

RNA-seq de novo ABiMS

Cleaning

1. **import des données d'entrée depuis Data Libraries :**
 - Shared Data → Data Libraries → RNA-seq de-novo
2. **lancement des programmes de nettoyage pas à pas**
 - **FastQC:Read QC**
 - **Short read data from your current history**
BlueLight.sample.read1.fastq
 - **Step1 : prinseq_lite**
 - **reads fastq file** BlueLight.sample.read1.fastq
 - **trim_ns_left** 1
 - **trim_ns_right** 1
 - **ns_max_n** 0
 - **trim_qual_right** 20
 - **min_qual_mean** 25
 - **min_len** 50
 - **noniupac** True
 - **FastQC:Read QC**
 - **Short read data from your current history**
BlueLight.sample.read1.fastq_good.fastq
 - **Step2 : Cutadapt**
 - **Fastq file to trim**
BlueLight.sample.read1.fastq_good.fastq
 - **3' Adapters** → **Enter custom 3' adapter sequence**
AGATCGGAAGAGCACACGTCTGAACTCCAG
 - **Minimum length** 50

Cleaning (suite)

- **FastQC:Read QC**
 - **Short read data from your current history**
BlueLight.sample.read1.fastq_good.fastq.cutadapt.fastq
- **Step3 : prinseq_lite**
 - **reads fastq file** BlueLight.sample.read1.fastq_good.fastq.cutadapt.fastq
 - **min_len** 50
 - **trim_tail_left** 5
 - **trim_tail_right** 5
 - **lc_method** entropy
 - **lc_threshold** 70
- **FastQC:Read QC**
 - **Short read data from your current history**
BlueLight.sample.read1.fastq_good.fastq.cutadapt.fastq_good.fastq
- **Step4 : riboPicker**
 - **from**
BlueLight.sample.read1.fastq_good.fastq.cutadapt.fastq_good.fastq
 - **Reference Database**
Non-redundant Ribosomal RNA Database (rrnadb)
- **FastQC:Read QC**
 - **Short read data from your current history**
BlueLight.sample.read1. [...] cutadapt.fastq_good.fastq.nonrrna.fastq

2. création du workflow via l'historique

-  → **Extract Workflow** : RNAseq – cleanning

3. visualisation du workflow

- Workflow → [RNAseq – cleanning] → Edit

Cleaning (suite 2)

4. lancement du workflow sur les 3 jeux restants

- Workflow → [RNAseq – cleanning] → Run
- paramétrages
 - **Step 1: Input dataset**
 - **Condition A - Read 1** BlueLight.sample.read1.fastq
 - or BlueLight.sample.read2.fastq
 - or Dark.sample.read1.fastq
 - or Dark.sample.read2.fastq
 - **Step3: prinseq_lite**
 - **Step5: Cutadapt**
 - **Step7: prinseq_lite**
 - **Step9: riboPicker**

Assemblage

Lancement des outils pas à pas

- **Step 1a: Get pairs**
 - **left reads fastq file** BlueLight.sample.read1.[...] .nonrrna.fastq
 - **right reads fastq file** BlueLight.sample.read2.[...] .nonrrna.fastq
- **Step 1b: Get pairs**
 - **left reads fastq file** Dark.sample.read1.[...] .nonrrna.fastq
 - **right reads fastq file** Dark.sample.read2.[...] .nonrrna.fastq
- **Step 2a: Concatenate datasets**
 - **Concatenate Dataset** BlueLight.sample.read1.[...] .paired.fastq
 - **Add new Dataset → Dataset 1** Dark.sample.read1.[...] .paired.fastq
- **Step 2b: Concatenate datasets**
 - **Concatenate Dataset** BlueLight.sample.read2.[...] .paired.fastq
 - **Add new Dataset → Dataset 1** Dark.sample.read2.[...] .paired.fastq
- **Step 3:**
 - Rename your datasets. Ex:
all.read1.cleaned.paired.fastq
all.read2.cleaned.paired.fastq
- **Step 4: normalize_by_kmer_coverage**
 - **single or paired reads** paired
 - **left reads fastq file** all.read1.cleaned.paired.fastq
 - **right reads fastq file** all.read2.cleaned.paired.fastq
 - **pairs_together** True
 - **max_cov** 30
 - **KMER_SIZE** 25
 - **min_kmer_cov** 1
 - **max_pct_stdev** 100
- **Step 5: Trinity**
 - **Left/Forward strand reads** all.read1.[...] K25_C30_pctSD100.fastq
 - **Right/Reverse strand reads** all.read2.[...] K25_C30_pctSD100.fastq
 - **Strand-specific Library type** None
 - **Group pairs distance** 500
- **Step 6:**
 - Rename your assembly file. Ex:
Trinity_assembly.fasta

- **Step 7: RSEM Align and Estimate**
 - **Trinity assembly** Trinity_assembly.fasta
 - **Left/Forward strand reads** all.read1.cleaned.paired.fastq
 - **Right/Reverse strand reads** all.read2.cleaned.paired.fastq
- **Step 8: Filter fasta by rsem values**
 - **Trinity Fasta File** Trinity_assembly.fasta
 - **RSEM output** RSEM.isoforms.results
 - **FPKM cutoff** 1
 - **Isopct cutoff** 1
- **Step 9:**
 - Rename your filtered assembly file. Ex:
Trinity_assembly.filtered.fasta

Analyse différentielle

Lancement des outils pas à pas

- **Step 1a: RSEM Align and Estimate**
 - **Trinity assembly** Trinity_assembly.filtered.fasta
 - **Left/Forward strand reads** BlueLight.sample.read1. [...] .paired.fastq
 - **Right/Reverse strand reads** BlueLight.sample.read2. [...] .paired.fastq
- **Step 1b: RSEM Align and Estimate**
 - **Trinity assembly** Trinity_assembly.filtered.fasta
 - **Left/Forward strand reads** Dark.sample.read1. [...] .paired.fastq
 - **Right/Reverse strand reads** Dark.sample.read2. [...] .paired.fastq
- **Step 2:**
 - Rename your datasets. Ex:
 - RSEM.BlueLight.isoforms.results
 - RSEM.BlueLight.genes.results
 - RSEM.Dark.isoforms.results
 - RSEM.Dark.genes.results
- **Step 3: Merging Tabular**
 - **With header** True
 - **Data column number** 5
 - **Tabular file** RSEM.BlueLight.isoforms.results
 - **Sample name** BlueLight
 - **Tabular file** RSEM.Dark.isoforms.results
 - **Sample name** Dark
- **Step 4: Trinity run DE analysis**
 - **Merge output file** Tabular merge
 - **Method** edgeR
 - **Replicate** No