Repeat-induced point mutation and the population structure of transposable elements in *Microbotryum violaceum.*

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ABSTRACT

Repeat-induced point mutations (RIP) is a genome defense in fungi that hyper-mutates repetitive DNA and is suggested to limit the accumulation of transposable elements. The genome of Microbotryum violaceum has a high density of transposable elements compared to other fungi, but there is also evidence of RIP activity. This is the first report of RIP in a basidiomycete, and was obtained by sequencing multiple copies of the integrase gene of a copia-type transposable element and the helicase gene of a Helitron-type element. In M. violaceum, the targets for RIP mutations are the cytosine residues of TCG trinucleotide combinations. Although RIP is a linkage-dependent process that tends to increase the variation among repetitive sequences, a chromosome-specific substructuring was observed in the transposable element population. The observed chromosome-specific patterns are not consistent with RIP, but rather suggest an effect of gene conversion, which is also a linkage-dependent process but results in a homogenization of repeated sequences. Particular sequences were found more widely distributed within the genome than expected by chance and may reflect the recently active variants. Therefore, sequence variation of transposable elements in M. violaceum appears to be driven by selection for transposition ability in combination with the context-specific forces of the RIP and gene conversion.
INTRODUCTION

Transposable elements make up a diverse group of molecular parasites found in nearly all organisms. Their activity is harmful to the host organism because insertion of new copies often disrupts gene expression (Wright et al. 2003) and because the similarity between copies at different loci promotes ectopic recombination and destabilizes chromosomes (Charlesworth et al. 1992; Petrov et al. 2003). Therefore, it is not surprising that parallels are drawn between transpositional activity and the virulence of conventional parasites (Jordan & McDonald, 1999). Moreover, the term “arms race” has been used when there is evidence that the host genome has evolved mechanisms to defend against transposable elements (Jordan et al. 1999; Lovsin et al. 2001).

A genome defense called *repeat-induced-point-mutation* (or “RIP”) is known to occur in ascomycete fungi (reviewed by Selker 2002). RIP is a homology-dependent gene silencing mechanism that directly mutates multi-copy DNA sequences. RIP has been credited with preventing transposable element accumulation in fungi, which generally have relatively small genomes that are said to be “streamlined” because they contain little repetitive DNA (Kidwell 2002; Selker 2002; Wöstemeyer and Kreibich 2002; Brookfield 2003; Galagan and Selker 2004).

The mechanistic details and specific protein-DNA interactions involved in target recognition and site specific mutation are not completely resolved, but the general properties of the RIP process have been well characterized in several ascomycete species (for reviews see Selker 2002, Galagan and Selker 2004). RIP induces C-to-T mutations in duplicated DNA sequences. Cytosine residues of particular nucleotide combinations are targeted; for example, in *Neurospora crassa* CA dinucleotides are the targets of RIP activity. RIP is known to occur
within haploid nuclei following mating but prior to meiosis (i.e. in the dikaryotic stage that precedes karyogamy). Cytosine methylation is frequently associated with RIP-mutated sequences, however it remains undetermined whether this is a required step in a deamination process to yield C-to-T mutations. RIP acts in a pair-wise manner on duplicated DNA sequences, such that they are not only altered but also become dissimilar because not all the same cytosine residues are changed in both copies. Finally, RIP occurs at a much higher rate between a pair of duplicated sequences when they are located on the same chromosome, and more so when they are in close linkage as compared to unlinked duplications. In addition to inactivating transposable elements by point mutations, RIP is thought to discourage ectopic recombination by decreasing the sequence similarity between copies at different loci (Cambareri et al. 1991).

RIP has been studied mostly from a mechanistic perspective and as a gene-silencing tool. However, there is little understanding of how RIP affects the intra-genomic “population” of transposable elements or their evolutionary history. The homology-dependent and linkage-dependent nature of RIP may cause the genetic changes in transposable elements to be largely influenced by the particular genomic or chromosomal contexts in which they reside. This presents a strong contrast to our paradigm of sequence evolution based on random mutation, and may seriously complicate the interpretation of transposable element evolution based on their sequence variation.

In this study, we determined the sequence variation for transposable elements in the fungus Microbotryum violaceum, which is unusual among fungi because this species contains a high density and diversity of transposable elements (Hood et al. 2004; Hood 2005). We provide evidence of RIP activity in M. violaceum by identifying the target site for C-to-T mutations, the
first such reported for a basidiomycete. We also compare the sequence variation of element copies within and between the different chromosomes, and provide evidence that gene-conversion acts in addition to RIP to determine the population structure of transposable elements in \textit{M. violaceum}.

\textbf{MATERIALS AND METHODS}

\textit{Microbotryum violaceum} (Basidiomycota; formerly \textit{Ustilago violacea}) is a pathogenic fungus of plants in the Caryophyllaceae. It is commonly studied as a model for disease ecology and fungal genetics (Antonovics et al. 2002, Garber and Ruddat 2002, and references therein). Collections of the fungus from different host plants probably represent cryptic species, but the taxonomy of these lineages is not resolved. The study reported here is based on a collection from the host \textit{Silene latifolia} from Italy (Lamole nr. Greve in Chianti; collection number IT00-15.1). This collection has been used in previous studies (Hood et al. 2004b; Hood 2005).

In order to isolate chromosome-specific DNA, karyotypes of the haploid meiotic products were separated by pulsed field gel electrophoresis following the isolation of linear tetrads by micromanipulation (Fig. 1A), as described in Hood (2002) and Hood et al. (2004). Electrophoresis conditions using a CHEF-DRII pulsed field system (BioRad) were 14 C, 96 hr, 2.8 v/cm, and 0.8% agarose, 200 sec initial and 1100 sec final switch times. DNA was extracted from each of the autosome bands and the A2 sex chromosome band using a culture of the A2 mating type. DNA from the A1 sex chromosome band was obtained from a culture from the same meiotic tetrad as the A2 culture. To extract DNA, an agarose plug was removed from the gel at each chromosome band using a glass Pasteur pipette, and DNA was isolated from the plugs by the phenol-based methods of Favre (1992). Most of the sampled locations in the gel corresponded to distinct single chromosome bands, but one sample (plug 7 of Fig. 1a) was in a
region where bands are not as well resolved; this extract may therefore have included DNA from multiple chromosomes.

PCR primers were designed for a retrotransposon fragment found previously by sequencing random AFLP products from *M. violaceum* (Hood 2002; Hood and Antonovics 2004): internal primers for the retrotransposon sequence were ECA/MGA700BR.2 Forward 5’ TGGAACCTGTACGTTGATGG and Reverse 5’ ATTTTCTGACCCGTTTGACG. The sequence is in the integrase region of a Ty1/copia type element, the most common type of transposable element in *M. violaceum*. The resulting 367 bp sequence begins approximately with the first histidine of the HHCC motif that characterizes the N-terminus of the Psuedoviridae integrase gene (as defined in Peterson-Burch and Voytas, 2002).

A separate PCR reaction was conducted with each chromosome-specific sample of DNA that was isolated from the karyotype gel. The resulting PCR products were then cloned using the TA Cloning Kit from Invitrogen, and approximately thirty clones per chromosome were sequenced using Dye Termination techniques with an ABI 377 automated DNA sequencer. Sequence alignment was carried out manually in Sequencher (Gene Codes Corporation, Inc.) and is available under accession numbers AY729078-AY729484 in the PopSet section of GenBank (www.ncbi.nlm.nih.gov). All deviations from the consensus sequence were confirmed to represent unambiguous base pair signals in the electropherograms.

As a test to determine whether evidence of RIP activity is also present in the sequence variation of the types of transposable elements in *M. violaceum*, PCR primers were designed for a Helitron-type sequence (accession BZ782234) identified in a previous study (Hood 2005). *Helitrons* are recently discovered DNA-based transposable elements that replicated by a rolling circle mechanism and are found in a broad range of organisms (Kapitonov and Jurka 2001;
Poulter et al. 2003. Internal primers for a 301 bp region of BZ782234 were 44-Forward 5’ CACGGTGAGTAGCCATTCC and 374-Reverse 5’ CCGTTTACTGCCTGATCTCC. PCR was conducted using whole genomic DNA, 12 products of which were then cloned and sequenced as described above, and sequences are available under accession numbers AY939912-AY939923 in GenBank.

As a control to confirm that DNA isolated from karyotype gels was chromosome-specific, the methods described above were replicated with a separate culture of the same fungal strain. DNA was isolated as before from chromosome bands corresponding to plugs 2 and 3 in Figure 1A, and were similarly used for PCR amplification and sequencing of thirty clones per chromosome. In this case sequencing was carried out by Genoscope-CNS (France). Also, considering that one PCR reaction may produce amplification products from the same locus, the results were analyzed using both the entire dataset as well as only the non-redundant sequences from within each chromosome (indicative of separate loci). In determining sequence redundancies, base pair ambiguities or gaps in the alignments were conservatively considered to be equivalent to base pairs in otherwise identical sequences.

RESULTS

A total of 359 cloned sequences were obtained for the 367 bp integrase region of the Ty1/copia-type retrotransposon. There were approximately 30 sequences from each of twelve isolated chromosome bands of the electrophoretic karyotype (Fig. 1): 32 sequences for chromosome band C1; 29 sequences for C5, C6, and C7; and 30 sequences for the other bands. Variations from the consensus sequence are shown in the alignment of Figure 1B. The consensus sequence was strongly supported. Specifically, there was complete identity across all
359 sequences at 62% of the nucleotide positions (sequence length = 367 bp), and there were very few (<1%) of the nucleotide positions with less than 80% identity across all the sequences.

**Pattern of Repeat-Induced Point Mutations**

The majority of deviations from the consensus copia-type retrotransposon sequence were C-to-T or G-to-A changes which is consistent with RIP activity (the latter are assumed to represent C-to-T mutations on the non-coding strand). In contrast to analyses of RIP in other fungi (Selker 2002; Clutterbuck 2004), the “target site” for hyper-mutation of cytosine residues was strictly the trinucleotide TCG rather than a dinucleotide. A partial match to this target site that was different from TCG immediately 3’ or 5’ to the cytosine residue did not show elevated C-to-T changes above the baseline for other cytosine residues (Fig. 2). There were 14 RIP target sites in the coding strand and 12 in the non-coding strand of the consensus sequence. The RIP target site was not present in the PCR primers used in this study.

For the region of the *Helitron*-type element, the 12 cloned sequences showed a bias in mutations to the same target site trinucleotide as shown in Fig 2. Specifically, 37% of TCG sites across the consensus sequence (including CGA sites for the complimentary strand) contained at least one sequence with transition mutations (n = 19), whereas the value for VCG was 11% (n = 37), for TCH was 3% (n = 34), and for VCH was 6% (n = 77) (using standard IUB codes for incompletely specified nucleotides).

RIP most often affects just one strand of the DNA duplex in a single event, and thus yields C-to-T mutation in only the coding strand or only the non-coding strand, the latter of which appear as G-to-A mutations in the coding strand (Cambareri et al. 1991; but see Watters et al. 1999). However, nearly half (47%) of cloned copia-type retrotransposon sequences contained
mutations in RIP target sites on both the coding and non-coding strands. This pattern results from multiple rounds of RIP that alternately affect the coding and non-coding strands. Nevertheless, sequences with RIP mutation only in the coding or only in the non-coding strands were more common than expected when they had less than ca. 20% of RIP target sites mutated. Specifically, 46 of 79 sequences with two RIP target sites mutated had the changes only in the coding or only in the non-coding strands (binomial distribution probabilities, \( P = 0.110 \)), as did 19 of 30 sequences with three RIP mutations (\( P=0.035 \)), and 23 of 44 sequences with four RIP mutations (\( P<0.001 \)). Sequences with higher numbers of mutated RIP sites did not differ from chance expectation as to whether all the changes were only on one strand. These results may indicate that a small percent (i.e. 10 to 20%) of TCG sites are mutated in a single round of RIP, but that many element copies have undergone multiple rounds of RIP activity.

RIP target sites (TCG) in the copia-type retrotransposon were distributed such that the induced changes would cause synonymous substitutions more often than expected by chance alone. A comparison of the consensus sequence with a copy mutated at all RIP sites (i.e. all instances of TCG substituted with TTG) resulted in a highly significant excess of synonymous over non-synonymous substitutions (Mega 2.1 Software, \( K_s > K_a \) P-value < 0.001). The tendency toward inducing synonymous substitutions is also true for RIP mutations in the model system of \( N. \ crassa \) (at CA target sites in that species), but only for C-to-T mutations in the coding strand (Selker 2002). However, in the transposable element of \( M. \ violaceum \), the distribution of RIP target sites in both the coding strand and the non-coding strand tended to produce synonymous substitutions (Fig. 3). This result was in contrast to a sample of house-keeping genes from \( M. \ violaceum \) that are not expected to be as greatly exposed to RIP; simulated RIP mutations induced more synonymous substitutions in the coding strand but not in the non-coding strand.
The more synonymous changes at TCG sites only in the coding strand of house-keeping genes was not due to a codon bias particular to *M. violaceum*, but instead was general to other fungal species. Moreover, a comparison of the integrase gene of Ty1 from yeast, where RIP has not been found (Selker 2002), also fit the pattern of house-keeping genes and not the pattern of the *M. violaceum* transposable element (Fig. 3).

**Chromosome-specific sequence variation.**

The copia-type retrotransposon copies showed chromosome-specific patterns in their deviations from the consensus sequence. Such patterns were apparent for mutations at both RIP-target sites as well as at other sites, and included transition and transversion mutations (Fig. 1B). Consequently, by including all sequences the chromosome-of-origin explained a significant amount of the sequence variation (AMOVA, Table 1; Arlequin 2.0 software; Schneider et al. 2000). Using the conservative approach of removing any identical (i.e. redundant) sequences from within the 30 sequences per chromosome, a smaller but still significant substructuring could be attributed to sequences being more similar within a chromosome than expected by chance alone (Table 2). The mean number of unique sequences per chromosome was 18 (S.E. = 1.11), with no chromosome differing significantly from the mean. Chromosome-specific patterns were confirmed by replicating the sampling methods with two chromosomes, which most closely resembled the same chromosomes from the first run of the experiment using a distance based on Fst statistics of sequence variation (Fig. 4). The depth of the branches containing the replicated chromosomes reflects a measure of error present in the sampling method, and furthermore suggests a relationship between the transposable element variation on other chromosomes, specifically chromosomes 9 and 10.
A few particular copia-type retrotransposon sequences however tended to be isolated from multiple chromosomes (Fig. 5). Among 139 distinct sequences found in the Microbotryum genome, 110 were isolated from only one chromosome, 12 from two chromosomes, and 3 from three chromosomes. However, assuming a Poisson distribution, sequences found on four or more chromosomes were more common than expected by chance alone at a significance level of \( P < 0.01 \). Specifically, 4 sequences were isolated from four chromosomes, 5 from five chromosomes, 3 from six chromosomes, and 2 from seven chromosomes. The consensus sequence was found on four chromosomes. Furthermore, those sequences that were isolated from larger numbers of chromosomes were also more likely to deviate from the consensus only by mutations at RIP target sites or only by silent substitutions (Fig. 5) (PROC CORR procedures in SAS, \( P = 0.01, N = 7 \)). One sequence containing a single base pair deletion was isolated from five chromosomes (Fig. 5).

**DISCUSSION**

**Repeat-Induced Point Mutations in *M. violaceum***.

This study has provided the first evidence of a RIP genome defense against transposable elements in a basidiomycete. The process has been sought in a small number of basidiomycetes other than *M. violaceum* (Selker 2002), but not necessarily by similar methods and its distribution throughout the phylum remains to be determined. There are clear similarities between our data with *M. violaceum* and the RIP process in ascomycetes, specifically that there is hyper-mutation of cytosine residues at particular nucleotide combinations (Hamann et al. 2000; Anderson et al. 2001; Daboussi et al. 2002, Clutterbuck 2004). However, in contrast to most ascomycetes where RIP is known to occur, the RIP target site in *M. violaceum* is a
trinucleotide (TCG) rather than a dinucleotide. Moreover, the specificity of RIP activity appears higher in *M. violaceum* than in the other fungi. In particular, the hyper-mutation was strictly limited to the TCG combination, whereas other systems show only a general preference toward cytosine residues of one or multiple dinucleotide combinations (Clutterbuck 2004). It remains to be determined whether the RIP mechanism in *M. violaceum* operates by the same molecular pathways as in ascomycetes. However, an important distinction must be drawn with regards to the dikaryotic stage during which RIP is active. The dikaryon is very transient in ascomycetes and occurs in a brief period between mating and karyogamy, but it is the dominant vegetative condition of basidiomycetes. This developmental difference may provide much greater opportunity for RIP activity in basidiomycetes. Transformation studies, such as those conducted in *N. crassa* and other ascomycetes (e.g. Bhat et al. 2004), can establish when precisely RIP mutations occur in the *M. violaceum* genome.

Because RIP occurs in a broader range of taxa than previously recognized, it is of greater interest to resolve its evolutionary history and adaptive significance in combating molecular parasites. For example, despite the apparent activity of RIP, the high density of transposable element in *M. violaceum* is inconsistent with previously studied ascomycetes (Hood et al. 2004, Hood 2005). Moreover, the mutational studies of Garber and Ruudat (1994, 1998, 2000, 2002) indicate that transposable elements continue to be active in *M. violaceum*. Why then does it appear that RIP is less effective at defending the *M. violaceum* genome than suggested for ascomycetes (Selker 2002)?

The first of four possible explanations outlined below is that active transposable elements have been repeatedly introduced into the *M. violaceum* genome by horizontal transfer. For example, the presence of an active *Tad* element in a RIP-competent strain of *N. crassa* was
explained by a recent genome invasion (Anderson et al. 2001). Also consistent with horizontal transfer is the observation that *M. violaceum* is particularly rich in the diversity of transposable elements for a species that exhibits a highly selfing mating system (i.e. automixis or intra-tetrad mating; Hood and Antonovics 2004). There are non-LTR, gypsy, and copia-type retrotransposons, as well as *Helitron*-type DNA-based elements (Hood 2005). Moreover, copia elements, as investigated in this study, are the most rare type of transposable elements in fungi, especially among basidiomycetes (Kempken and Kück 1998; Goodwin and Poulter 2001; Wöstemeyer and Kreibich 2002; Díez et al. 2003). However, copia-type elements are the most common type in *M. violaceum*. Whether these elements have originated from sources external to the fungus requires additional studies.

The frequent karyotypic rearrangements in *M. violaceum* may provide a second explanation for the apparent inability of RIP to effectively defend against transposable elements (Perlin et al. 1997; Hood et al. 2003). In *N. crassa*, translocations that result in the duplication of large chromosomal regions can act as sinks for the RIP machinery and allow shorter “gene-sized” duplications to go unaffected. As suggested by Bhat and Kasbekar (2001), transposable elements may indirectly contribute to their further proliferation by promoting ectopic recombinations that overwhelm the host’s genome defense.

It is also possible that RIP is no longer an active defense in *M. violaceum*. We may have observed the signatures of ancient RIP mutations in the background of actively proliferating lineages. In fact, RIP-like mutations are reported for some ascomycete fungi that are supposedly asexual in nature, and therefore do not experience the dikaryotic stage that is required for RIP activity. It has been suggested that evidence of RIP in the asexual ascomycetes may indicate
mutations that occurred either in the dikaryon of a sexual ancestor or during cryptic sex in extant populations (Neveuglise et al. 1996; Hua-Van et al. 1998; Ikeda et al. 2002).

A final possibility to explain the proliferation of transposable elements in *M. violaceum* is their adaptation to tolerate the RIP genome defense. Ikeda et al. (2002) suggested that the variation for preferred RIP target sites in ascomycetes contains a phylogenetic signal corresponding to the relationships between the species. Therefore, it may be possible to evaluate the changes in RIP specificity as a co-evolutionary arms-race between the genome defense and transposable elements. To further this hypothesis, this study showed that substitutions at the TCG target sites tended to cause synonymous changes not only on the coding strand but also on the non-coding strand of the copia-type retrotransposon in *M. violaceum*. In contrast, simulating RIP on the non-coding strands of genes not expect to be under selection by RIP tended to produce non-synonymous changes, including house-keeping genes and even the same integrase region of the Ty1 transposable element from yeast. A broader survey of genes is needed to confirm these results. However, if the pattern observed in *M. violaceum* is the result of selection against the RIP-sensitive variants, there may be constant pressure to adapt the preferred RIP target site to most effectively inactivate the elements. This co-evolutionary dynamic remains to be fully explored from theoretical and empirical perspectives, but may represent an important force in the interactions between molecular parasites and the host genome.

**Chromosome-specific sequence variation.**

Because chromosomes of fungi can be separated electrophoretically, methods used in this study provide a powerful approach to the intra-genomic population structure of molecular parasites. While the transposable element sequences in *M. violaceum* appear hyper-mutated by
RIP, they showed a statistically significant similarity among copies from the same chromosome. This is contrary to expectations because RIP acts most often between linked copies to diversify them due to the independent nature of mutations on the two copies involved. Furthermore, not all of the chromosome-specific variants were at RIP target sites, and some were transversion-type mutations that are not indicative of RIP. Therefore, a process other than RIP is likely responsible for the chromosomal substructuring of the variation in *M. violaceum*.

Gene conversion (Elder & Turner 1995) is the probable cause for the chromosome-specific patterns because it acts to homogenize repetitive sequences and does so in a linkage-dependent manner. However, Cambareri et al. (1991) suggested that the hyper-mutations caused by RIP may diversify repetitive sequences to the point that gene conversion is unable to act upon them. Moreover, theoretical studies suggested that gene conversion is insufficient to counter even normal mutation rates in transposable elements (Slatkin 1985). Recent evidence indicates that this is not always the case. The influence of gene conversion is seen in the MITE elements of *Arabidopsis thaliana* (Le et al 2000), S2 elements of *Drosophila melanogaster* (Maside et al. 2003), Ty1 elements of *Saccharomyces cerevisiae*, and human Alu elements (Roy et al. 2000). Our sequence data from *M. violaceum* are consistent with gene conversion between the elements on the same chromosome. Specifically, some elements contain two different chromosome-specific mutation patterns at opposite ends of the sequence (data not shown). It remains possible that such mosaic sequences are the result of reciprocal ectopic recombinations rather than gene conversion, although this is less likely due to fitness costs normally associated with such chromosomal rearrangements. Longer DNA sequences than used here may identify exchanged sequence “tracts” that are indicative of gene-conversion events (e.g. Sawyer 1989, Betran et al. 1997).
It is conceivable that processes other than gene conversion give rise to chromosome-specific patterns, but this is unlikely. Thus, it is possible that descendent copies might integrate differentially into their chromosomes-of-origin in order to create the similarity within chromosomes. However, transcripts of retrotransposons leave the nucleus for reverse transcription prior to integration (Havecker et al. 2004). Although some targeting of elements is know for particular chromosomal regions, such as in telomeric repeats (Anzai et al. 2001; Xie 2001), the local specificity needed to explain our results has never been observed. Furthermore, the control repeat sampling of chromosome bands confirmed that the patterns were chromosome-specific, as did the statistics using non-redundant sequences within a chromosome.

The sequence alterations caused by RIP and gene conversion can obscure the evolutionary history of transposable elements. First, conventional approaches to inferring phylogenies from DNA sequence data are precluded because genetic changes are in part induced by the local genomic environment. Second, because RIP tends to cause mutations at synonymous sites, it may create a false signature of purifying selection in the ratio of non-synonymous to synonymous substitutions (i.e. dN/dS statistics). Previous studies have taken steps to avoid the complications of RIP by excluding the nucleotide combinations that are targeted for mutation (e.g. Hua-Van et al. 2001). However, it appears from the current study that gene conversion can also have a major influence upon the transposable element population even in the presence of RIP. Further studies are needed to resolve the interplay between these two homology-dependent and linkage-dependent forces, particularly as they have opposite effects upon sequence variation.

Because evolution in the genomic context of RIP and gene conversion is likely to be complicated, the dendrogram of transposable elements from M. violaceum (Fig. 5) must be
interpreted cautiously. Here, we suggest only that sequences found on more chromosomes than expected by chance probably represent the recently proliferating variants. The similarity in amino acids between the consensus and the most widely-distributed sequences further suggests that these are active elements, particularly for the two sequences found on seven chromosomes. Finding a sequence with a base pair deletion on five chromosomes may contribute to the rising evidence of defective retrotransposons that hyper-parasitize variants for which the genes are still intact (e.g. Sanz-Alferez et al. 2003; Havecker et al. 2004).

Understanding transposable elements evolution in fungi requires not only further comparative studies on a wider range of taxa, but also a theory that integrates the effects of neutral substitution and selection with the context-specific effects of RIP and gene conversion.

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Table 1. AMOVA using all cloned sequences, with chromosome-of-origin as population substructure.

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<th>Sum of squares</th>
<th>Variance</th>
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<td>Among Chromosomes</td>
<td>288.49</td>
<td>0.76</td>
<td>16.82*</td>
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<tr>
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<td>1289.47</td>
<td>3.74</td>
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1000 permutations. *Probability of a larger value by chance < 0.001

Table 2. AMOVA using only non-redundant cloned sequences from within each chromosome, with chromosome-of-origin as population substructure.

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<td>Within Chromosomes</td>
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1000 permutations. *Probability of a larger value by chance < 0.001
FIGURE LEGENDS

Figure 1. Genome sampling of copia-type retrotransposon sequences from *M. violaceum*. **A.** Electrophoretic karyotypes of *M. violaceum* (collection number IT00-15.1). Lanes represent haploid cultures of opposite mating types (A1 and A2) isolated from the same meiotic tetrad. In these cultures, the autosomes are homozygous for size, but the sex chromosomes are dimorphic. Chromosome bands were sampled and used to isolate transposable element sequences as indicated by arrows. **B.** Alignment of coding strand sequences of integrase gene fragment of copia-type retrotransposon; ca. 30 clones per chromosome. Deviations from consensus shown in color: red = C to T mutation; orange = G to A mutations; green = A to G or T to C mutations (other transitions); blue = transversion mutations; black = deletion mutations.

Figure 2. Hyper-mutations at RIP target site in the integrase gene fragment of copia-type retrotransposon of *M. violaceum*. Mutation rates shown for cytosine residues in various 3' to 5' nucleotide combinations. Coding and non-coding strands combined. Standard IUB codes used for incompletely specified nucleotides; V = not T; H = not G. Numbers of nucleotide combinations are determined using the consensus sequence.

Figure 3. Effects of simulating RIP-induced mutations on amino acid sequences. All TCG sites were substituted with TTC in coding and non-coding strands. The difference between the substitutions per synonymous site (Ks) and per nonsynonymous site (Ka) is shown (analyses conducted in Mega 2.1 software). The analyzed integrase sequences were the consensus sequence copia-type retrotransposon fragment from *M. violaceum* (Mv) and an aligned sequence.
of Ty1 from *Saccharomyces cerevisiae* (Sc, NCBI accession number M18706). House keeping genes analyzed over aligned coding regions: carbamyl phosphate synthase from *M. violaceum* (Mv, BZ782437), *Trichosporon cutaneum* (Tc, L08965), *Neurospora crassa* (Nc, XM_331875), and dynein heavy chain from *M. violaceum* (Mv, BZ782578), *Ustilago maydis* (Um, AF403740), and *N. crassa* (Nc, XM_327261). Bars = standard error.

Figure 4. Dendrogram based upon Fst statistics of *M. violaceum* copia-type retrotransposon sequences grouped by chromosomes-of-origin (Fst distance analysis in DNAsp 4.0 software; tree constructed in Mega 2.1 software). Sex chromosome (A1 and A2) and autosome karyotype samples (C1 through C10) as indicated in Figure 1A. Rep2 and Rep3 correspond to methodological replication of samples C2 and C3.

Figure 5. Dendrogram of copia-type retrotransposon sequences from *M. violaceum* (neighbor-joining tree constructed in Mega 2.1 software). Colors indicate chromosome-of-origin as shown in Figure 1. Only non-redundant sequences within a chromosome sample are included. Numbers reflect from how many chromosomes a sequence was isolated (four or more are shown); asterisk indicates that this sequence is identical to the consensus (black). Sequences are marked that differ from consensus only by silent mutations (black dot) or only by RIP mutations (blue dots). Sequences marked with red dots contain a single base pair deletion.
Figure 1
Figure 2

Proportion of Nucleotide Combinations Exceeding Mutation Frequency

Mutations Frequency Among Sequences

- TCG, n = 26
- VCG, n = 45
- TCH, n = 29
- VCH, n = 87
Figure 3

Figure showing bar graphs comparing synonymous and non-synonymous (Ks-Kn) values for different regions of the genome: Integrase, Carbamyl Phosphate Syn., and Dynnein Heavy Chain. The bars are differentiated by Coding Strand and Non-Coding regions.