

Experimental Design

4 Main Stages

1

Biological
Question

2

Design &
Conduct
Experiment

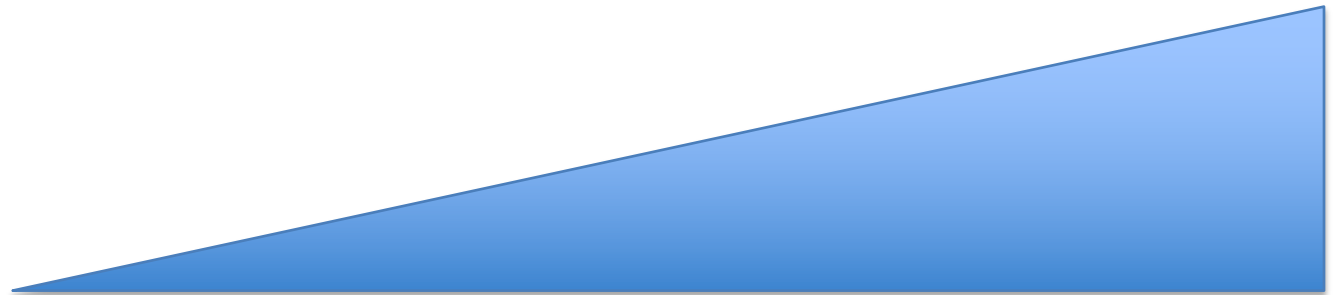
3

Prepare NGS
Library &
Sequence

4

Sequencing
Analysis

Class time



2

Design &
Conduct
Experiment

2. DESIGN & CONDUCT EXPERIMENT

Types of sequencing

- Clonal sequencing
 - Expect a single genome to be present. Highly resilient to sequencing errors.
- Population sequencing
 - Multiple subpopulations and genomes expected to be present. Highly susceptible to sequencing errors especially when making quantitative analysis.
- Amplicon/targeted sequencing
 - Know all/most reads correspond to specific genomic locations. Can be dominated by sequencing errors.

Nice...

We have so many options!

- Deep whole genome
- Low pass whole genome
- Deep whole exome
- Genomewide array
- Exome array

How would you like to be sequenced?

Genomics
Core
Facility

3

Prepare NGS
Library &
Sequence

3. PREPARE NGS LIBRARY & SEQUENCE

Standard Library Prep

1. Fragment DNA
 1. Enzymatic, sonication, acoustic, nebulization
2. Blunt DNA
3. “A”-Tail DNA
4. Ligate adapters
5. PCR

Standard library prep sufficient

- Clonal samples
 - Each base 0 or 100%
 - 50% possible in diploids
- Low to moderate coverage depth populations (<100)
 - Standard Illumina error rate 1%
 - Much more on this later
- Good reference
 - Typical want DNA fragment sizes 350 – 700 bp
 - miSeq: 700 – 1500 bp

Standard library prep lacking

- High coverage populations (>100)
 - Error rate of 1% sets limit of detection at 1% regardless of depth
 - Error rate reduction
- Non-model organism
 - Difficult to generate good reference assemblies using 300 - 700bp
 - Mate-pair libraries several kb inserts
 - Combine with other long read sequencing solutions
- Repeat-mediated rearrangements
 - Repeats often 1.5kb+ long, difficult to get reads on both sides
 - Mate-pair libraries again

3. Prepare NGS Library & Sequence

ERROR RATE REDUCTION

Basic Principle

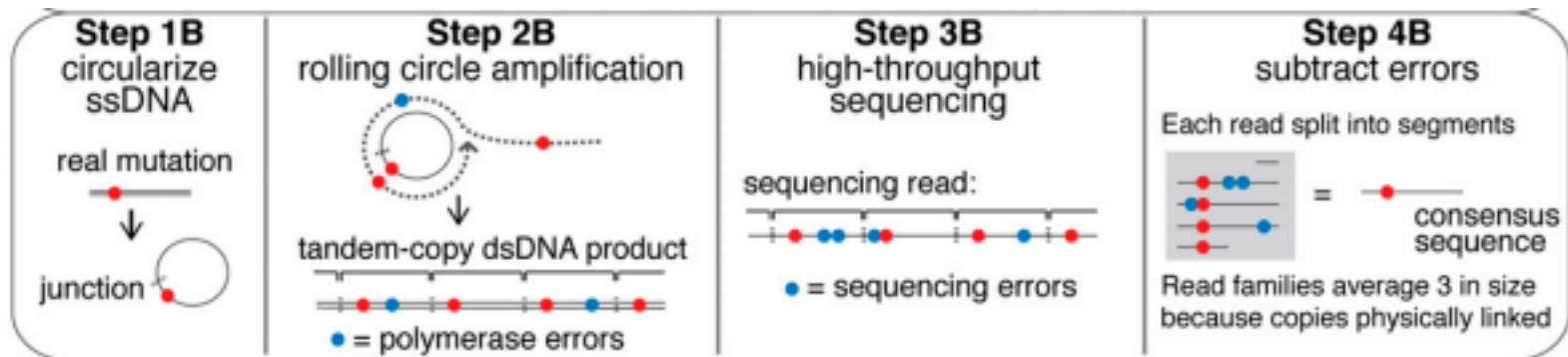
- Because majority of errors are randomly distributed along reads, several alternative library preparation methods exist to read the same original fragment of DNA multiple times to reduce error rates.
- 3 main ways.
 1. Molecular indexing
 2. Circle sequencing
 3. Short insert size approximately read length

Alternative Library Preparation

- Duplex sequencing ([Schmidt et al 2012 PNAS](#))
 - Molecular index to identify original DNA fragments

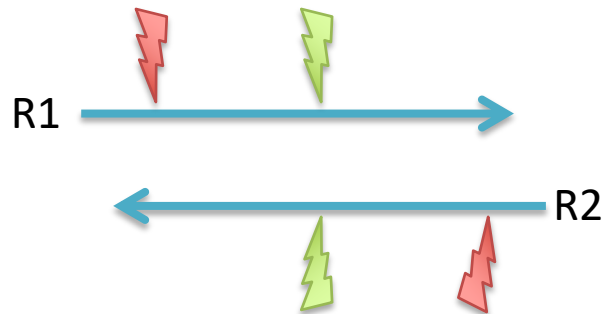


- Circle sequencing ([Lou et al 2013 PNAS](#))
 - Rolling circle amplification to reread same fragment



Alternative Library Preparation

- Double read
 - Fragment DNA size to \sim read length.
 - Can be done with any paired end read.



3. Prepare NGS Library & Sequence

“LONGER” READS

Illumina Options

- Technology:
 - Illumina: 2 x 300 miSeq runs
- Mate pair library:
 - Generates 2 outwardly facing reads separated by up to 25kb.
 - Allows connection of otherwise distant locations on a single read.



4

Sequencing
Analysis

Read Sequence Quality control

4. SEQUENCING ANALYSIS



Garbage In, Garbage Out



Data Preprocessing

- Massaging standard Illumina data is always less effective than better experimental planning and design using alternative library preparation methods.

Read Sequence Quality Control Questions

- Contaminated with other samples?
- Adapter dimers present?
 - Reads with no insert present.
- Adapters present on ends of reads?
 - Insert size smaller than read length.
- 3' end of reads quality decline?

About the Tutorials.

- Some more basic bash/linux interrogation about reads, and working with TACC.
- FastQC can answer all questions about raw read quality and is pretty much the only game in town.
- Cutadapt is a tool for quickly improving data when you identify a problem.

FASTQC WALK THROUGH