### ERGA Assembly Report

v24.10.15

Tags: ERGA-BGE

| TxID    | 387912                |  |
|---------|-----------------------|--|
| ToLID   | mCroZim1              |  |
| Species | Crocidura zimmermanni |  |
| Class   | Mammalia              |  |
| Order   | Eulipotyphla          |  |

| Genome Traits     | Expected             | Observed      |
|-------------------|----------------------|---------------|
| Haploid size (bp) | 2,224,927,356        | 2,571,728,606 |
| Haploid Number    | 17 (source: direct)  | 20            |
| Ploidy            | 2 (source: ancestor) | 2             |
| Sample Sex        | XX                   | XX            |

### EBP metrics summary and curation notes

Obtained EBP quality metric for collapsed: 7.8.Q62

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

. Observed Haploid Number is different from Expected

#### Curator notes

- . Interventions/Gb: 2
- . Contamination notes: ""
- . Other observations: "The assembly of Crocidura zimmermanni (mCroZiml.1) is based on 37X PacBio data and 189X Arima Hi-C data generated as part of the European Reference Genome Atlas (ERGA, https://www.erga-biodiversity.eu/) via the Biodiversity Genomics Europe project (BGE, https://biodiversitygenomics.eu/). The assembly process included the following steps: initial PacBio assembly generation with Hifiasm, removal of contaminant sequences using Context, removal of haplotypic duplications using purge\_dups, and Hi-C-based scaffolding with YaHS. In total, 8 contigs were identified as contaminants (bacterial), totaling 690.5 Kb (with the largest being 117.9 Kb); TAPAs has classified two scaffolds (scaffold\_102 & scaffold\_138) as bacteria, but they are host rDNA, therefore they are not removed. Additionally, 195 regions totaling 76.2 Mb (with the largest being 20.8 Mb) were identified as haplotypic duplications and removed. The mitochondrial genome was assembled using OATK. Finally, the primary assembly was analyzed and manually improved using Pretext. During manual curation, no regions were tagged as allelic duplications nor contaminants ; 24 sequences were tagged Unloc, their orientation is uncertain; the coverage of mCroZim1\_5 seems lower than the others and its alignment with Suncus\_etruscus (CM043992.1 chromosome\_X) suggests that mCroZim1\_5 is the X chromosome. the

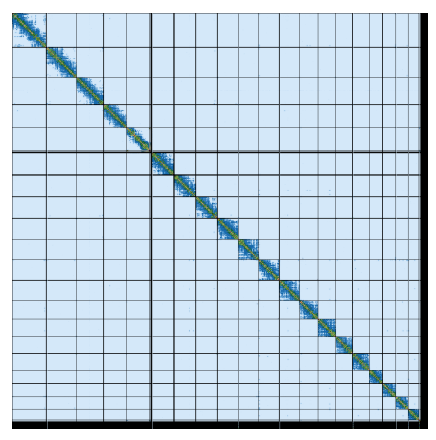
difference in expected vs observed haploid number is noteworthy, but there is a great variability of karyotypes within this genus and also between individuals of the same species; also, Crocidura zimmermanni is endemic (Crete) and shows a threatened status (Endangered), which could also contribute. Despite counting with karyotype information, that information is rather limited for this species; there is no evidence of false-joins with the contact map. Chromosome-scale scaffolds confirmed by Hi-C data were named in order of size. "

# Quality metrics table

| Metrics      | Pre-curation collapsed | Curated<br>collapsed |
|--------------|------------------------|----------------------|
| Total bp     | 2,571,745,479          | 2,571,728,606        |
| GC %         | 39.85                  | 39.85                |
| Gaps/Gbp     | 62.6                   | 62.6                 |
| Total gap bp | 16,100                 | 16,400               |
| Scaffolds    | 151                    | 146                  |
| Scaffold N50 | 127,229,103            | 127,229,103          |
| Scaffold L50 | 9                      | 9                    |
| Scaffold L90 | 18                     | 18                   |
| Contigs      | 312                    | 307                  |
| Contig N50   | 28,351,139             | 28,351,139           |
| Contig L50   | 25                     | 25                   |
| Contig L90   | 97                     | 97                   |
| QV           | 62.0833                | 62.0867              |
| Kmer compl.  | 92.9631                | 92.9629              |
| BUSCO sing.  | 90.6%                  | 90.6%                |
| BUSCO dupl.  | 2.2%                   | 2.2%                 |
| BUSCO frag.  | 0.9%                   | 0.9%                 |
| BUSCO miss.  | 6.3%                   | 6.3%                 |

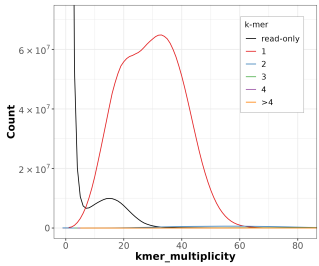
BUSCO: 5.4.3 (euk\_genome\_met, metaeuk) / Lineage: laurasiatheria\_odb10 (genomes:52, BUSCOs:12234)

HiC contact map of curated assembly

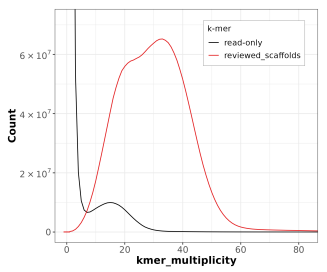


collapsed [LINK]

# K-mer spectra of curated assembly

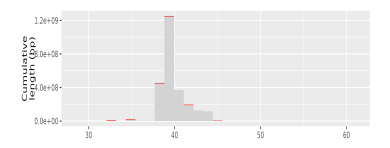


Distribution of k-mer counts per copy numbers found in asm

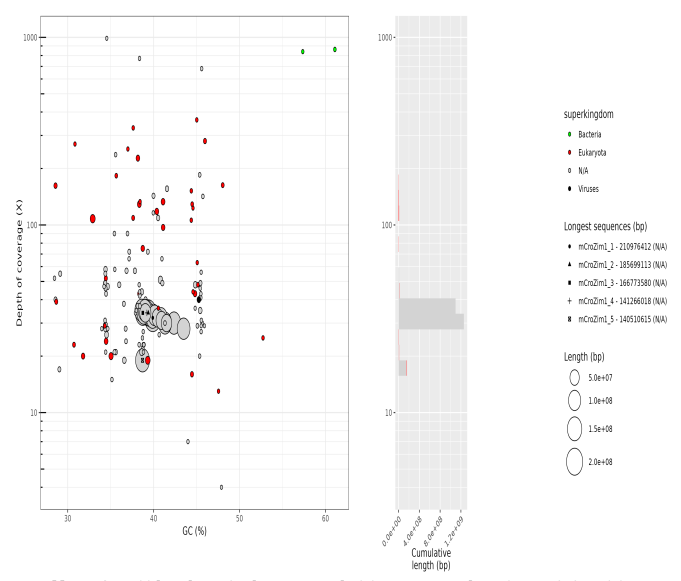


Distribution of k-mer counts coloured by their presence in reads/assemblies

# Post-curation contamination screening



## TAPAs summary Graph



**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     | PACBIO Hifi | Arima |
|----------|-------------|-------|
| Coverage | 37          | 189   |

### Assembly pipeline

```
- Hifiasm
```

|\_ ver: 0.19.5-r593 |\_ key param: NA

### - purge\_dups

|\_ ver: 1.2.5 |\_ key param: NA

#### - YaHS

|\_ ver: 1.2 |\_ key param: NA

### Curation pipeline

#### - PretextMap

|\_ ver: 0.1.9 |\_ key param: NA

#### - PretextView

|\_ ver: 0.2.5 |\_ key param: NA

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