DATA NOTE

The genome sequence of the two-banded wasp hoverfly, *Chrysotoxum bicinctum* (Linnaeus, 1758) [version 1; peer review: 2 approved]

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Abstract
We present a genome assembly from an individual female *Chrysotoxum bicinctum* (the two-banded wasp hoverfly; Arthropoda; Insecta; Diptera; Syrphidae). The genome sequence is 913 megabases in span. The majority of the assembly (98.81%) is scaffolded into five chromosomal pseudomolecules, with the X sex chromosome assembled.

Keywords
Chrysotoxum bicinctum, two-banded wasp hoverfly, genome sequence, chromosomal, Diptera

This article is included in the Tree of Life gateway.

Any reports and responses or comments on the article can be found at the end of the article.
Species taxonomy
Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha; Syrphoidea; Syrphidae; Syrphinae; Syrphini; Chrysotoxum; Chrysotoxum bicinctum (Linnaeus 1758) (NCBI: txid323313).

Background
Chrysotoxum bicinctum, the two-banded wasp hoverfly, is one of Britain’s most distinctive hoverflies. Its chocolate-coloured wing markings and bright yellow bars on the second and fourth abdominal segments make this fly unmistakable in the field (Ball et al., 2015; van Veen, 2010). The genus Chrysotoxum are large wasp-mimic hoverflies with long, elegant antennae and consist of more than 110 species (Masetti et al., 2006). This wasp mimicry likely gives protection against predation by birds through batesian mimicry (Leavey et al., 2021). Across their flight period of May to September, this species is common across southern Britain but its abundance decreases with northerly latitude.

Genome sequence report
The genome was sequenced from a single female C. bicinctum (Figure 1) collected from Wytham Great Wood, Oxfordshire, UK (latitude 51.769, longitude -1.33). A total of 29-fold coverage in Pacific Biosciences single-molecule long reads and 37-fold coverage in 10X Genomics read clouds (from molecules with an estimated N50 of 60 kb) were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. Manual assembly curation corrected 326 missing/misjoins and removed 87 haplotypic duplications, reducing the assembly length by 3.27% and the scaffold number by 64.06%, and increasing the scaffold N50 by 644.27%.

The final assembly has a total length of 913 Mb in 92 sequence scaffolds with a scaffold N50 of 118 Mb (Table 1). The majority, 98.81%, of the assembly sequence was assigned to 5 chromosomal-level scaffolds, representing 4 autosomes (numbered by sequence length), and the X sex chromosome (Figure 2–Figure 5; Table 2). The assembly has a BUSCO (Simão et al., 2015) completeness of 96.6% (single 95.5%, duplicated 1.1%) using the diptera_odb10 reference set. While not fully phased, the assembly deposited is of one haplotype. Contigs corresponding to the second haplotype have also been deposited.

Methods
Sample acquisition and nucleic acid extraction
A female (idChrBici1) C. bicinctum was collected from Wytham Great Wood, Oxfordshire, UK (latitude 51.769, longitude -1.33) by Will Hawkes, University of Exeter, who also identified the sample. A second sample of unknown sex (idChrBici2), was collected by Matt Smith from Hartslock Reserve, Oxfordshire, UK (latitude 51.511263, longitude -1.112222). The samples were collected using a net and snap-frozen on dry ice.

DNA was extracted from the whole organism of idChrBici1 at the Wellcome Sanger Institute (WSI) Scientific Operations core from head/thorax tissue using the Qiagen MagAttract HMW DNA kit, according to the manufacturer’s instructions. Following this, further DNA was extracted for a PacBio top-up. Tissue was cryogenically disrupted to a fine powder using a Covaris cryoPREP Automated Dry Pulveriser, receiving multiple...
Table 1. Genome data for *Chrysotoxum bicinctum*, idChrBici1.1.

| Project accession data                  |                                                                 |
|-----------------------------------------|------------------------------------------------------------------|
| Assembly identifier                     | idChrBici1.1                                                     |
| Species                                 | *Chrysotoxum bicinctum*                                           |
| Specimen                                | idChrBici1 (genome assembly, Hi-C); idChrBici2 (Hi-C, RNA-Seq)   |
| NCBI taxonomy ID                        | 323313                                                           |
| BioProject                              | PRJEB45198                                                       |
| BioSample ID                            | SAMEA7520032                                                     |
| Isolate information                     | Female, head/thorax, abdomen (idChrBici1); Unknown sex, head/thorax, abdomen (idChrBici2) |

| Raw data accessions                     |                                                                 |
|-----------------------------------------|------------------------------------------------------------------|
| PacificBiosciences SEQUEL II            | ERR6412376, ERR6558188                                           |
| 10X Genomics Illumina                   | ERR6054965-ERR6054968                                            |
| Hi-C Illumina                           | ERR6054969-ERR6054971                                            |
| Illumina polyA RNA-Seq                  | ERR6464930                                                       |

| Genome assembly                         |                                                                 |
|-----------------------------------------|------------------------------------------------------------------|
| Assembly accession                      | GCA_911387755.1                                                  |
| Accession of alternate haplotype        | GCA_911387745.1                                                  |
| Span (Mb)                               | 913                                                              |
| Number of contigs                       | 412                                                              |
| Contig N50 length (Mb)                  | 5.7                                                              |
| Number of scaffolds                     | 92                                                               |
| Scaffold N50 length (Mb)                | 265.8                                                            |
| Longest scaffold (Mb)                   | 269.7                                                            |
| BUSCO* genome score                     | C:96.6%(S:95.5%,D:1.1%),F:0.8%,M:2.6%,n:3285                    |

*BUSCO scores based on the diptera_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/idChrBici1.1/dataset/CAJVQW01/busco.

### Fragment size analysis

Fragment size analysis of 0.01–0.5 ng of DNA was then performed using an Agilent FemtoPulse. High molecular weight (HMW) DNA was again extracted using the Qiagen MagAttract HMW DNA extraction kit. 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 tissue of idChrBici2 in the Tree of Life Laboratory at the Wellcome Sanger Institute 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.
Sequencing
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 (10X) and Illumina HiSeq 4000 (RNA-Seq) instruments. Hi-C data were generated from abdomen tissue of idChrBici1, and head and abdomen tissue of idChrBici2 using the Arima Hi-C+ kit and sequenced on HiSeq X (idChrBici1) and Illumina NovaSeq 6000 instruments (idChrBici2).

Figure 2. Genome assembly of Chrysotoxum bicinctum, idChrBici1.1: 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 912,938,338 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 (269,711,166 bp, shown in red). Orange and pale-orange arcs show the N50 and N90 chromosome lengths (265,788,494 and 117,573,787 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 diptera_odb10 set is shown in the top right. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/idChrBici1.1/dataset/CAJVQW01/snaill.
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) with the -e flag. One round of polishing was performed by aligning 10X Genomics read data to the assembly with longranger align, calling variants with

Figure 3. Genome assembly of Chrysotoxum bicinctum, idChrBici1.1: 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/idChrBici1.1/dataset/CAJVQW01/blob.
The grey line shows cumulative length for all scaffolds. Coloured lines show cumulative lengths of scaffolds assigned to each phylum using the busco genes taxrule. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/idChrBici1.1/dataset/CAJVQW01/cumulative.

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 was performed using gEVAL, HiGlass (Kerpedjieva et al., 2018) and Pretext. The mitochondrial genome was assembled using MitoHiFi (Uliano-Silva et al., 2021) and annotated with 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.
Figure 5. Genome assembly of *Chrysotoxum bicinctum*, idChrBici1.1: Hi-C contact map. Hi-C contact map of the idChrBici1.1 assembly, visualised in HiGlass.

Table 2. Chromosomal pseudomolecules in the genome assembly of *Chrysotoxum bicinctum*, idChrBici1.1.

| INSDC accession | Chromosome | Size (Mb) | GC%  |
|-----------------|------------|-----------|------|
| OU426987.1      | 1          | 269.71    | 34.2 |
| OU426988.1      | 2          | 265.79    | 34.1 |
| OU426989.1      | 3          | 233.72    | 33.9 |
| OU426990.1      | 4          | 117.57    | 34   |
| OU426991.1      | X          | 14.54     | 33.6 |
| OU426992.1      | MT         | 0.02      | 17.5 |
| -               | Unplaced   | 11.59     | 36.8 |
Data availability
European Nucleotide Archive: Chrysotoxum bicinctum (two-bodied wasp hoverfly). Accession number PRJEB45198; https://identifiers.org/ena.embl/PRJEB45198. The genome sequence is released openly for reuse. The C. bicinctum 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 the RNA-Seq data and presented through the Ensembl pipeline at the European Bioinformatics Institute. Raw data and assembly accession identifiers are reported in Table 1.

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

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

Members of Wellcome Sanger Institute Scientific Operations: DNA Pipelines collective are listed here: https://doi.org/10.5281/zenodo.4790456.

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.

Table 3. Software tools used.

| Software tool   | Version | Source                        |
|-----------------|---------|-------------------------------|
| Hifiasm         | 0.15    | Cheng et al., 2021            |
| purge_dups      | 1.2.3   | Guan et al., 2020             |
| SALSA2          | 2.2     | Ghurye et al., 2019           |
| longranger 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     | Ulliano-Silva et al., 2021    |
| gEVAL           | N/A     | Chow et al., 2016             |
| HiGlass         | 1.11.6  | Kerpedjiev et al., 2018       |
| PretextView     | 0.2.x   | https://github.com/wtsi-hpag/PretextView |
| BlobToolKit     | 2.6.2   | Challis et al., 2020          |

References
Allio R, Schomaker-Bastos A, Romiguier J, et al.: MitoFinder: Efficient automated large-scale extraction of mitogenomic data in target enrichment phylogenomics. Mol Ecol Resour. 2020; 20(4): 892–905. PubMed Abstract | Publisher Full Text | Free Full Text

Ball S, Ball SG, Morris R: Britain's Hoverflies: A Field Guide - Revised and Updated Second Edition. Princeton University Press. 2015. Reference Source

Challis R, Richards E, Rajan J, et al.: BlobToolKit—Interactive Quality Assessment of Genome Assemblies. G3 (Bethesda). 2020; 10(4): 1361-74. PubMed Abstract | Publisher Full Text | Free Full Text

Cheng H, Concepcion GT, Feng X, et al.: Haplotype-Resolved de Novo Assembly Using Phased Assembly Graphs with Hifiasm. Nat Methods. 2021; 18(2): 170-75. PubMed Abstract | Publisher Full Text | Free Full Text

Chow W, Brugger K, Caccamo M, et al.: gEVAL — a Web-Based Browser for Evaluating Genome Assemblies. Bioinformatics. 2016; 32(16): 250810. PubMed Abstract | Publisher Full Text | Free Full Text

Garrison E, Marth G: Haplotype-Based Variant Detection from Short-Read Sequencing. arXiv: 1207.3907. 2012. Reference Source

Ghurye J, Rhie A, Walenz BP, et al.: Integrating Hi-C Links with Assembly Graphs for Chromosome-Scale Assembly. PLoS Comput Biol. 2019; 15(6): e1007273. PubMed Abstract | Publisher Full Text | Free Full Text
Guan D, McCarthy SA, Wood J, et al.: Identifying and Removing Haplotypic Duplication in Primary Genome Assemblies. Bioinformatics. 2020; 36(9): 2896–98. PubMed Abstract | Publisher Full Text | Free Full Text

Howe K, Chow W, Collins J, et al.: Significantly Improving the Quality of Genome Assemblies through Curation. GigaScience. 2021; 10(1): giaa153. PubMed Abstract | Publisher Full Text | Free Full Text

Kerpedjiev P, Abdennur N, Leksschas F, et al.: HiGlass: Web-Based Visual Exploration and Analysis of Genome Interaction Maps. Genome Biol. 2018; 19(1): 125. PubMed Abstract | Publisher Full Text | Free Full Text

Leavey A, Taylor CH, Symonds MRE, et al.: Mapping the Evolution of Accurate Batesian Mimicry of Social Wasps in Hoverflies. Evolution. 2021. PubMed Abstract | Publisher Full Text

Masetti A, Luchetti A, Sommaggio D, et al.: Phylogeny of Chrysotoxum Species (Diptera: Syrphidae) Inferred from Morphological and Molecular Characters. Ceske Budéjovice. 2006. Reference Source

Rao SSP, Huntley MH, Durand NC, et al.: A 3D Map of the Human Genome at Kilobase Resolution Reveals Principles of Chromatin Looping. Cell. 2014; 158(7): 1665–80. PubMed Abstract | Publisher Full Text | Free Full Text

Rotheray GE, Gilbert FS: The Phylogeny and Systematics of European Predacious Syrphidae (Diptera) Based on Larval and Puparial Stages. Zool J Linn Soc. 1989; 95(1): 29–70. Publisher Full Text

Simão FA, Waterhouse RW, Ioannidis P, et al.: BUSCO: Assessing Genome Assembly and Annotation Completeness with Single-Copy Orthologs. Bioinformatics. 2015; 31(19): 3210–12. PubMed Abstract | Publisher Full Text

Speight MCD: Puparium of Chrysotoxum Festivum (L.)(Diptera: Syrphidae). Entomologist’s Record and Journal of Variation. 1976. Reference Source

Uliano-Silva M, Nunes JGF, Krasheninnikova K, et al.: marcelauliano/MitoHIFI: mitohifi_v2.0. 2021. Reference Source

Van Veen MP: Hoverflies of Northwest Europe: Identification Keys to the Syrphidae. BRILL. 2010. Reference Source
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Hawkes and colleagues present a genome report for the de novo assembly of a reference genome for *Chrysotoxum bicinctum*, the two-banded wasp hoverfly. Authors provide a good description on the ecology and distribution of the species. Following a state-of-the-art procedure based on Pacbio HiFi, 10X genomics and Hi-C sequencing, the final assembly reaches chromosomal resolution including 4 autosomes and one sex chromosome. The assembly is partially phased.

Despite the availability of RNA-Seq data, the nuclear assembly has not been structurally annotated. This step might be valuable to add now or in close future. Data is available at the European Nucleotide Archive.

**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:** ecological genomics, population genomics

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.
This data note reports the genome of the two-banded wasp hoverfly *Chrysotoxum bicinctum*. The hoverflies are very interesting species of insects. Genomes of hoverflies are usually large and difficult to assemble. I am happy to see the publication of this hoverfly genome. The genome was assembled using long reads, linked reads, and Hi-C technologies. Data were well analyzed using available methods and software. The results will provide invaluable resources for hoverflies’ studies and references for other relative species’ genome assembly. I have some minor comments.

1. The genome was assembled into five pseudochromosomes with the X sex chromosome. What is the karyotype of this species or other relative species? How did the authors make sure the identified X chromosome is correct? There is a lack of analysis of the X chromosome to confirm the identification.

2. Sampling information was repeated in “Genome sequence report” and “Methods” sections. Remove one of the duplicated contents.

3. I noticed that many figures of this manuscript were generated by BlobToolKit Pipeline and used directly. However, there are many format problems in these figures. I suggest the author revise these figures carefully. Some typical issues: Figure 3, change “gc” to “GC content”, sum to “Sum”, “total” to “Total”, “no-hit” to “No-hit”. The same problems were found in other figures where the first letters were not capitalized for the first word. Figure 5, length in Mb should be marked in the axes.

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 insect genomics, population genetics, and pest control research. I used population genomics approaches to trace insects' origin, invasion routes, long-distance migration, and local adaptation to pesticide and environmental stresses.

**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.**