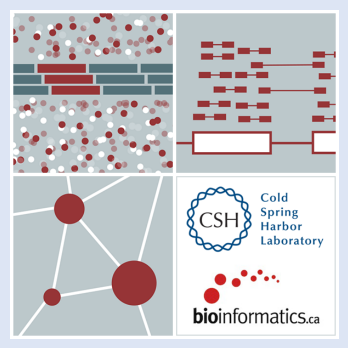
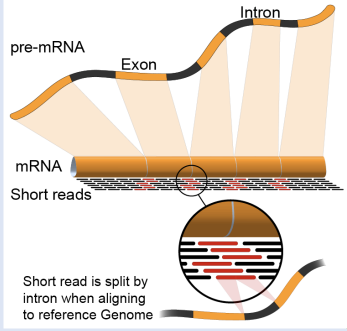


RNA-Seq Module 1 Indexing

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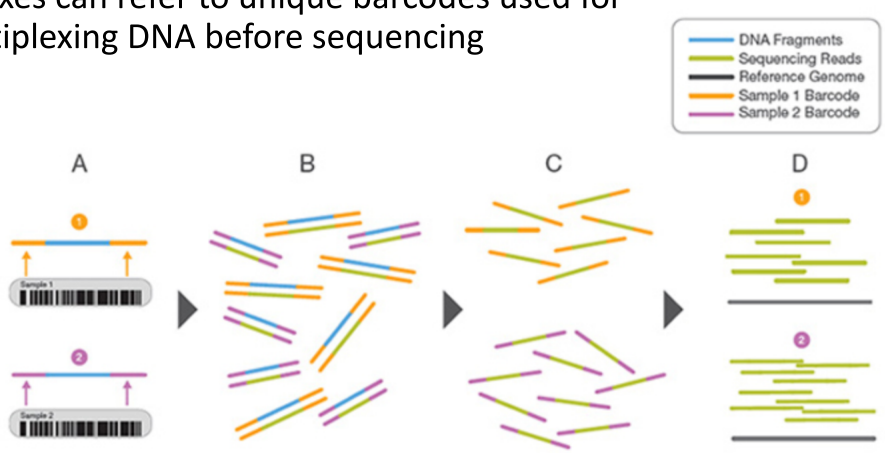






"Index" has many different meanings

• 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)

contig name	bases i	n contig	file v	index of the where the ig begins 	t t	oases	s per line	bytes per line
	chr1	248956422	2 6			60	61	
	chr2	24219352	9 2	53105708		60	61	
	chr3	19829555	9 4	99335802		60	61	
	chr4	19021455	5 7	00936293		60	61	
	chr5	181538259	9 8	94321097		60	61	
	chr6	17080597	9 1	078885000)	60	61	
	chr7	15934597	3 1	252537752	2	60	61	
	chr8	14513863	6 1	414539498	3	60	61	
	chr9	13839471	7 1	562097118	3	60	61	
	chr10	133797422	2 1	702798421	l	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