DATA NOTE

The genome sequence of the yellow-tail moth, *Euproctis similis* (Fuessly, 1775) [version 2; peer review: 2 approved, 1 approved with reservations]

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**Abstract**

We present a genome assembly from an individual male *Euproctis similis* (the yellow-tail; Arthropoda; Insecta; Lepidoptera; Lymantriidae). The genome sequence is 508 megabases in span. Over 99% of the assembly is scaffolded into 22 chromosomal pseudomolecules, with the Z sex chromosome assembled. The complete mitochondrial genome, 15.5 kb in length, was also assembled.

**Keywords**

*Euproctis similis*, yellow-tail, genome sequence, chromosomal

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This article is included in the Tree of Life gateway.
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Author roles: Boyes DH: Formal Analysis, Investigation, Resources; Holland PWH: Formal Analysis, Investigation, Supervision, Writing – Original Draft Preparation;

Competing interests: No competing interests were disclosed.

Grant information: This work was supported by Wellcome through core funding to the Wellcome Sanger Institute (206194) and the Darwin Tree of Life Discretionary Award (218328). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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How to cite this article: Boyes DH, Holland PWH, University of Oxford and Wytham Woods Genome Acquisition Lab et al. The genome sequence of the yellow-tail moth, Euproctis similis (Fuessly, 1775) [version 2; peer review: 2 approved, 1 approved with reservations] Wellcome Open Research 2021, 6:227 https://doi.org/10.12688/wellcomeopenres.17188.2

First published: 13 Sep 2021, 6:227 https://doi.org/10.12688/wellcomeopenres.17188.1
Species taxonomy

Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Lepidoptera; Glos-sata; Ditrysia; Noctuoidea; Erebidae; Lymantriinae; Euproctis; Euproctis similis Fuessly 1775 (NCBI:txid987935).

Introduction

Euproctis similis, the yellow-tail moth, is widespread across temperate Europe and Asia. In the UK, the moth is relatively common across much of England and Wales, with scattered records from southern Scotland and Northern Ireland. The larvae of E. similis feed on a range of deciduous trees and shrubs, including Crataegus, Prunus, and Betula, in some situations becoming a pest on ornamental and fruit trees. Larvae are also notable for bearing long hairs that can cause skin irritation in humans, although the effects are rarely as serious as those caused by larvae of the closely related Euproctis chrysorrhoea (brown-tail). A genome sequence for E. similis, therefore, may have agricultural and biomedical relevance, in addition to its use in evolutionary biology, ecology and genome biology. The karyotype of E. similis has been previously recorded as n=22 or 23 (Belyakova & Lukhtanov, 1994). This is not unexpected since Lepidoptera exhibit considerable variation in chromosome number, although n=31 is the most common karyotype (Ahola et al., 2014). The genome of E. similis was sequenced as part of the Darwin Tree of Life Project, a collaborative effort to sequence all of the named eukaryotic species in the Atlantic Archipelago of Britain and Ireland. Here we present a chromosomally complete genome sequence for E. similis, based on one male specimen from Wytham Woods, Oxfordshire (biological vice-country: Berkshire), UK.

Genome sequence report

The genome was sequenced from a single male E. similis (Figure 1) collected from Wytham Woods, Oxfordshire (biological vice-county: Berkshire), UK (latitude 51.772, longitude -1.338). A total of 70-fold coverage in Pacific Biosciences single-molecule long reads (N50 17 kb) and 78-fold coverage in 10X Genomics read clouds were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. Manual assembly curation corrected 40 missing/misjoins and removed 3 haplotypic duplications, reducing the assembly length by 0.10% and the scaffold number by 42.00%, and increasing the scaffold N50 by 14.24%. The final assembly has a total length of 508 Mb in 30 sequence scaffolds with a scaffold N50 of 24 Mb (Table 1). Over 99.9% of the assembly

Table 1. Genome data for Euproctis similis, ilEupSimi1.1.

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<td>Species</td>
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</tr>
<tr>
<td>Specimen</td>
<td>ilEupSimi1</td>
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<td>NCBI taxonomy ID</td>
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<td>BioSample ID</td>
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<td>Male, head/abdomen/thorax</td>
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<td>Span (Mb)</td>
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<tr>
<td>Contig N50 length (Mb)</td>
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<tr>
<td>Number of scaffolds</td>
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<tr>
<td>Longest scaffold (Mb)</td>
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<td>BUSCO* genome score</td>
<td>C:98.6%[S:97.7%,D:0.8%], F:0.3%,M:1.1%,n:5286</td>
</tr>
</tbody>
</table>

*BUSCO scores based on the lepidoptera_odb10 BUSCO set using v5.1.2. C= complete [S= single copy, D=duplicated]; F=fragmented, M=missing, n=number of orthologues in comparison. A full set of BUSCO scores is available at https://blobtoolkit.genomehubs.org/view/ilEupSimi1.1/dataset/CAJHUZ01/busc.
sequence was assigned to 22 chromosomal-level scaffolds, representing 21 autosomes (numbered by sequence length), and the Z sex chromosome (Figure 2–Figure 5; Table 2). The assembly has a BUSCO (Simão et al., 2015) v5.1.2 completeness of 98.6% using the lepidoptera_odb10 reference set. The complete, unbroken mitochondrial genome was assembled and is 15.5 kb in length. While not fully phased, the assembly deposited is of one haplotype. Contigs corresponding to the second haplotype have also been deposited.

Methods
A single male *E. similis*, ilEupSim1, was collected from Wytham Woods, Oxfordshire (biological vice-country: Berkshire), UK (latitude 51.772, longitude -1.338) by Douglas Boyes, University of Oxford, using a light trap. The specimen was snap-frozen in dry ice using a CoolRack before transferring to the Wellcome Sanger Institute (WSI).

DNA was extracted at the Tree of Life laboratory, WSI. The ilEupSim1 sample was weighed and dissected on dry ice with tissue set aside for RNA extraction and Hi-C sequencing. Thorax/abdomen tissue was cryogenically disrupted to a fine powder using a Covaris cryoPREP Automated Dry Pulveriser, receiving multiple impacts. Fragment size analysis of 0.01-0.5 ng of DNA was then performed using an Agilent FemtoPulse. High molecular weight (HMW) DNA was extracted using the Qiagen MagAttract HMW DNA extraction kit. Low molecular weight DNA was removed from a 200-ng aliquot of extracted DNA using 0.8X AMPure XP purification kit prior to 10X Chromium sequencing; a minimum of 50 ng

**Figure 2. Genome assembly of *Euproctis similis*, ilEupSim1.1: metrics.** The BlobToolKit Snailplot shows N50 metrics and BUSCO gene completeness. The main plot is divided into 1,000 size-ordered bins around the circumference with each bin representing 0.1% of the 507,648,621 bp assembly. The distribution of scaffold lengths is shown in dark grey with the plot radius scaled to the longest scaffold present in the assembly (29,627,438 bp, shown in red). Orange and pale-orange arcs show the N50 and N90 scaffold lengths (24,449,160 and 18,514,210 bp), respectively. The pale grey spiral shows the cumulative scaffold count on a log scale with white scale lines showing successive orders of magnitude. The blue and pale-blue area around the outside of the plot shows the distribution of GC, AT and N percentages in the same bins as the inner plot. A summary of complete, fragmented, duplicated and missing BUSCO genes in the lepidoptera_odb10 set is shown in the top right. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/ilEupSim1.1/dataset/CAJHUZ01/snail.
DNA was submitted for 10X sequencing. HMW DNA was sheared into an average fragment size between 12-20 kb in a MegaRuptor 3 system with speed setting 30. Sheared DNA was purified by solid-phase reversible immobilisation using AMPure PB beads with a 1.8X ratio of beads to sample to remove the shorter fragments and concentrate the DNA sample. The concentration of the sheared and purified DNA was assessed using a Nanodrop spectrophotometer and Qubit Fluorometer and Qubit dsDNA High Sensitivity Assay kit. Fragment size distribution was evaluated by running the sample on the FemtoPulse system.

RNA was extracted from thorax/abdomen tissue in the Tree of Life Laboratory at the WSI using TRIzol (Invitrogen).
according to the manufacturer’s instructions. RNA was then eluted in 50 µl RNAse-free water and its concentration assessed using a Nanodrop spectrophotometer and Qubit Fluorometer using the Qubit RNA Broad-Range (BR) Assay kit. Analysis of the integrity of the RNA was done using Agilent RNA 6000 Pico Kit and Eukaryotic Total RNA assay.

Pacific Biosciences HiFi circular consensus and 10X Genomics Chromium read cloud sequencing libraries were constructed according to the manufacturers’ instructions. Poly(A) RNA-Seq libraries were constructed using the NEB Ultra II RNA Library Prep kit. Sequencing was performed by the Scientific Operations core at the Wellcome Sanger Institute on Pacific Biosciences SEQUEL II (HiFi), Illumina HiSeq X

**Figure 4. Genome assembly of Euproctis similis, iEupSimi1.1: cumulative sequence.** BlobToolKit cumulative sequence plot. The grey line shows cumulative length for all chromosomes. Coloured lines show cumulative lengths of chromosomes assigned to each phylum using the buscogens taxrule. An interactive version of this figure is available at [https://blobtoolkit.genomehubs.org/view/iEupSimi1.1/dataset/CAJHUZ01/cumulative](https://blobtoolkit.genomehubs.org/view/iEupSimi1.1/dataset/CAJHUZ01/cumulative).
Table 2. Chromosomal pseudomolecules in the genome assembly of *Euproctis similis*, ilEupSimi1.1.

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Figure 5. Genome assembly of *Euproctis similis*, ilEupSimi1.1: Hi-C contact map. Hi-C contact map of the ilEupSimi1.1 assembly, visualised in HiGlass. Chromosomes are arranged in size order, from left to right and top to bottom.
(10X) and Illumina HiSeq 4000 (RNA-Seq) instruments. Hi-C data were generated from head tissue using the Qiagen EpTect Hi-C kit and sequenced on HiSeq X.

Assembly was carried out with HiCanu (Nurk et al., 2020); haplo-typic duplication was identified and removed with purge_dups (Guan et al., 2020). The assembly was polished with the 10X Genomics Illumina data by aligning to the assembly with longranger align, calling variants with freebayes (Garrison & Marth, 2012). One round of the Illumina polishing was applied. Scaffolding with Hi-C data (Rao et al., 2014) was carried out with SALSA2 (Ghurye et al., 2019). The assembly was checked for contamination and corrected using the gEVAL system (Chow et al., 2016) as described previously (Howe et al., 2021). Manual curation was performed using gEVAL, HiGlass (Kerpedjiev et al., 2018) and Pretext. The mitochondrial genome was assembled using MitoHiFi (Uliano-Silva et al., 2021). The genome was analysed and BUSCO scores generated within the BlobToolKit environment (Challis et al., 2020). Table 3 contains a list of all software tool versions used, where appropriate.

The materials that have contributed to this genome note have been supplied by a Darwin Tree of Life Partner. The submission of materials by a Darwin Tree of Life Partner is subject to the Darwin Tree of Life Project Sampling Code of Practice. By agreeing with and signing up to the Sampling Code of Practice, the Darwin Tree of Life Partner agrees they will meet the legal and ethical requirements and standards set out within this document in respect of all samples acquired for, and supplied to, the Darwin Tree of Life Project. Each transfer of samples is further undertaken according to a Research Collaboration Agreement or Material Transfer Agreement entered into by the Darwin Tree of Life Partner, Genome Research Limited (operating as the WSI), and in some circumstances other Darwin Tree of Life collaborators.

Data availability
European Nucleotide Archive: Euproctis similis (yellow-tail). Accession number PRJEB42127: https://identifiers.org/ena.embl:PRJEB42127

The genome sequence is released openly for reuse. The E. similis genome sequencing initiative is part of the Darwin Tree of Life (DToL) project. All raw sequence data and the assembly have been deposited in INSDC databases. The genome will be annotated using RNAseq data and presented through the Ensembl pipeline at the European Bioinformatics Institute.

Table 3. Software tools used.

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<tr>
<th>Software tool</th>
<th>Version</th>
<th>Source</th>
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</tr>
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Raw data and assembly accession identifiers are reported in Table 1.

Acknowledgements
Members of the University of Oxford and Wytham Woods genome Acquisition Lab are listed here: https://doi.org/10.5281/zenodo.4789929.

Members of the Darwin Tree of Life Barcoding collective are listed here: https://doi.org/10.5281/zenodo.4893704.

Members of the Wellcome Sanger Institute Tree of Life programme collective are listed here: https://doi.org/10.5281/zenodo.5377053.


Members of the Tree of Life Core Informatics collective are listed here: https://doi.org/10.5281/zenodo.5013542.

Members of the Darwin Tree of Life Consortium are listed here: https://doi.org/10.5281/zenodo.4783559.

References


Reference Source

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PubMed Abstract | Publisher Full Text | Free Full Text

PubMed Abstract | Publisher Full Text

Publisher Full Text
Open Peer Review

Current Peer Review Status: ✔️ ❓ ✔️

Version 2

Reviewer Report 09 December 2021

https://doi.org/10.21956/wellcomeopenres.19033.r47561

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✔️ Thomas Blankers
Institute for Biodiversity and Ecosystem Dynamics, University of Amsterdam, Amsterdam, The Netherlands

I have no further comments.

 Competing Interests: No competing interests were disclosed.

 Reviewer Expertise: Evolutionary genomics, currently working on crickets, moths, and C. elegans

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 09 December 2021

https://doi.org/10.21956/wellcomeopenres.19033.r47560

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✔️ Steven M. Van Belleghem
Department of Biology, University of Puerto Rico-Rio Piedras, San Juan, Puerto Rico

I am happy with the changes made.

 Competing Interests: No competing interests were disclosed.

 Reviewer Expertise: Genomics, functional genomics, insects
I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

**Version 1**

Reviewer Report 24 September 2021

https://doi.org/10.21956/wellcomeopenres.18991.r45902

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Thomas Blankers

Institute for Biodiversity and Ecosystem Dynamics, University of Amsterdam, Amsterdam, The Netherlands

This data note presents the genome assembly of *Euproctis similis* as part of the Darwin Tree of Life Project. The information about the molecular lab methods is sufficient and clear. The appropriate, protocols have been used, following recommendations of the manufacturers and sequencing platforms. The quality of the genome is high, owing to the use of complementary sequencing technologies that allow for contiguous assemblies.

I only have three small suggestions for additional information, although I also see that none of these are commonly supplied in the notes coming from the Darwin Tree of Life project. First, as a biologist, I would be interested in knowing a little bit more about the organism. For example that it is a night-active moth, wide-spread across the Eurasian continent, that they're active from August to June and that they are associated with both urban and non-urban habitats and with several host plants. Second, in the presentation of the methods there are no details about the bioinformatic analyses beyond the programs that were used. It would be good to specify any deviation from default settings or even to have a brief summary of the commands used to perform the analyses. This could be done in a separate file archived along with the note or in a Table, possibly integrated in Table 3. Third, the data presentation can benefit from brief expansion of the results. The interactive figures are nice, because some explanation of what is shown can also be found at the corresponding blobtoolkit repository. However, there is no text accompanying these figures beyond a single sentence referencing the number of scaffolds and citing figures 1 through 4. Some expansion of the genome assembly statistics seems desirable. And figure 4 could use a legend as well as axis labels with the chromosome numbers. Again, I do see that other examples of notes on genomes coming from this project also do not necessarily contain these additional pieces of information, so I guess it is up to the authors to decide whether that continuity matters more or whether the details are simply not necessary.

**Is the rationale for creating the dataset(s) clearly described?**

Yes
Are the protocols appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and materials provided to allow replication by others?  
Partly

Are the datasets clearly presented in a useable and accessible format?  
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Evolutionary genomics, currently working on crickets, moths, and C. elegans

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Niclas Backström
Evolutionary Biology Program, Department of Ecology and Genetics (IEG), Uppsala University, Uppsala, Sweden

Daria Shipilina
Evolutionary Biology Program, Department of Ecology and Genetics, Uppsala University, Uppsala, Sweden

The manuscript by Boyes and Holland describes an effort to assemble both the nuclear and mitochondrial genome of a male yellow-tailed moth. The procedure includes multiple sequencing techniques allowing for both primary assembly of contigs from long-read libraries (PacBio), polishing with linked reads (10X) and scaffolding with chromosome interaction information (HiC). The highly contiguous (seven very short scaffolds, that only sum up to 0.24 Mb in total, that are not linked to chromosomes) final assembly contains 21 autosomes, the Z-chromosome and the mitochondrial genome. This will be a valuable resource for both population genetic analyses within the species/genus and comparative genomics studies in insects in general and in Lepidoptera in particular. The study sets high standards for the methodology of genome assembly by using and efficiently combining state-of-the-art sequencing technologies and novel bioinformatic pipelines.

**Minor comments:**

Abstract:
1. Maybe omit ‘majority of the’.

2. Information about that the mitochondrial genome is also assembled should be included.

3. Perhaps mention briefly that the number of assembled chromosomes is lower than many previously sequenced lepidopterans – perhaps indicating fusions in the *Euproctis* lineage?

**Introduction:**
1. A brief description of the abundance and distribution range of this species in the AABI area and globally would be very informative.

2. Perhaps it could be of interest to characterize species specific and general biological questions that can be addressed using this genomic resource.

**Genome sequence report + Methods:**
1. The first sentence in these two sections is redundant. Perhaps it is sufficient to describe the sampling location, collector, sample ID and date of sampling (missing) in the methods section and omit the redundant details in the GSR section.

**Genome sequence report:**
1. Omit ‘The majority,’?

2. Add information about that the mitochondrial genome was assembled (including assessment of completeness, e. g. exact length of the scaffold)?

3. The finding that scaffolds are anchored to only 21 autosomes is a bit surprising given that most lepidopterans sequenced so far have 31 chromosome pairs. This could be another example of chromosome rearrangements (here potentially fusions, similar to what has been observed in e.g. *Heliconius melpomene*)? This might be worth mentioning briefly.

4. Were RNAseq sequencing reads used in the downstream analysis?

5. Table 1. Unclear if the “Number of contigs” corresponds to the number of contigs before or after the manual correction of the Hi-C map?

**Methods:**
1. ‘specimen was’ instead of ‘specimens were’.

2. Second paragraph. The sample ID can be given in the first part of the section where sample location, collector, date of sampling etc. is stated.

3. 4th line, left column under Figure 1. ‘its concentration RNA assessed’ should perhaps be ‘the RNA concentration assessed’.

4. Information about RNAseq library preparation method, sequencing technique applied and data availability of RNAseq reads is missing.

**Reflection:**
1. An annotation with the RNAseq data generated here in combination with previously available lepidopteran gene models would add useful information for downstream analyses using this resource.

**Is the rationale for creating the dataset(s) clearly described?**
Yes

**Are the protocols appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and materials provided to allow replication by others?**
Yes

**Are the datasets clearly presented in a useable and accessible format?**
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Comparative genomics, population genetics, molecular evolution.

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however we have significant reservations, as outlined above.

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**Abstract:**
- I suggest using a more precise alternative to the word ‘majority’.

**Introduction:**
- Is it possible to give more precise information on the geographic distribution of this
species?

Results:
○ I do not see any results on the RNA-seq data. Was a transcriptome assembly performed? I also do not see accession numbers to these data in Table 1.

○ An assessment of the repetitiveness (TE content) of the genome could be useful.

Is the rationale for creating the dataset(s) clearly described?
Yes

Are the protocols appropriate and is the work technically sound?
Yes

Are sufficient details of methods and materials provided to allow replication by others?
Yes

Are the datasets clearly presented in a useable and accessible format?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Genomics, functional genomics, insects

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Comments on this article

Author Response 07 Dec 2021
Jonathan Threlfall, Wellcome Trust Sanger Institute, UK

Thanks to each of the reviewers for their constructive feedback. Comments like this are being used to update and improve the template we use for each genome note, so are of considerable value to us.

Major updates:

We have updated the Introduction section to include information about abundance, habitat and distribution of the species, and a description of potential uses for the genome. Also included is a brief mention of lepidopteran karyotypes - a paper discussing lepidopteran chromosome evolution
using the sequences generated by this project is forthcoming.

We addressed the issue of the missing RNA-Seq data, which will be used for annotation by Ensembl in the near future as part of the Darwin Tree of Life project pipeline. We have also included details of the method of library preparation.

An image of the specimen used for genome sequencing has been included as Figure 1. The legends of Figures 2-5 have been expanded to make them easier to understand.

**Other comments:**
The number of contigs refers to those making up the final assembly, i.e. after manual curation of the genome using the Hi-C map and other information.

A step-by-step protocol describing the pipeline used for genome assembly is currently in preparation and should be released shortly.

**Competing Interests:** No competing interests were disclosed.