

Galaxy Workflow ' bowtie2-alignment-pe-dataset-collection'

Step	Annotation
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Step 1: Input dataset collection	
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input <i>select at runtime</i>	
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Step 2: Input dataset	
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input <i>select at runtime</i>	
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Step 3: FASTQ Groomer	
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File to groom Output dataset 'output' from step 1 Input FASTQ quality scores type Sanger & Illumina 1.8+ Advanced Options Hide Advanced Options	
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Step 4: Bowtie2	
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Is this single or paired library Paired-end Dataset Collection FASTQ Paired Dataset Output dataset 'output_file' from step 3 Write unaligned reads (in fastq format) to separate file(s) False Write aligned reads (in fastq format) to separate file(s) False Do you want to set paired-end options? No Will you select a reference genome from your history or use a built-in index? Use a genome from the history and build index Select reference genome Output dataset 'output' from step 2 Set read groups information? Do not set Select analysis mode 1: Default setting only Do you want to use presets? No, just use defaults Save the bowtie2 mapping statistics to the history False	
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Step 5: Flagstat

BAM File to report statistics of

Output dataset 'output' from step 4