Qualimap Analysis Results

BAM QC analysis Generated by Qualimap v.2.2.2-dev 2022/03/02 02:58:28



1. Input data & parameters

1.1. QualiMap command line

qualimap bamqc -bam output/UF802/UF802_f4_q37_sortc_markdup.rescaled.bam -nw 400 -hm 3

1.2. Alignment

Command line:	bwa samse -r @RG\tID:UF802\tSM:UF802\tLB:nan\ tPL:ILLUMINA /data/stonelab/references/Human_mit ochondrial/NC_012920.1.fasta output/UF802/UF802-ancient.sai output/UF802/UF802- ancient.trimmed.fq
Draw chromosome limits:	no
Analyze overlapping paired-end reads:	no
Program:	bwa (0.7.17-r1188)
Analysis date:	Wed Mar 02 02:58:27 MST 2022
Size of a homopolymer:	3
Skip duplicate alignments:	no
Number of windows:	400
BAM file:	output/UF802/UF802_f4_q37_sortc_ markdup.rescaled.bam



2. Summary

2.1. Globals

Reference size	16,569
Number of reads	3
Mapped reads	3 / 100%
Unmapped reads	0 / 0%
Mapped paired reads	0 / 0%
Secondary alignments	0
Read min/max/mean length	41 / 123 / 87
Duplicated reads (estimated)	0 / 0%
Duplication rate	0%
Clipped reads	0 / 0%

2.2. ACGT Content

Number/percentage of A's	93 / 35.63%		
Number/percentage of C's	81 / 31.03%		
Number/percentage of T's	68 / 26.05%		
Number/percentage of G's	19 / 7.28%		
Number/percentage of N's	0 / 0%		
GC Percentage	38.31%		

2.3. Coverage

Mean	0.0158
Standard Deviation	0.1245



2.4. Mapping Quality

Mean Mapping Quality	0.78
wear wapping Quality	0.70

2.5. Mismatches and indels

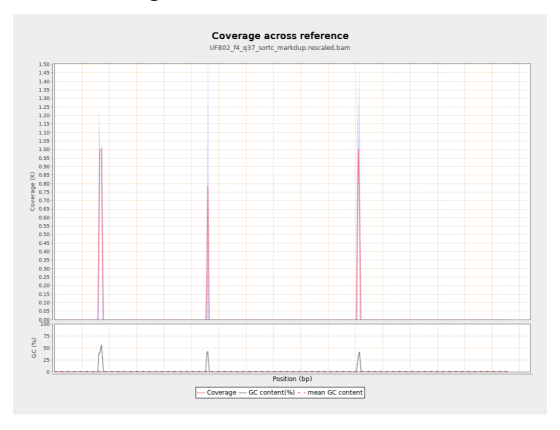
General error rate	3.45%
Mismatches	9

2.6. Chromosome stats

Name	Length	Mapped bases	Mean coverage	Standard deviation
NC_012920.1	16569	261	0.0158	0.1245

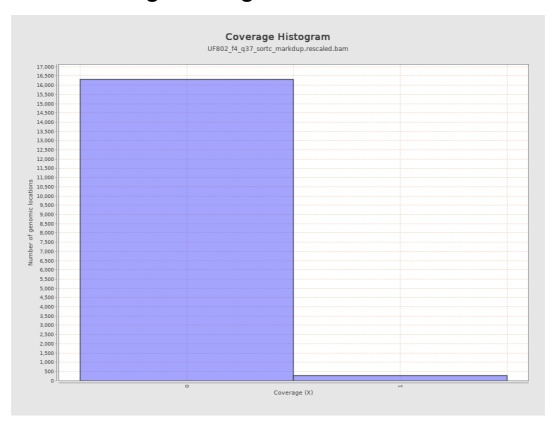


3. Results: Coverage across reference



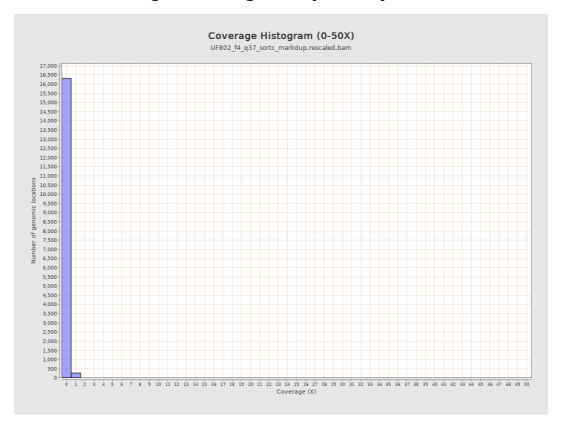


4. Results : Coverage Histogram



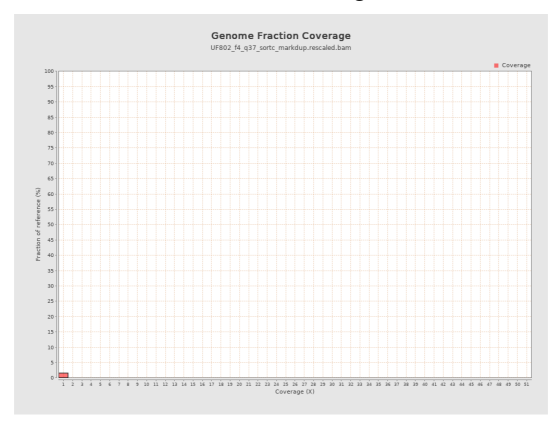


5. Results: Coverage Histogram (0-50X)



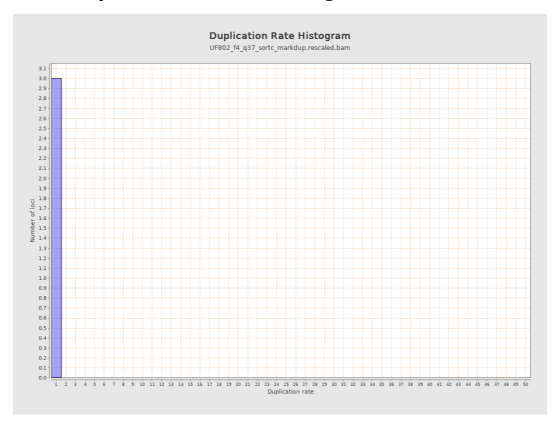


6. Results : Genome Fraction Coverage



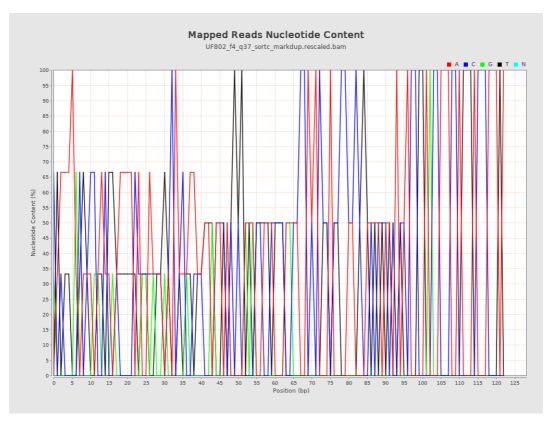


7. Results: Duplication Rate Histogram



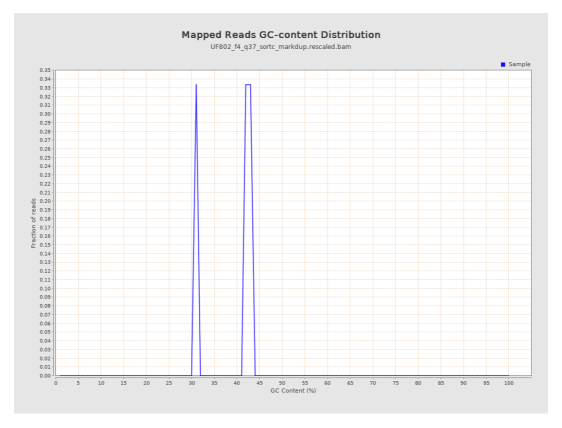


8. Results : Mapped Reads Nucleotide Content



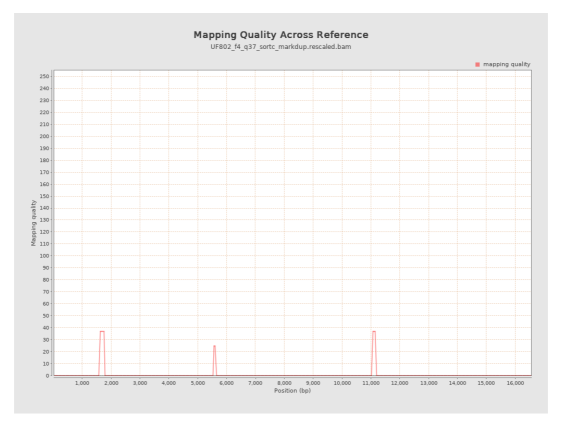


9. Results: Mapped Reads GC-content Distribution





10. Results: Mapping Quality Across Reference





11. Results: Mapping Quality Histogram

