The genome sequence of the common red soldier beetle, *Rhagonycha fulva* (Scopoli, 1763) [version 1; peer review: 2 approved]

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Abstract

We present a genome assembly from an individual female *Rhagonycha fulva* (the common red soldier beetle; Arthropoda; Insecta; Coleoptera; Cantharidae). The genome sequence is 425 megabases in span. The majority of the assembly is scaffolded into seven chromosomal pseudomolecules, with the X sex chromosome assembled.

Keywords

*Rhagonycha fulva*, common red soldier beetle, genome sequence, chromosomal
**Species taxonomy**

Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Coleoptera; Polyphaga; Elateriformia; Elateroidea; Cantharidae; Cantharinae; Rhagonycha; *Rhagonycha fulva* (Scopoli, 1763) (NCBI: txid100007).

**Introduction**

The common red soldier beetle, *Rhagonycha fulva*, is the most abundant and widespread soldier beetle (Coleoptera: Cantharidae) in the UK. They can be found in a variety of habitats, where adults are frequently encountered on the flowers of umbellifers (Apiaceae), thistles (Asteraceae) and ragwort (*Senecio jacobaea*) throughout the summer. It can be particularly abundant on the flowers of common hogweed, *Heracleum sphondylium* (Grace & Nelson, 1981), and their association with flowers indicates this species’ potential role as an important pollinator. Adults are predatory on small insects, but also feed extensively on floral resources. They are diurnal and fly readily, males in particular are highly mobile (Rodwell *et al.*, 2018). Mating occurs over a prolonged period of time, meaning female-male pairs are often encountered in copulation. Eggs are laid into the soil and the larvae are predatory, hunting amongst the leaf litter.

Adults can be easily recognised by the extensive reddish colour of the entire body with black tips to the elytra and black tarsi, antennae and palps.

**Genome sequence report**

The genome was sequenced from one female *R. fulva* collected from Wytham farm, Oxfordshire (biological vice-county: Berkshire), UK (latitude 51.779, longitude -1.317). A total of 41-fold coverage in Pacific Biosciences single-molecule long reads and 103-fold coverage in 10X Genomics read clouds were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. Manual assembly curation corrected 73 missing/misjoins and removed 12 haplotypic duplications, reducing the assembly length by 1.54% and the scaffold number by 84.62%, and increasing the scaffold N50 by 238.56%. The final assembly has a total length of 425 Mb in 13 sequence scaffolds with a scaffold N50 of 116 Mb (Table 1). The majority, 99.97%, of the assembly sequence was assigned to seven chromosomal-level scaffolds, representing six autosomes (numbered by sequence length), and the X sex chromosome (Figure 1–Figure 4; Table 2). The assembly has a BUSCO v5.1.2 (Simão *et al.*, 2015) completeness of 98.9% using the endopterygota_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**

A single female *R. fulva* was collected from Wytham farm, Oxfordshire (biological vice-county: Berkshire), UK (latitude 51.779, longitude -1.317) by Liam Crowley, University of Oxford, and snap-frozen on dry ice using a CoolRack. A second specimen of unknown sex, icRhaFulv4, was collected from Wigmore Park, Luton, UK (latitude 51.88378, longitude -0.3681422) by Olga Sivell, Natural History Museum, and snap-frozen on dry ice.

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**Table 1. Genome data for *Rhagonycha fulva*, icRhaFulv1.1.**

| **Project accession data** |  |
|----------------------------|---|
| Assembly identifier        | icRhaFulv1.1 |
| Species                   | *Rhagonycha fulva* |
| Specimen                  | icRhaFulv1 |
| NCBI taxonomy ID          | NCBI:txid41101 |
| BioProject                | PRJEB43742 |
| BioSample ID              | SAMEA7520319 |
| Isolate information       | Female, whole organism |

| **Raw data accessions** |  |
|-------------------------|---|
| PacificBiosciences SEQUEL II | ERR6606788 |
| 10X Genomics Illumina    | ERR6054565-ERR6054568 |
| Hi-C Illumina           | ERR6054569 |
| RNAseq PolyA Illumina   | ERR6286718 |

| **Genome assembly** |  |
|---------------------|---|
| Assembly accession  | GCA_905340355.1 |
| Accession of alternate haplotype | GCA_905340395.1 |
| Span (Mb)            | 425 |
| Number of contigs    | 82 |
| Contig N50 length (Mb) | 21 |
| Number of scaffolds  | 13 |
| Scaffold N50 length (Mb) | 116 |
| Longest scaffold (Mb) | 125 |
| BUSCO* genome score  | C:98.9%[S:96.0%,D:2.9%],F:0.6%,M:0.6%,n:2124 |

*BUSCO scores based on the coccidia_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/icRhaFulv1.1/dataset/CAJPIC01.1/buscio.
DNA was extracted from the whole organism of icRhaFulv1 at the Wellcome Sanger Institute (WSI) Scientific Operations core from the whole organism using the Qiagen MagAttract HMW DNA kit, according to the manufacturer’s instructions. RNA (from the whole organism of icRhaFulv4) was extracted in the Tree of Life Laboratory at the WSI using TRIzol, 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 read cloud DNA sequencing libraries, in addition to PolyA RNA-Seq libraries, were constructed according to the manufacturers’ instructions. Sequencing was performed by the Scientific Operations core at the WSI on Pacific
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

Figure 2. Genome assembly of *Rhagonycha fulva*, icRhaFulv1.1: 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/icRhaFulv1.1/dataset/CAJPIC01.1/blob.
Figure 3. Genome assembly of *Rhagonycha fulva*, icRhaFulv1.1: cumulative sequence. BlobToolKit cumulative sequence plot. The grey line shows cumulative length for all scaffolds. Coloured lines show cumulative lengths of scaffolds assigned to each phylum using the buscogenes taxrule. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/icRhaFulv1.1/dataset/CAJPIC01.1/cumulative.

assembly with longranger 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 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.
Figure 4. Genome assembly of *Rhagonycha fulva*, icRhaFulv1.1: Hi-C contact map. Hi-C contact map of the icRhaFulv1.1 assembly, visualised in HiGlass.

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
### Table 3. Software tools used.

| Software tool       | Version | Source                          |
|---------------------|---------|---------------------------------|
| Hifiasm             | 0.12    | 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            | 1.0     | Uliano-Silva et al., 2021       |
| gEVAL               | N/A     | Chow et al., 2016               |
| HiGlass             | 1.11.6  | Kerpedjiev et al., 2018         |
| PretextView         | 0.1.x   | https://github.com/wtsi-hpag/PretextView |
| BlobToolKit         | 2.6.1   | Challis et al., 2020            |

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: Rhagonycha fulva (common red soldier beetle). Accession number PRJEB43742; https://identifiers.org/ena.embl:PRJEB43742.

The genome sequence is released openly for reuse. The *R. fulva* 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.

### Acknowledgements

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 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.

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Members of the Tree of Life Core Informatics collective are listed here: https://doi.org/10.5281/zenodo.501354.

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

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Version 1

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✅ Yi-Ming Weng
University of Wisconsin Madison, Madison, WI, USA

Upon saying that the sufficient details of methods and materials were provided, I do want to learn more about the process of the assembly polishing. The author provided the aligning and variance calling tools for polishing, and I expected that the output of freebayes is a vcf file storing the variances. However, I can't directly depict the process of how the assembly was polished with the vcf file. It might be better to add this information in the manner of repeatability.

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 evolution and 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.

Reviewer Report 18 May 2022

https://doi.org/10.21956/wellcomeopenres.19002.r50422
The data note requires minor revision. The information on the species biology is sufficient, but I would recommend mentioning the distribution of the species as the researchers from other countries can use data and it is valuable to indicate where the species occurs. Geographic origin is mentioned twice with all coordinates, etc. If this is not a part of the report template, modify the text.

The description of applied methods is detailed, data access information is complete.

**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:** phylogenetics, phylogenomics

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.