Report	
	mini-ava-ont_100
# contigs (>= 0 bp)	4
# contigs (>= 1000 bp)	4
# contigs (>= 5000 bp)	4
# contigs (>= 10000 bp)	4
# contigs (>= 25000 bp)	4
# contigs (>= 50000 bp)	4
Total length (>= 0 bp)	4779061
Total length (>= 1000 bp)	4779061
Total length (>= 5000 bp)	4779061
Total length (>= 10000 bp)	4779061
Total length (>= 25000 bp)	4779061
Total length (>= 50000 bp)	4779061
# contigs	4
Largest contig	3997712
Total length	4779061
Reference length	4686137
GC (%)	50.76
Reference GC (%)	50.78
N50	
	3997712
NG50	3997712
N90	561933
NG90	561933
auN	3415224.8
auNG	3482947.3
L50	:
LG50	
L90	
LG90	-
# misassemblies	(
# misassembled contigs	(
Misassembled contigs length	(
# local misassemblies	(
# scaffold gap ext. mis.	(
# scaffold gap loc. mis.	(
# unaligned mis. contigs	(
# unaligned contigs	0 + 0 par
Unaligned length	(
Genome fraction (%)	99.944
Duplication ratio	1.020
# N's per 100 kbp	0.00
# mismatches per 100 kbp	0.00
# indels per 100 kbp	0.00
Largest alignment	3997712
Total aligned length	4779063
NA50	3997712
NGA50	3997712
NA90	561933
NGA90	561933
auNA	3415224.8
····	3482947.3
auNGA	J702371
auNGA	
LA50	

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

	mini-ava-ont_100
# misassemblies	0
# contig misassemblies	0
# c. relocations	0
# c. translocations	0
# c. inversions	0
# scaffold misassemblies	0
# s. relocations	0
# s. translocations	0
# s. inversions	0
# misassembled contigs	0
Misassembled contigs length	0
# local misassemblies	0
# scaffold gap ext. mis.	0
# scaffold gap loc. mis.	0
# unaligned mis. contigs	0
# mismatches	0
# indels	0
# indels (<= 5 bp)	0
# indels (> 5 bp)	0
Indels length	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).

Unaligned report

	mini-ava-ont_100
# fully unaligned contigs	0
Fully unaligned length	0
# partially unaligned contigs	0
Partially unaligned length	0
# 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).



















