ERGA Assembly Report

v24.04.03_beta

Tags: ERGA-Pilot

TxID	71183		
ToLID	fAnaHis1		
Species	Anaecypris hispanica		
Class	Actinopteri		
Order	Cypriniformes		

Genome Traits	Expected	Observed
Haploid size (bp)	968,863,844	984,733,391
Haploid Number	25 (source: direct)	25
Ploidy	2 (source: ancestor)	2
Sample Sex	unknown	unknown

EBP metrics summary and curation notes

Obtained EBP quality metric for hap1: 6.7.Q60 Obtained EBP quality metric for hap2: 6.7.Q59

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

Curator notes

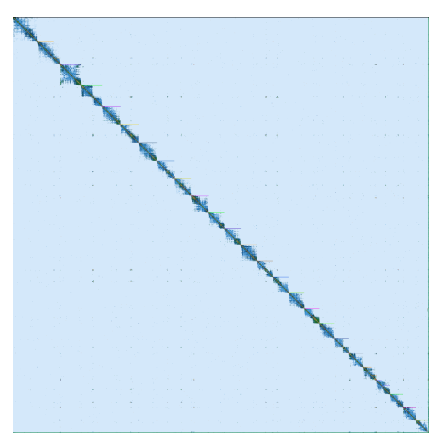
- . Interventions/Gb: None
- . Contamination notes: "No contaminants detected by FCS-GX. Those highlighted by blobtools are very short, repetitive contigs, which FCS either masked or marked as low-coverage hits to other taxa."

Quality metrics table

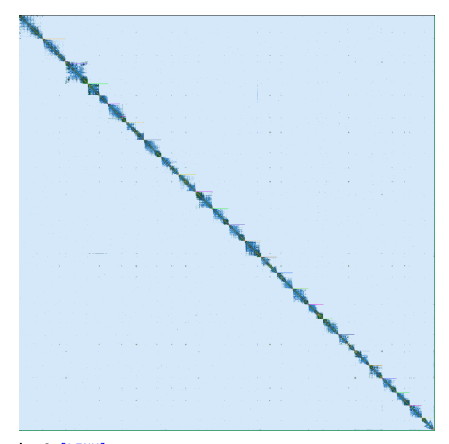
Metrics	Pre-curation hap1	Pre-curation hap2	Curated hap1	Curated hap2
Total bp	983,738,696	984,878,810	983,020,590	984,733,391
GC %	38.78	38.78	38.78	38.78
Gaps/Gbp	471.67	349.28	475.07	352.38
Total gap bp	191,962	36,080	189,282	36,340
Scaffolds	52	41	49	40
Scaffold N50	38,642,748	38,563,620	38,642,748	38,563,620
Scaffold L50	11	11	11	11
Scaffold L90	22	22	22	22
Contigs	516	385	516	387
Contig N50	4,199,513	5,445,377	4,199,513	5,445,377
Contig L50	75	57	75	57
Contig L90	262	189	263	189
QV	60.7852	59.6049	60.7852	59.6049
Kmer compl.	91.6665	91.5819	91.6665	91.5819
BUSCO sing.	97.4%	97.5%	97.4%	97.5%
BUSCO dupl.	0.7%	0.6%	0.7%	0.6%
BUSCO frag.	0.7%	0.6%	0.7%	0.6%
BUSCO miss.	1.2%	1.3%	1.2%	1.3%

BUSCO 5.7.1 Lineage: actinopterygii_odb10 (genomes:26, BUSCOs:3640)

HiC contact map of curated assembly

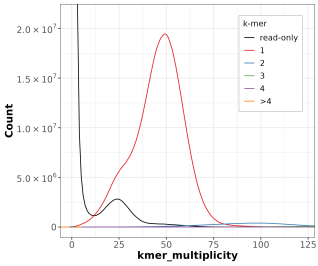


hap1 [LINK]

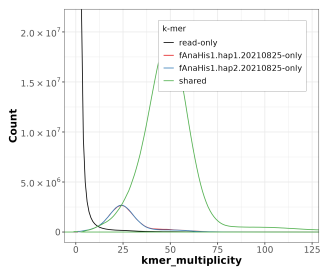


hap2 [LINK]

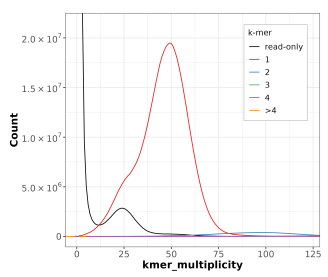
K-mer spectra of curated assembly



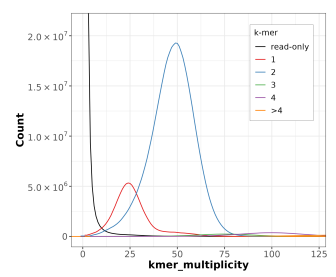
Distribution of k-mer counts per copy numbers found in **hap2** (hapl.)



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

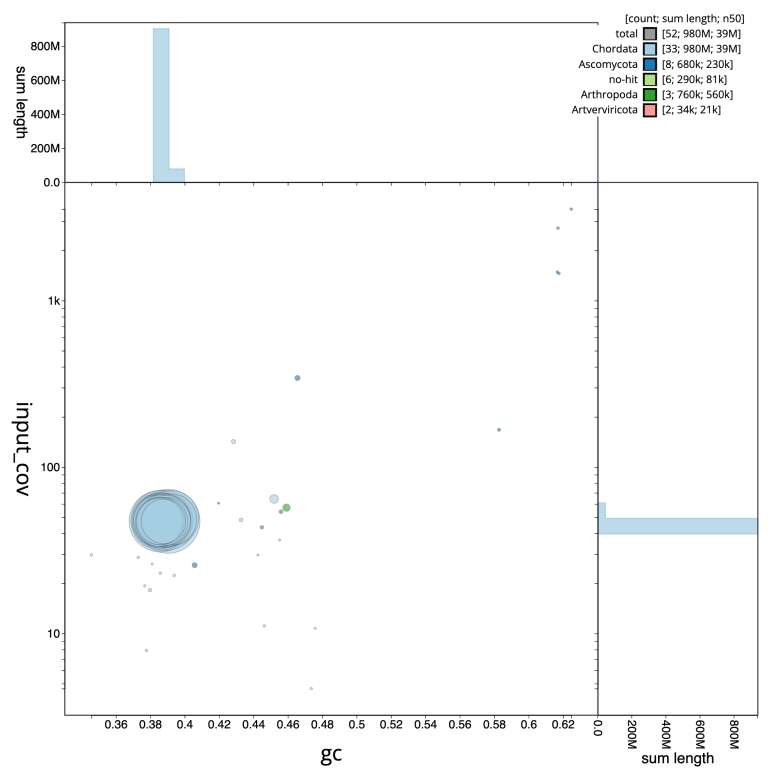


Distribution of k-mer counts per copy numbers found in **hap1** (hapl.)

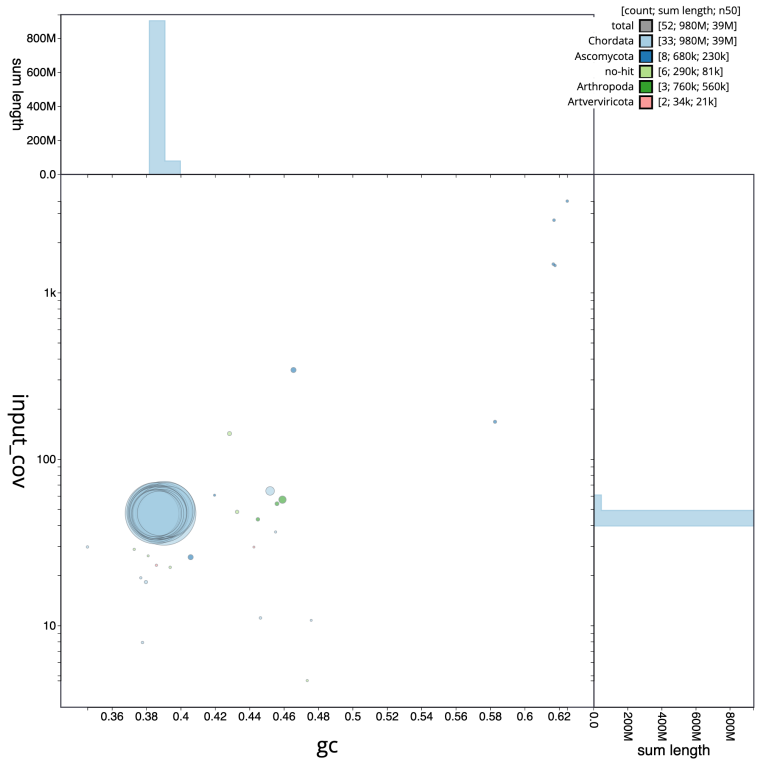


Distribution of k-mer counts per copy numbers found in asm (dipl.)

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.



hap2. 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 HiC
Coverage	50	50

Assembly pipeline

Curation pipeline

Submitter: Tom Brown

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Date and time: 2024-06-10 14:23:28 CEST