Diploidy within a Haploid Genus of Entomopathogenic Fungi

Knud Nor Nielsen (p^{1,*}, João Felipe Moreira Salgado (p^{2,3}, Myrsini Eirini Natsopoulou¹, Thea Kristensen^{1,†}, Jason E. Stajich (p², and Henrik H. De Fine Licht¹

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Abstract

Fungi in the genus *Metarhizium* are soil-borne plant–root endophytes and rhizosphere colonizers, but also potent insect pathogens with highly variable host ranges. These ascomycete fungi are predominantly asexually reproducing and ancestrally haploid, but two independent origins of persistent diploidy within the Coleoptera-infecting *Metarhizium majus* species complex are known and has been attributed to incomplete chromosomal segregation following meiosis during the sexual cycle. There is also evidence for infrequent sexual cycles in the locust-specific pathogenic fungus *Metarhizium acridum* (Hypocreales: Clavicipitaceae), which is an important entomopathogenic biocontrol agent used for the control of grasshoppers in agricultural systems as an alternative to chemical control. Here, we show that the genome of the *M. acridum* isolate ARSEF 324, which is formulated and commercially utilized is functionally diploid. We used single-molecule real-time sequencing technology to complete a high-quality assembly of ARSEF 324. K-mer frequencies, intragenomic collinearity between contigs and single nucleotide variant read depths across the genome revealed the first incidence of diploidy described within the species *M. acridum*. The haploid assembly of 44.7 Mb consisted of 20.8% repetitive elements, which is the highest proportion described of any *Metarhizium* species. The long-read diploid genome assembly sheds light on past research on this strain, such as unusual high UVB tolerance. The data presented here could fuel future investigation into the fitness landscape of fungi with infrequent sexual reproduction and aberrant ploidy levels, not least in the context of biocontrol agents.

Key words: genome duplication, ploidy, parasexual, Metarhizium, entomopathogen, ascomycete.

Significance

Increasing evidence suggests that chromosome copy number variation is an adaptive trait of many fungi and present at low prevalence in natural populations. Here, we describe a rare case of diploidy and the first case within the species *Metarhizium acridum*, which in contrast to more intensely studied haploid *Metarhizium* species with broad host ranges such as *Metarhizium robertsii* and *Metarhizium anisopliae*, is a locust-specific pathogen with infrequent sexual reproduction. Using long-read sequencing, we show that the diploid genome of *M. acridum* is markedly different in repeat content and signature of the fungus-specific repeat-induced point mutation defensive machinery compared with generalist *Metarhizium* species. This is relevant for studying evolutionary patterns of genome structure and ploidy levels within this genus of important biological control agents.

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¹Section for Organismal Biology, Department of Plant and Environmental Sciences, University of Copenhagen, Frederiksberg C, Denmark

²Department of Microbiology and Plant Pathology, University of California Riverside, California, USA

³Instituto de Microbiologia Paulo de Góes, Universidade Federal do Rio de Janeiro, Brazil

[†]Present address: Division for Global Surveillance, National Food Institute, Technical University of Denmark, Lyngby, Denmark

^{*}Corresponding author: E-mail: knn@plen.ku.dk.

Introduction

The duplication of single genes, genomic segments, chromosomes, and whole-genome duplications (WGD) has played a vital role in eukaryotic evolution by providing genetic material for adaptation. Throughout evolution, recurrent WGD events have been linked to the emergence, diversification, and survival of species (Crow et al. 2006). For example, WGD has been coupled with the emergence of angiosperms within plants (De Bodt et al. 2005), and with the emergence of vertebrates, gnathostomes, and teleosts within metazoan chordate evolution (Sacerdot et al. 2018).

A well-described ancient WGD event in the fungal kingdom is the ancient diploidization that led to the formation of the Saccharomyces genus (Wolfe and Shields 1997; Dietrich 2004; Kellis et al. 2004). Although some WGD events lead to stable polyploids, as in the case of Saccharomyces where an ancient hybridization event is thought to have provided stability and fertility to the new diploid (Marcet-Houben and Gabaldón 2015), duplications are usually perceived as transient. Duplicated genes undergo subsequent reductions by loss or mutational decay leading to pseudogenization or neofunctionalization (Lynch and Conery 2000; Levasseur and Pontarotti 2011). It should be noted that it can be difficult to distinguish ancient WGD events from cases of multiple segmental duplications caused by transposon activity (Roelofs et al. 2020). Regardless of origin, duplications have long been recognized as a driver of evolution by providing the material for natural selection to work on (Bridges 1919; Metz 1947; Ohno 1970; Crow et al. 2006).

WGD arise from abnormal cell cycles post chromosome duplication, either by the absence of mitotic division or cytokinesis. The duplication itself can occur by auto- or allopolyploidization. The latter involves the fusion of individual cells and subsequent fusion of nuclei bringing together the variants accumulated between the genetically distinct individuals. In contrast, autopolyploidization, or endoreplication, only involves a single cell and has been associated with environmental stress. Within plants, several studies have shown that endoreplication can be induced by abiotic stress, such as heat (Monjardino et al. 2006), drought (Cookson et al. 2006), elevated salinity (Barkla et al. 2018), and toxins (Biskup and Izmaiłow 2004). Duplications are considered to mediate an acute increase in the transcription of metabolic and stressmitigating genes (Scholes and Paige 2015). The human pathogenic fungi Candida albicans and Cryptococcus neoformans have been shown to repeatedly gain resistance to antifungal drugs such as fluconazole through chromosomal duplications (Sionov et al. 2010; Kronstad et al. 2011). The fungus C. neoformans can also evade phagocytosis within the lungs by creating titan cells (Okagaki et al. 2010; Zaragoza et al. 2010). The increased cell size is induced by multiple rounds of endoreplication and is observed in up to one-fifth of the C. neoformans cells residing in infected lung tissue (Okagaki et al. 2010). Transient somatic endoreplication has also been recognized as a driver of developmental change of some cell types in plants, insects and mammals (Fox and Duronio 2013). Examples of somatic endoreplication include the giant cells of the mammalian trophoblast that develop into a significant part of the placenta, and the $>1,000\times$ sister chromatids observed in the salivary glands of *Drosophila melanogaster* (Sher et al. 2013).

Within fungi, increasing evidence suggests widespread cryptic population-level ploidy variations. Whole-genome sequencing of an extensive collection of 794 natural (wild) isolates of Saccharomyces cerevisiae found alternative ploidy levels within 13% of the otherwise predominately diploid yeast (Peter et al. 2018). The same study found chromosomal copy number variation (CCNV or aneuploidy) in 19% of the isolates (193 of the 1,011), reiterating the importance of variation in chromosome numbers for organismal evolution (Mayrose and Lysak 2020). Although at least some fungi readily undergo endoreplication, such levels of genome plasticity are not without constraints. A long-term evolutionary experiment with S. cerevisiae showed a convergence from alternative ploidies to the ancestral diploid form (Gerstein et al. 2006). The chytrid fungus Batrachochytrium dendrobatidis threatening amphibians worldwide has a high rate of CCNV in nature (Rosenblum et al. 2013). This plasticity has been linked to pathogenicity through lab experiments, which showed a gain of chromosomal copies upon induced stress by a characteristic host antimicrobial peptide (Farrer et al. 2013).

Although the majority of known fungi are dikaryotic at some stage, diploidy is the exception. Filamentous ascomycetes (Pezizomycotina) are only dikaryotic in specific life stages, namely the ascogenous hyphae and ascocarp, whereas the bulk of the mycelium remains monokaryotic, and in most species, haploid. Exceptions to haploidy within this important group of fungi are found in the Phyllactinia, Stephensia, Xylaria, Botrytis, and Zygosaccharomyces genera (Albertin and Marullo 2012), and within the genus Metarhizium (Kepler et al. 2016). When these fungi contain two different idiomorphs of the mating-type genes within the same diploid, it indicates mating events that failed to complete meiosis, leading to the observed allopolyploidization. Diploidy may also arise from the parasexual cycle of fungi, which allows for nuclei fusion (karyogamy) following anastomosis of strains from somatically compatible groups. Contrary to the sexual cycle, fused nuclei of the parasexual cycle do not undergo meiosis, but continue to divide mitotically. The original ploidy level is restored by random chromosome loss, through a series of aneuploidy intermediates (Moore et al. 2011), which is known from the "asexual" human pathogen Candida albicans. Reversion is a gradual process with generations of aneuploidy. Sexual reproduction requires different mating-types, whereas the parasexual cycle normally requires compatible identities. The reproductive isolation of the

parasexual cycle is enforced by heterokaryon incompatibility protein domains that abort incompatible hyphal fusion attempts (Glass and Dementhon 2006). This entails that any parasexual fusion of nuclei will be of closely related and highly similar genomes.

Although sexual reproduction facilitates the recombination of genomes, it also opens a door for disseminating "selfish DNA" such as transposons and other mobile sequences throughout the genome. Selfish DNA elements can have detrimental effects on host fitness why Ascomycete genomes have different modes of defence to silence transposons and ensure genome integrity. Some ascomycetes have a repeatinduced point (RIP) mutation pathway (Selker and Garrett 1988) that induce cytosine to thymine mutations (Cambareri et al. 1991; Watters et al. 1999) in any kind of repetitive element, which are thereby degraded and deactivated. Because the RIP pathway is considered only to be active during the dikaryotic phase between fertilization and nuclear fusion within the sexual cycle (Selker and Garrett 1988), it is also considered exclusively available to sexually reproducing fungi (Galagan and Selker 2004; Gladyshev 2017). The genomic footprint of RIP in the form of cytosine to thymine transitions that preferentially affect the frequency of CpA dinucleotides (Hane and Oliver 2008), can therefore be used to indicate a sexual cycle within fungal species where sex otherwise is not known. It is important not to associate the presence of an active RIP pathway with the absence of recombination, because recombination also occurs during the parasexual cycle (Arnau and Oliver 1993). However, although the RIP pathway should not be associated with recombination per se, an active RIP pathway is considered linked to the sexual cycle (Galagan and Selker 2004; Gladyshev 2017).

For entomopathogenic fungi, infected insects can be hot spots of fungal parasexuality, and the formation of diploid conidia (Riba et al. 1980; Leal-Bertioli et al. 2000; Wang et al. 2011). Investigating the conidia harvested from an insect coinfected with two fluorescently labeled isolates of the same *Metarhizium robertsii* strain revealed that 24% of the conidia investigated were diploid (Wang et al. 2011) and 13% of conidia when coinfected with different *M. robertsii* isolates (Li et al. 2021). Similar observations have been reported from experiments with *Metarhizium anisopliae* and *Metarhizium majus* (Riba et al. 1980; Leal-Bertioli et al. 2000).

The entomopathogenic fungus *Metarhizium acridum* (Hypocreales: Clavicipitaceae) is a specialist with a narrow host range of insects of the order Orthoptera. Host specificity is enforced by the requirement of Orthopteran cuticle signals facilitating spore germination and appressorial formation (Wang and St. Leger 2005). The *M. acridum* strain ARSEF 324 is the active ingredient in the biocontrol agent "Green Guard" targeting grasshoppers (Acrididae), which is commercially deployed in Australia where these insects are agricultural pests of concern. Within the genus *Metarhizium*, *M. acridum*, as well as the specialist *Metarhizium album*, are considered to

be cryptic sexual species (Hu et al. 2014; St. Leger and Wang 2020). No teleomorphs are known, but genomic analyses have found signatures of an active RIP pathway as an indirect sign of meiosis and sexual recombination (Galagan and Selker 2004; Gladyshev 2017). This is in contrast to the co-generic generalists with wide host ranges, *Metarhizium pingshaense*, *M. anisopliae*, *M. robertsii*, and *Metarhizium brunneum* (the PARB clade, sensu Bischoff et al. 2009), where no RIP footprint is present, consequently causing these species to be considered primarily asexual.

The ascomycete genus Metarhizium is generally haploid, but Kepler et al. (2016) found two independent origins of stable diploid clades interspersed among lineages comprised entirely of haploid individuals within the larger Metarhizium quizhouense/majus/taii clade (MGT clade; sensu Bischoff et al. 2009). All isolates within these diploid taxa were consistently heterozygotic for the analyzed microsatellite markers and contained two different idiomorphs of the mating-type genes (MAT1 and MAT2). The presence of both mating-types was interpreted as a mating event that failed to complete meiosis, which lead to the observed allopolyploidization. The locustspecific M. acridum has so far been regarded as entirely haploid, and previously the only available genome for this species was from the haploid strain CQMa 102. This strain was seguenced and assembled into 241 scaffolds (>1 kb; N50, 329.5 kb) containing 1,609 contigs with a total genome size of 38.0 Mb (Gao et al. 2011).

In this study, we describe another instance of diploidy within the genus *Metarhizium* and the first within *M. acridum*. We present a de-duplicated haploid genome assembly of *M. acridum* strain ARSEF 324, consisting of 35 contigs representing a haploid genome of 44.7 Mb. Ploidy is established through k-mer analysis on unassembled sequencing reads, and intragenomic collinearity analysis by all-versus-all mapping of contigs. Euploidy was established through sequencing read depth comparison across each contig, and through estimation of heterozygosity ratios of single nucleotide variants (SNV), both in 10-kb nonoverlapping sliding windows. Finally, using DAPI fluorescence staining we demonstrate that conidia are uninucleate.

Results

K-mer Analysis

The k-mer analysis on the PacBio subreads predicted that the sequenced genome was diploid with a haploid length of 45 Mb (fig. 1*A*). The profile in figure 1*A* shows two peaks in the frequency of observed unique 21-mers within the sequencing data. The smaller peak has half the read coverage of the taller peak indicating the presence of heterozygous loci. The genome is estimated to have a heterozygosity rate of 0.5%. Analyzing the sum of k-mer pair coverages to coverage ratios confirmed the inferred diploidy of the data in figure 1*B*

GBE

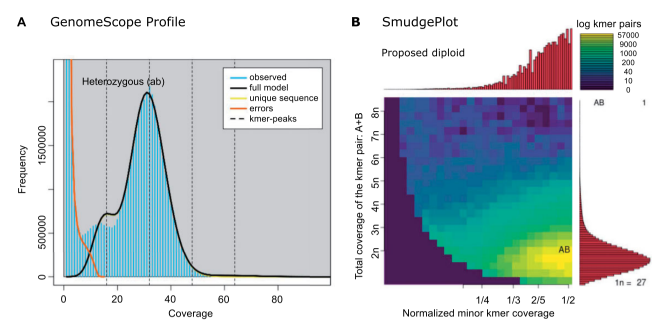


Fig. 1.—The k-mer analysis for *Metarhizium acridum* ARSEF 324 PacBio subread data revealed a diploid structure of heterozygous k-mer pairs. (*A*) Based on the number of unique 21-mers the genome is estimated to have a haploid length of 45 Mb, with 18.2% repeats. The two peaks of the k-mer frequency profile of the observed data, where the lesser have half the read coverage of the higher peak, indicate heterozygocity, consistent with diploidy. The low frequency of k-mers from heterozygous loci indicate that most 21-mers were heterozygous (0.5%). (*B*) The diploidy of the genome was confirmed by Smudgeplot analysis, comparing the sum of k-mer pair coverages (CovA + CovB) to their relative coverage (CovB/(CovA + CovB)). No higher-level ploidy was observed. Max and min estimates from the GenomeScope analysis are available in supplementary table S1, Supplementary Material online.

(for full GenomeScope results see supplementary table S1, Supplementary Material online).

Genome Assembly

A total of 478,156 filtered PacBio reads with an average length of 12.7 kb was used to assemble the genome into 35 primary contigs representing a haploid genome of 44.71 Mb. The primary assembly H1 has an L50 of 6, with a N50 of 23.37 Mb, and 4,276 complete BUSCO genes, out of the 4,494 BUSCO genes sought, or 95.1% (Scores in BUSCO format: C: 95.1% [S: 94.2%, D: 0.9%], F: 1.4%, M: 3.5%, n: 4494). Out of the 44.7 Mb in H1, 9.3 Mb (20.8%) was identified to be from repetitive elements, 74.2% of which had a strong GC-bias (<33.2%). The alternative haplotig (H2) is represented by 565 contigs with a collective size of 39.24 Mb, that is, 88% of the assumed haploid length. This assembly has an L50 of 144, with an N50 of 0.08 Mb. This set of contigs only contains loci sufficiently divergent from H1 to be forked out during the genome assembly. This implies that conserved loci between H1 and H2 will be present in the H1 primary assembly but largely absent from the H2 haplotigs. This is reflected in the associated BUSCO scores where 43.7% of the highly conserved BUSCO genes are missing from H2 (C: 54.1% [S: 51.7%, D: 2.4%], F: 2.2%, M: 43.7%, n: 4,494), likely because the genomic regions are highly conserved between the two haplotypes and therefore collapsed and only present in the H1 assembly. The outer band of the outer tracks of figure 2 depict the size of the 35 primary contigs representing the complete haploid genome. The second band depicts the mapping of the 565 H2 contigs to H1, indicating their redundancy (fig. 2). The mitochondrion genome was assembled to a length of 96,363 bp and contained 17 genes. The number and order of genes correspond with the conserved structure of mito-genomes within Hypocreales (Aguileta et al. 2014).

Ideally, the sequencing of an euploid genome should result in an equal read depth across the genome. We mapped both long and short-read data to the H1 assembly, to assess CCNV. The genome wide mean read depth was $54\times$ and $103\times$ (excluding $>500\times$ depth, at repetitive elements), for long and short-read data respectively (fig. 2). Smaller contigs ($<400\,\mathrm{kb}$) tended to have lower read depth, likely due to poor mapping in repeat dense regions. Contig 34 had a noticeable lower coverage than other contigs with a mean long-read depth of $52\times$. Contig 34 has an H2 haplotig that uniquely maps to it why it is unlikely that contig 34 constitutes a small haploid chromosome and more likely a haploid segment in the otherwise diploid chromosome. The equal depth across the genome is a strong indication of euploidy across all chromosomes represented in the assembled contigs.

The ploidy of each contig was also assessed by mean ratios of SNV read depth in 10-kb nonoverlapping windows. The expectation is that chromosomes with an even ploidy will

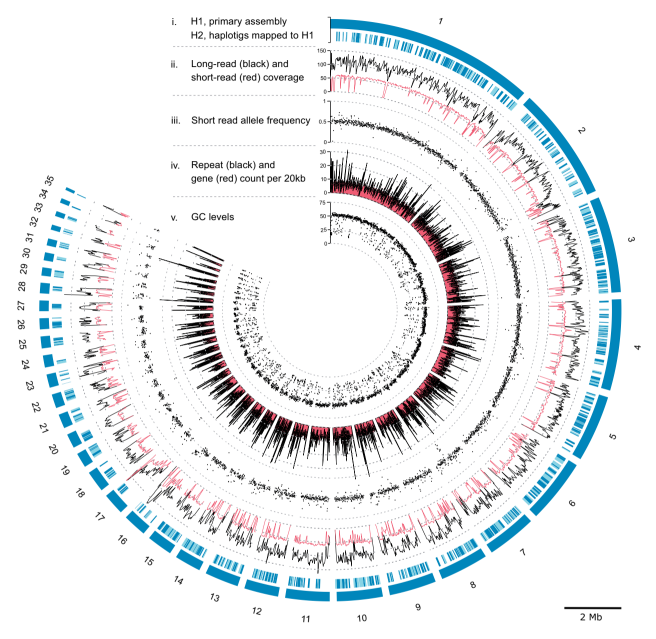


Fig. 2.—Genome-wide statistics of sequencing data, assembly and traits of the 35 primary haploid contigs of *Metarhizium acridum* strain ARSEF 324 genome. The plot consists of five major tracks; from the outside: (i) 35 contigs placed in the outer band (H1) represent the haploid genome. The remaining 565 contigs (H2) map to H1. H2 contigs are given in light and dark blue, where dark blue indicates the fraction of each contig mapped to H1. The following tracks, show (ii) the read depth of long reads (PacBio) and short reads (BGI DNBseq) mapped to H1 (mean depth within 20-kb windows), (iii) Allele frequencies calculated from short reads (BGI DNBseq) mapped to H1 (mean within 10-kb windows), (iv) Repeat and gene count within 20-kb windows shown in black and red respectively, (v) The GC levels within 10-kb windows. The contig length of the 35 primary contigs is indicated by the scale bar legend.

tend toward a 50:50 distribution across each SNV, whereas chromosomes with odd ploidy will tend toward a 33:66 or 33:33:33 ratio. All contigs have distributions of ratios with means of 0.5 (50:50) (fig. 2), confirming the diploidy of all contigs. Deviations from 0.5 correlate with read depth of mapped short-read DBNseq data on which the SNV data is based. Mapping of short-reads likewise correlates with the

repeat density (fig. 2), because high repeat densities impede the mapping.

The mean GC content across the 44.7 Mb of the H1 contigs is 45% (fig. 2, track 5). The GC-content analyzed in 100-bp windows shows a bimodal distribution with peaks at 17.6% and 51.3% (fig. 3A). In the following, we delimitate the low and high GC peaks at 33.2%. Repetitive elements explain 81.5% of the

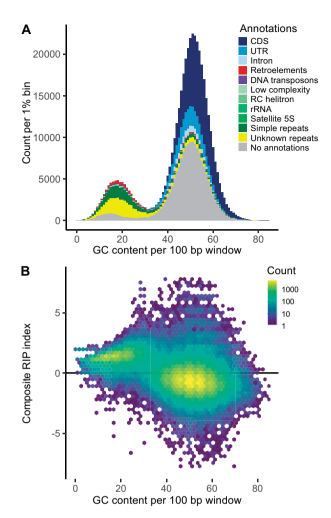


Fig. 3.—GC content calculated in 100-bp windows across the 44.7 Mb H1 haploid genome. (A) If a window overlapped with more than half of its length with an annotated feature this were signed to the window. The bimodal distribution has peak GC-levels of 17.6% and 51.3%. Each peak approximates normal distributions with standard deviations of 5.6 and 6.5, respectively. The fitted normal distributions intersect at 33.2%; this level is set to split high and low GC values. Genomic windows with high GC values are largely comprised CDS (37.3%), UTR (10.9%), introns (3.3%) and sequence that were without annotations (42.3%). Genomic windows with low GC values are dominated by repetitive elements. Of these repetitive elements 40.5% are unknown, 28.7% are simple repeats, 6.4% are retroelements, 5.9% are DNA transposons. Of the low GC windows 17% were without annotations. (B) Composite RIP index as a function of GC content. Regions with a composite RIP index >0 are considered RIP affected, which comprise 33.1% of the total genome, or 95.7% of the windows within the low GC peak. Combining different indices of RIP will provide a more conservative estimate of the proportion of the genome that is affected by RIP (17.9%, fig. 5).

low GC peak, 40.5% are unknown repetitive elements, simple repeats 28.7%, retroelements 6.4%, and DNA transposons explain 5.9%. Within low GC windows, 17% were without annotations (fig. 3A). Analyzing the dinucleotide composition across the 100-bp windows using the RIP composite index, showed

that 33.1% of the genome have a composite index value above zero and are thereby interpreted as affected by RIP. Only 19.3% of the windows with GC content above 33.2% are RIP affected, whereas 95.7% of the low GC peak are affected by RIP (fig. 3B). Across GC levels, 82% of transposable elements are affected by RIP. Similarly, 83% of unknown repeats are RIP affected.

Metarhizium acridum ARSEF 324 is homozygous for the MAT1-2 mating-type idiomorph (fig. 4). The two mating-type alleles of M. acridum ARSEF 324 and the mating-type allele of M. acridum CQMa 102 have notable differences in insertions. H1 contains a Mutator-like element (MULE) DNA transposon downstream of MAT1-2-3 not present in H2. In contrast, H2 harbors a LAGLIDADG homing endonuclease not present in H1. Metarhizium acridum CQMa 102 contains a WD-repeat domain that is not found in either of the two ARSEF 324 alleles.

Analysis of single-copy homolog gene pairs within the *M. acridum* ARSEF 324 diploid strain showed that homologous gene pairs on average were as divergent when compared with homologs of the Chinese *M. acridum* strain CQMa 102 (supplementary fig. S3, Supplementary Material online). The divergence observed between homologous genes within strain ARSEF 324 were likewise comparable to the intraspecific divergence between isolates of *M. bruneum* or *M. robertsii* (supplementary fig. S3, Supplementary Material online). Among the divergent single-copy homologs within ARSEF 324, no Gene Ontology functional annotations were notably enriched for synonymous substitutions (supplementary fig. S4B, Supplementary Material online) or correlated to the composite RIP index (supplementary fig. S4C, Supplementary Material online).

Genome Comparison

To assess if our haplotig assembly differed from other species within the genus Metarhizium, we compared the assembly of strain ARSEF 324 with genomes of eight Metarhizium species (fig. 5). For the phylogenetic analysis, two strains of the closely related Pochonia chlamydosporia was selected as outgroups. The Metarhizium genomes presented range in haploid size from 30.8 to 44.7 Mb. The 12.536 predicted genes within ARSEF 324 (H1), fall within the range predicted for other Metarhizium isolates, that is, 8,472–13,646. A notable difference can be seen in the proportion of repetitive elements between genomes. The median repeat level across the Metarhizium isolates is 6.1%, M. anisopliae strain JEF-290 has twice this, and the diploid M. acridum strain ARSEF 324 has three times this level, with 20.8% of the genome comprised repetitive elements. The proportion of the genome of M. acridum strain ARSEF 324 that is affected by RIP is likewise higher than observed for any other *Metarhizium* genome analyzed (fig. 5). A conservative estimate of the RIP affected region can be given by only including regions that are assessed as RIP affected with all three RIP calculating methods: RIP product, RIP substrate, and RIP composite as used above. With this conservative estimate, 17.9% of the M. acridum strain

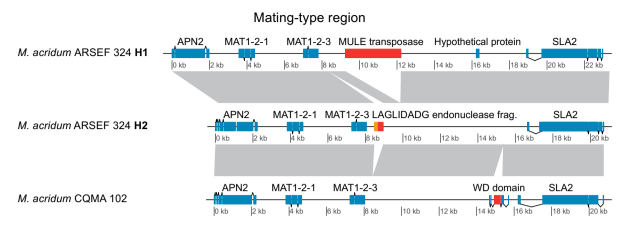


Fig. 4.—Synteny between mating-type loci of the two alleles *Metarhizium acridum* strain ARSEF 324 and *M. arcridum* strain CQMa 102. The *M. acridum* strain ARSEF 324 is homozygous for the MAT1-2 mating-type idiomorph. Down-stream of the ARSEF H1 MAT1-2-3 gene, a 3 kb DNA transposon of the "*MULE*" (*Mutator*-like elements) superfamily is found, this is not present in ARSEF H2. *Mutator* transposable elements are among the most mutagenic transposons known (Lisch 2015; Dupeyron et al. 2019). Conversely, H2 harbors a LAGLIDADG endonuclease fragment (red) together with a fragment of the mitochondrial COX3 gene (orange), not present in H1. Neither of these insertions is present in the CQMa 102 strain, which contains a WD domain missing in ARSEF 324.

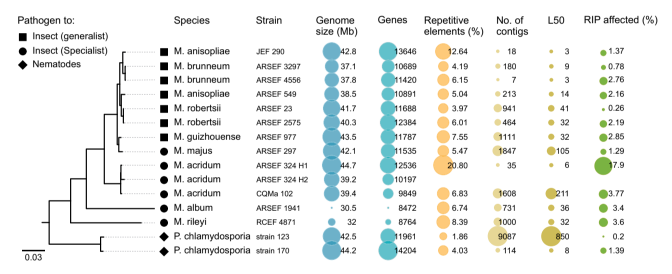


Fig. 5.—A consensus maximum-likelihood phylogeny based on 444 known single-copy orthologous genes for sequenced *Metarhizium* species and the related *Pochonia chlamydosporia*. All splits within the phylogeny had 100% bootstrap support, except the artificial split between H1 and H2. From the left: ML phylogeny, tip icons indicating the host adaptation, species names and strain id. Genome size and number of genes is retrieved from NCBI Genbank and the protein file available under the accession found in supplementary table S2, Supplementary Material online. The proportion of repetitive elements within each isolate was found as described in the method section. The figure highlights that the genome assembly of *M. acridum* ARSEF 324 generated in this study is the longest genome assembly of any *Metarhizium* isolate, and the one with the highest proportion of repetitive elements.

ARSEF 324 genome is affected by RIP (calculated in 1,000-bp window, in 500 bp steps). Evaluating the genomes ordered in descending order of RIP content, the diploid *M. acridum* strain is followed by *M. acridum* CQMa 102 and *M. rileyi* RCEF 4,871 with 3.8% and 3.6%, respectively (fig. 5). The effect of window size on total RIP proportions is shown in supplementary table S3, Supplementary Material online. DAPI staining of individual conidia supported that cells are homokaryotic diploid and could rule out that the strain was heterkaryotic by showing a single nucleus within each conidia (fig. 6).

Discussion

Metarhizium species are predominately haploid, with only two independent lineages within the M. majus complex described as stable diploids. Here we present a highly improved genome assembly of M. acridum based on long-read sequencing data assembled across repeat regions. This reveals a genome enriched in repetitive elements relative to other Metarhizium isolates or species studied. All presented analyses indicate that the sequences M. acridumstrain ARSEF324 show the genomic characteristics of a diploid. All 35 contigs of the representative assembled

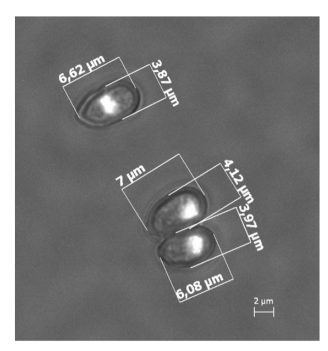


Fig. 6.—DAPI stained nuclei in *Metarhizium acridum* strain ARSEF 324 conidia. All inspected spores contained monokaryons.

haplotypewere subject to the mapping of smaller contigs, and all were heterozygous with mapped read allele frequencies around 0.5. The only exceptions are two minor contigs, contig 17 and contig 34, which could be examples of misassembly, or segmental an euploidy.

A growing body of evidence indicates that CCNV is a common adaptive trait of many fungi and has been observed at low prevalence in natural populations (Farrer et al. 2013). The effect on fitness induced by CCNVs has been shown to vary, and can be life-stage specific, and how often these variants are reproductively stable is unknown. We did not see any signs of chromosomal loss to indicate ongoing haploidization within this diploid strain of M. acridum. The absence of chromosome loss suggests that the polyploidization is recent, or that the strain constitutes a stable diploid. Kepler et al. (2016) found monophyletic clades of diploid isolates in a population genetic study of the M. majus species complex (the MGT clade, sensu Bischoff et al 2009). A similar population study could shed light on the prevalence of diploidy or CCNV within M. acridum. Alternatively, the diploid stability can be assessed by long-term evolution experiment focusing on possible reversions to the ancestral haploid state. Future research is needed to investigate gene evolution within this strain and identify concerted gene loss levels and rates of pseudo- and neofunctionalization. If the diploidization of this strain is not recent, it should be possible to trace divergence between homolog genes, where gene duplication did not result in increased fitness and purifying selection.

The highly continuous genome assembly ARSEF 324 includes the highest proportion of repetitive elements of

any Metharhizium species reported so far. Part of the reason for this is likely the fragmented character of many short read based assemblies and the inherent bias that spring from difficulties in assembling across repeats using these technologies. Our comparative analysis includes, besides the genome assembly of M. acridum ARSEF 324, three assemblies based on long-read technologies, that is, M. anisopliae JEF-290, M. brunneum ARSEF 4556, and P. chlamydosporia 170 (supplementary table S2, Supplementary Material online). Assuming that these assemblies give credible indications of their species' general repeat content, it is noticeable that the long-read assemblies of M. anisopliae and M. brunneum, both belonging to the presumable asexual PARB clade and have considerable less repetitive elements compared with ARSEF 324. This difference between the primarily asexual M. anisopliae and M. brunneum and the occasional sexual M. acridum corroborates the theory that sexual reproduction allows for the proliferation of transposable elements within populations like "a sexually transmitted nuclear parasite" (Hickey 1982). Another difference between asexual and sexual fungal species is in their defense against such selfish DNA elements. The fungal RIP pathway that mutates TE's is only active during the sexual cycle and should therefore only be observed in species that sexually recombine and be almost absent in primarily asexually reproducing species. Comparing the repeat content to the proportion of the genome affected by RIP across the Metarhizium species show that the two well-assembled genomes of M. anisopliae JEF-290 and M. brunneum ARSEF 4556 both have a higher ratio of "repeats %" to "RIP %" compared with M. acridum ARSEF 324, indicating a lack of an active RIP control of repeats. Both the high repeat content and the active RIP defence supports that M. acridum is at least occasionally sexually reproducing, even though no teleomorph is known (Mongkolsamrit et al. 2020). The impact of RIP defence on the genome of M. acridum has changed the nucleotide composition of approximately one-fifth of the genome. RIP affected regions have a reduced GC content that is less than half that of the background GC frequency (51%).

Cambareri et al. (1991) showed that both copies of duplicated gene-sized sequences within *Neurospora crassa* are affected by RIP in a single sexual cycle at frequencies of 50 – 100%. The effectiveness of the RIP machinery varies between species, but it generally affects homologous sequences longer than 400 bp within a nucleus, including selfish mobile elements, viruses and genes. The diploid nature of this *M. acridum* genome is therefore incompatible with an active RIP pathway and sexual reproduction because all duplicated gene regions would have been mutated. The activation of the RIP pathway must therefore have preceded the described genome duplication.

The presence of two identical mating-type ideomorphs within ARSEF 324 makes it unlikely that the diploidization results from a mating event as hypothesized of the diploid strains within the *M. majus* species complex. This supports a model where

either endoreplication, or allopolyploidization through parasexuality are responsible for diploidization. If the former were the case, the duplication event should be old enough for the measurable sequence divergence to have occurred between homologous genes. The phylogenetic distances between homologous genes within the diploid were comparable to the distance observed between this diploid strain and the conspecific strain QCMa 102, as well as the distances observed between haploid strains of *M. bruneum* and *M. robertsii* (supplementary fig. S3, Supplementary Material online). This indicates that the observed phylogenetic distance between ARSEF 324 homologs is consistent with allopolyploidization. The high prevalence of parasexuality observed in *M. robertsii* (Wang et al. 2011) renders alloploidization a likely mode of forming the observed diploid *M. acridum* isolate.

CCNV has thoroughly been linked with both stress and enhanced fitness (Hu et al. 2011; Farrer et al. 2013). In Metarhizium the high frequency of parasexuality observed in coinfected insects (Riba et al. 1980; Leal-Bertioli et al. 2000; Wang et al. 2011) supports that coinfected insects provides an opportunity for recombination between pathogenic isolates. If the diploidization of ARSEF 324 arose from a parasexual fusion, the question remains how stable or longlived this is. The diploid strain ARSEF 324 is likely to be phenotypically different from haploid M. acridum strains because higher ploidy can enable increased transcription of virulence factors or effectors, which could make the strain a more potent biocontrol agent. Diploidy can also work as a shield against detrimental mutation through functional redundancy (Haldane 1932, p. 110; Orr 1995) This could explain the strain-specific high tolerance to UV-B radiation of ARSEF 324 compared with other M. acridum isolates reported by Braga et al. (2001).

Although the diploidy reported here for an isolate within *M. acridum* is independent from the cases of diploidy in *M. majus* (Kepler et al. 2016), it is noteworthy that there are several instances of ploidy variation in the genus *Metarhizium*. It is tempting to speculate whether the complex species associations where many isolates are soil-dwelling plant—root endophytes and rhizosphere colonizers as well as potent insect pathogens influence the tendency for ploidy-level variation (St. Leger and Wang 2020). It certainly provides the potential for these fungi to experience several stressful microhabitats (Lovett and St. Leger 2015), which could be triggering ploidy-level variation among *Metarhizium* strains.

Materials and Methods

Strain and Culturing

The *Metarhizium acridum* strain ARSEF 324 was isolated in 1979 from a spur-throated locust (*Austracris guttulosa*) in Australia and obtained from the ARSEF collection (ARS Collection of Entomopathogenic Fungal Cultures, Ithaca,

New York). A single-spore isolate was obtained by plating serial-dilutions of conidia from the original culture. Conidia from the single-spore culture were grown in liquid Sabouraud dextrose broth with yeast extract (40 g/l dextrose, 10 g/l peptone, 10 g/l yeast extract, pH = 6.5) while being stirred at 170 rpm on a shaking table for 24 h before DNA extraction. Fluorescence staining with DAPI (blue, 4',6-diamidino-2-phenylindole) was used to visualize the number of nuclei within mature conidia, as described in Kepler et al. (2016). Imaging was done on a Zeiss Axioscope microscope (×100 objective) with an AxioCam ICm1, and spore sizes measured using ImageJ v1.53e (Schneider et al. 2012).

DNA and RNA Extraction and Sequencing

Liquid culture filtrate was ground with liquid nitrogen in a mortar and approximately 500 mg finely ground powder transferred to 50-ml Falcon tube. DNA was extracted using a CTAB method by adding 17.5 ml CTAB buffer with 0.1% 2-Mercaptoethanol and 125 µl Proteinase K and incubated for 1 h at 60 °C. After centrifugation for 20 min at 5.000 g at 4 °C, the supernatant was removed and washed with 1 v of Phenol/ chloroform/isoamyl alcohol (25:24:1) followed by two rounds of 1 v chloroform/isoamyl alcohol (24:1) and centrifuged for 10 min at 11,000 rpm at each step. To remove RNA, 150 µl RNAse was added to the aqueous phase and incubated for 120 min at 37 °C. To precipitate DNA, 0.6 v 2-propanol was added and incubated at -20 °C overnight, before centrifugation at 11,000 rpm at 4°C for 30 min. The resulting pellet was washed twice with 2 ml 70% ethanol and centrifuged for 10 min at 11,000 rpm at 4°C before being suspended in 500 µl TE-buffer. Purity was checked using NanoDrop reading and DNA quantity using a Qubit Broad-Range analysis kit.

A single SMART Cell library was sequenced on a PacBio Sequel platform using the Sequel Sequencing Kit 2.1 v2 (Sequencing Kit par number 101-309-500). Each continuous long read had one passage of sequencing; subsequently, subreads were generated by the removal of adapters and bases were called with the basecaller v5.0.0.6236. Sequencing was performed by Genewiz (Takeley, United Kingdom) with a yield from the single SMRT Cell equaling a coverage of 134×, assuming a 45 Mb genome, with a mean subread length of 11.2 kb. Short read sequence data were obtained using the PCR-free DBNseq platform at BGI-tech Copenhagen. Sequencing adapters were removed by the sequencing company, delivering a total of 15,111,206 reads of 150 bp, equaling a coverage of 50×, assuming a 45 Mb genome.

To obtain RNAseq samples to aid in the genome prediction, four replicate flasks with 20 ml media were grown for four days as described above but with 100 rpm agitation. After filtration through a Whatman filter paper (5 - 13 μ m), the fungal material was collected and flash-frozen in liquid nitrogen before being pulverized in a tissue lyzer and extracted

using a QIAGEN plant RNeasy kit following manufacturers specifications. The four samples were sequenced separately using the DBNseq platform at BGI-tech Copenhagen yielding approximately 16,000,000 paired-end 150 bp reads per sample.

K-mer Analysis and Genome Assembly

K-mers frequencies within PacBio subread data were obtained with KMC 3 (Kokot et al. 2017), and genome size and ploidy were inferred using GenomeScope v2.0 (Ranallo-Benavidez et al. 2020). Ploidy was further investigated comparing the sum of k-mer pair coverages using Smudgeplot implemented in GenomeScope. Sequence reads were both error-corrected and assembled with Canu v. 2.0, using the following parameters to co-assemble haplotypes: genomeSize = 45m correctedErrorRate = 0.03corOutCoverage = 200batOptions=-dg 3 -db 3 -dr 1 -ca 500 -cp 50. Two rounds of polishing were conducted to improve the assembly, each consisting of mapping raw-reads to the assembly using Minimap2 v.2.6 (Li 2018), and reducing remaining insertiondeletion and base substitution errors by polishing the consensus sequence using Arrow v2.3.3 (https://github.com/ PacificBiosciences/GenomicConsensus, last accessed April 30, 2021). Mirror-reads (putatively an effect of missing adapters during sequencing) were detected in the PacBio subreads set; therefore, reads longer than 40 kb were filtered out prior to assembly. Different levels of the correctedErrorRate parameter were tested (i.e., 0.020, 0.025, 0.030, 0.035, 0.040). Each of the five assemblies was assessed based on the complementarity of phased contigs. The value 0.030 were chosen for the final assembly (see supplementary fig. S1, Supplementary Material online).

Contigs were ordered according to length and grouped into primary contigs, forming a haploid representative genome assembly [Haplotig 1 (H1)] and shorter contigs (H2) that mapped to the primary contigs. Primary contigs were identified by an all-versus-all contig mapping on the repeat masked assembly with minimap2 v.2.17. The best match of each contig to a longer contig was assessed from the accumulated length of alignment between each pair of contigs. Primary contigs were manually curated; three contigs were not only subjects to mapping, but also query to other primary contigs. These contigs, and the regions they mapped to had approximately half the read depth of the mean read depth of the other primary contigs. Because the read depth of the subject regions could be raised to the mean read depth, if the query contigs were removed before mapping, these three contigs were assumed to be haplotigs and moved to the H2 group.

The mitochondrion was assembled from 2,806 reads mapping to the mitochondrion of *Metarhizium rileyi* strain RCEF 4871 (NCBI accession number: NC_047289.1; Zhang et al. 2020). The alignment was done with Minimap2 v2.17(r941)

(Li 2018). Only reads above 8 kb, which mapped with more than 70% of their length was kept. The mitochondrion was assembled using Canu v. 2.0 (Koren et al. 2017) with the settings: "genomeSize = 62k" and "corOutCoverage = 999." The mitochondrion was attempted circularized using circlator v1.5.5 (Hunt et al. 2015). To facilitate synteny analysis, the mt-genome fasta file was rearranged to start with rnl gene and end with nad6 gene.

Transcriptome Assembly

Adapter and low-quality sequences were trimmed from RNA-seq reads with Trimmomatic v.0.36 (Bolger et al. 2014) and poly-A sequences were clipped from transcripts using SeqClean (available at http://compbio.dfci.harvard.edu/tgi/software/, last accessed April 30, 2021). Reads were aligned to the complete assembled genome (H1 + H2) of *M. acridum* ARSEF 324 using HISAT2 (Kim et al. 2019), allowing for a maximum intron length of 3 kb. This was followed by clustering of aligned reads using Trinity v2.8.5 (Grabherr et al. 2011) in a genome-guided de novo transcriptomic assembly, using the jaccard clip parameter to reduce transcript fusion. The Trinity assembled transcripts were input to gene prediction training to support genome annotation.

Genome Annotation

A custom repeat library was created by adding de novo identified repeats from RepeatModeler (Hubley et al. 2016) to the repeat databases. Dfam Consensus-20170127 (Hubley et al. 2016), and RepBase-20170127 (Bao et al. 2015). RepeatModeler was run on the genome masked by these two public databases, and the iteratively growing custom library, using RepeatMasker v4.0.7 (Smit et al. 2015) with the option [-species] set to fungi. RepeatModeler relies on the three de novo repeat finding programs RECON (Bao and Eddy 2002), RepeatScout (Price et al. 2005), and LtrHarvester/Ltr_retriever (Ellinghaus et al. 2008). Repeats were identified using RepeatMasker [option: sensitive] and the generated custom repeat library. The custom library was built on repeat data from the following four Metarhizium strains: M. acridum CQMa 102, M. anisopliae JEF-290, M. brunneum ARSEF 4556, M. rileyi RCEF 4871.

Gene prediction and functional annotation of the polished assembly was conducted using the Funannotate pipeline v1.8.4 (https://github.com/nextgenusfs/funannotate, last accessed April 30, 2021). Repeats were identified with RepeatModeler and soft masked using RepeatMasker. Protein evidence from a UniProtKB/Swiss-Prot-curated database (v2021_01) (Bateman et al. 2021) was aligned to the genomes using TBlastN and Exonerate (Slater and Birney 2005). Two gene prediction tools were used: AUGUSTUS v3.3.3 (Stanke and Morgenstern 2005) and GeneMark-ES v4.33 (Besemer and Borodovsky 2005), with Fusarium

graminearum as the model for the AUGUSTUS gene predicter and BRAKER1 (Hoff et al. 2016) for the training of GeneMark-ES. tRNAs were predicted with tRNAscan-SE (Lowe and Eddy 1997). Consensus gene models were found with EvidenceModeler (Haas et al. 2008). Gene models were further updated with the "update" step in funannotate, which uses the tool PASA (Haas et al. 2008) to further extend untranslated regions and identify alternatively spliced isoforms based on alignments of RNA-Seq assembled transcripts. The completeness of the assembled genome was evaluated through comparison to the 4,494 single-copy ortholog genes of the hypocreales_odb10 data set (Creation date: 2020-08-05, https://busco-data.ezlab.org/v5/data/lineages/, last accessed April 30, 2021), using BUSCO v5 (Simão et al. 2015).

Functional annotations for the predicted proteins were obtained using BlastP to search the UniProt/SwissProt protein database (v2021_01). Protein families (Pfam) and Gene Ontology (GO) terms were assigned with InterProScan v5.48-83.0 (Jones et al. 2014). Functional predictions were also inferred by alignments to the eggNOG 5 orthology database (Huerta-Cepas et al. 2019) using emapper v2.0.1b-2g816e190 (Huerta-Cepas et al. 2017). The secretome was predicted using SignalP v5.0 (Almagro Armenteros et al. 2019) and Phobius v1.01 (Käll et al. 2007), identifying proteins carrying a signal peptide. To further characterize the secretome, putative CAZvmes were identified using HMMER v3.3 (Eddy 2011) and family-specific hidden Markov model profiles of dbCAN2 meta server (Lombard et al. 2014). Putative proteases and protease inhibitors were predicted using the MEROPS database (October 04, 2017) (Rawlings et al. 2016). Biosynthetic gene clusters were annotated using strict parameters of the antibiotics and Secondary Metabolites Analysis Shell v5 (antiSMASH) (Blin et al. 2019). All functional predictions and annotations were executed through the Funannotate pipeline.

The guanine-cytosine (GC) content was determined in nonoverlapping windows across the 35 H1 contigs, using seqkit v0.13.2 (seqkit sliding -s 100 -W 100 | seqkit fx2tab – name –gc). Each window was annotated with the gene or repeat annotation (or no annotation), which it primarily overlapped. This was achieved with bedtools v.2.28.0 option [intersect -wa -wb -f 0.51] (Quinlan 2014).

Variant Calling and Chromosome CCNV

The 150 bp PE DNBseq reads were mapped with the Burrow—Wheeler Aligner, BWA-MEM v.0.7.16a (Li 2013) to the primary haplotig (H1) of the assembled nuclear genome, and PCR duplicates were removed from the bam file using samtools v1.10 (markdup –r) (Li et al. 2009). Coverage and mean depth to reference was assessed using samtools (coverage). Allele frequency distribution across the genome was calculated based on SNV called using Bcftools v1.10.2 (Li 2011a). Variants were called using a combination of

BCFtools "mpileup" and "call" using a mapping quality filter of 30, a base quality filter of 20, and a minimum depth of 10, together with default parameters including BAQ (Li 2011b). Mean SNV allele ratios were summarized in nonoverlapping 10-kb windows using a custom R script (supplementary script 1, Supplementary Material online). Genome annotations and repeat annotations were combined for the mating-type loci including the flanking *APN2* and *SLA2* genes, for the two genome assemblies of *M. acridum* ARSEF 324 and *M. acridum* CQMa 102 (GenBank accession no. GCA_000187405.1). Synteny between loci was visualized using the R package genoPlotR v0.8.10 (Guy et al. 2010).

Genome Comparison

Single-copy homologous (SCH) genes were identified between H1 and H2, and 11 Metarhizium isolates representing eight species, along with two isolates of Pochonia chlamydosporia and two isolates used as outgroup, that is, Epichloe festucae strain FI1, and Villosiclava virens strain UV-8b. This was done using OrthoFinder v2.2.7 (Emms and Kelly 2019). The proteomes of the 16 isolates were obtained from NCBI GenBank; accession numbers are available in supplementary table S2, Supplementary Material online. Protein sequence alignment was generated with MAFFT v 7.453 as implemented in OrthoFinder. Substitution models for each gene were predicted using ModelFinder (Kalyaanamoorthy et al. 2017) as implemented in IQtree v2.1.2. A subset of 444 known genes with less than 10% gaps within the alignment was concatenated and a maximum likelihood phylogeny with gene-specific substitution models using IQtree (Nguyen et al. 2015; Chernomor et al. 2016). Per gene maximum likelihood distances were likewise calculated using IQtree and utilized in supplementary figure S3, Supplementary Material online, to summarize the distance between homologous genes.

SCH identified between H1 and H2 had their synonymous point substitution rates estimated with the pairwise alignment tool in subopt-kaks (Stajich 2019) (available at: https://github. com/hyphaltip/subopt-kaks, last accessed April 30, 2021), which implements yn00 pairwise dN/dS estimations (Yang and Nielsen 2000). Estimations are displayed in supplementary figure S4, Supplementary Material online, per gene, along with respective gene ontology annotations. RIP mutations indices were calculated using "The RIPper" (http://theripper.hawk.rocks). Three different indices were calculated, based on dinucleotide frequencies: RIP substrate [(CpA+TpG)/(ApC+GpT)]: $0.75 \ge x$, RIP product [TpA/ApT]: $x \ge 1.1$ and RIP composite [(TpA/ApT) – ((CpA + TpG)/(ApC + GpT))]: $x \ge 0$, where x are values indicating RIP on a given sequence. Conservative estimates of the RIP affected proportion of the genome are given as regions where all the above indices indicate RIP (fig. 5).

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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Author Contributions

H.H.D.F.L. initiated the project. M.N. cultured fungi, performed DNA extraction, and initial data QC. K.N.N. and T.K. performed genome assembly, J.F.M.S. and J.S. carried out genome annotation, K.N.N. and J.F.M.S. carried out all other data analyses with input from H.H.D.F.L. and J.S., K.N.N. drafted the manuscript and all authors were involved in the finalization of the manuscript.

Data Availability

Sequencing data and genome assemblies of Metarhizium acridum strain ARSEF 324 are available at DDBJ/ENA/GenBank under the BioProject accession numbers PRJNA714114 and PRJNA714115, and genome accession JAGROK00000000 and JAGROL00000000, for the primary genome assembly H1 and the secondary haplotig assembly H2, respectively. Gene expression data used to support genome annotation are available in the Short Read Archive under BioProject PRJNA708346. The DNA for the genome assembly is associated with BioSample accession number: SAMN18235592. Genbank accession numbers of all genomes used in this study are given in supplementary table S2, Supplementary Material online.

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