ERGA Assembly Report

v24.04.03_beta

Tags: ERGA-BGE

TxID	184164	
ToLID	rMalMon1	
Species	Malpolon monspessulanus	
Class	Lepidosauria	
Order	Squamata	

Genome Traits	Expected	Observed
Haploid size (bp)	1,613,706,619	1,752,148,786
Haploid Number	21 (source: direct)	22
Ploidy	3 (source: ancestor)	2
Sample Sex	ZW	ZW

EBP metrics summary and curation notes

Obtained EBP quality metric for collapsed: 7.8.Q48

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

- . Observed Haploid Number is different from Expected
- . Observed Ploidy is different from Expected

Curator notes

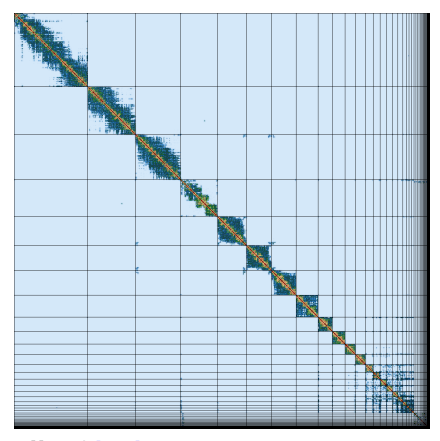
- . Interventions/Gb: 7
- . Contamination notes: "There seem to be no significant contamination of this assembly"
- . Other observations: "For this particular species we decided to scaffold and curate after the hypo polishing step rather than following purge_dups as we do for most species. We adopted this strategy because we noticed that we were missing all of the W chromosome from our final scaffolded assembly after running purge_dups. However upon alignment of the sequences eliminated by purge_dups against NR, we found that they all aligned to the W chromosome of another Sauropsida (Vipera ursinii) from the southern Mediterranean that diverged from the Montpellier snake roughly 38 MYA. We also used a recent publication on this snake from 2023 (https://www.iris.unina.it/retrieve/19087cf1-7fb4-4df2-8eb2-ff00e72c3809/2023_Mezzasalma%20et%20al_Eur%20Zool%20J_C omparative%20cytogenetics%20Hemorrhois%20and%20Malpolon.pdf) to help guide us on the number of chromosomes we should expect."

Quality metrics table

Metrics	Pre-curation collapsed	Curated collapsed	
Total bp	1,753,287,546	1,752,148,786	
GC %	42.26	42.27	
Gaps/Gbp	39.92	42.23	
Total gap bp	14,000	14,800	
Scaffolds	66	63	
Scaffold N50	121,923,953	121,923,953	
Scaffold L50	5	5	
Scaffold L90	16	16	
Contigs	136	137	
Contig N50	32,743,049	32,743,049	
Contig L50	13	13	
Contig L90	58	58	
QV	48.5441	48.5443	
Kmer compl.	95.4836	95.4648	
BUSCO sing.	95.0%	95.0%	
BUSCO dupl.	0.9%	0.9%	
BUSCO frag.	1.1%	1.1%	
BUSCO miss.	3.0%	3.0%	

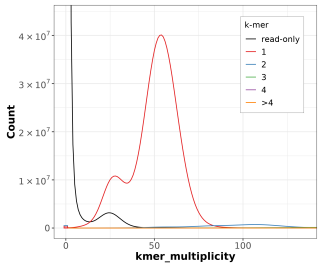
BUSCO 5.4.0 Lineage: tetrapoda_odb10 (genomes:38, BUSCOs:5310)

HiC contact map of curated assembly

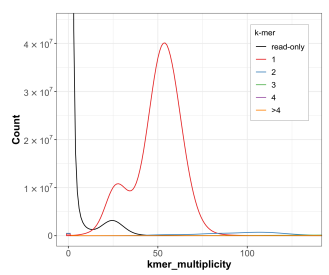


collapsed [LINK]

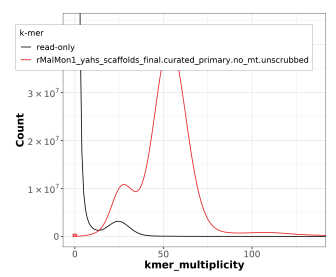
K-mer spectra of curated assembly



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

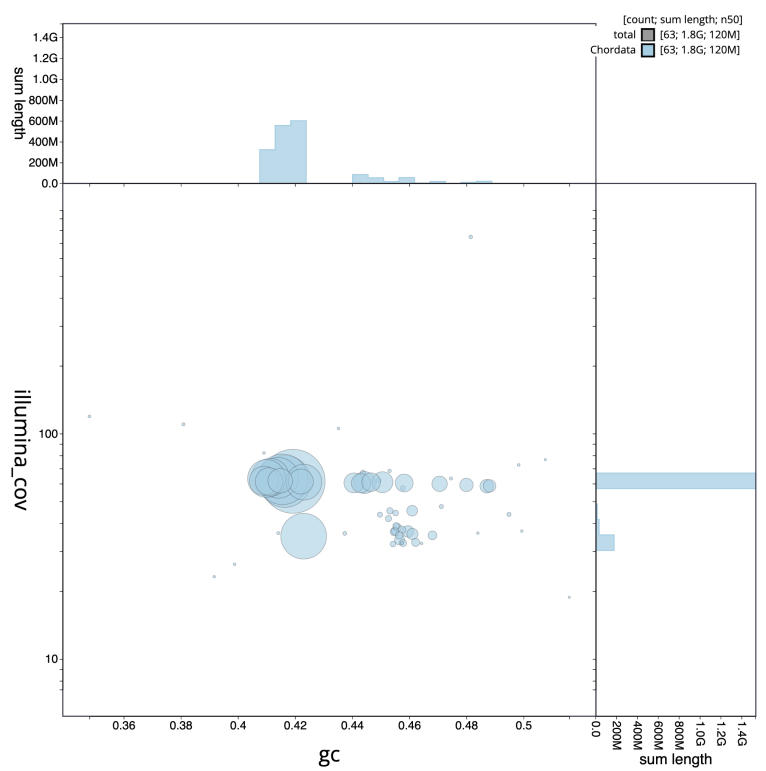


Distribution of k-mer counts per copy numbers found in asm



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

Post-curation contamination screening



collapsed. 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	OmniC
Coverage	93x	62x	51x

Assembly pipeline

```
- Trim_Galore
   |_ ver: 0.6.7
    |_ key param: "--gzip -q 20"
    _ key param: "--paired"
    |_ key param: "--retain_unpaired"
- Filtlong
   _ ver: 0.2.1
   _ key param: NA
- nextdenovo
   |_ ver: 2.5.0
    _ key param: NA
hypo
   _ ver: 1.0.3
   _ key param: NA
- YaHS
   _ ver: 1.2a
   _ key param: NA
```

Curation pipeline

PretextView

|_ ver: 0.2.5 |_ key param: NA

Submitter: Francisco Camara Affiliation: CNAG Barcelona

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