Copy Number and Structural Variation

Chris Miller

Some slides adapted from:

Malachi Griffith, Obi Griffith, Fouad Yousif High-Throughput Biology: From Sequence to Networks

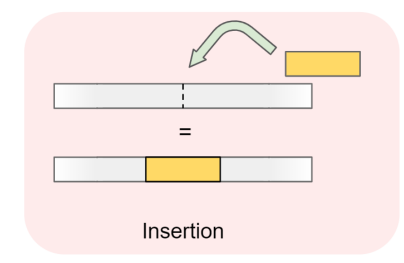
https://github.com/griffithlab/rnaseq_tutorial_wiki/blob/master/LectureFiles/cbw-cshl/2017/IGV Tutorial Brief.pptx

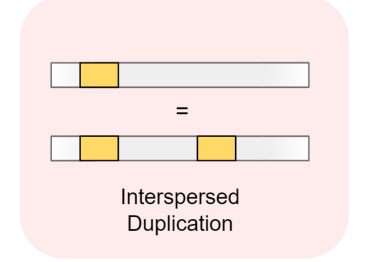
Tobias Rausch

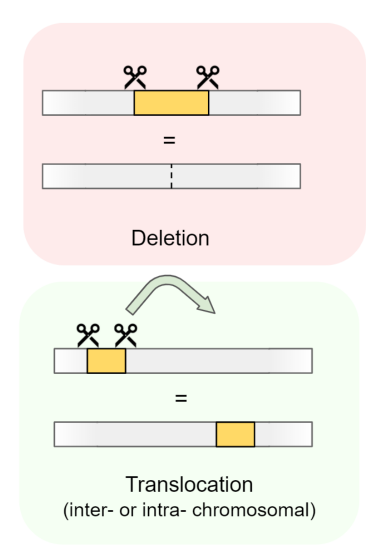
Structural and copy-number variation analysis

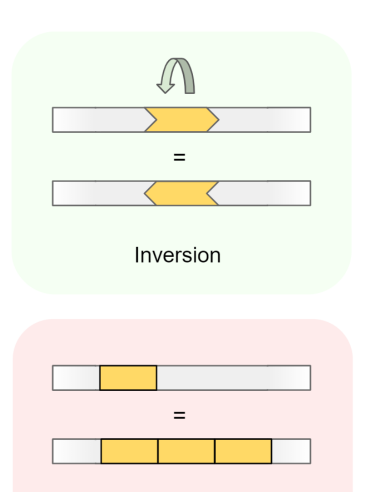
https://www.ebi.ac.uk/training/materials/cancer-genomics-materials/structural-and-copy-number-variation-mutational-signatures/structural-and-copy-number-variation-analysis/







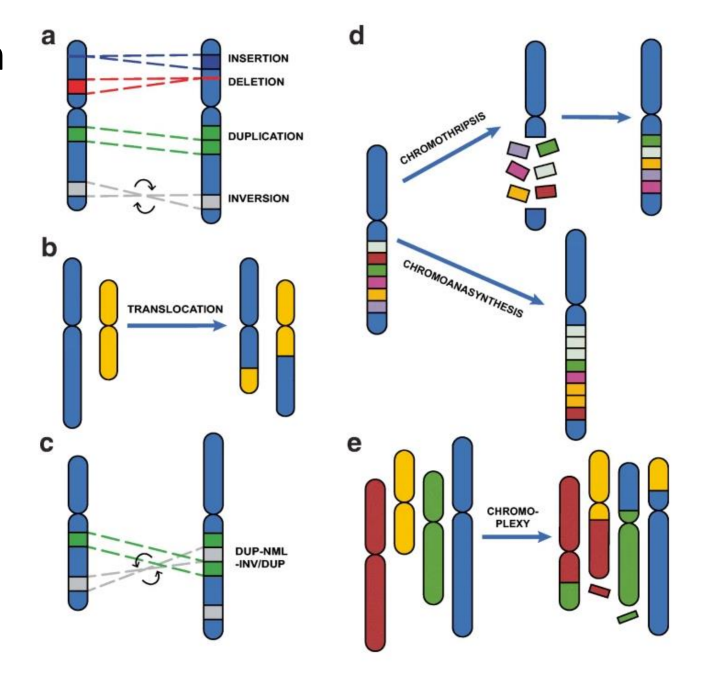




Tandem Duplication

Types of Structural Variation

SVs can get complicated!

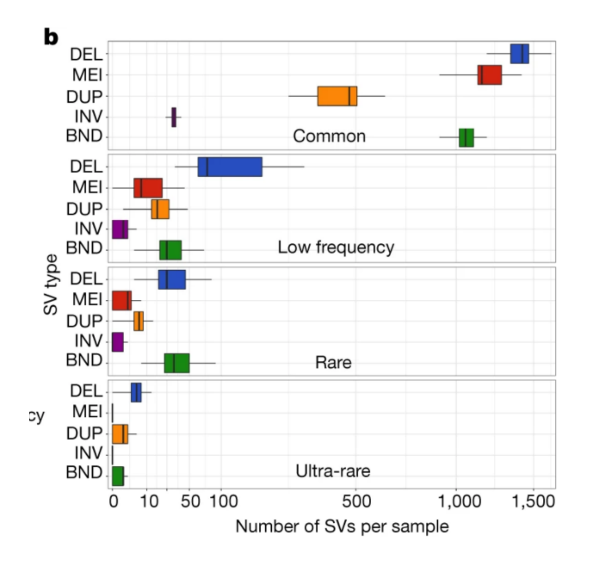


Genomic diversity from SVs

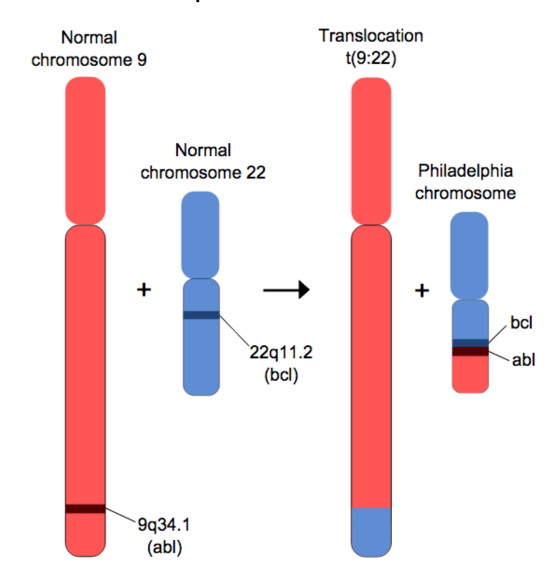
 Underappreciated due to past limitations of technology

- probably about 1% of each genome (by bp) differs from the reference
 - only 0.1% different by SNPs

SV calls from WGS of 14,623 samples



Somatic SVs are a frequent cause of cancer



Polyploidy

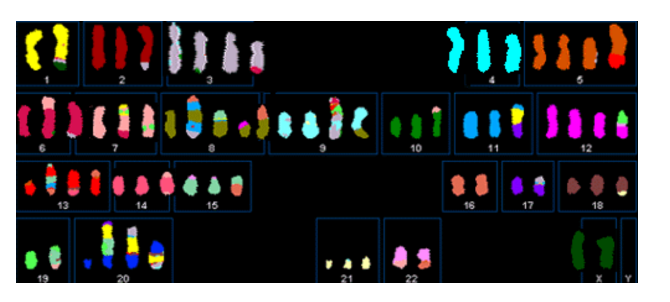
Haploid (N) Diploid (2N) Triploid (3N) Tetraploid (4N)

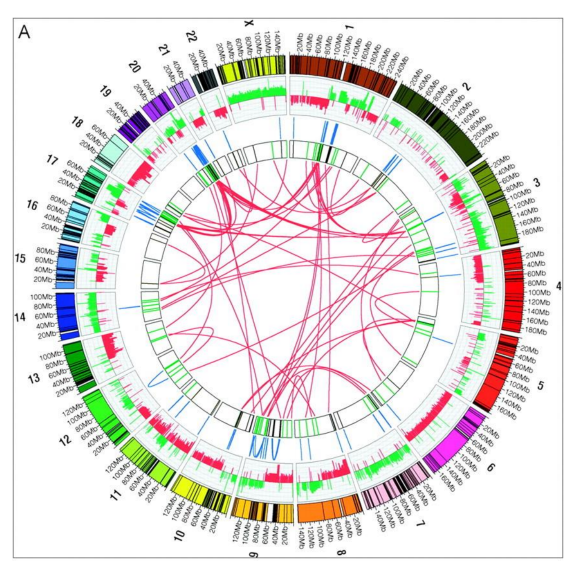
Whole Genome Doubling

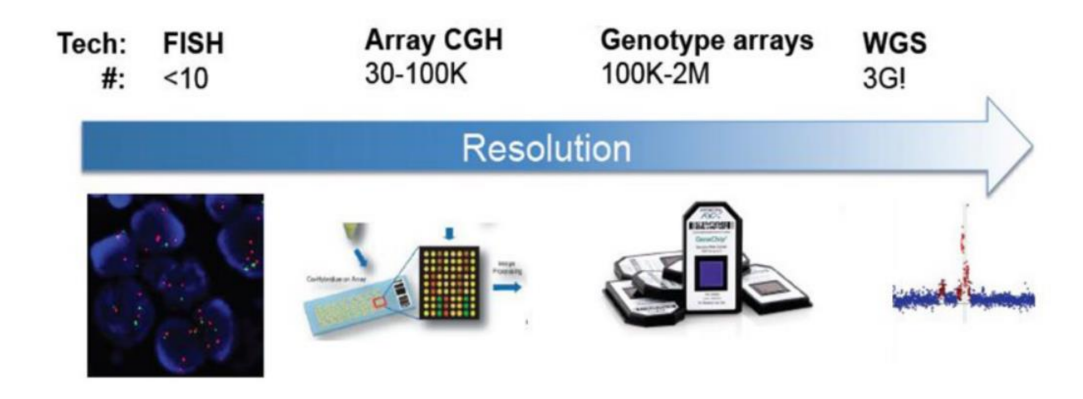


Somatic SVs are a frequent cause of cancer

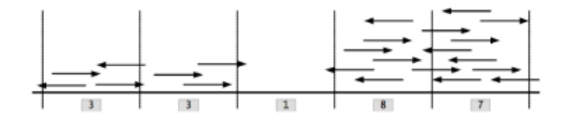
MCF7 Breast Cancer cell line





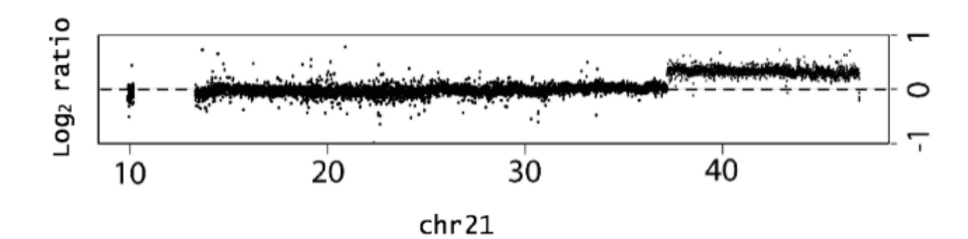


Read counting in windows for tumor and normal data

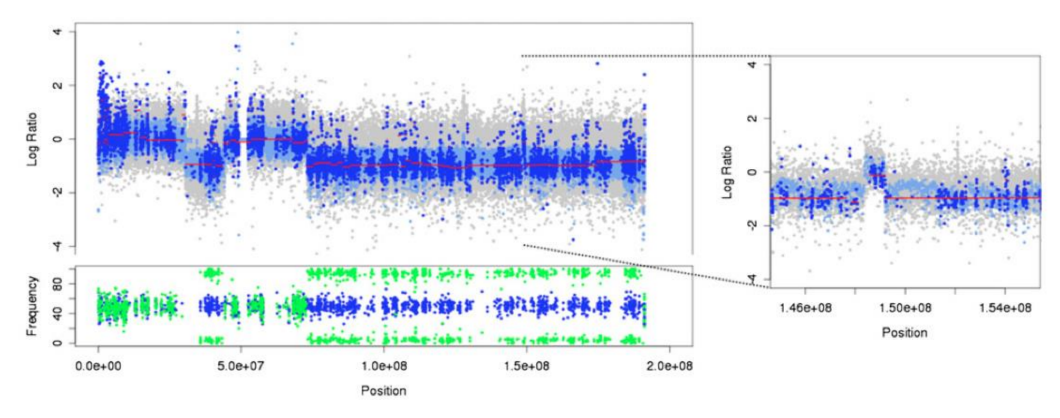


- Log2 ratio for each window
- Chromosome-wide plot

$$log_2 \frac{\# Reads_{Disease}}{\# Reads_{Normal}}$$



Gets more complicated with targeted sequencing, but still works!



B-allele frequency for CN-neutral Loss of Heterozygosity

- Other factors:
 - Sample prep
 - GC-bias
 - Probe affinities
 - Sample Purity
 - Subclonal populations

• Cleaner data, deeper data = higher resolution

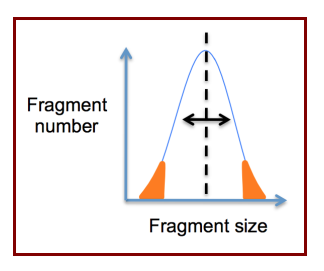
There are few decent packages for doing this

CNVkit is my go-to algorithm these days

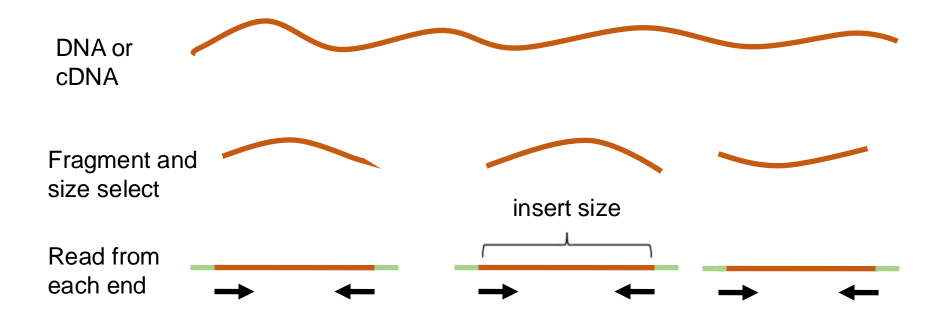
Paired-end sequencing

DNA or cDNA

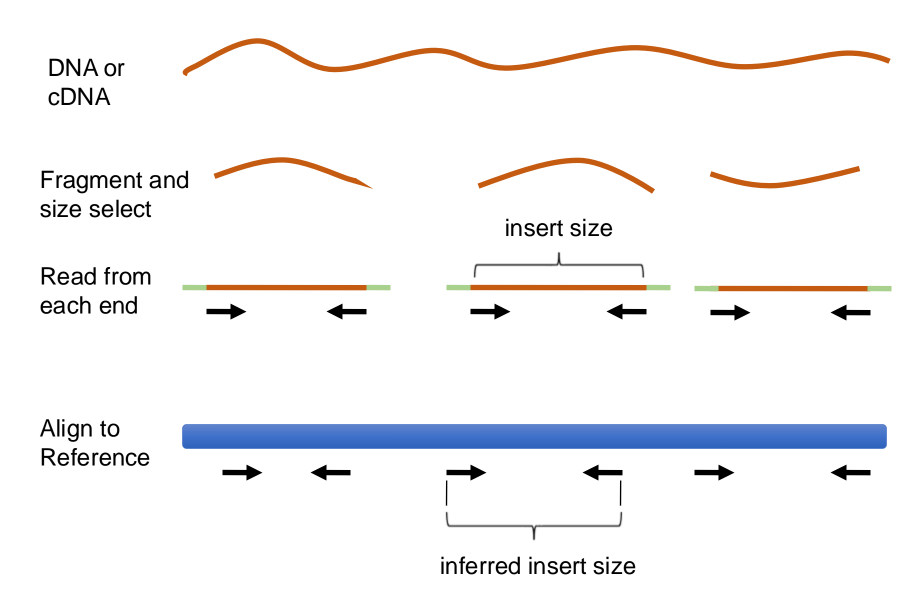
Fragment and size select



Paired-end sequencing



Paired-end sequencing

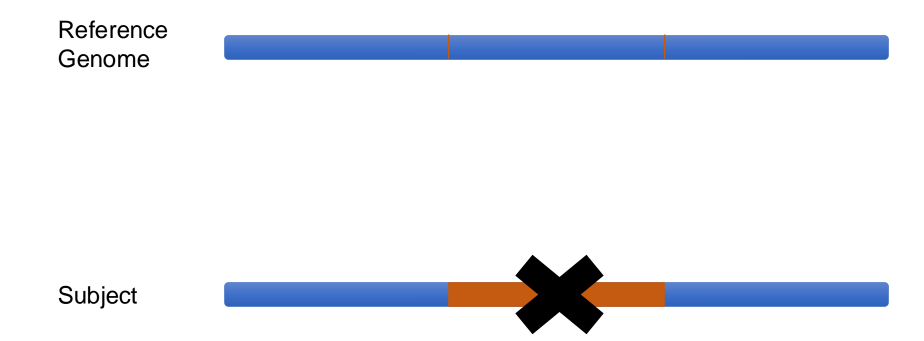


Interpreting inferred insert size

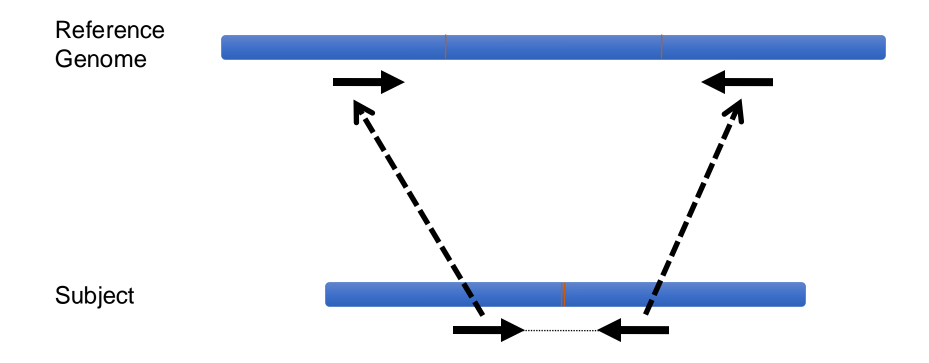
The "inferred insert size" can be used to detect structural variants including

- Deletions
- Insertions
- Inter-chromosomal rearrangements: (Undefined insert size)

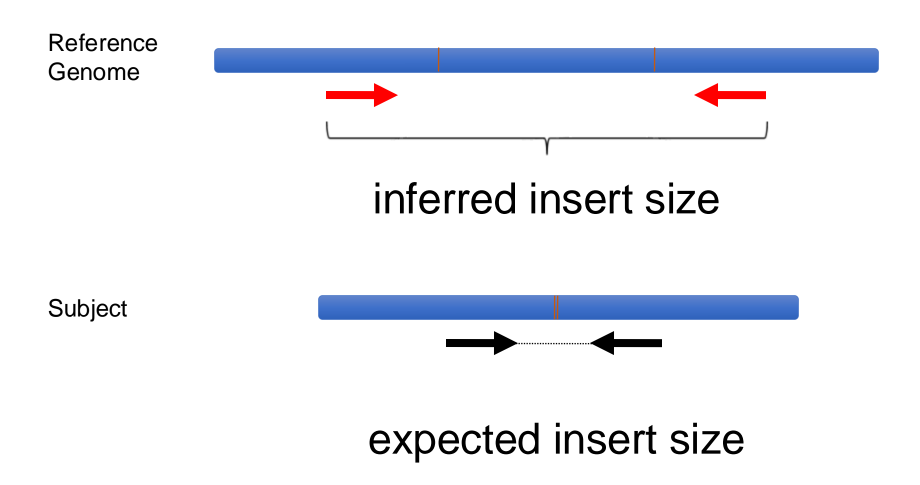
What is the effect of a deletion on inferred insert size?



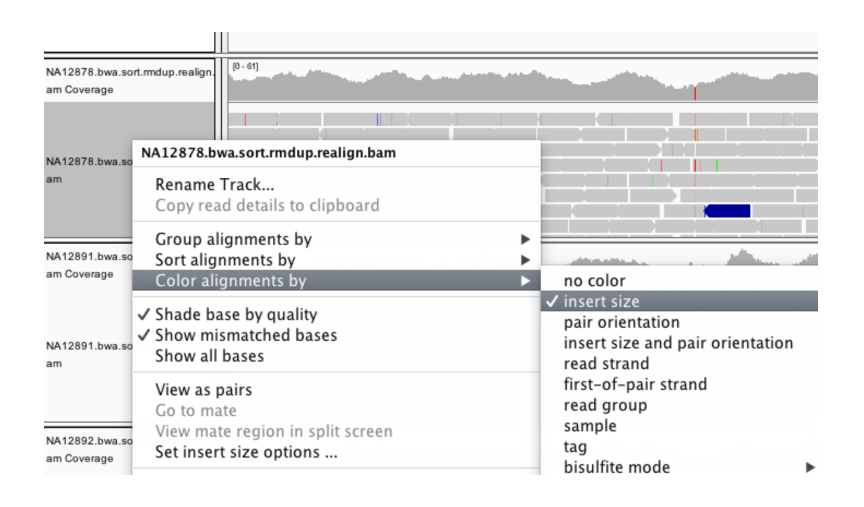




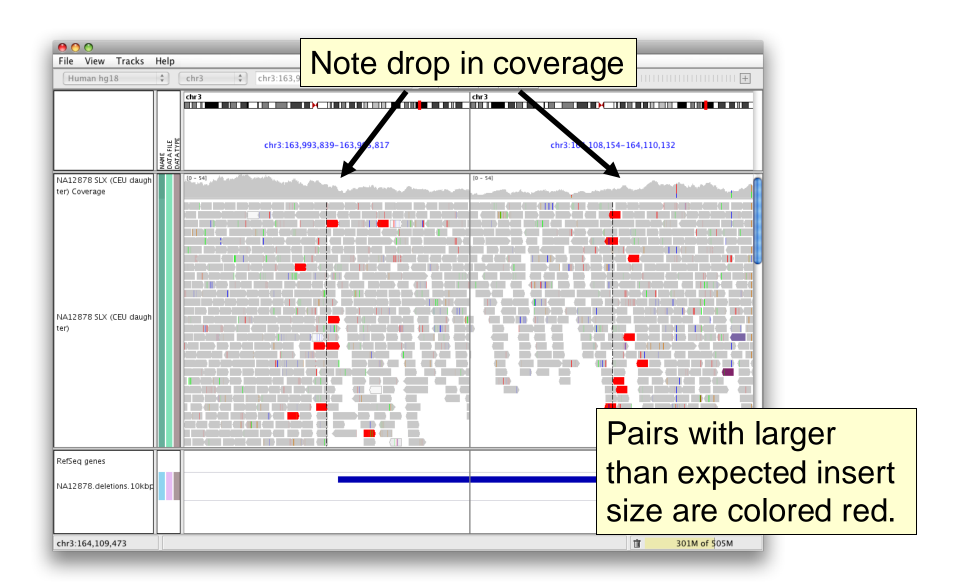
Inferred insert size is > expected value



Color by insert size







Insert size color scheme

• Smaller than expected insert size:

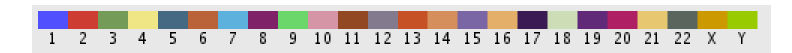


Larger than expected insert size:

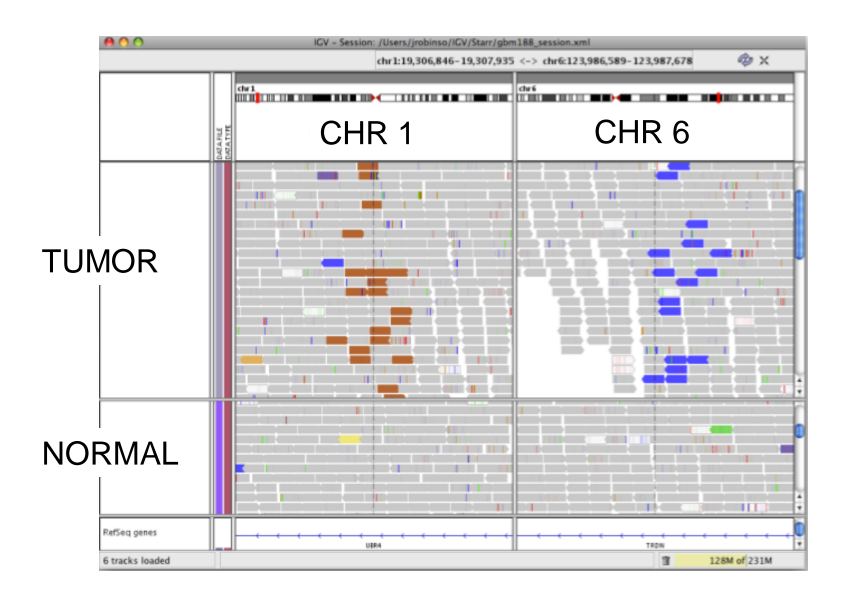


Pairs on different chromosomes

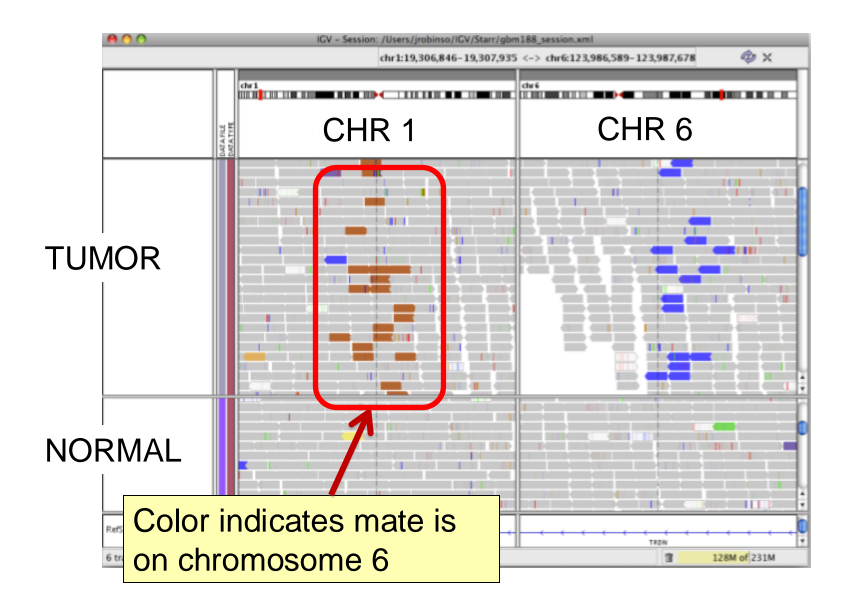
Each end colored by chromosome of its mate



Rearrangement



Rearrangement



Interpreting Read-Pair Orientations

Orientation of paired reads can reveal structural events:

- Inversions
- Duplications
- Translocations
- Complex rearrangements

Orientation is defined in terms of

- read strand, left vs right, and
- read order, first vs second

Interpretation of read pair orientations

LR Normal reads.

The reads are left and right (respectively) of the unsequenced part of the sequenced DNA fragment when aligned back to the reference genome.

LL,RR Implies inversion in sequenced DNA with respect to reference.

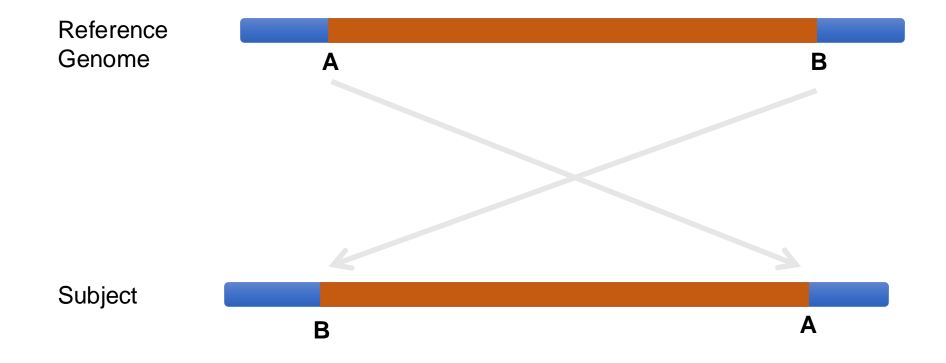
RL Implies duplication or translocation with respect to reference.

These categories only apply to reads where both mates map to the same chromosome.

Figure courtesy of Bob Handsaker

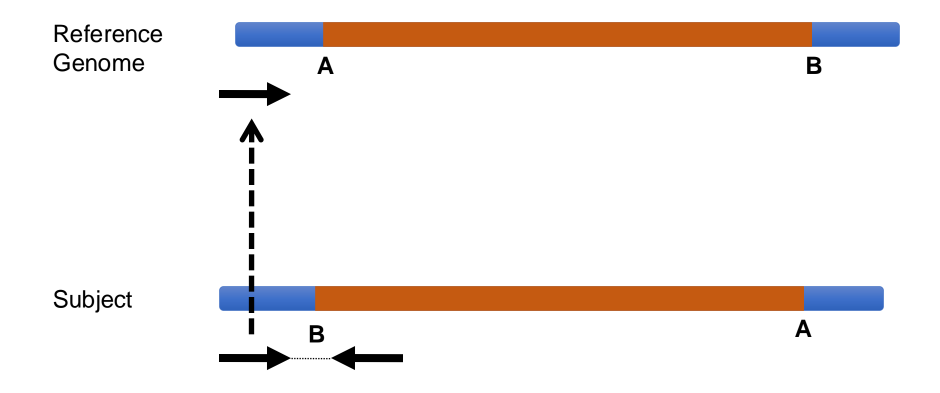
Reference genome

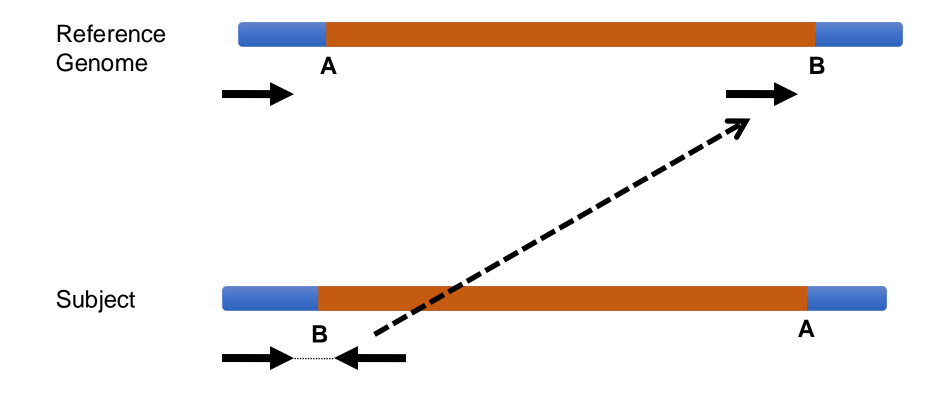




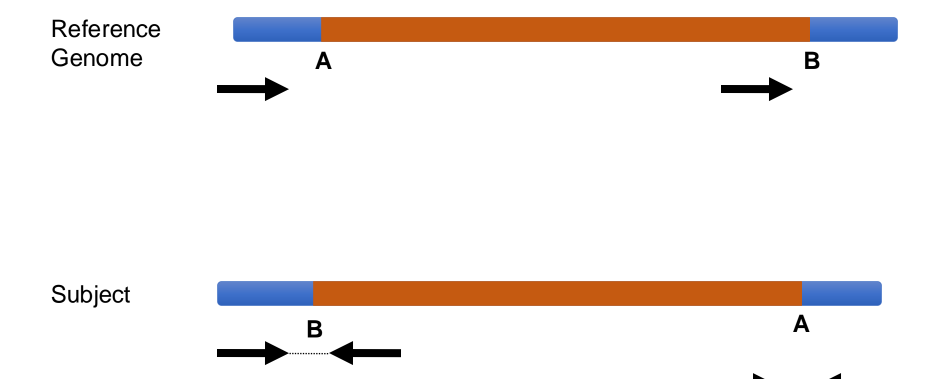


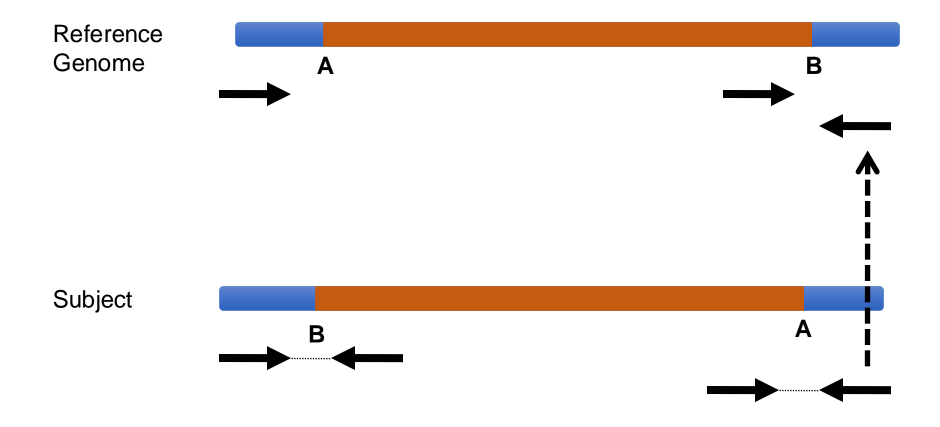


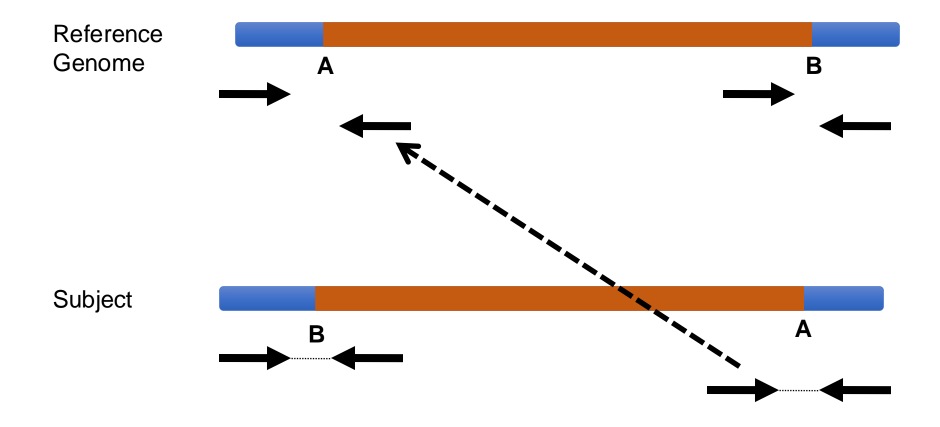


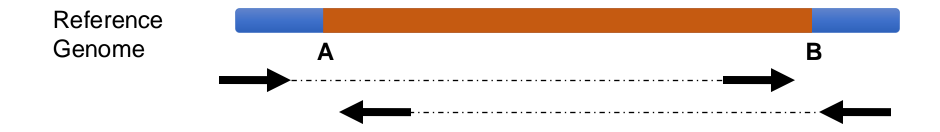










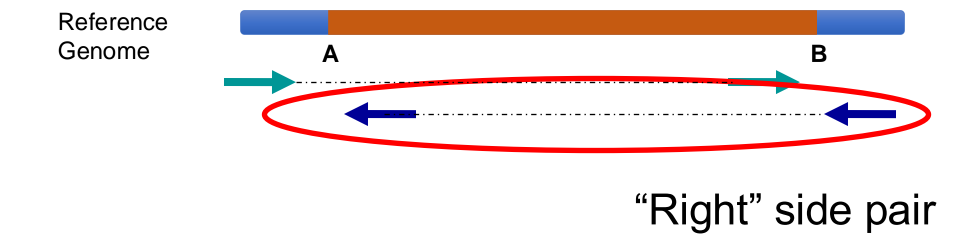




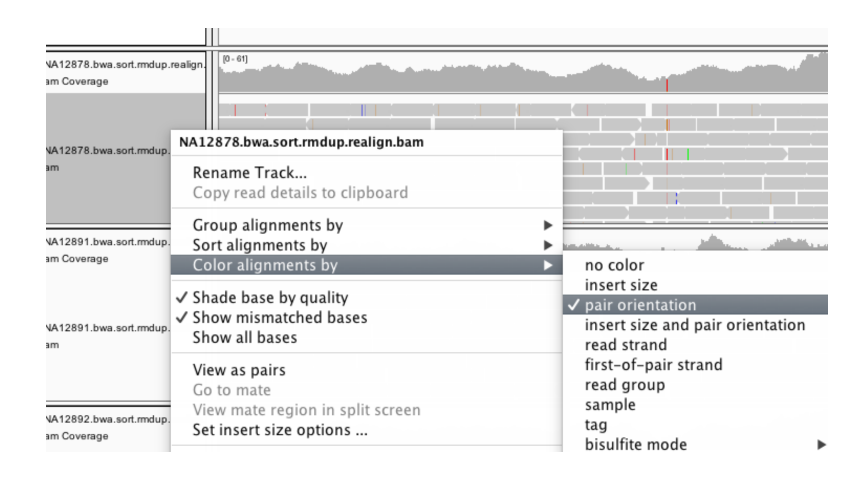
Anomaly: expected orientation of pair is inward facing (→ →)

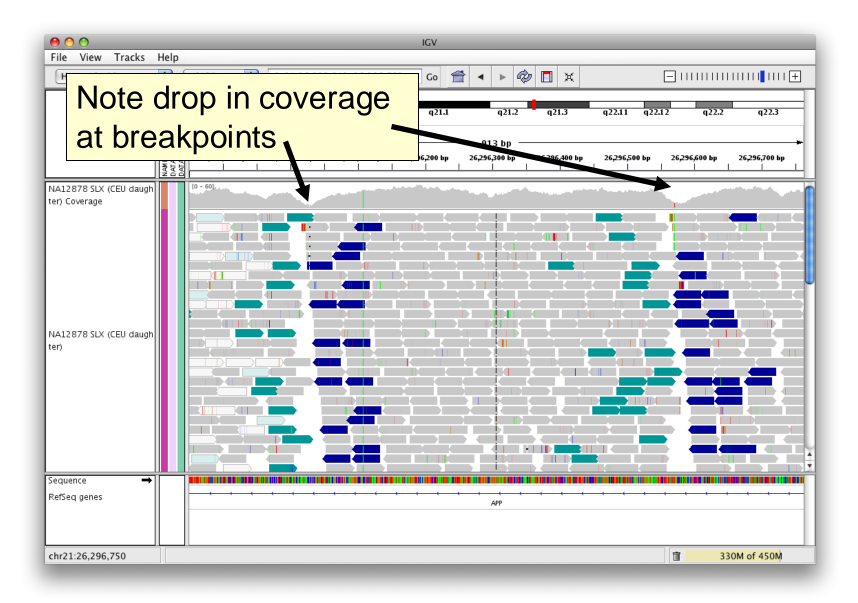


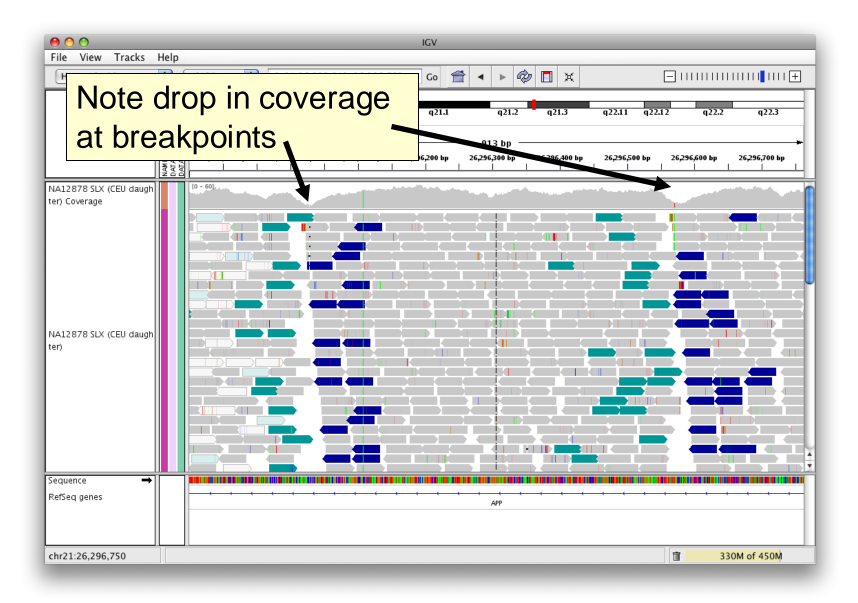
"Left" side pair



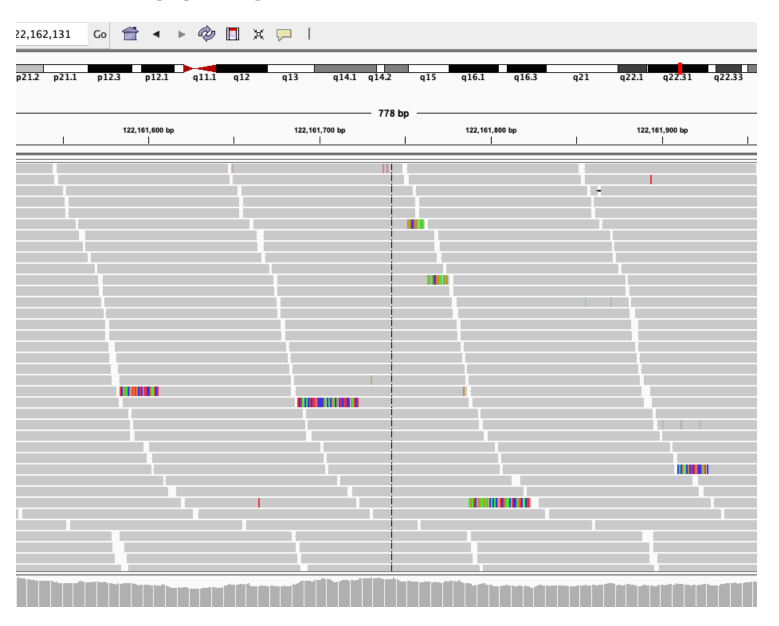
Color by pair orientation



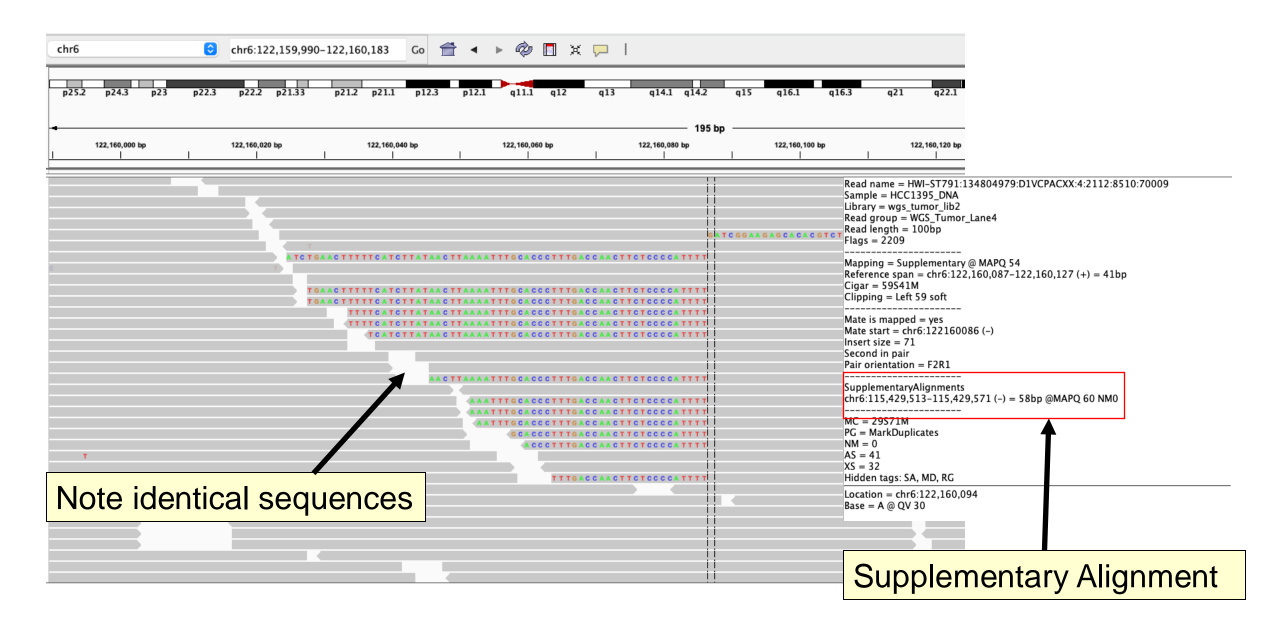




Soft-clipping



Soft-clipping



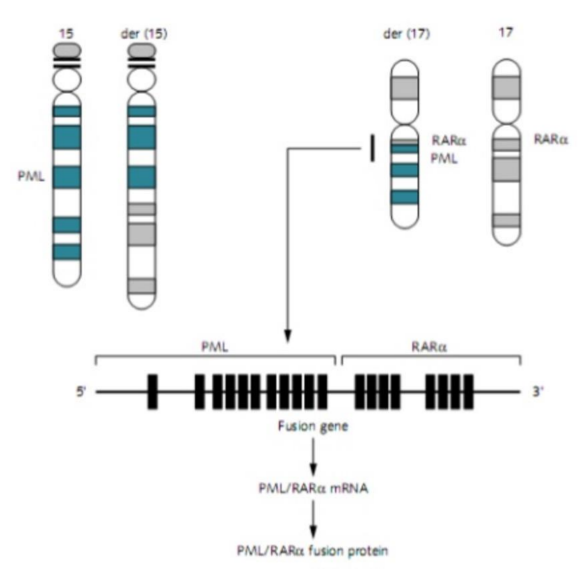
Assignment

https://gist.github.com/chrisamiller/1150bcdb1a269b6c32d1f2a77dccb9aa

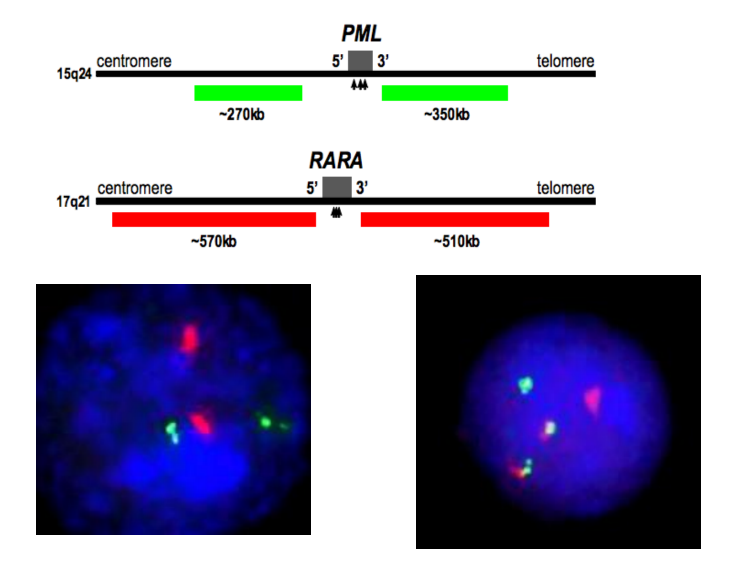


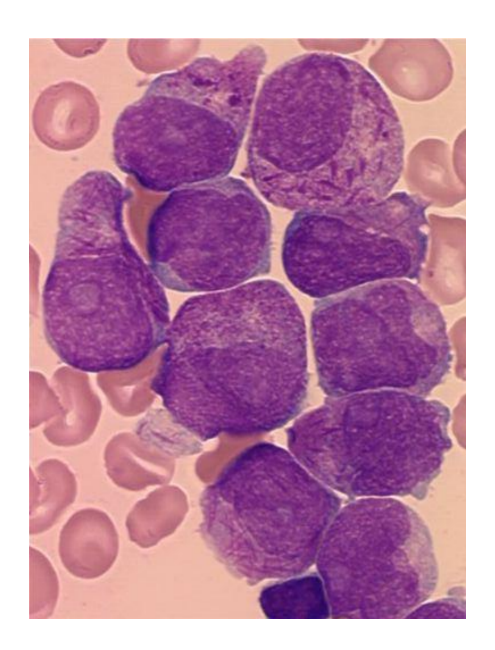
37 y.o. female with AML; M3 morphology

PML-RARA fusion



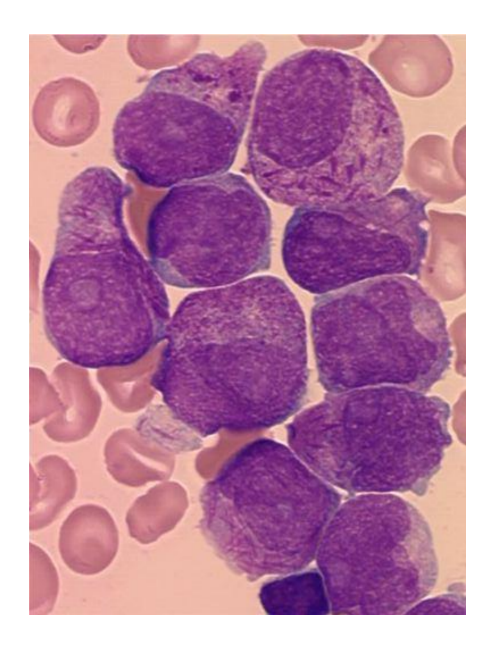
https://image.slidesharecdn.com/ sinhaematology2012-121128014851phpapp01/95/genetics-inhaematology2012-28-





37 y.o. female with AML; M3 morphology

Chemo + ATRA

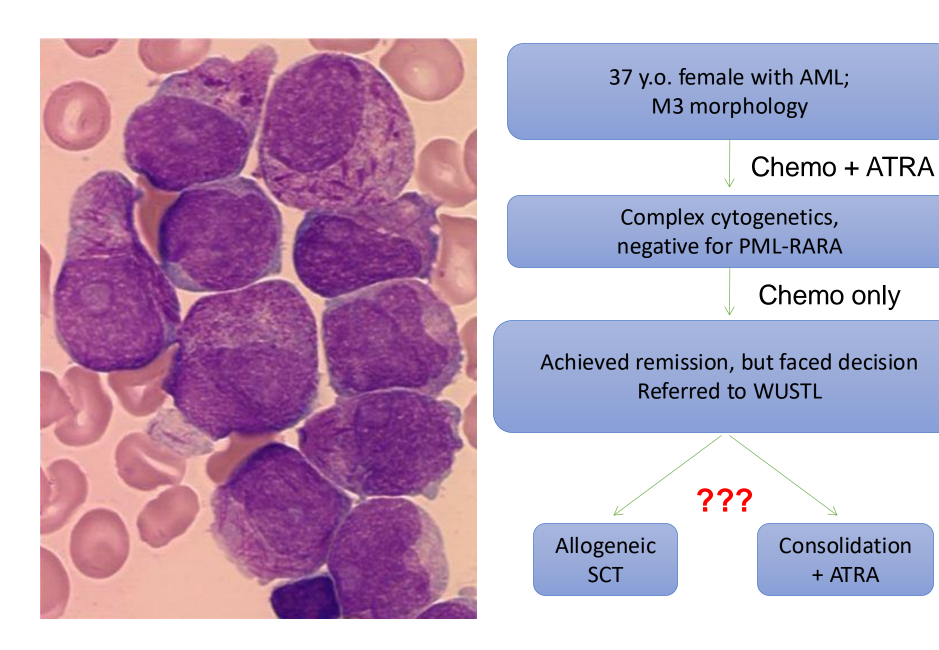


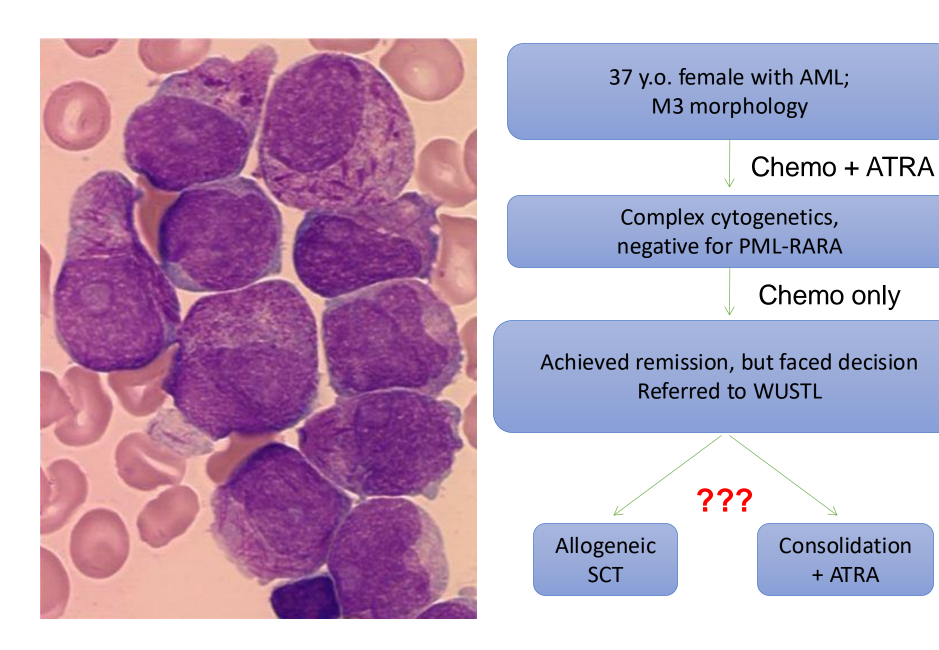
37 y.o. female with AML; M3 morphology

Chemo + ATRA

Complex cytogenetics, negative for PML-RARA

Chemo only



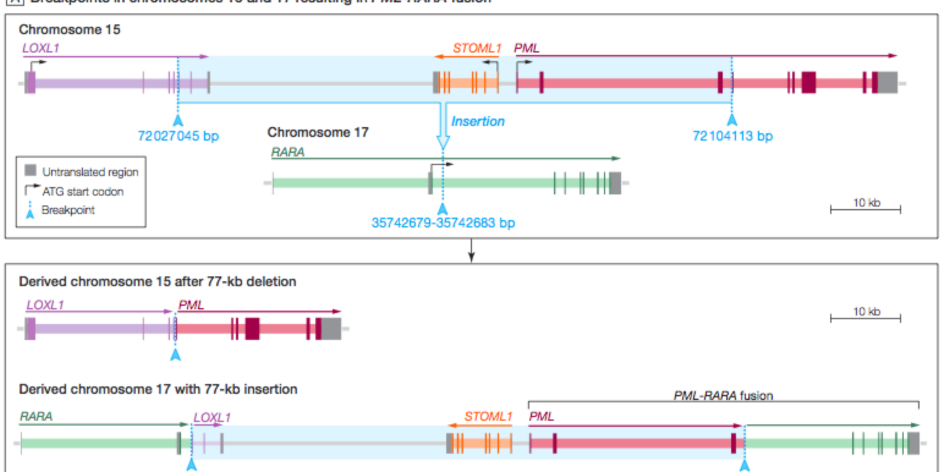




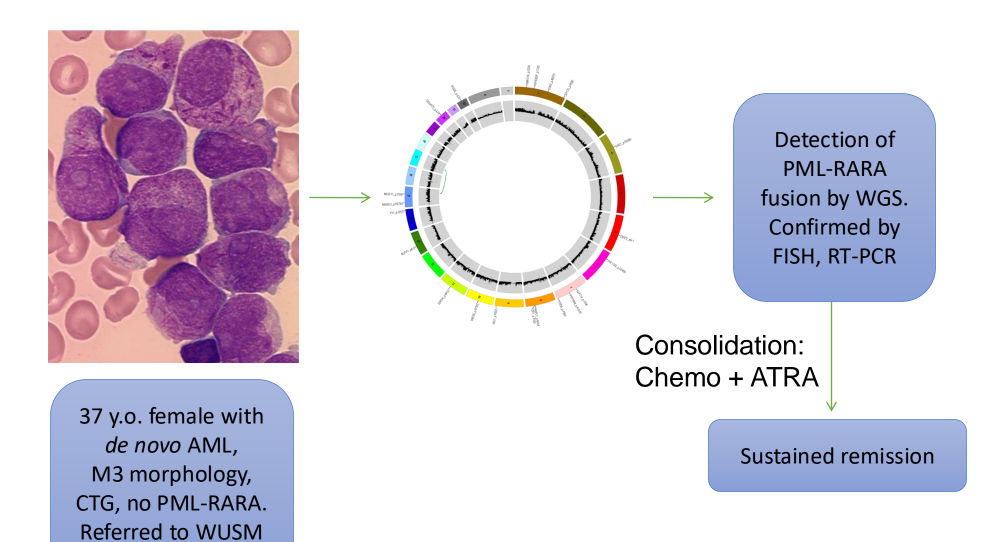
JAMA. 2011;305(15):1577-1584. doi: 10.1001/jama.2011.497

Use of Whole-Genome Sequencing to Diagnose a Cryptic Fusion Oncogene

A Breakpoints in chromosomes 15 and 17 resulting in PML-RARA fusion



for SCT.



Additional cryptic M3 AMLs

