

## Report

	final.contigs
# contigs (>= 0 bp)	844
# contigs (>= 1000 bp)	482
Total length (>= 0 bp)	1200225
Total length (>= 1000 bp)	930688
# contigs	844
Largest contig	8031
Total length	1200225
Reference length	615980
GC (%)	25.34
Reference GC (%)	25.33
N50	1711
NG50	2587
N75	1058
NG75	1999
L50	230
LG50	88
L75	453
LG75	156
# misassemblies	28
# misassembled contigs	27
Misassembled contigs length	66201
# local misassemblies	0
# unaligned contigs	0 + 2 part
Unaligned length	916
Genome fraction (%)	96.530
Duplication ratio	2.017
# N's per 100 kbp	0.00
# mismatches per 100 kbp	1652.02
# indels per 100 kbp	2.52
Largest alignment	8031
NA50	1666
NGA50	2425
NA75	1023
NGA75	1955
LA50	240
LGA50	92
LA75	466
LGA75	164

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

	final.contigs
# misassemblies	28
# relocations	28
# translocations	0
# inversions	0
# possibly misassembled contigs	1
# misassembled contigs	27
Misassembled contigs length	66201
# local misassemblies	0
# mismatches	9823
# indels	15
# short indels	15
# long indels	0
Indels length	15

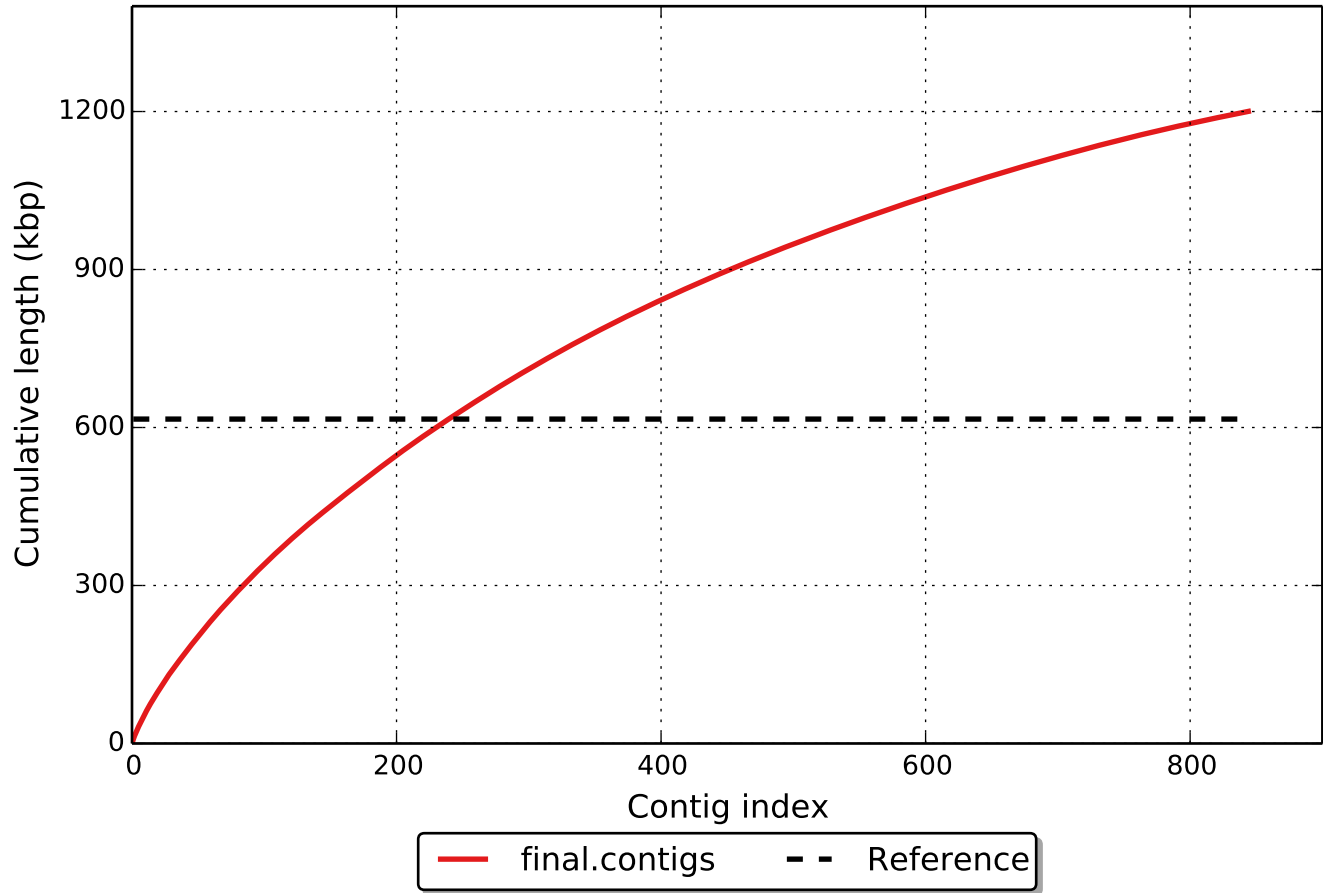
All statistics are based on contigs of size  $\geq 500$  bp, unless otherwise noted (e.g., "# contigs ( $\geq 0$  bp)" and "Total length ( $\geq 0$  bp)" include all contigs).

## Unaligned report

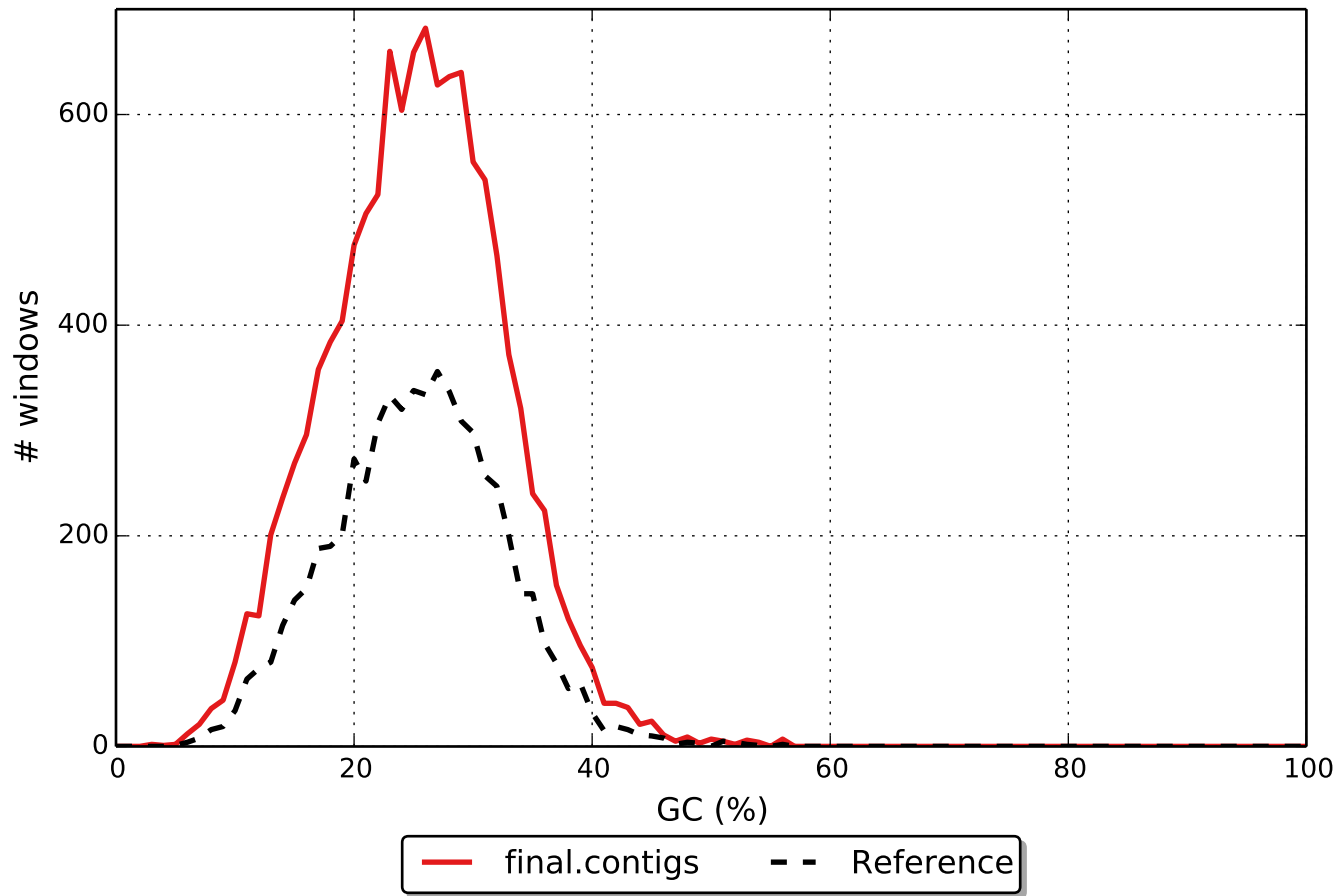
	final.contigs
# fully unaligned contigs	0
Fully unaligned length	0
# partially unaligned contigs	2
# with misassembly	0
# both parts are significant	1
Partially unaligned length	916
# N's	0

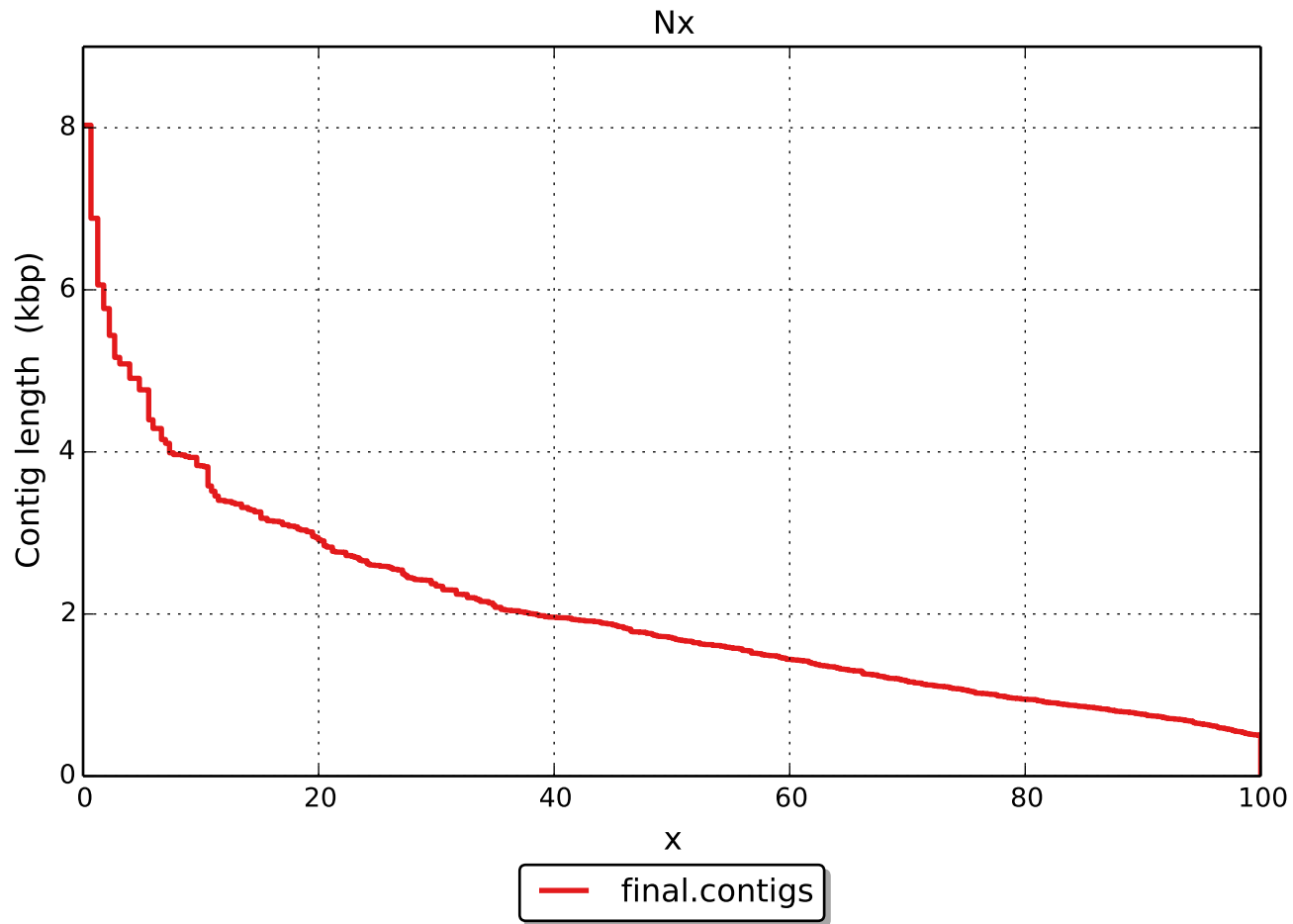
All statistics are based on contigs of size  $\geq 500$  bp, unless otherwise noted (e.g., "# contigs ( $\geq 0$  bp)" and "Total length ( $\geq 0$  bp)" include all contigs).

Cumulative length

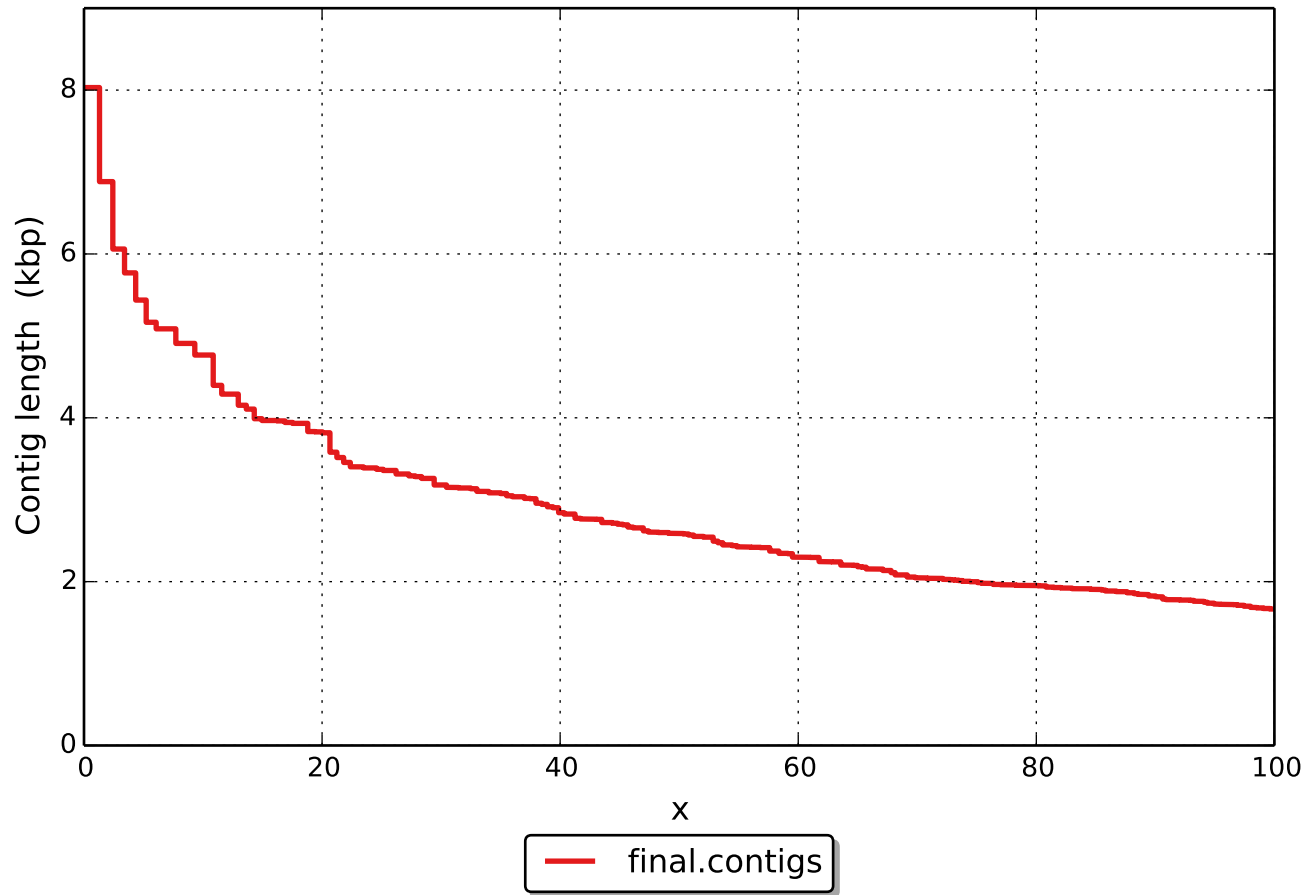


# GC content

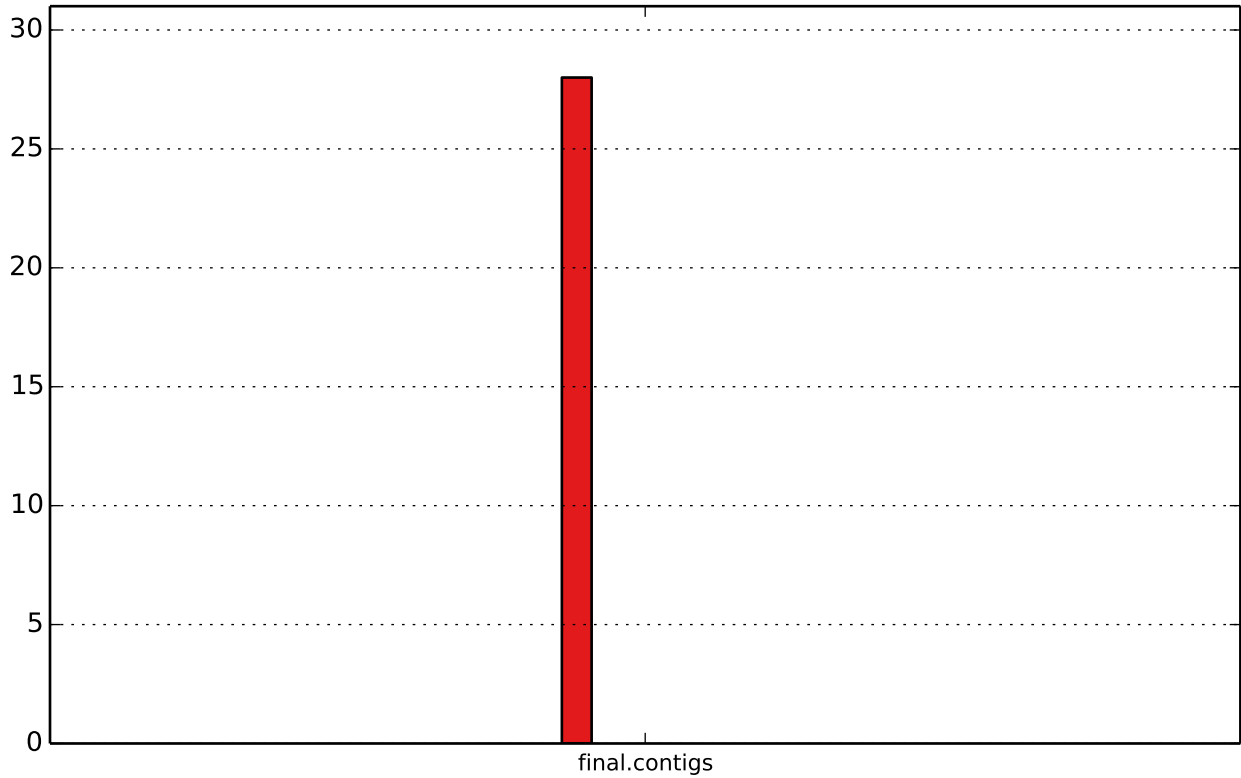




NGx



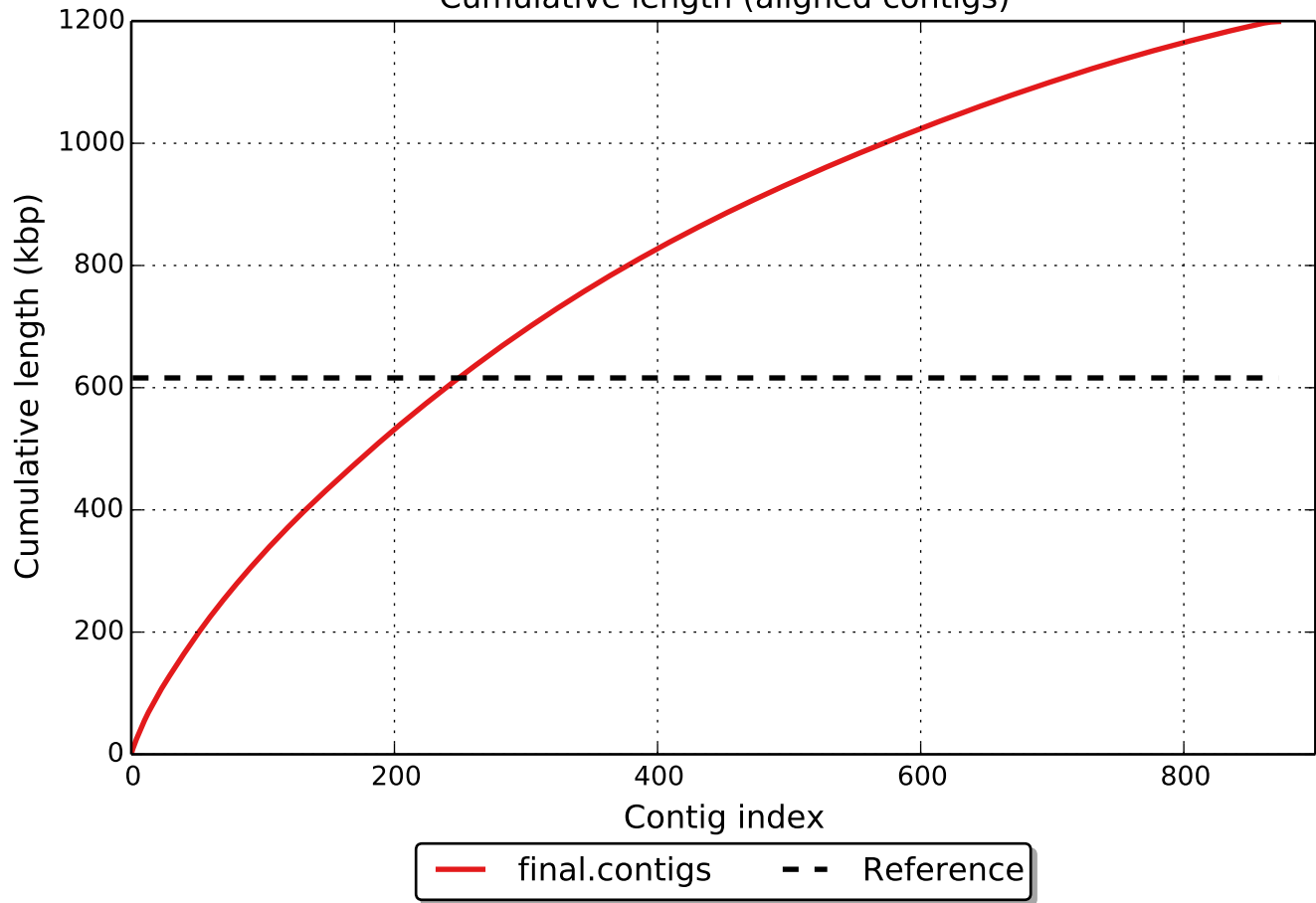
# Misassemblies



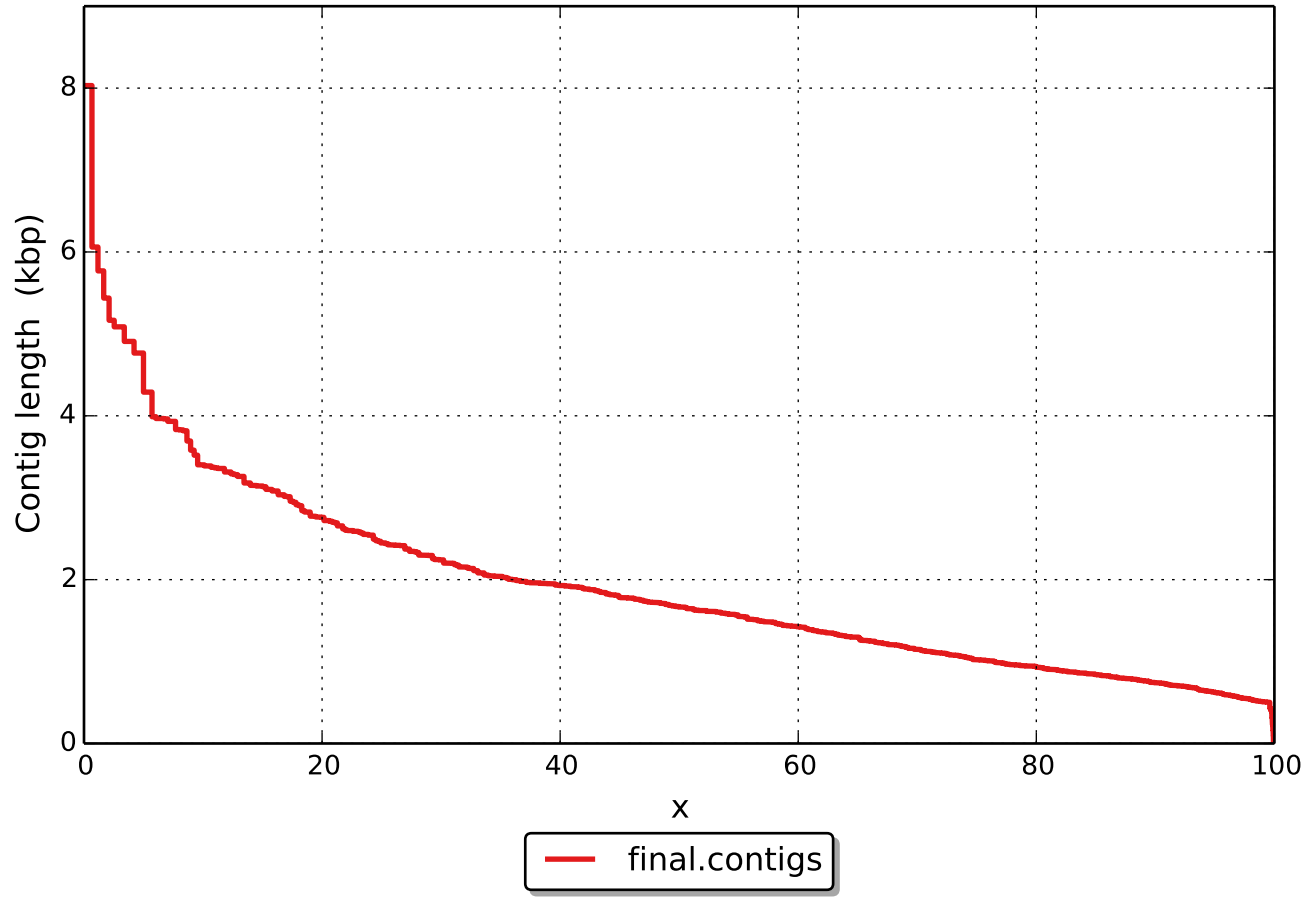
 # relocations



Cumulative length (aligned contigs)



NAx



# NGAx

