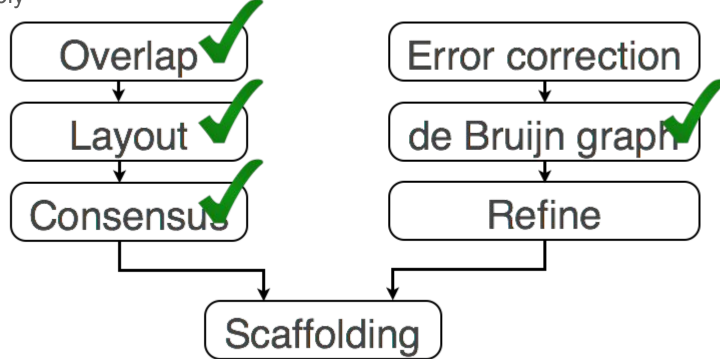
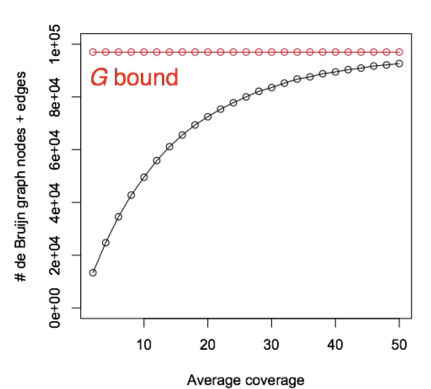
Assembly paradigms

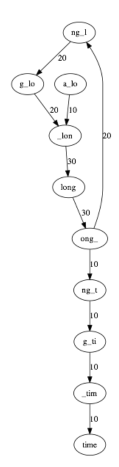
1: Overlap-Layout-Consensus (OLC) assembly 2: de Bruijn graph (DBG) assembly



When data is error-free, # nodes, edges in de Bruijn graph is $O(\min(G, N))$

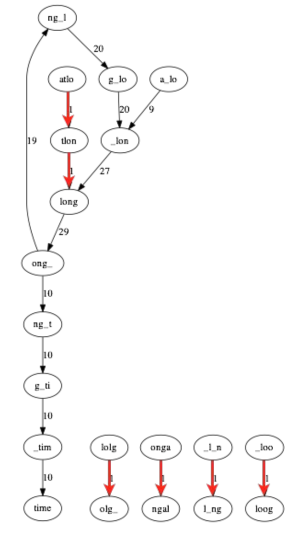


What about data with sequencing errors?

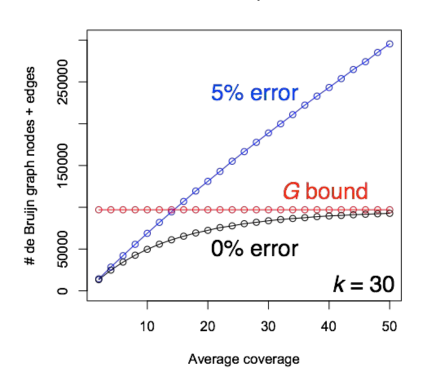


Take an example we saw (left) and mutate a *k*-mer character to a random other character with probability 1% (right)

6 errors result in 10 new nodes and 6 new *weighted* edges, all with weight 1



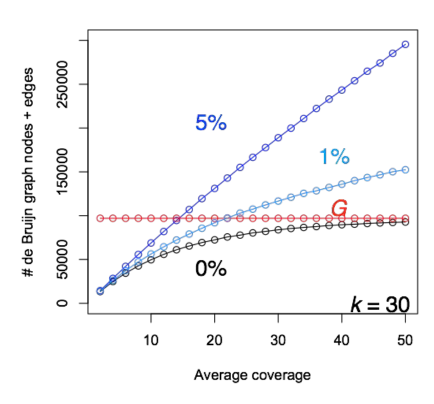
As more *k*-mers overlap errors, # nodes, edges approach *N*



Same experiment as before but with 5% error added

Errors wipe out much of the benefit of the *G* bound

Instead of $O(\min(G, N))$, we have something more like O(N)



If we can correct sequencing errors up-front, we can prevent De Bruijn graph from growing much beyond the *G* bound.

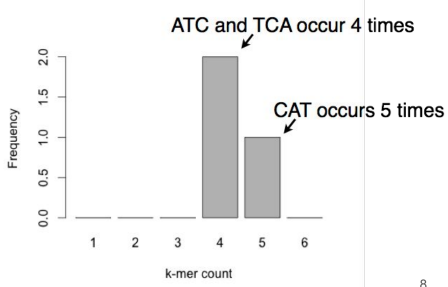
How do we correct errors?

Analogy: design a spell checker for a language you've never seen before. How do you come up with suggestions?

k-mer count histogram:

x axis is an integer k-mer count, y axis is # distinct k-mers with that count

Right: such a histogram for 3-mers of CATCATCATCATCAT:

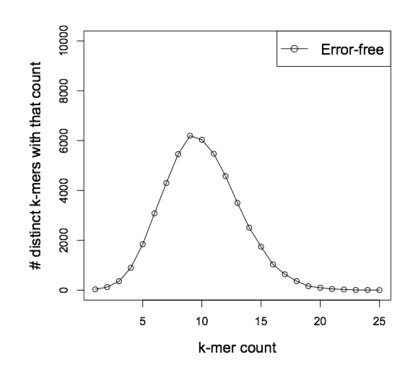


Say we have error-free sequencing reads drawn from a genome. The amount of sequencing

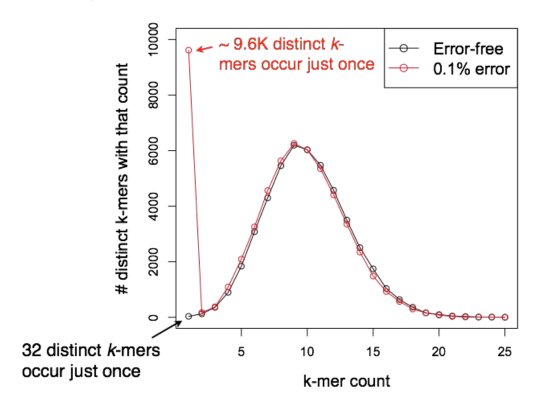
is such that average coverage = 200. Let k = 20.

How would the picture change for data with 1% error rate?

Hint: errors usually change high-count *k*-mer into low-count *k*-mer

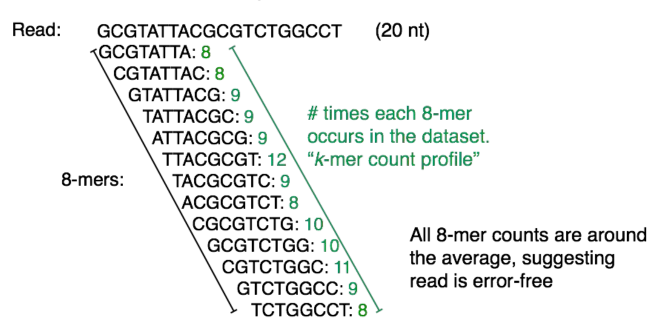


k-mers with errors usually occur fewer times than error-free *k*-mers



Idea: errors tend to turn frequent *k*-mers to infrequent *k*-mers, so corrections should do the reverse

Say we have a collection of reads where each distinct 8-mer occurs an average of ~10 times, and we have the following read:



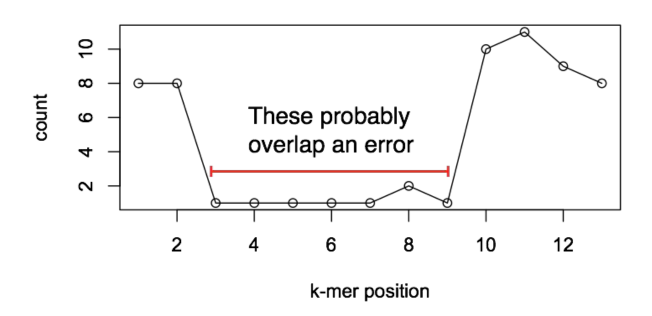
Suppose there's an error

```
Read:
       GCGTACTACGCGTCTGGCCT
        GCGTACTA: 1
                                        k-mer count profile has
         CGTACTAC: 3
                         Below average
                                        corresponding stretch of
          GTACTACG: 1
                                        below-average counts
            TACTACGC: 1
             ACTACGCG: 2
              CTACGCGT: 1
               TACGCGTC: 9
                ACGCGTCT: 8
                 CGCGTCTG: 10
                                  Around average
                   GCGTCTGG: 10
                     CGTCTGGC: 11
                      GTCTGGCC: 9
                        TCTGGCCT: 8
```

k-mer count profiles when errors are in different parts of the read:

```
GCGTACTACGCGTCTGGCCTGCGTATTACACGTCTGGCCT GCGTATTACGCGTCTGGTCT
GCGTACTA: 1
                        GCGTATTA: 8
                                                GCGTATTA: 8
 CGTACTAC: 3
                        CGTATTAC: 8
                                                CGTATTAC: 8
  GTACTACG: 1
                          GTATTACA: 1
                                                  GTATTACG: 9
    TACTACGC: 1
                           TATTACAC: 1
                                                   TATTACGC: 9
     ACTACGCG: 2
                            ATTACACG: 1
                                                    ATTACGCG: 9
      CTACGCGT: 1
                                                     TTACGCGT: 12
                             TTACACGT: 1
       TACGCGTC: 9
                              TACACGTC: 1
                                                      TACGCGTC: 9
        ACGCGTCT: 8
                               ACACGTCT: 2
                                                       ACGCGTCT: 8
         CGCGTCTG: 10
                                CACGTCTG: 1
                                                        CGCGTCTG: 10
          GCGTCTGG: 10
                                 GCGTCTGG: 10
                                                         GCGTCTGG: 10
           CGTCTGGC: 11
                                  CGTCTGGC: 11
                                                           CGTCTGGT: 1
             GTCTGGCC: 9
                                    GTCTGGCC: 9
                                                            GTCTGGTC: 2
              TCTGGCCT: 8
                                     TCTGGCCT: 8
                                                             TCTGGTCT: 1
```

k-mer count profile indicates where errors are



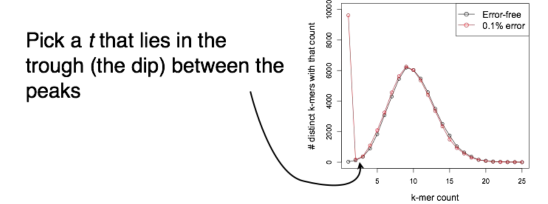
Simple algorithm: given a count threshold *t*:

For each read:

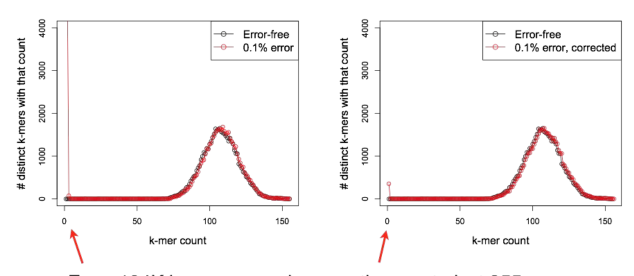
For each k-mer:

If k-mer count < t:

Examine k-mer's neighbors within certain Hamming/edit distance. If neighbor has count $\geq t$, replace old k-mer with neighbor.



Corrects 99.2% of the errors in the example 0.1% error dataset

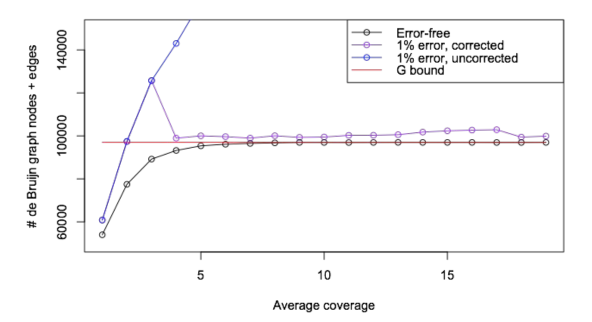


From 194K k-mers occurring exactly once to just 355

Error correction: results

For uncorrected reads, De Bruijn graph size is off the chart.

For corrected reads, De Bruijn graph size is near G bound.



For error correction to work well:

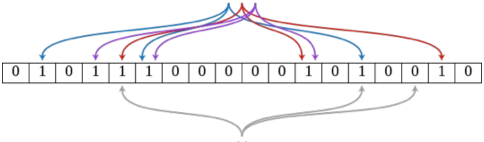
- Average coverage should be high enough and k should be set so we can distinguish infrequent from frequent k-mers
- *k*-mer neighborhood we explore must be broad enough to find frequent neighbors. Depends on error rate and *k*.
- Data structure for storing k-mer counts should be substantially smaller than the De Bruijn graph
 - Otherwise there's no point doing error correction separately
 - Counts don't have to be 100% accurate; just have to distinguish frequent and infrequent

Error correction: sketches

Sketch data structures are extremely compact, but fail sometimes

E.g. a Bloom Filter is like a hash set, but far smaller, and will sometimes say an object is in





CountMin sketches generalize Bloom Filters for histograms (sets where elements have associated counts); reported counts might be too high

These are candidates for compactly storing *k*-mer counts:

- http://en.wikipedia.org/wiki/Bloom_filter
- http://en.wikipedia.org/wiki/Count-Min_sketch