ERGA Assembly Report

v07.07.23_beta

Tags: ERGA-test

ToLID	rCarCar2		
Species	Caretta caretta		
Class	NA		
Order	Testudines		
Haploid Number	27 (source: direct)		
Ploidy	2		

Data profile

Data	Coverage
HiFi	40x
Bionano	NA
OmniC	90x

Pipeline summary

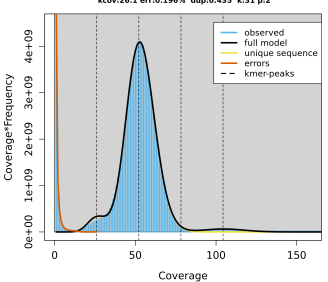
Tool	Version		
GenomeScope	2.0		
Smudgeplot	0.2.5		
Hifiasm	0.19.4		
purge_dups	1.2.6		
Bionano	Galaxy_3.7.0		
YaHS	1.1		

Genome profiling

Estimated Haploid Length	2,188,220,914		
Heterozygosity rate	0.171093%		
Kmer coverage	26.08		
Proposed ploidy	2		

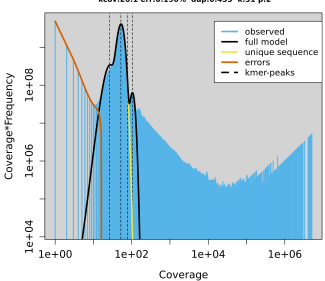
GenomeScope Profile

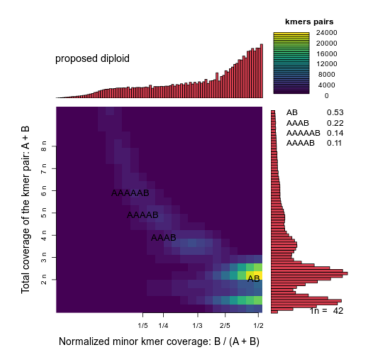
len:2,189,749,816bp uniq:82.2% aa:99.8% ab:0.176% kcov:26.1 err:0.196% dup:0.435 k:31 p:2

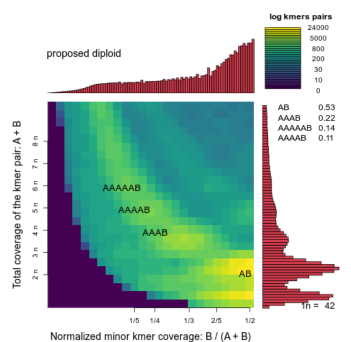


GenomeScope Profile

len:2,189,749,816bp uniq:82.2% aa:99.8% ab:0.176% kcov:26.1 err:0.196% dup:0.435 k:31 p:2





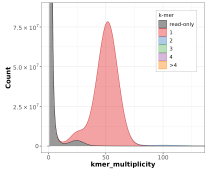


Genome assembly: contigging

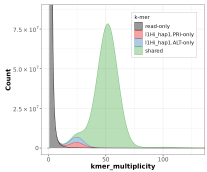
Metrics	Hifiasm hapl	Hifiasm hap2	purge_dups hap1	purge_dups hap2
Total bp	2257253306	2239458145	2216692574	2205561400
GC %	44.27	44.23	44.12	44.11
Gaps	0	0	0	0
Gaps/Gbp	0.0	0.0	0.0	0.0
Gaps bp	0	0	0	0
Scaffolds	172	144	95	84
Longest Scaf.	354291630	210263708	354291630	210263708
Scaffold auN	138591508.98	107233503.00	141094921.93	108830894.98
Scaffold N50	124322749	119638208	124322749	119638208
Scaffold L50	7	8	7	8
Scaffold L90	26	27	24	25
Contigs	172	144	95	84
Largest Cont.	354291630	210263708	354291630	210263708
Contig auN	138591508.98	107233503.00	141094921.93	108830894.98
Contig N50	124322749	119638208	124322749	119638208
Contig L50	7	8	7	8
Contig L90	26	27	24	25
QV	68.4529	67.9108	68.7149	67.9889
Kmer compl.	96.7687	96.8198	96.6829	96.6394
BUSCO sing.	96.4%	96.4%	96.4%	96.4%
BUSCO dupl.	0.9%	0.9%	0.9%	0.9%
BUSCO frag.	0.3%	0.3%	0.3%	0.3%
BUSCO miss.	2.4%	2.4%	2.4%	2.4%

Lineage: sauropsida_odb10 (genomes:76, BUSCOs:7480)

K-mer spectra: Hifiasm



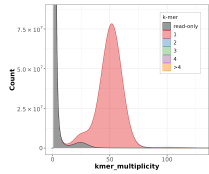
llHi_hap1_merqOutput.llHi_hap1.ALT.spectra-cn.st.png



11Hi_hap1_merqOutput.spectra-asm.st.png

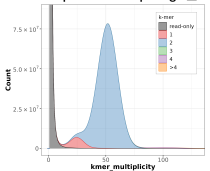
7.5 × 10⁷ 2.5 × 10⁷ 2.5 × 10⁷ 2.5 × 10⁷ 2.6 × 10⁷ 3 4 4 > 4

11Hi_hap1_merqOutput.spectra-cn.st.png

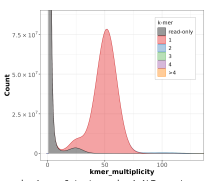


11Hi_hap1_merqOutput.11Hi_hap1.PRI.spectra-cn.st.png

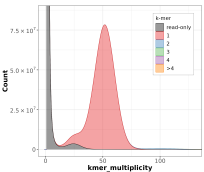
K-mer spectra: purge_dups



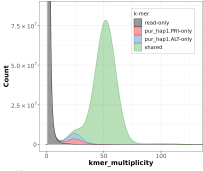
pur_hap1_merqOutput.spectra-cn.st.png



pur_hap1_merqOutput.pur_hap1.ALT.spectra-cn.st.png



pur_hap1_merqOutput.pur_hap1.PRI.spectra-cn.st.png



pur_hap1_merqOutput.spectra-asm.st.png

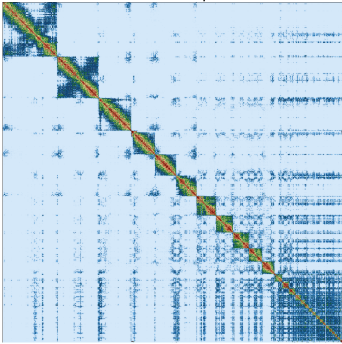
Genome assembly: scaffolding

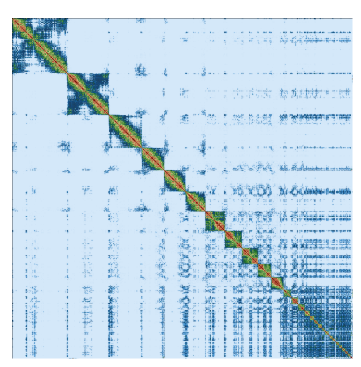
Metrics	Bionano hap1	Bionano hap2	YaHS hap1	YaHS hap2
Mecrics	парт	парг	парт	парг
Total bp	2220191759	2207617369	2220195358	2207621769
GC %	44.12	44.11	44.12	44.11
Gaps	38	27	55	49
Gaps/Gbp	17.12	12.23	24.77	22.2
Gaps bp	3499185	2055969	3502772	2060369
Scaffolds	57	57	41	35
Longest Scaf.	355489124	210263708	355489123	356830969
Scaffold auN	144780868.66	114892365.06	167596863.48	167403878.03
Scaffold N50	127105400	126644623	137541926	137610697
Scaffold L50	7	8	5	5
Scaffold L90	21	22	18	18
Contigs	95	84	96	84
Largest Cont.	354291630	210263708	355007410	210263708
Contig auN	141094921.93	108830894.98	141322292.78	108830894.98
Contig N50	124322749	119638208	124322749	119638208
Contig L50	7	8	7	8
Contig L90	24	25	24	25
QV	68.7149	67.9889	68.7149	67.9889
Kmer compl.	96.6829	96.6394	96.6829	96.6394
BUSCO sing.	96.4%	96.4%	96.4%	96.4%
BUSCO dupl.	0.9%	0.9%	0.9%	0.8%
BUSCO frag.	0.4%	0.4%	0.4%	0.4%
BUSCO miss.	2.3%	2.3%	2.3%	2.4%

Lineage: sauropsida_odb10 (genomes:76, BUSCOs:7480)

Pretext Full Map: Bionano Data not available

Pretext Full Map: YaHS





yahs_hap1FullMap.png

yahs_hap2FullMap.png

Submitter: Diego De Panis Affiliation: BeGenDiv

Date and time: 2023-07-07 22:38:40 CEST