# conting (>= 0 ha)	rt 1805	374
# contigs (>= 0 bp)	1805	376
# contigs (>= 1000 bp)	1805	376
# contigs (>= 5000 bp)	1805	376
# contigs (>= 10000 bp)	1805	376
# contigs (>= 25000 bp)	1800	372
# contigs (>= 50000 bp)	1773	361
Total length (>= 0 bp)	436920153	437273953
Total length (>= 1000 bp)	436920153	437273953
Total length (>= 5000 bp)	436920153	437273953
Total length (>= 10000 bp)	436920153	437273953
Total length (>= 25000 bp)	436829068	437196775
Total length (>= 50000 bp)	435805794	436784290
# contigs	1805	376
Largest contig	3341941	22675904
Total length	436920153	437273953
	430920133	43727393.
Reference length	490000000	40000000
Estimated reference length	 	49000000
GC (%)	44.99	44.99
Reference GC (%)	-	
N50	418614	14850352
NG50	346609	14090550
N90	97153	58328
NG90	-	
auN	668619.6	12187194.
auNG	596190.5	10875801.4
L50	264	13
LG50	334	15
L90	1171	6:
LG90		
# total reads	-	
# left	-	
# right	-	
Mapped (%)	-	
Reference mapped (%)	-	
Properly paired (%)	-	
Reference properly paired (%)	-	
Avg. coverage depth	- [
Reference avg. coverage depth	- 1	
Coverage >= 1x (%)	- 1	
Reference coverage >= 1x (%)	- 1	
# large blocks misassemblies	<u> </u>	
# misassemblies		
	-	
# misassembled contigs	-	
Misassembled contigs length	-	
# local misassemblies	-	
# scaffold gap ext. mis.	-	
# scaffold gap loc. mis.	-	
# structural variations	-	
# possible TEs	-	
# unaligned mis. contigs	-	
# unaligned contigs	- [
Unaligned length	- 1	
Genome fraction (%)	- 1	
Duplication ratio	- 1	
Avg contig read support	-	
# N's per 100 kbp	0.07	80.98
	0.07	
# mismatches per 100 kbp	-	
# indels per 100 kbp	-	
# genomic features	-	
# operons	-	
Complete BUSCO (%)	-	
Partial BUSCO (%)	-	
# predicted genes (unique)	-	
# predicted genes (>= 0 bp)	-	
# predicted genes (>= 300 bp)	- 1	
# predicted genes (>= 1500 bp)	- 1	
# predicted genes (>= 3000 bp)	-	
# predicted rRNA genes	-	
Largest alignment	-	
Total aligned length	-	
NA50	-	
NGA50	-	
NA90	<u> </u>	
NGA90	-	
auNA	-	
auNGA	-	
LA50	-	
LGA50	-	
LA90	-	
LGA90	_	
	-	
K-mer-based compl. (%)	 	
K mar based see les ett (00)		
K-mer-based cor. length (%) K-mer-based mis. length (%)	-	

All statistics are based on contigs of size >= 3000 bp, unless otherwise noted (e.g., "# contigs (>= 0 bp)" and "lotal length (>= 0 bp)" include all contigs).











