Introduction to RNA-Seq: Overview

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General Information

- Course page: https://sib-swiss.github.io/RNAseq-introduction-training/
- Slides, Data sets, Exercises, Solutions
- Optional exam, o.5 ECTS value
- Course from 09:00 to 17:00
- Lunch break 12:00 to 13:00
- 15min breaks around 10:30 and 15:00



Asking questions - Communication

Raise your hand anytime

• Done with an exercise?





Course Outline

Day 1

- 1. **Overview** of RNAseq
- 2. Getting started with the **cluster**
- 3. Quality Control of the raw data
- 4. Sequence trimming

Day 2

- 1. Reas mapping
- 2. Differential Expression Inference
- 3. Enrichment Analysis

slides Outline

- RNA and molecular biology
- Main challenges for RNAseq
- Major Sequencing technologies
- Planning your sequencing : choices, number of samples, ...
- Bioinformatics analysis overview

Introducing Ourselves



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alternative splicing adds a layer of complexity



Image credit: National Human Genome Research Institute - public domain



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What (and why) are we sequencing

Genomics

- Whole genome/exome sequencing (WGS/WES)
- Variant calling (SNPs, CNVs, structural variations)

Epigenomics

- Bisulphite sequencing : DNA methylation
- ATAC-Seq : chromatine opening
- ChIP-seq : TF binding sites

Transcriptomics

- Total RNA
- Poly-A tail selection : focus on mRNA
- Ribo depletion: mRNA + ncRNA
- 5'/3' RACE seq : isoform characterization for one gene
- scRNAseq
- Long read RNA sequencing
- ...

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Long read RNA sequencing

See : https://liorpachter.wordpress.com/seq/

Imagination is the limit

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Main challenges of RNAseq

Transcripts are diverse in size



Main challenges of RNAseq

Transcripts are diverse in size

Expression levels have a *high dynamic range*



From Gtex V8 – human tissue samples Data source : https://gtexportal.org/home/downloads/adult-gtex/bulk_tissue_expression

Main challenges of RNAseq

- Transcripts are diverse in size
- Expression levels have a *high dynamic range*
- RNA molecules are exposed to degradation enzyme:
- RNA integrity affects results
- Is there a reference genome.
- If yes,
- How good is it?
- How good is the gene annotation?

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Main sequencing technologies



lon torrent - reading pH changes



Oxford Nanopore - direct sequencing

DNA can be sequenced by threading it through a microscopic pore in a membrane. Bases are identified by the way they affect ions flowing through the pore from one side of the membrane to the other.



b

100

1D

2D

1D²

2,500

2,000

1,500

0

1,000 length

Yield

MinION/GridION

--- PromethION

Max

reac

(kb) - 500

Average

Maximum

2020

2020

2020

← 23.8 kb

Pacific Biosciences - Single Molecule Real Time



Fig. 1. Principle of single-molecule, real-time DNA sequencing. **(A)** Experimental geometry. A single molecule of DNA template-bound Φ 29 DNA polymerase is immobilized at the bottom of a ZMW, which is illuminated from below by laser light. The ZMW nanostructure provides excitation confinement in the zeptoliter (10^{-21} liter) regime, enabling detection of individual phospholinked nucleotide substrates against the bulk solution background as they are incorporated into the DNA strand by the polymerase. **(B)** Schematic event sequence of the phospholinked dNTP incorporation cycle,

with a corresponding expected time trace of detected fluorescence intensity from the ZMW. (1) A phospholinked nucleotide forms a cognate association with the template in the polymerase active site, (2) causing an elevation of the fluorescence output on the corresponding color channel. (3) Phosphodiester bond formation liberates the dye-linker-pyrophosphate product, which diffuses out of the ZMW, thus ending the fluorescence pulse. (4) The polymerase translocates to the next position, and (5) the next cognate nucleotide binds the active site beginning the subsequent pulse.

From Rhoads & Au. Genomics Proteomics Bioinformatics 2015

Pacific Biosciences - Circular Consensus Sequencing



(>99% accuracy)

Typical in isoseq

Illumina sequencing - cluster formation



Source: Fig1b of Metzker, M. Sequencing technologies — the next generation. Nat Rev Genet 11, 31–46 (2010). https://doi.org/10.1038/nrg2626

Illumina sequencing - sequencing by synthesis



Illumina sequencing - image analysis



Illumina sequencing - from image to sequence



The identity of each base of a cluster is read off from sequential images



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Paired-end sequencing

"Classical" paired end library (illumina)



"Nextera" paired end library



Source : France Genomique

Stranded vs Unstranded Sequencing

- Overlapping genes regions are substantial (~8% in *Homo sapiens*)
- Stranded sequencing allows us to quantify expression in these overlapping regions
- Achieved by ligating different adapters to 5' and 3' ends



M.Griffith et al. PLoS Comp Biol 2015 doi:10.1371/journal.pcbi.1004393

RNA purification

PolyA selection

- Commonly used and inexpensive
- 3' end bias when RNA is degraded
- Loses almost all non-polyA transcripts
- Gets rid of vast majority of ribosomal RNAs, but ncRNA too

Ribosomal RNA depletion

- Less popular, ~2x more expensive
- Higher proportion of rRNA than in polyA selection
- Bacterial data
- Allows identification of IncRNAs without polyA tails
- Retains more immature mRNAs (bad for gene expression quantification)

Sequencing depth

DE : usually aim for ~30-40 million reads

For rare events (isoforms, somatic mutations) much more depth is required

Not easy to know in advance



A User's Guide to the Encyclopedia of DNA Elements (ENCODE) PLoS 2011

Replicates - estimating a biological variance



What does this tell you about the number of replicates needed?

Replicates - estimating a biological variance



What does this tell you about the number of replicates needed ?

2 types of replicates:

- **Technical**: same RNA extract
- **Biological**: same biological condition

Technical replicates



Lauren M McIntyre et al., BMC Genomics 2011

Technical replicates



Article Published: 01 November 2013

Reproducibility of high-throughput mRNA and small RNA sequencing across laboratories

Peter A C 't Hoen ^I, Marc R Friedländer, Jonas Almlöf, Michael Sammeth, Irina Pulyakhina, Seyed Yahya Anvar, Jeroen F J Laros, Henk P J Buermans, Olof Karlberg, Mathias Brännvall, The GEUVADIS Consortium, Johan T den Dunnen, Gert-Jan B van Ommen, Ivo G Gut, Roderic Guigó, Xavier Estivill, Ann-Christine Syvänen, Emmanouil T Dermitzakis & Tuuli Lappalainen ^I

Nature Biotechnology 31, 1015–1022 (2013) Cite this article



Technical replicates



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Transcript level: not reproducible

Biological replicates



TPR at p_adj < 0.05, B₆₂₅



Samples per condition

Soneson, C., Delorenzi, M. A comparison of methods for differential expression analysis of RNA-seq data. *BMC Bioinformatics* **14**, 91 (2013).

More reads or more replicates?



From Liu et al. 2014. RNA-seq differential expression studies: more sequence or more replication?

More reads or more replicates?



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RNA sample preparation - RIN

- Sample quality is critically important: we cannot make up for poor data
- RNA Integrity Number (RIN)
- Minimums:
 - 7-8 : eukaryot mRNA
 - 9 : bacterial



RNA sample preparation - RIN

Effects of preprocessing analysis pipeline



Sigurgeirsson B, Emanuelsson O, Lundeberg J. Sequencing degraded RNA addressed by 3' tag counting. PLoS One. 2014 Mar 14;9(3):e91851.



Schroeder et al BMC Mol Biol 2006

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Basic RNAseq protocol overview











