Qualimap Analysis Results

BAM QC analysis Generated by Qualimap v.2.0 2016/05/19 11:52:47



1. Input data & parameters

1.1. QualiMap command line

qualimap bamqc -bam /export/home/amirh/novoalign_testing/outputBAM/p140sorted.bam.bam -c nw 400 -hm 3

1.2. Alignment

BAM file:	/export/home/amirh/novoalign_testing /outputBAM/p140sorted.bam.bam	
Program:	novoalign (V3.02.12)	
Size of a homopolymer:	3	
Number of windows:	400	
Analysis date:	Thu May 19 09:07:57 MYT 2016	
Draw chromosome limits:	yes	



2. Summary

2.1. Globals

Reference size	119,667,750	
Number of reads	11,947,007	
Mapped reads	11,503,465 / 96.29%	
Unmapped reads	443,542 / 3.71%	
Paired reads	0 / 0%	
Read min/max/mean length	100 / 100 / 100	
Clipped reads	383,018 / 3.21%	
Duplication rate	4.92%	

2.2. ACGT Content

Number/percentage of A's	367,472,787 / 31.96%		
Number/percentage of C's	207,547,875 / 18.05%		
Number/percentage of T's	367,120,401 / 31.93%		
Number/percentage of G's	207,555,495 / 18.05%		
Number/percentage of N's	0 / 0%		
GC Percentage	36.11%		

2.3. Coverage

Mean	9.61
Standard Deviation	3.58

2.4. Mapping Quality

Mean Manning Quality	68 64
Mean Mapping Quality	68.64



2.5. Mismatches and indels

General error rate	0.71%	
Mismatches	8,211,198	
Insertions	427	
Deletions	632	
Homopolymer indels	33.33%	

2.6. Chromosome stats

Name	Length	Mapped bases	Mean coverage	Standard deviation
Pt	154478	1014355	6.57	5.35
Mt	366924	1371937	3.74	4.9
4	18585056	180206267	9.7	3.46
2	19698289	190302897	9.66	3.51
3	23459830	227928554	9.72	3.45
5	26975502	260093204	9.64	3.54
1	30427671	288780574	9.49	3.71

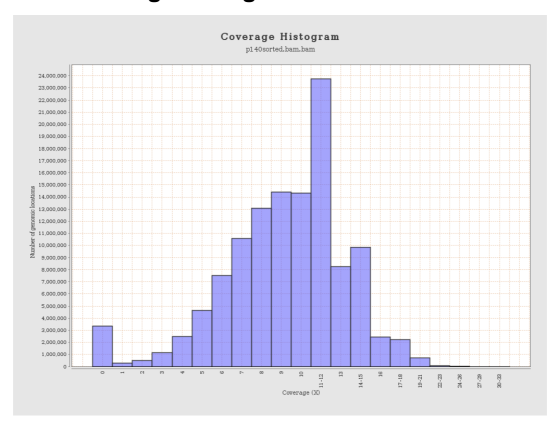


3. Results : Coverage across reference



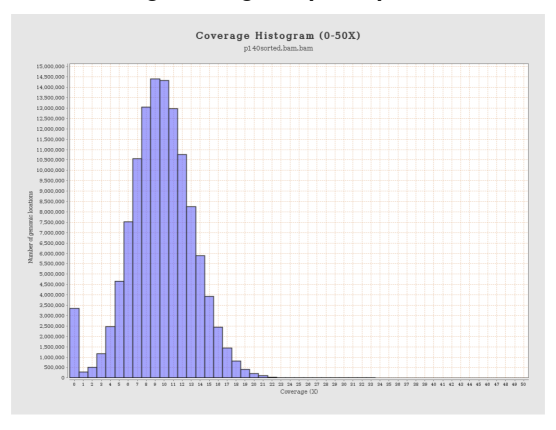


4. Results : Coverage Histogram



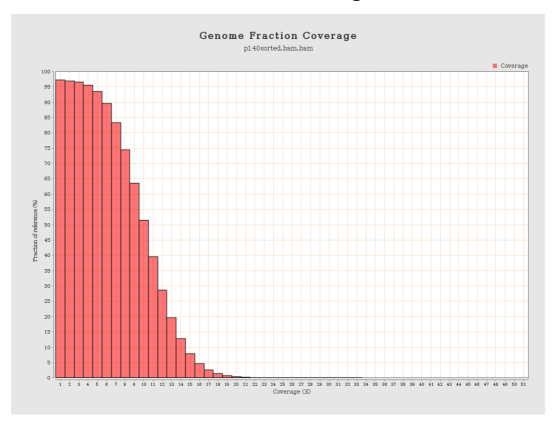


5. Results: Coverage Histogram (0-50X)



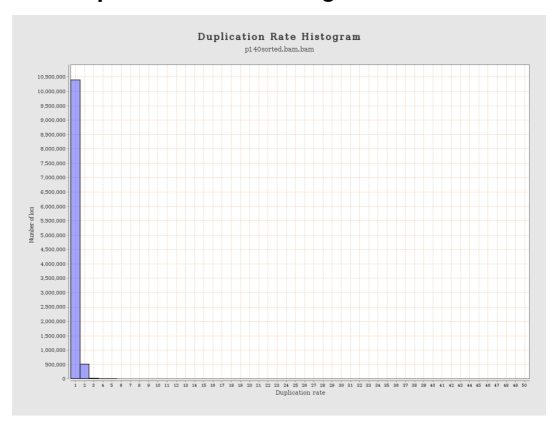


6. Results : Genome Fraction Coverage





7. Results : Duplication Rate Histogram



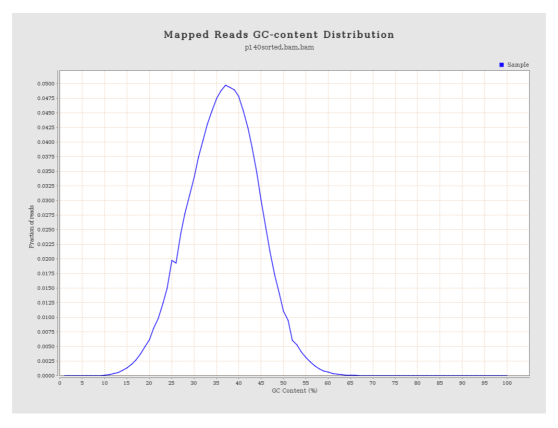


8. Results: Mapped Reads Nucleotide Content



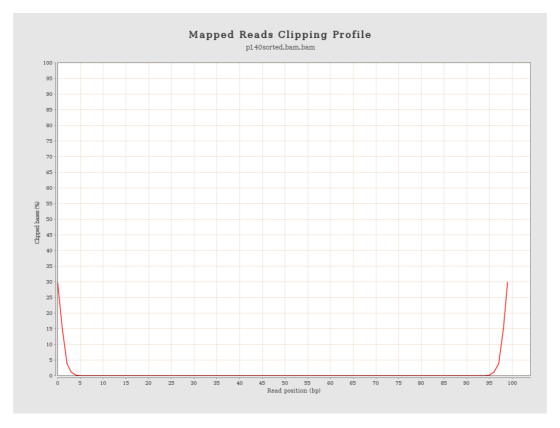


9. Results: Mapped Reads GC-content Distribution



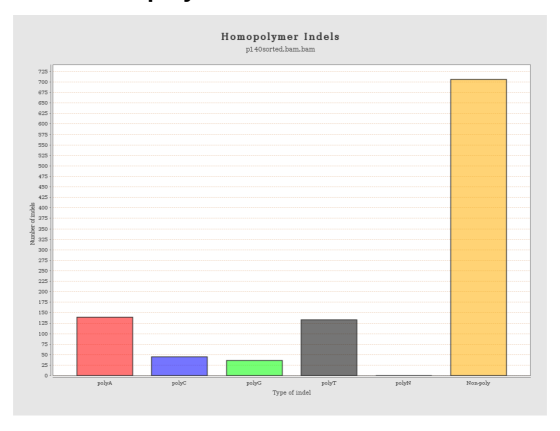


10. Results: Mapped Reads Clipping Profile



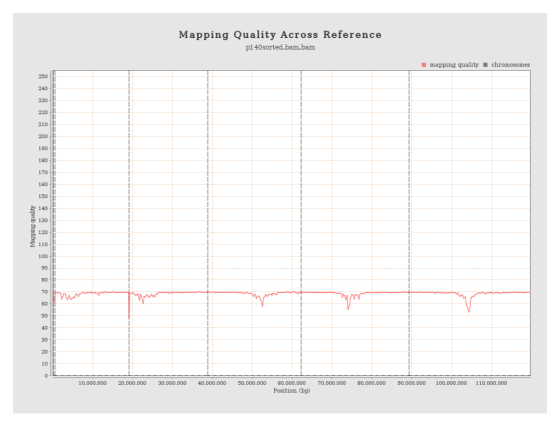


11. Results : Homopolymer Indels





12. Results: Mapping Quality Across Reference





13. Results: Mapping Quality Histogram

