## **Raw ChIP-Seq Data Processing**



Download and install the UGENE NGS package to use this pipeline.

Use this workflow sample to process raw ChIP-seq next-generation sequencing (NGS) data from the Illumina platform. The processing includes:

- Filtration:
  - Filtering of the NGS short reads by the CASAVA 1.8 header;
  - Trimming of the short reads by quality;
- Mapping:

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- Mapping of the short reads to the specified reference sequence (the BWA-MEM tool is used in the sample);
- Post-filtration:
   Filtering of the aligned short reads by SAMtools to remove reads with low mapping quality, unpaired/unaligned reads;
  - Removing of duplicated short reads.

The result of the data processing is provided in the BED format. Intermediate data files from the filtration and mapping steps are also available in the output.



If you haven't used the workflow samples in UGENE before, look at the "How to Use Sample Workflows" section of the documentation.

## Workflow Sample Location

The workflow sample "Raw ChIP-Seq processing" can be found in the "NGS" section of the Workflow Designer samples.

## Workflow Image

There are two versions of the workflow available. The workflow for single-end reads looks as follows:



The workflow for paired-end short appearance is the following:



## Workflow Wizard

The workflows have the similar wizards. The wizard for paired-end reads has 5 pages.

1. Input data: On this page you must input FASTQ file(s).

Raw ChIP-Seq Data Processing Wizard		/	8 X
	Input data Sequencing reads FASTQ files FASTQ files with pairs	Required	
Defaults			Next > Cancel

2. <u>Pre-processing:</u> On this page you can modify filtration parameters.

Raw ChIP-Seq Data Processing W	/izard		8
	Pre-processing		
	Reads filtration		
	Base quality	20	÷
	Reads length	1	÷
	Trim both ends	True	•
	3' adapters	r/ugene/data/adapters/adapters.fasta	
	5' adapters		
	5' and 3' adapters		
	Read pairs filtration		
	Base quality	20	÷
	Reads length	0	-
	Trim both ends	False	•
	3' adapters	r/ugene/data/adapters/adapters.fasta	
V	5' adapters		
	5' and 3' adapters		
Defaults		< <u>B</u> ack <u>N</u> ext > Cane	cel

The following parameters are available for reads and reads pairs filtration:

Base quality	Quality threshold for trimming.
Reads length	Too short reads are discarded by the filter.
Trim both ends	Trim the both ends of a read or not. Usually, you need to set True for Sanger sequencing and False for NGS
3'	

adapte rs	A FASTA file with one or multiple sequences of adapter that were ligated to the 3' end. The adapter itself and anything that follows is trimmed. If the adapter sequence ends with the '\$ character, the adapter is anchored to the end of the read and only found if it is a suffix of the read.
5' adapte rs	A FASTA file with one or multiple sequences of adapters that were ligated to the 5' end. If the adapter sequence starts with the charac ter ^, the adapter is 'anchored'. An anchored adapter must appear in its entirety at the 5' end of the read (it is a prefix of the read). A non-anchored adapter may appear partially at the 5' end, or it may occur within the read.
	If it is found within a read, the sequence preceding the adapter is also trimmed. In all cases, the adapter itself is trimmed.
5' and 3' adapte rs	A FASTA file with one or multiple sequences of adapter that were ligated to the 5' end or 3' end.

3. <u>Mapping:</u> On this page you must input reference and optionally modify advanced parameters.

Raw ChIP-Seq Data Processing Wizard		7	? ×
	Mapping		
	BWA-MEM mapping		
	Number of threads	8	••••
	Advanced	~	
	Min seed length	19	-
	Band width	100	<b>.</b>
	Dropoff	100	<b>*</b>
UGENE	Internal seed length	1.50000	*
	Skip seed threshold	10000	
	Drop chain threshold	0.50000	
	Rounds of mate rescues	100	
	Skip mate rescue	0	×
	Skip pairing	False	•
	Mismatch penalty	1	
	Mismatch penalty	4	
	Can anon nonalty		
Defaults		< Back Next >	Cancel

The following parameters are available:

Reference genome	Path to indexed reference genome.
Number of threads	Number of threads (-t).
Min seed length	Path to indexed reference genome (-k).
Band width	Band width for banded alignment (-w).
Dropoff	Off-diagonal X-dropoff (-d).
Internal seed length	Look for internal seeds inside a seed longer than {-k} (-r).
Skip seed threshold	Skip seeds with more than INT occurrences (-c).
Drop chain threshold	Drop chains shorter than FLOAT fraction of the longest overlapping chain (-D).
Rounds of mate rescues	Perform at most INT rounds of mate rescues for each read (-m).
Skip mate rescue	Skip mate rescue (-S).
Skip pairing	Skip pairing; mate rescue performed unless -S also in use (-P).
Mismatch penalty	Score for a sequence match (-A).
Mismatch penalty	Penalty for a mismatch (-B).

Gap open penalty	Gap open penalty (-O).
Gap extention penalty	Gap extension penalty; a gap of size k cost {-O} (-E).
Penalty for clipping	Penalty for clipping (-L).
Penalty unpaired	Penalty for an unpaired read pair (-U).
Score threshold	Minimum score to output (-T).

4. Post-processing: On this page you can modify post-processing parameters.

Raw ChIP-Seq Data Processing Wizard		7	? ×
Raw CruP-seq Data Processing Wizard	Post-process Filtration MAPQ threshold Skip flag Region Remove duplicates For single-end reads	1 The read is unmapped True	(A) (A) (A) (A) (A) (A) (A) (A) (A) (A)
Defaults		< Back Next >	Cancel

The following parameters are available:

MAPQ thresho Id	Minimum MAPQ quality score.
Skip flag	Skip alignment with the selected items. Select the items in the combobox to configure bit flag. Do not select the items to avoid filtration by this parameter.
Region	Regions to filter. For BAM output only. chr2 to output the whole chr2. chr2:1000 to output regions of chr 2 starting from 1000. chr2: 1000-2000 to ouput regions of chr2 between 1000 and 2000 including the end point. To input multiple regions use the space seprator (e.g. chr1 chr2 chr3:1000-2000).
For single- end reads	Remove duplicates for single-end reads.

5. Output data: On this page you must input output parameters.

Raw ChIP-Seq Data Processing Wizard		1	8 ×
	Aligned data		
	Output directory	output	
	Filtered FASTQ Show filtered fastqparame	ters	+
Ugene			
Defaults		< Back Apply Run	Cancel