

# Galaxy Workflow ' quality-control-se-dataset-collection'

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Step	Annotation
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Step 1: Input dataset collection

**Input Dataset Collection**

*select at runtime*

Step 2: FASTQ Groomer

**File to groom**

Output dataset 'output' from step 1

**Input FASTQ quality scores type**

Sanger & Illumina 1.8+

**Advanced Options**

Hide Advanced Options

Step 3: FastQC

**Short read data from your current history**

Output dataset 'output\_file' from step 2

**Contaminant list**

*select at runtime*

**Submodule and Limit specifying file**

*select at runtime*

Step 4: Trimmomatic

**Single-end or paired-end reads?**

Single-end

**Input FASTQ file**

Output dataset 'output\_file' from step 2

**Perform initial ILLUMINACLIP step?**

False

**Trimmomatic Operations**

**Trimmomatic Operation 1**

**Select Trimmomatic operation to perform**

Sliding window trimming (SLIDINGWINDOW)

**Number of bases to average across**

4

**Average quality required**

**Step 5: FastQC****Short read data from your current history**

Output dataset 'fastq\_out' from step 4

**Contaminant list**

*select at runtime*

**Submodule and Limit specifying file**

*select at runtime*