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1. Mapping summary report

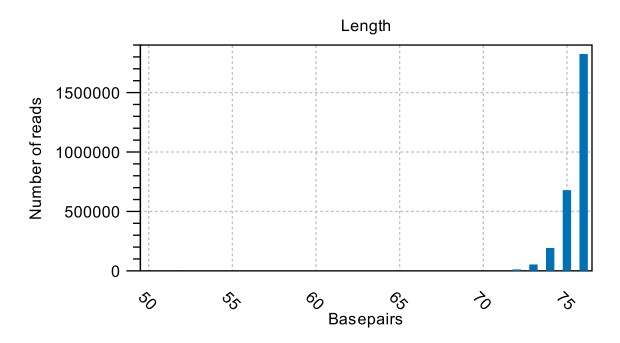
1.1 Summary statistics

Input type: Paired-end reads

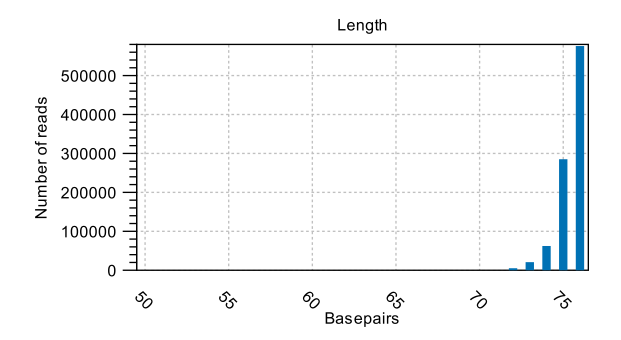
	Count	Percentage of reads (%)	Average length	Number of bases
References	1	-	29,903.00	29,903
Mapped reads	951,644	34.03	75.45	71,804,424
Not mapped reads	1,844,446	65.97	75.35	138,970,180
Reads in pairs	575,180	20.57	214.62	43,394,593
Broken paired reads	376,464	13.46	75.46	28,409,831
Total reads	2,796,090	100.00	75.38	210,774,604

Percentage of bases (%	
	-
	34.07
	65.93
	20.59
	13.48
	100.00

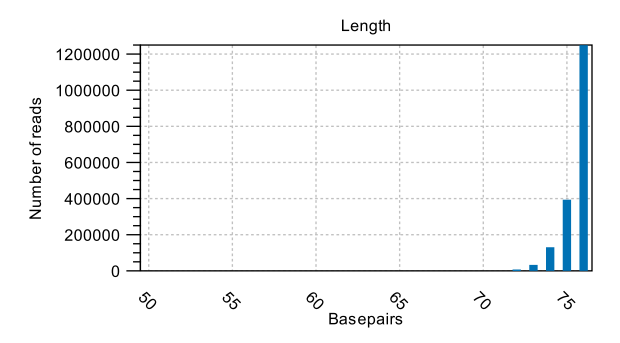
1.2 Distribution of read length



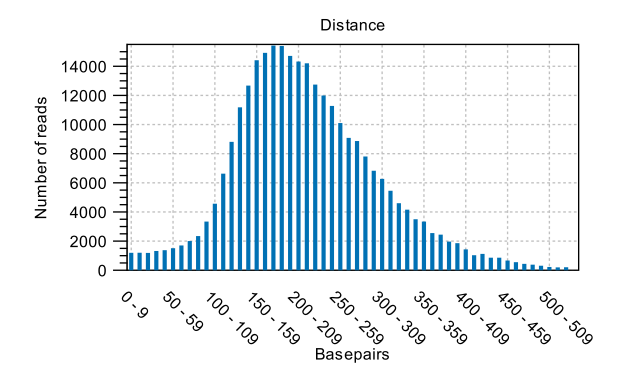
1.3 Distribution of mapped read length



1.4 Distribution of un-mapped read length



1.5 Paired reads distance distribution



History for:



Case B_Mapping_Report

Map Reads to Reference 1.7

15 Aug 2020 10:49:17

CLC Genomics Workbench 20.0.4 Version:

Modified by: sevin

References MN908947.3 (Genome)

Masking mode No masking

Match score Mismatch cost 2

Cost of insertions and deletions Linear gap cost

Insertion cost Deletion cost 3 Length fraction 0.5 Similarity fraction 8.0 Global alignment Nο Auto-detect paired distances Yes

Non-specific match handling Map randomly Output mode Create reads track

Create report Yes Collect unmapped reads No

Comments: Reads mapped: 951,644 of 2,796,090 Estimated paired distance range(s):

B (trimmed pairs): 0 to 530 bp

Originates from:

XX

MN908947.3 (Genome)

Trim Reads 2.4

15 Aug 2020 10:49:02

Version: CLC Genomics Workbench 20.0.4

Modified by: sevin Quality trim Yes Quality limit 0.05 Ambiguous trim Yes Ambiguous limit 2 Automatic read-through adapter Yes

trimmina

Remove 5' terminal nucleotides Nο Remove 3' terminal nucleotides Nο Fixed length trimming No Maximum length 150 Trim from side 3'-end Discard short reads Yes Minimum number of nucleotides in 50

reads

Discard long reads No Save discarded sequences Nο Save broken pairs No Create report

Comments: Processed a total of 2,821,770 sequences. 22,490 sequences have been

completely removed during trimming. 906,683 nucleotides have been trimmed,

altogether.

Originates from:		
	iF	Case B