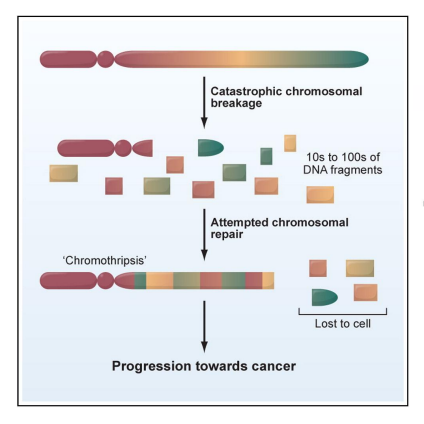
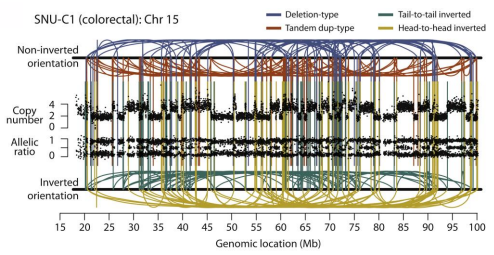
COMPLEX REARRANGEMENTS

Chromothripsis

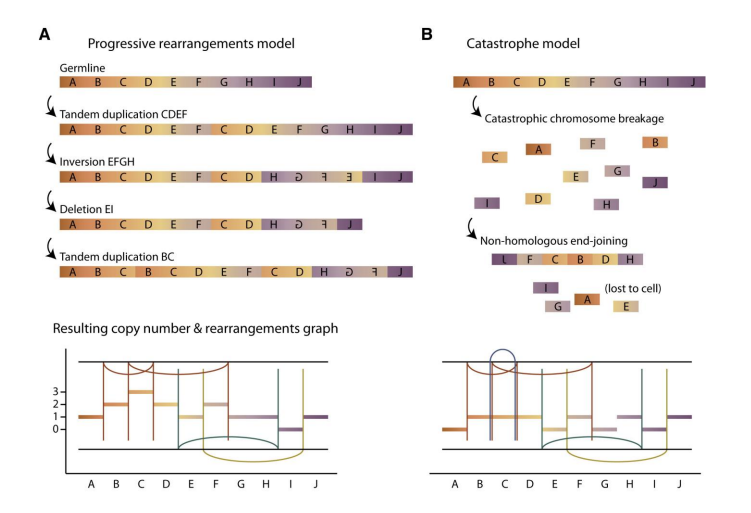




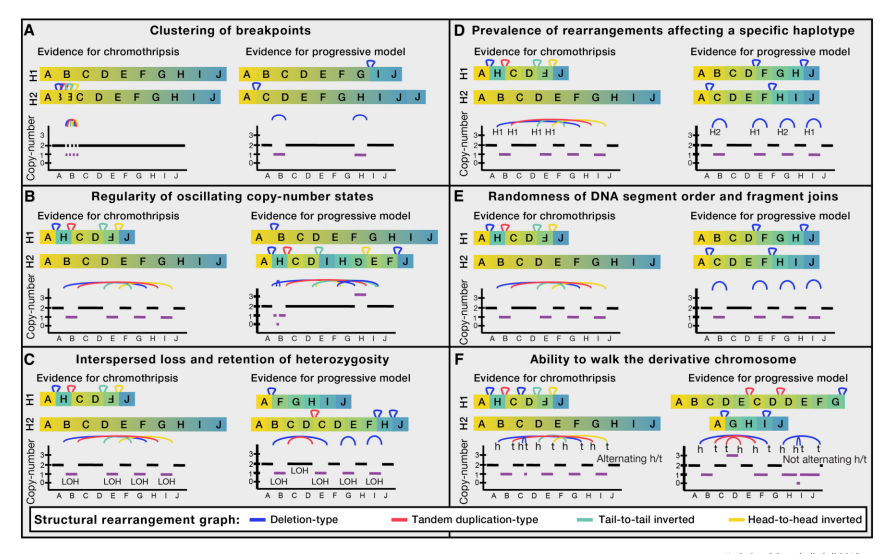
Stephens et al., Cell 2011

• prevalence in cancer varies (2-3% - 70% of cases in different cancers)

Chromothripsis



Criteria for inferring chromothripsis



Criteria for inferring chromothripsis: computational application

1. Clustering of breakpoints

Kolmogorov-Smirnov test for exponential distribution of breakpoint distances

2. Regularity of oscillating copy number states:

Calculated as percentage of consecutive 2-1/1-2 copy number steps in a chromosome

3. Interspersed loss and retention of heterozygosity

Calculated as percentage of consecutive retention/loss of fragments in a chromosome

4. Randomness of DNA segment order

Compare breakpoint distances with Monte Carlo simulations (t-test)

5. Randomness of DNA fragment joins

DEL, TanDUP, H2H, T2T-type rearrangement counts should follow a multinomial distribution (p=1/4)

6. Ability to walk the derivative chromosome

Alternating heads and tails (Wald-Wolfowitz test)

7. Prevalence of rearrangements affecting a specific haplotype

Chromosome-wide phasing data can be obtained when germline whole-genomic sequencing data from both parents or somatic genome sequencing data from aneuploid secondary tumors (which are common in the context of hereditary disorders such as Li-Fraumeni syndrome; Li and Fraumeni, 1969) are available for a patient sample in question. (Korbel and Campbell, 2013)

Criteria for inferring chromothripsis: in practice

No clear-cut rules:

> Stephens et al., 2011:

- 1. massive number of rearrangements on 1 or a few chromosomes (>10)
- 2. alternate copy number between 2 states only and alternate loss/retention of heterozygosity
- 3. clustering of breakpoints

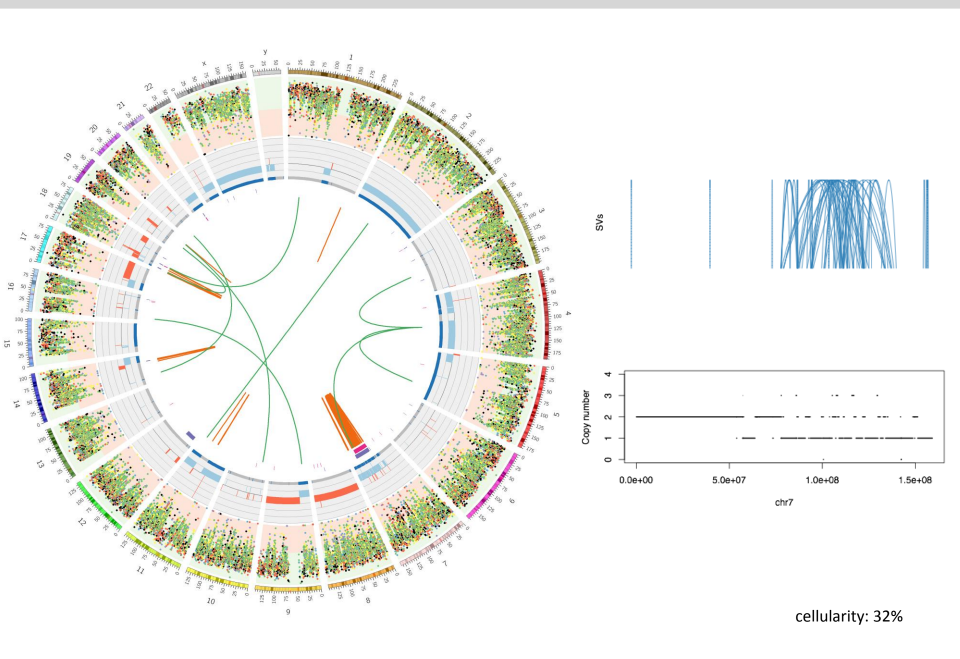
> Rausch et al., 2012:

10 changes in segmental copy number involving 2-3 distinct copy number states on a single chromosome

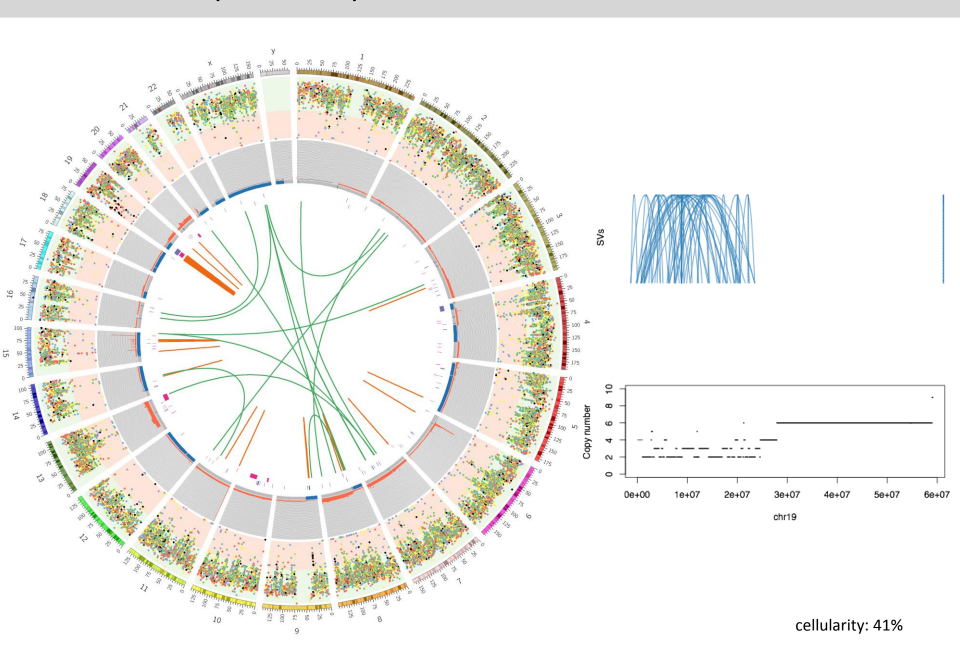
> Nones et al., 2014:

Evidence of clustering of breakpoints was estimated as proposed by Korbel and Campbell36. Chromosomes with evidence of clustering of breakpoints (P<0.001, Kolmogorov–Smirnov test—goodness of fit test) were reviewed for: (1) evidence of chromothripsis which included <u>oscillation of copy number, random joins</u> and <u>retention of heterozygosity</u> [...] A larger cohort of EACs (n=101) was screened for evidence of chromothripsis using SNP arrays (Illumina), chromothripsis was inferred in cases where one or few chromosomes showed <u>at least 10 switches in copy number states</u>, with retention of heterozygosity.

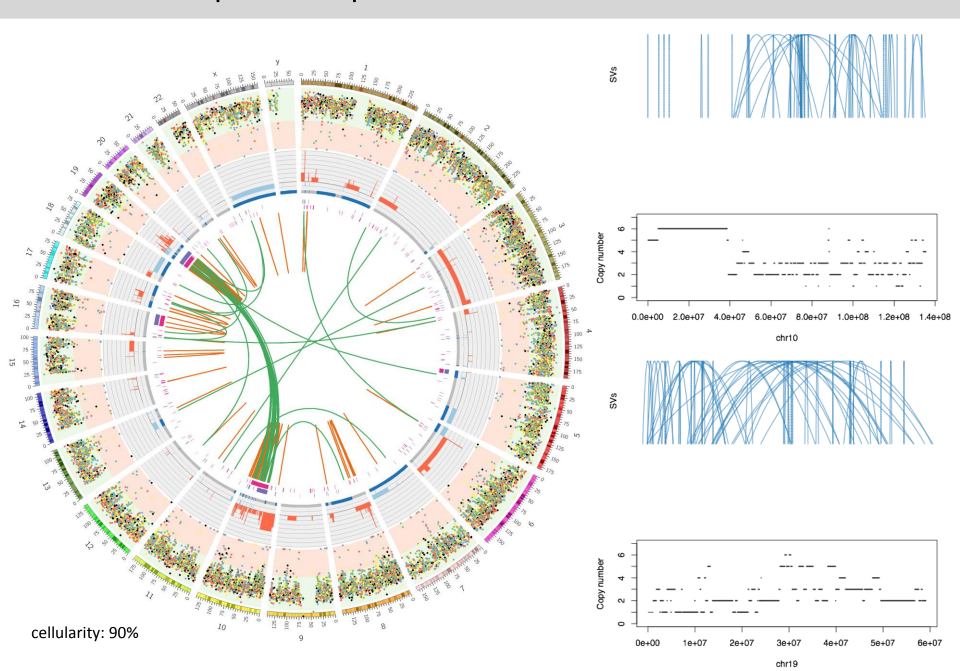
Chromothripsis example 1



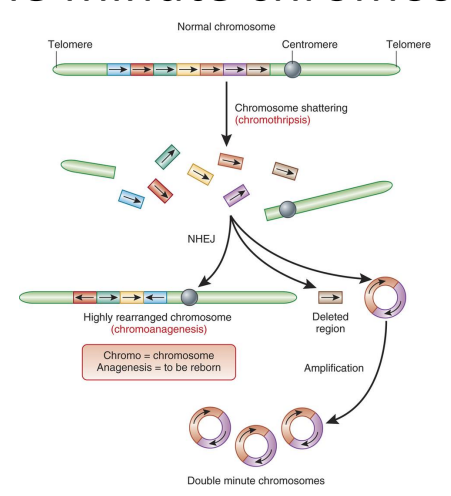
Chromothripsis example 2



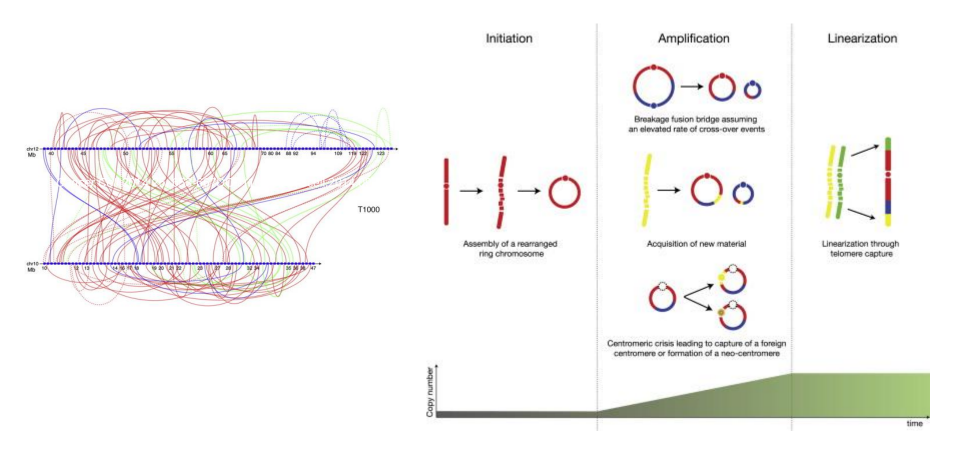
Chromothripsis example 3



Double minute chromosomes



Neochromosome characterisation



EXERCISE 5