SeqTrimNext Statistics of pre-processing

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1 Output Files

SeqTrimNext provides several files, the most interesting ones are in the following directories:

output_files

- output.less, containing an extensive information about the trimming of each sequence. It can be visualised on terminal using the command less -R.
- used_params.txt, containing the complete set of parameters used for execution of SeqTrimNext with your data
- rejected.txt, containing a list of rejected sequences together with the reason for their removal.
- initial_stats.json, containing statistics for raw sequences.
- stats.json, containing the statistics of the cleaning process.
- There is a collection of folders that gather sequences with the same MID; each folder contains a sequences file (in FASTQ format) with useful reads. There may also exists a file with reads containing low complexity regions. If you want to reconstruct a SFF with the useful segment of each pre-processed read, use sff_info file in combination with the original SFF file for the sfffile tool.

• graphs

- size_stats.png, a graph with the distribution of read lengths in raw data (see Fig. ??).
- qualities.png, a graph to inspect read qualities in raw data (see Fig. ??).
- PluginExtractInserts_insert_size.png, a graph with the distribution of read lengths after SeqTrimNext pre-processing (see Fig. 1).
- There are other graphs (mostly bar plots) that illustrate the quality of pre-processed reads. All are in PNG format.

• latex

 It is provided as a compressed file latex.zip containing all ".tex" files required to compile this document. Graphs are taken from the graph folder

2 Relevant parameters

In this section, the relevant parameters used in your experiment are shown. Full information about the parameters can be obtained from file used_params.txt





2.1 General

Plugins applied to every sequence, separated by commas. Order is important

- 1. PluginIndeterminations
- 2. PluginFindPolyAt
- 3. PluginAbAdapters
- 4. PluginUserContaminants
- 5. PluginContaminants
- 6. PluginVectors
- 7. PluginLowQuality
- 8. PluginLowComplexity
- $9. \ \ Plugin Extract Inserts$

Remove duplicated (clonal) sequences (using CD-HIT 454)

remove_clonality: false

Minimum insert size for every trimmed sequence

min_insert_size_trimmed: 30

Minimum insert size for each end of paired-end reads; true paired-ends have both single-ends longer than this value

min_insert_size_paired: 40

Seqtrim version

seqtrim_version: 2.0.67

min_sequence_size_raw:

2.2 Quality

Minimum quality value for every nucleotide

min_quality: 20

window_width:

2.3 Contaminants

Blast E-value used as cut-off when searching for contaminations

blast_evalue_contaminants: 1.0e-10

Minimum required identity (%) for a reliable contamination

blast_percent_contaminants: 85

Minimum hit size (nt) for considering a true contamination

min_contam_seq_presence: 40

Genus of input data: contaminations belonging to this genus will be ignored genus:





Is a contamination considered a source of sequence rejection? (setting to false will only trim contaminated sequences instead of rejecting the complete read)

contaminants_reject: true
Path for contaminants database
 contaminants.fasta
 cont_ribosome.fasta

3 Pre-processing statistics

Next figure is equivalent to Figure ?? but using output reads (useful sequences). The mode is expected to decrease but the shape of the plot should be similar.

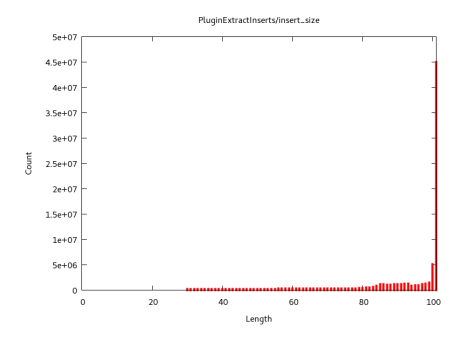


Figure 1: Size distribution of the output sequences. Short sequences (< min_insert_size_trimmed) were removed. [PluginExtractInsert_size.png]

Summary statistics of the SeqTrimNext analysis. Be careful and read all warnings that are indicating concerns about your data. In the files initial_stats.json and stats.json can be found a full statistics of your data and SeqTrimNext pre-processing





Input reads:	total	92960712
	Smallest read (bp)	101
	Largest read (bp)	101
	Mode (bp)	0
	Mean (bp)	0.0
Output results:	total	3509925
	Rejected	9210635
	Low complexity reads	63078
	Mode (bp)	91
	Mean (bp)	91.4
	Output paired reads	80177074
	Total output reads	83686999
Linkers:		

Table 1: List of the most frequent Vectors found among your reads

Vectors	sequences
Cloning vector pAAV-MCS, complete sequence.	461489
Cloning vector pKOHPRT complete sequence.	451615
Enterobacteria phage lambda	150978
Cloning vector pVLH/hsp	51755
Cloning vector pWormgate2, complete sequence.	43820

Table 2: List of the most frequent Adapters found among your reads

Adapters	sequences
ABI_Solid3_Adapter_A	263458
TruSeq_Universal_Adapter	239094
ABI_Solid3_GAPDH_Reverse_Primer	209976
Illumina_Single_End_Adapter_1	190056
ABI_Dynabead_EcoP_Oligo	180734





Table 3: List of the most frequent Contaminants found among your reads

Contaminants	sequences
Podospora anserina S mat+ unordered scaffolds, whole genome	62829
shotgun sequence	
rRNA_small_subunit_Metazoa_Myotis	56101
rRNA_long_subunit_Metazoa_Dasypus	54771
Saccharomyces cerevisiae S288c chromosome XII, complete se-	52719
quence	
Schizosaccharomyces pombe 972h- chromosome III, complete se-	38824
quence	

Table 4: Summary of nucleotides removed in every plugin.

Plugin	Nucleotides	Percent	Warnings
Low Quality	1226704310	Inf %	OK
Low Complexity	72151669	Inf $\%$	OK
Poly T	33060295	Inf $\%$	OK
Poly A	34861798	Inf $\%$	OK
Contaminants	42057730	Inf $\%$	OK
Adapters	84198916	Inf $\%$	OK
Vectors	54594834	Inf $\%$	OK
Indeterminations	218953	Inf $\%$	OK
Inserts	7654874036	Inf %	iW1

iW1 Warning!, only Inf% of nucleotides are useful

4 Rejected reads

Input sequences	92960712
Output sequences	3509925
Rejected sequences	9210635
Output paired sequences	80177074
Total output sequences	83686999
Low complexity sequences	63078





Table 5: Summary of reads removed in every plugin.

Case	Number of sequences	Percent	Warnings
	9210635	9.908 %	OK
Short inserts	6299033	6.776~%	OK
Empty Inserts	2423614	2.607~%	rdW3
Contaminants	456754	0.491~%	OK
No Valid Inserts	22539	0.024~%	OK
Low Complexity	6130	0.007~%	OK
Indeterminations	2435	0.003~%	OK
Unexpected Vector	130	0.000~%	OK
Total rejected	9210635	9.908 %	OK

rdW3 Warning!, a 2.607 % of your sequences are empty (without an insert)

References

- [1] Falgueras et al. SeqTrim: a high-throughput pipeline for preprocessing any type of sequence reads. *BMC Bioinformatics* 11:38 (2010)
- [2] Weizhong Li & Adam Godzik. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. *Bioinformatics* (2006) 22:1658-9

Thanks you for use SeqTrimNext! Send us any comment to scbi support



