Rapid Increase in Genome Size as a Consequence of Transposable Element Hyperactivity in Wood-White (Leptidea) Butterflies

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Abstract
Characterizing and quantifying genome size variation among organisms and understanding if genome size evolves as a consequence of adaptive or stochastic processes have been long-standing goals in evolutionary biology. Here, we investigate genome size variation and association with transposable elements (TEs) across lepidopteran lineages using a novel genome assembly of the common wood-white (Leptidea sinapis) and population re-sequencing data from both L. sinapis and the closely related L. reali and L. juvernica together with 12 previously available lepidopteran genome assemblies. A phylogenetic analysis confirms established relationships among species, but identifies previously unknown intraspecific structure within Leptidea lineages. The genome assembly of L. sinapis is one of the largest of any lepidopteran taxon so far (643 Mb) and genome size is correlated with abundance of TEs, both in Lepidoptera in general and within Leptidea where L. juvernica from Kazakhstan has considerably larger genome size than any other Leptidea population. Specific TE subclasses have been active in different Lepidoptera lineages with a pronounced expansion of predominantly LINEs, DNA elements, and unclassified TEs in the Leptidea lineage after the split from other Pieridae. The rate of genome expansion in Leptidea in general has been in the range of four Mb/Million year (My), with an increase in a particular L. juvernica population to 72 Mb/My. The considerable differences in accumulation rates of specific TE classes in different lineages indicate that TE activity plays a major role in genome size evolution in butterflies and moths.

Key words: butterfly, Lepidoptera, Leptidea, genome expansion, transposable elements, population.

Introduction
Causes and consequences of variation in genome size across taxa have been a matter of debate in the field of evolutionary biology (Petrov 2001; Cavalier-Smith 2005; Gregory 2005; Lynch 2007; Oliver et al. 2007). The observations that variation in genome size in extant species is on the level of ten- to hundred thousand folds (Gregory 2004) and that genome size correlates only weakly with the amount of coding sequence and the organism complexity (the C-value paradox; Thomas Jr. 1971) have rendered a lot of attention and attempts to understand the underlying mechanisms generating this variation (Petrov 2001; Cavalier-Smith 2005; Gregory 2005; Lynch 2007; Oliver et al. 2007; Fontdevila 2011). Key mutation classes that contribute to genome expansion are predominantly transposable element (TE) proliferations (e.g., Pritham 2009; Tenaillon et al. 2010), gene- (e.g., Ohno 1970; Lu et al. 2012) or genome duplications (Fontdevila 2011) and replication slippage of tandem repeat sequences (Ellegren 2004). Quantification of the rate and prevalence of these mutation types is essential to get a comprehensive understanding of the forces that shape genome size evolution in different lineages. Two major lines of argument have been put forward to...
explain the variation in genome size. In essence, these can be sorted under adaptive processes on the one hand (Gregory and Hebert 1999; Cavalier-Smith 2005; Gregory 2005), and stochastic gain or loss of genomic regions on the other (Petrov 2001, 2002a, b; Lynch 2007). The former make a case that addition of DNA can influence cell size and rate of cell division, which might have an effect on organism development, or that selection for increased cell size drives genome expansion to increase stability and/or enhance molecular transport from nucleus to cytoplasm (Cavalier-Smith 2005). Non-adaptive models rather focus on mutation and fixation biases of insertions and deletions as a consequence of random genetic drift driving genome expansion or contraction (Petrov 2001; Lynch 2007)—for example, that genome expansion is mainly driven by proliferation of TEs, ‘selfish-DNA’ that either copy and paste themselves passing through an RNA intermediary step (retrotransposons), or cut and paste themselves (DNA transposons) within the genome of the host organism (Orgel and Crick 1980; Kazazian 2004; Kidwell 2005). The role of TE proliferation for genome size variation is supported by the universal existence of TEs in virtually all eukaryotic genomes (Elliott and Gregory 2015) and genomic gigantism in salamanders is for example a result of hyperactivity of specific long terminal repeats (LTRs) (Sun et al. 2012). However, also amounts of non-repetitive DNA (non-coding, non-TE) tend to co-vary with genome size indicating that non-TE insertion/deletion forces might play an important role in genome size variation (Lynch 2007). Prokaryotes have compact genomes almost entirely consisting of protein coding genes, while the amount of non-coding DNA is higher in unicellular eukaryotes and constitutes the major part of the genome in multicellular eukaryotes (Lynch 2007). This has been attributed to be a consequence of an increased cost of replication of excess DNA that is more efficiently selected against in prokaryotes and unicellular eukaryotes that generally have very large effective population sizes, although there are currently no data at hand that directly verify this assumption (Lynch 2007). Interestingly, TE insertions have also been shown to be hindered by epigenetic mechanisms underlying inhibition of homologous recombination in prokaryotes (Fedoroff 2012).

In insects, observed genome size variation ranges from 0.09 to 16 Gb (Gregory et al. 2007; Hanrahan and Johnston 2011; Maumus et al. 2015), but sampling is sparse and sporadic with low representation of particular taxonomic groups, and there is indecisive support for natural selection on the one hand (Arnaqvist et al. 2015), and/or genetic drift on the other (Lefèbure et al. 2017) in driving genome size evolution (Sessegolo et al. 2016). Within the order Lepidoptera (butterflies and moths) there is considerable genome size variation (up to 7-fold) but, although these insects are among the most diverse groups of invertebrates, so far very few species have been studied in detail (Gregory and Hebert 2003). To investigate the impact of TEs on genome size variation in this understudied group of organisms, we here quantify the abundance of different classes of TEs across species and populations with considerable variation in genome sizes. We then use a phylogenetic approach to estimate genome expansion rates as a consequence of bursts of TE proliferation. Our main study system consists of three cryptic species of butterflies from the Eurasian genus Leptidea (Pieridae), namely the wood-whites Leptidea sinapis, L. reali, and L. juvernica. This triplet represents one of the most striking examples of cryptic species in Eurasian butterflies (Dincă et al. 2011, 2013) and has been widely studied in terms of ecology (e.g., Friberg and Wiklund 2010; Friberg et al. 2011), behavior (e.g. Wiklund 1977; Friberg and Wiklund 2007) and chromosome evolution (e.g., Lukhtanov et al. 2011; Šichová et al. 2016). The results from these studies call for a better understanding of the genomic particularities of Leptidea, but there are very limited data on genome architecture and genetic variation within and between species, both in this genus and as compared with other lepidopterans. Preliminary data suggest that there is considerable nucleus size differences between species in the Leptidea genus (Šichová et al. 2015), potentially indicating that there is substantial genome size variation between the species despite recent estimated divergence times (Dincă et al. 2011). To quantify the presence and accumulation of TEs, we assessed the interspersed repeat content of six populations from three cryptic Leptidea species and compared the results with lepidopteran taxa with previously available genome assemblies. Our results indicate that genome expansion can occur rather quickly (~70 Mb/Million years [My]) as a result of activity of specific repeat elements in certain lineages, potentially as a consequence of increased rate of TE proliferation during periods with reduced effective population size.

**Materials and Methods**

**Genome Assembly**

Offspring from one mated L. sinapis female collected in Sweden during June 2013 were recurrently inter-crossed to generate a five-generation full-sib inbred line. Larvae of the fifth generation were kept until the ultimate (fifth) instar before being harvested for DNA extraction. Genomic DNA was extracted using a standard phenol–chloroform protocol (Sambrook et al. 1989) from three full siblings from the fifth generation. Illumina paired-end libraries with insert sizes 180 and 650 bp were generated using genomic DNA obtained from one of the sampled individuals. Two Illumina mate-pair libraries with insert sizes 4 and 8 kb, respectively, were generated using the remaining two individuals (one for each library). All libraries were sequenced to deep coverage (>150X paired-end coverage, >100X mate-pair coverage) using Illumina HiSeq 2500 technology (Illumina, Inc., San Diego, USA) with 125 bp read length. The sequences were assembled using ALLPATHS-LG (Butler et al. 2008) to obtain a draft reference genome assembly that was 650 Mb in total length.
and consisted of 7,096 scaffolds. The draft assembly was then screened for potential contaminants using the NCBI BLAST database (Altschul et al. 1990). Five scaffolds were identified as potential contaminants based on high sequence similarity (e-value < e−10) to non-lepidopteran taxa and deviating overall GC content and were removed from the assembly. The removed scaffolds were in total 134,034 bp long with four shorter (2 kb each) and one 128 kb long. The longest and three of the shorter removed scaffolds were likely remaining host plant material since they had high sequence similarity to plant species Trifolium pratense and Lotus japonicus (Lepidoptera butterfly larvae feed on plants from the pea family, Fabaceae). One of the three shorter removed scaffolds was 100% identical to a plasmid (suicide vector pCD-Ras1). We assembled the L. sinapis mtDNA genome by mapping all L. sinapis reads to the previously available mitochondrial genome of the close relative Leptidea amurensis (Hao et al. 2014; Šíchová et al. 2016) using BWA (Li and Durbin 2010), and assembling the reads using ABySS (Simpson et al. 2009). The process was repeated once with the de novo assembled L. sinapis mtDNA genome as reference for mapping. The final mtDNA assembly matched one scaffold from the genome assembly to 100%. The mtDNA assembly was 15,171 bp long and was annotated using the Mitos web server (Bernt et al. 2013). After removing likely contaminant sequences and the mtDNA-derived scaffold, the final nuclear genome assembly used for downstream analysis consisted of 7,090 scaffolds spanning in total 643 Mb. The scaffold N50 was 857.2 kb and 95% of the assembly was covered by 1,083 scaffolds. The quality of the assembly was further assessed by reciprocal similarity searches of scaffolds with BLAST (Altschul et al. 1990) and to identify potentially false duplicated regions we screened the assembly for conserved genes using BUSCO (Challis et al. 2017). The rationale behind this was to visually inspect the distribution of k-mers to assess completeness, heterozygosity and potential collapsed regions in the genome assembly. To get an additional and independent estimate of the genome size in L. sinapis, we also generated a Supernova genome assembly (Weisenfeld et al. 2017) from 10X Genomics Chromium linked-read data. Paired-reads of 150 bp were sequenced on the Illumina HiSeq X technology (Illumina, Inc., San Diego, USA) using a Chromium library with maximized insert sizes from the same DNA as used for paired-end libraries of the Allpaths-LG assembly. The size of the Supernova assembly was the same as the ALLPATHS-LG assembly, further supporting that the genome size of L. sinapis is in the range of 650 Mb. The genome statistics and comparisons with other previously available butterfly and moth genomes are provided in table 1 and the comparison of the ALLPATHS-LG and Supernova assemblies are presented in supplementary table 1 and figure 1, Supplementary Material online [both assemblies are available at the European Nucleotide Archive (ENA) under project accession numbers ERZ468508 and ERS1830260, respectively].

Population Sampling and Re-Sequencing

Ten individuals from each of six different populations [L. sinapis central Sweden (LsSwe), L. sinapis northern Spain (LsSpa), L. sinapis eastern Kazakhstan (LsKaz), L. reali northern Spain (LrSpa), L. juvernica eastern Kazakhstan (LjKaz), and L. juvernica Ireland (LjIre)] were sampled in the field during seasons 2013–2015. DNA was extracted from head and thorax combined using standard phenol–chloroform procedures (Sambrook et al. 1989). Each sample was prepared for sequencing by generating individually barcoded, 380-bp paired-end Illumina libraries. Samples were multiplexed and sequenced using Illumina HiSeq technology (Illumina, Inc., San Diego, USA). Library preparations and sequencing were done by the SNP&SEQ Technology Platform at the Science for Life Laboratory (SciLife, Stockholm and Uppsala). Sequencing was performed twice, the first run included LsSpa, LsKaz, LrSpa, and LjKaz and was run on the Illumina HiSeq 2000 instrument (100 bp read length), multiplexing on two separate lanes (20 samples per lane). The second bout of sequencing was performed using updated chemistry (125 bp read length) and included all populations; LsSpa, LsKaz, LrSpa, and LjKaz were multiplexed on two lanes (20 samples per lane) and LsSwe and LjIre were multiplexed on two lanes (10 samples per lane) and ran on the Illumina HiSeq 2500 instrument.

<table>
<thead>
<tr>
<th>Species</th>
<th>Assembly</th>
<th>#Scaff</th>
<th>N50</th>
<th>LS</th>
<th>GC</th>
<th>BUSCO</th>
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<tr>
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<tr>
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<td>2.1</td>
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<td>33.1</td>
<td>55</td>
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<tr>
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<td>0.857</td>
<td>6.9</td>
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</table>

NOTE.—Assembly size (Assembly), N50 for scaffolds (N50) and longest scaffold (LS) are given in Mb, and GC-content (GC) and fraction of completely covered conserved arthropod genes (BUSCO) are given in %.

*The genome assemblies for these species were downloaded from the web repository: http://prodata.swmed.edu/LepDB/, last accessed June 9, 2017 (Cong et al. 2016; Shen et al. 2016).
final sequencing reads obtained for each individual were trimmed for remaining adapter sequences and low quality bases (Q-score < 30) using the tool Cutadapt (Martin 2011). The overall coverage of quality filtered reads was estimated to be 8–15X per individual given a *L. sinapis* reference genome size of 650 Mb (see above). All sequence reads for genome assembly and population re-sequencing have been deposited at the ENA under project accession number PRJEB21838 (for details see supplementary tables 1 and 2, Supplementary Material online).

### K-mer-Based Genome Size Estimates

To estimate the approximate genome size of each species/population, mean sequencing depth was estimated for each individual using the K-mer counting tool JELLYFISH (Marcais and Kingsford 2011). Histograms were generated using the “-histo” command in JELLYFISH to identify the distinct K-mer peak by plotting multiplicity of unique K-mers versus the total number of K-mers (cf. Guo et al. 2015). All uneven K-mer lengths from 15 to 35 were assessed and the optimal K-mer length was estimated to be 17 (most distinct peaks). Sequencing reads were then randomly sub-sampled to the coverage of 5X for each individual to reduce potential coverage bias on the genome size estimate. The randomly sampled reads were used to generate K-mer histograms (multiplicity of unique K-mers with size 17 vs. the total number of K-mers) using JELLYFISH (Marcais and Kingsford 2011). The genome size of each individual was estimated by dividing the total number of nucleotides in the read set with the estimated coverage. This procedure has previously been used to estimate genome size of, for example, *Bemisia tabaci* and the conclusion has been that K-mer-based estimates are inflated compared with what was expected from flow cytometry (Guo et al. 2015). To correct for this bias, we scaled the K-mer-based estimates for each sample with the genome size information we have from our *L. sinapis* reference genome (see above). All in-house developed scripts for the genome size...
Phylogenetic Reconstruction

Genome assemblies of ten butterfly and moth species (Bombyx mori, Calycopsis cecrops, Danaus plexippus, Heliconius melpomene, Lerema accius, Manduca sexta, Melitaea cinxia, Papilio glaucus, Papilio polytes, and Papilio xuthus) were obtained from the Lepbase database (http://www.lepbase.org; last accessed June 9, 2017, Challis et al. 2017) and the genome sequences of two additional species, Phoebis sennae and Pieris rapae, were obtained from the web repository: http://prodata.swmed.edu/LepDB/, last accessed June 6, 2017, (Cong et al. 2016; Shen et al. 2016). Our L. sinapis reference genome was used for the Swedish L. sinapis population and whole-genome consensus sequences were generated for the individuals with the highest coverage from each of the remaining five Leptidea populations using the “mpileup” command in SAMtools (Li et al. 2009). The set of 18 genome sequences was screened for core arthropod genes using BUSCO v.1.22 (Simao et al. 2015). We found 224 complete single copy genes common to all the 18 scanned genomes in the arthropod gene set. The protein sequences for each of these genes were obtained for each species and aligned using MAFFT (Katoh and Standley 2013). A global phylogeny was generated using RAxML v.8.2.4 (Stamatakis 2006) with the amino acid substitution model “PROTGAMMAGR” to allow for rate variation among sites and calculate the most accurate likelihood scores (Izquierdo-Carrasco et al. 2011), and node support was estimated using 10,000 bootstrap iterations.

Estimating Divergence Times

Protein coding nucleotide sequences were obtained using genomic coordinates for the previously identified 224 common single-copy genes and the sequences were aligned using PRANK (Löytynoja and Goldman 2005). The alignments were corrected by taking codon constraints into account using MACSE (Ranvez et al. 2011). The corrected alignments were used to estimate the divergence time between the focal (Leptidea) group and the most closely related taxa in the previously available genome dataset (P. sennae, P. rapae; Pieridae), and the divergence time between clades within the Leptidea species set using BEAST v.1.7.5 (Drummond et al. 2012). Prior probabilities were set using previously available divergence time estimates (Hedges and Kumar 2009). The prior for the time to most recent common ancestor (TMRCA) of families Papilionidae and Nymphalidae was set at 122 million years (My) with a standard deviation of 12 My (Espeland et al. 2015), the priors for the internal TMRCA s within these families were set to 31 ± 10 and 103 ± 12 My, respectively (Hedges and Kumar 2009; Wahlberg et al. 2009; Nazari et al. 2011; Condamine et al. 2012), and the prior for TMRCA of Pieridae was set to 79 ± 12 My (Heikilä et al. 2012). All priors were set to represent normal distributions with a log-normal relaxed clock and multiple hits were corrected for using the GTR substitution model as implemented in the software. The phylogeny established using RAxML (see above) was given as a guide tree. The divergence time estimates for Leptidea—Phoebis/Pieris and for LjKaz—Ljre were used to estimate genome expansion rates in Leptidea in general after the split from the other Pieridae butterflies (Phoebis and Pieris), and in LjKaz after the split from Ljre.

RepeatModeler and RepeatMasker Analyses

We conducted RepeatModeler version 1.0.8 (Smit and Hubley 2010) de-novo predictions of repetitive elements in each of the following 12 lepidopteran genome assemblies: B. mori, C. cecrops, D. plexippus, H. melpomene, L. accius, M. sexta, M. cinxia, P. glaucus, P. polytes, P. rapae, P. xuthus, and P. sennae. We merged the resultant raw libraries with curated in-depth repeat annotations of H. melpomene (Lavioie et al. 2013), Heliconius erato (Ray DA, unpublished data), and all Hexapoda repeats available in Repbase (Bao et al. 2015). Redundancies between the eleven RepeatModeler libraries and the existing curated libraries were removed using the ReannTE_mergeFasta.pl script available via https://github.com/4ureliek/ReannTE/, last accessed June 12, 2017 (Kapusta et al. 2017) while giving priority to retaining curated consensus sequences. We then annotated all sampled lepidopteran genome assemblies via RepeatMasker version 4.0.6 with the “ncbi” search algorithm (Smit et al. 1996–2010) using this specific library (supplementary data file 1, Supplementary Material online). Landscapes of relative TE activity were generated using the calcDivergenceFromAlign.pl and createRepeatLandscape.pl scripts of the RepeatMasker packages. Ages of TE copies were inferred by dividing each TE copy’s Kimura 2-parameter distance to respective TE consensus by the neutral substitution rate. Lineage-specific neutral substitution rates were based on 4-fold degenerate sites in 224 coding genes that were also used to reconstruct the phylogenetic relationship between all taxa in the study (see...
Repeat proliferation rates within lineages. To estimate within-population differences in TE abundances, we analyzed the Leptidea spp. re-sequencing data using dnaPipeTE (Goubert et al. 2015). DnaPipeTE performs de novo assembly of TEs from a low-coverage subsample of re-sequencing reads in Trinity (Grabherr et al. 2011), followed by automatic quantification and annotation of TEs in the re-sequencing reads together with Repeatbase data. DnaPipeTE thus allows the quantification of recently active repeat elements in re-sequencing data, unlike RepeatMasker which quantifies repeats with a wide range of ages across genome assemblies. To optimize the amount of re-sequencing data for dnaPipeTE subsampling of each population, we selected one individual per Leptidea spp. population and ran dnaPipeTE on subsamples ranging between 200,000 and 1,200,000 reads in intervals of 100,000 reads (11 runs). For each of the 11 runs per individual, we selected the subsample yielding the highest contig N50 metric in the Trinity assembly step of dnaPipeTE, as a measure of optimized read subsampling. The optimized read subsample (LsSwe = 500,000 reads, LsKaz = 400,000 reads, LsSpa = 500,000 reads, LrSpa = 600,000 reads, Ljre = 300,000 reads, LjKaz = 700,000 reads) was then used to run dnaPipeTE on the remaining nine individuals of each population, respectively. Similar to the aforementioned RepeatMasker annotations, TE families were grouped into TE subclasses "DNA elements", "SINEs", "LINEs", "LTRs", and "Unclassified".

To get time estimates of variation in TE activity over the course of butterfly divergence, the divergence levels between repeats within each class were time scaled using neutral mutation rate estimates (2.9 × 10⁻⁹ mutations per site per generation) from H. melpomene (Keightley et al. 2015), assuming one generation per year in general. Current generation times vary considerably across lepidopteran taxa (Boggs et al. 2003) and many species show regional variation in voltinism dependent on climatic conditions. Within Leptidea for example, populations inhabiting the regions in central and southern Europe are usually multivoltine while populations in the northern part of the distribution range are univoltine (Friberg and Wiklund 2007). In addition, over evolutionary time scales the generation times may have varied in lineages due to variation in climatic conditions and distribution ranges (Altermatt 2010). The estimated divergence times of different repeat classes may thus not be seen as absolute values comparable across all lineages but rather as relative temporal variation in repeat proliferation rates within lineages.

**Results**

**The L. sinapis Genome Assembly**

The size of the L. sinapis genome assembly was 643 Mb (table 1), considerably larger than the size of the closest relatives with sequenced and well-characterized genome, the cloudless sulphur P. sennae (Cong et al. 2016) and the small cabbage white P. rapae (Shen et al. 2016). Reciprocal similarity searches of all repeat masked scaffolds using BLAST (Altschul et al. 1990) did not reveal any significantly similar scaffolds indicating false duplication during the assembly process. At a sequence similarity level of ≥95%, the average scaffold proportion that aligned to other scaffolds was only 1.55%, similar to what is observed when running an identical analysis for H. melpomene scaffolds (1.69%).

**Reconstruction of Phylogenies and Rate Estimates**

Phylogenetic reconstruction using the concatenated 224 core single-copy arthropod genes from the 16 lepidopteran genomes verified previously established topologies (fig. 1), both for global Lepidoptera species relationships (Cong et al. 2016), and for the Leptidea cryptic species complex (Dincă et al. 2011). The phylogenetic analysis also supported
considerable genetic structuring within the *Leptidea* clade, LsSpa being reciprocally monophyletic to the combined populations of LsSwe and LsKaz, and LjIre being reciprocally monophyletic as compared to LjKaz (fig. 1). Our estimated divergence times corresponded well with previous estimates of divergence times within Lepidoptera. For instance, the split between *H. melpomene* and *M. cynthia* was estimated to ca. 68.1 My (95% CI 56.8–79.6 My) in our analysis, matching previous estimates well (64.7–78.5 My), and the divergence time between *P. xuthus* and *P. polypetes* has previously been estimated to 23–40 My which agrees with our estimate of ca. 28.8 My (95% CI 23.7–34.5 My) (Hedges and Kumar 2009; Wahlberg et al. 2009; Nazari et al. 2011). Our main interest was to get approximate divergence time estimates for the *Leptidea* clade as compared with other Pieridae (represented by *P. sennae* and *P. rapae*) and for the species within the *Leptidea* genus to estimate the rate of genome size changes. We found that the TMRCA of the *Leptidea* and the *P. sennae/ P. rapae* lineage was 80.6 My (95% CI 67.6–93.2 My) and the TMRCA for the whole *Leptidea* species complex was ~3.0 My (95% CI 2.5–3.6 My). Within the *Leptidea* clade, the estimates of TMRCA for *L. sinapis* (LsSwe, LsKaz, LsSpa) and *L. juvernica* (LjKaz, LjIre) populations were 1.5 My (95% CI 1.1–1.8 My) and 1.6 My (95% CI 1.3–2.0 My), respectively (fig. 1).

**Association between Genome Size and Repeat Content in Lepidoptera**

To assess the fractions of interspersed repeat elements, we scanned 13 representative lepidopteran genome assemblies for interspersed repeat content. There was a very strong association (Pearson’s; *n* = 13, *r* = 0.800, *P* = 0.001) between genome size and proportion of repeat elements across species and *L. sinapis* was at the extreme end with a considerably larger proportion of repeats in the genome than other lepidopteran species (38%; fig. 2). To further investigate the contribution of specific repeat groups to genome size variation across Lepidoptera we counted the abundance of SINEs, LINES, DNA elements, LTR elements, and unclassified interspersed repeats using the RepeatModeler/RepeatMasker tools and found a difference in occurrence of specific repeat classes between taxa (fig. 3, supplementary table 3, Supplementary Material online). Specifically, the *L. sinapis* genome contained a larger proportion of LINES (5.4%), DNA elements (5.7%), and unclassified repeats (22.8%) than most other taxa, while fractions of other repeat subclasses (SINES andLTRs) were within the main range of other lepidopterans (fig. 3, supplementary table 3, Supplementary Material online). When counting the number of repeat subfamilies identified by RepeatModeler (Smit and Hubley 2010) and Repeat Masker (Smit et al. 1996–2010), present with ≥1,000 copies in a specific genome, the range was between six (*D. plexippus*) and 235 (*L. sinapis*) (supplementary table 4a, Supplementary Material online). *L. sinapis* had the highest LINE subfamily repeat count of any species, 4.8 times higher than the average across all other lineages, and comparatively high counts of DNA element (3.8 times higher than the mean in *L. sinapis*, highest in *C. cecrops*), LTR (3.4 times higher than mean, highest in *C. cecrops*), and unclassified subfamily repeats (2.5 times higher than the mean, highest in *C. cecrops*). The number of SINE subfamily repeats observed in *L. sinapis* was more similar to the range observed in other lineages (1.8 times higher than mean, highest in *B. mori*) (supplementary table 4b, Supplementary Material online).

**Association between Genome Size and Repeat Content within Leptidea**

The k-mer-based estimates of genome sizes within *Leptidea* showed that LjKaz had considerably larger genome size (776.7 Mb, ~13% larger, Kruskal–Wallis test: *χ*² = 22.97, df 5, *P* < 0.001, see supplementary table 5, Supplementary Material online, for details on the multiple population comparisons) than any other *Leptidea* population (on average 686.3 Mb), including the conspecific Irish population LjIre (656.0 Mb; fig. 4). We applied the dnaPipeTE pipeline (Goubert et al. 2015) on optimized subsampled re-sequencing reads (300,000–700,000 reads per individual) to de novo assemble the “repeatome” for each individual. This permitted the estimation of the abundance and relative age of recently active repeat elements in the re-sequencing data. There was a strong correlation between the fraction of both LINES (Pearson’s: *n* = 6, *r* = 0.42, *P* < 0.001) and LTRs (*n* = 6, *r* = 0.57, *P* < 0.001) with genome size within *Leptidea* but the correlation was driven only by the considerably higher repeat content and larger genome size in LjKaz (fig. 4). Upon removal of the LjKaz samples from the dataset we observed no significant correlations (Pearson’s: *n* = 6, *r* = 0.029, *P* = 0.84, and *n* = 6, *r* = 0.085, *P* = 0.56, for LINES and LTRs.
respectively) between specific repeat fractions and genome sizes estimates, indicating that the genome size and repeat content is similar across all other *Leptidea* populations (fig. 4).

The fractions of LINEs and SINEs were negatively correlated in *Leptidea* (Pearson’s: \( n = 60, r = -0.36, P < 0.005 \)), predominantly as an effect of a higher than average LINE content and lower than average SINE content in LjKaz (supplementary fig. 3, Supplementary Material online). However, the total amount of SINEs in LjKaz is at the same level as in other *Leptidea* populations, indicating that the genome expansion in LjKaz did not involve SINEs (supplementary fig. 3, Supplementary Material online). To further illustrate the discrepancy between LjKaz and other *Leptidea* populations, we performed a principal component analysis (PCA) based on the repeat fractions in the genome of all ten individuals in each respective population. The results clearly show that LjKaz deviates from the other populations (fig. 5).

**Reconstruction of Transposable Element Activity in Lepidoptera Lineages**

To obtain further information about timing and rates of proliferation of specific repeat elements, the aforementioned 13 genome assemblies were scanned for repeat content and sequence divergence between individual copies of repeat sub-classes LINEs, SINEs, LTRs, DNA transposons, and unclassified TEs using RepeatMasker. By applying a neutral mutation rate of \( 2.9 \times 10^{-9} \), estimated from de novo mutations in *H. melipomene* (Keightley et al. 2015), and the previously established tree topology, we modeled the divergence time of specific repeats in each lineage from the per-copy distance to consensus and visualized the activities of each respective repeat class in the butterfly tree of life. We found that the activity of TEs has been modest in butterflies in general but that particular elements have experienced a higher proliferation rate in the lineage leading to the *Leptidea* clade after the split from the other pierids around 80 Mya, with a considerable burst of activity in the time period 10–20 Mya (fig. 6). A notable observation is that the patterns of proliferation vary extensively between lineages (fig. 6) with particular repeat element classes being active at different time points during the radiation. Here, we point to some examples to illustrate this phenomenon. In almost all sampled lineages, for the identified LINE repeat families we observed a more or less constant rate of proliferation, but with an overall higher rate in the *Leptidea* lineage after the split from the other pierids around 80 Mya, with a considerable burst of activity in the time period 10–20 Mya (fig. 6). SINEs show varying patterns with an activity peak coinciding with the general TE activity peak in the *Leptidea* lineage and a similar activity pattern in the *Bombyx* lineage, while there is a more constant low rate in *Calycopis, Pieris, Melitaea*, and *P. glaucus* and more ancestral activity peaks in *P. xuthus* and *P. polytes* (fig. 6). LTRs are generally found at low frequency...
and the only observable pattern is a recent activity increase in *B. mori* and a higher ancestral activity in the *Papilio* lineage prior to the split of the three species (fig. 6). Unclassified repeats are abundant in several lineages, except in *H. melpomene*, the only lepidopteran species with a completely curated in-depth TE annotation (Lavoie et al. 2013), and in *D. plexippus*, which has a minor proportion of the genome consisting of repeats in general. Unclassified repeats also show varying patterns of activity across lineages where they are abundant (figs. 3 and 6). The most obvious pattern observed is the comparatively recent activity burst in *B. mori* that contrasts with constant, low rates in *P. xuthus*, *P. glaucus*, *P. sennae*, *M. cinxia*, *L. accius*, *P. rapae*, and *M. sexta*, comparatively high rate over the past 50 My in *Calycopis* and peaks of activity ~10 and 30 Mya in *L. sinapis* and *P. glaucus*, respectively (fig. 6). It should be noted that the observed general low rate of repeat activity far back in time (>50 My) in TE classes and lineages at least partly reflects the systematic problems in inferring activity for old repeats as a consequence of difficulties in identification when sequence divergence gets too high.

The frequency analyses of different repeat subclasses across *Leptidea* populations indicate that the activity of especially LINEs, LTRs, and DNA elements has continued or even increased in the *LjKaz* lineage after the split from *LjIre*.

**Discussion**

**The *L. sinapis* Genome Assembly**

In this study, we quantify the prevalence of TEs in butterflies and moths with extensive variation in genome size and estimate the rate of genome expansion in the *Leptidea* lineage in general and between different populations in the species *L. juvernica* in particular. These estimates point towards a mean expansion rate of 4.3 Mb/My in the *Leptidea* lineage after the split from other pierid butterflies ca. 80.6 Mya, and a mean expansion rate of 72 Mb/My in the *LjKaz* lineage after the split from *LjIre* ca. 1.6 Mya. These estimates can be translated to roughly 4 bp expansion rates per year in *Leptidea* in general, and 72 bp per year in *LjKaz* after the split from *LjIre*. (supplementary fig. 3, Supplementary Material online).

**Genome Size Expansion in *Leptidea* Butterflies**

Their genome assembly of *L. sinapis* was found to be 643 Mb which is one of the largest genome assembly of any lepidopteran taxon currently available (Challis et al. 2017), for example 2.8 times
larger than the smallest (227 Mb, P. polytes), and almost two times larger than the average lepidopteran genome (356.6 Mb) among the 12 investigated species for which relatively high-quality genome assemblies are available. In perspective, the 80 Lepidoptera species (91 entries in total) listed in the animal genome size database (www.genomesize.com; last accessed July 4, 2017, 2017-07-04; Gregory 2006) have a mean C-value of 0.62 pg [±0.29 s.d.; i.e., 0.61 ± 0.28 Gb following the conversion of Doležel et al. (2003) and the range varies from 0.23 to 1.94 pg (i.e., 0.22–1.90 Gb), with B. mori at a C-value of 0.52 pg (i.e., 0.51 Gb; genome assembly size = 482 Mb)]—this indicates that the genome size of L. sinapis is by no means extreme within the group. Furthermore, comparisons between genome size estimates based on flow-cytometry and DNA sequence assemblies consistently point toward that the latter tend to underestimate the genome size since repetitive sequences (especially centromeric and telomeric regions and non-recombining parts of sex-limited sex chromosomes) are always difficult to assemble (Chaisson et al. 2015; Kapusta and Suh 2017). This general feature of the assembly process, in combination with the observed high repeat content (38%) indicates that the L. sinapis genome might be larger than our current estimate—preliminary data from flow-sorting actually points toward a genome size of >800 Mb in L. sinapis, and even larger in L. juvernica, although these are estimates based on one single male specimen from each species, respectively (Petr Nguyen, personal communication).

To rule out the possibility that the comparatively large genome assembly of L. sinapis was an assembly artefact generating false duplication of a potentially highly heterozygous genome (Zhang and Backstroom 2014), we scanned the assembly for highly conserved eukaryotic (n = 434 entries in total) and arthropod (2,718 entries) gene sets and found only a minor fraction of duplicated genes (5 and 43 in each respective gene set which corresponds to 1.1 and 1.6% of all

![Fig. 6](https://example.com/fig6.png)
Genome Size Expansion in Leptidea Butterflies


Phylogenetic Analyses

When comparing our nuclear gene-based divergence time estimates to previously available data, there was good agreement regarding the deeper nodes corresponding to the split between Papilionidae and Nymphalidae, and nodes within these families associated with the splits between H. melpomene and M. cinxia and between P. xuthus and P. polytes, respectively (Hedges and Kumar 2009; Wahlberg et al. 2009; Nazari et al. 2011). However, the nuclear gene-based divergence time estimates within the Leptidea complex (e.g., TMRCA = 2.4–3.6 My) were considerably higher than previous estimates based on a small set of mitochondrial and nuclear genes (TMRCA = 0.3 My; Dincă et al. 2011). Given the comparatively small data set used by Dincă et al. (2011), the extensive variation in the ratio of mtDNA to nuclear mutation rates (Allio et al. 2017), and the good agreement between ours and previous estimates of other divergence times in the Lepidoptera phylogeny (Hedges and Kumar 2009; Wahlberg et al. 2009; Nazari et al. 2011; Condamine et al. 2012; Heiklää et al. 2012; Espeland et al. 2015; ), the divergence times within Leptidea presented in this study are likely more robust.

Variation in TE Activity, Genome Size and Karyotype Structure in Lepidoptera

To get an amalgamated view of the causes and consequences of genome size variation it is crucial to have detailed information about the genomic architecture of the organisms (Petrov 2001; Gregory 2005). Therefore, we set out to investigate if the genome size differences in Lepidoptera could be explained by the expansion of specific selfish genetic elements. In line with previous research from a wide range of taxonomic groups—for example pufferfish (Aparicio et al. 2002); rice and thale cress (Bennetzen et al. 2005); threespine stickleback (Blass et al. 2012); birds/mammals (Kapusta et al. 2017); Norway spruce (Nystedt et al. 2013); Hordeum grasses (Vicient et al. 1999), and migratory locust (Wang et al. 2014)—our analyses in Lepidoptera show that genome size variation across lineages can be largely explained by differences in overall content of TEs. Across all lineages, the investigated portion of the genomes contained between seven and 38% interspersed repeats, and different lineages displayed considerable differences in proportions and numbers of specific elements and TE subfamilies, with L. sinapis containing both the highest proportion of TEs and the largest number of TE subfamilies. The overall largest group of repeats in our data was in the partition that could not be accurately classified in automatic de novo repeat predictions. This has also been observed in a recently developed genome assembly of the squinting bush brown butterfly (Bicyclus anyana) that contained 18% unclassified and 7% classified repeats (Nowell et al. 2017). A recent study in birds where unclassified repeats were manually curated showed that those predominantly represented LTRs (Kapusta and Suh 2017), but it is not clear if this also applies to Lepidoptera and tedious manual curation of unclassified repeats will be necessary to get a detailed picture of the composition of TEs in this category. There was a negative relationship between the fractions of LINEs and SINEs when comparing LjKaz to the other populations in the Leptidea clade but the amount of SINE sequences in all Leptidea genomes were similar. This suggests that the activity of SINEs has been very low in LjKaz after the split from the other Leptidea populations. This is in line with the observation that SINEs mainly proliferate by hijacking the enzymatic machinary of LINEs and, if so, often outnumber their LINE counterparts (Ohshima et al. 1996; Ohshima and Okada 2005).

By comparing the average genome sizes of Leptidea to other lepidopteran lineages we could estimate the rate of genome size increase over time. The closest relatives to Leptidea available in the sample set is P. sennae and P. rapae. We estimated these two lineages (both within family Pieridae) to have a divergence time of ~80 My and this translates to an overall net expansion rate of four Mb/My which is ~4 bp/year. If we assume a generation time of one generation per year, this yields a net expansion rate in Leptidea of four bp per generation (see the Methods section for comments on assumed generation times and how that may affect rate estimates). Since the proliferation mechanism for TEs involves entire repeat sequences ranging from ~100 bp in SINEs to over 10 kb in LTRs (Sotero-Caio et al. 2017) and repeat sequences underlie the overwhelming part of the observed genome size differences, the increase in genome size should however rather be seen as a punctuated process rather than a steady, stepwise addition of small DNA fragments. Nonetheless, the low overall expansion rate in Leptidea suggests hyperactivity of TEs and that there is limited power for natural selection to act against their proliferation unless TE insertions occur in functional regions.

Our phylogenetic analysis based on nuclear genes clearly separates L. sinapis populations with distinct karyotypes into
monophyletic clades, a novel finding that has not previously been detected with smaller marker sets (Dincă et al. 2011, 2013). It is tempting to speculate about the association between a high repeat content and the high fission/fusion rates resulting in dramatic variation in chromosome numbers within and across species within the *Leptidea* species complex (Dincă et al. 2011; Lukhtanov et al. 2011; Šíchová et al. 2015). Previous analyses indicate that interspersed repeats may mediate chromosomal rearrangements via for example non-homologous recombination (Völker et al. 2010; Zhang et al. 2011) which means that TE activity may not only have an impact on genome size, but also on the karyotype evolution. Interestingly, several Lepidoptera lineages show evidence for recurrent chromosome number changes. Within the subgenus *Agrodiaetus* chromosome numbers vary from 2*n*=20 to 2*n*=268 (Lukhtanov et al. 2005) and the species *Lysandra condon* displays a chromosome number cline similar to *L. sinapis* but less dramatic (Talavera et al. 2013). In addition, *Polymnatus atlanticus* has the highest number of chromosomes (2*n*=448–452) recorded in any metazoan organism (Lukhtanov 2015). This highlights the suitability of Lepidoptera in general, and *Leptidea* butterflies with their extreme karyotype variation in particular (Dincă et al. 2011; Lukhtanov et al. 2011; Šíchová et al. 2015), as a study system for investigating the role of repetitive sequences on karyotype structure, why chromosome number changes seem to be spurted in specific lineages and how rearrangements affect patterns of genome differentiation, adaptation and speciation.

**Genome Size Variation and TE Activity in Leptidea**

An intriguing observation related to the observed larger genome size in *Leptidea* as compared with other lepidopteran lineages was that cell nucleus size varies across *Leptidea* species with a considerably larger nucleus in *L. juvernica* than in *L. reali* and *L. sinapis* (Šíchová et al. 2015). The *L. juvernica* populations sampled in the comparison were collected in the Czech Republic and samples from other parts of the distribution range were not included. Previous analyses have not detected any morphological differences, including male or female genitalia, that suggest the existence of a cryptic species within *L. juvernica* (Dincă et al. 2011), and no prezygotic barriers among LjKaz, LjIre, and LjSwe populations have been found (which exist between *L. sinapis*, *L. reali*, and *L. juvernica* because of female choice) (Dincă et al. 2013). Genetic analyses based on the generally fast evolving mtDNA genes (*COI* and *ND1*) and a handful of nuclear genes have shown that the LjKaz population is undistinguishable from all other populations studied from European mainland (Dincă et al. 2011, 2013; Lukhtanov et al. 2011; Šíchová et al. 2015). Given our results, showing that LjKaz has a considerably larger genome size than LjIre (and all other sampled *Leptidea* populations), it is plausible that a genome size expansion has happened in “continental” Eurasian *L. juvernica* after the split from LjIre. This could potentially be a consequence of lower effective population size and less efficient purging of slightly deleterious TE insertions in LjKaz (Lynch 2007). In support of that, genome-wide analyses of nucleotide diversity (θ) show that *L. juvernica* in general have a reduced level of polymorphism (θ=0.11–0.17%) compared with other *Leptidea* populations (θ=0.33–0.39%), and the reduction in LjKaz (θ=0.11%) is particularly pronounced (Talla V, Dincă V, Vila R, Wiklund C, Backström N, Unpublished data). Since the divergence time between LjKaz and LjIre was estimated to 1.6 My and the genome size difference was roughly 115 Mb, the net genome expansion rate is in the range of 72 Mb/My. Again, when translated to per generation (year) the expansion rate is rather modest (<100 bp/year). Recent, detailed proliferation rate analyses of LTRs in *Drosophila melanogaster* (one insertion in 1,000 to 1,000,000 generations; Huang et al. 2012) and LINEs (one insertion in 212 births; Xing et al. 2009) and Alu-elements (one insertion in 20 births; Cordaux et al. 2006) in humans indicate that proliferation rates vary across taxa and between types of repeats; at the extreme, novel insertions of TEs have been observed to occur with up 20–100 transposition events (0.1–1 Mb in total) in a single generation in some taxa (Petrov 2001 and references therein). Hence, the observed overall genome expansion rates as a consequence of TE activity in *Leptidea* in general and in LjKaz in particular are not spectacular, but high enough to generate considerable genome size differences over short evolutionary time. Again, we cannot specifically rule out that the increase in genome size is an adaptive response in LjKaz as compared to other *Leptidea* lineages, but the observation of stochastic variation in TE proliferation rates, likely accompanied by reduced efficiency of purifying selection during periods of limited population size, suggests that genetic drift has contributed considerably to genome size evolution in *Leptidea* (Lynch 2007). Notably, differences in TE proliferation rates might even be exacerbated under environmental stress (Kim et al. 2014; Migicovsky and Kovalchuk 2014), indicating that variation in environmental conditions during population isolation—for example, under allopatic separation during glacial periods—may also contribute to genome size evolution.

**Perspectives**

Even if accumulation of repetitive DNA might be associated with a cost due to insertion in functional regions and potentially increased energetic demands for the replication machinery, novel inserted DNA may also provide a template for evolution of novel functions. It has, for example, been suggested that the gain of introns in eukaryotes might have had a non-adaptive origin, but once present they have allowed for the evolution of novel functions. It has, for example, been suggested that the gain of introns in eukaryotes might have had a non-adaptive origin, but once present they have allowed for the evolution of novel functions. It has, for example, been suggested that the gain of introns in eukaryotes might have had a non-adaptive origin, but once present they have allowed for the evolution of novel functions.
involving a classic textbook example of a rapid evolutionary response due to natural selection is that a large TE insertion in the first intron of the gene cortex underlies the melanistic phenotype in *Biston betularia*, the peppered moth (Van’t Hof et al. 2016). This is potentially just a beginning of what will be unearthed in the upcoming years. Although we are still far from having a detailed understanding of potential functional gains or deleterious effects of proliferation of repetitive sequences in the Lepidoptera, the increasing availability of high-quality genome assemblies and accumulating annotation information and functional genomic studies will likely shed light on costs and benefits of repetitive DNA proliferation in general and in specific cases also in this group of organisms.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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