SAMPLE3 SE.AF0.75.consensus.masked SAMPLE1 PE.AF0.75.consensus.masked SAMPLE2\_PE.AF0.75.consensus.masked # contigs (>= 0 bp) 1 # contigs (>= 1000 bp) 1 1 1 1 1 1 # contigs (>= 5000 bp) # contigs (>= 10000 bp) 1 1 1 # contigs (>= 25000 bp) 1 1 1 # contigs (>= 50000 bp) 0 0 0 Total length (>= 0 bp) 29903 29903 29903 29903 29903 29903 Total length (>= 1000 bp) 29903 29903 29903 Total length (>= 5000 bp) Total length (>= 10000 bp) 29903 29903 29903 29903 Total length (>= 25000 bp) 29903 29903 Total length (>= 50000 bp) 0 0 0 # contigs 1 1 Largest contig 29903 29903 29903 29903 29903 Total length 29903 Reference length 29903 29903 29903 37.98 38.12 GC (%) 37.95 Reference GC (%) 37.97 37.97 37.97 29903 29903 N50 29903 NG50 29903 29903 29903 N75 29903 29903 29903 NG75 29903 29903 29903 L50 1 1 1 LG50 1 1 1 L75 1 1 1 LG75 1 1 1 # misassemblies 0 0 0 0 0 # misassembled contigs 0 0 0 Misassembled contigs length 0 # local misassemblies 0 0 0 # scaffold gap ext. mis. 0 0 0 2 9 # scaffold gap loc. mis. 1 0 0 # unaligned mis. contigs 0 # unaligned contigs 0 + 0 part 0 + 0 part 0 + 0 part Unaligned length 0 0 0 89.804 Genome fraction (%) 97.432 98.094 1.026 **Duplication ratio** 1.019 1.114 # N's per 100 kbp 3100.02 2193.76 10497.27 20.45 22.34 # mismatches per 100 kbp 0.00 # indels per 100 kbp 0.00 0.00 0.00 # genomic features 23 + 7 part 23 + 7 part 14 + 14 part Largest alignment 29135 29333 26854 Total aligned length 29135 29333 26854 29135 29333 26854 NA50 29135 29333 26854 NGA50 NA75 29135 29333 26854 NGA75 29135 29333 26854 LA50 1 1 1 LGA50 1 1 LA75 1 1 1 LGA75 1 1

Report

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).

## Misassemblies report

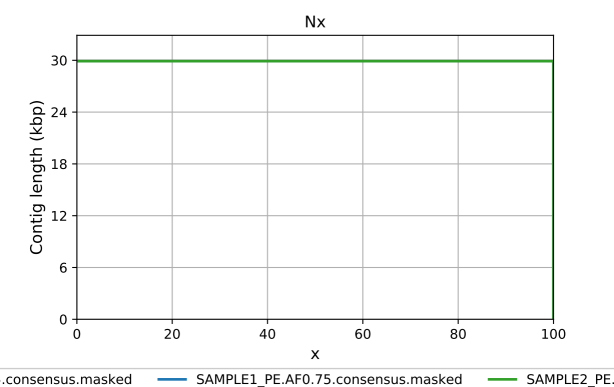
	SAMPLE3_SE.AF0.75.consensus.masked	SAMPLE1_PE.AF0.75.consensus.masked	SAMPLE2_PE.AF0.75.consensus.masked
# misassemblies	0	0	0
# contig misassemblies	0	0	0
# c. relocations	0	0	0
# c. translocations	0	0	0
# c. inversions	0	0	0
# scaffold misassemblies	0	0	0
# s. relocations	0	0	0
# s. translocations	0	0	0
# s. inversions	0	0	0
# misassembled contigs	0	0	0
Misassembled contigs length	0	0	0
# local misassemblies	0	0	0
# scaffold gap ext. mis.	0	0	0
# scaffold gap loc. mis.	2	1	9
# unaligned mis. contigs	0	0	0
# mismatches	0	6	6
# indels	0	0	0
# indels (<= 5 bp)	0	0	0
# indels (> 5 bp)	0	0	0
Indels length	0	0	0

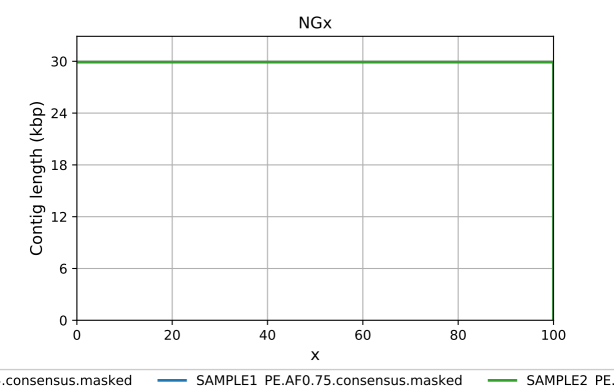
All statistics are based on contigs of size >= 500 bp, unless otherwise noted (e.g., "# contigs (>= 0 bp)" and "Total length (>= 0 bp)" include all contigs).

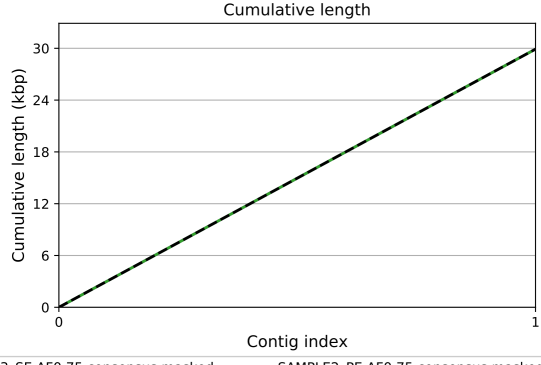
## Unaligned report

	SAMPLE3_SE.AF0.75.consensus.masked	SAMPLE1_PE.AF0.75.consensus.masked	SAMPLE2_PE.AF0.75.consensus.masked
# fully unaligned contigs	0	0	0
Fully unaligned length	0	0	0
# partially unaligned contigs	0	0	0
Partially unaligned length	0	0	0
# N's	927	656	3139

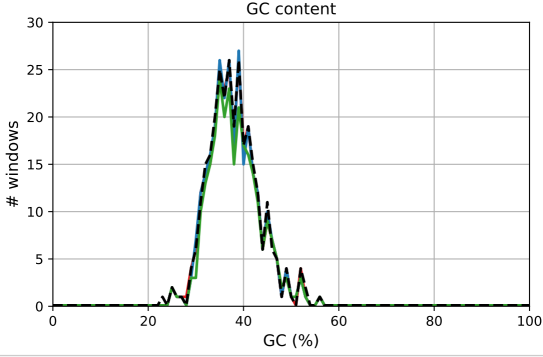
All statistics are based on contigs of size  $\geq$  500 bp, unless otherwise noted (e.g., "# contigs ( $\geq$  0 bp)" and "Total length ( $\geq$  0 bp)" include all contigs).





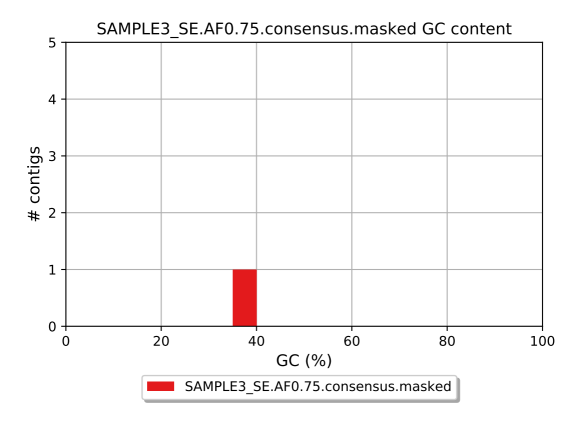


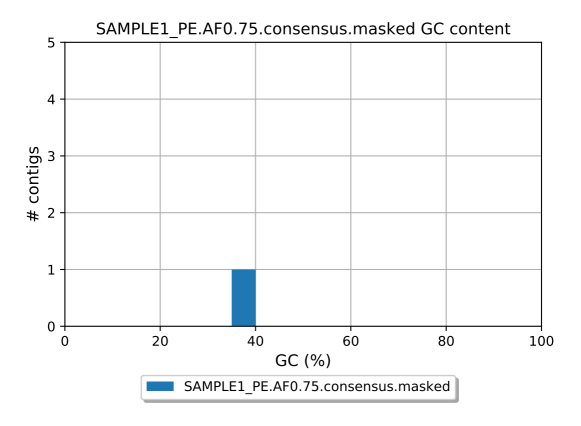
PLE3\_SE.AF0.75.consensus.masked —— SAMPLE2\_PE.AF0.75.consensus.masked PLE1\_PE.AF0.75.consensus.masked

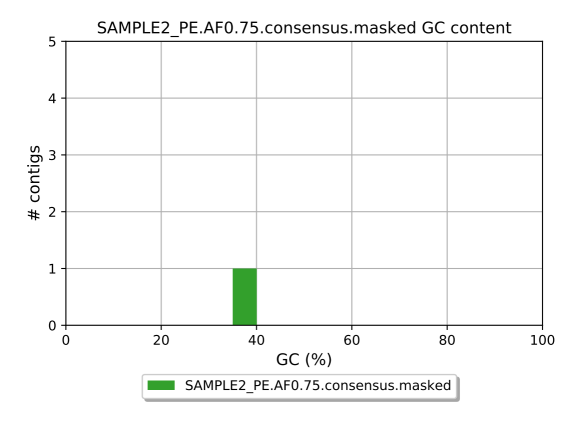


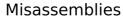
PLE3\_SE.AF0.75.consensus.masked PLE1 PE.AF0.75.consensus.masked

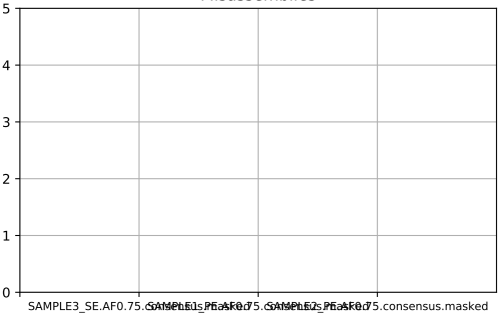
SAMPLE2 PE.AF0.75.consensus.masked

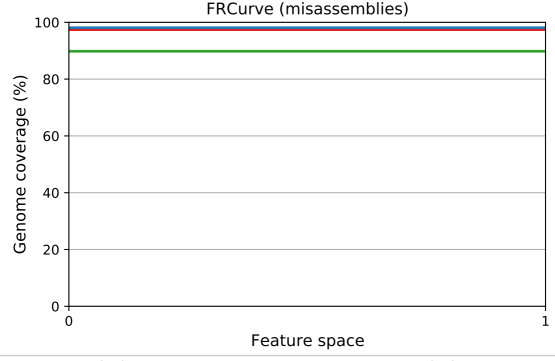




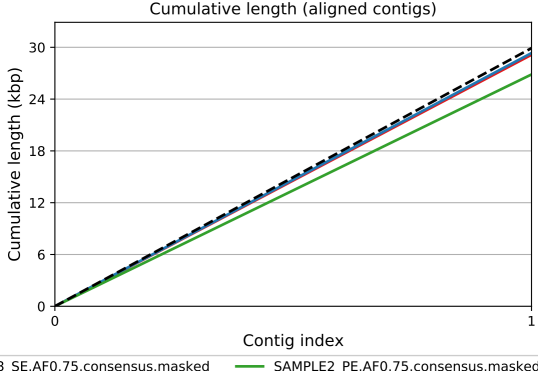




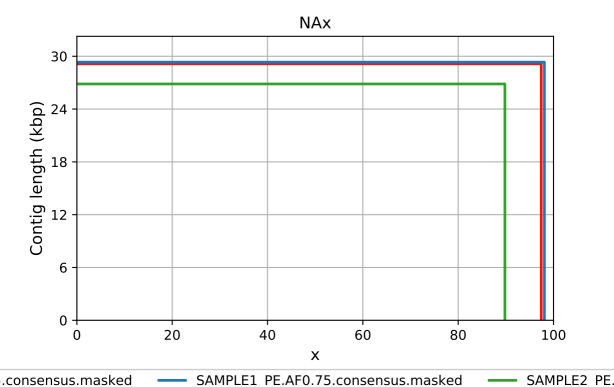


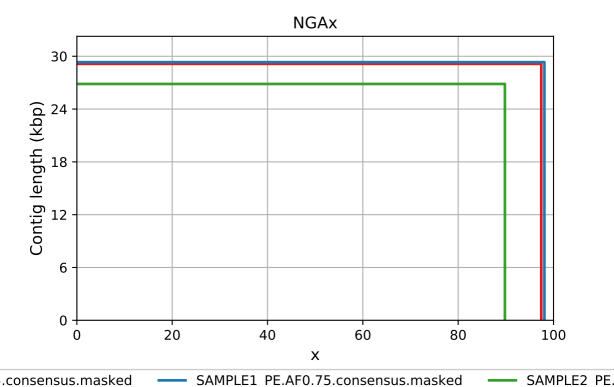


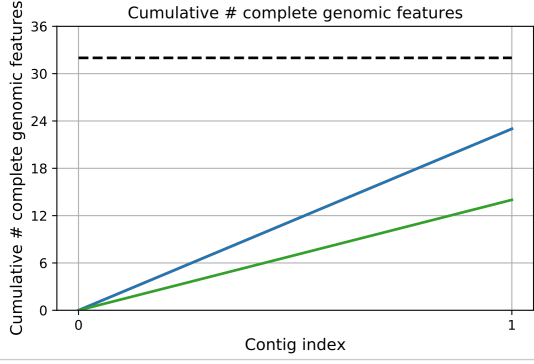
.consensus.masked —— SAMPLE1\_PE.AF0.75.consensus.masked —— SAMPLE2\_PE.



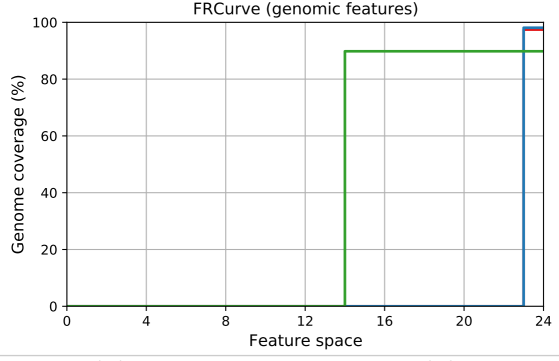
PLE3\_SE.AF0.75.consensus.masked —— SAMPLE2\_PE.AF0.75.consensus.masked PLE1 PE.AF0.75.consensus.masked







PLE3\_SE.AF0.75.consensus.masked —— SAMPLE2\_PE.AF0.75.consensus.masked PLE1\_PE.AF0.75.consensus.masked



.consensus.masked —— SAMPLE1\_PE.AF0.75.consensus.masked —— SAMPLE2\_PE.



