#### ERGA Assembly Report

v24.10.15

Tags: ERGA-BGE

TxID	190562	
ToLID	tfGorWolt8	
Species	Gordionus wolterstorffii	
Class	Gordioida	
Order	Chordodea	

Genome Traits	Expected	Observed
Haploid size (bp)	142,426,324	210,545,467
Haploid Number	5 (source: ancestor)	4
Ploidy	2 (source: ancestor)	2
Sample Sex	unknown	unknown

#### EBP metrics summary and curation notes

Obtained EBP quality metric for pri: 7.7.Q70

The following metrics were automatically flagged as below EBP recommended standards or different from expected:

- . Observed Haploid size (bp) has >20% difference with Expected
- . Observed Haploid Number is different from Expected
- . Kmer completeness value is less than 90 for pri
- . BUSCO single copy value is less than 90% for pri

#### Curator notes

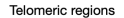
- . Interventions/Gb: None
- . Contamination notes: "FCS-GX identified 15 bacterial sequences that were removed from the base assembly (3,998,988 bp in total)"
- . Other observations: "It was not possible to get a good Hi-C library for this species but still we strongly believe that we have achieved a chromosome level assembly. Hifiasm assembled 4 long sequences that seem to be full chromosomes. Due to the lack of Hi-C data, we have posted below an image of the distribution of telomeres along the 4 chromosomes."

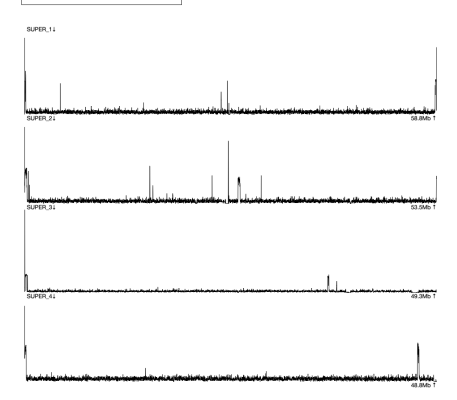
## Quality metrics table

Metrics	Pre-curation pri	Curated pri
Total bp	214,544,449	210,545,467
GC %	29.86	29.51
Gaps/Gbp	0	0
Total gap bp	0	0
Scaffolds	23	8
Scaffold N50	53,486,554	53,486,554
Scaffold L50	2	2
Scaffold L90	4	4
Contigs	23	8
Contig N50	53,486,554	53,486,554
Contig L50	2	2
Contig L90	4	4
QV	60.8148	70.0004
Kmer compl.	73.9189	72.3129
BUSCO sing.	54.8%	55.0%
BUSCO dupl.	1.9%	0.9%
BUSCO frag.	11.8%	12.1%
BUSCO miss.	31.5%	32.0%

Warning! BUSCO versions or lineage datasets are not the same across results:
BUSCO: 5.4.0 (euk\_genome\_met, metaeuk) / Lineage: metazoa\_odb10 (genomes:65, BUSCOs:954)
BUSCO: 5.5.0 (euk\_genome\_met, metaeuk) / Lineage: metazoa\_odb10 (genomes:65, BUSCOs:954)

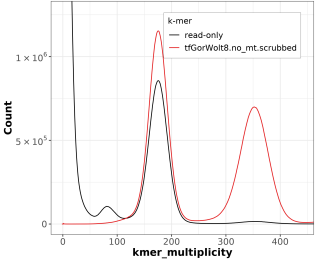
# HiC contact map of curated assembly



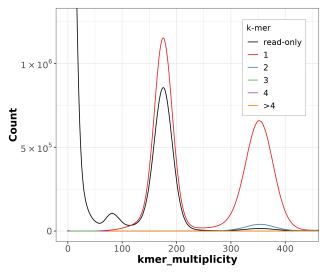


pri File link is missing!

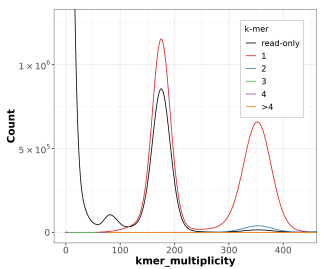
## K-mer spectra of curated assembly



Distribution of k-mer counts coloured by their presence in reads/assemblies



Distribution of k-mer counts per copy numbers found in asm



Distribution of k-mer counts per copy numbers found in asm

## Post-curation contamination screening

#### Data profile

Data	ONT	Illumina
Coverage	86	85

### Assembly pipeline

## Curation pipeline

```
- FCS-GX
|_ ver: None
|_ key param: NA
```

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Date and time: 2025-09-18 11:20:10 CEST