

# **Genomic Architecture and Inheritance of Human Ribosomal RNA Gene Arrays**

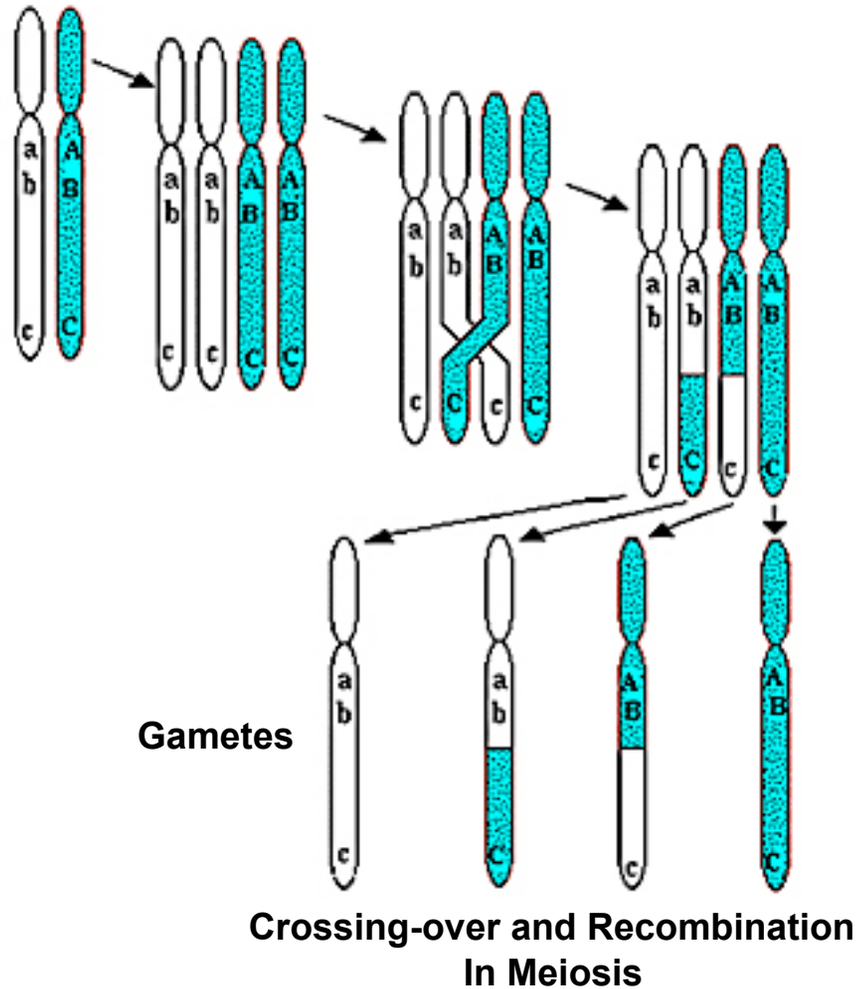
## **Meiotic and Mitotic Instability**

**Andrew J. Pierce**

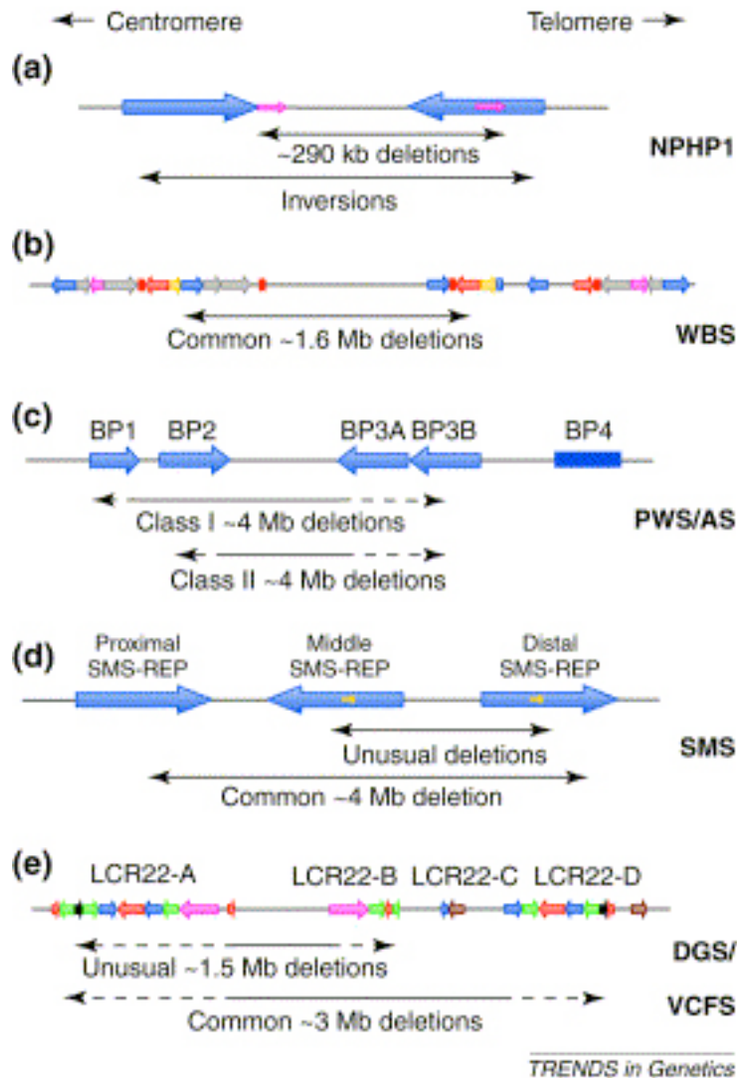
**Microbiology, Immunology and Molecular Genetics  
Graduate Center for Toxicology  
Markey Cancer Center  
University of Kentucky**

**MI615**

# Meiosis Requires Genomic Restructuring



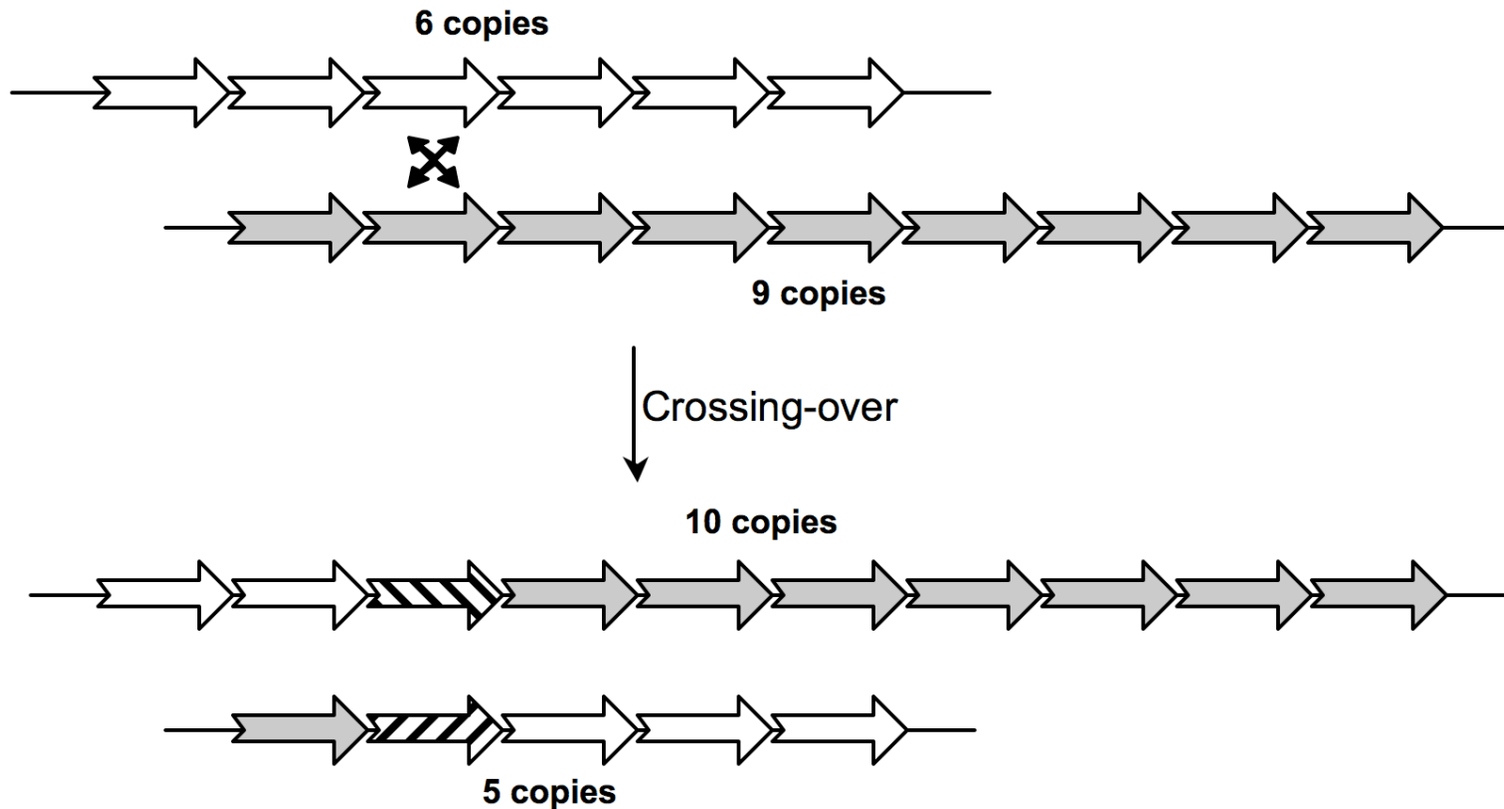
# Some Genomic Disorders Mediated by Repetitive Sequences



Complex structure of selected low-copy repeats (LCRs). Horizontal lines represent specific genomic regions with the centromere toward the left and telomere to the right. At the right are listed abbreviations for the disease manifested through common deletions of the regions. The colored regions refer to LCRs with the orientation given by the arrowhead. Note complex structure of LCRs consisting of both direct and inverted repeats. (a) LCRs in chromosome 2q13 responsible for rearrangements associated with familial juvenile nephronophthisis 1 (NPHP1). (b) LCRs7 flanking the Williams–Beuren syndrome (WBS) chromosome region 7q11.23. (c) LCRs15 within the Prader–Willi syndrome/Angelman syndrome (PWS/AS) chromosome region 15q11.2. (d) Smith–Magenis syndrome (SMS) repeats within 17p11.2. (e) LCRs22 within the DiGeorge syndrome (DGS) chromosome 22q11.2.

# Instability Mechanism: Homologous Recombination

Figure 1. Intermolecular Crossing-over



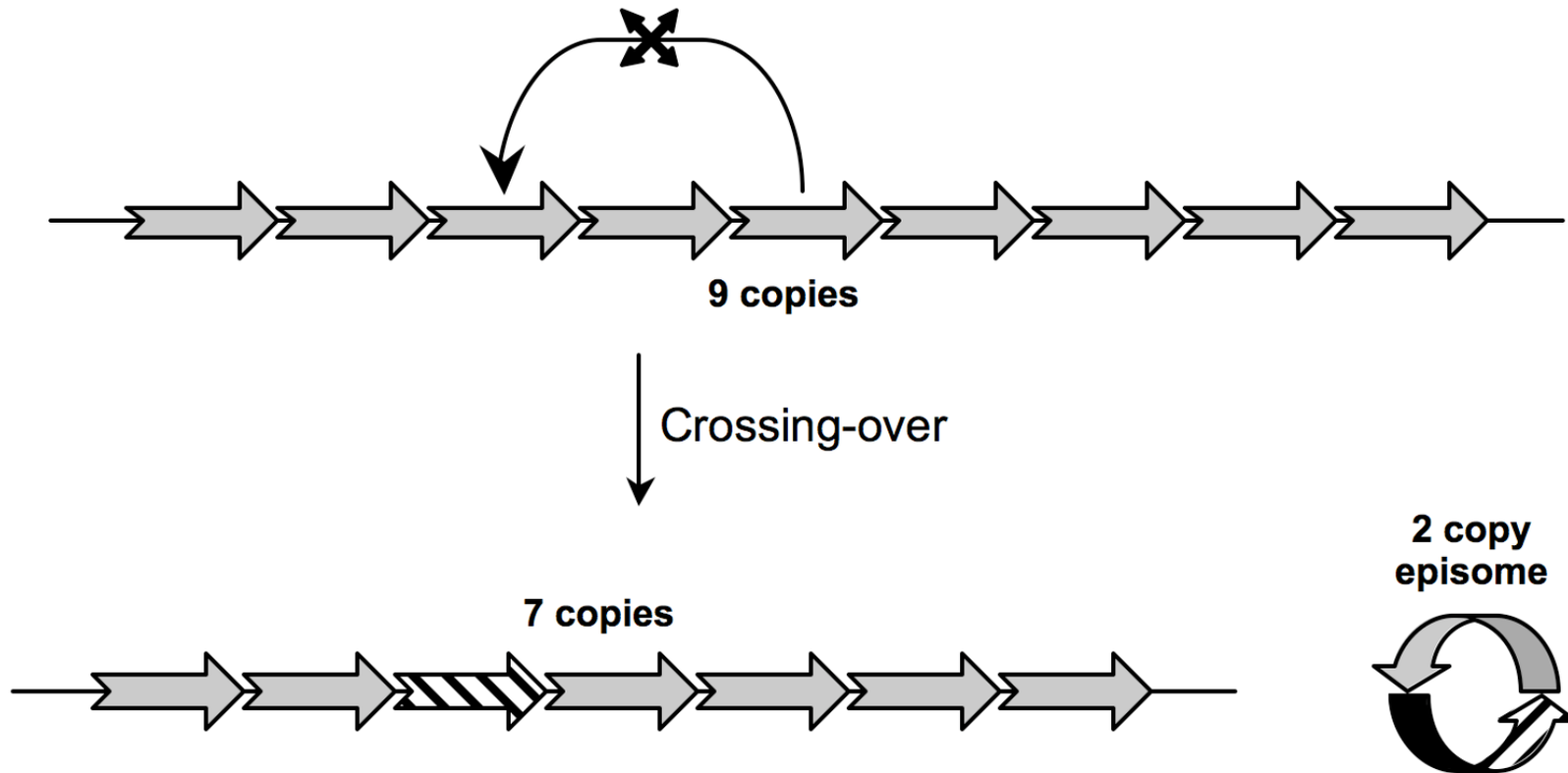
white arrows and gray arrows: unit repeated elements  
(arrow direction indicates relative orientation)

crossed arrows: site of crossing-over

hatched arrows: repeats in which crossover exchange has occurred

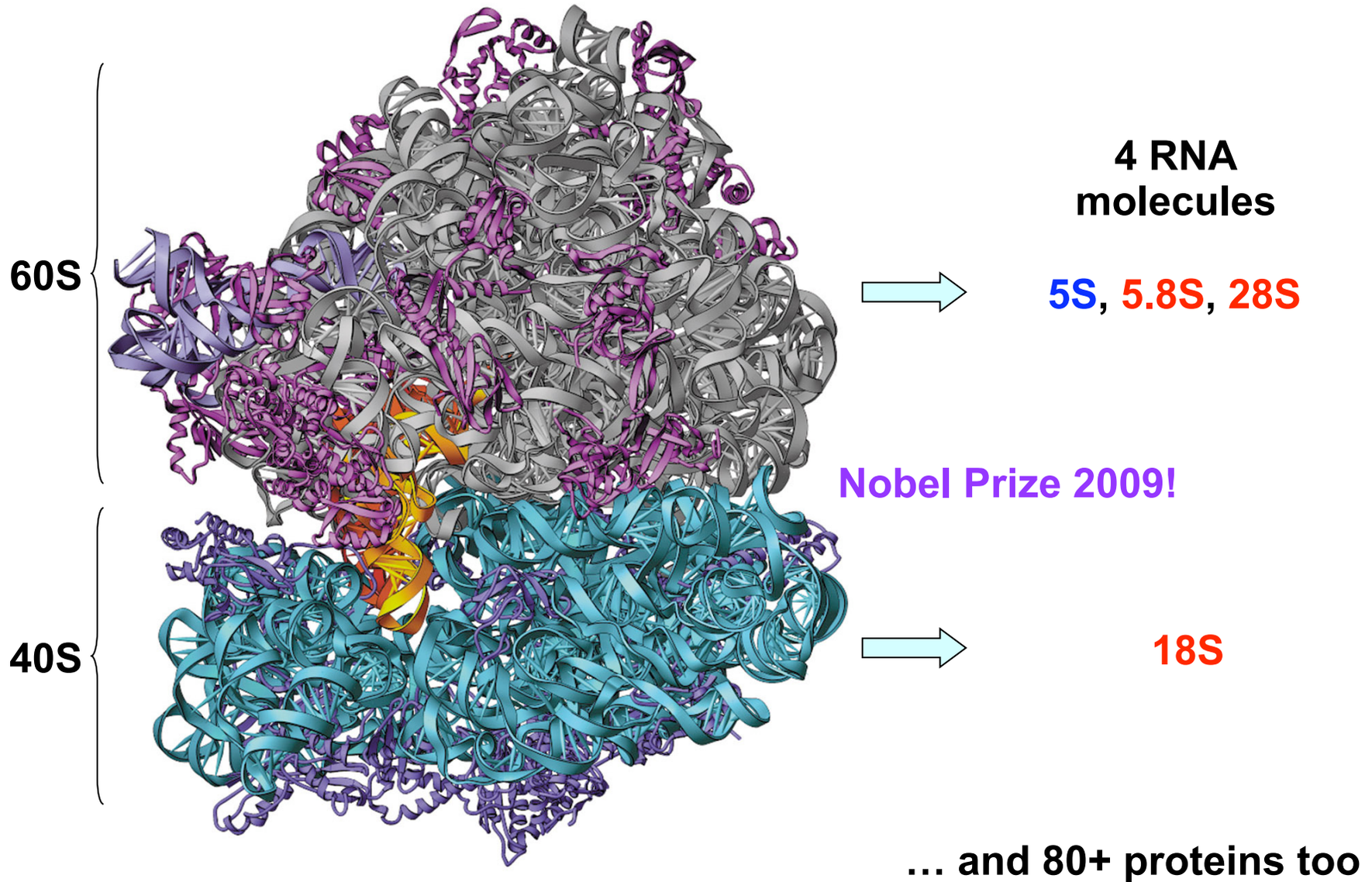
# Instability Mechanism: Homologous Recombination

Figure 2. Intramolecular Crossing-over



gray arrows: unit repeated elements  
(arrow direction indicates relative orientation)  
crossed arrows: site of crossing-over  
hatched arrows: repeats in which crossover exchange has occurred

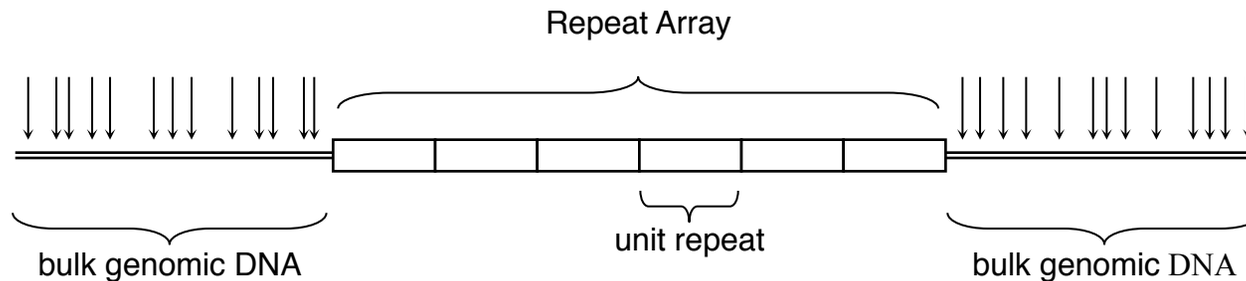
# Ribosomal Composition





# Experimental Strategy

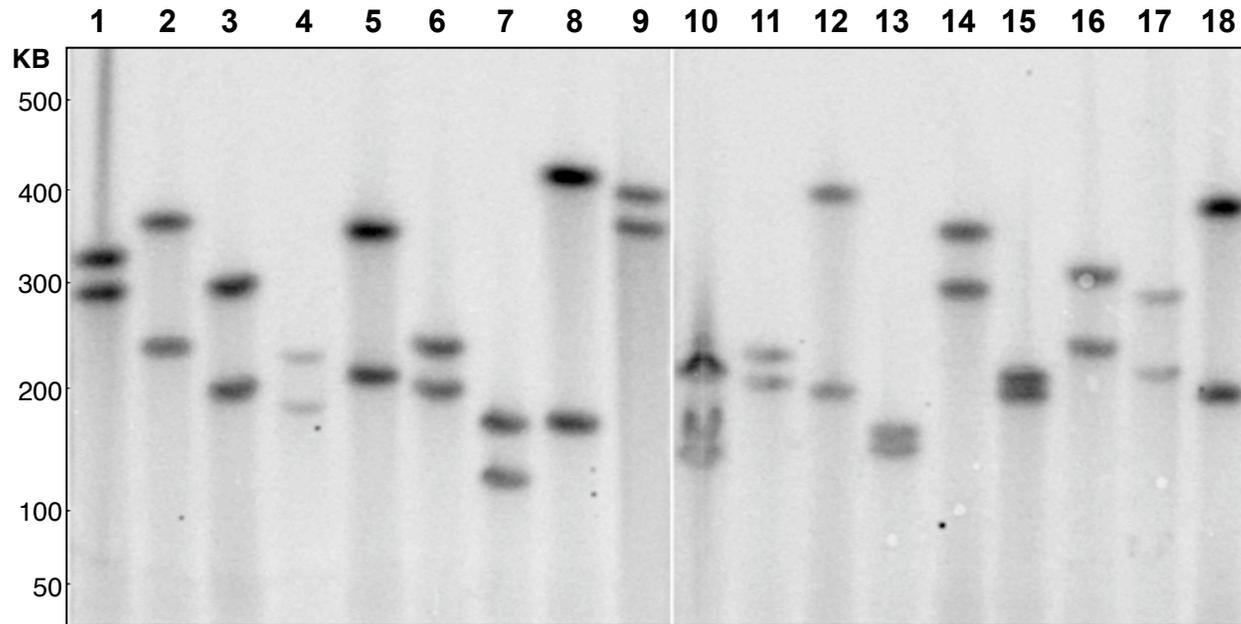
1. Isolate very high molecular weight genomic DNA from human peripheral blood
2. Liberate rDNA arrays by digesting with restriction enzymes that cut external to the array
3. Separate arrays by size on pulsed-field gels
4. Detect rDNA specific bands by Southern blotting



**Band intensity is proportional to:**

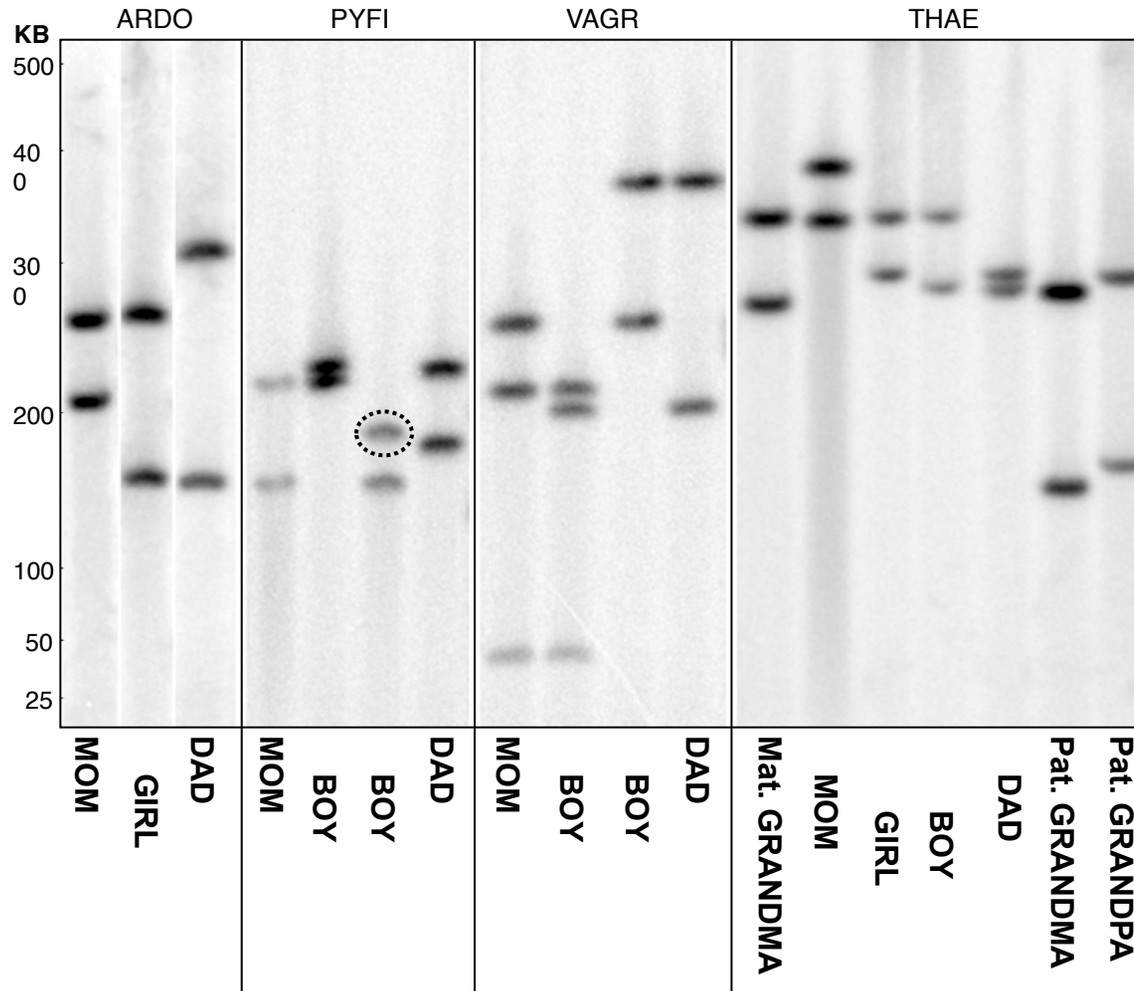
- i. Length of the array
- ii. Fraction of cells in the population containing any given sized array

# 5S rDNA Arrays from Anonymous Human Donors



- Everyone has two different sized arrays
- No two people have the same array lengths

# 5S rDNA Arrays from Human Families – Inheritance



Sample Source:		Mother	(Other)	Child	(Focus)	Father	(Alleged)	Exclusion Status*	Direct Index	Column Not Used
Name:		PYFI-10-MA		PYFI-02-TY		PYFI-20-BR				
Racial Database Used:		Caucasian-American		Caucasian-American		Caucasian-American				
Date Collected:		Unknown		Unknown		Unknown				
STR Locus	Allele Range	Alleles Called		Alleles Called		Alleles Called				
D3S1358	(12 - 20)	14	15	15	16	15	16	OK	2.072	---
TH01	(4 - 13.3)	6	10	6	9.3	9.3		OK	2.820	---
D21S11	(24 - 38)	32.2		29	32.2	29	32.2	OK	1.933	---
D18S51	(8 - 27)	12	16	12	13	13	16	OK	3.510	---
Penta E	(5 - 24)	7	12	5	12	5	12	OK	7.478	---
D5S818	(7 - 16)	11	12	11	12	12		OK	1.357	---
D13S317	(7 - 15)	11	12	12		10	12	OK	1.552	---
D7S820	(6 - 14)	10	13	12	13	11	12	OK	4.719	---
D16S539	(5 - 15)	11	13	9	13	9	11	OK	5.333	---
CSF1PO	(6 - 15)	10	11	10	12	11	12	OK	1.667	---
Penta D	(2.2 - 17)	9	10	9		9	13	OK	2.915	---
Amelogenin	(XX - XY)	Female (XX)		Male (XY)		Male (XY)		N/A	N/A	---
vWA	(10 - 22)	15	19	17	19	17		OK	3.800	---
D8S1179	(7 - 18)	12	15	12	14	14		OK	5.375	---
TPOX	(6 - 13)	8	11	8	11	8		OK	1.265	---
FGA	(16 - 46.2)	20	23	20	21	21	23	OK	2.730	---
NOT USED										
NOT USED										
NOT USED										
NOT USED										
NOT USED										
NOT USED										
NOT USED										
NOT USED										

Laboratory Batch Number: 63613130C1 Notable Events: Exclusions-None; Infrequent Events-None. See COMMENTS section for additional information.

**Statement of Results:** Alleged relationship is not excluded.

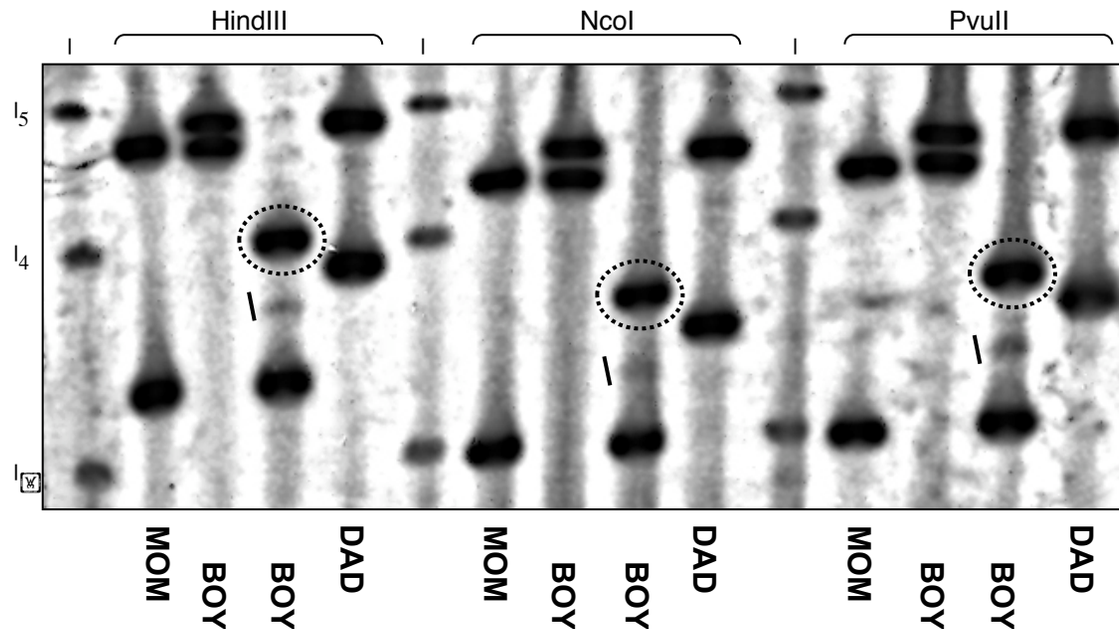
Based on the DNA analysis, the alleged Father, PYFI-20-BR, cannot be excluded as the biological Father of the Child, PYFI-02-TY, because they share genetic markers. Of the genetic identity systems tested, 15 of 15 match. (99.999945964304% of the Caucasian-American male population is excluded from the possibility of being the biological Father). The probability of the stated relationship is indicated below, as compared with an untested, unrelated Caucasian-American male. Analyses, with the exception of sample collection, were conducted in accordance with the Standards for PCR DNA analysis set forth by the AABB.

**Statistical Results:**

Combined Direct Index 5,383,421 .....Probability = 99.999981424455%

Prior Probability 0.50 .....Statistical Constant

# 5S rDNA Arrays – Inheritance and Somatic Mosaicism



**Meiotic rearrangement: Band length not represented in either parent ( dotted circle )**

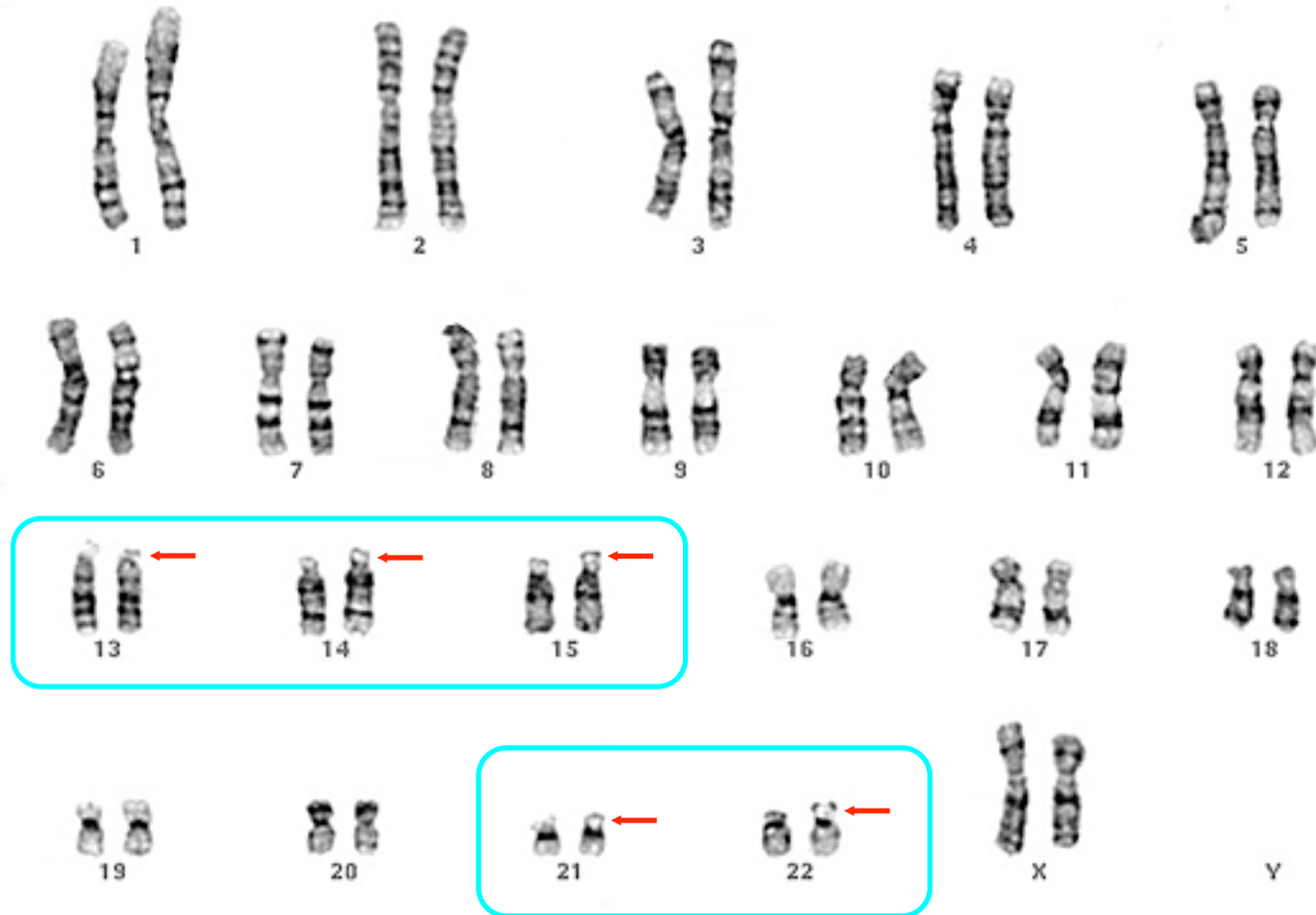
**Somatic mosaicism: Band of reduced intensity relative to length ( square bracket )**

## 5S rDNA Arrays – Human Statistics ( N = 27 )

	<b>Each Haploid Array</b>	<b>Diploid Totals</b>
<b>Average number of repeats</b>	<b>98 ± 35</b>	<b>195 ± 51</b>
<b>Fewest repeats observed</b>	<b>35</b>	<b>51</b>
<b>Most repeats observed</b>	<b>175</b>	<b>299</b>

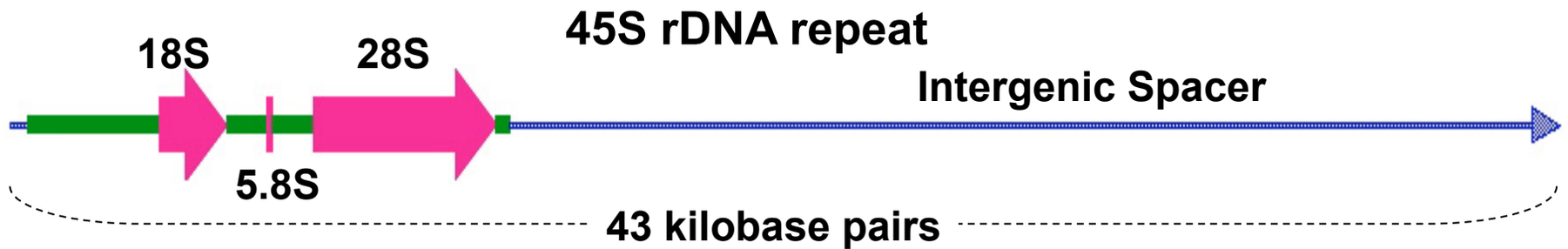
# 45S Ribosomal DNA in Human Karyotypes

Human female  
G-bands

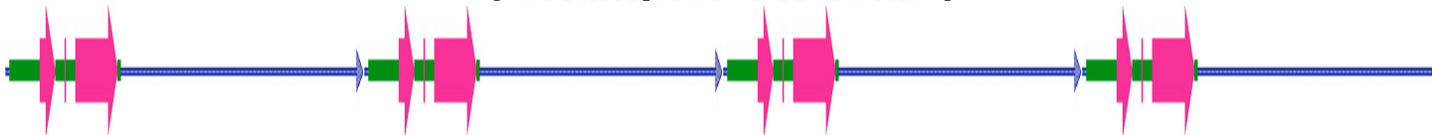


# Ribosomal DNA Repeats

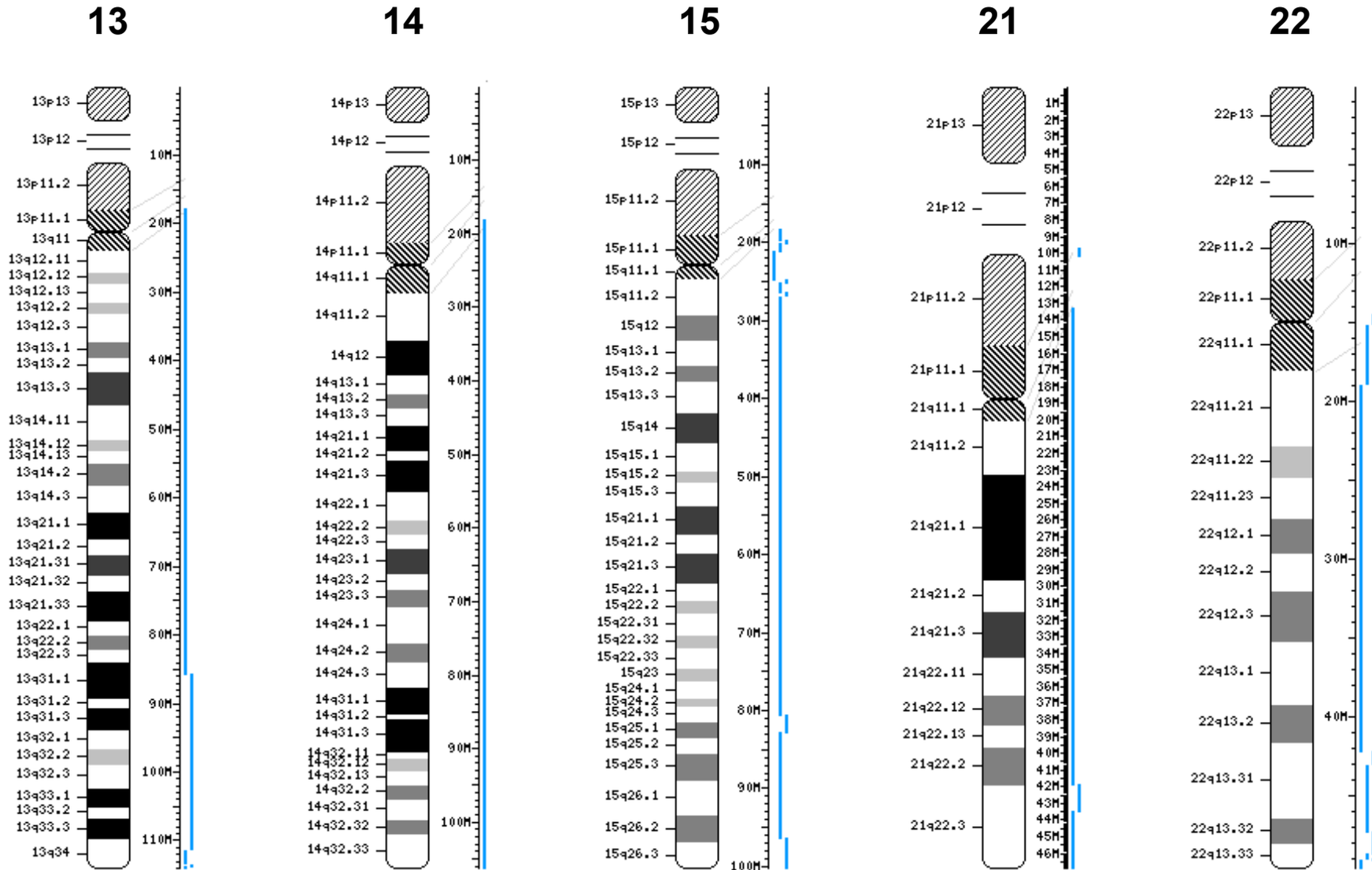
- generally structured in tandem head-to-tail arrays
- unit repeats completely sequenced



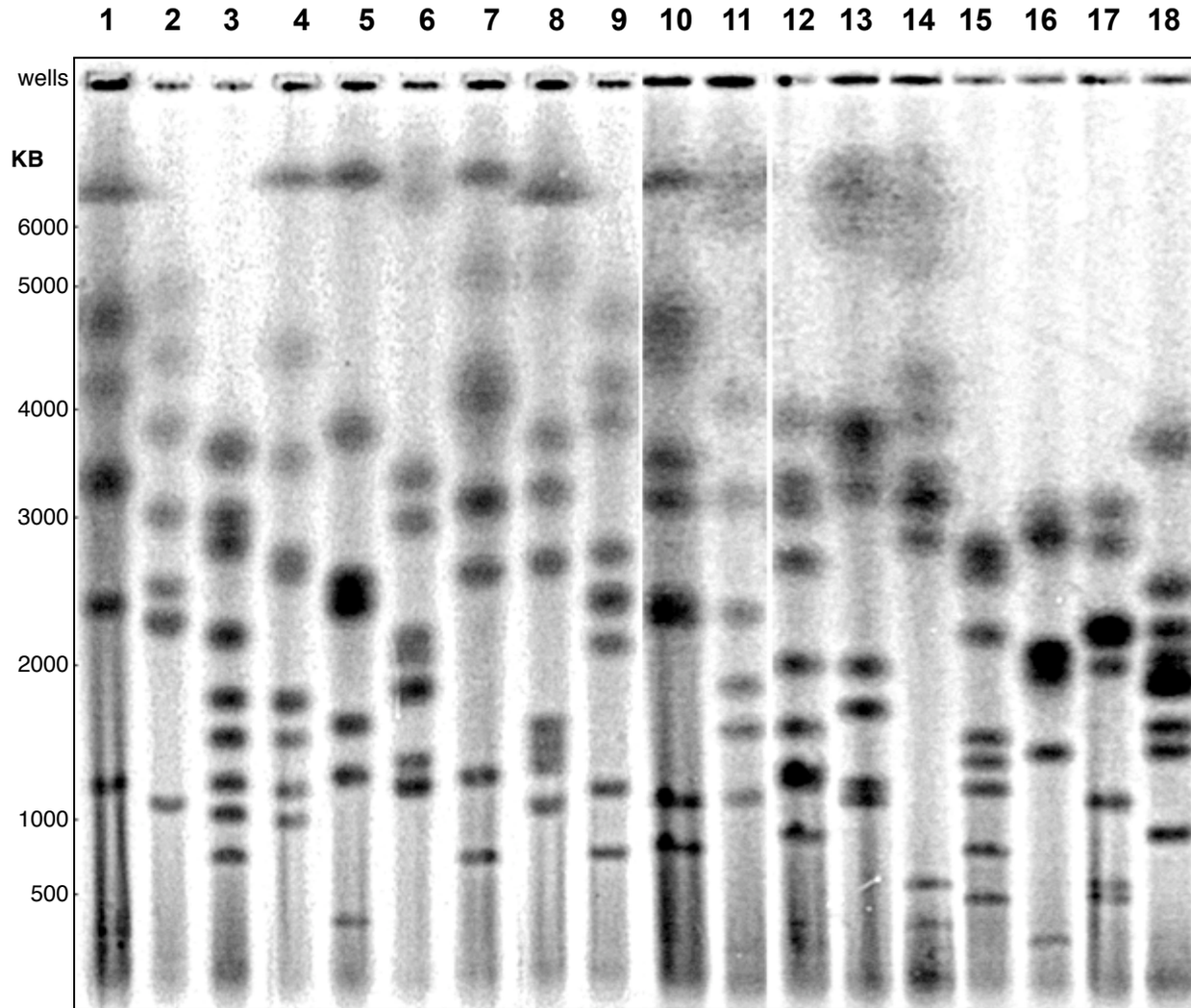
**Tandem Array of Repeats**  
( example: 4 x 43kb )



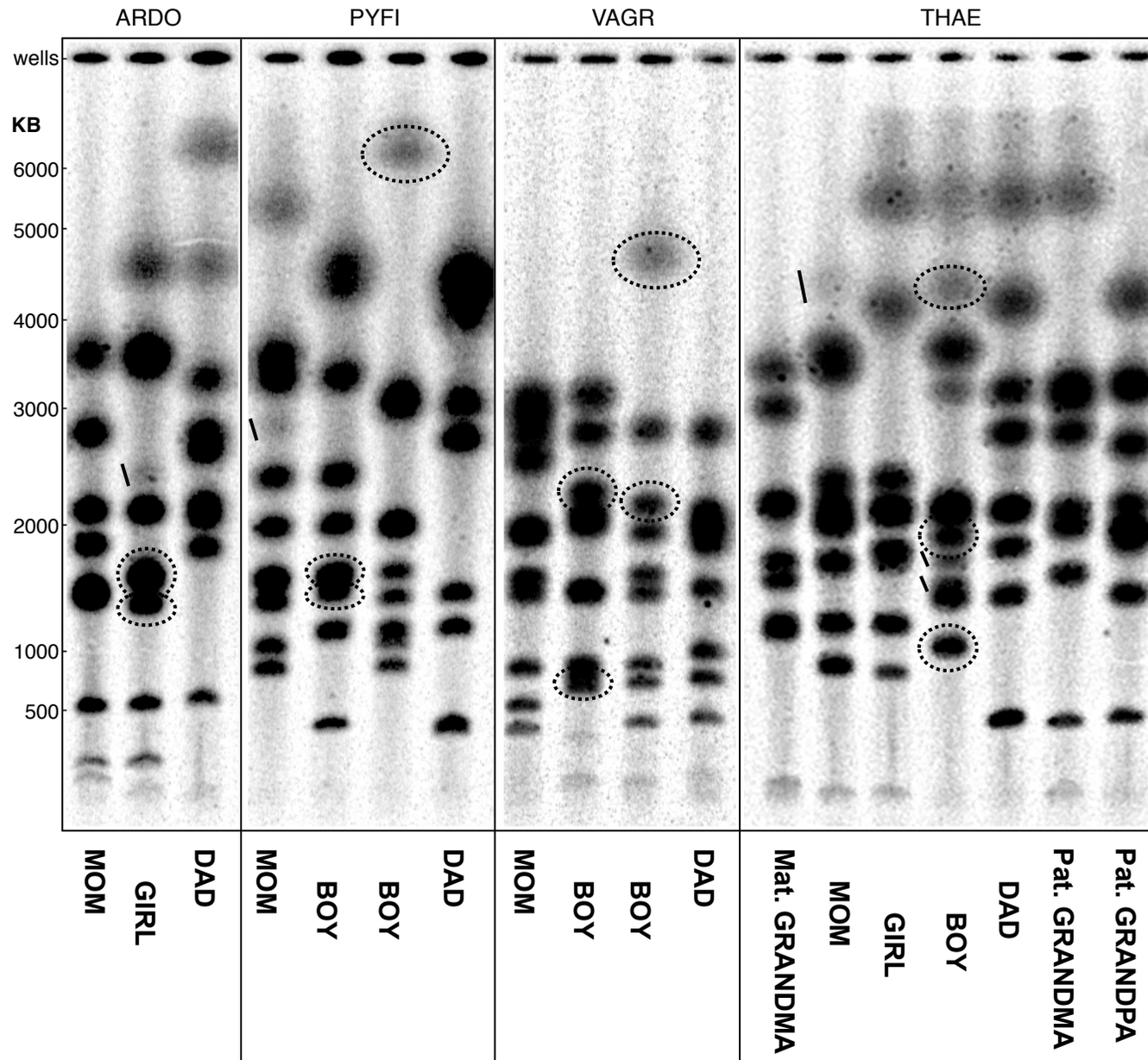
# 45S rDNA Arrays and the Human Genome Project



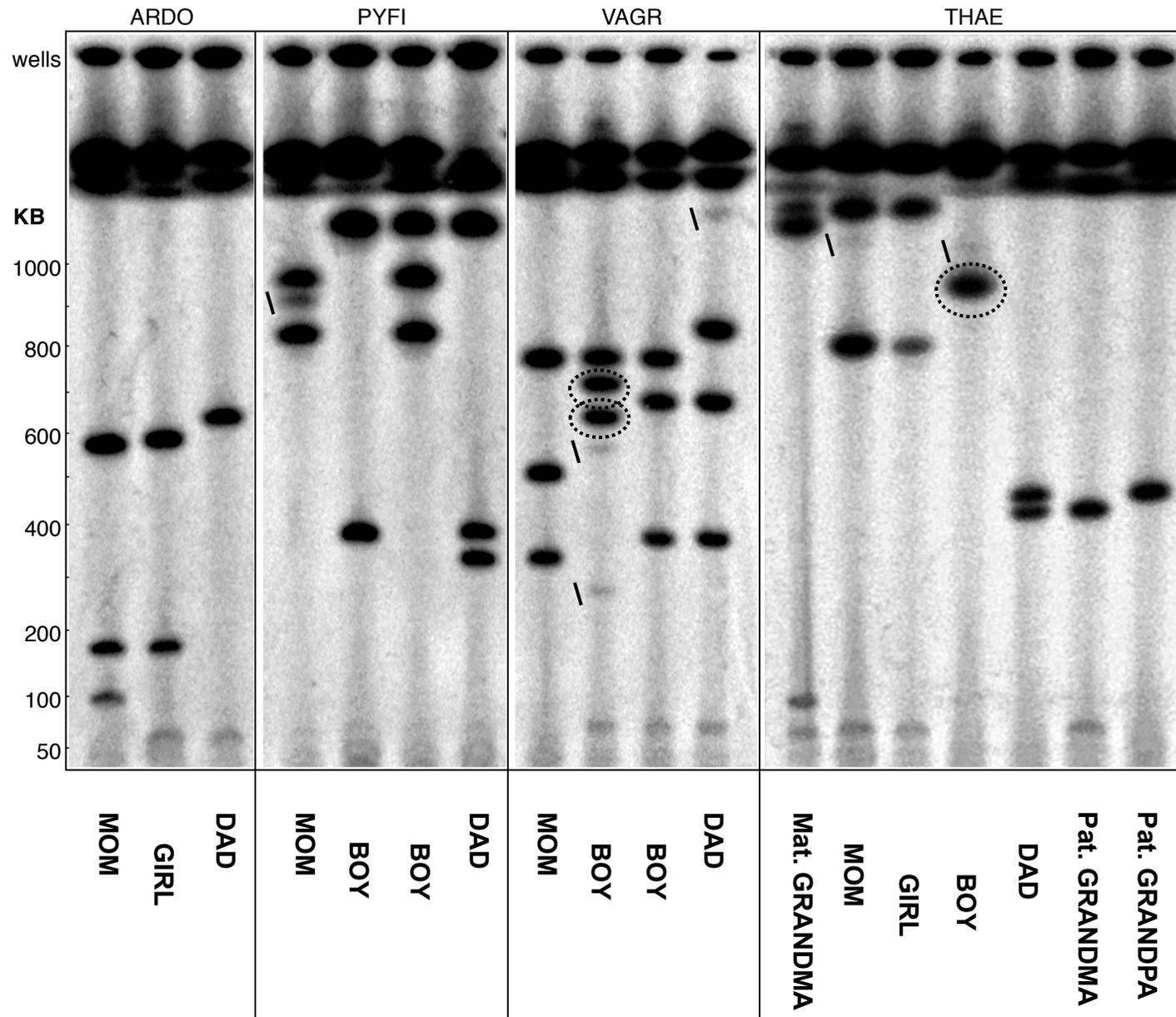
# 45S rDNA Arrays from Anonymous Human Donors



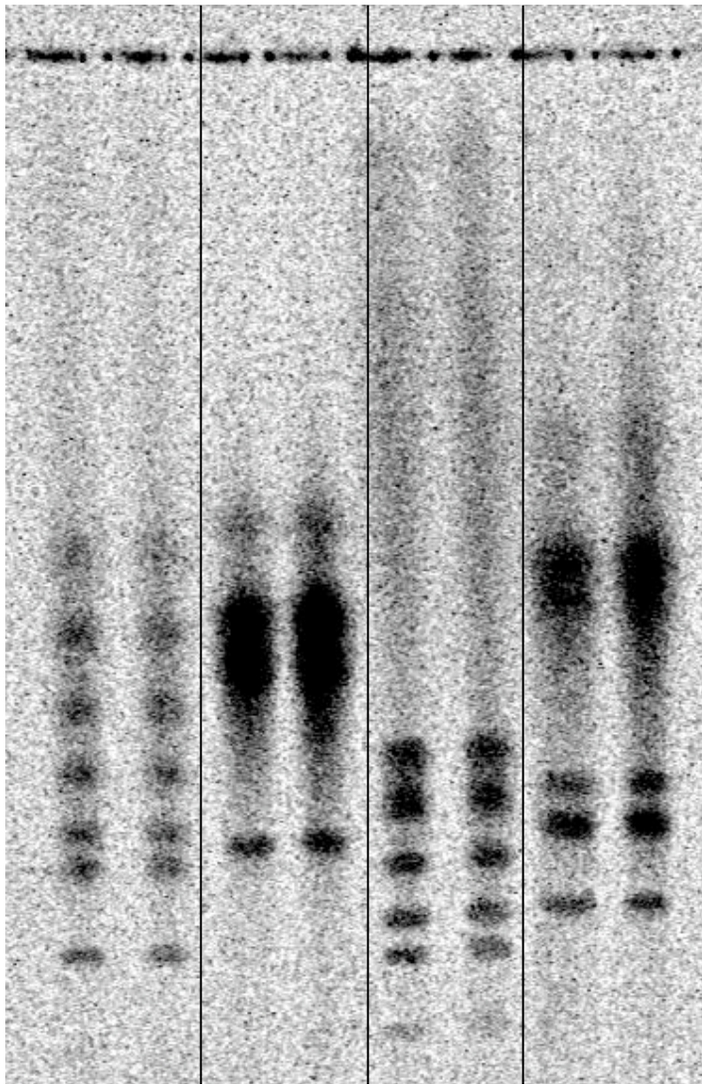
# 45S rDNA Arrays from Human Families – Inheritance



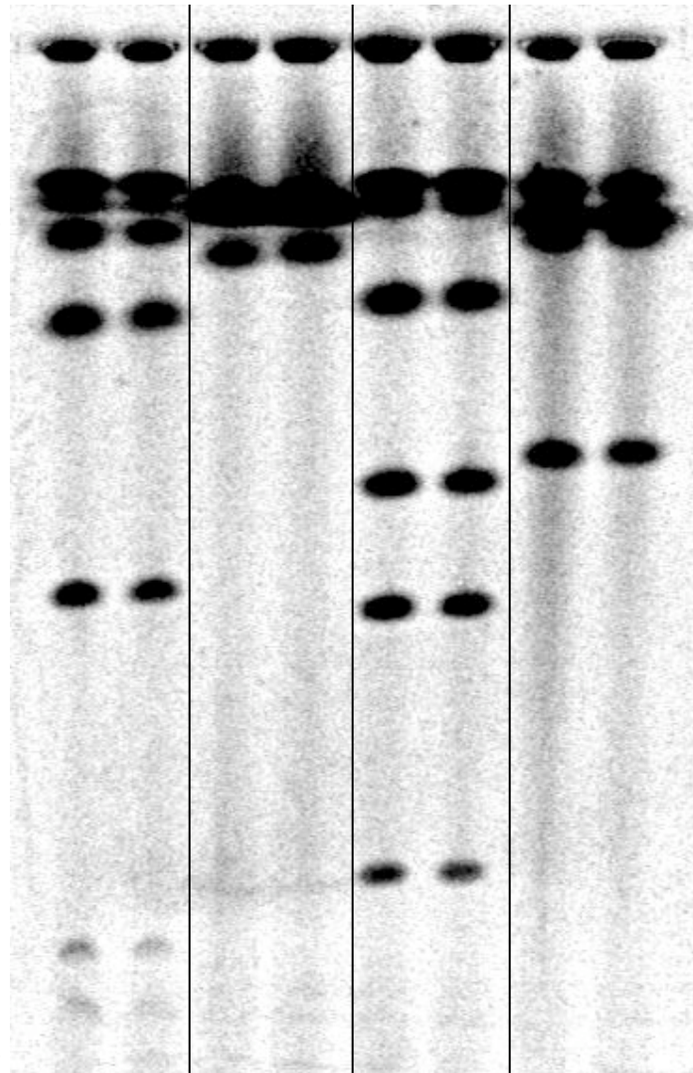
# High Resolution 45S Arrays – Inheritance and Mosaicism



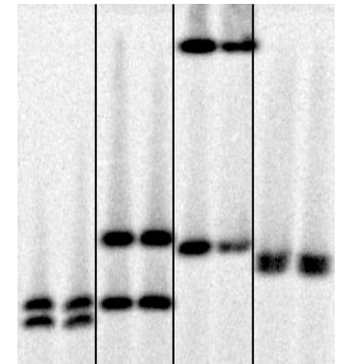
# Four Sets of Identical Twins



**45S rDNA 1Mbp - 6Mbp**



**45S rDNA 50kb - 1Mbp**

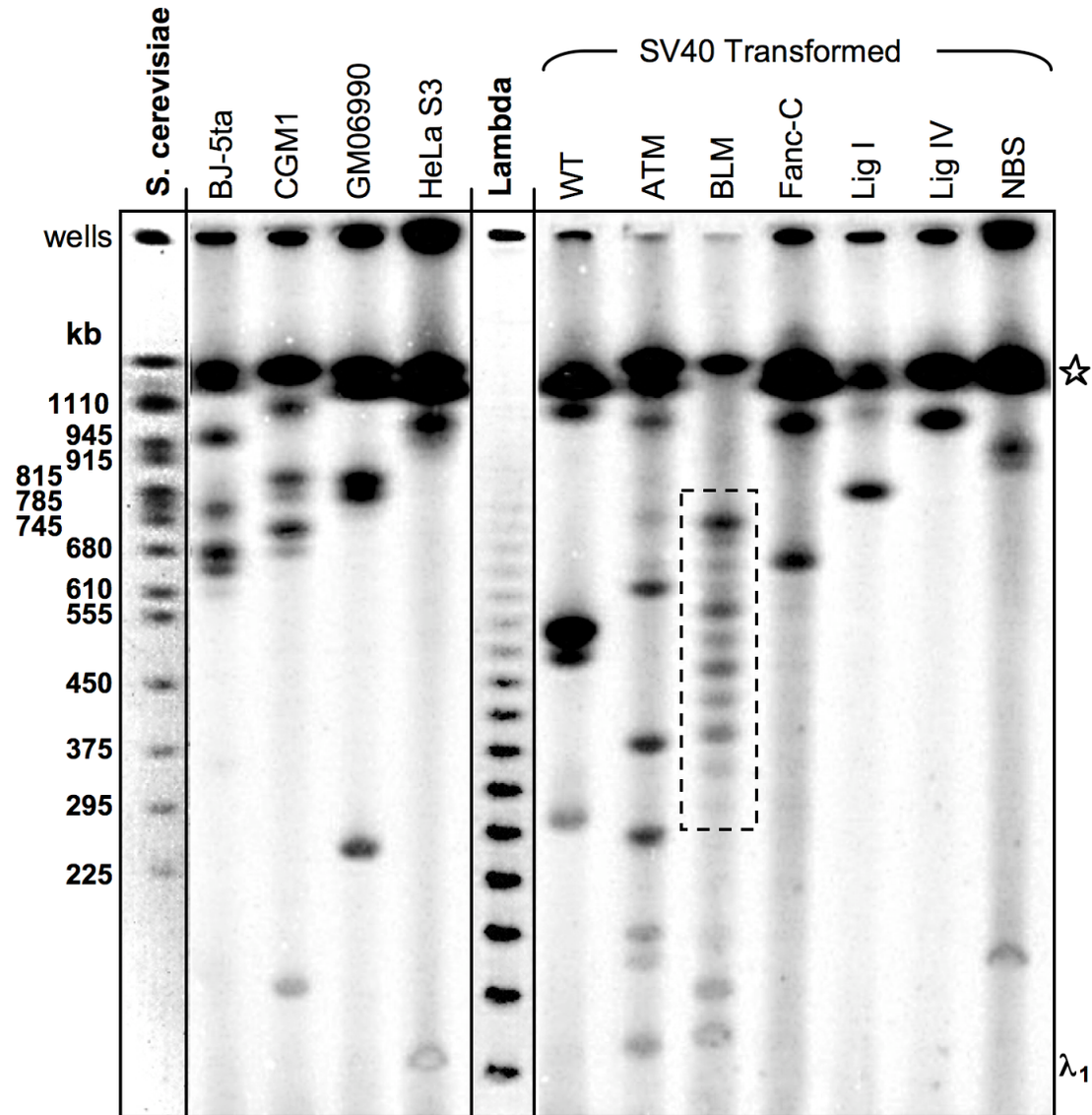


**5S rDNA**

## Conclusions

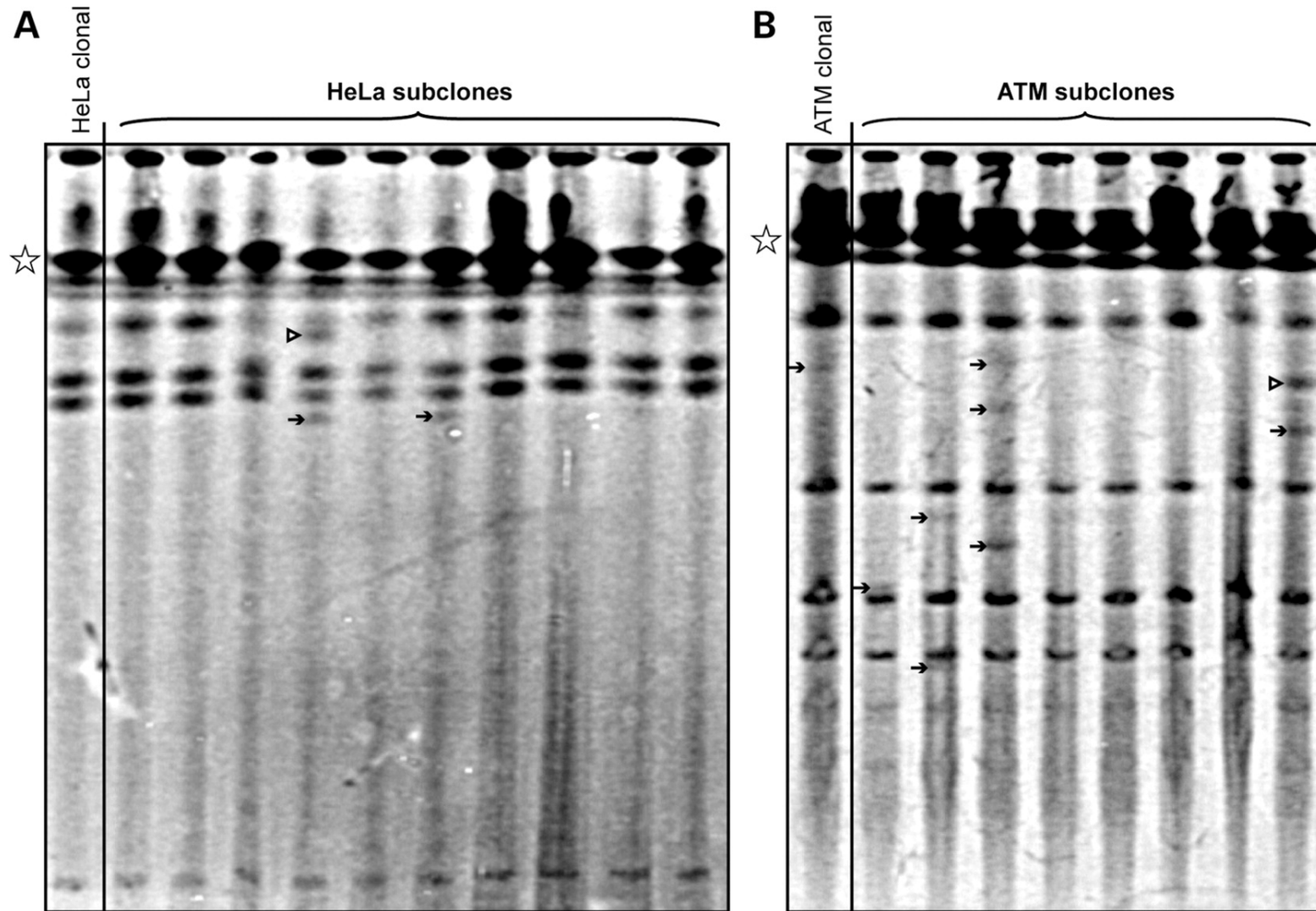
- **Array lengths vary from person to person**
- **Array lengths are not coordinated between chromosome homologs**
- **Individual arrays are usually inherited in Mendelian manner but undergo meiotic recombination with a frequency of ~10% per array**
- **Mitotic recombination is capable of altering rDNA architecture in different cells in the same person**

# Mitotic Recombination: Chromosomal Instability Syndromes

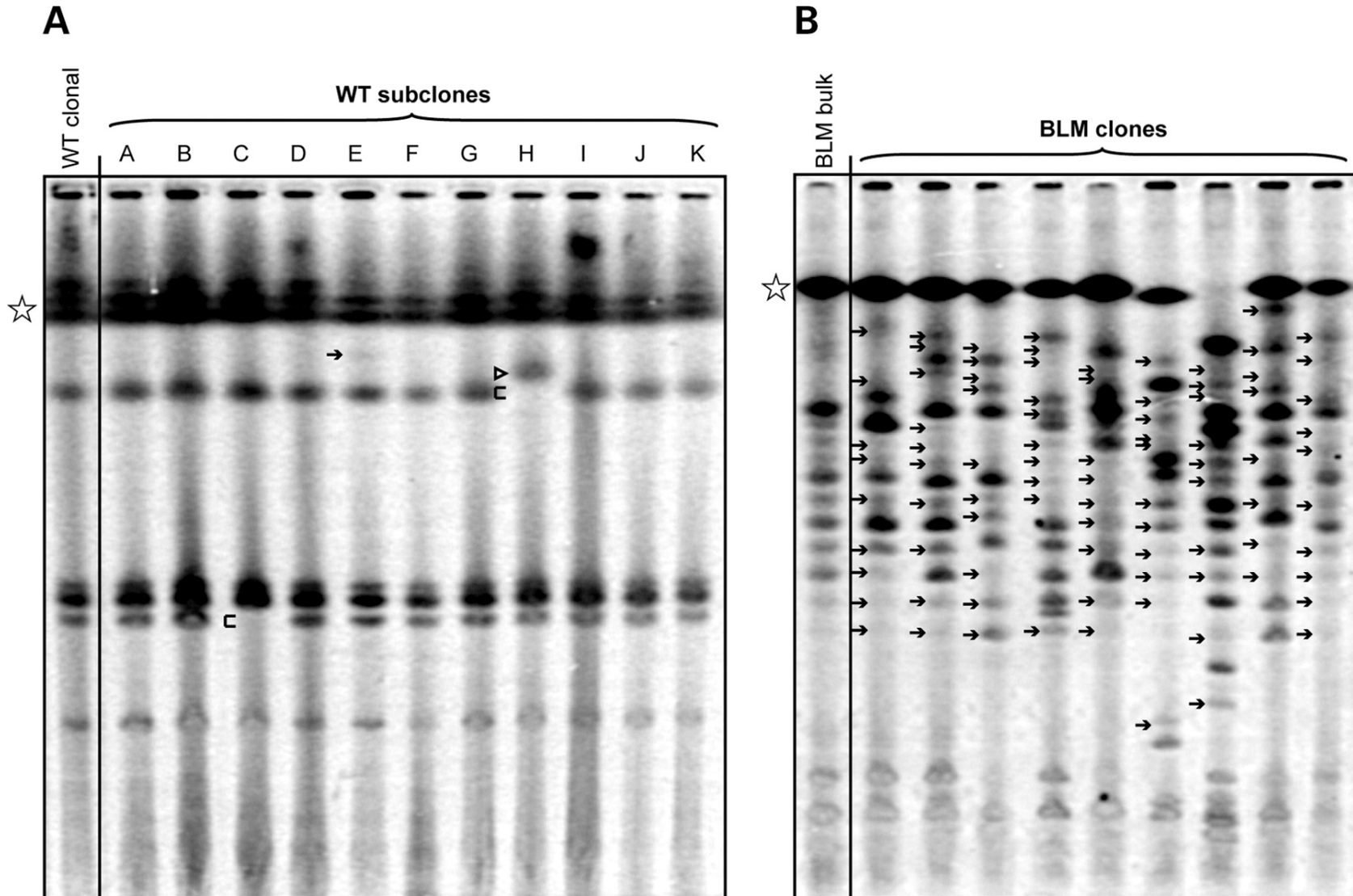


S. cerevisiae: molecular weight markers  
Lambda: lambda phage concatemer ladder  
 $\lambda_1$ : lambda monomer (48.5 kb)  
star: unresolved DNA

# Human Cells are (Mostly) Stable



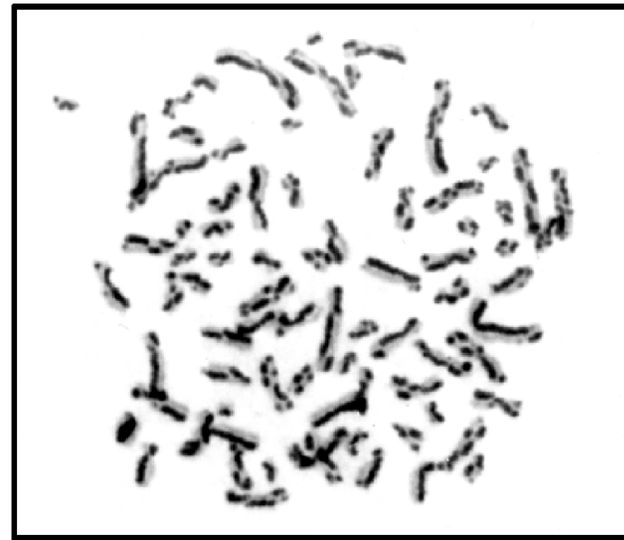
# BLM Cells Restructure rDNA Every Cell Division



# Bloom's Syndrome



Normal cell metaphase



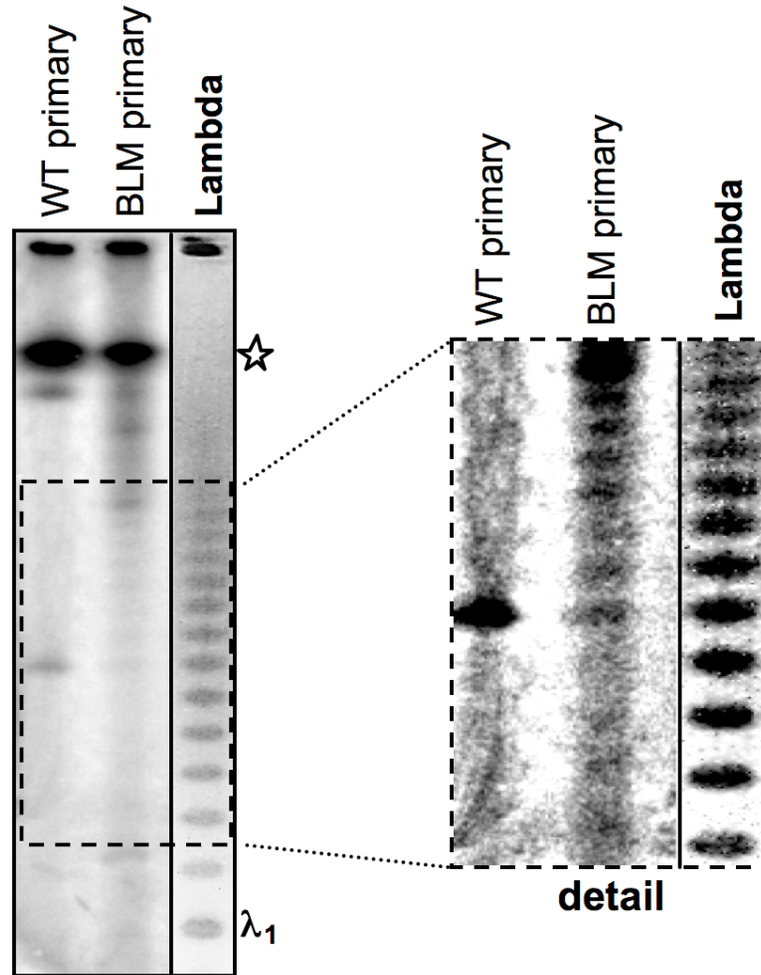
Bloom's syndrome cell metaphase

Amor-Gueret M. **Bloom's syndrome**. Orphanet Encyclopedia. February 2004

- proportionate pre- and postnatal growth deficiency
- sun-sensitive
- telangiectatic
- hypo- and hyperpigmented skin
- **chromosomal instability, predisposition to malignancy**

BLM protein in RecQ-helicase family with Werner Syndrome (WRN), Rothmund-Thomson Syndrome (RecQ4), and two other non-disease-associated members RecQL, RecQ5b

# BLM Primary Cells Show Instability



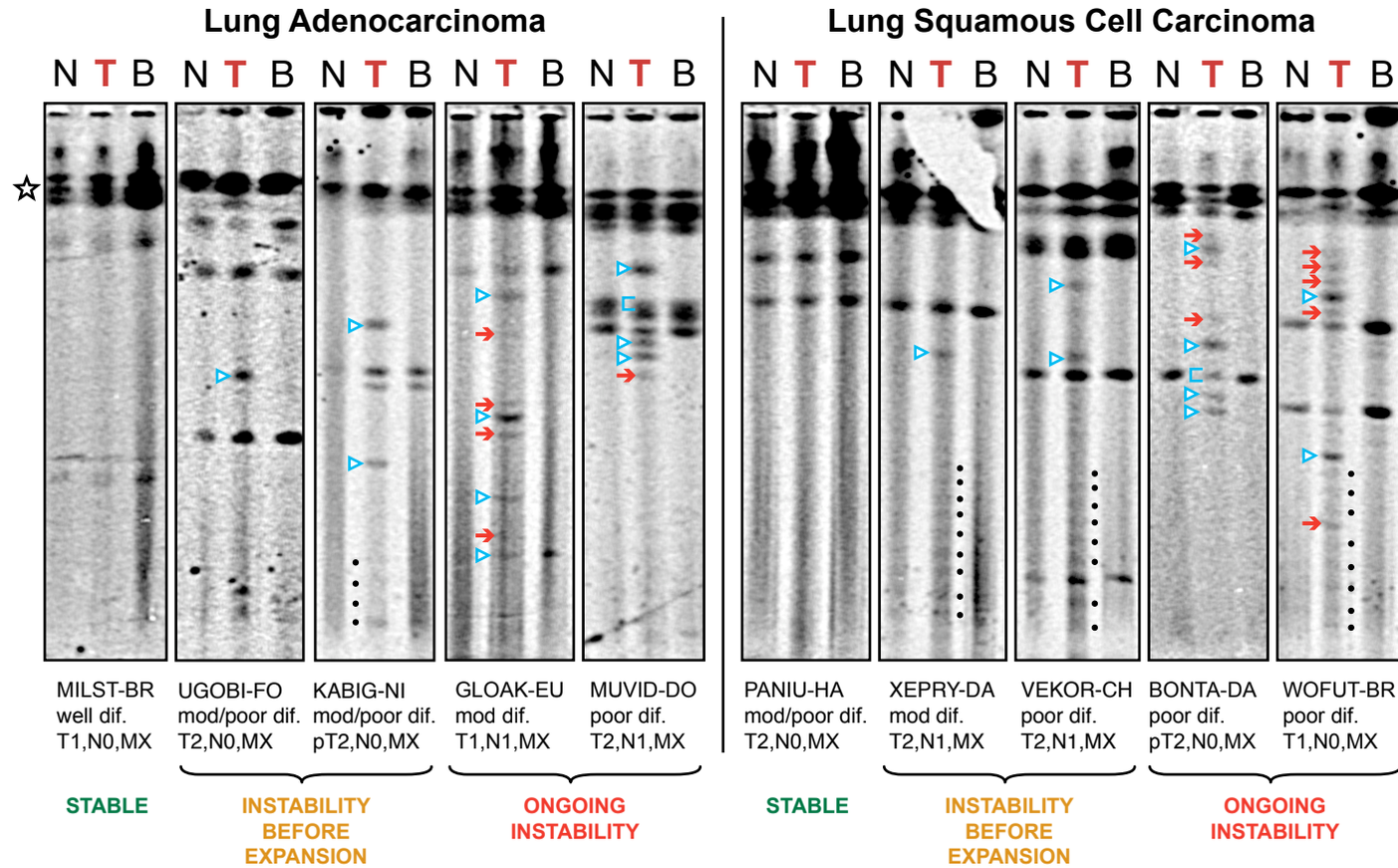
Lambda: lambda phage concatemer ladder  
 $\lambda_1$ : lambda monomer (48.5 kb)  
star: unresolved DNA

## MSI



Thibodeau SN, Bren G, Schaid D  
Microsatellite instability in cancer of the proximal colon.  
Science. 1993 May 7;260(5109):816-9.

# rDNA Clusters Restructured in Half of Solid Tumors



**Lung & Colorectal  
Cancer:  
(39 tumors analyzed)**

**45% stable  
30% preclonal alteration  
25% active restructuring**