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

# **Step** Annotation

## 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 specifing 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 specifing file select at runtime