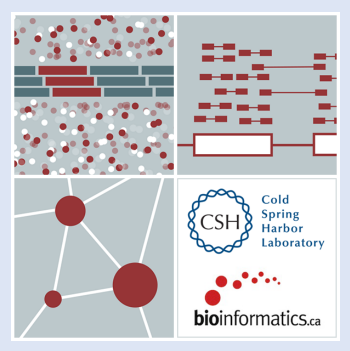
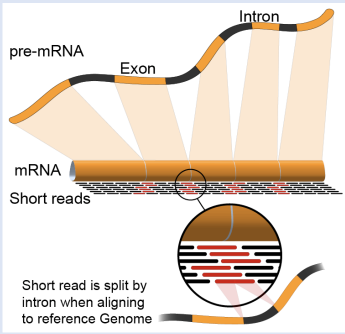


#### RNA-Seq Module 1 Indexing

Kelsy Cotto, Felicia Gomez, Obi Griffith, Malachi Griffith, Allegra Petti, Megan Richters, Huiming Xia Advanced Sequencing Technologies & Bioinformatics Analysis November 16-20, 2020

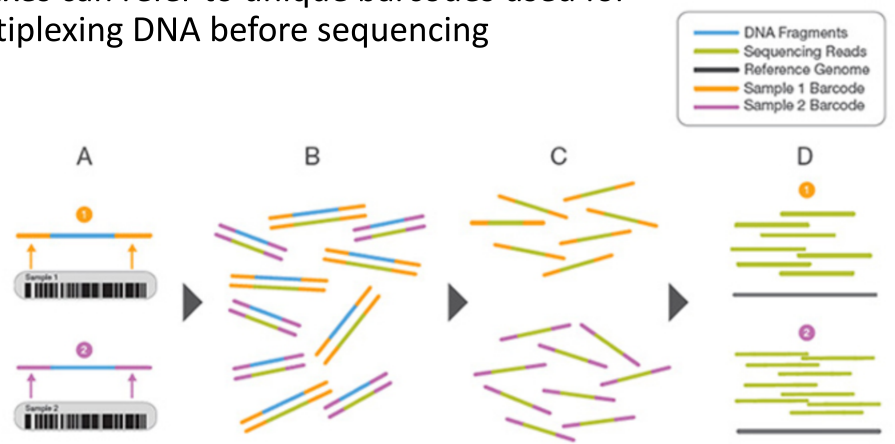






### "Index" has many different meanings

• Indexes can refer to unique barcodes used for multiplexing DNA before sequencing



https://www.illumina.com/science/technology/next-generation-sequencing/multiplex-sequencing.html

### Indexing in bioinformatics/CS enables rapid access

- Indexing is a recurring theme in genome analysis
- Files are \*big\* scanning through them can take a long time
- Indexing builds a table-of-contents so that we can jump directly to specific positions

- Indexing may require significant compute/time but typically only occurs once
- Each application may require a different indexing strategy

## What's inside a fasta's index file? (.fai)

|             | bases in contig |           | byte index of the file where the |             | bases per line |    |                |
|-------------|-----------------|-----------|----------------------------------|-------------|----------------|----|----------------|
| contig name |                 |           | con                              | itig begins |                | /  | bytes per line |
|             |                 |           |                                  |             |                |    |                |
|             | chr1            | 248956422 | 2                                | 6           | 60             | 61 |                |
|             | chr2            | 242193529 | 9 :                              | 253105708   | 60             | 61 |                |
|             | chr3            | 19829555  | 9 (                              | 499335802   | 60             | 61 |                |
|             | chr4            | 19021455  | 5 '                              | 700936293   | 60             | 61 |                |
|             | chr5            | 181538259 | 9                                | 894321097   | 60             | 61 |                |
|             | chr6            | 170805979 | 9                                | 1078885000  | 60             | 61 |                |
|             | chr7            | 159345973 | 3                                | 1252537752  | 60             | 61 |                |
|             | chr8            | 14513863  | 6                                | 1414539498  | 60             | 61 |                |
|             | chr9            | 13839471  | 7                                | 1562097118  | 60             | 61 |                |
|             | chr10           | 133797422 | 2                                | 1702798421  | 60             | 61 |                |

# Example index applications and associated files

| Source file | Indexed file | Indexing tool  | Use case                                   |
|-------------|--------------|----------------|--|
| .bam        | .bai         | samtools index | Visualize bam in IGV                       |
| .fasta      | .fai         | faidx          | Extract specific sequences from ref genome |
| .vcf        | vcf.gz.tbi   | bgzip/tabix    | Pull out specific variants                 |
| .bed        | .bed.gz.tbi  | bgzip/tabix    | extract specific genomic regions           |

#### Indexing is also essential for alignment

• Finding out where to place a read in the genome is impractical unless matches can be quickly found

All read aligners use some kind of indexing

 These indices must be "built" once for a reference genome, but can then be used every time the aligner is run

 Different aligners use different indexing schemes that are not compatible

# We are on a Coffee Break & Networking Session