



BIO634 - Day 2: RNA sequencing technologies

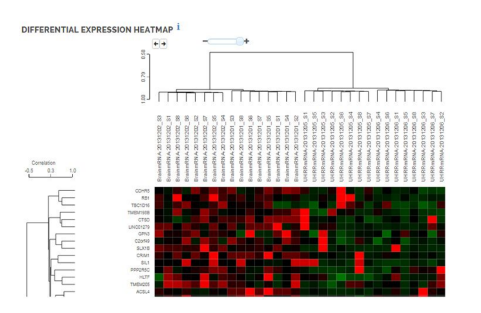
Carla Bello, carla.bello@ieu.uzh.ch
December 3-4th, 2020

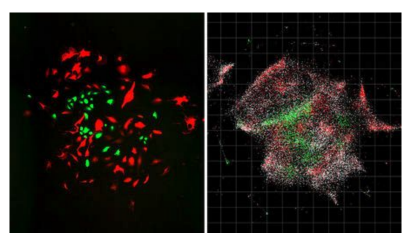
Zürich, CH

Overview

- Introduction to RNA sequencing
- RNA sequencing workflow: steps to analyze the data
- Important considerations: technical, biological replicates, etc.
- Abundance quantification: Gene, exons, transcripts
- **Differential expression analyses:** DEseq, edgeR, etc

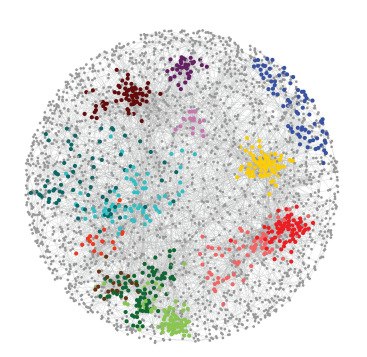
RNAseq reveals the presence and quantity of RNA



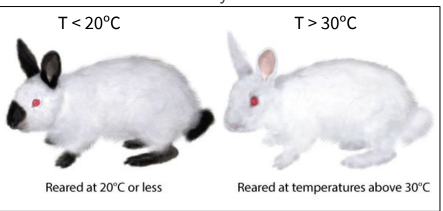


NGS technology that quantifies the transcription / expression of the genome

Gene expression and phenotype



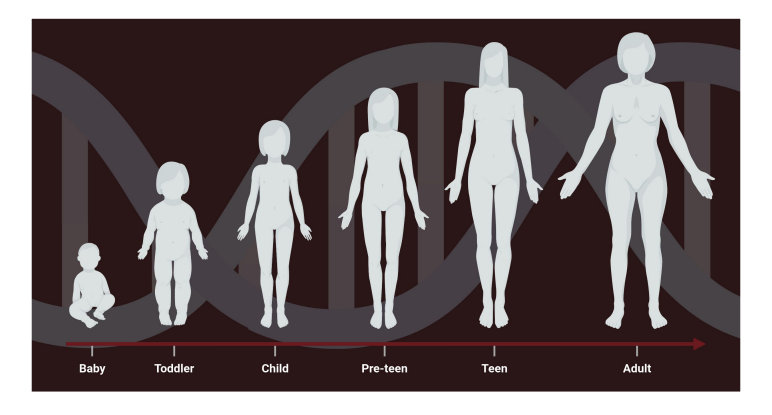
Himalayan rabbit



Example: A pigment gene is influenced by temperature. When the temperature is < 20°C the gene is inactive

Nothing in the genome makes sense except in the light of the transcriptome

RNA sequencing of whole genomes

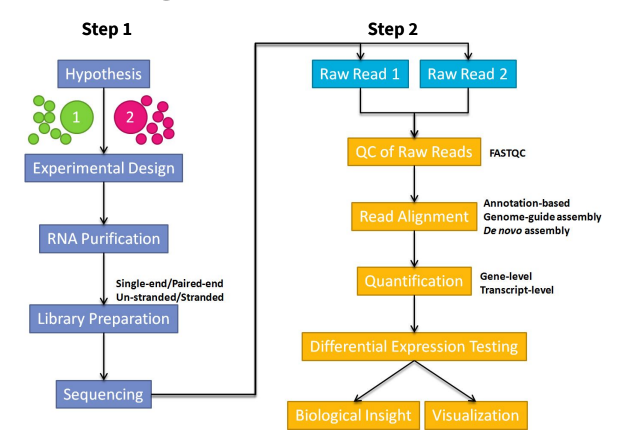


Expression of all the genes in the genome at different stages of aging

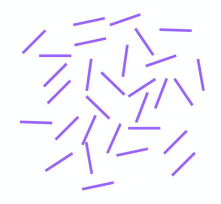
Applications of RNA sequencing

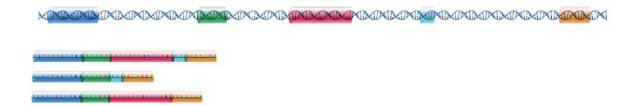
- Gene expression/differential gene expression
- Detecting novel or alternative transcripts
- De-novo transcriptome assembly
- SNP analysis, e.g disease association studies
- Allele-specific expression
- RNA studies: miRNA, tRNAs, snRNA, lncRNAs, etc.

RNA sequencing workflow



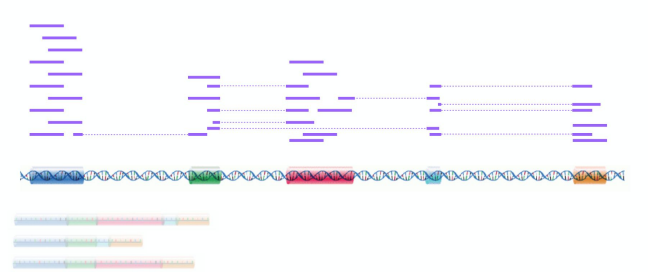
Abundance quantification





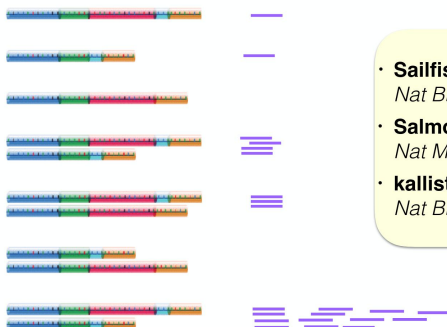
Abundance quantification

Genome alignment of RNA-seq requires a spliceaware aligner (STAR, HISAT2)



Abundance quantification

Transcript-level counts, e.g. obtained by "alignment-free" estimation methods



- Sailfish (Patro et al, Nat Biotechnol 2014)
- Salmon (Patro et al, Nat Methods 2017)
- kallisto (Bray et al, Nat Biotechnol 2016)

Pre-indexed datasets of transcripts

Important considerations before sequencing

- 3 different samples
 3 different conditions
- r_3

3-12 replicates per sample/condition

Technical and biological replicates are important and should be taken into consideration when planning the experiments

Technical replicates

1. <u>Technical replicates:</u> Biological material is the same but the technical steps used to measure gene expression are repeated.

In particular **RNA-seq library preparation** (RNA fragmentation, cDNA synthesis and PCR amplification) **may introduce biases in the data**.

Biological replicates and statistical power

2. <u>Biological replicates</u>: Are different biological samples that are processed separately. They are <u>required if inference on the population is to be made</u>, with <u>three</u> biological replicates <u>being the minimum for any inferential analysis</u>.

3. <u>Desired statistical power</u>, that is the capacity for detecting statistically significant differences in gene expression between experimental groups.

Different methods to analyze RNA-seq data

 There are different packages for differential expression analysis, such as edgeR and DESeq based on negative binomial (NB) distributions or baySeq and EBSeq which are Bayesian approaches based on a negative binomial model.

 These packages work mostly by estimating the variance mean dependance in count data.

Factors to consider for RNAseq analysis

1) Within sample

- Gene/transcript length
- Relative expression (a few highly expressed genes)

2) Between samples

- Sequencing depth (library size)
- Sequencing biases
- 3) Raw read counts are <u>NOT</u> directly comparable between samples: Solution: Normalize read counts

RNA-Seq Read Count Normalization

- RPKM: Reads per kilobase of transcript per million reads of library (single-end reads)
- <u>FPKM</u>: Fragments per kilobase of transcripts per million reads of library (paired-end reads)
- <u>TPM</u>: Transcripts per million reads of library

RNA-Seq Read Count Normalization

RPKM:

Reads Per Kilobase and Million mapped reads

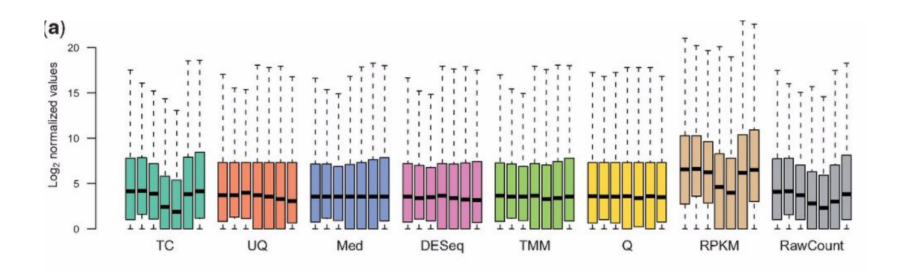
⊌nit of measurement

$$RPKM = \#MappedReads* \frac{1000bases*10^6}{length\ of\ transcript* Total\ number\ of\ mapped\ reads}$$

- RPKM reflects the molar concentration of a transcript in the starting sample by normalizing for
 - RNA length
 - Total read number in the measurement
- This facilitates transparent comparison of transcript levels within and between samples

RNA-Seq Read count normalization

RPKM/FPKM are normalized counts. **DESeq/edgeR** requires <u>raw counts</u> as input as they have their **own normalisation methods**



Differential expression analyses

- Many statistical methods available
 - T-test
 - Poisson Distribution
 - Negative binomial
- No clear consensus yet.
- Tools shown to perform well (under certain circumstances):
 - LIMMA (TMM)
 - DESeq (RLE)
 - edgeR (TMM)
 - Cuffdiff (FPKM)
 - RSEM (EM)
 - Trinity

Identify genes that show differences in expression level between conditions (samples)

Differential expression analyses

- 1. Mapping RNAseq data with Salmon
- 2. Importing transcript abundance with the tximport library
- Differential analysis: A comparison between DEseq and edgeR

Later in the afternoon

- 1. List of differentially expressed genes
- 2. Biological context
- 3. Pathway Analysis (differentially expressed biological pathways)
- 4. Gene Set Enrichment Analysis (GSEA) (functional enrichment between two biological groups)
- 5. Co-expression analysis

Hands-on session - Part II: RNA-seq

Please, go here and follow the instructions:

https://github.com/carlalbc

https://github.com/carlalbc/BIO634_2020/

~~ Lunch break from 12:00 to 13:30 ~~





Enjoy! See you in a bit!:)