



# Intro to NGS

# Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics

November 4th, 2019

Selene L. Fernández-Valverde regRNAlab.github.io @SelFdz

# Learning objectives

In this class we will learn

- How high-throughput (NGS) sequencing technologies arose
- How NGS technologies transformed our capacity to acquire large amounts of genomic information '
- Get acquainted with the common NGS techniques available in the market



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### The sequencing revolution



**Figure 1: Sequencing Cost and Data Output Since 2000**—The dramatic rise of data output and concurrent falling cost of sequencing since 2000. The Y-axes on both sides of the graph are logarithmic.









THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



# High-throughput sequencing techniques

- **Pyrosequencing** ٠
- Sequencing by synthesis
- Sequencing by ligation
- Ion semiconductor •
- Nanopore sequencing ٠
- **Single Molecule Real Time** • Sequencing (SMRT)





THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



## Pyrosequencing - 1





5



#### Pyrosequencing - 2





THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



# Pyrosequencing

#### **Advantages**

- Reasonable cost
- Long sequences (500 nts)

#### Disadvantages

- Few sequences produced
- High number of errors in regions with the same nucleotide (homopolymers)
- With the rise of other • technologies and given its high level of errors it was ultimately discontinued



7





The process starts by joining ulletadapters to the DNA or RNA fragments that we want to sequence.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





- The templates are immobilized on a flow cell
- In the case of RNA-Seq, complementarity with the adapter is used to synthesize a new cDNA chain in order to preserve information about the directionality of the transcript.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





 A chain of DNA complementary to the DNA template is synthesized on the flow cell surface.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





 A chain of DNA complementary to the DNA template is synthesized on the flow cell surface.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





• The templates are separated using high temperature.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





 This process is repeated hundreds of times until generating a "colony" or cluster of identical transcripts.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde







 Primers and fluorescent nucleotides (reversible terminators) are added in order (first A, then T, etc.) along with polymerase. When a nucleotide is incorporated a laser pulse coupled with imaging are used to identify which base was incorporated in each position.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde







This process is continued for ulletall bases.



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde









→ GCTGA...

Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



# Sequencing by Synthesis

#### **Advantages**

- Undoubtedly the leader in the market = strong scientific support network
- Produces large amounts of sequences (Up to 20 billion for NovaSeq)
- Low error rate compared with other technologies

#### Disadvantages

- The sequences are short (150 to 300 bp)
- The cost is high
- Relatively slow sequencing (13–44 hr for NovaSeq)



Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### Nanopore 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.





THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde





#### Nanopore sequencing



Kate Rubins









THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### Nanopore whale watching



"Both jaws, like enormous shears, bit the craft completely in twain."  $-Page_{fic}$ .

- Nanopore is capable of generating very very long reads or "whales"
- The longest read detected to date has a length of 2,272,580 bases





## Nanopore sequencing

#### **Advantages**

- Real-time sequencing
- You can stop sequencing when you have enough data
- Very portable useful for work in difficult areas
- Simple preparation
- Low cost \$ 80 USD per sample

#### Disadvantages

- High number of errors although they have had a drastic increase in accuracy in the last year
- Pores failed sequence loss



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### Sources of error

- There are two main sources of error:
  - Human error: mixing of samples (in the laboratory or when the files were received), errors in the protocol
  - Technical error: Errors inherent to the platform (e.g., mononucleotide sequences in pyrosequencing) All platforms have some level of error that must be taken into account when designing the experiment.



**HE ROYAL** Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### Errors in sample preparation

- User error (e.g. mistakenly labeling a sample)
- DNA / RNA degradation by preservation methods
- Contamination with external sequences
- Low amount of DNA start



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



## Errors in library preparation

- User error (e.g. polluting one sample with another, contaminate with previous reactions, errors in the protocol)
- PCR amplification errors
- Bias for primers (binding bias, methylation bias, primer dimers [first dimers])
- Bias for capture (Poly-A, Ribozero)
- Machine errors (misconfiguration, reaction interruption)
- Chimeras
- Index errors, adapter (contamination of adapters, lack of index diversity, incompatible codes (barcodes), overload)

THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics SOCIETY Selene L. Fernandez-Valverde



# Sequencing and image errors

- User error (e.g. cell overload) •
- Delay (e.g., incomplete extension, addition of multiple nucleotides)
- Dead fluorophores, damaged nucleotides and overlapping signals
- Context of the sequence (e.g. high GC content, homologous) and low complexity sequences, homopolymers).
- Machine errors (e.g. laser, hard disk, programs)
- Chain biases

SOCIETY



The challenge - differentiate biological signals from noise/errors

- Negative and positive controls What do I expect?
- Technical and biological replicas help determine the noise rate
- Know the types of common errors in a certain platform



THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde



#### Now what?





THE ROYAL Theoretical and Practical HiC Workshop: Wet-lab and Bioinformatics Selene L. Fernandez-Valverde









#### Practical - Fastq format and QC of NGS data

https://liz-fernandez.github.io/HiC-Workshop/01-quality.html



