DATA NOTE

The genome sequence of the garden grass-veneer, *Chrysoteuchia culmella* (Linnaeus, 1758) [version 1; peer review: 2 approved, 1 approved with reservations]

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**Abstract**

We present a genome assembly from an individual male *Chrysoteuchia culmella* (the garden grass-veneer; Arthropoda; Insecta; Lepidoptera; Crambidae). The genome sequence is 645 megabases in span. The majority of the assembly (99.81%) is scaffolded into 31 chromosomal pseudomolecules with the Z sex chromosome assembled. The complete mitochondrial genome was also assembled and is 15.4 kilobases in length. Gene annotation of this assembly on Ensembl has identified 21,251 protein coding genes.

**Keywords**

Chrysoteuchia culmella, garden grass-veneer, genome sequence, chromosomal, Crambidae

This article is included in the Tree of Life gateway.
Corresponding author: Darwin Tree of Life Consortium (mark.blaxter@sanger.ac.uk)

Author roles: Boyes D: Investigation, Resources; Parkerson L: Writing – Original Draft Preparation;

Competing interests: No competing interests were disclosed.

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Species taxonomy
Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia; Pyraloidea; Crambidae; Crambinae; Chrysoteuchia; Chrysoteuchia culmella (Linnaeus, 1758) (NCBI:txid1594250).

Background
The garden grass-veneer, Chrysoteuchia culmella (Linnaeus, 1758), is a micro moth of the Crambinae subfamily. It is common in grassland, rough meadows and gardens throughout much of Europe, including the British Isles (“Chrysoteuchia Culmella (Linnaeus, 1758)” 2010). It is recognised by its angled subterminal line, golden metallic cilia and size, with a wing-span of 20–24mm (“Chrysoteuchia Culmella (Garden Grass-Veneer)” 2012). The eggs are laid on various grasses and, after hatching, the larvae feed from September to April on the stem bases of grasses. After pupating in May from a cocoon near the ground, the species is on the wing from mid-May to mid-September (and occasionally until late October). During this time, it can be readily disturbed from grasses during the day and attracted to light during the night (Langmaid et al., 2018).

C. culmella larvae are frequent hosts of the endoparasitic larvae of Eriothrix rufomaculata, a parasitoid species of fly (Paston & Rotheray, 2009). We present a complete genome assembly for C. culmella as part of the Darwin Tree of Life project, Wellcome Sanger Institute, aiming to sequence the genomes of 70,000 species of eukaryotic organisms in Britain and Ireland.

Genome sequence report
The genome was sequenced from a single male C. culmella (ilChrCulm1) collected from Wytham Woods, Berkshire, UK (Figure 1). A total of 43-fold coverage in Pacific Biosciences single-molecule HiFi long reads and 56-fold coverage in 10X Genomics read clouds were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. Manual assembly curation corrected 18 missing/misjoins and removed 2 haplotypic duplications, reducing the assembly size by 0.64% and the scaffold number by 17.39%, and increasing the scaffold N50 by 4.73%.

The final assembly has a total length of 645 Mb in 57 sequence scaffolds with a scaffold N50 of 22.8 Mb (Table 1). The majority, 99.81%, of the assembly sequence was assigned to 31 chromosomal-level scaffolds, representing 30 autosomes (numbered by sequence length) and the Z sex chromosome (Figure 2–Figure 5; Table 2). The assembly has a BUSCO v5.13.2 (Manni et al., 2021) completeness of 98.6% (single 98.3%, duplicated 0.3%) using the

| Table 1. Genome data for Chrysoteuchia culmella, ilChrCulm1.1. |
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| **Project accession data** |
| Assembly identifier | ilChrCulm1.1 |
| Species | Chrysoteuchia culmella |
| Specimen | ilChrCulm1 (genome assembly); ilChrCulm2 (Hi-C, RNA-Seq) |
| NCBI taxonomy ID | 1594250 |
| BioProject | PRJEB45126 |
| BioSample ID | SAMEA7701502 |
| Isolate information | Male, whole organism (ilChrCulm1); Whole organism tissue, unknown sex (ilChrCulm2) |
| **Raw data accessions** |
| PacificBiosciences SEQUEL II | ERR6406210 |
| 10X Genomics Illumina | ERR6054801-ERR6054804 |
| Hi-C Illumina | ERR6054805 |
| PolyA RNA-Seq Illumina | ERR9434977 |
| **Genome assembly** |
| Assembly accession | GCA_910589605.1 |
| Accession of alternate haplotype | GCA_910589405.1 |
| Span (Mb) | 645 |
| Number of contigs | 89 |
| Contig N50 length (Mb) | 16.4 |
| Number of scaffolds | 57 |
| Scaffold N50 length (Mb) | 22.8 |
| Longest scaffold (Mb) | 26.47 |
| BUSCO* genome score | C:98.6%;S:98.3%,D:0.3%,F:0.5%,M:0.9%,n:5,286 |

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Figure 1. Image of the Chrysoteuchia culmella specimen taken prior to preservation and processing.
lepidoptera_odb10 reference set (n=5,286). While not fully phased, the assembly deposited is of one haplotype. Contigs corresponding to the second haplotype have also been deposited.

### Genome annotation report

The ilChrCulm1.1 genome has been annotated using the Ensembl rapid annotation pipeline (Table 1; https://rapid.ensembl.org/Chrysoteuchia_culmella_GCA_910589605.1). The resulting annotation includes 21,475 transcribed mRNAs from 21,251 protein-coding genes.

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**Figure 2.** Genome assembly of *Chrysoteuchia culmella*, ilChrCulm1.1: 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 645,226,651 bp assembly. The distribution of chromosome lengths is shown in dark grey with the plot radius scaled to the longest chromosome present in the assembly (31,066,094 bp, shown in red). Orange and pale-orange arcs show the N50 and N90 chromosome lengths (22,805,156 and 13,759,558 bp), respectively. The pale grey spiral shows the cumulative chromosome 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/ilChrCulm1.1/dataset/CAJUUR01/snail.

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**Methods**

### Sample acquisition and nucleic acid extraction

Two *C. culmella* specimens (ilChrCulm1, genome assembly; ilChrCulm2, Hi-C and RNA-Seq) were collected using a light trap from Wytham Woods, Berkshire, UK (latitude 51.772, longitude -1.338) by Douglas Boyes (University of Oxford). The specimens were identified by Douglas Boyes snap-frozen on dry ice.

DNA was extracted at the Tree of Life laboratory, Wellcome Sanger Institute. The ilChrCulm1 sample was weighed and
dissected on dry ice. Whole organism tissue was disrupted using a Nippi Powermasher fitted with a BioMasher pestle. 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 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

**Figure 3.** Genome assembly of *Chrysoteuchia culmella*, ilChrCulm1.1: GC coverage. BlobToolKit GC-coverage plot. Scaffolds are coloured by phylum. Circles are sized in proportion to scaffold length. Histograms show the distribution of scaffold length sum along each axis. An interactive version of this figure is available at [https://blobtoolkit.genomehubs.org/view/ilChrCulm1.1/dataset/CAJUUR01.1/blob](https://blobtoolkit.genomehubs.org/view/ilChrCulm1.1/dataset/CAJUUR01.1/blob).
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 whole organism tissue of ilChrCulm2 in the Tree of Life Laboratory at the WSI using TRIZol,
Table 2. Chromosomal pseudomolecules in the genome assembly of *Chrysoteuchia culmella*, ilChrCulm1.1.

| INSDC accession | Chromosome | Size (Mb) | GC%  |
|-----------------|------------|-----------|------|
| OU342642.1      | 1          | 26.47     | 36.5 |
| OU342643.1      | 2          | 26.39     | 36.4 |
| OU342644.1      | 3          | 26.29     | 36.2 |
| OU342645.1      | 4          | 25.12     | 36   |
| OU342646.1      | 5          | 25.07     | 36.4 |
| OU342647.1      | 6          | 24.95     | 36.1 |
| OU342648.1      | 7          | 24.75     | 36.1 |
| OU342649.1      | 8          | 24.7      | 36.2 |
| OU342650.1      | 9          | 24.46     | 36.4 |
| OU342651.1      | 10         | 23.8      | 36.3 |
| OU342652.1      | 11         | 23.69     | 36.8 |
| OU342653.1      | 12         | 22.81     | 36.2 |
| OU342654.1      | 13         | 22.6      | 36.2 |
| OU342655.1      | 14         | 21.77     | 36.4 |
| OU342656.1      | 15         | 21.63     | 36.3 |

Figure 5. Genome assembly of *Chrysoteuchia culmella*, ilChrCulm1.1: Hi-C contact map. Hi-C contact map of the ilChrCulm1.1 assembly, visualised in HiGlass. Chromosomes are arranged in size order from left to right and top to bottom. The interactive Hi-C map can be viewed at https://genome-note-higlass.tol.sanger.ac.uk/l/?d=PizMaC2sTVqZKsjMxz2y9A.
according to the manufacturer’s instructions. RNA was then eluted in 50 μl RNase-free water and its concentration RNA 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.

Sequencing
Pacific Biosciences HiFi circular consensus and 10X Genomics Chromium read cloud sequencing libraries were constructed according to the manufacturers’ instructions. Sequencing was performed by the Scientific Operations core at the Wellcome Sanger Institute on Pacific Biosciences SEQUEL II (HiFi). Illumina NovaSeq 6000 (10X) and Illumina HiSeq 4000 (RNA-Seq) instruments. Hi-C data were generated in the Tree of Life laboratory from remaining tissue of iChrCulm2 using the Arima v2 kit and sequenced on a NovaSeq 6000 instrument.

Genome assembly
Assembly was carried out with Hifiasm (Cheng et al., 2021); haplotypic duplication was identified and removed with purge_dups (Guan et al., 2020). One round of polishing was performed by aligning 10X Genomics read data to the assembly with longeranger align, calling variants with freebayes (Garrison & Marth, 2012). The assembly was then scaffolded with Hi-C data (Rao et al., 2014) using 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 (Howe et al., 2021) was performed using gEVAL, HiGlass (Kerpedjiev et al., 2018) and Pretext. The mitochondrial genome was assembled using MitoHiFi (Uliano-Silva et al., 2021), which performs annotation using Mitofinder (Allio et al., 2020). 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.

Table 3. Software tools used.

| Software tool | Version | Source |
|---------------|---------|--------|
| Hifiasm       | 0.14    | Cheng et al., 2021 |
| purge_dups    | 1.2.3   | Guan et al., 2020 |
| SALSA2        | 2.2     | Ghurye et al., 2019 |
| longeranger align | 2.2.2   | https://support.10xgenomics.com/genome-exome/software/pipelines/latest/advanced/other-pipelines |
| freebayes     | 1.3.1-17-gaa2ace8 | Garrison & Marth, 2012 |
| MitoHiFi      | 2.0     | Uliano-Silva et al., 2021 |
| HiGlass       | 1.11.6  | Kerpedjiev et al., 2018 |
| PretexView    | 0.2.x   | https://github.com/wtsi-hpag/PretexView |
| BlobToolKit   | 3.2.6   | Challis et al., 2020 |

Genome annotation
The Ensembl gene annotation system (Aken et al., 2016) was used to generate annotation for the Chrysoteuchia culmella assembly (GCA_910589605.1). Annotation was created primarily through alignment of transcriptomic data to the genome, with gap filling via protein-to-genome alignments of a select set of proteins from UniProt (UniProt Consortium, 2019).

Ethics/compliance issues
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 Wellcome Sanger Institute), and in some circumstances other Darwin Tree of Life collaborators.

Data availability
European Nucleotide Archive: Chrysoteuchia culmella (garden grass-veneer). Accession number PRJEB45126; https://identifiers.org/ena.embl/PRJEB45126.

The genome sequence is released openly for reuse. The C. culmella 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. Raw data and assembly accession identifiers are reported in Table 1.

Author information
Members of the University of Oxford and Wytham Woods Genome Acquisition Lab are listed here: https://doi.org/10.5281/zenodo.6418202.

Members of the Darwin Tree of Life Barcoding collective are listed here: https://doi.org/10.5281/zenodo.6418156.

Members of the Wellcome Sanger Institute Tree of Life programme are listed here: https://doi.org/10.5281/zenodo.6866293.

Members of Wellcome Sanger Institute Scientific Operations: DNA Pipelines collective are listed here: https://doi.org/10.5281/zenodo.5746904.

Members of the Tree of Life Core Informatics collective are listed here: https://doi.org/10.5281/zenodo.6125046.

Members of the Darwin Tree of Life Consortium are listed here: https://doi.org/10.5281/zenodo.6418363.
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Shuai Zhan
University of the Chinese Academy of Sciences,, Beijing, China

In this manuscript, the authors report a genoe reference for the garden grass-veneer moth, *Chrysoteuchia culmella*. The genome reference was assembled in a high level of contiguity and fully anchored to 31 chromosomes. The mitochondrial genome was also assembled in a reasonable length. Overall, the manuscript was clearly written. I have only concern in the method, in which the authors did not provide detailed information and some of them were simply given a previous reference. No other audience could repeat the assembly with currently provided methods. Also, it's unclear to me how chromosome Z was assigned.

*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?*
No

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

*Competing Interests:* No competing interests were disclosed.

*Reviewer Expertise:* Genomics; Evolution.

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.
Ma Cristina Del Rincón-Castro
Universidad de Guanajuato,, Campus Irapuato-Salamanca, Mexico

The article deals with genome assembly from a male Chrysoteuchia culmella (the garden grass-veen; Arthropoda; Insecta; Lepidoptera; Crambidae). The authors report that the genome sequence is 645 megabases in span. The majority of the assembly (99.81%) is scaffolded into 31 chromosomal pseudomolecules with the Z sex chromosome assembled. The article presents the complete information for a database of genomic data on this insect, its approval is recommended.

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?
Yes

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

**Reviewer Expertise:** Molecular biology of baculoviruses as biological control agents

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.
Museum für Naturkunde Berlin, Leibniz-Institut für Evolutions- und Biodiversitätsforschung
Invalidenstr, Berlin, Germany

Josh Penalba
Museum für Naturkunde Berlin, Leibniz-Institut für Evolutions- und Biodiversitätsforschung
Invalidenstr, Berlin, Germany

This short and concise data-release article provides a complete genome of the garden grass-veneer. The molecular and data analysis methods look sound to me (but as I am no expert in genomics I leave this part to my colleague). I would only have two comments here:

1. Identification of the specimens should be confirmed through blasting the COI DNA barcode against BOLD or GenBank, as worn specimens could be misidentified (similar to Thysanotia, Chrysocrambus or some Catoptria). It looks to me as C. culmella indeed but a double-check is worth it.

2. The "genome annotation" part is a bit short, and it is not stated whether all genes the Ensembl gene set were found in the present genome. This part could be completed for the reader.

Here are the comments of the co-reviewer:

1. Page 4. Col. 1. “While not fully phased, the assembly deposited is of one haplotype” – This isn’t entirely clear. Does this mean for SNPs which were not phased a random allele was chosen for the haplotype?

2. Page 5. Col. 1. “Fragment size analysis of 0.01-0.5ng of DNA...” The placement of this statement is not clear because it precedes the DNA extraction. Is there another source of DNA this is referring to or should this be later?

3. Page 8. Col1. Genome assembly. “Assembly was carried out with Hifiasm” Specify that the only data that was used here was the PacBio HiFi and the 10X Genomics was only used for the polishing.

We both agree that the manuscript can be considered for indexing once these points are addressed.

**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?**
Yes

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

**Reviewer Expertise:** I work on the systematics and phylogeny of pyraloid moths
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.