Differential Gene Expression between Fungal Mating Types Is Associated with Sequence Degeneration

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Data deposition: We used published gene expression data to investigate the association of sequence degeneration and differential gene expression in Microbotryum lychnidis-dioicae (Fontanillas et al. 2015; Perlin et al. 2015; accession number: PRJNA246470). We used published genome assembly, gene predictions, and assignments to genomic compartments (Branco et al. 2017, 2018). We also used published transposable elements identification in M. lychnidis-dioicae (Hartmann et al. 2018). All relevant scripts and data files to perform these analyses have been deposited in Zenodo at DOI: 10.5281/zenodo.3672080, and Github at https://github.com/Wen-Juan/Differential_expression_associateswith_degeneration_Microbotryum_fungus.

Abstract

Degenerative mutations in non-recombining regions, such as in sex chromosomes, may lead to differential expression between alleles if mutations occur stochastically in one or the other allele. Reduced allelic expression due to degeneration has indeed been suggested to occur in various sex-chromosome systems. However, whether an association occurs between specific signatures of degeneration and differential expression between alleles has not been extensively tested, and sexual antagonism can also cause differential expression on sex chromosomes. The anther-smut fungus Microbotryum lychnidis-dioicae is ideal for testing associations between specific degenerative signatures and differential expression because 1) there are multiple evolutionary strata on the mating-type chromosomes, reflecting successive recombination suppression linked to mating-type loci; 2) separate haploid cultures of opposite mating types help identify differential expression between alleles; and 3) there is no sexual antagonism as a confounding factor accounting for differential expression. We found that differentially expressed genes were enriched in the four oldest evolutionary strata compared with other genomic compartments, and that, within compartments, several signatures of sequence degeneration were greater for differentially expressed than non-differentially expressed genes. Two particular degenerative signatures were significantly associated with lower expression levels within differentially expressed allele pairs: upstream insertion of transposable elements and mutations truncating the protein length. Other degenerative mutations associated with differential expression included nonsynonymous substitutions and altered intron or GC content. The association between differential expression and allele degeneration is relevant for a broad range of taxa where mating compatibility or sex is determined by genes located in large regions where recombination is suppressed.

Key words: mating-type chromosomes, differential gene expression, sequence degeneration, transposable elements, premature stop codon, sexual antagonism.

Introduction

In plants and animals with differentiated sex chromosomes, recombination is largely suppressed in the sex chromosome that is always in a heterozygous state, that is, the Y or the W chromosome (Charlesworth et al. 2005; Bachtrog et al. 2014; Wright et al. 2016; Muyle et al. 2017). A consequence of such a lack of recombination is the diminished efficacy of selection, allowing deleterious mutations to accumulate (Charlesworth 1991; Rice 1996; Bachtrog 2006, 2008; Wright et al. 2016). The absence of recombination indeed reduces the effective population size, promotes the genetic hitchhiking of deleterious mutations with beneficial ones (Rice 1987), and prevents the purging of deleterious mutations (Rice 1996; Bachtrog 2005; Wright et al. 2016). Degenerative changes in the non-recombining regions include transposable element (TE) insertions, mutations causing alteration of protein function.

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such as early stop codons, and even gene copy loss. Such degenerative signatures are commonly found on the non-recombining Y or W chromosomes in many plants and animals (Charlesworth 2002; Bachtrog 2008; Kaiser et al. 2017; Muyle et al. 2017).

Degenerative changes have, in particular, the potential to lead to suboptimal gene expression. Indeed, reduced allelic expression has been reported as a form of degeneration on sex chromosomes (Bachtrog et al. 2008; Bachtrog 2013; Konuma et al. 2013; Hough et al. 2014; White et al. 2015; Pucholt et al. 2017; Xu et al. 2019), although most prior studies have focused on altered expression levels in relation to dosage compensation for differences in gene copy number on sex chromosomes between sexes (Ohno 1966; Charlesworth 1996; Mank 2009, 2013; Mank et al. 2011; Darolti et al. 2019). Studies that test whether various signatures of sequence degeneration are associated with differences in expression between alleles are generally lacking. Furthermore, sexually antagonistic selection is an alternative and frequently cited cause to explain differential expression (DE) on sex chromosomes and can render differences in gene expression between alleles challenging to interpret in the context of degenerative mutations (Rice 1987; Connallon and Knowles 2005; Ellegren and Parsch 2007; Parsch and Ellegren 2013).

Testing the association between allelic expression levels and specific degenerative signatures, particularly in systems without sexually antagonistic selection as a confounding factor, could yield insights into the evolution of gene expression in non-recombining regions. For instance, an early stop codon in an allele that truncates protein length can lead to post-transcriptional regulatory negative feedbacks upon expression (e.g., nonsense mediated decay; Montgomery et al. 2013). TE insertion in upstream promoter regions, or internal to genes, has long been recognized for effects on expression (McCintock 1942; Feschotte 2008; Cordaux and Batzer 2009; Tiross et al. 2009; Lee and Young 2013). Although less well recognized, base pair substitutions and in-frame indels (insertion or deletion mutations) can cause changes in amino acid sequence that affect gene expression through modulation of the mRNA translation (Kimball and Jefferson 2004) or disrupt promoter regions that impact transcriptional regulation (Wray et al. 2003). Epigenetic modifications, particularly cytosine methylation, contribute to both heterochromatin formation and elevated mutation rates that reduce GC content (Bird 1980; Grummt and Pikaard 2003); thus reduced GC content could represent a signature of methylation-induced gene silencing, among multiple other factors (Galtier et al. 2001; Meunier and Duret 2004). Shorter introns are more efficient for correcting transcription (Ma et al. 2015), such that changes in introns can influence transcription rates, nuclear export, and transcript stability (Heyn et al. 2015). These forms of degenerative changes are expected to accumulate under the reduced selection efficacy in non-recombining regions, and DE may occur where the mutations by chance affect one allele more than the other.

Fungal mating-type chromosomes share many features with sex chromosomes (Fraser and Heitman 2004; Hood et al. 2004) and can provide valuable insights into the relationship between various degenerative signatures and differential gene expression in non-recombining regions. Particular benefits of many fungi relative to other types of organisms are that antagonistic selection is not a confounding factor, an easy access to the haploid phase where alternate mating types are expressed, and the existence of young events of recombination suppression in successive evolutionary strata in linkage to the mating-type loci (Giraud et al. 2008; Fontanillas et al. 2015; Branco et al. 2017, 2018; Bazzicalupo et al. 2019). The anther-smut fungi, in the genus Microbotryum, undergo mating in the haploid phase via isogamous yeast-like cells of opposite mating types (a1 and a2), which can be cultured separately to analyze expression levels of alleles (Perlin et al. 2015). The species Microbotryum lycnthis-dioicae, causing anther-smut disease on the plant Silene latifolia, carries dimorphic mating-type chromosomes that have been assembled at the chromosome-level scale (Hood 2002; Hood et al. 2013; Branco et al. 2017). These mating-type chromosomes (a1, ~3.3 Mb, and a2, ~4.0 Mb) lack recombination across 90% of their length (Hood 2002; Hood et al. 2013) and are enriched in signatures of sequence degeneration compared to autosomes (Fontanillas et al. 2015). Because mating type is determined at the haploid phase, both mating-type chromosomes are always heterozygous and non-recombining, so that both degenerate (Fontanillas et al. 2015). Importantly, evolutionary strata of different ages have been identified, that is, regions with different levels of differentiation between mating types as a result of an expanding process of recombination suppression over the past 1.5 Myr (Branco et al. 2017, 2018). The non-recombining regions of the mating-type chromosomes in M. lycnthis-dioicae are flanked by small recombining pseudoautosomal regions (PARs).

In the isogamous fungus M. lycnthis-dioicae, there is no male or female function, so that there cannot be any sexual antagonism. Any analogous “mating-type antagonistic selection” (sensu Abbate and Hood 2010) would require fitness differences associated with mating-type dimorphic traits, which are necessarily expressed at the haploid stage when cells are of different mating types. However, there is only a very brief haploid stage in M. lycnthis-dioicae, as mating occurs readily after meiosis, within a tetrad, often before the haploid cells separate from the meiotic divisions (Day 1979; Garber and Day 1985; Hood and Antonovics 2000, 2004; Giraud et al. 2008). Moreover, a recent study on gene expression and positive selection detected no evidence for mating-type antagonistic selection (Bazzicalupo et al. 2019). This model system is therefore ideal for investigating the impact of degeneration on differential gene expression between chromosomes determining reproductive...
compatibility, notably without the confounding effect of sexual antagonism.

In this study, we therefore investigated whether genes that are differentially expressed between mating types were more often associated with various signatures of degeneration than non-differentially expressed genes in the genome of M. lychnidis-dioicae. We determined whether differential gene expression varied among genomic compartments defined as autosomes, PARs, youngest evolutionary strata on non-recombining regions of the mating-type chromosomes (i.e., the previously identified red and green evolutionary strata; Branco et al. 2017), and oldest evolutionary strata on non-recombining regions (i.e., the blue, purple, orange, and black evolutionary strata; Branco et al. 2017). We studied differential gene expression between mating types only in the haploid stage because the a1 and a2 mating types are determined at the haploid stage: Mating can only occur between haploid sporidia of opposite mating types. We determined whether DE was associated with greater differences in degenerative mutations between alleles, including comparisons for each degenerative trait between differentially and non-differentially expressed genes within each genomic compartment. We then assessed the possibility that the allele showing lower expression levels would have higher levels of degeneration footprints. The investigated degeneration signatures included differences between alleles in the levels of nonsynonymous sequence divergence, TE insertions, alteration of predicted protein length (via mechanisms including acquisition of indels and/or early stop codons), intron content, and GC content (Hoof and Green 1996; Marais et al. 2005; Feschotte 2008; Bedford and Hartl 2009; Cordaux and Batzer 2009). Associations between mating-type-specific DE and signatures of degeneration, while requiring further study to establish the nature of causality, would reflect an important component of gene evolution.

Materials and Methods

Allele Identification between a1 and a2 Haploid Genomes

In order to quantify differentially expressed genes between the two haploid genomes, we first identified the alleles between a1 and a2 haploid genomes for those genes. We used the a1 and a2 predicted coding gene sequences of the two haploid genome assemblies (accession number: a1—PRJEB12080, ERS459551, ERZ250722 and a2—PRJEB12080, ERS1013678, ERZ250721) (Branco et al. 2017, 2018). To identify 1:1 single-copy homologs in each haploid genome, the Reciprocal Best BLAST(p) Hits (RBBH) python script (github.com/peterjc/galaxy_blast/tree/master/tools/blast_rbh) was applied (Camacho et al. 2009), with 50% of length coverage. RBBH scripts also identified paralogs within each haploid genome. A number of protein sequence alignment identity thresholds were tested, in order to identify the best strategy for maximizing the number of allele pair identification on the non-recombining regions while avoiding spurious BLAST results with low identity percent. Increasing the percent of protein sequence identity threshold from >70% to >85% resulted in a decrease from 12.2% to 9.9% of single-copy genes on the mating-type chromosomes being identified as differentially expressed genes (detailed below), whereas decreasing the threshold from >70% to >30% resulted in only a marginal increase from 12.2% to 12.7%. The change in the percentages of identified alleles that were differentially expressed on autosomes was negligible, being 1.0%, 1.1% and 1.1%, respectively, for 80%, 70% and 30% thresholds (supplementary fig. S1, Supplementary Material online).

Therefore, the threshold of >70% protein sequence identity was used. Additionally, we also have detected orthologs between a1 and a2 genomes using OrthoMCL (Li et al. 2003). As the set of identified single-copy orthologous genes and their alleles was almost identical, differing in only 5 out of 9,396 genes compared with reciprocal best BLASTp analysis, we used the ortholog list generated by reciprocal best BLASTp for downstream analysis. To avoid potential bias due to paralogs for identifying differential gene expression and other downstream analysis, genes with paralogs within each haploid genome were filtered out and only single-copy allele pairs were retained for downstream analysis. Genes were assigned to genomic compartments, that is, autosomes, PARs, youngest evolutionary strata of the mating-type chromosomes (i.e., the previously identified red and green strata; Branco et al. 2017) and oldest evolutionary strata (i.e., the blue, purple, orange, and black strata; Branco et al. 2017). Finally, we confirmed the lack of small RNAs in our predicted coding gene sequences by checking the absence of noncoding RNAs in the M. lychnidis-dioicae predicted coding genes, using BLAST searches of ncRNA, tRNA, and rRNA sequences from the Rfam database (https://rfam.xfam.org; Griffiths-Jones et al. 2003; Kalvari et al. 2018). No ncRNA was found in the M. lychnidis-dioicae predicted coding gene set; the rRNA sequences detected in the genome did not have BLAST hits in the predicted coding gene set; only one tRNA sequence returned a partial hit (17.5% alignment), which had, however, not been retained in our 1:1 ortholog list.

TE Filtering

TE annotation of both haploid genomes of M. lychnidis-dioicae was published previously (Hartmann et al. 2018) and was used for analysis in this study. The coding sequence of each gene from both a1 and a2 haploid genomes was searched by BLASTn against the published annotated TE consensus sequences of the same species, and alignment >80% of query coverage (coding sequences) was used for identifications of TEs. The BLASTn output was parsed using BASH scripts, and the coding sequences identified as TEs were removed from the gene list for all further downstream analyses.
Identification of Differentially Expressed Genes

We studied differential gene expression between mating types only in the haploid stage because the \( a_1 \) and \( a_2 \) mating types are determined at the haploid stage: mating can only occur between haploid sporeidia of opposite mating types. In addition, almost all genes on autosomes and PARs are homozygous due to a high selfing rate (Badouin et al. 2017; Branco et al. 2017), so that we could assign expression levels to \( a_1 \) or \( a_2 \) mating types in the diploid or dikaryotic stages only for highly differentiated alleles, that is, on oldest evolutionary strata, which would profoundly limit and likely bias the analyses by taking into account only degenerated genes.

RNAseq samples and data sets were described previously (Perlin et al. 2015). Briefly, haploid sporeidial strains of the original isolate (the same as the reference genome “Lamole strain”) were generated from the meiotic products of a single tetrad. Then haploid fungal cells of either haploid \( a_1 \) or \( a_2 \) strain were grown separately on 2% water agar, each strain being grown in two replicates, with nutrient-free environment without the mating partner for 2 days, which essentially mimicked the natural conditions on the plant before mating and infection (Perlin et al. 2015). For RNAseq, polyA RNA was purified and a strand-specific library was constructed for each RNA sample; each library was sequenced with Illumina technology, generating on average 34.786 million 76-bp paired-end reads for \( a_1 \) libraries, and 35.017 million paired-end reads for \( a_2 \) libraries. The raw data of haploid culture growing separately in water agar conditions were downloaded from the deposited NCBI database (accession number: PRJNA246470). The RNAseq raw reads were quality assessed using FastQC v0.11.2 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and quality trimmed using Trimmomatic v0.33 with default parameters for paired-end reads for \( a_1 \) libraries, and was used to map the trimmed reads of haploid samples to the trimmed homolog sequences of each haploid genome, respectively, with Kallisto v.0.43.0 (Bray et al. 2016). Read counts of the output from Kallisto mapping (e.g., using pseudoalignment) were imported for gene expression analysis in EdgeR v3.4 (Robinson et al. 2010; McCarthy et al. 2012). We filtered low counts and kept genes with average LogCPM >0 per sample, and CPM (count per million) >1 in half of the total samples per haploid culture. We then normalized the expression using the weighted trimmed mean of M-values implemented in EdgeR, which is a scaling factor for library sizes that minimizes the log-fold change between samples. We explored the libraries of both haploid cultures in pairwise correlation of raw counts between replicates (supplementary fig. S2, Supplementary Material online), and two dimensions using multidimensional scaling plots (supplementary fig. S3, Supplementary Material online). Normalized expression counts for each sample were used to calculate DE between mating types using standard measures. We first identified genes with DE between mating types based on overall expression of the comparison group, and using Benjamini–Hochberg correction for multiple-testing with false discovery rate (FDR) of 5%. DE between mating types was classified into four categories of fold changes, namely \(<2, 2–4 \) (mild), \(4–8 \) (high), and \(>8 \) (very high), and expressed as \(\log_2 \) ratio of \(a_1\)-to-\(a_2\) expression (which has negative values for genes with higher \(a_2\) expression and positive values for higher \(a_1\) expression). We only considered genes with fold changes \(>1\) (i.e., \(|\log_2FC| > 0\) ), as recommended (Montgomery and Mank 2016), because we worked on haploid cell cultures and there were no possible scaling nor allometry issues due to whole-body sampling. Thus, unless stated otherwise, both conditions FDR \(< 0.05 \) and \(\log_2FC > 0\) were met when calling mating-type bias. Finally, to investigate the expression level of differentially (with \( a_1 \) or \( a_2 \) mating-type bias) and non-differentially expressed genes, we compared normalized read counts (transcripts per million, Log2TPM, obtained from EdgeR v3.4) of significantly expressed genes at autosomal and mating-type chromosomes (filtering criteria is the same as described above) from \( a_1 \) and \( a_2 \) samples (supplementary fig. S4, Supplementary Material online).

The classification of genes as having DE between mating types was based on the absolute values of gene expression ratio \(\log_2(a_1/a_2)\) and was used to assess relationships to various forms of mutational changes. A generalized linear model (GLM) analysis was used to assess the predictors of the absolute values of expression ratio \(\log_2(a_1/a_2)\), with the following main effect variables: genomic compartments, the absolute value of differences between alleles for sequence divergence (dN), TE insertions number within 20 kb (up and downstream), predicted protein length, intron content, and GC
Differential Gene Expression Correlates with Sequence Degeneration

content; we also included all two-way interaction terms in this model. The absolute value of the differences between alleles was calculated for each trait as detailed below. Model family comparison was based upon minimizing Akaike’s information criterion and over-/underdispersion using ratio of deviance/df; Tweedie, power 1.7 (approaching gamma distribution) provided the best available fit for the expression ratio response variable. A best-fit model was selected using stepwise model selection, following removal of non-significant interaction terms. Other post-hoc tests evaluating individual degeneration trait are described below. All statistical analyses were conducted in SPSS v23 (IBM Corp 2015) and R v3.4.3 (R Core Team 2017).

Relationship between DE and Elevated Substitution Rates

Each pair of allele was aligned with PRANK (v170427), using the codon model (Löytynoja and Goldman 2010), and each alignment was then analyzed with yn00 in PAML (Yang 2007) (runmode -2) to calculate the number of nonsynonymous substitutions per nonsynonymous site (dN), the number of synonymous substitutions per synonymous site (dS), and the ratio of the two (dN/dS), the latter excluding genes with dS value of zero. We then compared sequence divergence between alleles using nonparametric Wilcoxon rank sum tests for DE versus non-DE genes within genomic compartments.

We also compared the allele sequences of M. lychnidis-dioicae with their orthologs in Microbotryum lagerheimii, which carries mating-type chromosomes inferred to be largely collinear and a good proxy for the ancestral state in the Microbotryum genus (Branco et al. 2017, 2018). Pairs of a1 or a2 orthologs present in M. lychnidis-dioicae and M. lagerheimii were aligned with PRANK (v170427) using the codon model, then each ortholog alignment was analyzed with codeml (runmode -2) in PAML (Yang 2007). The single-copy orthologs for a1 or a2 genomes between M. lychnidis-dioicae and M. lagerheimii were identified using RBBH with 70% protein sequence coverage identity (github.com/peterjc/galaxy_blast/tree/master/tools/blast_rbbh; Camacho et al. 2009). Wilcoxon rank sum test was used to assess dN between orthologs in M. lychnidis-dioicae and M. lagerheimii to evaluate the hypothesis that the alleles with lower expression levels would have greater sequence divergence.

Relationship between DE and TE Insertions

The TE annotation of the M. lychnidis-dioicae genome published previously (Hartmann et al. 2018) was used for the analysis in this study. First, the TE insertion sites were assessed for each given focal gene, upstream 0–5 kb, 5–10 kb, 10–15 kb, 15–20 kb distance intervals, and downstream 0–5 kb, 5–10 kb, 10–15 kb and 15–20 kb distance intervals using Bedtools window function for each indicated distance window (https://bedtools.readthedocs.io/en/latest/content/tools/ window.html). Both annotation GFF3 files of gene models and TE annotations of M. lychnidis-dioicae were provided as input files. The output files were parsed using Bash scripts. Wilcoxon rank sum tests were used to compare TE insertions for DE and non-DE genes within genomic compartments.

A limited GLM model was used to assess the hypothesized directional association of TE insertions and reducing allelic expression (\(\log_2(a_1/a_2)\)); this model contained genomic compartment and oriented TE differences between alleles as main effects and their interaction term. Oriented TE differences between alleles were calculated as the TE number for the allele with lower expression minus the TE number for the higher expressed allele; a positive value thus represented an excess of TEs in the lower expressed allele. A sliding-window approach was used with a window size of three adjacent intervals, progressing from upstream to downstream of the genes.

Relationship between DE and Differences in Predicted Protein Length

We first tested whether there was no bias issue in the gene prediction model across genomic compartments, as degenerative mutation accumulation in the non-recombining region may decrease the accuracy of coding sequence prediction. The ratio of the predicted coding sequence length divided by three times the protein sequence length was consistently close to 1 and did not differ among genomic compartments (autosome, PAR, youngest strata, and oldest evolutionary strata; linear model, \(R^2 = -5.50e-05\), F-statistic = 0.869, \(P = 0.530\); supplementary fig. S5, Supplementary Material online). We therefore calculated the ratio of the predicted protein length between allele pairs, and compared the proportions of genes in DE and non-DE categories that had unequal lengths between alleles using two-proportion Z-test for genes within genomic compartments. The mutational causes of unequal protein lengths was assessed by manually quantifying premature stop codons or indels using Geneious v8.1.7 (Kearse et al. 2012). A limited GLM model was used to assess the hypothesized directional association of protein truncation and reducing allelic expression (\(\log_2(a_1/a_2)\)); this model contained genomic compartment and oriented predicted protein length differences between alleles as main effects and their interaction term. Oriented predicted protein length differences between alleles were calculated as the ratio for the allele with higher expression divided by the allele with lower expression; a larger ratio thus represented a shorter length for the allele with lower expression.

Relationship between DE and Intron Content

Using the published annotation gene models and coding sequences, we extracted the intron number and mean intron length information from the annotation gff3 file, using Perl script (https://bioops.info/2012/11/intron-size-gff3 perl). We investigated the differences in the proportional intron content for both DE and non-DE genes within genomic compartments.
using Wilcoxon rank sum test. We also used a limited GLM model to test the hypothesized directional association of greater intron content and reducing allelic expression ($\log_2(a_2/a_1)$); this model contained genomic compartments and oriented intron content differences between alleles as main effects and their interaction term. Oriented intron content differences between alleles were calculated as the value for the allele with lower expression minus the value for the allele with higher expression; a positive value thus represented greater intron content for the lower expressed allele.

**Relationship between DE and GC Content**

We calculated the total GC percentage (GC0) and the GC percentage at the third position of amino acid (GC3) for alleles of each gene coding sequence using homemade awk scripts. We investigated the differences of GC0 and GC3 for both DE and non-DE genes within genomic compartments using Wilcoxon rank sum test. We also used a limited GLM model to test the hypothesized directional association of reduced GC content and reducing allelic expression ($\log_2(a_2/a_1)$); this model contained genomic compartments and oriented GC content differences between alleles as main effects and their interaction term. Oriented GC content differences between alleles were calculated as the value for the allele with higher expression minus the value for the allele with lower expression; a positive value thus representing reduced GC content for the allele with lower expression.

**Results**

**Allele Identification and Differential Gene Expression between $a_1$ and $a_2$ Haploid Genomes**

We used the two available haploid genomes of *M. lychnidis-dioicae*, sequenced from separated haploid $a_1$ and $a_2$ mating-type cells derived from a single diploid individual, assembled at the chromosome-level scale and annotated (Branco et al. 2017, 2018). We investigated whether differential gene expression varies among genomic compartments defined as autosomes, PARS, youngest evolutionary strata, and oldest evolutionary strata of the mating-type chromosomes, and whether the greater degeneration signatures of several types were associated with the lower expressed allele. For this goal, we used whole-genome RNAseq data from each of two replicate cultures for separate $a_1$ and $a_2$ haploid mating-type sporidia of *M. lychnidis-dioicae*, under low nutrient conditions, that resemble the natural haploid growth environment (Schäfer et al. 2010; Perlin et al. 2015). Alleles of single-copy genes in *M. lychnidis-dioicae* were identified using the criterion of 1:1 reciprocal best BLASTp between the $a_1$ and $a_2$ haploid genomes, which was highly consistent with the results by the OrthoMCL approach, as the two methods gave nearly (99.95%) identical results. After filtering out TE-related gene sequences, we identified 371 single-copy allelic pairs in mating-type chromosomes and 9,025 in autosomes (supplementary table S1, Supplementary Material online). We retained 8,549 single-copy genes with significantly detectable expression (see filtering criteria and details in Materials and Methods section) for further analysis (342 on mating-type chromosomes and 8,207 on autosomes). We analyzed 8,549 genes and used the threshold for differential gene expression of $\log_2(a_2/a_1)$ being significantly greater than zero, with FDR < 0.050. This revealed 392 genes (4.59%) that were significantly more highly expressed in the $a_1$ haploid culture and 203 (2.37%) that were significantly more highly expressed in the $a_2$ haploid culture (fig. 1 and supplementary table S2 and fig. S4, Supplementary Material online).

**Differential Gene Expression and Multiple Signatures of Sequence Degeneration**

Regression analysis (GLM) revealed that the degree of DE between allele pairs of the two haploid mating types significantly increased with increasing differences between alleles (using absolute values) in the various degeneration traits examined (table 1). We found no asymmetry in reduced allelic expression or degeneration features between $a_1$ and $a_2$ mating-type chromosomes, so we combined genes with $a_1$ or $a_2$ mating-type biased expression as the set of DE genes in the following analyses. The significant main-effect predictors of DE included genomic compartment and differences between alleles in non-synonymous divergence (dN), TE insertion number within 20 kb (up and downstream of genes), intron content (proportional to coding sequence length), and overall GC content (GC0). Differences between alleles in predicted protein length (via mechanisms including acquisition of indels and/or early stop codons) were not a significant main-effect predictor but strongly significant as an interaction term with genomic compartment and all other traits except intrinsic content (table 1 and fig. 2). DE indeed increased with differences between alleles in protein length but only in the oldest evolutionary strata and when associated with higher differences between alleles in dN, TE content, and GC0 (table 1 and fig. 2). The oldest evolutionary strata were significantly enriched in genes with DE between mating types compared with autosomes, whereas the youngest evolutionary strata or the PARS were not (table 2). Similar patterns were observed in the comparisons using each of the $a_1$ or $a_2$ haploid genomes separately (supplementary table S3, Supplementary Material online). Further post-hoc assessments of degenerative traits are presented in the following sections, including whether the difference between alleles is oriented such that the more affected allele is less expressed.

**Relationship between DE and Elevated Substitution Rates**

Differentially expressed genes had greater sequence divergence between alleles than non-differentially expressed (non-DE) genes within genomic compartments, specifically...
within the oldest evolutionary strata of the mating-type chromosomes. DE genes had significantly higher nonsynonymous mutation rate ($dN$) and synonymous mutation rate ($dS$) between alleles than non-DE genes within the oldest evolutionary strata (Wilcoxon rank sum test for independent samples, $dN$: $W = 1,433, P < 0.001$, $dS$: $W = 1,422, P < 0.001$) (fig. 3A and supplementary fig. S6 and table S4, Supplementary Material online). There was almost no sequence divergence ($dN$ or $dS$) between alleles on either autosomes or PARs for DE or non-DE genes. The youngest evolutionary strata had only one DE gene, precluding comparison to non-DE genes within this compartment. The finding that DE genes had significantly higher $dN$ and $dS$ in the oldest strata held for both genes with higher expression in $a_1$ cells ($dN$: $W = 1,156.5, P = 0.019$) and those with higher expression in $a_2$ cells ($dN$: $W = 1,968.5, P = 0.003$; $dS$: $W = 2,108, P = 1.539e-4$) when considered separately, except the $dS$ in $a_1$ ($W = 1,028.5, P = 0.172$; supplementary figs. S7A and B, Supplementary Material online). In addition, there were no significant differences in $dS$ or $dN$ between $a_1$-biased and $a_2$-biased gene sets (supplementary fig. S7, Supplementary Material online, $dS$: $W = 154$, $P = 0.122$; $dN$: $W = 226, P = 0.811$). A tendency of higher $dN/dS$ for DE genes compared with non-DE genes was not significant within the oldest strata ($W = 1,946, P = 0.611$) (supplementary fig. S8, Supplementary Material online).

As DE genes were found to be associated with higher $dN$ than non-DE genes, we then tested within DE genes whether the allele with lower expression was associated with greater accumulation of nonsynonymous (or synonymous) changes. This was assessed by computing the sequence divergence between each allele of DE genes in *M. lychnidiis-dioicae* with their ortholog in *M. lagerheimii*, which has retained largely collinear and homozygous mating-type chromosomes, and was inferred to have retained an ancestral chromosomal state in the *Microbotryum* genus (Branco et al. 2017, 2018). For DE genes, the allele that had lower expression levels in *M. lychnidiis-dioicae* did not show significantly greater accumulation of nonsynonymous changes than the allele with higher expression. The $dN$ divergence between *M. lychnidiis-dioicae* alleles (either $a_1$ or $a_2$) and their ortholog in *M. lagerheimii* was not
greater for alleles having lower expression levels than alleles with higher expression levels in *M. lychnidis-dioicae* ($W = 1,267$, smallest $P = 0.909$) (supplementary fig. S9 and table S5, Supplementary Material online).

**Relationship between DE and TE Insertions**

Differentially expressed genes were associated with greater differences between alleles for TE insertions (within 20 kb upstream and downstream of genes) than alleles of non-DE genes across genomic compartments. However, the difference was significant only in the autosomes ($W = 313,879$, $P < 0.001$; fig. 3B), not in the PARs ($W = 546$, $P < 0.192$) or the oldest evolutionary strata ($W = 4,062$, $P = 0.173$); the comparison was not possible in the youngest evolutionary strata, as noted above.

The alleles with lower expression had more TE insertions than the alleles with higher expression. This was assessed by calculating the differences in TE insertion numbers (upstream and downstream of genes) between alleles, as the TE number
for the allele with lower expression minus the TE number for the allele with higher expression; a positive value thus represented an excess of TEs in the less expressed allele. This oriented TE number difference between alleles was tested as a predictor of the expression ratio $\log_2(a_1/a_2)$ using a sliding window approach with a 15-kb window size overlapping by 5 kb. Among DE genes, oriented TE insertion difference was a significant predictor of the expression ratio only in the window covering from 10 kb upstream to the gene (fig. 4A and supplementary fig. S10, Supplementary Material online); alleles with more TE insertions had reduced expression (for this window, Wald $X^2 = 6.674$, $P = 0.010$, statistics of remaining windows in supplementary table S6, Supplementary Material online). Among non-DE genes, none of the windows was a significant predictor of variation in the expression ratio (supplementary table S6, Supplementary Material online).

Relationship between DE and Differences in Predicted Protein Length

Differential gene expression was associated with the mutational changes that affect the predicted protein length, including altered stop codon positions, indels (including those causing frameshifts). Within genomic compartments, alternate alleles of DE genes were significantly more likely to produce proteins of different lengths than alleles of non-DE genes, particularly within the oldest evolutionary strata (two-proportion $Z$-test, $Z = 2.186$, $P = 0.029$) and autosomes ($Z = 4.64$, $P = 8.78\times 10^{-06}$; fig. 3C and supplementary table S7, Supplementary Material online); there were too few DE genes on PARs and youngest evolutionary strata for statistical comparisons.

The various types of mutational changes that caused protein length variation between alleles differed between DE and non-DE genes, as well as among genomic compartments. Among the 258 genes with different protein sequence lengths between alleles, all had indels. However, DE genes in the oldest evolutionary strata and autosomes had significantly more indels than non-DE genes; mean indel number in oldest strata differed between alleles by 2.64 for DE genes and by 1.85 for non-DE genes ($W = 2,453.5$, $P = 0.013$), and in autosomes alleles differed by a mean of 1.19 indels for DE genes and 1.03 for non-DE genes ($W = 490.5$, $P = 0.025$; fig. 5A); PARs and youngest evolutionary strata could not be analyzed.

Similarly, differences in the positions of stop codons contributed to protein length variation more for DE genes than for non-DE genes. Among genes with different protein

### Table 1

Output of a Reduced Best-Fit GLM with Differential Gene Expression ($\log_2(a_1/a_2)$) As the Response Variable and the Following Predictable Variables: Genomic Compartment and Various Degeneration Traits, That Is, Nonsynonymous Substitution Rate (dN), TE Insertions, Protein Length, Intron Content, and GC Content

| Explanatory Variables and Interaction Terms | GLM Model Output Parameter |
|---------------------------------------------|----------------------------|
|                                            | Wald Chi-Square | Degree of Freedom (df) | P-value | Regression Coefficient |
| (Intercept)                                 | 496.78         | 1                       | <0.001  | NA                     |
| Compartment                                 | 20.151         | 3                       | <0.001  | NA                     |
| dN                                          | 13.21          | 1                       | <0.001  | 5.081                  |
| TE insertions                               | 8.405          | 1                       | 0.004   | 10.612                 |
| Protein length                              | 0.41           | 1                       | 0.522   | 0.768                  |
| Intron content                              | 10.209         | 1                       | 0.001   | 0.499                  |
| GC content                                  | 4.233          | 1                       | 0.040   |                       |
| Compartment * Protein length                | 24.662         | 3                       | <0.001  | NA                     |
| dN * Protein length                         | 13.36          | 1                       | <0.001  | 50.726                 |
| TE insertions * Protein Length              | 8.398          | 1                       | 0.004   | –0.37                  |
| GC content * Protein length                 | 10.801         | 1                       | 0.001   | –3.962                 |

**NOTE.**—$P$-values < 0.05 are in bold. NA, not applicable.

### Table 2

Numbers and Percentages of Genes with DE on the Different Genomic Compartments on Mating-Type Chromosomes and Autosomes, and Fisher’s Exact Test for Even Distribution between DE Genes on Autosomes and Other Genomic Compartments, Including PARs, Youngest Evolutionary Strata (previously identified red and green strata; Branco et al. 2017) and Oldest Evolutionary Strata (blue, purple, orange, and black strata; Branco et al. 2017)

|                | Autosomes | PAR   | Youngest Strata | Oldest Strata |
|----------------|-----------|-------|-----------------|---------------|
| DE gene number | 507       | 12    | 1               | 74            |
| Total number   | 8,207     | 114   | 29              | 198           |
| Percentage     | 6.18%     | 10.53%| 3.45%           | 37.37%        |
| Fisher’s exact test ($P$-value) | NA | 0.085 | 1 | 2.20E-16 |

**NOTE.**—$P$-values < 0.05 are in bold. NA, not applicable.
lengths between alleles in the oldest evolutionary strata, 44.6% ($N = 56$) of DE genes had different stop codon positions between alleles, which was significantly higher than the 24.0% ($N = 75$) of non-DE genes (two-proportion Z-test, $P = 0.018$; fig. 5B). Similarly, DE genes in the autosomes were marginally significantly more likely to have different stop codon positions between alleles than non-DE genes, with 33.3% ($N = 21$) versus 10% ($N = 40$), respectively.

**FIG. 3.**—Comparisons of differentially expressed (DE) versus nondifferentially expressed (non-DE) genes between mating types of *Microbotryum lychnids-dioicae* for various degeneration-associated traits within genomic compartments. (A) Nonsynonymous sequence divergence, $d_N$, between alleles of DE and non-DE genes. (B) TE insertion number differences between alleles within 20 kb (upstream and downstream) of DE and non-DE genes. (C) Proportions of differentially expressed (DE) and non-differentially expressed (non-DE) genes with different protein lengths between alleles. (D) Intron content proportional differences between alleles of DE and non-DE genes. (E) Total GC content (GC0) proportional differences between alleles of DE and non-DE genes. Analyzed allele differences represent absolute value comparisons (i.e., unoriented with regard to allelic expression levels). Comparisons in panels (A), (C)–(E) reflect Wilcoxon rank sum tests; panel (B) reflects a two-proportion Z-test. Significance levels shown as ***$P < 0.001$, *$P < 0.05$; non-significant test results shown in supplementary tables S4 and S6–S9, Supplementary Material online. Genomic compartments include autosomes, PARs, youngest and oldest evolutionary strata. The notation “a” indicates that the youngest evolutionary strata contained only one DE gene, precluding comparisons to non-DE genes within this compartment. For boxplot, the horizontal bars (from bottom to top) represent the 25% quartile, median, and 75% quartile, respectively.
Only three frameshift mutations were observed among the 258 genes with different protein/coding sequence lengths, and thus frameshifts were not a distinguishing feature of DE versus non-DE genes. The alleles with lower expression did show a pattern of truncation of protein length compared with the alleles with higher expression (i.e., by early stop codons or deletions). To assess this pattern, differences in protein lengths between alleles were calculated as the ratio of protein length for the allele with higher expression divided by the allele with lower expression; a larger ratio thus represented a shorter length for the allele with lower expression. Among DE genes, this oriented metric of protein length differences was a significant predictor of the DE degree as the ratio $\log_2(a_1/a_2)$, with alleles producing shorter proteins being less expressed ($\chi^2 = 19.326$, two-tailed $P < 0.001$; fig. 4B). Among non-DE genes, in contrast, we found no significant relationship between the expression level ratio and the oriented protein length ratio ($\chi^2 = 0.222$, $P = 0.638$; fig. 4B).

### Relationship between DE and Intron Content

Differential gene expression was associated with differences between alleles in intron content (the ratio of intron to coding sequence length), lower intron content being considered to be favored by selection (Marais et al. 2005). There were significantly greater intron content differences between alleles for DE genes than for non-DE genes; considering the ratio of intron to coding sequence lengths, alleles of DE genes overall differed on average by 0.008 and alleles of non-DE genes differed by 0.002 ($W = 2,102,758$, $P < 0.001$). Alleles differed in intron content more for DE than non-DE genes within the autosomes ($W = 1,920,124$, $P = 0.033$) and oldest evolutionary strata ($W = 3,205$, $P = 0.001$) (fig. 3D and supplementary table S8, Supplementary Material online) but not within the PARs ($W = 605$, $P = 0.888$); the comparison in youngest evolutionary strata was not possible.

The alleles with lower expression did not show a greater intron content than the alleles with higher expression. This was assessed by calculating differences between alleles as the value for the less expressed allele minus the value for the more expressed allele; a positive value thus represented greater intron content for the less expressed allele. This oriented metric of intron content differences between alleles was not a significant predictor of DE level among DE genes ($\chi^2 = 0.350$, $P = 0.554$) or among non-DE genes ($\chi^2 = 0.216$, $P = 0.642$).

### Relationship between DE and GC Content

DE genes had significantly greater overall GC0 differences between their alleles than non-DE genes within the autosomes ($W = 1,907,831$, $P < 0.001$) and oldest evolutionary strata ($W = 2,102,758$, $P < 0.001$). Alleles of DE genes overall differed on average by 0.008 and alleles of non-DE genes differed by 0.002 ($W = 2,102,758$, $P < 0.001$). Alleles differed in intron content more for DE than non-DE genes within the autosomes ($W = 1,920,124$, $P = 0.033$) and oldest evolutionary strata ($W = 3,205$, $P = 0.001$) (fig. 3D and supplementary table S8, Supplementary Material online) but not within the PARs ($W = 605$, $P = 0.888$); the comparison in youngest evolutionary strata was not possible.

The alleles with lower expression did not show a greater intron content than the alleles with higher expression. This was assessed by calculating differences between alleles as the value for the less expressed allele minus the value for the more expressed allele; a positive value thus represented greater intron content for the less expressed allele. This oriented metric of intron content differences between alleles was not a significant predictor of DE level among DE genes ($\chi^2 = 0.350$, $P = 0.554$) or among non-DE genes ($\chi^2 = 0.216$, $P = 0.642$).
In addition to antagonistic selection, the general phenomenon in nonrecombining sex chromosomes that includes degenerative mutations, which is likely a consequence of the studied degenerative mutations or from relaxed sequence degeneration, which can result either from a direct effect of the studied degenerative mutations or from relaxed selection following changes in expression levels. Our results, neither the oriented GC0 nor GC3 differences between alleles were significant predictors of the level of DE (GC0: Wald $X^2 = 1.039$, $P = 0.308$, and GC3: Wald $X^2 = 2.226$, $P = 0.136$).

**Discussion**

In the anther-smut fungus *M. lychnidis-dioicae*, the mating-type chromosomes harbor evolutionary strata of various ages and lack sexually antagonistic selection, while sharing nonrecombining and heterogametic characteristics with sex chromosomes (Hood and Antonovics 2004; Giraud et al. 2008; Branco et al. 2017, 2018). We found no asymmetry in reduced allelic expression or degeneration features between $a_1$ and $a_2$ mating-type chromosomes, as expected given their lack of heterozygosity asymmetry and their lack of ecological differences (Hood 2002). Only the oldest strata of the mating-type chromosomes were enriched in genes differentially expressed between the haploid mating types, as found previously (Bazzicalupo et al. 2019). Most importantly, our study provides evidence for associations between DE and several different signatures of degenerative changes in the *M. lychnidis-dioicae* mating-type chromosomes. The differentially expressed genes displayed various forms of sequence (dN, dS, or GC content) and structural (TE insertions, intron content, or protein length) heterozygosity at levels higher than nondifferentially expressed genes within genomic compartments (i.e., autosomes, PARs, the youngest strata, and the oldest strata). These results show that differential gene expression is strongly associated with sequence degeneration, which can result either from a direct effect of the studied degenerative mutations or from relaxed selection following changes in expression levels. Our results thus support the view that DE should be interpreted in a context that includes degenerative mutations, which is likely a general phenomenon in nonrecombining sex chromosomes in addition to antagonistic selection.

**Differential Gene Expression between Haploid Mating Types**

The proportion of genes with DE between haploid mating types of *M. lychnidis-dioicae* was low but slightly higher than in a previous study based on the same data set (Fontanillas et al. 2015), likely due to an improved genome assembly and non-recombining region identification. The proportion of genes with DE between mating types in *M. lychnidis-dioicae* (6.96%) was similar to the proportion of genes with DE between sexes in plant and animal non-reproductive tissues, for example, liver, spleen, leaves, and roots (Yang et al. 2006; Ayroles et al. 2009; Perry et al. 2014; Haselman et al. 2015; Meisel et al. 2017; Ma, Veltsos, Sermier, et al. 2018; Ma, Veltsos, Toups, et al. 2018). However, this proportion was much lower than in reproductive tissues (e.g., ovaries or testes) of most animals and plants (Ellegren and Parsch 2007; Parsch and Ellegren 2013).

Differentiated sex chromosomes in anisogamous animals and plants are often enriched in DE genes (reviewed by Dean and Mank 2014; Mank et al. 2014). In fungi other than *M. lychnidis-dioicae*, such as *Neurospora tetrasperma* and *Podospora anserina*, DE genes were also more frequently detected on non-recombining regions of the mating-type chromosomes than in autosomes, which has been interpreted as resulting from ecological differences particular to those species, that is, differences in terms of vegetative or sexual growth between mating types (Samils et al. 2013; Grognet et al. 2014). In animals and plants, sexual antagonism occurs when trait values that increase gene transmission through the male function then decrease gene transmission through the female function, or conversely (Lande 1980; Rice 1987; Charlesworth et al. 2014; Dean and Mank 2014). The linkage of sexually antagonistic genes to the sex-determining genes in non-recombining regions is considered fundamental to the formation of evolutionary strata and the resolution of sexual conflict, for example, by allowing for sex-specific or sex-biased gene expression (Rice 1987; Charlesworth et al. 2005; Otto et al. 2011; Lipinsk et al. 2017). Differentially expressed genes on differentiated sex chromosomes are therefore often considered to resolve sexual antagonism or sexual conflicts between females and males, even if several studies have shown that differentially expressed genes could also be associated with degenerative mutations (Bachtrog et al. 2008; Wright et al. 2012; Hough et al. 2014; White et al. 2015; Xu et al. 2019). Our findings support the view that degenerative mutations likely contribute to levels of DE in sex-related chromosomes, whereas it is important to note that this is not exclusive of, and should be considered in addition to, the hypothesis invoking the role of sexual antagonism.

Our results thus encourage a broader view of the evolutionary forces related to DE genes and their enrichment on chromosomes determining reproductive compatibility. In *M. lychnidis-dioicae*, any differential gene expression between the alternative haploid mating types is unlikely due to “mating-type antagonistic selection” given the lack of female and male functions and the lack of haploid phase beyond the tetrad stage. Little evidence was found for a role of antagonistic selection in driving the evolution of new evolutionary...
strata in mating-type chromosomes of *M. lychnidis-dioicae* (Bazzicalupo et al. 2019). Young evolutionary strata were indeed found not to be enriched in genes upregulated in the haploid mating phase compared with the dikaryotic stage, nor in genes differentially expressed between mating types, nor in genes displaying footprints of specialization (i.e., high \(dN/dS\)) between mating types (Bazzicalupo et al. 2019). Importantly, gene degeneration is expected to occur commonly on nonrecombining sex or mating-type chromosomes due to the reduced efficacy of selection caused by the absence of recombination (Rice 1987; Charlesworth et al. 2005; Ellegren and Parch 2007; Otto et al. 2011; Fontanillas et al. 2015; Lipinska et al. 2017). The resulting mutation accumulation may generate contrasting expression levels between differently affected alleles or may accumulate where DE decreases purifying selection on the less expressed allele. Again, sexually antagonistic selection and degeneration are not mutually exclusive processes for explaining sex-biased gene expression. Still, the potential role of degenerative mutations remains worth highlighting in all systems. The association found between multiple types of degeneration footprints and DE in *M. lychnidis-dioicae*, in the absence of sexual antagonism, suggests the possibility of similar associations across diverse types of organisms. Finally, previous studies showed that there was overall very low genetic variation within *M. lychnidis-dioicae*, both on autosomes and within each of \(a_1\) and \(a_2\) mating-type chromosomal systems, due to high selfing rates and small effective population sizes (Badouin et al. 2017; Branco et al. 2017). Therefore, we expect most \(a_1\) mating types to be similar in expression and most \(a_2\) mating types likewise. Further studies using more individuals and species would nevertheless be interesting to see the degree to which these associations form general and consistent patterns.

Various Forms of Degeneration

One of the main insights gained in the present study was the directional associations found between DE and specific signatures of degeneration. The properties of nonrecombining regions that reduce the efficacy of selection (reduced effective population size, hitchhiking, and lack of deleterious mutation purging) can lead to the fixation of various mutations having degenerative effects, several of which were significant predictors in the overall regression model of DE between mating types of *M. lychnidis-dioicae*. Even within genomic compartments, DE genes had significantly higher levels of degenerative mutations distinguishing the alleles than non-DE genes. Some signatures of degeneration, such as TE insertions, indels, and/or premature stop codons, may be most plausibly conceived as mechanisms that reduce transcription levels. In particular, TE insertions into genes or upstream have long been recognized to alter gene expression (McClintock 1942; Britten and Davidson 1971). TEs can disrupt promoter regions or other regulatory sequences internal to genes (Feschotte 2008; Cordaux and Batzer 2009). In addition, epigenetic silencing, as a defense against TE proliferation, can tighten local chromatin structure and inhibit access of transcriptional machinery (Eichten et al. 2014; King 2015). Consistent with a direct effect upon DE, the relative excess of TE insertions between alleles, specifically upstream of genes, was associated with a lower expression level between alleles of DE genes. Similarly, the introduction of early stop or nonsense codons may reduce expression. Transcripts from alleles with premature stop codons are affected by nonsense-mediated decay, involving the degradation of mRNA and further components of the RNAi pathway that downregulate expression (Hoof and Green 1996). We found that an allele with a shorter predicted protein length in DE genes was indeed more likely to have lower expression. Although TE insertions, indels or premature stop codons are potentially important mutations affecting DE between alleles, it is difficult to establish causal relationship from our current data sets, and further studies are needed to directly test the nature of causality between DE and specific degenerative mutations. For instance, in investigating the degeneration of the neo-Y chromosome of *Drosophila albomarginis*, Zhou and Bachrög (2012) provided evidence that TEs accumulated mostly after regulatory changes in gene expression. Furthermore, the interplay between mutations and expression may be complex; for example, an early stop codon might not directly reduce expression, but it still may cause partial loss in protein function that reduces the strength of selection to maintain the expression of the allele.

Other signatures of degeneration in *M. lychnidis-dioicae* were not directionally predictive of lower allelic expression but were nevertheless more associated with DE than non-DE genes. Most substantial among these characteristics was the degree of sequence divergence between alleles. Alleles of DE genes were distinguished by markedly more nonsynonymous and synonymous base pair differences than alleles of non-DE genes. Similar results were obtained in the anisogamous, hermaphroditic ascomycete *N. tetrasperma*, showing that differential gene expression was positively correlated with sequence divergence between alleles on mating-type chromosomes (Samils et al. 2013).

Degeneration across Genomic Compartments

The different forms of genetic degeneration in *M. lychnidis-dioicae* were not equally represented across genomic compartments, perhaps reflecting the history of recombination suppression. In this system, enrichment of DE genes on the mating-type chromosomes is unlikely to be due to antagonistic selection. As a matter of fact, enrichment of DE genes was significant only in the oldest evolutionary strata and not in the younger strata, indicating that it is a consequence and not a driver of recombination suppression.

Mating between different haploid sexes or mating types ensures that all diploids are heterogametic (Bull 1978), and it
has long been recognized that regions linked to mating-type loci can preserve heterozygosity (Mather 1942). In *M. lycnids-dioicae*, the large non-recombining regions are in fact highly heterozygous (Branco et al. 2017). In contrast, the autosomes and PARs are largely homozygous, due to the selfing mating system of *M. lycnids-dioicae* (Hood and Antonovics 2000, 2004; Giraud et al. 2008). Consistent with mating-type linkage preserving heterozygosity, nearly the full range of mutational changes or footprints of degeneration showed lowest levels in the autosomes and PARs and increasing from lowest levels in the youngest evolutionary strata to highest levels in the oldest evolutionary strata. Importantly, however, comparisons within genomic compartments repeatedly showed that allele-distinguishing mutations occurred more in association with DE genes than non-DE genes or in the manner positively associated with levels of DE. This represents strong evidence for these degenerative changes being associated with changes in expression levels between alleles. Finally, the recent discovery of multiple independent mating-type linkage events across the *Microbotryum* genus (Branco et al. 2018) should allow further assessment of mutation accumulation and its consequences for gene functions.

**Conclusions**

Our findings on differential gene expression, being more frequent in oldest evolutionary strata and being associated with various types of sequence degeneration, and without sexually antagonistic selection as confounding factor, shed new lights on how differentially expressed genes might evolve in non-recombining regions in general, such as sex chromosomes or mating-type chromosomes. Our study showed that the accumulation of degenerative mutations between alleles was significantly associated with the degree of differential gene expression, in a system where sexually antagonistic selection is unlikely to occur as a confounding factor. Furthermore, the genes with DE were highly enriched on mating-type chromosomes, as in diverse organisms where the separate sex functions have been cited as the primary cause. We further found evidence of a directional relationship between differential gene expression and some types of mutational changes, in particular TE insertions and premature stop codons, being greater in the alleles with lower expression levels, although a causal relationship remains to be demonstrated. Our results suggest an important relationship between mutation accumulation and DE between alleles, which is relevant to a broad range of taxa where reproductively compatible, sex or other complex traits are determined in extensive regions of recombination suppression.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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