	GCF_003367295.1_ASM336729v1_genomic
# contigs (>= 0 bp)	16
# contigs (>= 1000 bp)	16
# contigs (>= 5000 bp)	16
# contigs (>= 10000 bp)	16
# contigs (>= 25000 bp)	9
# contigs (>= 50000 bp)	2
Total length (>= 0 bp)	1280240
Total length (>= 1000 bp)	1280240
Total length (>= 5000 bp)	1280240
Total length (>= 10000 bp)	1280240
Total length (>= 25000 bp)	1161007
Total length (>= 50000 bp)	962310
# contigs	16
Largest contig	908512
Total length	1280240
Reference length	1521208
GC (%)	28.24
Reference GC (%)	28.18
N50	908512
NG50	908512
N90	26507
NG90	-
auN	653097.9
auNG	549643.5
L50	1
LG50	1
L90	9
LG90	-
# misassemblies	16
# misassembled contigs	6
Misassembled contigs length	133618
# local misassemblies	7
# scaffold gap ext. mis.	0
# scaffold gap loc. mis.	0
# unaligned mis. contigs	3
# unaligned contigs	
	0 + 12 part
Unaligned length	0 + 12 part 123725
Unaligned length Genome fraction (%)	
	123725
Genome fraction (%)	123725 75.032
Genome fraction (%) Duplication ratio	123725 75.032 1.009
Genome fraction (%) Duplication ratio # N's per 100 kbp	123725 75.032 1.009
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp	123725 75.032 1.009 0.00 758.69
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp	123725 75.032 1.009 0.00 758.69 44.05
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50 NA90	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50 NA90 NGA90	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648 904648
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50 NA90 NGA90 auNA	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648 904648
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50 NA90 NGA90 auNA auNGA	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648 904648 643463.5 541535.2
Genome fraction (%) Duplication ratio # N's per 100 kbp # mismatches per 100 kbp # indels per 100 kbp # genomic features Largest alignment Total aligned length NA50 NGA50 NA90 NGA90 auNA auNGA	123725 75.032 1.009 0.00 758.69 44.05 1183 + 49 part 904648 1151059 904648 904648 643463.5 541535.2

All statistics are based on contigs of size >= 500 bp, unless otherwise noted (e.g., "# contigs (>= 0 bp)" and "Total length (>= 0 bp)" include all contigs).

Misassemblies report

Г	GCF_003367295.1_ASM336729v1_genomic
# misassemblies	16
# contig misassemblies	16
# c. relocations	3
# c. translocations	13
# c. inversions	0
# scaffold misassemblies	0
# s. relocations	0
# s. translocations	0
# s. inversions	0
# misassembled contigs	6
Misassembled contigs length	133618
# local misassemblies	7
# scaffold gap ext. mis.	0
# scaffold gap loc. mis.	0
# unaligned mis. contigs	3
# mismatches	8733
# indels	507
# indels (<= 5 bp)	462
# indels (> 5 bp)	45
Indels length	1797

All statistics are based on contigs of size >= 500 bp, unless otherwise noted (e.g., "# contigs (>= 0 bp)" and "Total length (>= 0 bp)" include all contigs).

Unaligned report

	GCF_003367295.1_ASM336729v1_genomic
# fully unaligned contigs	0
Fully unaligned length	0
# partially unaligned contigs	12
Partially unaligned length	123725
# N's	0

All statistics are based on contigs of size >= 500 bp, unless otherwise noted (e.g., "# contigs (>= 0 bp)" and "Total length (>= 0 bp)" include all contigs).

























