Qualimap Analysis Results

BAM QC analysis Generated by Qualimap v.2.0 2016/05/19 11:50:16



1. Input data & parameters

1.1. QualiMap command line

qualimap bamqc -bam /export/home/amirh/novoalign_testing/outputBAM/p70sorted.bam -c -nw 400 -hm 3

1.2. Alignment

| BAM file: | /export/home/amirh/novoalign_testing /outputBAM/p70sorted.bam | |
|-------------------------|--|--|
| Program: | novoalign (V3.02.12) | |
| Size of a homopolymer: | 3 | |
| Number of windows: | 400 | |
| Analysis date: | Thu May 19 09:23:20 MYT 2016 | |
| Draw chromosome limits: | yes | |



2. Summary

2.1. Globals

| Reference size | 119,667,750 |
|--------------------------|---------------------|
| Number of reads | 11,947,007 |
| Mapped reads | 11,485,040 / 96.13% |
| Unmapped reads | 461,967 / 3.87% |
| Paired reads | 0 / 0% |
| Read min/max/mean length | 100 / 100 / 100 |
| Clipped reads | 378,500 / 3.17% |
| Duplication rate | 4.91% |

2.2. ACGT Content

| Number/percentage of A's | 366,892,174 / 31.96% | | |
|--------------------------|----------------------|--|--|
| Number/percentage of C's | 207,212,498 / 18.05% | | |
| Number/percentage of T's | 366,541,425 / 31.93% | | |
| Number/percentage of G's | 207,218,815 / 18.05% | | |
| Number/percentage of N's | 0 / 0% | | |
| GC Percentage | 36.1% | | |

2.3. Coverage

| Mean | 9.59 |
|--------------------|------|
| Standard Deviation | 3.57 |

2.4. Mapping Quality

| Mean Mapping Quality | 68.64 | |
|----------------------|-------|--|
|----------------------|-------|--|



2.5. Mismatches and indels

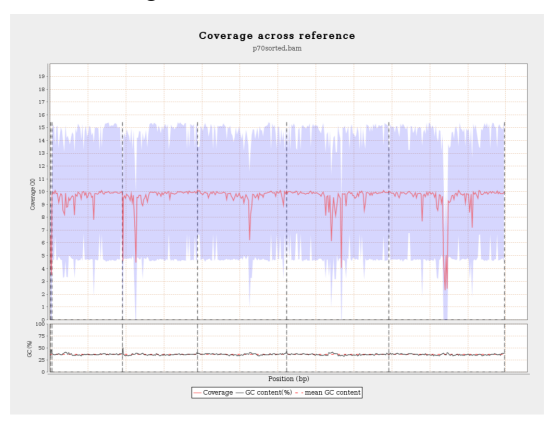
| General error rate | 0.71% |
|--------------------|-----------|
| Mismatches | 8,139,231 |
| Insertions | 357 |
| Deletions | 510 |
| Homopolymer indels | 33.22% |

2.6. Chromosome stats

| Name | Length | Mapped bases | Mean coverage | Standard deviation |
|------|----------|--------------|------------------|--------------------|
| Pt | 154478 | 1012456 | 6.55 | 5.34 |
| Mt | 366924 | 1369351 | 3.73 | 4.9 |
| 4 | 18585056 | 179908154 | 9.68 | 3.46 |
| 2 | 19698289 | 189996349 | 9.65 | 3.51 |
| 3 | 23459830 | 227566481 | 9.7 | 3.44 |
| 5 | 26975502 | 259684668 | 9.63 | 3.54 |
| 1 | 30427671 | 288328419 | 9.48 | 3.71 |

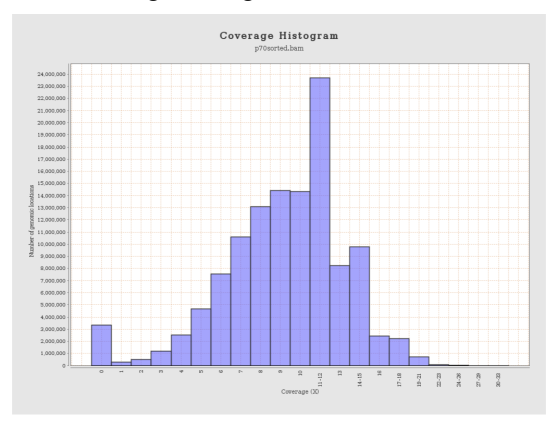


3. Results : Coverage across reference



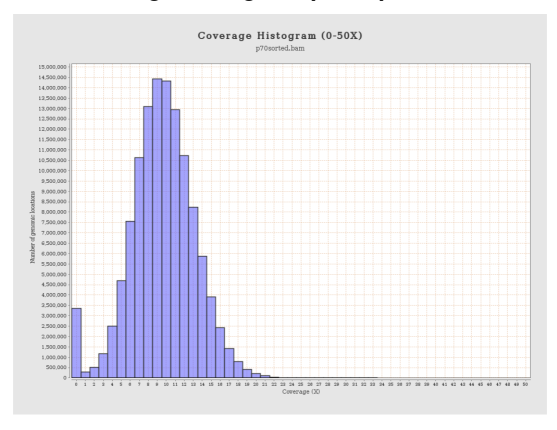


4. Results : Coverage Histogram



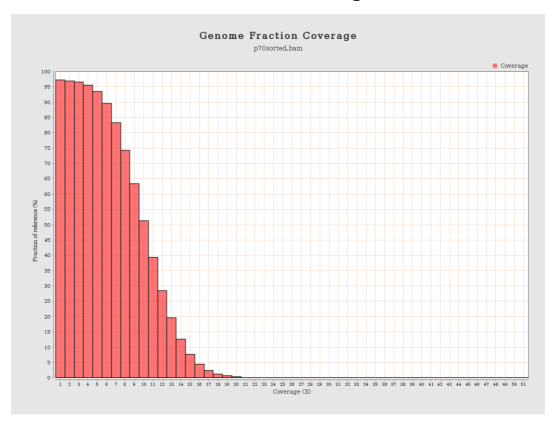


5. Results: Coverage Histogram (0-50X)



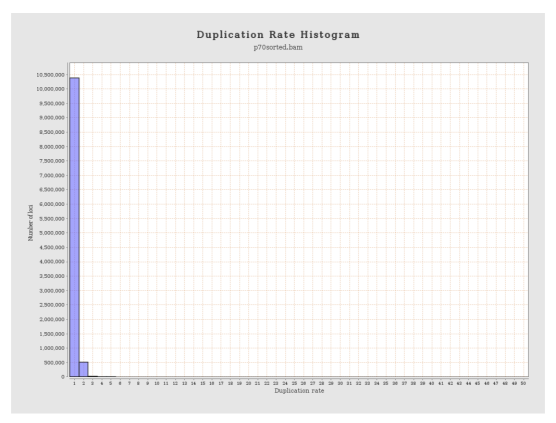


6. Results : Genome Fraction Coverage





7. Results : Duplication Rate Histogram



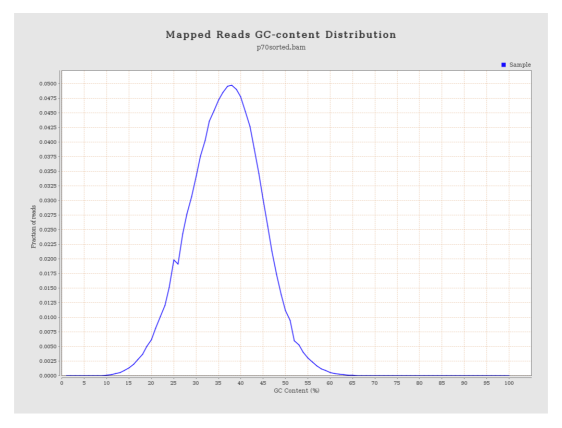


8. Results: Mapped Reads Nucleotide Content





9. Results: Mapped Reads GC-content Distribution



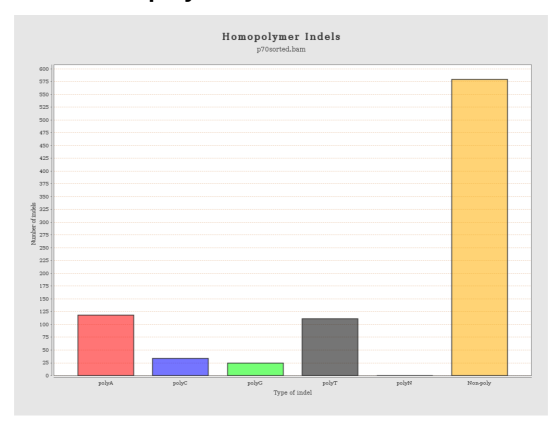


10. Results: Mapped Reads Clipping Profile



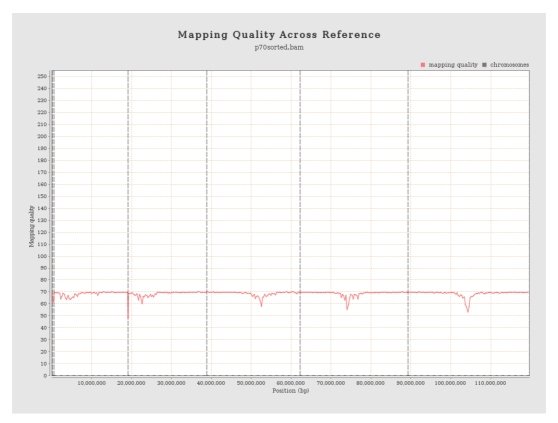


11. Results : Homopolymer Indels





12. Results: Mapping Quality Across Reference





13. Results: Mapping Quality Histogram

