#### ERGA Assembly Report

v24.10.15

Tags: ERGA-BGE

TxID	522842		
ToLID	ilMarQuer1		
Species	Marumba quercus		
Class	Insecta		
Order	Lepidoptera		

Genome Traits	Expected	Observed
Haploid size (bp)	596,365,935	596,998,386
Haploid Number	28 (source: ancestor)	28
Ploidy	2 (source: ancestor)	2
Sample Sex	ZZ	ZZ

#### EBP metrics summary and curation notes

Obtained EBP quality metric for hap1: 7.7.Q70 Obtained EBP quality metric for hap2: 7.7.Q70

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

#### Curator notes

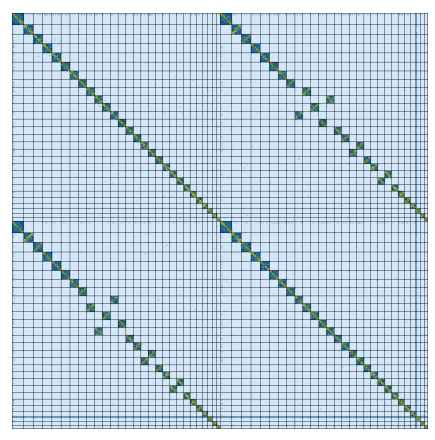
- . Interventions/Gb: 7
- . Contamination notes: "No contamination was detected by Blobtoolkit which was run on the diploid assembly"
- . Other observations: "This assembly was obtained with Hifiasm from ONT data. Given the high quality of both haplotypes, curation was done in diploid mode. We have shared the curated and painted pretext map corresponding to the diploid assembly (after curating each hap separately and combining them). The sex SUPER was initially mistakenly labelled as X, but this is corrected in the round corresponding to diploid curation. The Z was confirmed by alignment to other evolutionarily close species."

# Quality metrics table

Metrics	Pre-curation hap1	Pre-curation hap2	Curated hap1	Curated hap2
Total bp	596,105,788	594,728,036	596,998,386	593,835,838
GC %	39.93	39.91	39.94	39.9
Gaps/Gbp	6.71	6.73	1.68	15.16
Total gap bp	800	800	200	1,800
Scaffolds	32	30	32	29
Scaffold N50	22,445,820	22,081,071	22,445,820	21,960,911
Scaffold L50	12	12	12	12
Scaffold L90	24	24	24	24
Contigs	36	34	33	38
Contig N50	21,511,819	21,256,303	22,445,820	20,976,931
Contig L50	13	13	12	13
Contig L90	26	26	24	28
QV	70.5765		70.5765	70.5873
Kmer compl.	96.8579		96.8579	96.7613
BUSCO sing.	99.0%	98.9%	99.0%	98.9%
BUSCO dupl.	0.2%	0.2%	0.2%	0.2%
BUSCO frag.	0.5%	0.5%	0.5%	0.5%
BUSCO miss.	0.3%	0.4%	0.3%	0.4%

BUSCO: 5.5.0 (euk\_genome\_met, metaeuk) / Lineage: arthropoda\_odb10 (genomes:90, BUSCOs:1013)

# HiC contact map of curated assembly

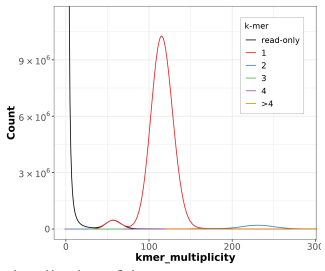


hap1 [LINK]

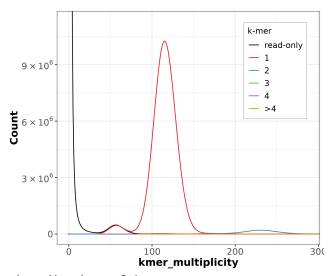
hap2 HiC PNG is missing!

hap2 File link is missing!

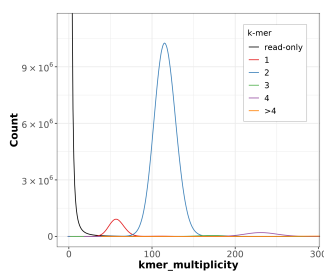
# K-mer spectra of curated assembly



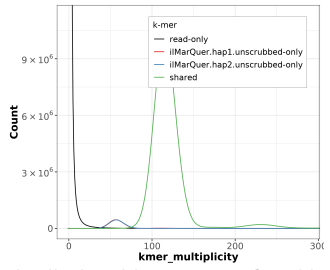
Distribution of k-mer counts per copy numbers found in **hap1** (hapl.)



Distribution of k-mer counts per copy numbers found in **hap2** (hapl.)

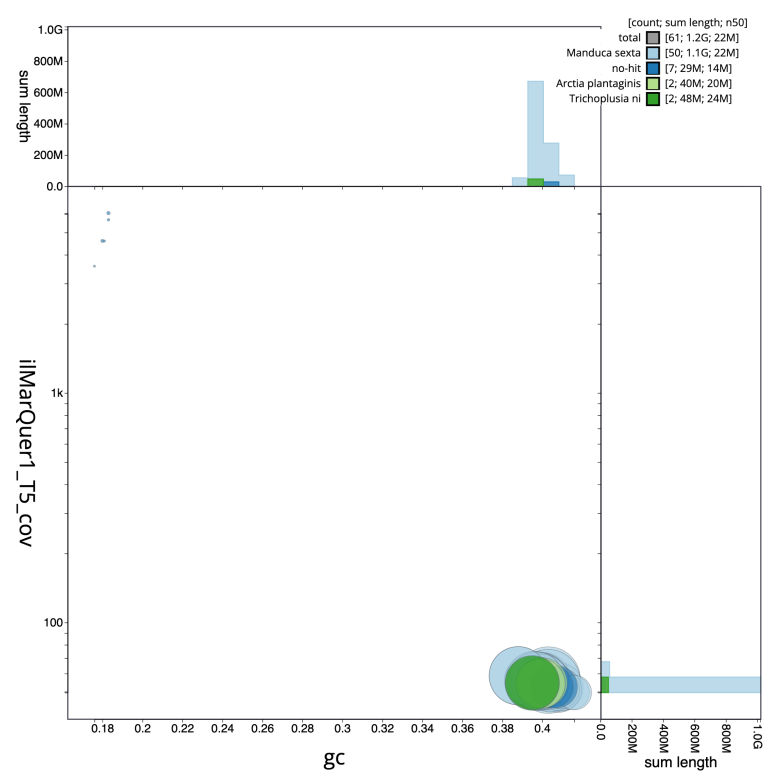


Distribution of k-mer counts per copy numbers found in asm (dipl.)



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

## Post-curation contamination screening



hap1. Bubble plot circles are scaled by sequence length, positioned by coverage and GC proportion, and coloured by taxonomy. Histograms show total assembly length distribution on each axis.

## Data profile

Data	ONT	Illumina	Omni-C
Coverage	117	72	139

# Assembly pipeline

# Curation pipeline

```
- Blobtoolkit Nextflow pipeline
|_ ver: 0.6
|_ key param: NA
- PretextViewAI
```

|\_ ver: 1.0.3 |\_ key param: NA

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