



Data Article

Complete mitogenome data from a European specimen of *Ostrinia scapularis* (Walker, 1859) (Lepidoptera, Pyraloidea, Crambidae, Pyraustinae)



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ABSTRACT

We present an assembly and annotation of the mitogenome of a European specimen of the Adzuki bean borer, *Ostrinia scapularis* (Walker, 1859). The data were obtained by combining WGS data issue of a *de novo* and a previously published sequence library (Gschloessl et al., 2018). We also provide the phylogenetic positioning of the mitogenome within the *Ostrinia* genus, the *Crambidae* family and with more distant Lepidoptera species.

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**Specifications Table**

Subject	Insect Science
Specific subject area	Lepidoptera, Crambidae, Mitogenomics
Type of data	<ul style="list-style-type: none"> <li>• FASTQ: DNA sequence reads</li> <li>• FASTA: mitochondrial genome assembly</li> <li>• GFF: mitogenome gene annotations</li> <li>• TABLE: gene annotations, base composition, Lepidoptera species used for phylogenetic analysis</li> <li>• FIGURE: mitogenomic circular map, AT-rich region sequence, phylogenetic tree</li> </ul>
How data were acquired	Whole-genome shotgun sequencing on Illumina NextSeq 500 v2 platform using the paired-end protocol (2 × 150 base pairs (bp)).
Data format	Raw sequence reads and analysed (assembled and annotated) mitogenome.
Parameters for data collection	Genomic DNA was extracted from one <i>Ostrinia scapularis</i> larva using the DNeasy Tissue Kit from Qiagen (Hilden, Germany). A 2 × 150 bp shot-gun paired-end library with an insert size of 300 bp was generated and sequenced by LGC Genomics GmbH (Berlin, Germany) on an Illumina NextSeq 500 v2 platform.
Description of data collection	The <i>Ostrinia scapularis</i> mitogenome was assembled with MITObim v1.9. Genomic regions were annotated with MITOS v2. The circular mitochondrial genome map was generated with OGDRAW. Multiple sequence alignments were generated separately for each protein-coding and rRNA gene with MAFFT v7.471. Phylogenetic trees were built on the concatenated gene alignments with PHYML v3.0 and MrBayes v3.2.7a and visualized with FigTree v1.4.4.
Data source location	Samples were collected in 2015 near Abbeville/France (50° 8' 13.57"N; 1° 50' 23.62"E).
Data accessibility	Repository name: NCBI BioProject Data identification number: PRJNA637835 Direct URL to data: <a href="http://www.ncbi.nlm.nih.gov/bioproject/637835">http://www.ncbi.nlm.nih.gov/bioproject/637835</a> Repository name: NCBI Genbank Data identification number: MT801073 Direct URL to data: <a href="http://www.ncbi.nlm.nih.gov/nucleotide/MT801073">http://www.ncbi.nlm.nih.gov/nucleotide/MT801073</a>

**Value of the Data**

- This data represents the mitogenome of a European specimen of *Ostrinia scapularis*.
- The data can benefit to researchers working on Lepidoptera evolution and higher systematics.
- The present data can be used for phylogenetic studies of Lepidoptera mitogenomes especially for getting deeper insights in diversification dynamics of *Ostrinia* species.

**1. Data Description**

The *O. scapularis* (hereafter named OSCA) mitogenome had a size of 15,305 bp and an A + T content of 80.9%. All 38 genetic regions commonly known for arthropod mitogenomes [2] were identified (Supplementary GFF annotation file): 13 protein-coding genes, 22 tRNA and 2 rRNA (12S rRNA and 16S rRNA) genes, and one A + T-rich control region. The gene order and the composition of start on stop codons of the protein-coding genes corresponded to those observed for other *Ostrinia* spp. [3,4] and Crambidae [5] mitogenomes. Fifteen intergenic spacer regions (see Table 1: positive intergenic nucleotide values) of a total length of 341 bp were identified

**Table 1**Gene features of the *O. scapularis* mitogenome.

Gene (anticodon)	Start	Stop	Strand	Size	Intergenic nucleotides	Start/stop codon
trnM(cat)	1	68	+	68	-1	
trnI(gat)	68	135	+	68	0	
trnQ(ttg)	136	204	-	69	92	
ND2	297	1289	+	993	-15	ATT/TAA
trnW(tca)	1275	1341	+	67	-8	
trnC(gca)	1334	1398	-	65	0	
trnY(gta)	1399	1465	-	67	8	
COX1	1474	3009	+	1536	-5	CGA/TAA
trnL2(taa)	3005	3071	+	67	48	
COX2	3120	3788	+	669	-35	ATA/TAA
trnK(ctt)	3754	3824	+	71	-1	
trnD(gtc)	3824	3892	+	69	0	
ATP8	3893	4054	+	162	-7	ATA/TAA
ATP6	4048	4722	+	675	-1	ATG/TAA
COX3	4722	5513	+	792	2	ATG/TAA
trnG(tcc)	5516	5582	+	67	0	
ND3	5583	5936	+	354	10	ATT/TAA
trnA(tgc)	5947	6013	+	67	-1	
trnR(tcg)	6013	6077	+	65	-1	
trnN(gtt)	6077	6143	+	67	2	
trnS1(gct)	6146	6211	+	66	1	
trnE(ttc)	6213	6279	+	67	-2	
trnF(gaa)	6278	6346	-	69	-17	
ND5	6330	8057	-	1728	24	ATT/TAA
trnH(gtg)	8082	8148	-	67	-1	
ND4	8148	9488	-	1341	7	ATG/TAA
ND4L	9496	9789	-	294	9	ATG/TAA
trnI(tgt)	9799	9865	+	67	0	
trnP(tgg)	9866	9930	-	65	32	
ND6	9963	10,469	+	507	-1	ATT/TAA
CYTB	10,469	11,617	+	1149	-1	ATG/TAA
trnS2(tga)	11,617	11,684	+	68	38	
ND1	11,723	12,661	-	939	1	ATG/TAG
trnL1(tag)	12,663	12,730	-	68	21	
16S rRNA	12,752	14,023	-	1272	46	
trnV(tac)	14,070	14,135	-	66	0	
12S rRNA	14,136	14,914	-	779	0	
A + T-rich region	14,915	15,305	+	391	0	

The gene names, the coordinates, the orientation, the size, the gene overlaps/intergenic spacers (column 6) and the start/stop codons of the *O. scapularis* mitogenome are listed.

**Table 2**Base composition and skewness indices for each specific gene region of the *O. scapularis* mitogenome.

Type	Size [bp]	A%	G%	T%	C%	A + T%	AT skew	GC skew
Entire mitogenome	15,305	41.7	7.7	39.2	11.4	80.9	0.031	-0.197
Protein-coding genes	11,139	34.5	10.8	44.9	9.9	79.4	-0.131	0.045
tRNA	1480	41.7	10.8	39.9	7.6	81.6	0.022	0.172
rRNA	2051	41.7	9.9	43.4	4.9	85.1	-0.021	0.338
A + T-rich region	391	42.7	2.0	49.6	5.6	92.3	-0.075	-0.467

which ranged from 1 to 92 bp with the longest being located between the trnQ and ND2 genes. Furthermore, region overlaps (see Table 1: negative intergenic nucleotide values) of 1 to 35 bp could be observed for fifteen gene pairs. More annotation details can be found in Table 1 and are also represented as mitogenome map in Fig. 1. Base compositions, AT and GC skews to each genomic region type are shown in Table 2. The tRNA secondary structures showed the typical



**Table 3**

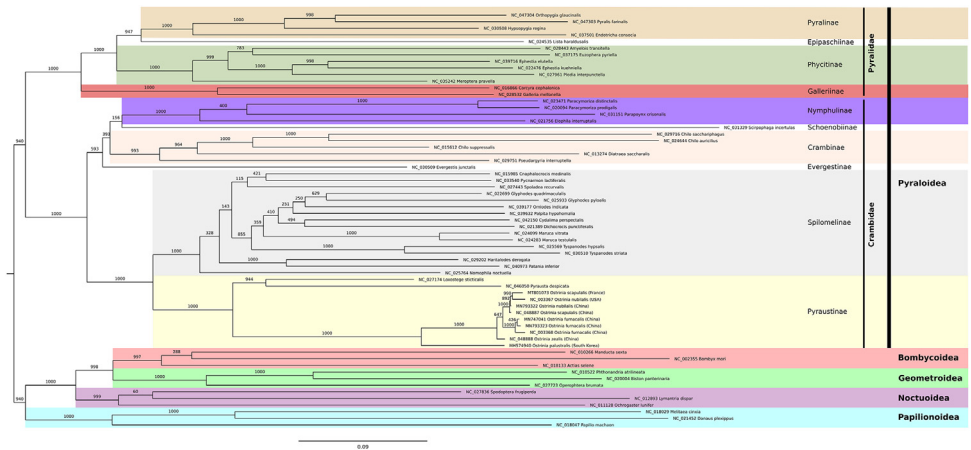
Lepidoptera mitogenomes which were used to build the phylogenetic trees. For *Ostrinia* spp. the countries are also given in which the samples were taken.

Super-family	Family	Subfamily	Species	Genbank no.
Bombycoidea	Bombycidae	Bombycinae	<i>Bombyx mori</i>	NC_002355
Bombycoidea	Saturniidae	Saturniinae	<i>Actias selene</i>	NC_018133
Bombycoidea	Sphingidae	Sphinginae	<i>Manduca sexta</i>	NC_010266
Geometroidea	Geometridae	Ennominae	<i>Biston panterinaria</i>	NC_020004
Geometroidea	Geometridae	Ennominae	<i>Phthonandria atrilineata</i>	NC_010522
Geometroidea	Geometridae	Larentiinae	<i>Operophtera brumata</i>	NC_027723
Noctuoidea	Erebidae	Lymantriinae	<i>Lymantria dispar</i>	NC_012893
Noctuoidea	Noctuidae	Noctuidae	<i>Spodoptera frugiperda</i>	NC_027836
Noctuoidea	Notodontidae	Thaumetopoecinae	<i>Ochrogaster lunifer</i>	NC_011128
Papilionoidea	Nymphalidae	Danainae	<i>Danaus plexippus</i>	NC_021452
Papilionoidea	Nymphalidae	Nymphalinae	<i>Melitaea cinxia</i>	NC_018029
Papilionoidea	Papilionidae	Papilioninae	<i>Papilio machaon</i>	NC_018047
Pyraloidea	Crambidae	Crambinae	<i>Chilo auricilius</i>	NC_024644
Pyraloidea	Crambidae	Crambinae	<i>Chilo sacchariphagus</i>	NC_029716
Pyraloidea	Crambidae	Crambinae	<i>Chilo suppressalis</i>	NC_015612
Pyraloidea	Crambidae	Crambinae	<i>Diatraea saccharalis</i>	NC_013274
Pyraloidea	Crambidae	Crambinae	<i>Pseudargyria interruptella</i>	NC_029751
Pyraloidea	Crambidae	Evergestinae	<i>Evergestis junctalis</i>	NC_030509
Pyraloidea	Crambidae	Nymphulinae	<i>Elophila interruptalis</i>	NC_021756
Pyraloidea	Crambidae	Nymphulinae	<i>Paracymoriza distinctalis</i>	NC_023471
Pyraloidea	Crambidae	Nymphulinae	<i>Paracymoriza prodigalis</i>	NC_020094
Pyraloidea	Crambidae	Nymphulinae	<i>Parapopynx crisonalis</i>	NC_031151
Pyraloidea	Crambidae	Pyraustinae	<i>Parostege sticticalis</i>	NC_027174
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia furnacalis</i> (China)	NC_003368
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia furnacalis</i> (China)	MN793323
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia furnacalis</i> (China)	MN747041
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia nubilalis</i> (USA)	NC_003367
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia nubilalis</i> (China)	MN793322
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia palustralis</i> (South Korea)	MH574940
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia scapularis</i> (France)	MT801073
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia scapularis</i> (China)	NC_048887
Pyraloidea	Crambidae	Pyraustinae	<i>Ostrinia zealis</i> (China)	NC_048888
Pyraloidea	Crambidae	Pyraustinae	<i>Pyrausta despicata</i>	NC_046050
Pyraloidea	Crambidae	Schoenobiinae	<i>Scirpophaga incertulas</i>	NC_031329
Pyraloidea	Crambidae	Spilomelinae	<i>Cnaphalocrocis medinalis</i>	NC_015985
Pyraloidea	Crambidae	Spilomelinae	<i>Cydalima perspectalis</i>	NC_042150
Pyraloidea	Crambidae	Spilomelinae	<i>Dichocrocis punctiferalis</i>	NC_021389
Pyraloidea	Crambidae	Spilomelinae	<i>Glyphodes pyloalis</i>	NC_025933
Pyraloidea	Crambidae	Spilomelinae	<i>Glyphodes quadrimaculalis</i>	NC_022699
Pyraloidea	Crambidae	Spilomelinae	<i>Haritalodes derogata</i>	NC_029202
Pyraloidea	Crambidae	Spilomelinae	<i>Maruca testulalis</i>	NC_024283
Pyraloidea	Crambidae	Spilomelinae	<i>Maruca vitrata</i>	NC_024099
Pyraloidea	Crambidae	Spilomelinae	<i>Nomophila noctuella</i>	NC_025764
Pyraloidea	Crambidae	Spilomelinae	<i>Omiodes indicata</i>	NC_039177
Pyraloidea	Crambidae	Spilomelinae	<i>Palpita hypohomalia</i>	NC_039632
Pyraloidea	Crambidae	Spilomelinae	<i>Patania inferior</i>	NC_040973
Pyraloidea	Crambidae	Spilomelinae	<i>Pycnarmon lactiferalis</i>	NC_033540
Pyraloidea	Crambidae	Spilomelinae	<i>Spoladea recurvalis</i>	NC_027443
Pyraloidea	Crambidae	Spilomelinae	<i>Tyspanodes hypsalis</i>	NC_025569
Pyraloidea	Crambidae	Spilomelinae	<i>Tyspanodes striata</i>	NC_030510
Pyraloidea	Pyrallidae	Epipaschiinae	<i>Lista haraldusalis</i>	NC_024535
Pyraloidea	Pyrallidae	Galleriinae	<i>Corcyra cephalonica</i>	NC_016866
Pyraloidea	Pyrallidae	Galleriinae	<i>Galleria mellonella</i>	NC_028532
Pyraloidea	Pyrallidae	Phycitinae	<i>Amyelois transitella</i>	NC_028443
Pyraloidea	Pyrallidae	Phycitinae	<i>Ephestia elutella</i>	NC_039716
Pyraloidea	Pyrallidae	Phycitinae	<i>Ephestia kuehniella</i>	NC_022476
Pyraloidea	Pyrallidae	Phycitinae	<i>Euzophera pyriella</i>	NC_037175
Pyraloidea	Pyrallidae	Phycitinae	<i>Meroptera pravella</i>	NC_035242
Pyraloidea	Pyrallidae	Phycitinae	<i>Plodia interpunctella</i>	NC_027961
Pyraloidea	Pyrallidae	Pyralinae	<i>Endotricha consocia</i>	NC_037501
Pyraloidea	Pyrallidae	Pyralinae	<i>Hyposopygia regina</i>	NC_030508
Pyraloidea	Pyrallidae	Pyralinae	<i>Orthopygia glaucinalis</i>	NC_047304
Pyraloidea	Pyrallidae	Pyralinae	<i>Pyralis farinalis</i>	NC_047303

12S rRNA-14,915-

TAATACTTAAAATTTTAACTTAGATTTTTTTTTTTTTTTTTTATATTA  
 AATATTTAATATAAATTATTAATTTTAAATATTTATTTCTTTTCTTT  
 TTCATAATATTAATATTAATAATTAATATCATTTATACAGCGATTTATAA  
 TCATTGAAATAAATAATTAATTATTAAGTTTAATAATTAATTAATTAATGA  
 TTTATTAATATATTTATAAATAAATTTCTATTATATATATATATATA  
 TTAAATATAATAATTAATTAATATTTAATATATATATATATATATA  
 AACCATTCTAATTTTTTTTCTTTTAATAATAAATTAATAAATAAGCTAA  
 ATAAAGCTTATATAAACCATTCCTAATTTTTTTTCTTTTAATAATAAA-  
 15,305-trnM

**Fig. 2.** Characteristics of the A+T-rich region of the *O. scapularis* mitogenome. The AT-rich region consists of a TTAGA (red) motif which is followed by a 18 bp poly-T stretch (brown). Subsequently, it contains five ATTTA motifs (blue) of which the last one is followed by a (AT)<sub>10</sub> microsatellite stretch (purple) and terminates with a polyT/polyA region (green).



**Fig. 3.** Phylogenetic tree representing 63 Lepidoptera mitogenomes which was built on concatenated alignments of the nuclear sequences of the 13 protein-coding and both rRNA genes applying a Maximum Likelihood approach with PHYML. For *Ostrinia* spp. the countries are also given in which the samples were taken.

an Illumina NextSeq 500 v2 platform. In total, 372,260,856 genomic reads were generated of which 306,203,188 remained after quality trimming with Trimmomatic v0.36 [6] (specific parameters were: PE -pred33, ILLUMINACLIP:\$ADAPTERF:2:26:10 HEADCROP:5 LEADING:26 TRAILING:26 SLIDINGWINDOW:5:28 MINLEN:30, with \$ADAPTERF containing all Illumina adapters and their reverse complement, respectively).

2.2. Mitogenome assembly and feature analyses

The OSCA mitogenome was assembled in two steps as follows. First, the scaffolds of a previously published *O. scapularis* genome draft [1], OSCA v1.2 (BioProject PRJNA390510), were searched for putative mitochondrion sequences by using *blastn* v2.2.28 [7] against the mitochondrial genome sequence of the closely related Asian corn borer *Ostrinia furnacalis* (Guenée) (NCBI

accession number NC\_003368). Genome scaffolds with high quality mitochondrion hits were selected and assembled by LGC with CAP3 v2013 [8] into a mitogenome sequence of 14,864 bp. In a second step, this draft mitogenome was refined using MITObim v1.9 [9] (using the default k-mer size of 31 bp and the `-pair` parameter) and the Trimmomatic-cleaned paired-end Illumina reads. Gene identification as well as tRNA structure prediction were carried out on the MITOS web server v2 [10], using as reference set 'RefSeq 81 Metazoa' and the genetic code '5' for invertebrates. Nucleic base contents were determined with the `wordcount` program of the EMBOSS toolkit v6.6.0.0 [11]. AT/GC skewness indices describing the genomic compositional asymmetry of a mitogenome or specific sequence region were calculated as:  $AT\ skew = [A - T] / [A + T]$  and  $GC\ skew = [G - C] / [G + C]$ , where each base letter represents the respective base count. The circular map of the OSCA mitogenome was generated with the web tool OGDRAW v1.3.1 [12].

### 2.3. Phylogenetic analyses

In order to perform comparative analyses of *O. scapularis* with other Lepidoptera the nucleotide sequences of the 13 PCGs and 2 rRNA genes of 63 mitogenomes - comprising 59 Lepidoptera species - (Table 3) were downloaded from the NCBI 'nucleotide' database (<https://www.ncbi.nlm.nih.gov/nucleotide>, May 2020). Our taxon sampling was focused on the superfamily Pyraloidea, encompassing 34 species belonging to the Crambidae (including *O. scapularis*) and 13 species belonging to the Pyralidae. Regarding *Ostrinia* spp., we included in total nine mitogenomes belonging to five *Ostrinia* species. As outgroups we used 12 representative species of the lepidopteran superfamilies Bombycoidea, Geometroidea, Noctuoidea and Papilionoidea; based on the comprehensive study of Wahlberg et al. [13], we used the three Papilionoidea representatives to root our trees. The PCG and rRNA nucleotide sequences of all species were grouped into 15 gene-specific sequence sets. Each gene set was in separate globally aligned with the program MAFFT v7.471 [14] by applying the Needleman-Wunsch algorithm (`mafft-ginsi` for PCGs and `mafft-qinsi` for rRNA sequences) with a maximum of 1000 refinement iterations (`-maxiterate 1000`), leaving the other parameters set to their default value. Subsequently, for each species all gene-specific alignments were concatenated following the same gene order using in-house Python and Perl scripts.

Phylogenetic trees were generated with the PHYML web server v3.0 [15] and a local installation of MrBayes v3.2.7a [17]. PHYML was run using the Smart Model Selection [16] option with the Akaike Information Criterion and the tree searching method BIONJ being set and by applying 1000 bootstraps. Concerning the MrBayes analysis, a separate partition was set for each gene. For each partition the evolutionary model was set to the GTR substitution model (`nst=6`) with gamma-distributed rate variation (`rates=invgamma`) across sites and a proportion of invariable sites (`GTR +I +Γ`). The standard nucleotide substitution model (`nucmodel=4by4`) was chosen and priors settings were left as by default. Trees were built using 100,000 samples (`ngen=1,000,000`, `samplefreq=10`). Further parameters were: `printfreq=100`, `diagnfreq=1000`, `nchains=4`, `savebrlens=yes`, `starttree=random`, `startparams=reset`, `outgroup pamac` (corresponding to *Papilio machaon*), `sump burnin=25,000`, `sumt burnin=25,000`. The phylogenetic tree was visualized using the program FigTree v1.4.4 [18].

### CRedit Author Statement

BG conceived the study. RS led the ANR research program on European *Ostrinia* genomics. RS, PA, SN collected the samples in the field. SN, PA prepared the samples and extracted the DNA. BG was responsible for the bioinformatics analyses, i.e. improvement of the mitogenome assembly, annotation, characterization, comparative analyses. BG and GJK did the phylogenetic analyses. All authors wrote and approved the manuscript.

## Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships which have, or could be perceived to have, influenced the work reported in this article.

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## Supplementary Materials

Supplementary material associated with this article can be found, in the online version at doi:[10.1016/j.dib.2020.106427](https://doi.org/10.1016/j.dib.2020.106427).

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