Qualimap Analysis Results

BAM QC analysis Generated by Qualimap v.2.3 2024/10/15 10:11:04



1. Input data & parameters

1.1. QualiMap command line

qualimap bamqc -bam
./Results_from_Pipeline/ggal_gut.Aligned.sortedByCoord.out.bam -nw 400 hm 3

1.2. Alignment

| Command line: | /home/Ashutosh.Yadav/anaconda3/e nvs/starAligner/bin/STAR-avx2 runThreadN 16genomeDir indexreadFilesIn trimmed_ggal_gut_1.fastq trimmed_ggal_gut_2.fastq outFileNamePrefix ggal_gut outSAMtype BAM SortedByCoordinate |
|---------------------------------------|---|
| Draw chromosome limits: | no |
| Analyze overlapping paired-end reads: | no |
| Program: | STAR (2.7.11b) |
| Analysis date: | Tue Oct 15 10:11:03 IST 2024 |
| Size of a homopolymer: | 3 |
| Skip duplicate alignments: | no |
| Number of windows: | 400 |
| BAM file: | ./Results_from_Pipeline/ggal_gut.Alig ned.sortedByCoord.out.bam |



2. Summary

2.1. Globals

| Reference size | 171,001 | |
|------------------------------|-------------------|--|
| Number of reads | 4,667 | |
| Mapped reads | 4,667 / 100% | |
| Unmapped reads | 0 / 0% | |
| Mapped paired reads | 4,667 / 100% | |
| Mapped reads, first in pair | 2,333 / 49.99% | |
| Mapped reads, second in pair | 2,334 / 50.01% | |
| Mapped reads, both in pair | 4,666 / 99.98% | |
| Mapped reads, singletons | 1 / 0.02% | |
| Secondary alignments | 204 | |
| Read min/max/mean length | 37 / 105 / 107.67 | |
| Duplicated reads (estimated) | 1,501 / 32.16% | |
| Duplication rate | 28.08% | |
| Clipped reads | 1,215 / 26.03% | |

2.2. ACGT Content

| Number/percentage of A's | 128,310 / 27.6% |
|--------------------------|------------------|
| Number/percentage of C's | 104,023 / 22.37% |
| Number/percentage of T's | 127,769 / 27.48% |
| Number/percentage of G's | 104,839 / 22.55% |
| Number/percentage of N's | 1 / 0% |
| GC Percentage | 44.92% |



2.3. Coverage

| Mean | 29.0722 |
|--------------------|---------|
| Standard Deviation | 28.589 |

2.4. Mapping Quality

| 21.08 |
|-------|
| 21.00 |
| |

2.5. Insert size

| Mean | 1,638.42 | |
|--------------------|-----------------|--|
| Standard Deviation | 7,504.89 | |
| P25/Median/P75 | 114 / 155 / 945 | |

2.6. Mismatches and indels

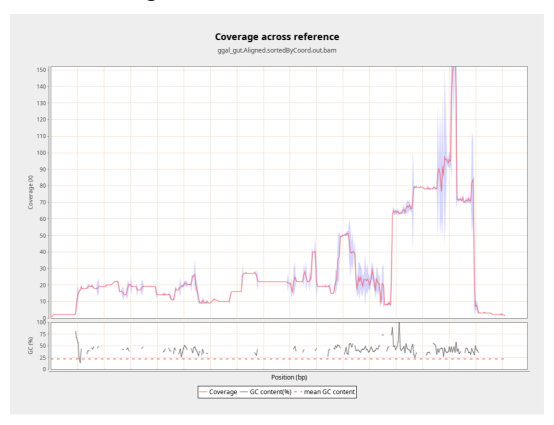
| Insertions | 162 |
|--|-------|
| Mapped reads with at least one insertion | 3.26% |
| Deletions | 38 |
| Mapped reads with at least one deletion | 0.73% |
| Homopolymer indels | 50.5% |

2.7. Chromosome stats

| Name | Length | Mapped bases | Mean coverage | Standard deviation |
|------|--------|--------------|------------------|--------------------|
| 1 | 171001 | 4971369 | 29.0722 | 28.589 |

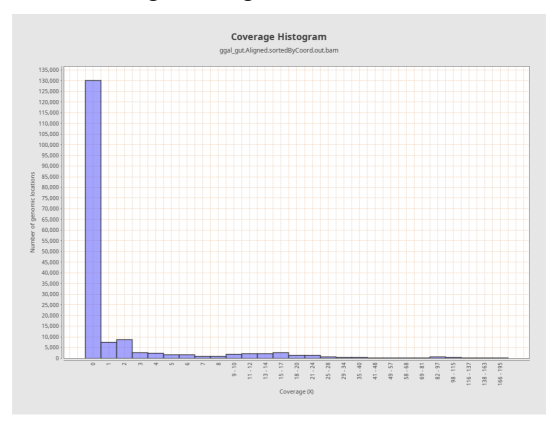


3. Results : Coverage across reference



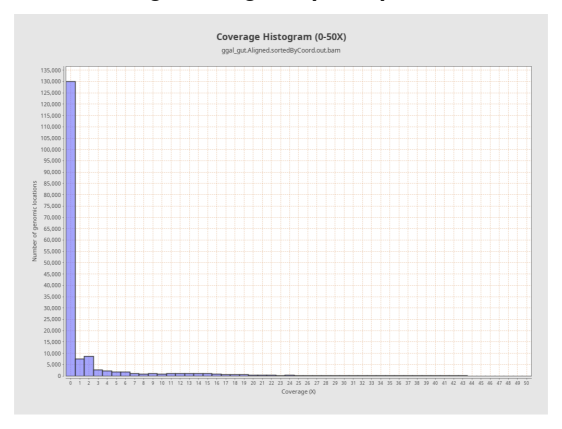


4. Results : Coverage Histogram



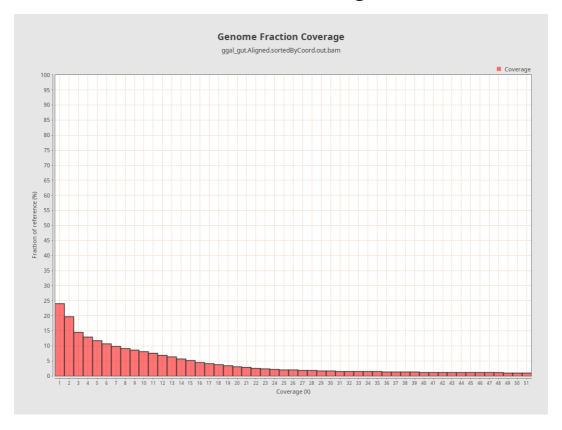


5. Results: Coverage Histogram (0-50X)



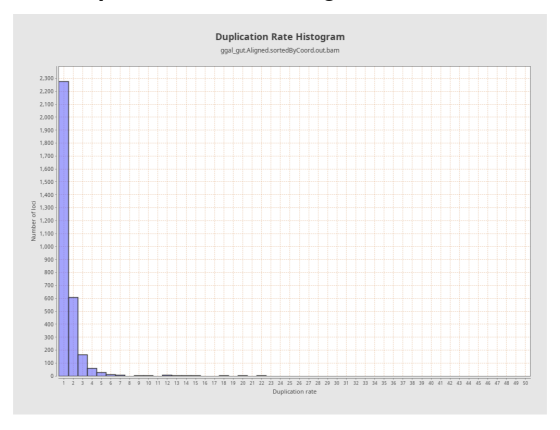


6. Results : Genome Fraction Coverage



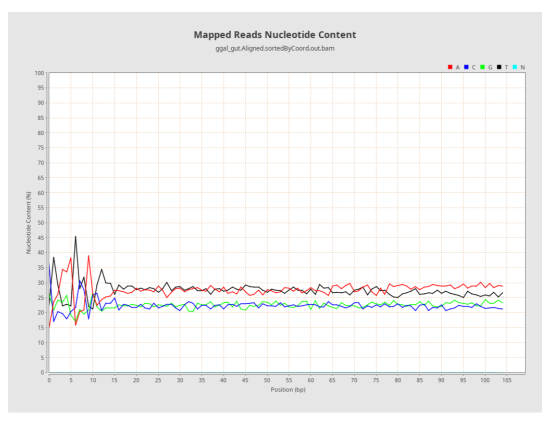


7. Results: Duplication Rate Histogram



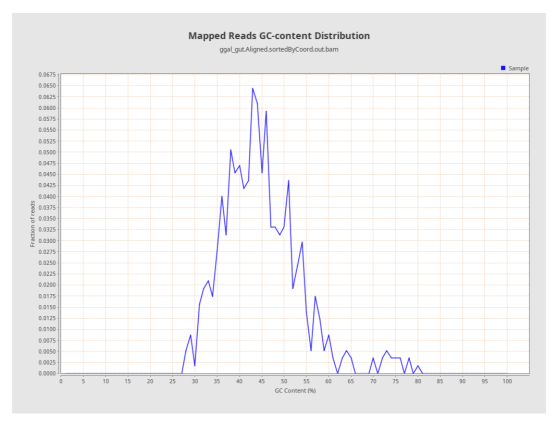


8. Results: Mapped Reads Nucleotide Content



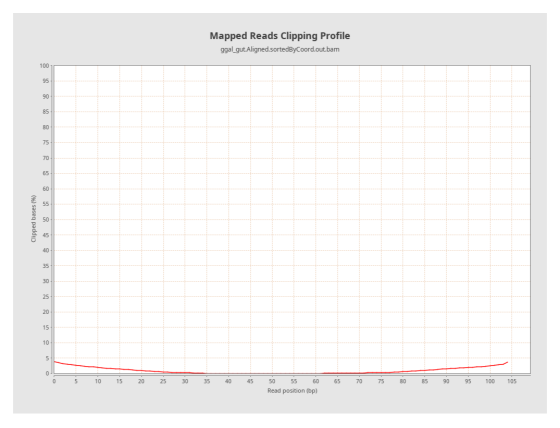


9. Results: Mapped Reads GC-content Distribution



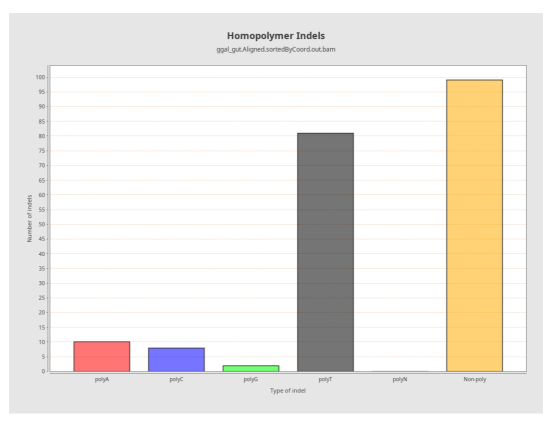


10. Results: Mapped Reads Clipping Profile



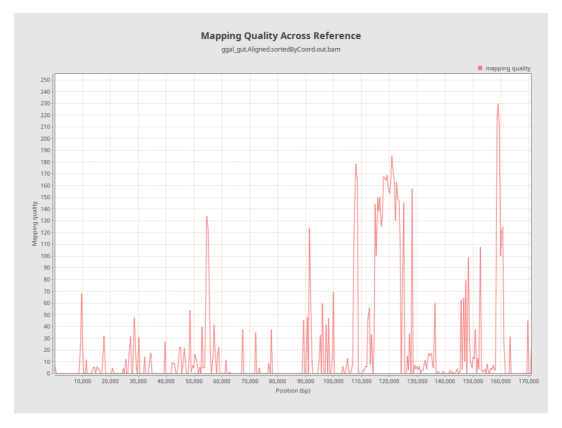


11. Results : Homopolymer Indels



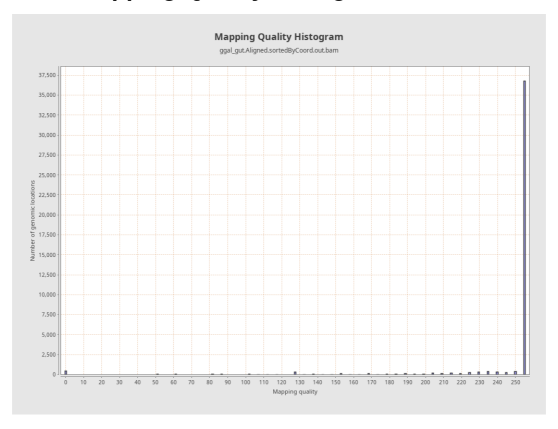


12. Results: Mapping Quality Across Reference



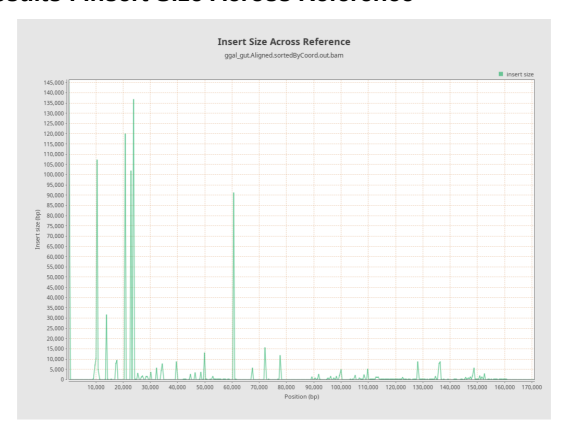


13. Results: Mapping Quality Histogram





14. Results: Insert Size Across Reference





15. Results: Insert Size Histogram

