



DATA NOTE

The genome sequence of a cave spider, *Troglohyphantes excavatus* Fage, 1919

[version 1; peer review: 3 approved]

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V1 First published: 08 Apr 2025, 10:177
<https://doi.org/10.12688/wellcomeopenres.23920.1>
Latest published: 08 Apr 2025, 10:177
<https://doi.org/10.12688/wellcomeopenres.23920.1>

Abstract
We present a genome assembly from a female *Troglohyphantes excavatus* (cave spider; Arthropoda; Arachnida; Araneae; Linyphiidae). The genome sequence has a total length of 1,028.29 megabases. Most of the assembly (94.43%) is scaffolded into 13 chromosomal pseudomolecules. The mitochondrial genome has also been assembled, with a length of 14.82 kilobases.

Keywords
Troglohyphantes excavatus, cave spider, Dinaric karst, genome sequence, chromosomal, Araneae



This article is included in the [Tree of Life gateway](#).

Open Peer Review

Approval Status   

	1	2	3
version 1			
08 Apr 2025	view	view	view

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Author roles: **Pavlek M:** Investigation, Resources, Writing – Original Draft Preparation, Writing – Review & Editing; **Rožman T:** Investigation, Resources;

Competing interests: No competing interests were disclosed.

Grant information: This work was supported by Wellcome through core funding to the Wellcome Sanger Institute (220540).

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How to cite this article: Pavlek M, Rožman T, Wellcome Sanger Institute Tree of Life Management, Samples and Laboratory team *et al.*

The genome sequence of a cave spider, *Troglohyphantes excavatus* Fage, 1919 [version 1; peer review: 3 approved] Wellcome Open Research 2025, 10:177 <https://doi.org/10.12688/wellcomeopenres.23920.1>

First published: 08 Apr 2025, 10:177 <https://doi.org/10.12688/wellcomeopenres.23920.1>

Species taxonomy

Eukaryota; Opisthokonta; Metazoa; Eumetazoa; Bilateria; Protostomia; Ecdysozoa; Panarthropoda; Arthropoda; Chelicerata; Arachnida; Araneae; Araneomorphae; Entelegynae; Orbiculariae; Araneoidea; Linyphiidae; Micronetinae; *Troglohyphantes*; *Troglohyphantes excavatus* Fage, 1919 (NCBI:txid1977180)

Background

The spider *Troglohyphantes excavatus* (Figure 1) is a cave-dwelling spider distributed in south-east Europe. With a range of approximately 16,000 square kilometres, extending through Croatia, Slovenia, Italy, and Austria, it is one of the species with the largest distribution area in the genus *Troglohyphantes* (Milano *et al.*, 2022). Although primarily found in caves, at its northern distribution border in Austria it is found in surface localities like in moss or in spruce or mixed forests (Milano *et al.*, 2022; Thaler, 1986). Occasionally it can be found in surface habitats in the south of its distribution also, but always in moist and shadowed places (Pavlek *et al.*, 2022). It is considered as troglophile (Mammola *et al.*, 2022), meaning that although it is primarily a cave species, it can survive in certain surface habitats and use them for dispersal.

Troglohyphantes excavatus is a relatively small spider with body size up to 3 mm, and leg length up to 9 mm (Deeleman-Reinhold, 1978; Mammola *et al.*, 2022), brownish in colour, and with functional eyes. It spins a thin and gentle sheet web from which it hangs upside down. It preys on other small cave invertebrates like springtails, beetles, isopods and similar, or, when positioned near a cave entrance, on small flying insects that seek shelter in caves. The species is not listed on the IUCN red list, but its conservation status has been evaluated and is considered as least concern (Milano *et al.*, 2022).

This is the first chromosome level genome of the genus *Troglohyphantes*. Most of its 138 species are found in caves, and they exhibit some level of cave adaptation, so this genome will be very important resource for studying evolution of cave adaptation in spiders and in invertebrates in general. Here we present the genome sequence of *Troglohyphantes excavatus*, based on a specimen from Špilja u kamenolomu Otruševac Cave, Otruševac Village.



Figure 1. Photograph of *Troglohyphantes excavatus* from Donja Baraćeva špilja cave. Photo: Martina Pavlek.

Genome sequence report

Sequencing data

The genome of a specimen of *Troglohyphantes excavatus* was sequenced using Pacific Biosciences single-molecule HiFi long reads, generating 48.75 Gb from 3.96 million reads. GenomeScope analysis of the PacBio HiFi data estimated the haploid genome size at 839.34 Mb, with a heterozygosity of 1.08% and repeat content of 32.10%. These values provide an initial assessment of genome complexity and the challenges anticipated during assembly. Based on this estimated genome size, the sequencing data provided approximately 29.0x coverage of the genome. Chromosome conformation Hi-C sequencing produced 113.52 Gb from 751.79 million reads. Table 1 summarises the specimen and sequencing information.

Assembly statistics

The primary haplotype was assembled, and contigs corresponding to an alternate haplotype were also deposited in INSDC databases. The assembly was improved by manual curation, which corrected 58 misjoins or missing joins and removed 15 haplotypic duplications. These interventions reduced the total assembly length by 0.53%, decreased the scaffold count by 4.29%, and increased the scaffold N50 by 0.63%. The final assembly has a total length of 1,028.29 Mb in 133 scaffolds, with 464 gaps, and a scaffold N50 of 74.87 Mb (Table 2).

The snail plot in Figure 2 provides a summary of the assembly statistics, indicating the distribution of scaffold lengths and other assembly metrics. Figure 3 shows the distribution of scaffolds by GC proportion and coverage. Figure 4 presents a cumulative assembly plot, with separate curves representing different scaffold subsets assigned to various phyla, illustrating the completeness of the assembly.

Most of the assembly sequence (94.25%) was assigned to 13 chromosomal-level scaffolds. These chromosome-level scaffolds, confirmed by Hi-C data, are named according to size (Figure 5; Table 3). During curation, it was noted that the specimen is the homogametic sex (female), but no X chromosome could be identified.

The mitochondrial genome was also assembled. This sequence is included as a contig in the multifasta file of the genome submission and as a standalone record.

Assembly quality metrics

The estimated Quality Value (QV) and *k*-mer completeness metrics, along with BUSCO completeness scores, were calculated for each haplotype and the combined assembly. The QV reflects the base-level accuracy of the assembly, while *k*-mer completeness indicates the proportion of expected *k*-mers identified in the assembly. BUSCO scores provide a measure of completeness based on benchmarking universal single-copy orthologues.

The combined primary and alternate assemblies achieve an estimated QV of 57.4. The *k*-mer recovery for the primary haplotype is 88.78%, and for the alternate haplotype

Table 1. Specimen and sequencing data for *Troglohyphantes excavatus*.

Project information			
Study title	Troglohyphantes excavatus		
Umbrella BioProject	PRJEB71877		
Species	<i>Troglohyphantes excavatus</i>		
BioSpecimen	SAMEA14515608		
NCBI taxonomy ID	1977180		
Specimen information			
Technology	ToLID	BioSample accession	Organism part
PacBio long read sequencing	qqTroExca7	SAMEA14515645	whole organism
Hi-C sequencing	qqTroExca2	SAMEA10458248	whole organism
RNA sequencing	qqTroExca3	SAMEA10458249	whole organism
Sequencing information			
Platform	Run accession	Read count	Base count (Gb)
Hi-C Illumina NovaSeq 6000	ERR12507424	7.52e+08	113.52
PacBio Sequel IIe	ERR12510606	1.66e+06	18.7
PacBio Sequel IIe	ERR12510605	2.30e+06	30.04
RNA Illumina NovaSeq 6000	ERR12507425	7.20e+07	10.88

74.06%; the combined primary and alternate assemblies have a *k*-mer recovery of 98.45%. BUSCO analysis using the arachnida_odb10 reference set (*n* = 2,934) identified 98.1% of the expected gene set (single = 93.1%, duplicated = 5.0%).

Table 2 provides assembly metric benchmarks adapted from Rhie *et al.* (2021) and the Earth BioGenome Project (EBP) Report on Assembly Standards September 2024. The assembly achieves the EBP reference standard of 6.C.Q57.

Methods

Sample acquisition

A female adult *Troglohyphantes excavatus* (specimen ID ERGA MP HR 06, ToLID qqTroExca7) was collected from Špilja u kamenolomu Otruševac cave, Otruševac Village (latitude 45.8199, longitude 15.6791) on 2022-03-26. The specimen was collected by Tin Rožman (Croatian Biospeleological Society, Zagreb, Croatia) and identified by Martina Pavlek (Ruder Boskovic Institute, Zagreb, Croatia).

The specimens used for Hi-C sequencing (specimen ID SAN0001794, ToLID qqTroExca2) and RNA sequencing (specimen ID SAN0001795, ToLID qqTroExca3) were collected from Donja Baračeva špilja cave, Rakovica Village (latitude 44.9842, longitude 15.7232) on 2021-04-11. The specimens were collected and identified by Martina Pavlek (Ruder Boskovic Institute, Zagreb, Croatia).

Nucleic acid extraction

The workflow for high molecular weight (HMW) DNA extraction at the Wellcome Sanger Institute (WSI) Tree of Life Core Laboratory includes a sequence of procedures: sample preparation and homogenisation, DNA extraction, fragmentation and purification. Detailed protocols are available on protocols.io (Denton *et al.*, 2023b). The qqTroExca7 sample was prepared for DNA extraction by weighing and dissecting it on dry ice (Jay *et al.*, 2023). Tissue from the whole organism was homogenised using a PowerMasher II tissue disruptor (Denton *et al.*, 2023a). HMW DNA was extracted using the Automated MagAttract v2 protocol (Oatley *et al.*, 2023a). DNA was sheared into an average fragment size of 12–20 kb in a Megaruptor 3 system (Bates *et al.*, 2023). Sheared DNA was purified by solid-phase reversible immobilisation, using AMPure PB beads to eliminate shorter fragments and concentrate the DNA (Oatley *et al.*, 2023b). The concentration of the sheared and purified DNA was assessed using a Nanodrop spectrophotometer and Qubit Fluorometer using the 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 qqTroExca3 in the Tree of Life Laboratory at the WSI using the RNA Extraction: Automated MagMax™ mirVana protocol (do Amaral *et al.*, 2023). The RNA concentration was assessed using a

Table 2. Genome assembly data for *Troglohyphantes excavatus*.

Genome assembly		
Assembly name	qqTroExca7.1	
Assembly accession	GCA_963932185.1	
Alternate haplotype accession	GCA_963932215.1	
Assembly level for primary assembly	chromosome	
Span (Mb)	1,028.29	
Number of contigs	597	
Number of scaffolds	133	
Longest scaffold (Mb)	82.79	
Assembly metric	Measure	Benchmark
Contig N50 length	3.43 Mb	≥ 1 Mb
Scaffold N50 length	74.87 Mb	= chromosome N50
Consensus quality (QV)	Primary: 57.8; alternate: 57.0; combined 57.4	≥ 40
k-mer completeness	Primary: 88.78%; alternate: 74.06%; combined: 98.45%	$\geq 95\%$
BUSCO*	C:98.1%[S:93.1%,D:5.0%], F:0.7%,M:1.2%,n:2,934	$S > 90\%$; $D < 5\%$
Percentage of assembly mapped to chromosomes	94.25%	$\geq 90\%$
Sex chromosomes	Not identified	localised homologous pairs
Organelles	Mitochondrial genome: 14.82 kb	complete single alleles

* BUSCO scores based on the arachnida_odb10 BUSCO set using version 5.5.0. C = complete [S = single copy, D = duplicated], F = fragmented, M = missing, n = number of orthologues in comparison.

Nanodrop spectrophotometer and a Qubit Fluorometer using the Qubit RNA Broad-Range Assay kit. Analysis of the integrity of the RNA was done using the Agilent RNA 6000 Pico Kit and Eukaryotic Total RNA assay.

Hi-C sample preparation

Tissue from the whole organism of the qqTroExca2 sample was processed for Hi-C sequencing at the WSI Scientific Operations core, using the Arima-HiC v2 kit. In brief, 20–50 mg of frozen tissue (stored at -80°C) was fixed, and the DNA crosslinked using a TC buffer with 22% formaldehyde concentration. After crosslinking, the tissue was homogenised using the Diagenode Power Masher-II and BioMasher-II tubes and pestles. Following the Arima-HiC v2 kit manufacturer's instructions, crosslinked DNA was digested using a restriction enzyme master mix. The 5'-overhangs were filled in and labelled with biotinylated nucleotides and proximally ligated. An overnight incubation was carried out for enzymes to digest remaining proteins and for crosslinks to reverse. A clean up was performed with SPRIselect beads prior to library preparation. Additionally, the biotinylation percentage was

estimated using the Qubit Fluorometer v4.0 (Thermo Fisher Scientific) and Qubit HS Assay Kit and Arima-HiC v2 QC beads.

Library preparation and sequencing

Library preparation and sequencing were performed at the WSI Scientific Operations core.

PacBio HiFi

At a minimum, samples were required to have an average fragment size exceeding 8 kb and a total mass over 400 ng to proceed to the low input SMRTbell Prep Kit 3.0 protocol (Pacific Biosciences, California, USA), depending on genome size and sequencing depth required. Libraries were prepared using the SMRTbell Prep Kit 3.0 (Pacific Biosciences, California, USA) as per the manufacturer's instructions. The kit includes the reagents required for end repair/A-tailing, adapter ligation, post-ligation SMRTbell bead cleanup, and nuclease treatment. Following the manufacturer's instructions, size selection and clean up was carried out using diluted AMPure PB beads (Pacific Biosciences, California, USA). DNA concentration was

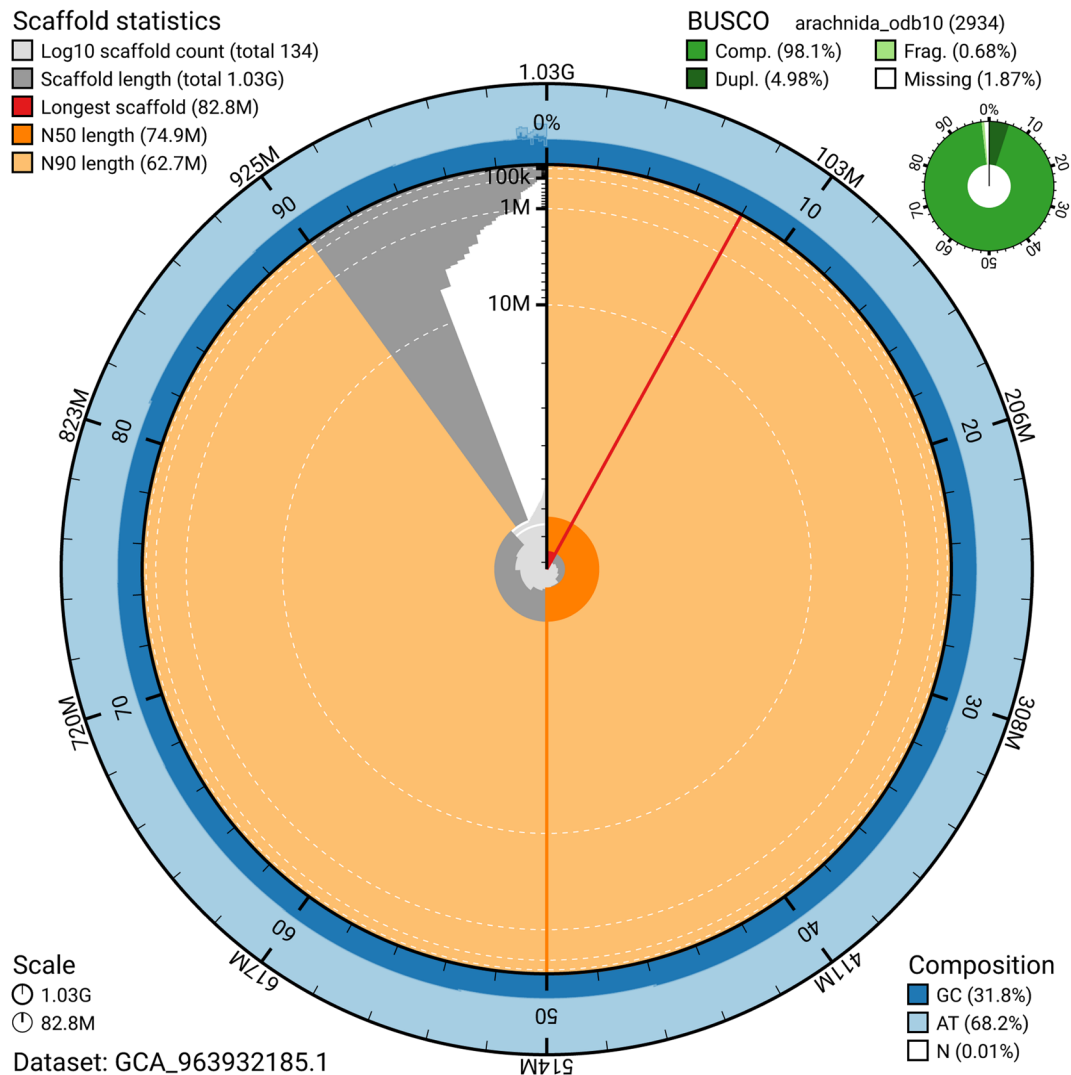


Figure 2. Genome assembly of *Troglodyphantes excavatus*, qqTroExca7.1: metrics. The BlobToolKit snail plot provides an overview of assembly metrics and BUSCO gene completeness. The circumference represents the length of the whole genome sequence, and the main plot is divided into 1,000 bins around the circumference. The outermost blue tracks display the distribution of GC, AT, and N percentages across the bins. Scaffolds are arranged clockwise from longest to shortest and are depicted in dark grey. The longest scaffold is indicated by the red arc, and the deeper orange and pale orange arcs represent the N50 and N90 lengths. A light grey spiral at the centre shows the cumulative scaffold count on a logarithmic scale. A summary of complete, fragmented, duplicated, and missing BUSCO genes in the arachnida_odb10 set is presented at the top right. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/GCA_963932185.1/dataset/GCA_963932185.1/snail.

quantified using the Qubit Fluorometer v4.0 (Thermo Fisher Scientific) with Qubit 1X dsDNA HS assay kit and the final library fragment size analysis was carried out using the Agilent Femto Pulse Automated Pulsed Field CE Instrument (Agilent Technologies) and gDNA 55kb BAC analysis kit.

Samples were sequenced using the Sequel IIe system (Pacific Biosciences, California, USA). The concentration of the library loaded onto the Sequel IIe was in the range 40–135 pM. The SMRT link software, a PacBio web-based end-to-end workflow

manager, was used to set-up and monitor the run, as well as perform primary and secondary analysis of the data upon completion.

Hi-C

For Hi-C library preparation, DNA was fragmented using the Covaris E220 sonicator (Covaris) and size selected using SPRISelect beads to 400 to 600 bp. The DNA was then enriched using the Arima-HiC v2 kit Enrichment beads. Using the NEBNext Ultra II DNA Library Prep Kit (New England

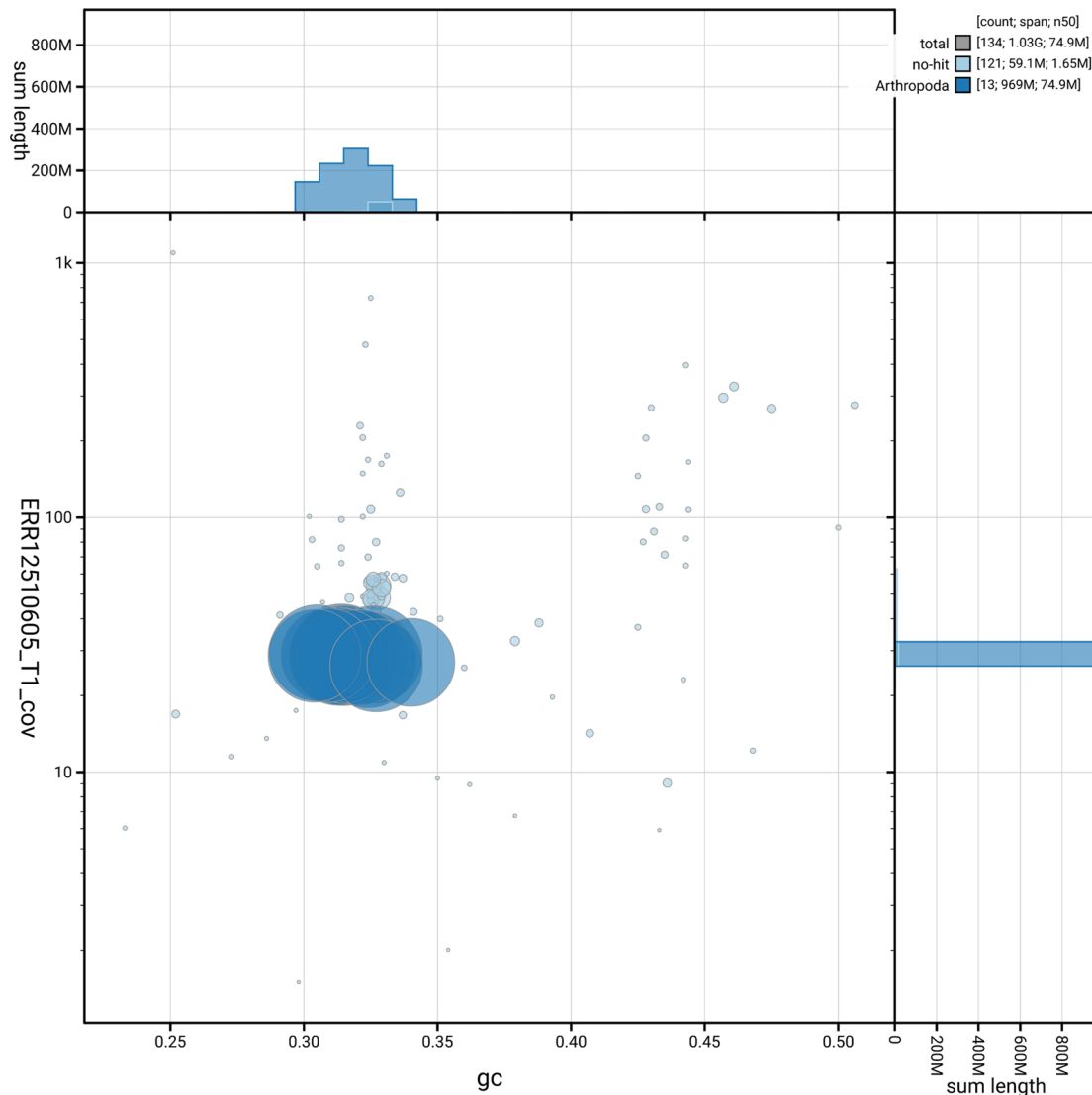


Figure 3. Genome assembly of *Troglohyphantes excavatus*, qqTroExca7.1: BlobToolKit GC-coverage plot. Blob plot showing sequence coverage (vertical axis) and GC content (horizontal axis). The circles represent scaffolds, with the size proportional to scaffold length and the colour representing phylum membership. The histograms along the axes display the total length of sequences distributed across different levels of coverage and GC content. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/GCA_963932185.1/dataset/GCA_963932185.1/blob.

Biolabs) for end repair, a-tailing, and adapter ligation. This uses a custom protocol which resembles the standard NEBNext Ultra II DNA Library Prep protocol but where library preparation occurs while DNA is bound to the Enrichment beads. For library amplification, 10 to 16 PCR cycles were required, determined by the sample biotinylation percentage. The Hi-C sequencing was performed using paired-end sequencing with a read length of 150 bp on an Illumina NovaSeq 6000 instrument.

RNA

Poly(A) RNA-Seq libraries were constructed using the NEB Ultra II RNA Library Prep kit, following the manufacturer's

instructions. RNA sequencing was performed on the Illumina NovaSeq 6000 instrument.

Genome assembly, curation and evaluation

Assembly

Prior to assembly of the PacBio HiFi reads, a database of k -mer counts ($k = 31$) was generated from the filtered reads using FastK. GenomeScope2 (Ranallo-Benavidez *et al.*, 2020) was used to analyse the k -mer frequency distributions, providing estimates of genome size, heterozygosity, and repeat content.

The HiFi reads were assembled using Hifiasm (Cheng *et al.*, 2021) with the --primary option. Haplotypic duplications were

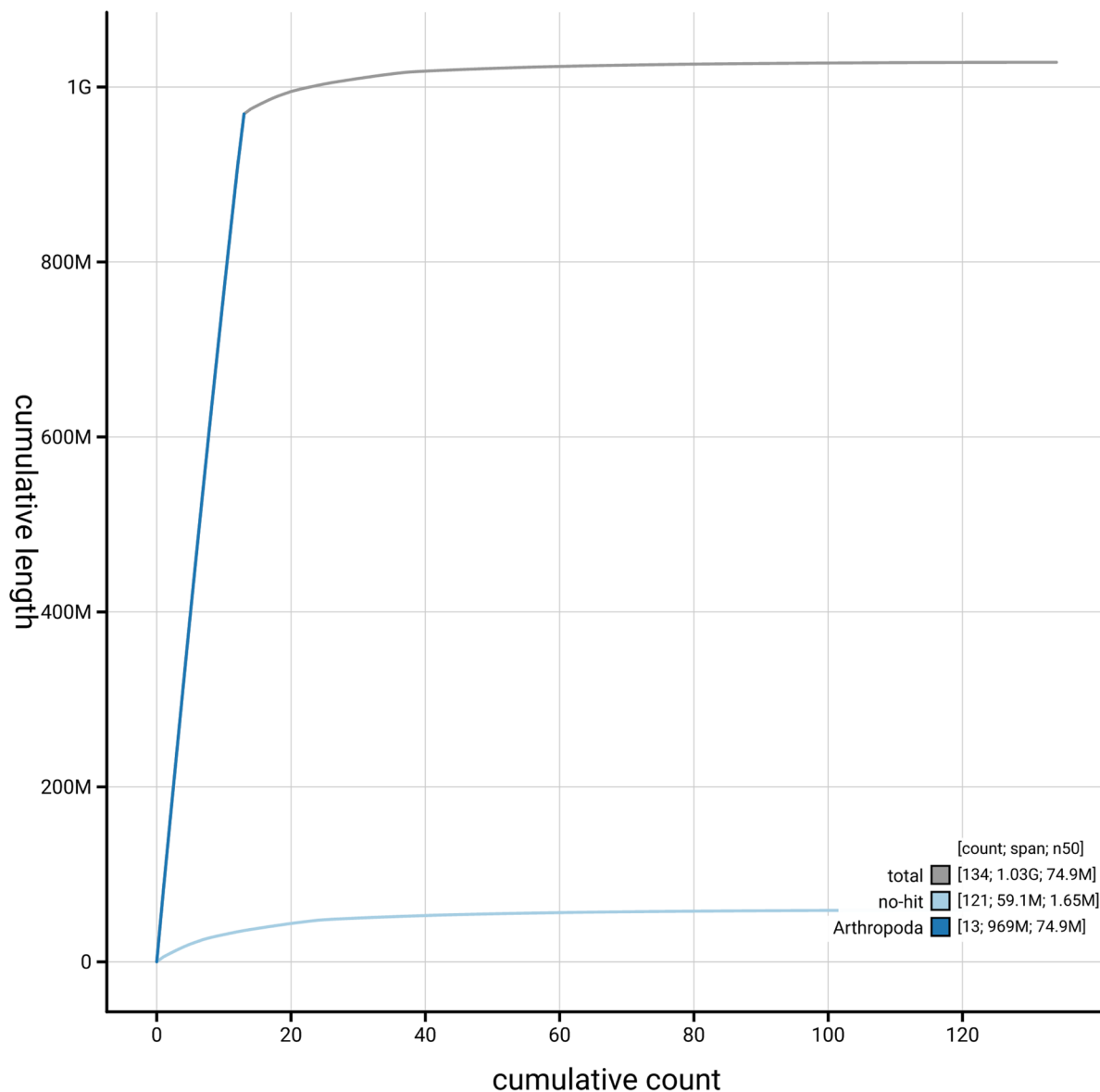


Figure 4. Genome assembly of *Troglodyphantes excavatus*, qqTroExca7.1: BlobToolKit cumulative sequence plot. The grey line shows cumulative length for all scaffolds. Coloured lines show cumulative lengths of scaffolds assigned to each phylum using the buscogenes taxrule. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/GCA_963932185.1/dataset/GCA_963932185.1/cumulative.

identified and removed using `purge_dups` (Guan *et al.*, 2020). The Hi-C reads were mapped to the primary contigs using `bwa-mem2` (Vasimuddin *et al.*, 2019). The contigs were further scaffolded using the provided Hi-C data (Rao *et al.*, 2014) in YaHS (Zhou *et al.*, 2023) using the `--break` option for handling potential misassemblies. The scaffolded assemblies were evaluated using Gfastats (Formenti *et al.*, 2022), BUSCO (Manni *et al.*, 2021) and MERQUERY.FK (Rhie *et al.*, 2020).

The mitochondrial genome was assembled using MitoHiFi (Uliano-Silva *et al.*, 2023), which runs MitoFinder (Allio *et al.*, 2020) and uses these annotations to select the final

mitochondrial contig and to ensure the general quality of the sequence.

Assembly curation

The assembly was decontaminated using the Assembly Screen for Cobionts and Contaminants (ASCC) pipeline. Flat files and maps used in curation were generated via the TreeVal pipeline (Pointon *et al.*, 2023). Manual curation was conducted primarily in PretextView (Harry, 2022) and HiGlass (Kerpedjiev *et al.*, 2018), with additional insights provided by JBrowse2 (Diesh *et al.*, 2023). Scaffolds were visually inspected and corrected as described by Howe *et al.* (2021).

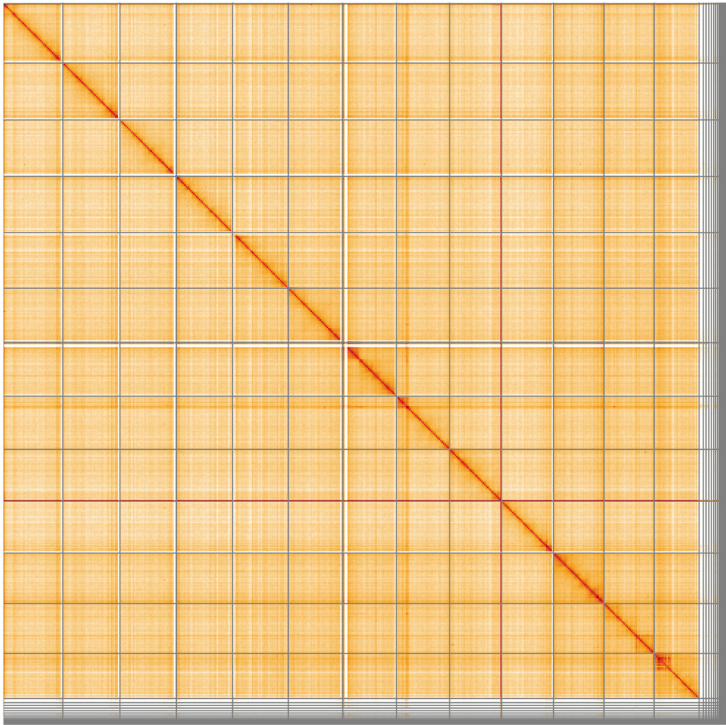


Figure 5. Genome assembly of *Troglohyphantes excavatus*: Hi-C contact map of the qqTroExca7.1 assembly, visualised using HiGlass. Chromosomes are shown in order of size from left to right and top to bottom. An interactive version of this figure may be viewed at <https://genome-note-higlass.tol.sanger.ac.uk/l/?d=f10fgFnaR8OaMMV4jWUjhq>.

Table 3. Chromosomal pseudomolecules in the genome assembly of *Troglohyphantes excavatus*, qqTroExca7.

INSDC accession	Name	Length (Mb)	GC%
OZ010666.1	1	82.79	31.5
OZ010667.1	2	79.46	31.5
OZ010668.1	3	78.79	31.5
OZ010669.1	4	78.27	31.5
OZ010670.1	5	77.59	32.5
OZ010671.1	6	75.81	32.5
OZ010672.1	7	74.87	30.5
OZ010673.1	8	74.11	32
OZ010674.1	9	72.31	31.5
OZ010675.1	10	72.27	31
OZ010676.1	11	70.42	30.5
OZ010677.1	12	69.77	32.5
OZ010678.1	13	62.71	34
OZ010679.1	MT	0.01	25

Any identified contamination, missed joins, and mis-joins were amended, and duplicate sequences were tagged and removed. The curation process is documented at <https://gitlab.com/wtsi-grit/rapid-curation>.

Assembly quality assessment

The Merquy.FK tool (Rhie *et al.*, 2020), run in a Singularity container (Kurtzer *et al.*, 2017), was used to evaluate *k*-mer completeness and assembly quality for the primary and alternate haplotypes using the *k*-mer databases (*k* = 31) that were computed prior to genome assembly. The analysis outputs included assembly QV scores and completeness statistics.

A Hi-C contact map was produced for the final version of the assembly. The Hi-C reads were aligned using bwa-mem2 (Vasimuddin *et al.*, 2019) and the alignment files were combined using SAMtools (Danecek *et al.*, 2021). The Hi-C alignments were converted into a contact map using BEDTools (Quinlan & Hall, 2010) and the Cooler tool suite (Abdennur & Mirny, 2020). The contact map was visualised in HiGlass (Kerpedjiev *et al.*, 2018).

The blobtoolkit pipeline is a Nextflow port of the previous Snakemake Blobtoolkit pipeline (Challis *et al.*, 2020). It aligns the PacBio reads in SAMtools and minimap2 (Li, 2018) and generates coverage tracks for regions of fixed size. In parallel,

it queries the GoAT database (Challis *et al.*, 2023) to identify all matching BUSCO lineages to run BUSCO (Manni *et al.*, 2021). For the three domain-level BUSCO lineages, the pipeline aligns the BUSCO genes to the UniProt Reference Proteomes database (Bateman *et al.*, 2023) with DIAMOND blastp (Buchfink *et al.*, 2021). The genome is also divided into chunks according to the density of the BUSCO genes from the closest taxonomic lineage, and each chunk is aligned to the UniProt Reference Proteomes database using DIAMOND blastx. Genome sequences without a hit are chunked using seqtk and aligned to the NT database with blastn (Altschul *et al.*, 1990). The blobtools suite combines all these outputs into a blobdir for visualisation.

The blobtoolkit pipeline was developed using nf-core tooling (Ewels *et al.*, 2020) and MultiQC (Ewels *et al.*, 2016), relying

on the Conda package manager, the Bioconda initiative (Grüning *et al.*, 2018), the Biocontainers infrastructure (da Veiga Leprevost *et al.*, 2017), as well as the Docker (Merkel, 2014) and Singularity (Kurtzer *et al.*, 2017) containerisation solutions.

Table 4 contains a list of relevant software tool versions and sources.

Wellcome Sanger Institute – Legal and Governance

The materials that have contributed to this genome note have been supplied by a Tree of Life collaborator. The Wellcome Sanger Institute employs a process whereby due diligence is carried out proportionate to the nature of the materials themselves, and the circumstances under which they have been/are to be collected and provided for use. The purpose

Table 4. Software tools: versions and sources.

Software tool	Version	Source
BEDTools	2.30.0	https://github.com/arq5x/bedtools2
BLAST	2.14.0	ftp://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/
BlobToolKit	4.3.9	https://github.com/blobtoolkit/blobtoolkit
BUSCO	5.5.0	https://gitlab.com/ezlab/busco
bwa-mem2	2.2.1	https://github.com/bwa-mem2/bwa-mem2
Cooler	0.8.11	https://github.com/open2c/cooler
DIAMOND	2.1.8	https://github.com/bbuchfink/diamond
fasta_windows	0.2.4	https://github.com/tolkit/fasta_windows
FastK	666652151335353eef2fcd58880bcef5bc2928e1	https://github.com/thegenemyers/FASTK
Gfastats	1.3.6	https://github.com/vgl-hub/gfastats
GoAT CLI	0.2.5	https://github.com/genomehubs/goat-cli
Hifiasm	0.19.8-r603	https://github.com/chhy123/hifiasm
HiGlass	44086069ee7d4d3f6f3f0012569789ec138f42b84aa44357826c0b6753eb28de	https://github.com/higlass/higlass
MerquyFK	d00d98157618f4e8d1a9190026b19b471055b22e	https://github.com/thegenemyers/MERQUERY.FK
Minimap2	2.24-r1122	https://github.com/lh3/minimap2
MitoHiFi	3	https://github.com/marcelauliano/MitoHiFi
MultiQC	1.14, 1.17, and 1.18	https://github.com/MultiQC/MultiQC
NCBI Datasets	15.12.0	https://github.com/ncbi/datasets
Nextflow	23.04.1	https://github.com/nextflow-io/nextflow
PretextView	0.2.5	https://github.com/sanger-tol/PretextView
purge_dups	1.2.5	https://github.com/dfguan/purge_dups
samtools	1.19.2	https://github.com/samtools/samtools
sanger-tol/ascc	-	https://github.com/sanger-tol/ascc
sanger-tol/blobtoolkit	0.5.1	https://github.com/sanger-tol/blobtoolkit
Seqtk	1.3	https://github.com/lh3/seqtk
Singularity	3.9.0	https://github.com/sylabs/singularity
TreeVal	1.2.0	https://github.com/sanger-tol/treeval
YaHS	1.2a.2	https://github.com/c-zhou/yahs

of this is to address and mitigate any potential legal and/or ethical implications of receipt and use of the materials as part of the research project, and to ensure that in doing so we align with best practice wherever possible.

The overarching areas of consideration are:

- Ethical review of provenance and sourcing of the material
- Legality of collection, transfer and use (national and international)

Each transfer of samples is undertaken according to a Research Collaboration Agreement or Material Transfer Agreement entered into by the Tree of Life collaborator, Genome Research Limited (operating as the Wellcome Sanger Institute) and in some circumstances other Tree of Life collaborators.

Data availability

European Nucleotide Archive: *Troglohyphantes excavatus*. Accession number PRJEB71877; <https://identifiers.org/ena.embl/PRJEB71877>. The genome sequence is released openly for reuse. The *Troglohyphantes excavatus* genome sequencing initiative is part of the European Reference Genome Atlas Pilot Project (<https://www.erga-biodiversity.eu/pilot-project>). All

raw sequence data and the assembly have been deposited in INSDC databases. The genome will be annotated using available 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](#) and [Table 2](#).

Author information

Members of the Wellcome Sanger Institute Tree of Life Management, Samples and Laboratory team are listed here: <https://doi.org/10.5281/zenodo.12162482>.

Members of Wellcome Sanger Institute Scientific Operations: Sequencing Operations are listed here: <https://doi.org/10.5281/zenodo.12165051>.

Members of the Wellcome Sanger Institute Tree of Life Core Informatics team are listed here: <https://doi.org/10.5281/zenodo.12160324>.

Members of the Tree of Life Core Informatics collective are listed here: <https://doi.org/10.5281/zenodo.12205391>.

Acknowledgements

Members of The Croatian Biospeleological Society from Zagreb, Croatia.

References

- Abdennur N, Mirny LA: **Cooler: scalable storage for Hi-C data and other genomically labeled arrays.** *Bioinformatics*. 2020; **36**(1): 311–316. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- 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](#)
- Altschul SF, Gish W, Miller W, et al.: **Basic Local Alignment Search Tool.** *J Mol Biol*. 1990; **215**(3): 403–410. [PubMed Abstract](#) | [Publisher Full Text](#)
- Bateman A, Martin MJ, Orchard S, et al.: **UniProt: the Universal Protein Knowledgebase in 2023.** *Nucleic Acids Res*. 2023; **51**(D1): D523–D531. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- Bates A, Clayton-Lucey I, Howard C: **Sanger Tree of Life HMW DNA fragmentation: diagenode Megaruptor[®]3 for LI PacBio.** *protocols.io*. 2023. [Publisher Full Text](#)
- Buchfink B, Reuter K, Drost HG: **Sensitive protein alignments at Tree-of-Life scale using DIAMOND.** *Nat Methods*. 2021; **18**(4): 366–368. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- Challis R, Kumar S, Sotero-Caio C, et al.: **Genomes on a Tree (GoaT): a versatile, scalable search engine for genomic and sequencing project metadata across the eukaryotic Tree of Life [version 1; peer review: 2 approved].** *Wellcome Open Res*. 2023; **8**: 24. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- Challis R, Richards E, Rajan J, et al.: **BlobToolKit – interactive quality assessment of genome assemblies.** *G3 (Bethesda)*. 2020; **10**(4): 1361–1374. [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): 170–175. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- da Veiga Leprevost F, Grüning BA, Alves Afritas S, et al.: **BioContainers: an open-source and community-driven framework for software standardization.** *Bioinformatics*. 2017; **33**(16): 2580–2582. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- Danecek P, Bonfield JK, Liddle J, et al.: **Twelve years of SAMtools and BCFtools.** *GigaScience*. 2021; **10**(2): giab008. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- Deeleman-Reinhold CL: **Revision of the cave-dwelling and related spiders of the genus *Troglohyphantes* Joseph (Linyphiidae), with special reference to the Yugoslav species.** *Razpr Slov Akad Znan Umet (Prirod. Vede)*. 1978; **23**: 1–220. [Reference Source](#)
- Denton A, Oatley G, Cornwell C, et al.: **Sanger Tree of Life sample homogenisation: PowerMash.** *protocols.io*. 2023a. [Publisher Full Text](#)
- Denton A, Yatsenko H, Jay J, et al.: **Sanger Tree of Life wet laboratory protocol collection V.1.** *protocols.io*. 2023b. [Publisher Full Text](#)
- Diesh C, Stevens GJ, Xie P, et al.: **JBrowse 2: a modular genome browser with views of synteny and structural variation.** *Genome Biol*. 2023; **24**(1): 74. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)
- do Amaral RJV, Bates A, Denton A, et al.: **Sanger Tree of Life RNA extraction: automated MagMax[™] mirVana.** *protocols.io*. 2023. [Publisher Full Text](#)
- Ewels PA, Peltzer A, Fillinger S, et al.: **The nf-core framework for community-curated bioinformatics pipelines.** *Nat Biotechnol*. 2020; **38**(3): 276–278. [PubMed Abstract](#) | [Publisher Full Text](#)
- Ewels P, Magnusson M, Lundin S, et al.: **MultiQC: summarize analysis results for multiple tools and samples in a single report.** *Bioinformatics*. 2016; **32**(19): 3047–3048. [PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Formenti G, Abueg L, Brajuka A, *et al.*: **Gfastats: conversion, evaluation and manipulation of genome sequences using assembly graphs.** *Bioinformatics*. 2022; **38**(17): 4214–4216.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Grüning B, Dale R, Sjödin A, *et al.*: **Bioconda: sustainable and comprehensive software distribution for the life sciences.** *Nat Methods*. 2018; **15**(7): 475–476.

[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): 2896–2898.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Harry E: **PretextView (Paired REad TEXTure Viewer): a desktop application for viewing pretext contact maps.** 2022.

[Reference Source](#)

Howe K, Chow W, Collins J, *et al.*: **Significantly improving the quality of genome assemblies through curation.** *GigaScience*. 2021; **10**(1): g1aa153.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Jay J, Yatsenko H, Narváez-Gómez JP, *et al.*: **Sanger Tree of Life sample preparation: triage and dissection.** *protocols.io*. 2023.

[Publisher Full Text](#)

Kerpedjiev P, Abdennur N, Lekschas 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](#)

Kurtzer GM, Sochat V, Bauer MW: **Singularity: scientific containers for mobility of compute.** *PLoS One*. 2017; **12**(5): e0177459.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Li H: **Minimap2: pairwise alignment for nucleotide sequences.** *Bioinformatics*. 2018; **34**(18): 3094–3100.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Mammola S, Pavlek M, Huber BA, *et al.*: **A trait database and updated checklist for European subterranean spiders.** *Sci Data*. 2022; **9**(1): 236.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Manni M, Berkeley MR, Seppay M, *et al.*: **BUSCO update: novel and streamlined workflows along with broader and deeper phylogenetic coverage for scoring of eukaryotic, prokaryotic, and viral genomes.** *Mol Biol Evol*. 2021; **38**(10): 4647–4654.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Merkel D: **Docker: lightweight Linux containers for consistent development and deployment.** *Linux J*. 2014; **2014**(239): 2, [Accessed 2 April 2024].

[Reference Source](#)

Milano F, Borio L, Komposch C, *et al.*: **Species conservation profiles of the endemic spiders *Traglohyphantes* (Araneae, Linyphiidae) from the Alps and the north-western Dinarides.** *Biodivers Data J*. 2022; **10**: e87261.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Oatley G, Denton A, Howard C: **Sanger Tree of Life HMW DNA extraction: automated MagAttract v.2.** *protocols.io*. 2023a.

[Publisher Full Text](#)

Oatley G, Sampaio F, Howard C: **Sanger Tree of Life fragmented DNA clean up: automated SPRI.** *protocols.io*. 2023b.

[Publisher Full Text](#)

Pavlek M, Gauthier J, Tonzo V, *et al.*: **Life-history traits drive spatial genetic structuring in Dinaric cave spiders.** *Front Ecol Evol*. 2022; **10**: 910084.

[Publisher Full Text](#)

Pointon DL, Eagles W, Sims Y, *et al.*: **sanger-tol/treeval v1.0.0 – Ancient Atlantis.** 2023.

[Publisher Full Text](#)

Quinlan AR, Hall IM: **BEDTools: a flexible suite of utilities for comparing genomic features.** *Bioinformatics*. 2010; **26**(6): 841–842.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Ranallo-Benavidez TR, Jaron KS, Schatz MC: **GenomeScope 2.0 and Smudgeplot for reference-free profiling of polyploid genomes.** *Nat Commun*. 2020; **11**(1): 1432.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

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; **159**(7): 1665–1680.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Rhie A, McCarthy SA, Fedrigo O, *et al.*: **Towards complete and error-free genome assemblies of all vertebrate species.** *Nature*. 2021; **592**(7856): 737–746.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Rhie A, Walenz BP, Koren S, *et al.*: **Merqury: reference-free quality, completeness, and phasing assessment for genome assemblies.** *Genome Biol*. 2020; **21**(1): 245.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Thaler K: **Über einige Funde von Troglolyphantes-Arten in Kärnten (Österreich) (Arachnida, Aranei: Linyphiidae).** *Carinthia II*. 1986; **176**(96): 287–302.

[Reference Source](#)

Ulliano-Silva M, Ferreira JGRN, Krashennikova K, *et al.*: **MitoHiFi: a python pipeline for mitochondrial genome assembly from PacBio high fidelity reads.** *BMC Bioinformatics*. 2023; **24**(1): 288.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Vasimuddin M, Misra S, Li H, *et al.*: **Efficient architecture-aware acceleration of BWA-MEM for multicore systems.** In: *2019 IEEE International Parallel and Distributed Processing Symposium (IPDPS)*. IEEE, 2019; 314–324.

[Publisher Full Text](#)

Zhou C, McCarthy SA, Durbin R: **YaHS: yet another Hi-C scaffolding tool.** *Bioinformatics*. 2023; **39**(1): btac808.

[PubMed Abstract](#) | [Publisher Full Text](#) | [Free Full Text](#)

Open Peer Review

Current Peer Review Status:   

Version 1

Reviewer Report 10 May 2025

<https://doi.org/10.21956/wellcomeopenres.26389.r122437>

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Ethan Briggs

University of Idaho, Moscow, USA

Reviewer report for "The genome sequence of a cave spider, *Troglohyphantes excavatus* Fage, 1919"

In this manuscript the authors assemble a *de novo* genome of a female *Troglohyphantes excavatus*, using PacBio HiFi, HiC and RNAseq data. The background information on this species as well as the rationale for sequencing this spider is well explained. The protocols used for this study are appropriate and technically sound, following the current, generally accepted protocols for genome assembly and assessment. The authors provide sufficient details to allow for replication. I have assumed that when a program is used, the default settings were implemented unless otherwise stated. Datasets in this study appear to be accessible, the accession numbers for the genome and raw data take the reader to the appropriate online depositories.

A minor comment that I would like to see added to this paper, as well as other similar studies, is the citation of the paper the species was originally described. Not citing such papers overlooks the important work taxonomists provide. In this specific case, I ask the authors to cite: Fage 1919 Biospelogica XL. Etudes sur les araignées cavernicoles. III. Sur le genre *Troglohyphantes*. Archives de Zoologie Expérimentale et Générale 58: 55-148, pl. 2-8.

I recommend this manuscript be accepted and believe it will provide valuable genomic information for both arachnid and troglomorphic evolution.

References

1. Fage 1919 Biospelogica XL. Etudes sur les araignées cavernicoles. III. Sur le genre *Troglohyphantes*. Archives de Zoologie Expérimentale et Générale 58: 55-148, pl. 2-8.

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: Genome assembly, phylogenomics, systematics, species delimitation

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 07 May 2025

<https://doi.org/10.21956/wellcomeopenres.26389.r122018>

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Marc Domènech 

Genètica, Microbiologia i Estadística, Universitat de Barcelona, Barcelona, Spain

Jesus Lozano-Fernandez 

Universitat de Barcelona, Barcelona, Catalonia, Spain

This article provides the first chromosome-level genome for a spider in the genus *Troglohyphantes*. Although there are other genomes available for Araneae, even in the same family Linyphiidae, having the genome of a troglophile species can help study the genomic adaptations to life in caves.

-In the second paragraph, "excavates" should be replaced by "excavatus".

-In the third paragraph, according to the World Spider Catalog (as checked in 16 April 2025), there are 139 species of *Troglohyphantes*.

-Although it would be optimal that all specimens used were collected in the same locality, especially when dealing with cave organisms, these species are so widespread in south-east Europe that it should not be a problem.

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: Arachnology, phylogenomics

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.

Reviewer Report 28 April 2025

<https://doi.org/10.21956/wellcomeopenres.26389.r122446>

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Feng Zhang 

Nanjing Agricultural University, Nanjing, China

This Data Note effectively reports the generation of a high-quality, chromosome-level genome assembly for the cave spider *Troglohyphantes excavatus*. This represents the first such resource for this genus, which is relevant for studies of cave adaptation, speciation in fragmented habitats like the Dinaric Karst, and invertebrate evolution more broadly. The use of state-of-the-art sequencing (PacBio HiFi, Hi-C) and assembly methods, combined with thorough quality assessment (BUSCO, QV, k-mer completeness, Hi-C contact maps) and adherence to reporting standards (e.g., EBP standards), are major strengths. The data release is timely and valuable for the community. While the report is generally clear and comprehensive for its format.

A minor comment:

The k-mer based estimate (GenomeScope) suggested a haploid size of ~839 Mb, while the final primary assembly is ~1.03 Gb. This ~23% difference is not insignificant and is not commented upon. Increase the parameter 'm'?

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: genomics, 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.
