Microsporidia with Vertical Transmission Were Likely Shaped by Nonadaptive Processes

Karen L. Haag1,*†, Jean-François Pombert2,†, Yukun Sun2, Nathalia Rammé M. de Albuquerque1, Brendan Batliner2, Peter Fields3, Tiago Falcon Lopes1, and Dieter Ebert3

1Department of Genetics and Post-Graduation Program of Genetics and Molecular Biology, Federal University of Rio Grande do Sul, Porto Alegre, Brazil
2Department of Biology, Illinois Institute of Technology
3Department of Environmental Sciences, Zoology, Basel University, Switzerland

†These authors contributed equally to this work.
*Corresponding author: E-mail: karen.haag@ufrgs.br.
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Abstract

Microsporidia have the leanest genomes among eukaryotes, and their physiological and genomic simplicity has been attributed to their intracellular, obligate parasitic life-style. However, not all microsporidia genomes are small or lean, with the largest dwarfing the smallest ones by at least an order of magnitude. To better understand the evolutionary mechanisms behind this genomic diversification, we explore here two clades of microsporidia with distinct life histories, Ordospora and Hamiltosporidium, parasitizing the same host species, Daphnia magna. Based on seven newly assembled genomes, we show that mixed-mode transmission (the combination of horizontal and vertical transmission), which occurs in Hamiltosporidium, is found to be associated with larger and AT-biased genomes, more genes, and longer intergenic regions, as compared with the exclusively horizontally transmitted Ordospora. Furthermore, the Hamiltosporidium genome assemblies contain a variety of repetitive elements and long segmental duplications. We show that there is an excess of nonsynonymous substitutions in the microsporidia with mixed-mode transmission, which cannot be solely attributed to the lack of recombination, suggesting that bursts of genome size in these microsporidia result primarily from genetic drift. Overall, these findings suggest that the switch from a horizontal-only to a mixed mode of transmission likely produces population bottlenecks in Hamiltosporidium species, therefore reducing the effectiveness of natural selection, and allowing their genomic features to be largely shaped by nonadaptive processes.

Key words: microsporidia, genome evolution, population genomics, transmission modes, neutral evolution, genetic drift.

Introduction

Streamlining in the context of microbial evolution refers to the minimization of cell and genome size and complexity (Novichkov et al. 2009; Giovannoni et al. 2014). In populations with large effective population sizes, \( N_e \), natural selection enables even mutations with small positive selection coefficients, that is, “slightly” beneficial mutations to go to fixation. Likewise, mutations with small negative selection coefficients (mildly deleterious mutations) can rapidly be eliminated. Considering that the maintenance of any genomic segment involves a cost, and that the efficiency of selection depends on \( N_e \), theory predicts a negative correlation between population and genome sizes (Lynch and Conery 2003; Lynch 2007b). With huge \( N_e \) at the order of \( 10^9 \) and compact genomes tightly packed with protein-encoding genes, free-living bacteria are the primary examples for genome streamlining due to natural selection (Bobay and Ochman 2017).

In contrast, natural selection is less effective in populations with low \( N_e \), where there is greater influence of genetic drift (Lynch 2007a). An example for this is bacterial lineages that turned endosymbionts. They are believed to suffer a strong reduction in population size due to host restriction (Mira and Moran 2002; Moran and Plague 2004), and for obligate
symbionts, their transmission is exclusively vertical. The evolution of the small genomes of many endosymbionts is believed to be mostly driven by genetic drift at low $N_e$—not natural selection—with the random fixation of mildly deleterious mutations. Although loss of function by natural selection can occur if selection coefficients are greater than about $1/2N_e$ (Hottes et al. 2013), metabolic degeneration of mutualistic symbiotic bacteria is caused by a process of genomic erosion that reflects both a mutualistic bias toward deletions and the reduced efficacy of natural selection in maintaining gene functionality (Mira et al. 2001; Kuo and Ochman 2009; Nowack and Weber 2018). Bacterial populations that have been serially passaged to simulate recurrent bottlenecks revealed that extensive genome reduction can occur on a short evolutionary time scale (Nilsson et al. 2005). This reduction is often accompanied by an overall degeneration of the DNA repair machinery, leading to increased mutations rates and a decrease in GC content, as DNA damage such as cytosine deamination and guanine oxidation bias mutations toward A and T bases (Moran 1996; McCutcheon and Moran 2012).

In eukaryotes, the most extreme genomic reduction is found in some microsporidia, obligate intracellular parasites related to fungi. Because most microsporidia have no mitochondria (but see Haag et al. [2014] for an exception), they rely heavily on their hosts for their energetic requirements. Microsporidian genomes are poor in genes involved in resource-producing metabolic pathways—such as ATP synthesis—and rich in genes that enhance transport mechanisms and enable the hijacking of resources from the host (Keeling et al. 2010; Nakjang et al. 2013; Boakye et al. 2017). This adaptation has been associated with their intracellular lifestyle (Peyretailleade et al. 2011; Corradi and Slamovits 2011). With genomes smaller than 3 Mb and coding for about 2,000 proteins, microsporidian species from the genus *Encephalitozoon* are considered paragons of eukaryote streamlining (Katinka et al. 2001; Corradi et al. 2010; Pombert et al. 2012). However, this streamlining did not occur in all microsporidians and several species harbor genomes larger by an order of magnitude (Williams et al. 2008). Genome size variability in microsporidia has been attributed in part to the differential accumulation of transposable elements (Xu et al. 2006; Parisot et al. 2014) but this accumulation cannot, by itself, explain such large differences, suggesting that different evolutionary forces are at play.

An often-overlooked aspect of microsporidian evolution is the diversity present in their life cycles, with some species even relying on more than one host (Becnel and Andreadis 2014). Transmission in microsporidian species can be horizontal, vertical, or both (mixed mode), and their capacity to perform standard meiosis is questionable (Lee et al. 2014). In a few species with well-characterized life cycles, abortive meiosis is observed (Canning et al. 1999), and population genetic studies suggest that some microsporidia are clonal (Haag, Traunecker, et al. 2013). *Daphnia magna* is the host of a diversity of microsporidia with divergent life-styles and genomic features (Ebert 2005; Corradi et al. 2009; Haag et al. 2014; Pombert et al. 2015) offering the unique opportunity to investigate the influence of microsporidian life history strategies on genome evolution while keeping the host factor constant. Here, to investigate the impact of transmission modes on the evolution of genomic architectures of microsporidia, we sequenced seven genomes from two clades of microsporidia that specifically parasitize *D. magna* with contrasting life histories: *Hamiltosporidium* and *Ordospora*. The two *Hamiltosporidium* species analyzed in our study, *H. tvaerminnensis* and *H. magnivora*, are diploid (Haag, Traunecker, et al. 2013). *Hamiltosporidium tvaerminnensis* is asexual and transmitted both horizontally and vertically, whereas *H. magnivora* is possibly sexual but only transmitted vertically from mother to offspring (Haag et al. 2011; Haag, Traunecker, et al. 2013) and possibly horizontally to a second, yet unknown, host. In contrast, vertical transmission was never observed for *Ordospora colligata*, which belongs to a derived group of microsporidia with highly reduced genomes (Pombert et al. 2015). Its ploidy and sexuality are unclear, but its closest known relatives have been found diploid, albeit with very low levels of heterozygosity, and postulated to feature a sexual cycle (Selman et al. 2013). In our comparisons, we focus on the relative roles of natural selection and genetic drift in genome evolution of microsporidian parasites with known modes of transmission.

**Materials and Methods**

**Host and Parasite Collection**

*Daphnia magna* are planktonic Crustaceans inhabiting standing fresh- and brackish-water bodies with a Holarctic distribution. In the course of diverse sampling efforts (e.g., Roulin et al. 2013; Yampolsky et al. 2014; Fields et al. 2018), *Daphnia* were brought to the laboratory either in form of planktonic samples or in form of resting eggs (=ephippia), collected from the sediment surface of the water body. Field collected planktonic females were cloned, that is, individual females were allowed to reproduce asexually. Resting eggs were washed and stimulated to hatch by exposure to continuous light under room temperature in well-oxygenated medium. Hatchlings were isolated and were allowed to produce clonal lines by asexual reproduction. All clones were kept in the laboratory under conditions of continuous asexual reproduction with standard laboratory conditions (20°C, 16:8 h light:dark cycles, green algae *Scenedesmus* sp. as food in artificial *Daphnia* medium (Ebert et al. 1998). Animals from each clone were tested for microsporidian infections. Clones with infections of *O. colligata*, *H. magnivora*, and *H. tvaerminnensis* were mass...
propagated to produce enough parasite material for DNA isolation using standard protocols.

DNA Isolation and Sequencing

Reduction of nonfocal DNA, mainly from host microbiota and food items, followed the protocol of Dukić et al. (2016). In short, infected D. magna were treated for 72 h with three antibiotics (streptomycin, tetracycline, and ampicillin). At the same time, all animals were fed with dextran beads (Sephadex beads, 50-μm diameter, Sigma Aldrich) to aid gut evacuation. Animals were then placed in 1.5-ml Eppendorf tubes and excess fluids were removed. We added extraction buffer (Qiagen GenePure DNA Isolation Kit) to the tubes and disrupted the tissue using sterile and DNA-free plastic pestles. The sample was incubated overnight with Proteinase K at 55°C, followed by RNA degradation using RNase treatment (1 h, 37°C). Protein removal and DNA precipitation, including the addition of glycogen (Qiagen) to aid DNA precipitation, were done using the Qiagen GenePure DNA Isolation Kit instructions. Resultant DNA was suspended in 40 µl of Qiagen DNA hydration solution and subsequently tested for purity and concentration using a Nanodrop and Qubit 2.0, respectively. Libraries were prepared using Kapa PCR-free kits and sequenced by the Quantitative Genomics Facility service platform at the Department of Biosystem Science and Engineering (D-BSE, ETH), in Basel, Switzerland, on an Illumina HiSeq 2000 (upgraded to 2500) with the HiSeq 2500 v4 kit in paired ends mode (2 × 126 bp [i.e., 125 bp + 1 bp for final quality score assessment]; data sets FI-SK-17-1, GB-EP-1, NO-V-7, IL-G-3) or the HiSeq 2500 Rapid Run v2 kit in paired ends mode (2 × 301 bp [300 bp + 1 bp]; data sets BE-OM-2, IL-BN-2, FI-OER-3-3).

Genome Assembly

Ordospora

Daphnia magna clones collected from populations in Finland, England, and Norway were found to be infected with O. colligata (sample codes: FI-SK-17-1, GB-EP-1, and NO-V-7). Low quality bases in the data sets were discarded with Trimmomatic 0.27 (Bolger et al. 2014) using default parameters. Host sequences were filtered out from the Ordospora data sets before the initial assemblies by read mapping with Bowtie2 2.2.9 against the D. magna (dmagna-v2.4-20100422-assembly.fna; http://arthropods.eugenes.org/EvidentialGene/daphnia/daphnia_magna/Genome/?). Contigs from the host were filtered out postassembly of two H. magnivora isolates from Belgium and Israel (BE-OM-2, IL-BN-2) and two H. tvaerminnensis isolates from Finland and Israel (FI-OER-3-3, IL-G-3). The Hamiltosporidium spp. genomes were assembled iteratively with Ray 2.3.2 using kmers 31, 41, 51, 61, 71, 81, 91, 101, and 111, and the best contiguous assemblies were selected with QUAST 4.4. Contaminants were then filtered out in successive steps. Contigs from Daphnia were first filtered by mapping the Hamiltosporidium-free BE-OM-1 data set containing sequenc- ing reads from the host and Ordospora (in very low proportion) using Bowtie2 2.2.9 as implemented in the SSRG pipeline (https://github.com/PombertLab; last accessed December 18, 2019). Contigs showing an average sequencing depth of at least 1× in the resulting coverage files were discarded with the filter_by_coverage.pl custom Perl script (all custom scripts are available on the Pombert lab GitHub page). Contigs were then filtered by GC content with filter_by_GC.pl using minimum and maximum binning values set to 35 and 60, respectively, and the contigs averaging <35% GC kept thereafter. Potential contaminants remaining in these contigs were identified by taxonomized BLASTN searches with the megablastn algorithm against the NCBI nucleotide NT database (NT) and the Taxonomy databases, and then discarded from the final assemblies. To ensure that the read mapping approach and the 35% GC criterion above did not end up removing true Hamiltosporidium contigs, taxonomized BLASTN searches were also performed independently on the full BE-OM-2 assembly.

Genome Annotation

Ordospora

Annotations from the Ordospora OC4 reference genome were transferred to the final FI-SK-17-1, GB-EP-1, and NO-V-7 assemblies with Geneious 9.1.2 (Biomatters, Auckland, New Zealand) using the “Annotate & Predict -> Transfer annotations” built-in function. Transferred annotations were manually curated with Artemis 16.0.0 (Rutherford et al. 2000).
to correct incomplete or missing features. Curated EMBL annotations were converted to TBL format with the custom Perl script EMBLtoTBL.pl, and GenBank-compatible annotations were generated from the TBL files with the NCBI tool TBL2ASN.

**Hamiltosporidium**

Initial gene prediction in the *Hamiltosporidium* assemblies were performed with the eukaryotic gene predictors GeneMark-ES 4.33 (Ter-Hovhannisyan et al. 2008) and Augustus 3.2.3 (Keller et al. 2011) as implemented in MAKER 2.31 (Holt and Yandell 2011), with Augustus independently using the built-in Encephalitozoon cuniculi gene model and *Rozella allomycis* gene model from James et al. (2013). Additional intron-less putative open reading frames were also positioned on the contigs with Prodigal 2.6.3 (Hyatt et al. 2010). In parallel, microsporidia proteins from *Rozella allomycis* (James et al. 2013) and from MicrosporidiaDB (Aurrecoechea et al. 2017) data sets EcuniculiGBM1, AalgeraePRA109, AalgeraePRA399, EaedisUSNM41457, MdaphniaUGP3, NausubeliERTm2, NausubeliERTm6, NbbombicsCQ1, NceraeaeBLR01, NparisiEERTm1, NparisiEERTm3, OcolligataOC4, PneurophiliaMK1, Slophi42_110, ThomisinUnknown, VcorneaeATCC50505, and Vculicisfloridensis were queried against the SwissProt, TREMBL, InterProScan 5.26-65.0 (Jones et al. 2014) searches and with EMBLtoPROT.pl. Protein functions were predicted using WebApollo 2.0.5 browser (Lee et al. 2013), and the annotations were curated manually from the sum of the independent searches. The curated annotations were exported in GFF3 format using WebApollo’s built-in tools, the gff3 files were split per contig with splitGFF3.pl and then converted to EML format with WebApolloGFF3toEMBL.pl, and the predicted protein sequences were exported to FASTA format with EMBLtoPROT.pl. Protein functions were predicted using InterProScan 5.26-65.0 (Jones et al. 2014) searches and BLAST homology searches against the SwissProt, TREMBL, and UniProt databases. The functions inferred from these predictions were compared with the parse_annotators.pl and curate_annotations.pl Perl scripts, and the predicted functions were annotated based on the consensus with ambiguous functions further checked using NCBI’s CDD searches (Marchler-Bauer et al. 2015). EMBL files were converted to TBL format with EMBLtoTBL.pl, and the ASN files generated with NCBI’s TBL2ASN. Miscellaneous annotation issues caused by the gene predictors (e.g., peptides shorter than 50 aa) not automatically fixed by WebApolloGFF3toEMBL.pl and detected in the TBL2ASN validations (.val) files were corrected in the EMBL files by manual curation using the check_errors_1.sh, check_errors_2.sh and check_errors_3.sh Bash scripts and Artemis 16.0.0 (Rutherford et al. 2000). Corrected EMBL files were then converted to TBL and ASN, as described above. Intron/exon junctions in Figure SX and SY were annotated and corrected manually using Artemis 16.0.0; introns (nucleotide sequence) and exons (amino acid sequence) were aligned using MAFFT v7.407 (Katoh and Standley 2013).

**Duplication Analyses**

During the manual curation phase of the *Hamiltosporidium* spp. genomes, we noticed that some of the contigs appeared duplicated to some extent in the assemblies. The extent of these duplications was evaluated by performing BLASTN homology searches of each assembly against itself, then by parsing the results with parse_BLAST_selftest.pl with thresholds of 80% and 90% nucleotide identity between candidate duplicates. *Hamiltosporidium* is known to be diploid (Haag, Traunecker, et al. 2013); therefore in order to test whether these redundant contigs were real duplicates or instead arose from allelic differences representing assembly artifacts, the sequencing reads were mapped against the assemblies with Bowtie 2.2.9 in pair-ends mode as implemented in the SSRG pipeline (Pombert lab, GitHub), then the relative sequencing depth of each contig (Cdepth) was compared with the average depth (Adepth) of the assembly with seq_depth.pl. Statistics on duplicated contigs were further evaluated by t-tests using StatPlus for Excel (AnalystSoft, Walnut, CA) assuming unequal variances.

**Repetitive Elements Analyses**

A de novo identification of repetitive elements was performed with RepeatModeler 1.0.11 as implemented in RepeatMasker (Smit et al. 2013) with default parameters. RepeatModeler created genome-specific libraries for the seven de novo assemblies generated in this study, as well as ten other assemblies downloaded from NCBI (MdaphniaeUGP3 GCA_0000760515.1, Edha_aedisV4b GCA_000230595.3, NosBomCq1v01 GCA_000838757.1, NapisbRLv01 GCA_000447185.1, NceraeaeBLR01 GCA_000182985.1, OcolligataOC4 GCA_000803265.1, EcuniculiGBM1 GCF_000091225.1, EistnesinialATCC50506 GCA_000146465.1, EhellemATCC50504 GCA_000277815.3, and Eromalae SJ2008 GCA_000280035.2). Each genomic-specific library was combined with representative transposable elements (TEs) from fungi obtained from Repbase (Bao et al. 2015). Assemblies were screened for the repetitive elements contained in the genome-specific libraries using BLAST as search engine, as implemented in RepeatMasker, with default parameters. Low complexity DNA sequences and simple repeats were not masked. Kimura distances between genome copies and TE consensus from the library were determined using RepeatLandscape (Jallabbaro 2012) on alignments included in *.align files after genome masking with RepeatMasker. The rates of transitions and transversions
were calculated on alignments and transformed to Kimura distance (Kimura 1980).

Phylogenetic Analysis

Orthologous sequences of 27 proteins were extracted from the above-mentioned genome assemblies (supplementary file 1, Supplementary Material online). Orthologs (E-value cutoff 1e-10) were confirmed by manual inspection of postalignment similarity. Each individual protein was aligned using MAFFT (Katoh and Standley 2013), concatenated, and the ambiguously aligned regions removed with Geneious 11.0.4 (Biomatters) using the “Tools -> Mask Alignment” command. The final alignment contained 13,409 sites. The maximum likelihood phylogeny was estimated using the LG model in RAxML version 8 (Stamatakis 2014) with empirically determined amino acid frequencies, and support estimated by 500 bootstrap pseudo-replicates.

Recombination Analyses

The population mutation ($\Theta$) and recombination ($\rho$) rates were assessed directly with mlRho 2.9 (Haubold et al. 2010) as follows (see mlRho.sh on GitHub for the full commands). Briefly, reads were filtered using Trimomatic 0.39 (Bolger et al. 2014) with the ILLUMINACLIP and SLIDINGWINDOW:4:28 command line switches to clip out Illumina adapters and to discard low quality bases, respectively. Reads passing the quality filters were concatenated in a single FASTQ file, then mapped against the corresponding genomes with minimap2, as implemented in get_SNPs.pl from the SSRG pipeline (https://github.com/PombertLab; last accessed December 18, 2019) with the -bam command line switch to keep the resulting BAM alignments files. The BAM alignments were indexed with samtools 1.9-50, and used as input for formatPro 0.5 and mlRho 2.9, iteratively using minimum coverage values ($\gamma$) of 4, 8, and 16.

The effect of recombination in the *Hamiltosporidium* and *Ordospora* genomes was further assessed indirectly by calculating GC-biased gene conversion (gBGC) values with phastBias from the PHAST 1.5 package (Capra et al. 2013). Orthologs within each data set were identified with OrthoFinder 2.2.6 using default parameters and single-copy orthologs were separated from multicopy genes in the OrthoFinder output with split_csv.pl. Single-copy ortholog data sets were generated with make_phast_datasets.pl from the corresponding predicted mRNAs, and the nucleotide data sets were aligned at the amino acid level with MACSE 2.01 using the multithreaded run_mace.pl Perl script. Neutral models for each alignment and gBGC values were generated and calculated with the phyloFit and phastBias programs from the PHAST package, respectively, as implemented in the Perl scripts run_phyloFit_ORDOSPORA.pl and run_phyloFit_HAMIL.pl tailored for each data set. These analyses were repeated using each isolate/species iteratively as outgroup for the gBGC calculations.

Data Availability

All custom scripts and software are available on the Pombert lab GitHub page (https://github.com/PombertLab; last accessed December 18, 2019). This project was deposited in NCBI under BioProject accession number PRJNA419750. Sequencing data sets were deposited in the NCBI Sequence Read Archive under the same accession number (PRJNA419750). The *Ordospora* genomes were deposited in GenBank under accession numbers PITH00000000 (FI-SK-17-1), PITG00000000 (GB-EP-1), and PTF00000000 (NO-V-7). The *H. magnivora* genomes were deposited under accession numbers PITU00000000 (BE-OM-2) and PixR00000000 (IL-BN-2). The *H. tvaerminnensis* genomes were deposited under accession numbers PITJ00000000 (FI-OER-3-3) and PTK00000000 (IL-G-3).
Results

Contrasting Genome Architectures of *Ordospora* and *Hamiltosporidium*

We de novo assembled and annotated seven new genomes of microsporidian parasites of *D. magna* with different modes of transmission belonging to two clades: *Ordospora* and *Hamiltosporidium*. Table 1 shows that their genomes differ by one order of magnitude in size. *Ordospora*—an exclusively horizontally transmitted parasite of the gut epithelium—bears a compact genome with less than half of the coding capacity of *Hamiltosporidium*—a parasite with horizontal and vertical transmission (mixed-mode transmission) infecting the fat tissue and ovaries of its host. Predicted proteins, on average, were not found to be larger in *Hamiltosporidium* genomes compared with those found in *Ordospora* isolates (table 1) and we identified only one clear expansion in the family of genes encoding a minichromosome maintenance (MCM) protein (data not shown). In *Hamiltosporidium*, these MCM proteins feature in-frame expansions rich in asparagine residues that split conserved MCM domain motifs. However, we could not ascertain if those expansions are inteins that are spliced post-translationally—in teins often interrupt MCM proteins in archaea and bacteria—or simply expansions in variable regions that do not disrupt the function of the protein. Although ab initio predictions with GeneMark suggest the presence of numerous and longer inteons in *Hamiltosporidium* compared with *Ordospora*, manual investigation of inteons inserted at cognate and ectopic sites (supplementary file 2, Supplementary Material online) rather suggests that the inteons that are present in *Hamiltosporidium* spp. are similarly small. In *Trachipleistophora hominis* (Heinz et al. 2012) for which RNAseq data are available, further investigation of the GeneMark predicted inteons revealed that these are not genuine inteons (Whelan et al. 2019), and we believe that it is also the case here. The main difference we observed in the genomes of the two clades is that the *Hamiltosporidium* genomes contain much longer intergenic regions, a large proportion of repetitive elements, and long segmental duplications, which are absent in *Ordospora*. Interestingly, the *Hamiltosporidium* genomes presented here vary regarding their accumulation patterns of repetitive elements. Differently from *H. magnivora* (H.m.) BE-OM-2, H.m. IL-BN-2 and *H. tvaerminnensis* (H.t.) FI-OER-3-3, which show a unimodal distribution of TE age classes, H.t. IL-G-3 is unique in having a slightly bimodal distribution, suggesting two instances of genomic bursts caused by TE expansion in the latter (supplementary file 3, Supplementary Material online). Furthermore, *Hamiltosporidium* genomes are strongly AT biased (26% GC, on average) as compared with *O. colligata* (38% GC, on average).

To investigate whether assembly artifacts due to allelic divergence generated segmental duplications, the four *Hamiltosporidium* assemblies were searched for inteons showing regions of at least 80% of sequence identity. Such highly similar segments range from about 0.9 to 59 kb (supplementary file 4, Supplementary Material online) and correspond to 5–15% of the assembled genomes (table 1). Because *Hamiltosporidium* is known to be diploid (Haag, Traunecker, et al. 2013), the duplicated segments may represent allelic versions of the same genomic regions, or genuine duplications. Therefore, the ratio of contig sequence depth to average sequencing depth (Cdepth/Adepth) from the two *H. tvaerminnensis* assemblies showing Adepth larger than 300× (Adepth = 389× for H.t. FI-OER-3-3, and 645× for H.t. IL-G-3) was used as a proxy for their “ploidy.” Sequencing depths from genuine genomic duplications should match that of the assembly average; allelic inteons should yield Cdepth/Adepth ratios near 0.5. Duplicated inteons show a skewed distribution toward a lower Cdepth/Adepth ratios (supplementary file 5, Supplementary Material online); their mean ratio significantly differs from nonduplicated inteons (unequal variance t-test, P = 0) by about 0.3 in both assemblies (supplementary file 3, Supplementary Material online), suggesting that at least some segmental duplications may in fact represent alleles, not duplications.

Genomic Features Associated with the Mode of Transmission of Microsporidia

Our hypothesis to explain the divergent genome architectures in these two *D. magna* parasite taxa is that vertical transmission causes population bottlenecks and thus reduces Ne, and consequently the power of natural selection. If this hypothesis is correct, phylogenetically unrelated microsporidia that exploit similar routes of transmission should evolve similar genome architectures. Therefore, a phylogeny was built including *Hamiltosporidium* spp., *O. colligata*, as well as nine other microsporidian species with known genomes and modes of transmission (fig. 1). The tree contains a subset of some highly divergent microsporidia and is in agreement with previously published microsporidian phylogenies built with a larger number of species using different molecular markers (Vossbrinck and Debrunner-Vossbrinck 2005; Pombert et al. 2015). Horizontal transmission is an ancestral feature in our phylogeny and associated with a high proportion of protein-coding sequences within genomes. In contrast, the distantly related microsporidia with vertical transmission (in addition to horizontal transmission) show enlarged, less dense genomes with a reduced proportion of protein-encoding sequences (fig. 1). *Nosema* and *Hamiltosporidium*, in particular, are both characterized by the accumulation of repetitive elements. The compact genomes of the exclusively horizontally transmitted microsporidia, on the other hand, are largely devoid of repetitive sequences.

If the accumulation of noncoding and repetitive elements reflects the lack of power of purifying selection due to genetic
drift, then vertically transmitted species are expected to show the accumulation of mildly deleterious mutations in their protein-encoding genes. To test for the differential strength of purifying selection in microsporidia with distinct modes of transmission, genome-wide estimates of the ratio of nonsynonymous versus synonymous substitutions (d_{NS}/d_S) were obtained. We found a 4-fold increase in average d_{NS}/d_S ratios of single-copy orthologous genes from species with mixed-mode transmission (HN data set) in relation to species with horizontal transmission only (OE data set; fig. 2A and supplementary file 6, Supplementary Material online). The difference is less clear, and not statistically significant, for multicity genes (fig. 2B and supplementary file 6, Supplementary Material online). However, these multicity genes include cases where “paralogous” copies might represent alleles. There is large variation in d_{NS}/d_S estimates of genes encoding isoforms of similar proteins within genomes, reaching extreme values in the HN group with mixed-mode transmission (see supplementary file 6, Supplementary Material online). In order to control for the large variation in evolutionary distances between taxa and to account for differences in behavior of subsets of genes within our data set, we plotted the distribution of d_N versus d_S (supplementary file 7, Supplementary Material online) for a sample of microsporidian genomes from the OE and HN data sets with comparable d_S estimates. Because synonymous substitutions are considered neutral, d_S can be used as a proxy for their evolutionary distance. For microsporidia relying exclusively on horizontal transmission (OE data set), d_N distributions are flat, independently of the evolutionary distance (fig. 3), with regression slopes of 0.02–0.2 and intercept around 0.0 (fig. 3), whereas for those with mixed-mode transmission (HN data set), the regression slopes are much larger (0.4–0.65, and intercepts around 0.0), but only at shorter evolutionary distances (top graphs in fig. 4). For the highly divergent genomes of Nosema spp. (bottom graphs in fig. 4), the regression slopes are equivalent to those obtained for the OE data set, but the intercepts are much larger, suggesting that d_N might saturate at high d_S. Furthermore, the large d_N versus d_S correlation coefficients of the Hamiltosporidium spp. comparisons indicate that the excess of nonsynonymous substitutions in vertically transmitted microsporidia are not biased by small subset of genes with extremely high d_N, but instead represent a genomic pattern.

Reduced Levels of Recombination in Hamiltosporidium spp.

Accumulation of nonsynonymous substitutions as well as assembly artifacts caused by allelic divergence within Hamiltosporidium genomes could result from the lack of recombination (Muller 1964; Welch and Meselson 2000). Assuming that each Hamiltosporidium isolate corresponds to a single diploid individual, we calculated the recombination rate \( \rho \) for genomic regions considering three different levels of coverage (\( c = 4.8 \) and 16; supplementary file 8, Supplementary Material online). Recombination estimates are slightly above the sequencing error rate, averaging 158 events per 100 kb. Although we also applied the same approach to Ordospora spp., the zygosity correlation \( \theta \) could not be calculated—in line with our current expectations that these species are...
either haploids or show very low levels of heterozygosity like their Encephalitozoon relatives (Selman et al. 2013)—and as such the values reported for Ordospora spp. in supplementary file 8, Supplementary Material online (1,114 events per 100 kb on average) should only be considered with caution. To enable a comparison between Hamiltosporidium and Ordospora, we assessed recombination indirectly, searching for genomic regions with increased GC content (gBGC tracts). In the absence of recombination, gBGC should occur at greatly reduced rates, eliminating such regions. We employed a Bayesian approach to identify gBGC tracts within all orthologous single-copy genes of the seven new genome assemblies plus the previously published Ordospora genome; 1,227 genes for Hamiltosporidium spp. and 1,748 for O. colligata). Overall, we found 141 gBGC tracts with probability >0.5, ranging from 10 to 2,168 bp in the four Hamiltosporidium genomes, but none for O. colligata (supplementary file 9, Supplementary Material online). The vast majority (111) of those tracts were found in H.m. BE-OM-2, whereas only 12, 10, and 8 tracts were found in H.t. FI-OER-3-3, H.t. IL-G-3, and H.m. IL-BN-2, respectively. Overall, these results suggest that some form of recombination (either mitotic or meiotic) occurs in Hamiltosporidium, favoring the hypothesis that the excess of nonsynonymous substitutions observed in the vertically transmitted microsporidia results from genetic drift, and not from the lack of recombination.

Discussion

Exploring the genomic landscape of seven new microsporidian genomes, we found several differences between Hamiltosporidium and Ordospora taxa: The former have more protein-coding genes, a larger proportion of repetitive sequences relative to the total assembled sequence, longer intergenic regions, and reduced GC content. These differences are associated with a 10-fold larger genome size of Hamiltosporidium compared with Ordospora, suggesting a possible link between the genomic features and genome transmission modes.
Microsporidian genome size variation has remained elusive due to insufficient genomic data from species with sufficiently well described life histories. The two taxa sequenced here are good models for studying the mechanisms shaping genome evolution, given that our knowledge about their ecology, evolution, and epidemiology is better than for other microsporidia (Ebert et al. 2000, 2001; Ebert 2005; Lass and Ebert 2006; Sheikh-Jabbari et al. 2014; Urca and Ben-Ami 2018; Kirk et al. 2019).

**Natural Selection for Smaller or Larger Coding Capacity**

It is suggested that the divergence of microsporidia from their fungal relatives coincided with a dramatic bottleneck that resulted in the evolutionary loss of gene families, and proceeded with the gain and subsequent adaptive enlargement of other gene families, resulting in an expanded core proteome (Nakjang et al. 2013). Sharp contrasts in microsporidian genomic architectures are proposed to be derived from divergent factors causing differential selection and being associated with variable degrees of host specificity (Desjardins et al. 2015), such that genomes from microsporidia that use a range of different host species would be selected for having a larger and more flexible protein repertoire than microsporidia restricted to a single host species. However, neither gene family expansions nor host adaptation seems satisfying explanations for their genome size variation, as much of the variation remains unexplained. For example, all encephalitozoaans have extremely reduced genomes of about 2.5 Mb, but their host specificities vary. Furthermore, the increase in genome size in

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**Fig. 2.**—Box plots of mean dNdS estimates (A) for all orthologous genes identified in OE and HN data sets, based on 112,055 and 8,846 pairwise comparisons, respectively and (B) for all copies of similar genes within genomes based on 7,790 comparisons of data set OE and 64,437 of data set HN. The statistical significance of the difference between dNdS means was assessed by unequal variance t-tests (**P < 0.01; NS = not significant; supplementary file 4, Supplementary Material online).
Fig. 3.—Linear regression analyses of dN versus dS estimates for all orthologous genes identified in a subset of comparisons from the OE data set (see supplementary file 7, Supplementary Material online, for details). The regression equations describing the best fit of the dN versus dS distributions are given. Correlation coefficients are indicated. Oc = O. colligata; Eh = E. hellem; Ec = E. cuniculi.
Linear regression analyses of $d_N$ versus $d_S$ estimates for all orthologous genes identified in a subset of comparisons from the HN data set (see supplementary file 7, Supplementary Material online, for details). The regression equations describing the best fit of the $d_N$ versus $d_S$ distributions are given. Correlation coefficients are indicated. Hm = H. magnivora; Ht = H. tvaerminnensis; Nb = N. bombycis; Na = N. apis; Nc = N. ceranae.
microsporidia is associated with a proportional increase in noncoding sequence, whereas the associated increase of the proteome is much smaller. The here studied microsporidia are highly host specific, with *D. magna* being the only known host (Green 1974; Ebert et al. 2001; Ebert 2005). Still *Hamiltosporidium* spp. and *O. coligata* genomes differ by a factor of 10, whereas their proteomes only differ by a factor of 2–3 (table 1). Perhaps a valid generalization on microsporidian genomes is that there are those that we might call large and “gene sparse,” whereas others are smaller and “gene dense” (Keeling et al. 2014). Taking into account that the “gene sparse” genomes are scattered across the microsporidian phylogeny (see Pombert et al. 2015), and if natural selection would be the only mechanism shaping genome architecture, then the accumulation of noncoding and potentially deleterious sequences would remain unexplained.

Accumulation of Selfish TEs

It is believed that the wide variation in genome size observed among eukaryotic species is more closely correlated with the amount of repetitive DNA than with the number of coding genes (Kidwell and Lisch 2000). In microsporidia, the smallest encephalitozoan genomes are devoid of TEs, and large genomes, such as from *Anncalia algerae* (23 Mb), contain hundreds of copies of different families of TEs (Parsot et al. 2014). Similarly, the *Hamiltosporidium* genomes (17–25 Mb) that we have sequenced contain a large proportion of repetitive elements, some of which identified as known TE families. The age-distribution of different TEs in *Hamiltosporidium* genomes suggests the occurrence of cyclical TE expansions; the largest genome of strain H.t. IL-G-3 shows two peaks in the abundance of TEs from distinct age classes (supplementary file 3, Supplementary Material online). However, the largest known microsporidian genome, from the mosquito parasite *Edhazardia aedis* (assembly size of 51.34 Mb; fig. 1), is rather poor in TEs, but enriched with intergenic AT-biased sequences (Desjardins et al. 2015). This finding is not in direct support of a simple relationship between total TE-derived DNA and microsporidian genome size.

Reduced Recombination and the Accumulation of TEs

Reduced recombination has been associated with TE accumulation and larger genomes (Tiley and Burleigh 2015). Two potential mechanisms for a negative association between recombination rate and genome size are that recombination either deletes TEs by chance or facilitates selection against TE insertions (Langley et al. 1988). However, recent evidence suggests that TEs might actively contribute to the spread of recombination suppression (Kent et al. 2017). Nevertheless, the absence of recombination reduces effectiveness of natural selection on short and long evolutionary timescales and is predicted to be associated with reduced GC content because gBGC does not occur (Capra et al. 2013). Searching our assemblies for regions suggestive of gBGC and by estimating recombination rates directly, we detected recombination in *Hamiltosporidium*, though at very low rates (0.0006–0.00158; supplementary file 8, Supplementary Material online). These results are lower than recombination estimates obtained for other asexual fungi such as *Candida glabrata* (0.003–0.008, Carreté et al. 2018), which may engage in parasexual processes that allow them to reshuffle DNA, and which have been postulated to achieve gBGC by mitotic recombination (Marsolier-Kergoat 2013). In contrast, recombination rates calculated for *Ordospora* spp. were found higher by a 10-fold, congruent with the idea that the *Hamiltosporidium* spp. underwent fewer recombination events. Overall, the apparent lower recombination rates in *Hamiltosporidium* spp. are compatible with the idea that these genomes acquired TE at least in part due to reduced recombination pressure.

Genetic Drift Caused by Vertical Transmission

A general model of genome evolution must account for the accumulation of neutral and maladaptive sequences, as well as their elimination. The two microsporidian taxa that we compared are specific to *D. magna* but differ in their modes of transmission. Mixed-mode transmitted *Hamiltosporidium* spp. show larger and “gene sparse” genomes containing duplicated segments and other repetitive elements. These are common features of eukaryotic genomes that have been affected by genetic drift. Mixed-mode transmission is a strategy used by many microsporidia (Dunn et al. 2001), enabling them to achieve very high prevalence at equilibrium (Lipsitch et al. 1995; Lass and Ebert 2006). Vertical transmission in addition to horizontal transmission involves a fitness tradeoff (Vízoso and Ebert 2005) but is probably an important survival strategy when opportunities to transmit horizontally are temporarily reduced or absent, for example, under low host density and during a phase of adverse environmental conditions (Lucarotti and Andreidis 1995; Ebert 2013; Sheikh-Jabbari et al. 2014). Vertical transmission under such conditions may cause transmission bottlenecks and thus reduces *N_e*. As a consequence, the strength of natural selection is reduced, because genetic drift adds stochastic noise to the fate of variants.

In microsporidia with mixed-mode transmission, two morphologically different spore types are often observed, and vertically transmitted spores resemble those at early stages of development, with a shorter polar filament and a thinner endospore wall (Dunn et al. 2001). Horizontally transmitted spores, on the other hand, show a thick cell wall and are produced in large numbers, creating conditions for multiple infections of a single host with a diversity of parasite genotypes (Dunn and Smith 2001). This is less likely with vertical transmission, because only a small proportion of spores translocate into reproductive cells. *Hamiltosporidium*
*tvaerminnensis* produces two types of spores with different shapes, but they do not seem to differ with regard to their roles in horizontal transmission (Urca and Ben-Ami 2018). Although the ratio of vertical to horizontal transmission is not known for microsporidia with mixed-mode transmission, studies using microsatellite polymorphisms in *Hamiltosporidium* suggest that multiple infections are rare (Haag, Sheikh-Jabbari, et al. 2013). Once the *Daphnia* host is born vertically infected, the chances of becoming horizontally reinfected by a different *Hamiltosporidium* genotype may be low. Theory predicts that the tradeoff between horizontal and vertical transmission is controlled by the proportion of available uninfected hosts, with vertical transmission being selectively favored when uninfected hosts are rare (Turner et al. 1998). In Finland, where *H. tvaerminnensis* is common in a *D. magna* metapopulation, parasite prevalence is cyclical, reaching up to 100% during summer, and decreasing every year after winter and summer host diapausas (Lass and Ebert 2006). Hence, it is reasonable to assume that vertical transmission is favored when *H. tvaerminnensis* prevalence is high, thus reducing *N_v* and facilitating the accumulation of neutral and mildly deleterious mutations by genetic drift.

**Accumulation of Mildly Deleterious Mutations**

To further investigate the hypothesis that microsporidian genomes expand through a reduced effectiveness of purifying selection, we compared the ratio of nonsynonymous versus synonymous substitutions in the microsporidian genomes. We found that mixed-mode transmission, and larger genomes, are associated with significantly larger \( d_{SNP}/d_{S} \) ratios between single-copy orthologous genes of different genomes (fig. 2A). We speculate that the excess of nonsynonymous substitutions in orthologous genes results from the accumulation of mildly deleterious mutations due to genetic drift. Variable \( d_{SNP}/d_S \) ratios among multiple copy sequences (fig. 2B) indicate variable functional constraints (Ohta 1992). Thus, it seems that the reduced power of natural selection associated with the addition of vertical transmission, may contribute to the accumulation of noncoding sequences, TE, and duplicated genes, leading to microsporidian genome expansions. In the long term, the opposite outcome occurs in exclusively vertically transmitted bacterial endosymbionts, which have extremely miniaturized genomes (Mira and Moran 2002; Moran and Bennett 2014). Genome miniaturization associated with genetic drift in endosymbiotic bacteria is believed to be caused by a mutational bias toward deletions (Mira et al. 2001; Kuo and Ochman 2009).

**Relative Roles of Reduced Recombination and Vertical Transmission**

Accumulation of mildly deleterious mutations might be generated both by reduced recombination via Muller’s ratchet, or by reduced \( N_v \) due to recurrent population bottlenecks.
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