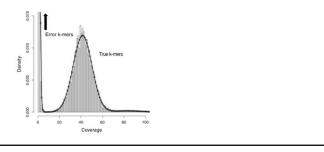


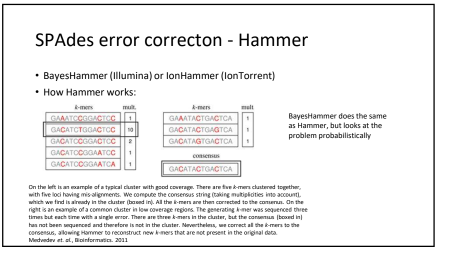
Velvet vs SPAdes

- SPAdes developed to be able to assemble single-cell sequence data
- Single-cell data:
 - Not uniform coverage
- Three main differences between Velvet and SPAdes
 - Error correction
 - Graph construction
 - Graph simplification/resolution
- Other differences too, but won't go into that here

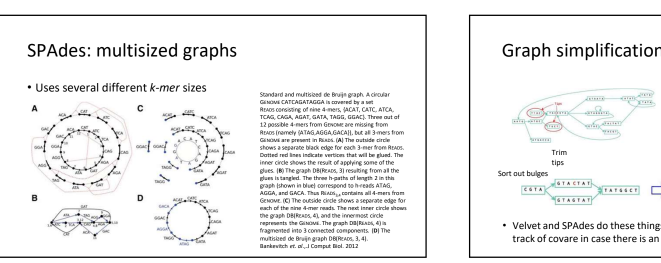
Velvet error correction

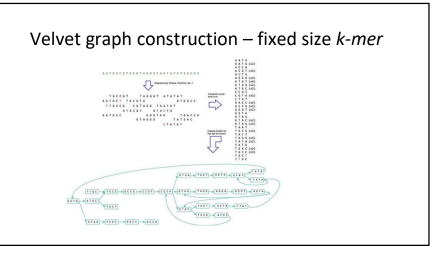
- Velvet: expects uniform coverage
- Uses high coverage k-mers to error correct low coverage k-mers

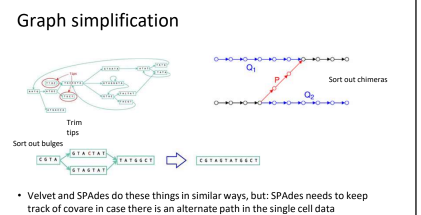


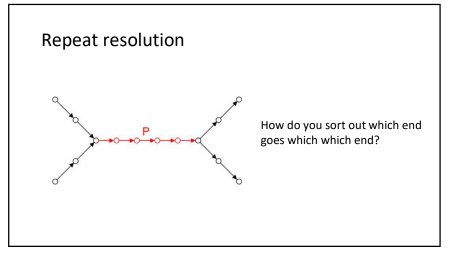


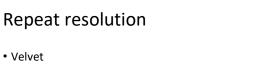












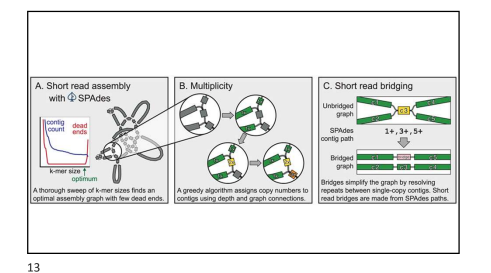
- Looks at the reads connecting longer contigs
- Uses paired read information to "straighten" out the repeats
- SPAdes
 - Uses read pair information
 - Creates a paired de Bruijn graph each node a pair
 - Much sparser than the "normal" graph

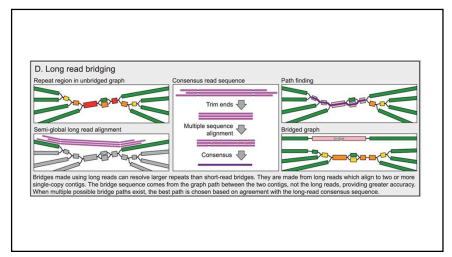
SPAdes options

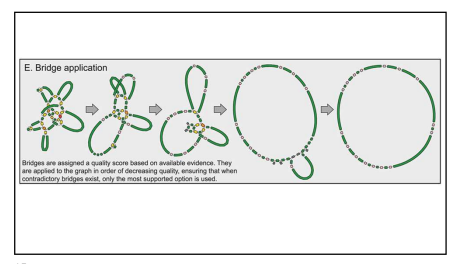
- · hybrid spades short and long read data
- metaSPAdes a pipeline for metagenomic data sets
- plasmidSPAdes a pipeline for extracting and assembling plasmids from WGS data sets
- metaplasmidSPAdes a pipeline for extracting and assembling plasmids from metagenomic data sets
- rnaSPAdes a de novo transcriptome assembler from RNA-Seq data.
- truSPAdes a module for TruSeq barcode assembly
- biosyntheticSPAdes a module for biosynthetic gene cluster assembly with paired-end reads

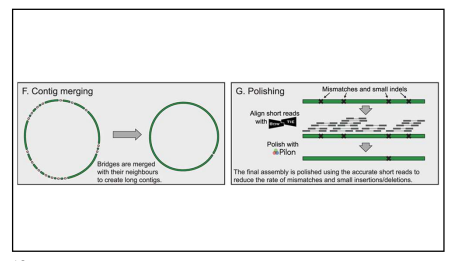
Unicycler - hybrid assembler

- Short read, long read and hybrid assembler
- Short only SPAdes optimizer
- Long reads map (miniasm), assemble w. overlap, polish (racon)
- Hybrid:
 - Create SPAdes assembly
 - Scaffold with long reads
- Note: only for bacterial genomes!





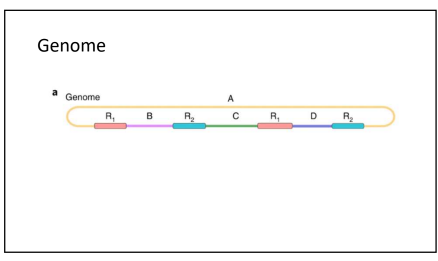




Flye - long read assembly

- Only works on long reads
- Follows the assemble-then-correct approach
- Quite fast, and especially good for resolving repeats
- Alternative: canu
- Correct-assemble
- $\circ~$ Thorough, but takes a long time

17



Flye procedure Find k-mers in reads, find reads with shared k-mers Find reads that have overlaps Assemble contigs from overlaps - called disjointigs Reconstruct graph Resolve repeats Polish Output assembly

18

