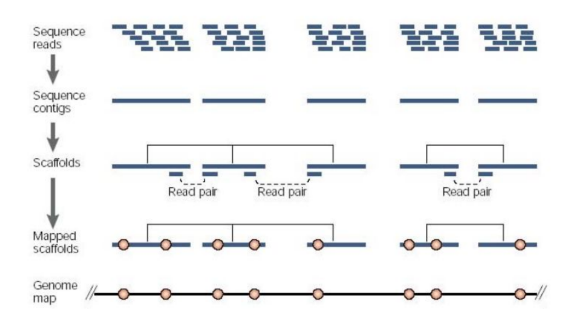
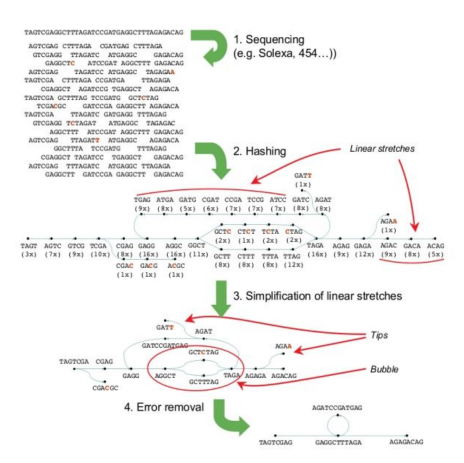
Scaffolding

de novo whole-genome shotgun assembly



Contig construction



Scaffolding

Both OLC and DBG are concerned with constructing the longest, most accurate *contigs* possible

Scaffolding orders and orients *contigs* with respect to each other

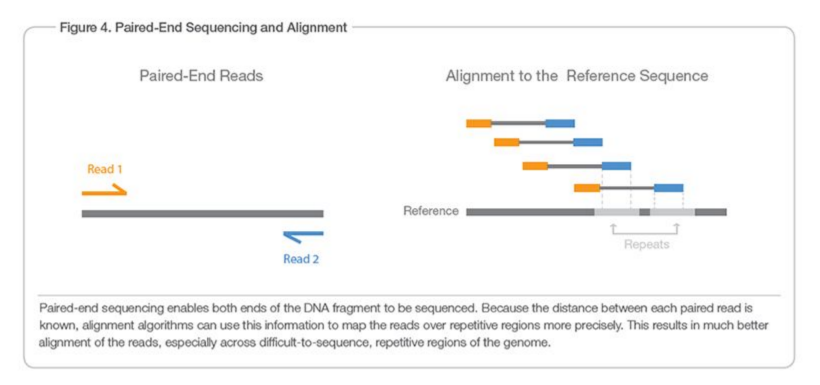
For this we can use data from various sources, especially paired ends.

Contig: is a stretch of unambiguously assembled sequence.

Scaffold: may contain gaps.

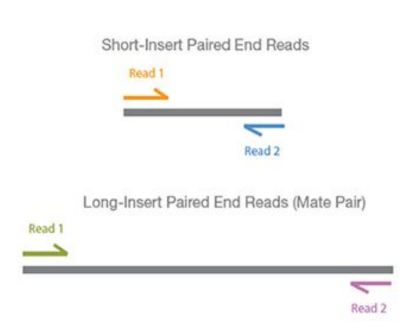
Paired-end sequencing





Vocabulary

- Paired end reads: read1, insert < 500 bp, read2
- Mate pair reads: read1,
 insert > 1 kbp, read2
- k-mer: any sequence of length k
- Contig: gap-less assembled sequence
- Scaffold: sequence which may contain gaps



Alternative protocol produces a *pair* of reads taken from either end of a longer fragment

Paired reads are also called *mates* to distinguish them from the *unpaired* reads we've been discussing

GCATCATTG

GCATCATTG

GCATCATTG

GCATCATTG

Mate 1

Fragment

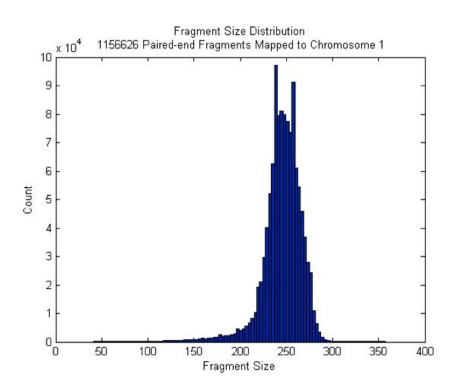
GCATCATAAAAACC

Mate 2

Depending on lengths, mates might overlap in the middle of the fragment

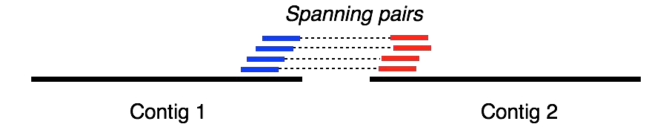
Example fragment length distribution

Fragments are not exactly the same length, but there's a clear peak around 250 nt, very few < 150 nt or > 300 nt



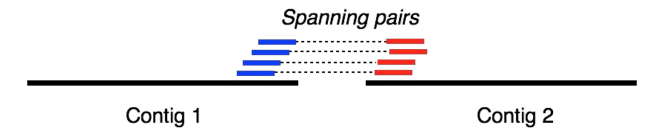
Say we have a collection of pairs and we assemble them as usual.

Assembly yields two contigs:



...and we discover that some of the mates at one edge of contig 1 are paired with mates in contig 2

Call these spanning pairs

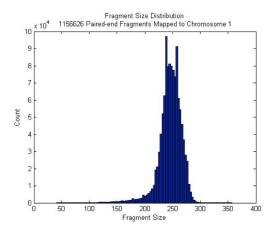


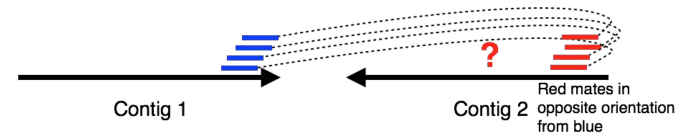
What does this tell us?

Contig 1 is close to contig 2 in the genome.

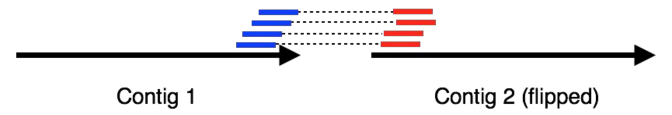
In fact, we can *estimate distance between contigs* using what we know about fragment length distribution.

The more spanning pairs we have, the better our estimate.





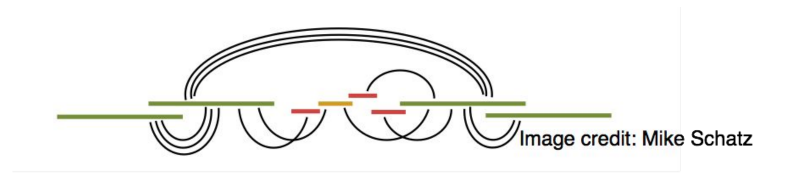
What does the picture look like if contigs 1 and 2 are close, but we assembled contig 2 "backwards" (i.e. reverse complemented)



Pairs also tell us about contigs' relative *orientation*

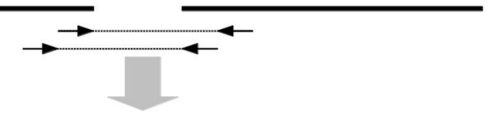
Scaffolding

Scaffolding output: collection of *scaffolds*, where a scaffold is a collection of contigs related to each other with high confidence using pairs.



Scaffolds construction in practice (lab)

- Scaffolding using pairing information
- This is why libraries containing multiple insert sizes are used:
 - Gnerre 2011:
 - 45x overlapping PE reads (insert size: 180 bp, reads: >100 bp)
 - 45x short jump MP reads (insert size: 3 kb)
 - 5x (optional) long jump MP reads (insert size: 6 kb)
 - 1x (optional) fosmid jump MP reads (insert size: 40 kb)
 - Ribeiro 2012:
 - 50x overlapping PE reads (insert size: 180 bp, reads: >100 bp)
 - 50x PacBio reads (reads: 1-3 kb)



Scaffolds construction

- Scaffolding using pairing information
- Techniques:
 - Jumping libraries
 - Linked reads (10x Genomics, Dovetail Genomics Chicago libraries)
 - Long reads (PacBio, Oxford nanopore)
 - Structural maps (BioNano)

Gap closing / contig extension

