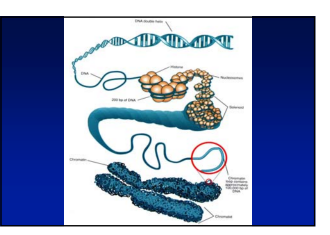
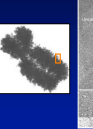
# The Human Genome

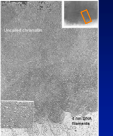
- 6 billion base pairs of DNA
   ~ 3 meters of DNA
- Approximately 30,000 70,000 genes
   Approximately 80-100,000 proteins
- These genes are spread across 24 different chromosomes
- One chromosome each from each parent, for a total of 23 pairs (24 different chromosomes) or 46 chromosomes per somatic cell



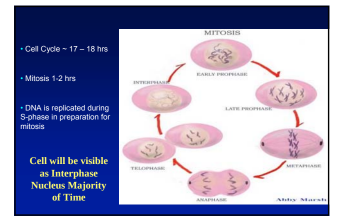
### Chromatin Compaction

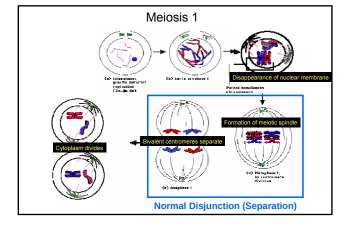


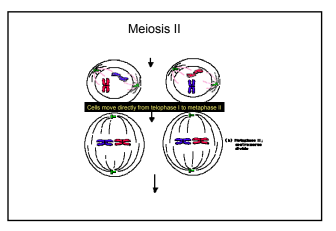


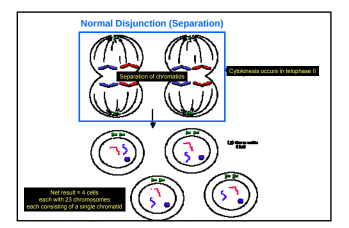


Metaphase chromosome is compacted into a structure that is 50,000 times shorter than its extended length









#### Male Meiosis

- · Begins at puberty
- Continues throughout life
- Spermatocytes continually replaced by mitosis
- Sperm maturation involves loss of histones and highly condensed DNA
- Each cycle from spermatocyte to sperm takes about 40 days
- Each meiotic division produces 4 sperm

#### **Female Meiosis**

- All oocytes are formed during fetal life: continually lost by apoptosis throughout life
- Meiotic prophase begins at 14 weeks of gestation
- Meiosis is arrested after diplotene and resumes only at the time of ovulation, when meiosis I is completed
- Meiosis II is completed only after fertilization
- Cell division is asymmetrical, producing one large egg and 3 nonfunctional polar bodies

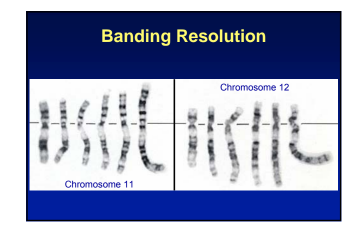
# Cytogenetics

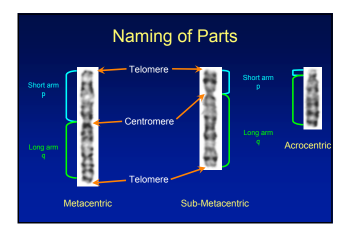
Banding techniques enable identification of chromosomes

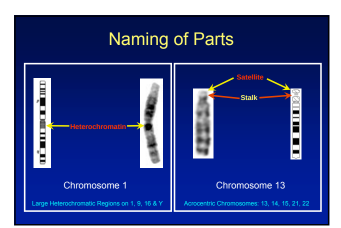
#### Chromosome Identification

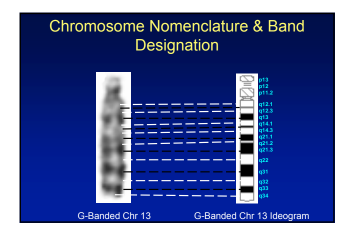
Chromosomes are identified by their size, banding pattern and the position of the centromere

									772			
1	2	3	4	5	6	7	8	9	10	11	12	
13	14	15	16	17	18	19	20	21	22	۷	x	





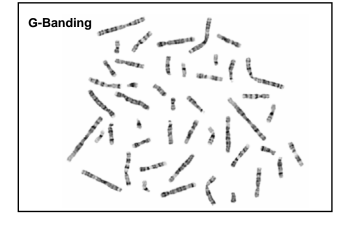


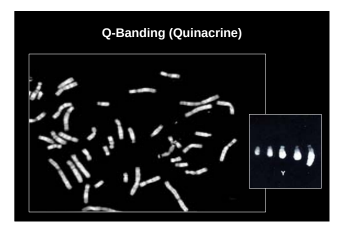


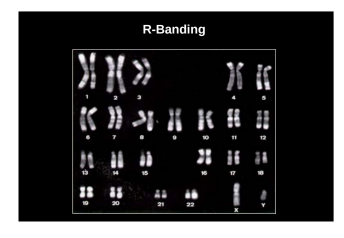
#### **Chromosome Banding**

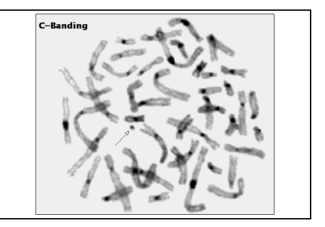
- Various banding patterns can be produced by using different enzymes, chemicals and stains
- G-Banding: Routine banding method in USA
   GTG: <u>G</u> bands produced with <u>Trypsin using Giemsa</u>
- Q-banding: First banding method developed for Human Chromosomes
   Certain fluorochromes, such as quinacrine dihydrochloride, will bind DNA & produce distinct banding patterns of bright & dull fluorescence. Requires fluorescence microscope for analysis
- C-banding: Stains constitutive heterochromatin around the centromeres and other heterochromatic regions (1, 9, 16, Y)
   CBG: <u>C</u> bands produced with <u>Barium hydroxide using Giemsa</u>

• R-banding: Banding pattern produced is the Reverse/Opposite of G-banding









#### **Studying Human Chromosomes**

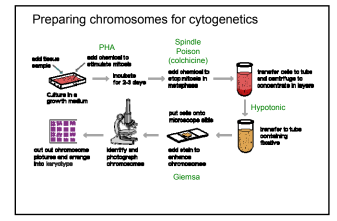
Cell will be visible as an Interphase Nucleus Majority of Time

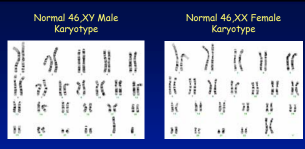
Have to catch cell during active division (METAPHASE) in order to view chromosomes



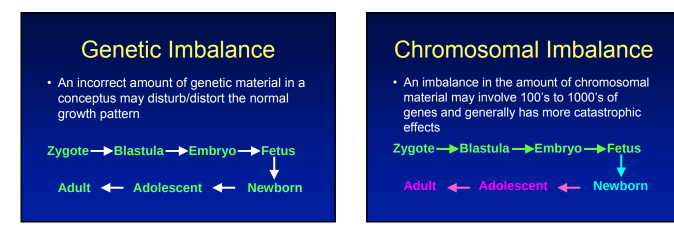
#### **Technical Advances**

- Mitogens (PHA) to push cell into active cell division
- Spindle poisons (colchicine) produce metaphase arrest
- Hypotonic solution to rupture nucleus
- Differential staining





 A precise amount of genetic material is required for normal development & functioning



# **Chromosomal Imbalance**

- May involve the gain or loss of a <u>whole</u> chromosome (complete aneuploidy) or of part of a chromosome (partial
- · The abnormality may occur in the non-mosaic or mosaic state (Mosaicism = Various chromosome complements in
- Monosomy (one missing) is generally more devastating than trisomy (one extra)

# Chromosomal Imbalance

- Most (complete) autosomal trisomies & all (complete) autosomal monosomies are so catastrophic that their presence in a conceptus is not compatible with survival
- Trisomies, monosomy X and polyploids are the most common abnormalities observed in spontaneous abortions
- ~ 66% of first trimester spontaneous abortions
- ~20% of 2nd trimester spontaneous abortions

#### Frequency of Chromosome Abnormalities

Source	%
Sperm	8
Oocytes	20-30
"Good" preimplantation embryos	30-40
"Poor" preimplantation embryos	85
Early recognized conceptions (>4 wks)	8-10
Early miscarriages	60-70
Late fetal deaths and stillbirths	7
Livebirths	0.3
Children with mental retardation	12-15
Infants with congenital heart disease	25

#### Chromosomal Imbalance and Pregnancy\* Loss

Trisomies

- 65% • 11% Monosomies
- 11% Triploidies
- 7.5% **Multiple Aneuploidies**
- 5.5% Tetraploid and structural

Most common trisomy observed in POC studies

• Never seen in liveborn

- Next most common (equally)

	Chromosomal
Imbalance	

### Survivable Chromosomal Imbalance

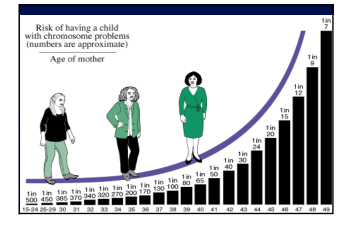
- · Only a few complete non-mosaic aneuploidies are observed in liveborns.
  - Down syndrome (Tri 21), Edward Syndrome (Tri 18), Patau Syndrome (Tri13), Turner Syndrome (Mono X)
- · All other imbalances will contain much smaller chromosomal regions (partial aneuploidy) that would allow for the organisms to survive .... Albeit with clinical abnormalities (in most cases)

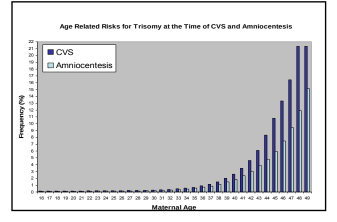
#### Numerical abnormalities

- Ploidy: The category of chromosome changes which involve the addition or loss of complete sets of chromosomes.
- Triploidy
  - The possession of one complete extra set of chromosomes. Usually caused by polyspermy, the fertilisation of an egg by more than one sperm. Such embryos will usually spontaneously abort.
- Tetraploidy
  - Usually the result of a failure of the first zygotic division. It is also lethal to the embryo. Any other cell division may also fail to complete properly and in consequence a very small proportion of tetraploid cells can sometimes be found in normal individuals (mosaicism).

#### Autosomal Numerical abnormalities

- Aneuploidy
  - The category of chromosome changes which do not involve whole sets. It is usually the consequence of a failure of a single chromosome (or bivalent) to complete division.
- Monosomies
  - All autosomal monosomies are lethal in very early embryogenesis. Most abort too early even to be recognised as a conception.
- Down syndrome, trisomy 21
  - The incidence of trisomy 21 rises sharply with increasing maternal age.





#### Clinical Phenotypes of Chromosomal Abnormalities

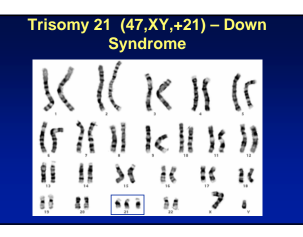
- Associated with Developmental Delay/MR
- Alteration of facial morphogenesis to produce characteristic facial features
- Growth delay
- · Malformations of the internal organs especially cardiac

Indication for chromosome analysis = MCA/MR

# Survivable Chromosomal Imbalance

 Only a few full non-mosaic aneuploidies are observed in liveborns

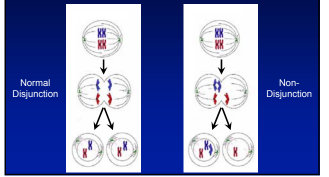
Trisomy 13 (47,XY,+13)	Trisomy 18 (47,XY,+18)	Trisomy 21 (47,XY,+21)			
12 15 21 31 1	KIIHHHH	N K K K K G			
KRNNSN	is if if if if if if if	(5 )( )  Ic    ); ))			
128 13 24 42 23	18 49 45 16 18 63 888	HIMKKKK			
		8 9 50 9 7 6			



#### Down syndrome: Cytogenetics

95% caused by nondisjunction = Trisomy 21
 95% of trisomies due to maternal nondisjunction
 75% of errors occur during meiosis 1
 MATERNAL MEIOSIS 1 NONDISJUCTION

# MEIOSIS-1 NONDISJUCTION





#### Trisomy 21 Down Syndrome

- Hypotonia
- · Redundant neck fold/flat occiput
- Low set ears with characteristic pinnae
- Protruding/large tongue
- Abnormal dermatoglyphics
  - Simian line and clinodactyly
  - Wide space between 1<sup>st</sup> & 2<sup>nd</sup> toes

#### **Down Syndrome**



Small overfolding of Angulated upper helix. Small/absent ear lobes



**Epicanthal fold** 



Simian crease





#### Down syndrome: Medical **Problems**

3%

- Gastrointestinal obstruction
- Respiratory infections Common 15-20 X
- Leukemia
- Congenital heart defect 40%
- Moderate to severe mental retardation 100%
- Development:
  - Early intervention program

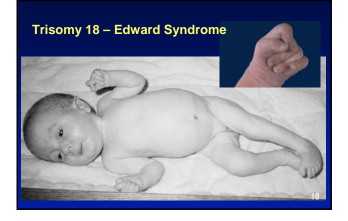
Trisomy 18 (47,XY,+18) – Edward Syndrome							
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#### Trisomy 18

- Incidence 1:3333 live births
- Most common abnormality in stillbirths with multiple congenital abnormalities
- Prenatal growth deficiency resulting in a small for gestational age infant (SGA)
- 90% congenital heart defect VSD
- 10% alive at one year
- Marked developmental disability

#### **Trisomy 18 - Physical Features**

- Prominent occiput
- Micrognathia
- Microcephaly
- Low set malformed ears
- Characteristic clenched fists
- Rocker-bottom feet
- Short big toe that is dorsiflexed



#### Trisomy 18

- Prominent Occiput
- Low-set malformed ears
- Small chin
- Clenched fists





#### Trisomy 18 – Rocker-Bottom Feet



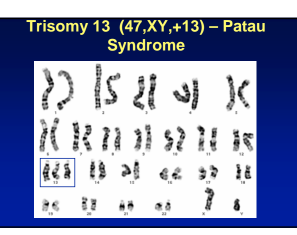
#### Trisomy 18

- Ocular manifestations in 10%
- · Low-arch dermal ridge pattern
- Underdeveloped nails
- Congenital anomalies of lungs, diaphragm, and kidneys
- Hernias, cryptorchidism, rectus muscle separation



# Trisomy 18 - Medical Management

- Feeding difficulties GE reflux
- Apnea
- Seizures
- Slow post natal growth
- Developmental disability/ mental retardation
- Scoliosis



#### Trisomy 13

- Incidence 1:5,000 births
- Distinctive malformation pattern (Craniofacial and Central Nervous System)
- 95% spontaneously aborted
- Survival rate and development similar to Trisomy 18

#### Trisomy 13 Patau Syndrome

- Microcephaly with sloping forehead
- Holoprosencephaly
- Ophthalmologic abnormalities
  - microphthalmia or anophthalmia
    Colobomata of iris and ciliary body
- Cleft lip +/- palate
- · Low set ears with abnormal helices

#### Trisomy 13 Patau Syndrome

- Cardiac defects: ASD, PDA, VSD
- Males: cryptorchidism ; Females: Bicornuate uterus
- Polycystic kidneys
- Aplasia cutis congenita
- Polydactyly of hands +/- feet
- Rockerbottom feet

#### Trisomy 13 -Cytogenetics

- 75% due to meiotic nondisjunction
- 20% arise from translocations
   25% are due to familial translocations
- 5% due to mosaicism
  - Mitotic nondisjunction

#### Trisomy 13 - Cleft Lip & Palate



Trisomy 13







# Trisomy 13 – Polydactyly



#### Numerical Abnormalities of the Sex Chrms

Sex Chromosome Aneuploidies

 Because of X inactivation and because of the paucity of genes on the Y chromosome, aneuploidies involving the sex chromosomes are far more common than those involving autosomes.

#### Sex Chromosome Abnormalities

- Turner syndrome
- Klinefelter syndrome (XXY)
- Triple X
- XYY

#### Numerical Abnormalities of the Sex Chrms

- Turner syndrome 45,X
  - The incidence is about 1 in 5000 female births but this is only the tip of the iceberg because 99% of Turner syndrome embryos are spontaneously aborted.
  - Individuals are very short, they are usually infertile. Characteristic body shape changes include a broad chest with widely spaced nipples and may include a webbed neck.
  - IQ and lifespan are unaffected.

#### Turner syndrome: Phenotype

20%

50%

- Facies- Triangular shape
   Posteriorly rotated ears
- Webbed neck
- Shield chest
- Lymphedema at birth
- Coarctation of aorta
- Structural kidney defects
- Rx: Growth hormone and estroger



Turner Syndrome



#### Turner syndrome

- SHORT STATURE
- OVARIAN DYSGENESIS
   INFERTILITY
- LEARNING DISABILITIES
   SPATIAL PERCEPTION



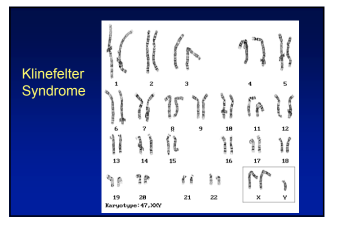
#### Turner syndrome: Cytogenetics

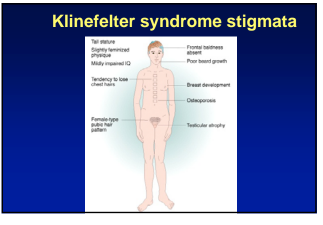
- 45, X 50%
   45,X/46,XX Mosaics 30-40%
   Structural X abnormalities 10-20%
   80% due to paternal meiotic error
- 15-20% spontaneous abortions due to 45,X



#### Numerical Abnormalities of the Sex Chrms

- Klinefelter Syndrome 47,XXY
  - The incidence at birth is about 1 in 1000 males.
  - Testes are small and fail to produce normal levels of testosterone which leads to breast growth (gynaecomastia) in about 40% of cases and to poorly developed secondary sexual characteristics. There is no spermatogenesis (Sterility).
  - These males are taller and thinner than average and generally have a slight reduction in IQ (10-15 points below sibs). Many Kleinfelter males lead a normal life.
  - Very rarely more extreme forms of Kleinfelter's syndrome occur where the patient has 48, XXXY or even 49, XXXXY karyotype. These individuals are generally severely retarded.







- 1:1000 female births
- No phenotypic abnormalities

 Variable symptoms: REPRODUCTIVE LOSS/ STERILITY LEARNING DISABILITIES/ SPEECH

LANGUAGE

Multi-X: 48,XXXX 49,XXXXX

#### Numerical Abnormalities of the Sex Chrms

- XXX females
  - About one woman in 1000 has an extra X chromosome. It seems to do little harm, individuals are fertile and do not transmit the extra chromosome.
  - They do have a reduction in IQ comparable to that of Kleinfelter's males (10-15 point below sibs).

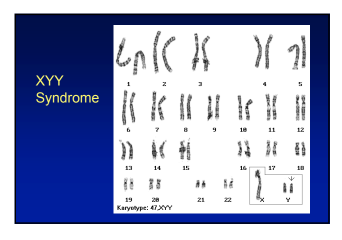
#### Variants

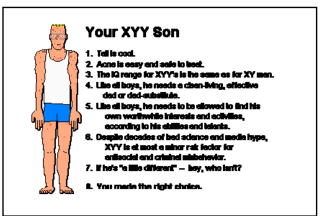
- Multi-X: 48,XXXX 49,XXXXX
- Mild to moderate MR
- Variable dysmorphic features

#### Numerical Abnormalities of the Sex Chrms

#### • 47,XYY males

- Incidence 1 in 1000 male births. May be without any symptoms.
- Males are tall but normally proportioned.
- 10 15 points reduction in IQ compared to sibs? (/Q.93-109/109-147)
- More common in high security institutions than chance would suggest? (Problems with impulse control?)

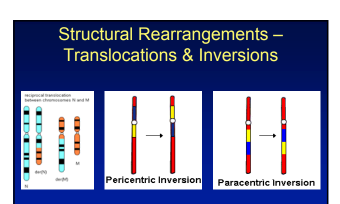




#### Structural Rearrangements

- Translocations
- Inversions

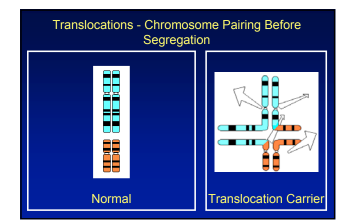
Multiple options for Gametes only 2 of which are balanced

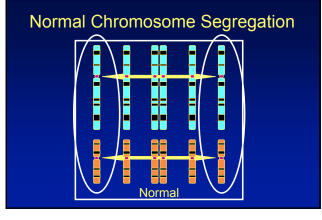


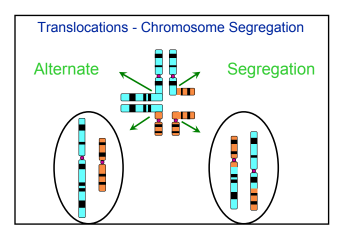
#### Structural Abnormalities - Translocations

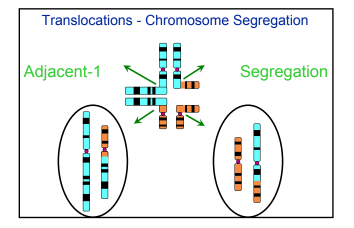
- RECIPROCAL/BALANCED = Mutual exchange after two breaks Most balanced reciprocal translocations have no phenotype

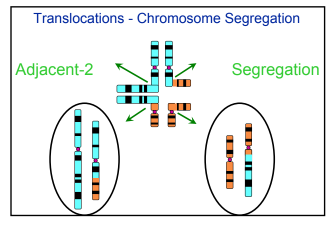
  - Clinical phenotype may result if there is a disruption of critical genes at the breakpoint regions. Clinical phenotype tends to be more like those observed in single gene defects
- UNBALANCED DERIVATIVE = Partial monosomy & Partial trisomy

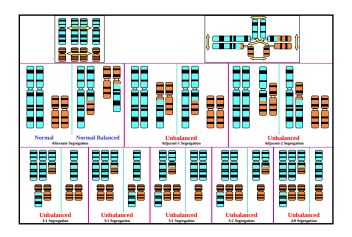


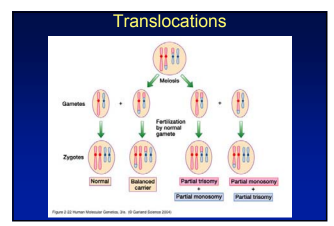


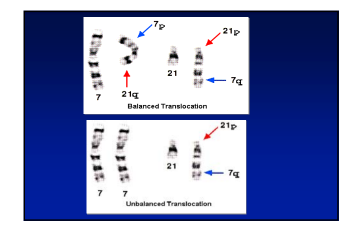


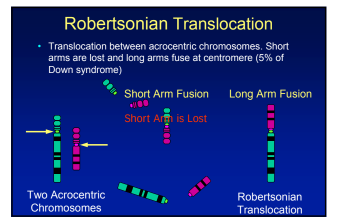


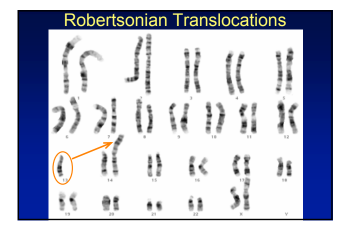


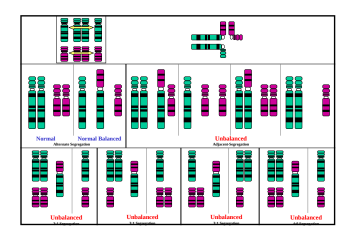


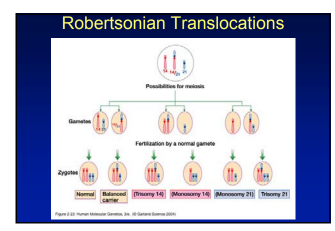




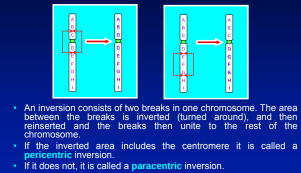


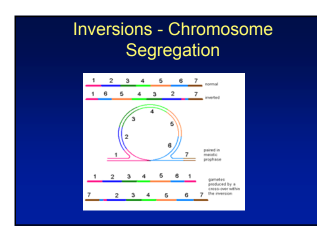


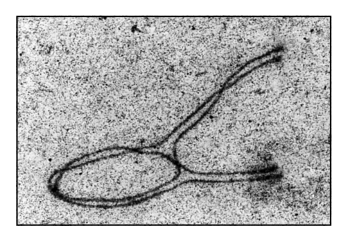


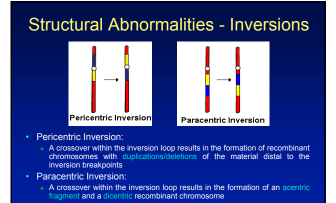


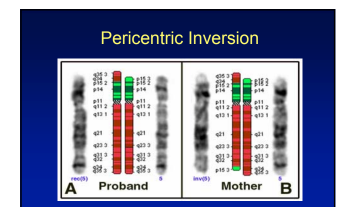
#### Structural Abnormalities - Inversions

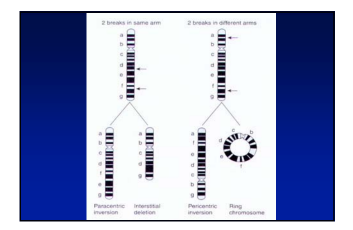


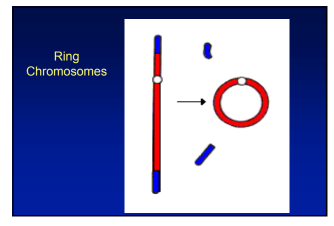












# Translocations & Inversions

#### BALANCED

- Most balanced rearrangements have no phenotype
- Clinical phenotype may result if there is a disruption of critical genes at the breakpoint regions. Clinical phenotype tends to be more like those observed in single gene defects
- UNBALANCED
  - Partial Monosomy/Trisomy

#### Deletions

- 5p- Cri-du Chat
- 4p- Wolf-Hirschhorn

## Deletion 5p - Cri Du Chat Syndrome

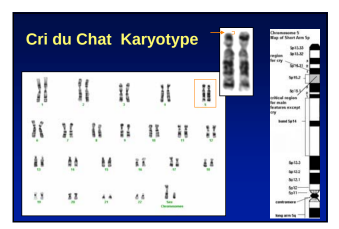




# Cri Du Chat Syndrome - Older childre





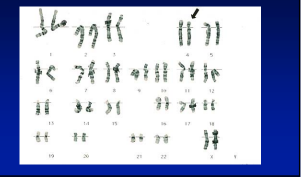




#### Deletion 4p Cytogenetics Wolf Hirschhorn Syndrome

- Deletion in the terminal band 4p16.3
- 87% of cases due to de novo interstitial deletion of paternal origin
- 13% due to unbalanced product of a parental reciprocal translocation

#### Deletion 4p Karyotype



### "Viable" Chromosome Imbalance

- Unbalanced Translocations Partial Monosomy & Partial Trisomy
- Deletions Partial Monosomy
- Duplications Partial Trisomy
- Ring & marker chromosomes Partial Trisomy
- Recombinant Inversion derivatives Partial Monosomy & Partial Trisomy
- Isochromosomes Partial Trisomy/Tetrasomy or Partial Monosomy & Partial Trisomy

# "Viable" Chromosome Imbalance

- 1 in 150 Livebirths
- 10-15% Mentally retarded population
  - Higher percentage when cryptic rearrangements are included
- Most case reports involve partial aneuploidy