# Qualimap Analysis Results

BAM QC analysis Generated by Qualimap v.2.1.1 2015/09/02 09:37:29



#### 1. Input data & parameters

#### 1.1. QualiMap command line

qualimap bamqc -bam /data/qualimap\_release\_data/alignments/ERR089819.bam -c -nw 400 -hm 3

#### 1.2. Alignment

BAM file:	/data/qualimap_release_data/alignme nts/ERR089819.bam
Program:	Bowtie (0.12.7)
Size of a homopolymer:	3
Number of windows:	400
Skip duplicated alignments:	no
Analysis date:	Wed Sep 02 09:37:28 CEST 2015
Draw chromosome limits:	yes



#### 2. Summary

#### 2.1. Globals

Reference size	100,286,002
Number of reads	35,576,180
Mapped reads	30,983,200 / 87.09%
Unmapped reads	4,592,980 / 12.91%
Paired reads	30,983,200 / 87.09%
Mapped reads, only first in pair	15,491,600 / 43.54%
Mapped reads, only second in pair	15,491,600 / 43.54%
Mapped reads, both in pair	30,983,200 / 87.09%
Mapped reads, singletons	0 / 0%
Read min/max/mean length	100 / 100 / 100
Clipped reads	0 / 0%
Duplicated reads (estimated)	5,125,380 / 14.41%
Duplication rate	16.39%

#### 2.2. ACGT Content

Number/percentage of A's	1,003,585,776 / 32.43%
Number/percentage of C's	541,692,060 / 17.5%
Number/percentage of T's	1,006,351,599 / 32.52%
Number/percentage of G's	543,136,344 / 17.55%
Number/percentage of N's	0 / 0%
GC Percentage	35.05%



#### 2.3. Coverage

Mean	30.89
Standard Deviation	25.01

#### 2.4. Mapping Quality

Mean Mapping Quality	257.71
Hingail Mapping Quality	ZJ4.1
1. 0	

#### 2.5. Insert size

Mean	407.69	
Standard Deviation	92.69	
P25/Median/P75	342 / 400 / 469	

#### 2.6. Mismatches and indels

General error rate	0.22%
Mismatches	6,868,749

#### 2.7. Chromosome stats

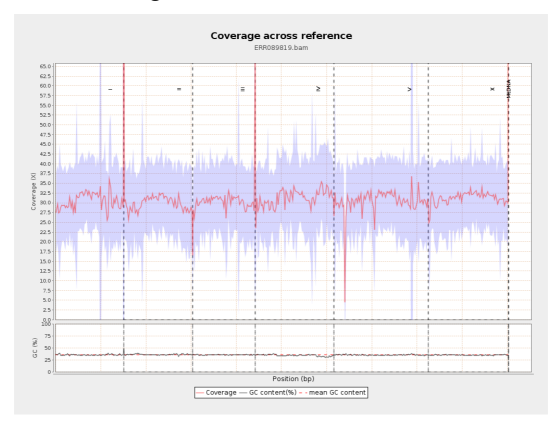
Name	Length	Mapped bases	Mean coverage	Standard deviation
I	15072421	467935600	31.05	36.49
II	15279324	457196800	29.92	9.13
III	13783682	417175000	30.27	9.81
IV	17493784	546120800	31.22	10.46
V	20924143	635772800	30.38	18.85
Х	17718854	554131800	31.27	8.75



MtDNA	13794	19987200	1,448.98	346.72
	13/34	13301200	1,440.90	340.12

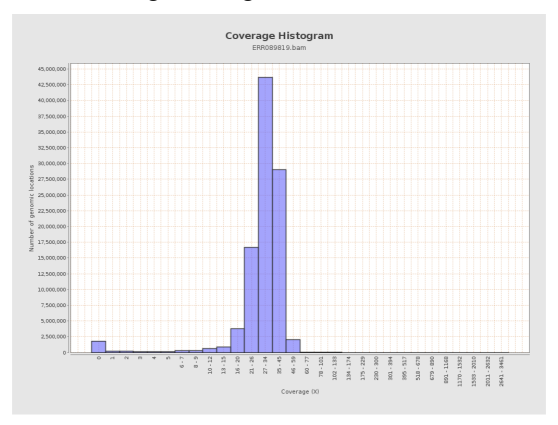


#### 3. Results: Coverage across reference



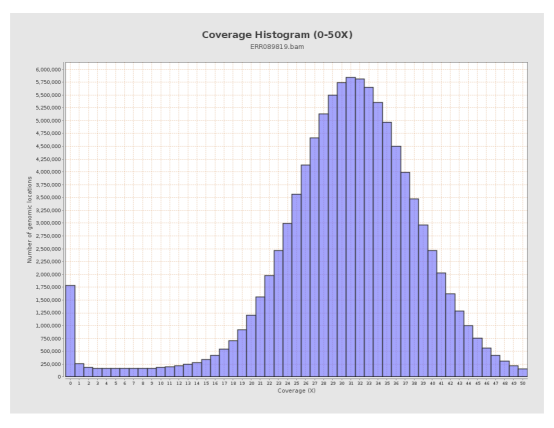


## 4. Results : Coverage Histogram



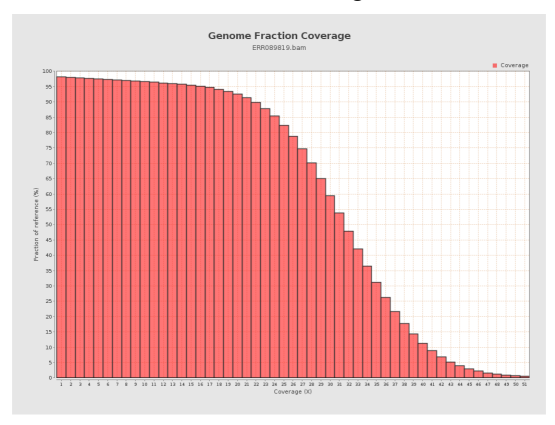


## 5. Results: Coverage Histogram (0-50X)



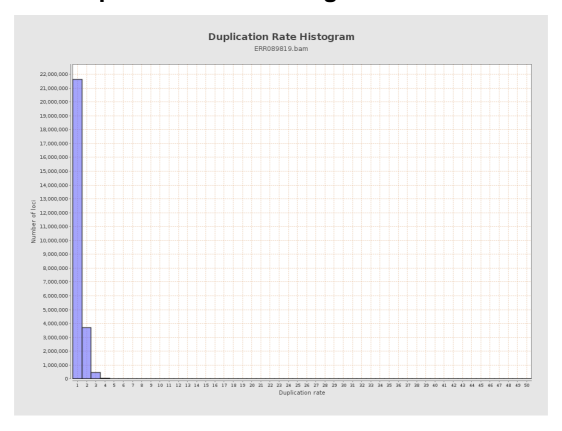


## 6. Results : Genome Fraction Coverage



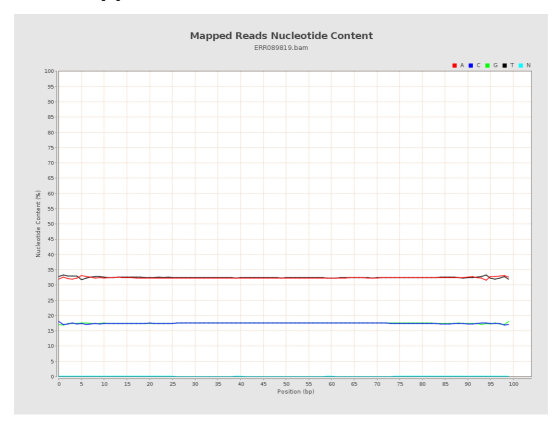


## 7. Results: Duplication Rate Histogram



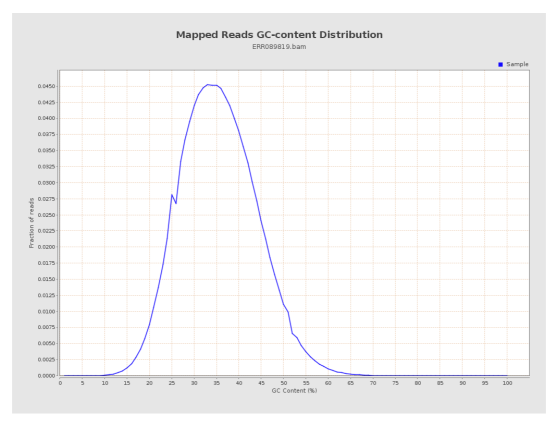


## 8. Results: Mapped Reads Nucleotide Content



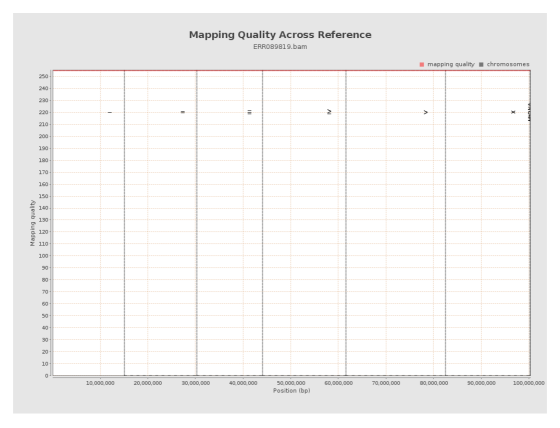


#### 9. Results: Mapped Reads GC-content Distribution



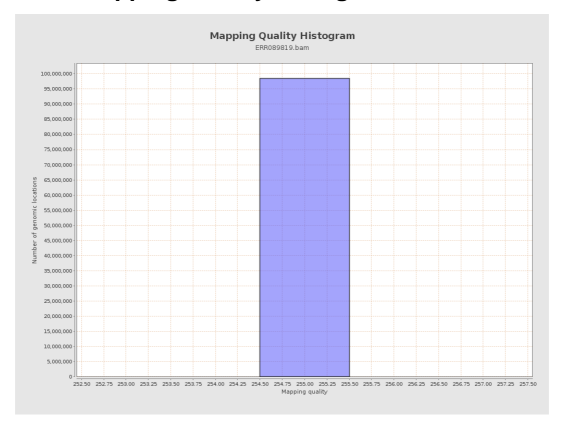


## 10. Results: Mapping Quality Across Reference



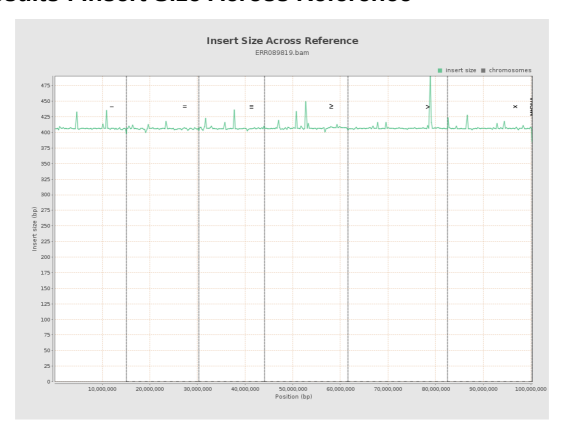


## 11. Results: Mapping Quality Histogram





#### 12. Results: Insert Size Across Reference





## 13. Results : Insert Size Histogram

