2 assemblies and the phased assembly of isolate A5, which is divided in a primary assembly and an haplotig assembly. The long-read assemblies were more complete and contain longer scaffolds than the short reads assemblies (S3B, S1C, S1E and S1F Fig). The A5 phased assembly consisted of a primary

---

**Fig 5. Recombination events in nucleotype-specific genes in A5 isolate.**

- **a** Schematic representation of possible outcomes after fusion of two different isolates. Please note the schema illustrates different contigs, separated by blank lines and no different chromosomes. **b** Phylogenetic relationship of different nucleotype-specific alleles among different nuclei from A1, A5 and C2 isolates. Cases where no recombination was detected. Nuclei having the same MAT-type clustered together. **c** Phylogenetic relationship of different nucleotype-specific alleles among different nuclei from A1, A5 and C2 isolates. Cases where recombination was detected. Nuclei having the same MAT-type did not clustered together. We performed 100 bootstraps for the branch support of all phylogenetic constructions.

https://doi.org/10.1371/journal.pone.0270481.g005
assembly of 392 scaffolds and an haplotig assembly on 292 of the primary scaffolds. The primary assembly was 115Mb long and the haplotig assembly was 19.2Mb [36].

We identified 1250 single-copy orthologs among isolates A1, C2, the A5 primary assembly and the A5 haplotig assembly (S9 Table). We then focused only in the orthologs were one of the genes of A5 clustered with either A1 or C2. The majority of the gene sequences were very similar among all the isolates, however we identified 44 orthologous groups where one of the alleles of A5 isolate cluster with an allele of isolate A1 or C2 (23 genes A1-A5haplotig and 21 genes A1-A5primary). We observed that for several contiguous phased regions, a gene displayed the A5primary allele clustering with A1 allele and in another gene, contained in the same contiguous region, the A5 primary allele clustered with the C2 allele (Fig 6), confirming that reciprocal recombination signatures are found within continuous haplotypes (S9 Table).

Discussion

In this study, we used published short and long read sequencing data to demonstrate that nucleotypes from the dikaryon isolate are more genetically related to monokaryon isolates displaying the same putative MAT-type than between each other. We also demonstrated that signatures of reciprocal recombination can be detected in genomes of dikaryon AMF isolates with the help of related monokaryon isolates that display the same putative MAT-type as the dikaryon. This study suggests that in the AMF life cycle there could be a transition between monokaryon and dikaryon isolates sharing the same putative MAT-type. It also suggests that a
genetic exchange, involving reciprocal recombination, could be another mechanism generating genetic variability in AMF.

In the absence of experimental evidence that isolates with different putative MAT-types are involved in sexual reproduction, genomic signatures of recombination can help us to understand whether sexual or parasexual reproduction are involved in the AMF life cycle. On one hand, parasexual reproduction genomic signatures involves gene-conversion, which result is the loss of heterozygosity [51]. On the other hand, reciprocal recombination associated with a putative sex-determining region (MAT-type) is a genomic signature of sexual reproduction, although gene-conversion could also happen in sexual reproduction [52]. It could be plausible that parasexual recombination could lead to a reciprocal recombination pattern by fusion of two monokaryons, mitotic reciprocal recombination, followed by reciprocal chromosome loss, however this cascade of events seems less plausible.

In this manuscript, we used single-nucleus short read sequence data which produced highly fragmented and incomplete single-nucleus assemblies. We were able to work with this data because it served as a confirmatory dataset. We would not be able to identify recombination signatures relying only on the single-nucleus sequence dataset. The identification of genomic regions to test the hypothesis was made on the bulk short-read genome assemblies. However, with the bulk genome assemblies we were not able discriminate which of the two copies of the dikaryon is more similar to A1 or C2, because we did not have phasing information. It was at this step when the single-nucleus assemblies are useful. The single nucleus assemblies contained phasing information. Each nucleus had been sequenced independently and the information of the putative MAT-locus was associated to the nucleus ID. Consequently, there is an ID number (e.g. SN01) of a given nucleus and then its assembly is associated with that ID. Then the nucleus ID can be associated with the sequence of the putative MAT-locus. This makes it possible to associate other sequences at other genomic regions to the putative MAT-locus ID. We then were able to test our hypothesis by querying each nucleotype-specific gene detected with the bulk short-read data on the different types of nucleus (which are defined by their putative MAT locus).

We identified few regions that display recombination patterns within continuous haplotypes in the dikaryon isolate A5. The principal reason, is that we did not evaluate all the different genes present, but only the genes that display only 1 copy in a haploid phased contig. We only compared single-copy orthologs among the different assemblies, to avoid the confounding effect of paralogy on our results. In consequence, the examples of reciprocal recombination shown in this study, are only a small subset of the total plausible recombination spots present in their genomes. With the information of nucleotype-specific haplotypes identified with the drop in coverage analysis, we found that nucleotypes of isolate A5 were as little 1% diverged from isolate A1 and 4% diverged from isolate C2, suggesting that isolates sharing the same MAT-type as the monokaryon isolates A1 and C2 are closely related ancestors from which the dikaryon A5 arose. Interestingly, with the long-read data, we identified several loci where the A1 and C2 genes were more closely related among them than to the alleles of isolate A5. This suggests, that there could be other evolutionary forces shaping AMF genome evolution. AMF genomes could be influenced by horizontal gene transfer from the host plant or bacteria [53]. Transposable elements (TE) have been reported to be responsible of genome duplications, inversions, insertions and deletions [18, 54, 55]. In nature, AMF co-exist with different conspecifics and they co-exists as well with different host-plants. We could speculate, that gene trajectories within AMF populations could be also driven by TE-mediated horizontal gene transfer events. However, an important sampling of AMF genomes is necessary to better understand the genome architecture features and life-history of the different gene families [56].
An intriguing question, is how only two coexisting nucleotypes and a well-orchestrated mechanism such as meiosis could be achieved without the presence of a single-cell stage in the AMF life cycle. A genetic bottleneck, where only 1 or 2 nuclei co-occur has never been detected in AMF. Despite the last feature, two studies that evaluated nuclear imbalance in dikaryon isolates show that nuclei ratio among single-spore lines are conserved despite multiple culturing for years [57] or are stable when exposed to different host plants [26]. These studies suggest that a nuclei regulation mechanism should exist and could be responsible of the stability of the nucleotypes in the dikaryons.

In this study, we compared haplotypes within the dikaryon A5 to monokaryon isolates (A1 and C2) that contain the same putative MAT-types as the dikaryon A5 to identify the recombination patterns. We were able to analyze these signatures on the A5 dikaryon, but not on the dikaryon A4, because there are no available genomic assemblies of monokaryon isolates displaying the same putative MAT-type as in Isolate A4. We hypothesized that MAT-type compatibility exists between the two different putative MAT-types identified in the dikaryon isolates. However, to date there is no direct experimental evidence that different putative MAT-types could be compatible. An indirect evidence of compatibility between different putative MAT-types was found, when two different isolates harboring different putative MAT-types elicited a putative fungal mating response [22, 23]. However, Mateus et al., did not test if the mating of two co-existing isolates took place [22]. Consequently, in order to experimentally identify mating and MAT-type compatibility, crossing experiments between strains harboring different putative MAT-types, including transcriptome and recombinant progeny analyses should be performed.

One of the main findings in this study is that in dikaryon short-read assemblies, if the two alleles of the same gene are highly divergent (i.e. like idiomorphs), the alleles are dispatched in two separate contigs. One allele can be found in a large contig containing the neighboring region of the allele and displaying a drop in coverage, whereas the second allele, without its neighboring region, is present in another short contig. We were able to demonstrate this by looking at the orthologous relationships between the two alleles of the gene present in the two different contigs. Our approach is reliable because we also detect the putative MAT-locus which is present in two separate contigs in the Illumina assemblies.

Our “drop in coverage” approach allowed us to identify divergent nucleotide-specific alleles in dikaryon isolates situated in different contigs of the short-read whole genome assemblies. This approach differs from previous approaches of global intra-isolate divergence assessment that measured the number of SNPs [58] or poly-allelic sites [9]. Although the comparison of both types of measurements gave similar information (intra-isolate divergence), their comparison should be carefully addressed as their methodology and the types of sequences compared are different. While SNPs are best identified in low divergence regions, where reads can be confidently mapped to the same contig, the sequences with large genetic differences, detected with the drop in coverage approach, are highly divergent to the point that they are assembled in different contigs in the same genome assembly. Consequently, genetic divergence between alleles detected in different genomic locations should be higher than when the two alleles are collapsed in the genome assembly. Contrary, to the drop in coverage analysis using short-read data, the analysis made on phased haplotypes from long-read sequencing, display genes that differ in several SNPs between the two dikaryon haplotypes.

Inter-nucleus recombination has been previously reported [24], although the robustness of the analysis has been questioned. The application of strict filtering parameters such as removal of heterozygous sites in haploid nuclei, duplicated regions of the genome, and low-coverage depths base calls results in an extreme loss of the signal of recombination [59]. Although some of these limitations, as coverage depth and filtering-out heterozygous sites were addressed
other limitations such as replicability (recombination events shown in several nuclei sharing the same putative MAT-type) and issues related to the whole-genome amplification step as the formation of chimeric sequences [60], allelic drop-out [61] and SNP miscalling [62] are inherent limitations of the analysis of any single nucleus amplification data. Furthermore, the previously reported evidence of large inter-nuclei recombination [24, 58] does not fit the observations about a monokaryon–dikaryons organization in AMF. In Chen et al., 2018a, the selected examples of genotypes presented, show 4 nuclei per isolate, which represent four different nucleus genotypes for SL1, A4 and three different nucleus genotypes on A5. The same pattern can be observed in Chen et al., 2020, the examples of genotypes show 7 out of 8 different nucleus genotypes in isolate A5, at least 7 different nucleus genotypes in isolate A4 and at least 5 different nucleus genotypes in SL1. The presence of repeated inter-nuclear recombination in dikaryons without prior crossing with other isolates, as observed in the single-nucleus genotypes shown in Chen et al. (2018a, 2020) would result in heterokaryons with more than two types of nucleus genotypes. But this does not appear to be the case for *R. irregularis* [16, 24, 59, 63].

Maintaining monokaryon and dikaryon isolates within the same natural population suggests that both forms are stable over time. Rather than a promiscuous mixing between isolates via anastomosis, a mechanism of recognition that involves a putative MAT-locus could regulate which isolates can form a dikaryon [64]. The fact that nucleotype-specific haplotypes from isolate A5 are more closely related to isolates A1 and C2, and that A5 nucleotypes display recombination, suggests that isolates sharing the same putative MAT-type as A1 and C2 could be the origin of a recombining dikaryon isolate. However, we cannot discard that these findings could apply to another step of the AMF life cycle. It could also be possible that a stable A5 isolate could segregate producing recombined monokaryons that share the same putative MAT-type as isolates A1 and C2, that can disperse and then fuse again to form stable dikaryons and complete a life cycle which involves recombination. It then becomes crucial to experimentally confirm if monokaryon isolates having different putative MAT-types could generate a dikaryon-like form and if a dikaryon isolate could segregate into recombining monokaryon isolates. It is difficult to establish in-vitro crosses mainly due to the difficulty of spore identification arising from the hyphae that fuse from the two parental isolates (an in-vitro culture produces thousands of spores). Hyphal fusion between different parental isolates has been reported [29, 65, 66]. We think that with technologies such as microfluidic channels developed for the study of filamentous fungi [67], it could be easier to identify hyphal fusions and the resulting spores from individual fusions. However, if recombination is a rare event between AMF genomes, which is a possibility, then it may be difficult to observe it in the laboratory.

Understanding the life cycle of AMF could have an enormous impact in the generation of AMF genetic variability. The generation of diverse AMF monokaryons or dikaryons could be used to generate variants that enhance plant growth and have an enormous potential in agriculture [11].

**Supporting information**

**S1 Fig. Number of copies of each orthologous group within each isolate.** Number of copies of different orthologous groups found within the genome of each isolate.

(PDF)

**S2 Fig. Synteny among candidate nucleotype specific regions in isolates A1, A4, A5 and C2.** a, b We show examples of orthologous genes on different isolates that are situated in the same genomic location in isolate A4 and isolate A5 respectively. This is evidenced by the blast homology (Grey zones linking the different contigs) shown in surrounding regions of the focal
gene (in red). c, d Examples of paralogous genes on different isolates where there genomic location of the gene is not the same. Evidenced by the lack of homology (grey zones linking the different contigs) of the surrounding regions of the focal gene (in red).

S3 Fig. Total length and N50 statistics of the genome assemblies used in this study. a, d Single-nuclei assemblies, b, e short-reads assemblies, c, f long-reads genome assemblies.

S4 Fig. Multiple sequence alignment of nucleotype-specific alleles of isolate A5 and isolates A1 and C2. Sequences are issued from the short-reads genome assemblies. The sequences shown are collapsed and do not represent the total length of the genes.

S5 Fig. Multiple sequence alignment of nucleotype-specific alleles of single-nuclei of isolate A5 and isolates A1 and C2. This example represents the case when no recombination is identified. Please note that MAT-3 or MAT-6 sequences cluster together. The sequences are issued from the single-nuclei genome assemblies. The sequences shown are collapsed and do not represent the total length of the genes.

S6 Fig. Multiple sequence alignment of nucleotype-specific alleles of single-nuclei of isolate A5 and isolates A1 and C2. This example represents the case when recombination is identified. Please note that MAT-3 or MAT-6 sequences do not cluster together. The sequences are issued from the single-nuclei genome assemblies. The sequences shown are collapsed and do not represent the total length of the genes.

S1 Table. Public available data used in this study.

S2 Table. Drop in coverage regions.

S3 Table. Genes present in drop in coverage regions.

S4 Table. Orthology inference among isolates A1, A4, A5 and C2.

S5 Table. Putative nucleotype specific genes in isolate A4 and their orthologs in isolates A1, A5 and C2.

S6 Table. Total length, number of contigs and N50 value of the different de novo single-nuclei genome assemblies.

S7 Table. Confirmation of equivalent nucleotype specific genes between bulk-sequencing data and single-nuclei data on isolates A4 and A5.

S8 Table. Confirmation of recombinant nucleotype specific genes in single-nuclei data from isolates A1, C2 and A5.
S9 Table. Identification of single-copy orthologues in long-reads data and recombination examples.
(XLSX)

S1 File. Script used for the identification of drop in coverage regions.
(TXT)

Acknowledgments
We would like to thank Daniel Croll for his feedback on a preliminary version of this manuscript. We specially thank Bart Nieuwenhuis and his group for the constructive feedback made on a bioRxiv version of this manuscript.

Author Contributions
Conceptualization: Ivan D. Mateus.
Data curation: Ivan D. Mateus, Joaquim Cruz.
Formal analysis: Ivan D. Mateus, Ben Auxier, Mam M. S. Ndiaye, Joaquim Cruz.
Funding acquisition: Ian R. Sanders.
Investigation: Ivan D. Mateus.
Methodology: Ivan D. Mateus.
Software: Mam M. S. Ndiaye.
Supervision: Ivan D. Mateus, Ian R. Sanders.
Validation: Soon-Jae Lee.
Writing – original draft: Ivan D. Mateus.
Writing – review & editing: Soon-Jae Lee, Ian R. Sanders.

References
1. Harrison MJ. The arbuscular mycorrhizal symbiosis: An underground association. Trends Plant Sci. 1997; 2: 54–60. https://doi.org/10.1016/S1360-1385(97)82563-0
2. van der Heijden M, Klironomos J, Ursic M, Moutoglis P, Streitwolf-Engel R, Boller T, et al. Mycorrhizal fungal diversity determines plant biodiversity, ecosystem variability and productivity. Nature. 1998; 74: 69–72. Available: http://www.nature.com/nature/journal/v396/n6706/abs/396069a0.html
3. Antunes PM, Koch AM, Morton JB, Rillig MC, Klironomos JN. Evidence for functional divergence in arbuscular mycorrhizal fungi from contrasting climatic origins. New Phytol. 2011; 189: 507–14. https://doi.org/10.1111/j.1469-8137.2010.03480.x PMID: 20880038
4. Thygesen K, Larsen J, Bødker L. Arbuscular mycorrhizal fungi reduce development of pea root-rot caused by Aphanomyces euteiches using oospores as pathogen inoculum. Eur J Plant Pathol. 2004; 110: 411–419. https://doi.org/10.1023/B:EJPP.0000021070.61574.8b
5. Augé RM. Water relations, drought and vesicular-arbuscular mycorrhizal symbiosis. Mycorrhiza. 2001; 11: 3–42. https://doi.org/10.1007/s005720100097
6. Smith SE, Read DJ. Mycorrhizal Symbiosis. Third edit. London: Elsevier Ltd; 2008.
7. Koch AM, Sanders IR, Kuhn G, Fontanillas P, Fumagalli L. High genetic variability and low local diversity in a population of arbuscular mycorrhizal fungi. PNAS. 2004; 101: 2369–2374. https://doi.org/10.1073/pnas.0306441101 PMID: 14983016
8. Croll D, Wille L, Gamper H a, Mathimaran N, Lammers PJ, Corradi N, et al. Genetic diversity and host plant preferences revealed by simple sequence repeat and mitochondrial markers in a population of the arbuscular mycorrhizal fungus Glomus intraradices. New Phytol. 2008; 178: 672–67. https://doi.org/10.1111/j.1469-8137.2008.02381.x PMID: 18298433
9. Wyss T, Masclaux FG, Rosikiewicz P, Pagni M, Sanders IR. Population genomics reveals that within-fungus polymorphism is common and maintained in populations of the mycorrhizal fungus Rhizophagus irregularis. ISME J. 2016; 10: 2514–2526. https://doi.org/10.1038/ismej.2016.29 PMID: 26953600
10. Angelard C, Colard A, Niculita-Hirzel H, Croll D, Sanders IR. Segregation in a mycorrhizal fungus alters rice growth and symbiosis-specific gene transcription. Curr Biol. 2010; 20: 1216–21. https://doi.org/10.1016/j.cub.2010.05.031 PMID: 20541408
11. Sanders IR. Designer mycorrhizas: Using natural genetic variation in AM fungi to increase plant growth. ISME J. 2010; 4: 1081–1083. https://doi.org/10.1038/ismej.2010.109 PMID: 20613789
12. Spatafora JW, Chang Y, Benny GL, Lazarus K, Smith ME, Berbee ML, et al. A phylum-level phylogenetic classification of zygomycete fungi based on genome-scale data. Mycologia. 2016; 108: 1028–1046. https://doi.org/10.3852/16-042 PMID: 27738200
13. Remy W, Taylor TN, Hass H, Kerh H. Four hundred-million-year-old vesicular arbuscular mycorrhizae. Proc Natl Acad Sci U S A. 1994; 91: 11841–11843. https://doi.org/10.1073/pnas.91.25.11841 PMID: 11607500
14. Marleau J, Dalpé Y, St-Arnaud M, Hijri M. Spore development and nuclear inheritance in arbuscular mycorrhizal fungi. BMC Evol Biol. 2011; 11: 51. https://doi.org/10.1186/1471-2148-11-51 PMID: 21349193
15. Lin K, Limpen E, Zhang Z, Ivanov S, Saunders DGO, Mu D, et al. Single nucleus genome sequencing reveals high similarity among nuclei of an endomycorrhizal fungus. PLoS Genet. 2014; 10: e1004078. https://doi.org/10.1371/journal.pgen.1004078 PMID: 24415955
16. Ropars J, Toro KS, Noel J, Pelin A, Charron P, Farinelli L, et al. Evidence for the sexual origin of heterokaryosis in arbuscular mycorrhizal fungi. Nat Microbiol. 2016; 1: 16033. https://doi.org/10.1038/nmicrobiol.2016.33 PMID: 27572831
17. Kobayashi Y, Maeda T, Yamaguchi K, Kameoka H, Tanaka S, Ezawa T, et al. The genome of Rhizophagus clarus HR1 reveals a common genetic basis for auxotrophy among arbuscular mycorrhizal fungi. BMC Genomics. 2018; 19: 1–11. https://doi.org/10.1186/s12864-017-4368-0
18. Morin E, Miyauuchi S, Can Clemente H, Chen ECH, Pelin A, Providencia I, et al. Comparative genomics of Rhizophagus irregularis, R. cerebriforme, R. diaphanus and Gigaspora rosea highlights specific genetic features in Glomeromycota. New Phytol. 2019; 222: 1584–1598. https://doi.org/10.1111/nph.15687 PMID: 30636349
19. Judson OP, Normark BB. Ancient asexual scandals. Trends Ecol Evol. 1996; 11: 41–46. https://doi.org/10.1016/0169-5347(96)81040-8 PMID: 21237759
20. Halary S, Malik S-B, Lildhar L, Slamovits CH, Hijri M, Corradi N. Conserved meiotic machinery in Glomus spp., a putatively ancient asexual fungal lineage. Genome Biol Evol. 2011; 3: 950–8. https://doi.org/10.1093/gbe/evr089 PMID: 21876220
21. Croll D, Sanders IR. Recombination in Glomus intraradicis, a supposed ancient asexual arbuscular mycorrhizal fungus. BMC Evol Biol. 2009; 9: 13. https://doi.org/10.1186/1471-2148-9-13 PMID: 19146661
22. Mateus ID, Rojas EC, Savary R, Dupuis C, Masclaux FG, Aletti C, et al. Coexistence of genetically different Rhizophagus irregularis isolates induces genes involved in a putative fungal mating response. ISME J. 2020. https://doi.org/10.1038/s41396-020-0694-3 PMID: 32514118
23. Mateus ID, Lee S, Sanders IR, Sanders IR. Co-existence of AMF with different putative MAT-alleles induces genes homologous to those involved in mating in other fungi: a reply to Malar et al. ISME J. 2021. https://doi.org/10.1038/s41396-021-00979-x PMID: 33941891
24. Chen EC, Mathieu S, Hoffrichter A, Sedzielewksa-Toro K, Peart M, Pelin A, et al. Single nucleus sequencing reveals evidence of inter-nucleus recombination in arbuscular mycorrhizal fungi. Elife. 2018; 7: 1–17. https://doi.org/10.7554/eLife.39813 PMID: 30516133
25. Masclaux FG, Wyss T, Mateus-Gonzalez ID, Aletti C, Sanders IR. Variation in allele frequencies at the bg112 locus reveals unequal inheritance of nuclei in a dikaryotic isolate of the fungus Rhizophagus irregularis. Mycorrhiza. 2018; 28: 369–377. https://doi.org/10.1007/s00572-018-0834-z PMID: 29675619
26. Kokkoris V, Chagnon P-L, Yildirir G, Clarke K, Goh D, MacLean AM, et al. Host identity influences nuclear dynamics in arbuscular mycorrhizal fungi. Curr Biol. 2021; 31: 1531–1538.e6. https://doi.org/10.1016/j.cub.2021.01.035 PMID: 33540434
27. Giovannetti M, Azzolini D, Citemesi AS. Anastomosis Formation and Nuclear and Protoplasmic Exchange in Arbuscular Mycorrhizal Fungi. Appl Environ Microbiol. 1999; 65: 5571–5575. https://doi.org/10.1128/AEM.65.12.5571-5575.1999 PMID: 10584019
28. Bever JD, Morton J. Heritable variation and mechanism of inheritance of spore shape within a population of Scutellosporea pellucida, An arbuscular mycorrhiz. Am J Bot. 1999; 86: 1209–1216. PMID: 10487808
29. Croll D, Giovannetti M, Koch AM, Sbrana C, Ehinger M, Lammers PJ, et al. Nonself vegetative fusion and genetic exchange in the arbuscular mycorrhizal fungus Glomus intraradices. New Phytol. 2009; 181: 924–37. https://doi.org/10.1111/j.1469-8137.2008.02726.x PMID: 19140939

30. Pontecorvo G. The Parasexual Cycle in Fungi. Annu Rev Microbiol. 1956; 10: 393–400. https://doi.org/10.1146/annurev.mi.10.100156.002141 PMID: 13363369

31. Andersen SL, Sekelsky J. Meiotic versus mitotic recombination: Two different routes for double-strand break repair. BioEssays. 2010; 32: 1058–1066. https://doi.org/10.1002/bies.201000087 PMID: 20967781

32. Yildirir G, Malar CM, Kokkoris V, Corradi N. Parasexual and Sexual Reproduction in Arbuscular Mycorrhizal Fungi: Room for Both. Trends Microbiol. 2020; xx: 1–3. https://doi.org/10.1016/j.tim.2020.03.013 PMID: 32360097

33. Helber N, Requena N. Expression of the fluorescence markers DsRed and GFP fused to a nuclear localization signal in the arbuscular mycorrhizal fungus Glomus intraradices. New Phytol. 2008; 177: 537–48. https://doi.org/10.1111/j.1469-8137.2007.02257.x PMID: 17999519

34. Yoon S, Xuan Z, Makarov V, Ye K, Sebat J. Sensitive and accurate detection of copy number variants using read depth of coverage. Genome Res. 2009; 19: 1586–1592. https://doi.org/10.1101/gr.092981.109 PMID: 19657104

35. Li F, Upadhya N, Sperschneider J, Matny O, Nguyen-Phuc H, Magro R, et al. Emergence of the Ug99 lineage of the wheat stem rust pathogen through somatic hybridisation. Nat Commun. 2019; 10: 5068. https://doi.org/10.1038/s41467-019-12927-7 PMID: 31699975

36. Chaturvedi A, Cruz Corella J, Robbins C, Loha A, Menin L, Gasilova N, et al. The methylome of the model arbuscular mycorrhizal fungus, Rhizophagus irregularis, shares characteristics with early diverging fungi and Dikarya. Commun Biol. 2021; 4: 901. https://doi.org/10.1038/s42003-021-02414-5 PMID: 34294866

37. Leinonen R, Sugawara H, Shumway M. The sequence read archive. Nucleic Acids Res. 2011; 39: 2010–2012. https://doi.org/10.1093/nar/gkq1019 PMID: 21062823

38. Krueger F. Trim galore. A wrapper tool around Cutadapt FastQC to consistently apply Qual Adapt trimming to FastQ files. 2015; 516: 517.

39. Li H. Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM. 2013; 00: 1–3. Available: http://arxiv.org/abs/1303.3997

40. Quinlan AR, Hall IM. BEDTools: A flexible suite of utilities for comparing genomic features. Bioinformatics. 2010; 26: 841–842. https://doi.org/10.1093/bioinformatics/btq33 PMID: 21012078

41. Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, et al. SPAdes: A new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol. 2012; 19: 455–477. https://doi.org/10.1089/cmb.2012.0021 PMID: 22506599

42. Mikheenko A, Prjibelski A, Saveliev V, Antipov D, Gurevich A. Versatile genome assembly evaluation with QUAST-LG. Bioinformatics. 2018; 34:i142–i150. https://doi.org/10.1093/bioinformatics/bty266 PMID: 29949999

43. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, et al. BLAST+: Architecture and applications. BMC Bioinformatics. 2009; 10: 1–9. https://doi.org/10.1186/1471-2105-10-421

44. Sullivan MJ, Petty NK, Beatson SA. Easyfig: A genome comparison visualizer. Bioinformatics. 2011; 27: 1009–1010. https://doi.org/10.1093/bioinformatics/btr396 PMID: 21278367

45. Katoh K, Rozewicki J, Yamada KD. MAFFT online service: multiple sequence alignment, interactive sequence choice and visualization. Brief Bioinform. 2017; 1–7. https://doi.org/10.1093/bib/bbx108

46. Kumar S, Stecher G, Li M, Knyaz C, Tamura K. MEGA X: Molecular evolutionary genetics analysis across computing platforms. Mol Biol Evol. 2018; 35: 1547–1549. https://doi.org/10.1093/molbev/msy096 PMID: 29722887

47. Savary R, Masclaux FG, Wyss T, Droh G, Cruz Corella J, Machado AP, et al. A population genomics approach shows widespread geographical distribution of cryptic genomic forms of the symbiotic fungus Rhizophagus irregularis. ISME J. 2018; 12: 17–30. https://doi.org/10.1038/ismej.2017.153 PMID: 29027999

48. Chen ECH, Morin E, Beaudet D, Noel J, Yildirir G, Ndikumana S, et al. High intraspecific genome diversity in the model arbuscular mycorrhizal symbiont Rhizophagus irregularis. New Phytol. 2018; 220: 1161–1171. https://doi.org/10.1111/nph.14989 PMID: 29355972
51. Forche A, Alby K, Schaefer D, Johnson AD, Berman J, Bennet RJ. The Parasexual Cycle in Candida albicans Provides an Alternative Pathway to Meiosis for the Formation of Recombinant Strains. Heitman J, editor. PLoS Biol. 2008; 6: e110. https://doi.org/10.1371/journal.pbio.0060110 PMID: 18462019

52. Sun S, Hsueh YP, Heitman J. Gene conversion occurs within the mating-type locus of Cryptococcus neoformans during sexual reproduction. PLoS Genet. 2012;8. https://doi.org/10.1371/journal.pgen.1002810 PMID: 22792079

53. Li M, Zhao J, Tang N, Sun H, Huang J. Horizontal Gene Transfer From Bacteria and Plants to the Arbuscular Mycorrhizal Fungus Rhizophagus irregularis. 2018; 9: 1–13. https://doi.org/10.3389/fpls.2018.00701

54. Daboussi M-J, Capy P. Transposable Elements in Filamentous Fungi. Annu Rev Microbiol. 2003; 57: 275–299. https://doi.org/10.1146/annurev.micro.57.030502.091029 PMID: 14527280

55. Venice F, Ghignone S, Salvioli A, Armselem J, Novero M, Xianan X, et al. At the nexus of three kingdoms: the genome of the mycorrhizal fungus Gigaspora margarita provides insights into plant, endobacterial and fungal interactions. 2020; 22: 122–141. https://doi.org/10.1111/1462-2920.14827 PMID: 31621176

56. Badet T, Croll D. The rise and fall of genes: origins and functions of plant pathogen pangenomes. Curr Opin Plant Biol. 2020; 56: 65–73. https://doi.org/10.1016/j.pbi.2020.04.009 PMID: 32480355

57. Robbins C, Cruz Corella J, Aletti C, Seiler R, Mateus ID, Lee S, et al. Generation of disproportionate nuclear genotype proportions in Rhizopus irregularis progeny causes allelic imbalance in gene transcription. New Phytol. 2021. https://doi.org/10.1111/nph.17530

58. Chen ECH, Mathieu S, Hoffrichter A, Ropars J, Dreissig S, Fuchs J, et al. More Filtering on SNP Calling Does Not Remove Evidence of Inter-Nucleus Recombination in Dikaryotic Arbuscular Mycorrhizal Fungi. Front Plant Sci. 2020; 11: 1–9. https://doi.org/10.3389/fpls.2020.00912

59. Auxier B, Bazzicalupo A. Comment on “Single nucleus sequencing reveals evidence of inter-nucleus recombination in arbuscular mycorrhizal fungi.” Elife. 2019; 8: 1–9. https://doi.org/10.7554/eLife.47301 PMID: 31650958

60. Yilmaz S, Singh AK. Single cell genome sequencing. Curr Opin Biotechnol. 2012; 23: 437–43. https://doi.org/10.1016/j.copbio.2011.11.018 PMID: 22154471

61. Lauri A, Lazzari G, Galli C, Lagutina I, Genzini E, Braga F, et al. Assessment of MDA efficiency for genotyping using cloned embryo biopsies. Genomics. 2013; 101: 24–9. https://doi.org/10.1016/j.ygeno.2012.09.002 PMID: 22982297

62. Ning L, Liu G, Li G, Hou Y, Tong Y, He J. Current challenges in the bioinformatics of single cell genomics. Front Oncol. 2014;4 JAN: 1–7. https://doi.org/10.3389/fonc.2014.00007

63. Masclaux FG, Wyss T, Pagni M, Rosikiewicz P, Sanders IR. Investigating unexplained genetic variation and its expression in the arbuscular mycorrhizal fungus Rhizophagus irregularis: A comparison of whole genome and RAD sequencing data. PLoS One. 2019; 14: 1–20. https://doi.org/10.1371/journal.pone.0226497 PMID: 31881076

64. Corradi N, Brachmann A. Fungal Mating in the Most Widespread Plant Symbionts? Trends Plant Sci. 2017; 22: 175–183. https://doi.org/10.1016/j.tplants.2016.10.010 PMID: 27876487

65. Avio L, Pellegrino E, Bonari E, Giovannetti M. Functional diversity of arbuscular mycorrhizal fungal isolates in relation to extraradical mycelial networks. 2002; 347–357.

66. Pepe A, Giovannetti M, Sbrana C. Different levels of hyphal self-incompatibility modulate interconnectedness of mycorrhizal networks in three arbuscular mycorrhizal fungi within the Glomeraceae. Mycorrhiza. 2016; 26: 325–332. https://doi.org/10.1007/s00572-015-0671-2 PMID: 26630971

67. Held M, Lee AP, Edwards C, Nicolau D V. Microfluidics structures for probing the dynamic behaviour of filamentous fungi. Microelectron Eng. 2010; 87: 786–789. https://doi.org/10.1016/j.mee.2009.11.096