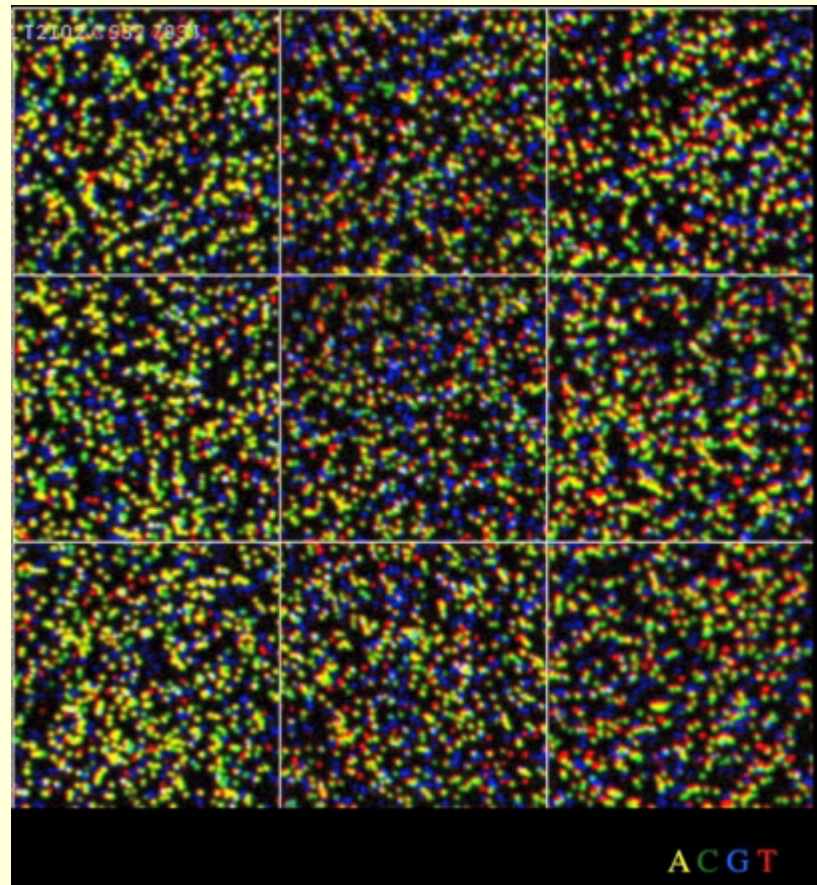


PLNT2530

2025

Unit 6e

# DNA Sequencing



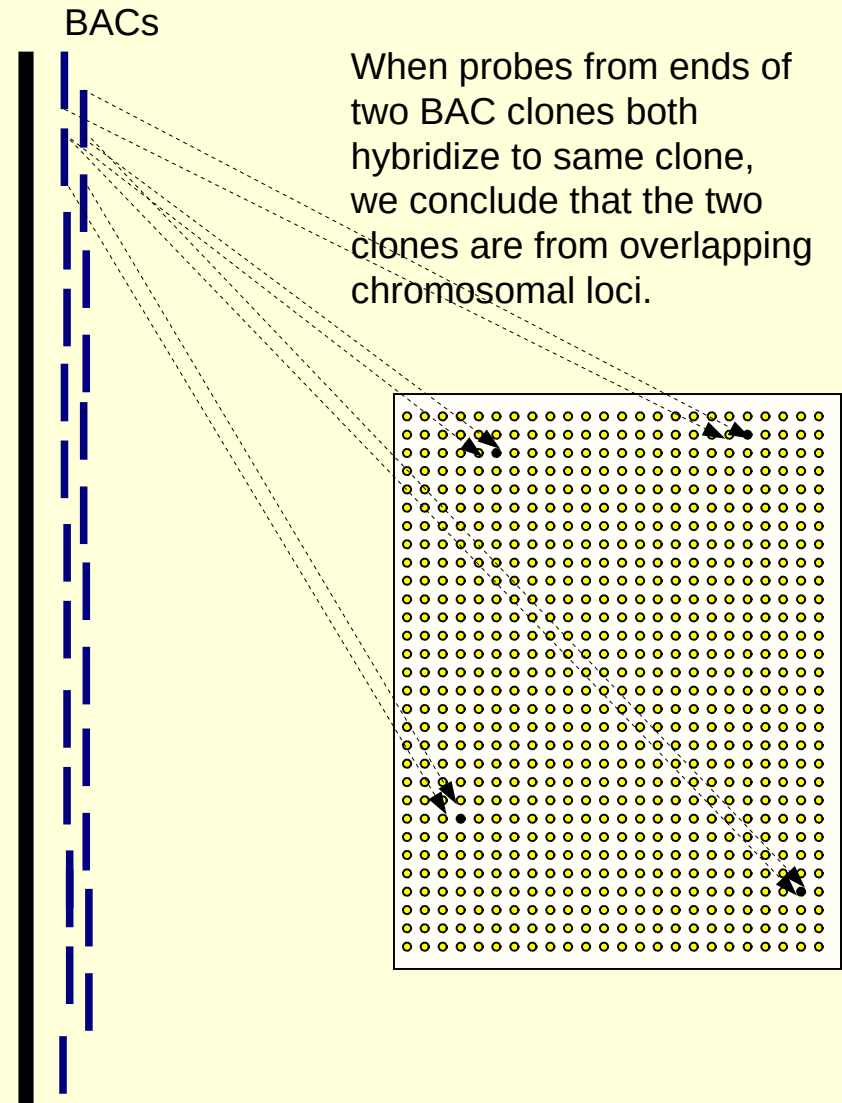
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# Genomic Sequencing - Divide and Conquer Strategy

1. Create a BAC Library
2. Create an ordered library by finding out which BACs cross-hybridize. Essentially, this is a chromosome walk along the entire genome.  
It may take hundreds or thousands of hybridizations to find at least one BAC for every part of every chromosome.
3. Sequence at least one BAC at each chromosomal location.

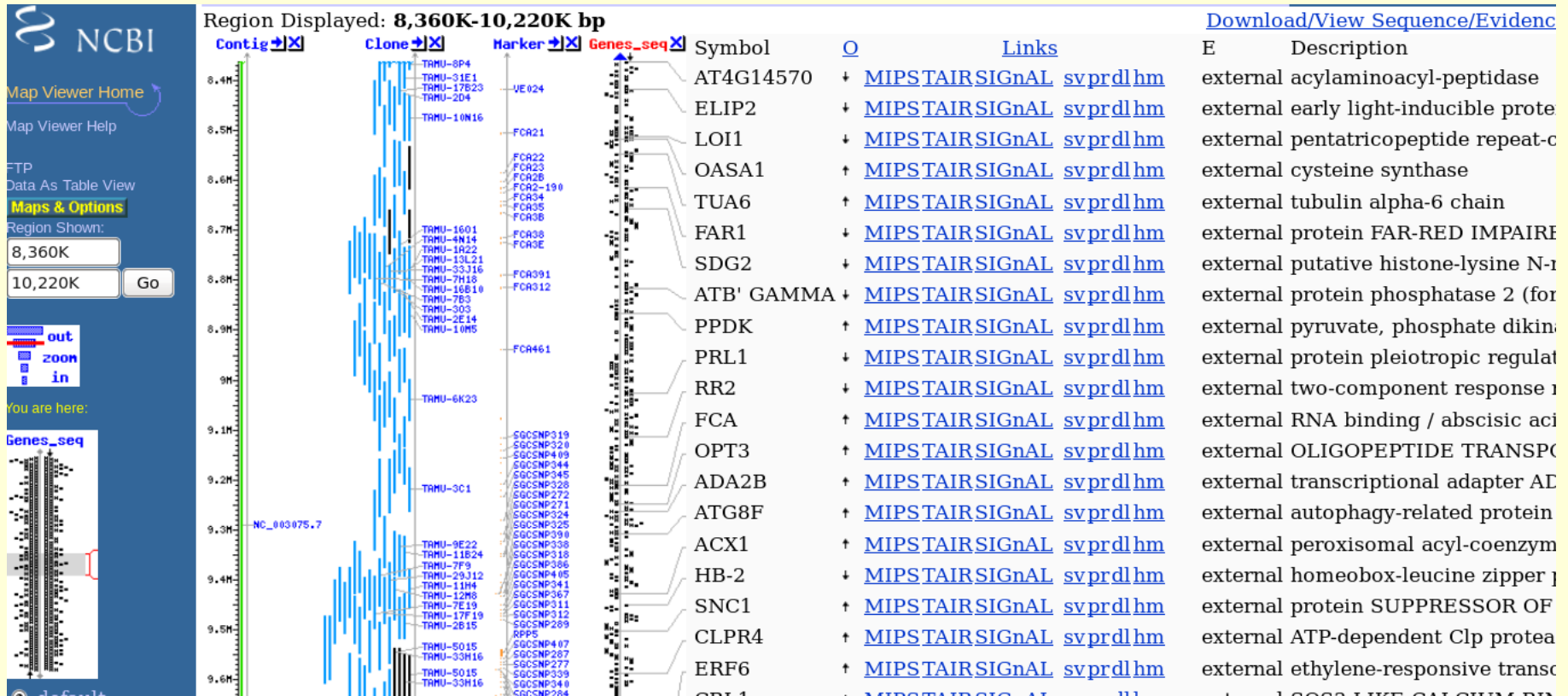
**ADVANTAGE** - This is the only way to be sure you have the entire genome.

**DISADVANTAGE** - This method takes years for most genomes.



# 2 Mb region of *A. thaliana* chromosome 4

This output from the genome map viewer at NCBI gives you some idea of how many BAC clones (light blue) were sequenced in a 2 million base region.



[http://www.ncbi.nlm.nih.gov/projects/mapview/maps.cgi?TAXID=3702&CHR=4&MAPS=cntg-r,clone,tair\\_ma rker,genes\[1.00%3A18585056.00\]&zoom=10](http://www.ncbi.nlm.nih.gov/projects/mapview/maps.cgi?TAXID=3702&CHR=4&MAPS=cntg-r,clone,tair_ma rker,genes[1.00%3A18585056.00]&zoom=10)

# Whole Genome Shotgun Sequencing (WGS)

A single BAC clone is usually 100 kb - 200 kb.

A plant genome may be  $10^7$  to  $10^{12}$  bp.

How do we sequence such large genomes?

In principle, the answer is: sequence millions of very short DNA fragments and use sophisticated software to find the overlaps to assemble complete chromosomal sequences.

In practice, this ideal is not possible for lots of reasons. Nonetheless, we can assemble many thousands of runs to create long contigs that span several thousand to several hundred thousands of bases each.

# High-throughput “Next Generation Sequencing” (NGS)

NGS technologies (eg. Illumina, IonTorrent) can generate several million short sequences in a single run. The short sequences can be overlapped, to assemble a sequence thousands, or even hundreds of thousands of bases long.

NGS technologies now make it possible to:

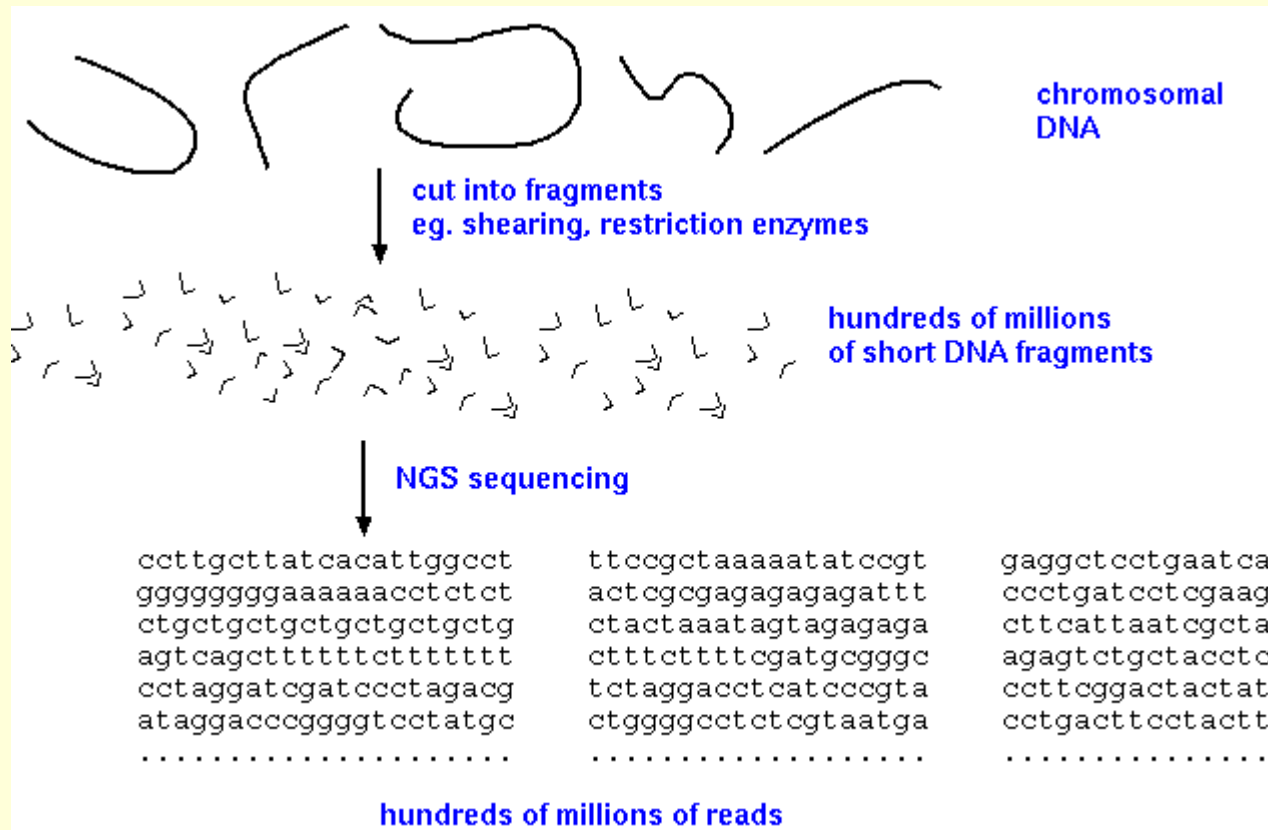
- generate almost complete genomic sequences for several thousand dollars
- generate sequences for DNA samples from mixed populations of microbes, allowing measurement of biodiversity (Metagenomics)
- generate sequences for RNA populations, making it possible to directly measure gene expression levels for thousands of genes simultaneously.

# Next Generation DNA sequencing

There are numerous NGS sequencing methods. Essentially all of them follow the same steps for sequencing a genome:

- 1) **fragment** genomic DNA into small fragments of a few hundred bases
- 2) **immobilize** individual DNA molecules onto a solid surface
- 3) **amplify** each molecule by PCR many thousands of times
- 4) perform **DNA synthesis** using nucleotides that emit a characteristic wavelength of light each time a base is added
- 5) **read** the sequence by imaging the emission of light in real time
- 6) from the millions of reads, **assemble** overlapping reads into long contiguous segments known as “contigs”

# Whole Genome Shotgun Sequencing (WGS)



# WGS by Illumina Sequencing

## **Principal**

sequencing by synthesis

as each base is added, a photon is emitted at a characteristic wavelength for A,G,C and T

## **Method**

Illumina video (5:03)

<https://youtu.be/fCd6B5HRaZ8>

Argonne Lab Video showing real Illumina images

<http://youtu.be/tuD-ST5B3QA>



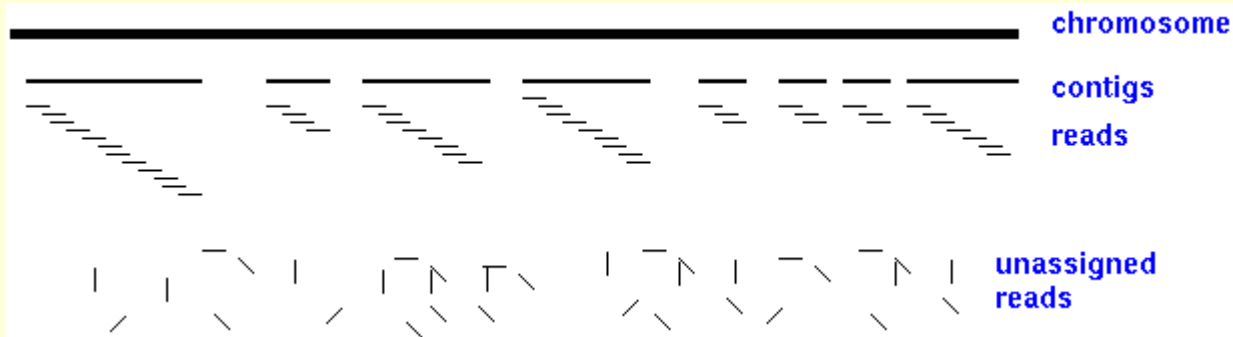
# Sequence Assembly

The screenshot shows a software window titled "Contig Editor: +21.2277 SRR006330.349400". The window contains a table of sequence reads with columns for read ID, SRR ID, offset, and sequence. A red text overlay reads: "Reads are assembled using programs that find overlaps between sequences from either strand." The table shows reads from SRR006330 and SRR006465. The bottom status bar indicates "Tag type:R454 Direction:- Comment: """.

Read ID	SRR ID	Offset	Sequence
+229822	SRR006330	.1168	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229823	SRR006330	.1122	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGT
-229824	SRR006330	.4625	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229825	SRR006330	.3062	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229826	SRR006330	.3381	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229827	SRR006330	.2563	GGTGCCATTCAACGTCAGCTTCCTTCCAAA
-229828	SRR006330	.3341	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229829	SRR006330	.2625	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229831	SRR006330	.1114	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229833	SRR006330	.2546	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCA
-229834	SRR006330	.2904	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTG
-229835	SRR006330	.3334	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229836	SRR006330	.3417	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229837	SRR006330	.3496	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229838	SRR006330	.3989	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
+229840	SRR006330	.4461	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
+229841	SRR006465	.5363	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCAT
-229842	SRR006330	.5115	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
+229843	SRR006330	.4208	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229844	SRR006330	.2371	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229845	SRR006465	.3088	GGTGCCATTCAACGTCAG
-229846	SRR006330	.1341	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229847	SRR006330	.2000	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229848	SRR006330	.1736	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229849	SRR006330	.2221	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCA*AGGTTTTGGTCAGTGACGTATTGGCCA
+229850	SRR006330	.3732	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229851	SRR006330	.5007	TCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
-229852	SRR006330	.3026	TCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA
<>	CONSENSUS	----	GGTGCCATTCAACGTCAGCTTCCTTCCAAACTCATCTTGGACTATCCATCAG*AGGTTTTGGTCAGTGACGTATTGGCCA

# Whole Genome Shotgun Sequencing (WGS)

High performance computers and software assemble millions of reads into contigs



- Contigs are built by overlapping reads.
- There are always gaps between contigs where the software cannot extend contigs any further.
- There are always large numbers of reads left over that cannot be assigned to a contig.
- The main reason: when reads contain parts of repetitive sequences, they may overlap thousands of other reads, making it impossible to uniquely determine overlaps. Consequently: **Very few eukaryotic genomes are ever completely sequenced.**

# Illumina Sequencing

## Advantages

- Very large number of reads
- Low cost
- Hundredfold coverage or greater

## Disadvantages

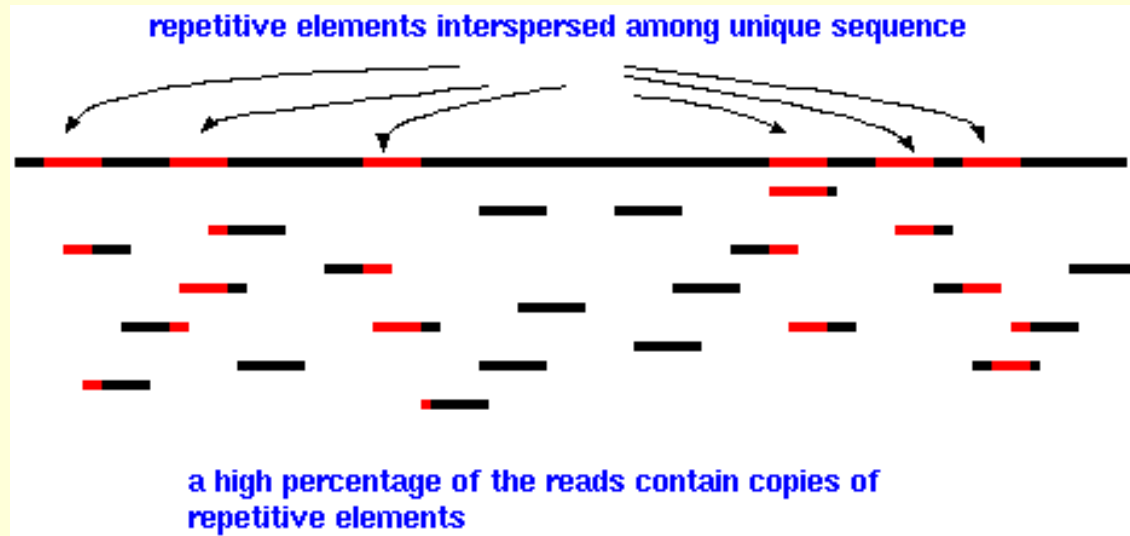
- Reads are short, 150 - 250 bp
- Error rate < 0.1 per 100 bases
- Short reads present many problems during sequence assembly

Example: 100 kb BAC clone:  $(100,000 \text{ bp}/150 \text{ bp per read}) \times 100 \text{ fold coverage}$   
= 40,000 reads.

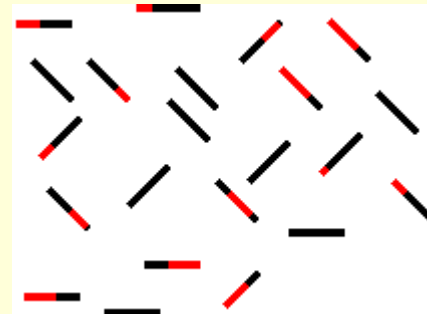
Example: Soybean Genome  $(1.1 \times 10^9 \text{ bp}/150 \text{ bp per read}) \times 100 \text{ fold coverage}$   
= 440 million reads

# Sequencing Very Large Genomes using Long Read Sequencing

Problem: Plant Genomes are mostly repetitive DNA.

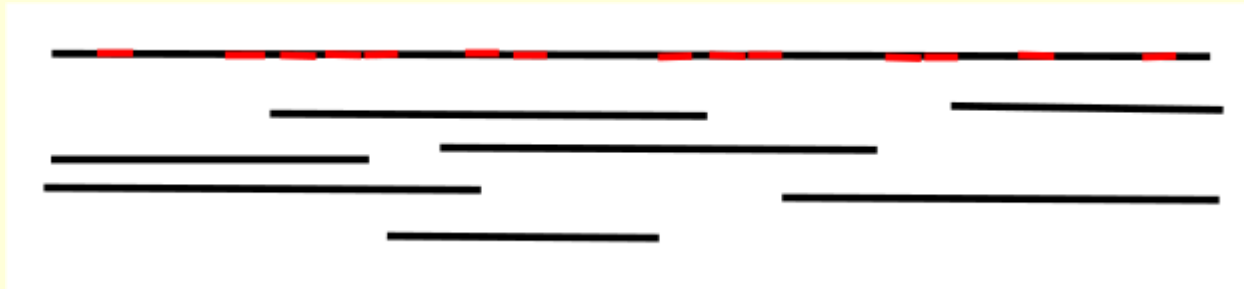


When the software tries to assemble from short reads, one copy of a repetitive sequence (red) will match every other copy of that sequence. There is no way to decide which reads overlap.



# Sequencing Very Large Genomes using Long Read Sequencing

Solution: Sequence longer reads!



Reads that are several kb in length will always have substantial amounts of unique DNA on either side of repetitive DNA sequences. The unique DNA gives sequence assemblers enough information to find a unique path to assemble overlapping reads into contigs.

# Long Read Sequencing

Long read methods sequence single DNA molecules. Instead of immobilizing the DNA fragments, long read methods pull long single-stranded DNAs through an enzyme. As the strand is pulled through, sequence is read. There are currently two competing technologies: PacBio SMRT sequencing and Oxford Nanopore.

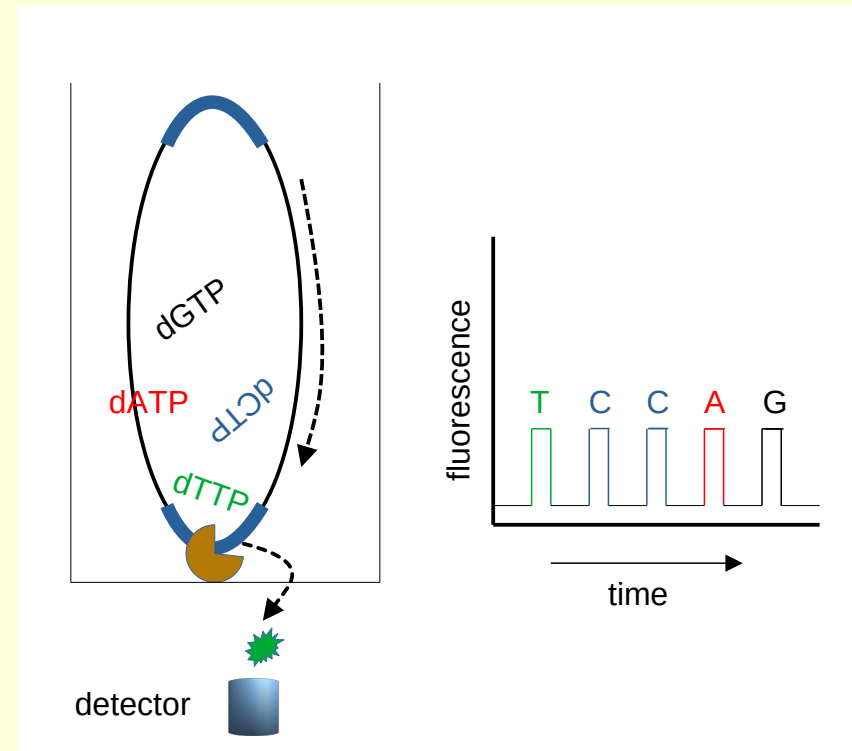
Amarashinge SL et al. (2020) Opportunities and challenges in long-read sequencing data analysis. *BMC Genome Biology* 21:30 <https://doi.org/10.1186/13059-020-1935-5>

# PAC Bio - SMRT sequencing

By adding adapters to both ends of a double-stranded fragment, a single-stranded circular molecule is created. The circular molecule is captured by a DNA polymerase which is anchored to the transparent floor of the cell. As each nucleotide is added, a photon is produced at a wavelength characteristic for A, G, C or T. Each nucleotide is therefore detected as a peak of fluorescence over time.

High accuracy is achieved by cycling the circular DNA through the DNA polymerase many times. The redundant sequence serves to confirm the nucleotide read at each position.

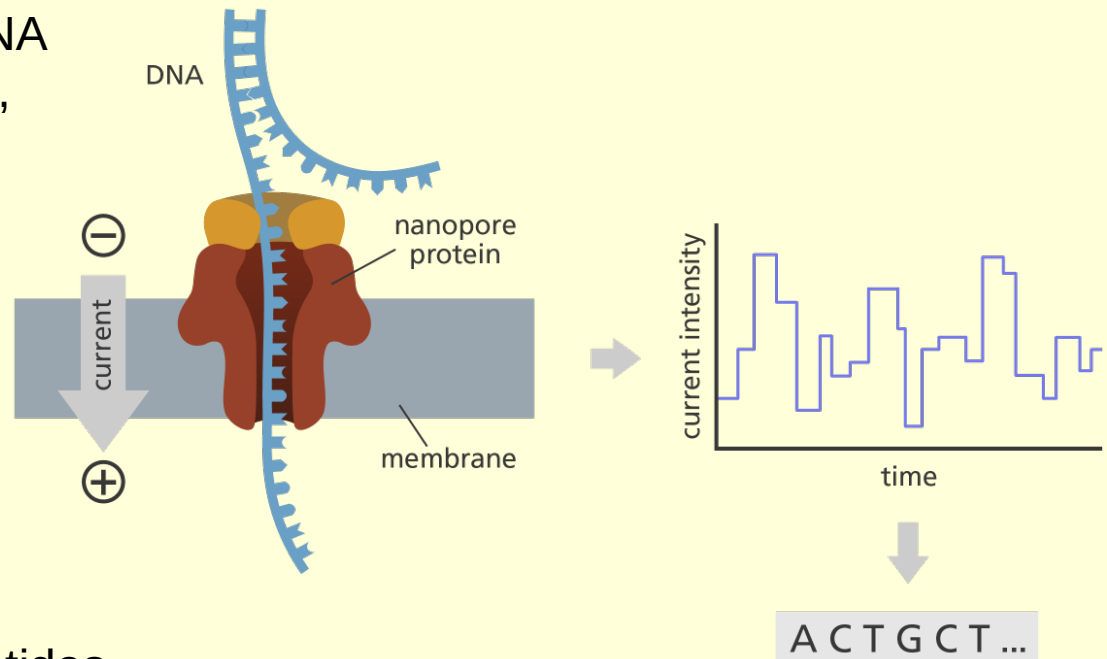
2021 - Average read lengths are around 30 kb.



# Oxford nanopore sequencing

Nanopore utilizes a membrane bound motor protein,  $\alpha$ -hemolysin, which has been genetically engineered to pull a single strand of DNA through a membrane pore.

As the single-stranded DNA enters the lower chamber, the conductivity of the solution at the inner surface of the membrane changes. Because each of the four nucleotides changes the conductivity to a different degree, a plot of current intensity over time gives a reading of the sequence of nucleotides as they emerge from the pore.



As of 2021, the average read lengths range between 15 - 20 kb. The longest reads often exceed 1000 - 2000 kb.