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

Chromatin Compaction

- Cell Cycle ~ 17 – 18 hrs
- Mitosis 1-2 hrs
- DNA is replicated during S-phase in preparation for mitosis

Cell will be visible as Interphase Nucleus Majority of Time

Meiosis

Meiosis I

- Normal Disjunction (Separation)
- Telophase I
- Nuclear membrane reforms
- Daughter cells each receive one chromosome from each homologous pair

Meiosis II

- Cell moves directly from metaphase I to metaphase II
- Telophase II
- Daughter cells receive 23 chromosomes from each parent

Meiosis I

- Disappearance of nuclear membrane
- Metaphase I
- Meiotic spindle forms
- Bivalent centromeres separate
- Chromosomes move to opposite poles

Meiosis II

- Cell moves directly from metaphase I to metaphase II
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Meiosis I

- Disappearance of nuclear membrane
- Metaphase I
- Meiotic spindle forms
- Bivalent centromeres separate
- Chromosomes move to opposite poles
Normal Disjunction (Separation)

Separation of chromatids
Cytokinesis occurs in telophase II

Net result = 4 cells
each with 23 chromosomes
each consisting of a single chromatid

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

Banding Resolution

Chromosome 11
Chromosome 12
**Naming of Parts**

- Telomere
- Centromere
- Short arm (p)
- Long arm (q)
- Metacentric
- Sub-Metacentric
- Acrocentric

**Chromosome Nomenclature & Band Designation**

Chromosome 13

- G-Banding: Routine banding method in USA
  - GTG: G bands produced with Trypsin using Giemsa
- 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: C bands produced with Barium hydroxide using Giemsa
- R-banding: Banding pattern produced is the Reverse/Opposite of G-banding

**Chromosome Banding**

- Various banding patterns can be produced by using different enzymes, chemicals and stains
- **G-Banding:** Routine banding method in USA
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**G-Banding**

**Q-Banding (Quinacrine)**
R-Banding

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

Preparing chromosomes for cytogenetics

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

Studying Human Chromosomes

Normal 46,XY Male Karyotype

Normal 46,XX Female Karyotype
Genetic Imbalance

• An incorrect amount of genetic material in a conceptus may disturb/distort the normal growth pattern.

Zygote → Blastula → Embryo → Fetus

Adult ← Adolescent ← Newborn

Chromosomal Imbalance

• An imbalance in the amount of chromosomal material may involve 100’s to 1000’s of genes and generally has more catastrophic effects.

Zygote → Blastula → Embryo → Fetus

Adult ← Adolescent ← Newborn

Chromosomal Imbalance

• May involve the gain or loss of a whole chromosome (complete aneuploidy) or of part of a chromosome (partial aneuploidy).

• The abnormality may occur in the non-mosaic or mosaic state (Mosaicism = Various chromosome complements in different cells).

• 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

<table>
<thead>
<tr>
<th>Source</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sperm</td>
<td>6.0</td>
</tr>
<tr>
<td>Dicentrics</td>
<td>20–30</td>
</tr>
<tr>
<td>&quot;GOOD&quot; postimplantation embryos</td>
<td>20–40</td>
</tr>
<tr>
<td>&quot;POOR&quot; postimplantation embryos</td>
<td>0.0</td>
</tr>
<tr>
<td>Early recognized conceptions (&gt;4 wks)</td>
<td>0.05</td>
</tr>
<tr>
<td>Early miscarriages</td>
<td>0.5</td>
</tr>
<tr>
<td>Late fetal deaths and stillbirths</td>
<td>0.5</td>
</tr>
<tr>
<td>Livebirths</td>
<td>6.5</td>
</tr>
<tr>
<td>Children with mental retardation</td>
<td>10–15</td>
</tr>
<tr>
<td>Infants with congenital heart disease</td>
<td>5.0</td>
</tr>
</tbody>
</table>

Chromosomal Imbalance and Pregnancy* Loss

• 65% Trisomies

• 11% Monosomies

• 11% Triploidies

• 7.5% Multiple Aneuploidies

• 5.5% Tetraploid and structural

Trisomy 16

• Most common trisomy observed in POC studies

• Never seen in liveborn

Trisomy 21 & 22

• Next most common (equally)

*Recognized pregnancies
Chromosomal Imbalance

- Only a few complete non-mosaic aneuploidies are observed in liveborns.
  - Down syndrome (Tri 21), Edward Syndrome (Tri 18), Patau Syndrome (Tri 13), 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.

Risk of having a child with chromosomal problems (numbers are approximate)

Age Related Risks for Trisomy at the Time of CVS and Amniocentesis
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 21 (47,XY,+21) – Down Syndrome

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

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 NONDISJUNCTION

MEIOSIS-1 NONDISJUNCTION

Normal Disjunction

Non-Disjunction

Trisomy 21 – Down Syndrome
Trisomy 21 Down Syndrome

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

Down Syndrome

- Epicanthal fold
- Small overfolding of Angulated upper helix.
- Small/absent ear lobes
- Simian crease

Down Syndrome

- Small overfolding of Angulated upper helix.
- Small/absent ear lobes
- Simian crease

Down syndrome: Medical Problems

- Gastrointestinal obstruction 3%
- Respiratory infections Common
- Leukemia 15-20 X
- Congenital heart defect 40%
- Moderate to severe mental retardation 100%

- Development:
  - Early intervention program

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

- Early intervention program
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 – Edward Syndrome

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

Trisomy 18 – Rocker-Bottom Feet

- 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 – High Morbidity & Mortality

- Feeding difficulties
- GER reflux
- Apnea
- Seizures
- Slow postnatal growth
- Developmental disability/mental retardation
- Scoliosis

Trisomy 18 - Medical Management

Trisomy 13 (47,XY,+13) – Patau Syndrome

- 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

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

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**

- Facies- Triangular shape
- Posteriorly rotated ears
- Webbed neck
- Shield chest
- Lymphedema at birth
- Coarctation of aorta 20%
- Structural kidney defects 50%
- Rx: Growth hormone and estrogen

**Turner Syndrome**

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

**Height**

- Turner Syndrome
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 Klinefelter males lead a normal life.
  - Very rarely more extreme forms of Klinefelter's syndrome occur where the patient has 48, XXXY or even 49, XXXXY karyotype. These individuals are generally severely retarded.

Klinefelter syndrome stigmata

- 1:1000 female births
- No phenotypic abnormalities
- Variable symptoms:
  - REPRODUCTIVE LOSS/ STERILITY
  - LEARNING DISABILITIES/ SPEECH LANGUAGE

Triple X – 47,XXX

- 48, XXXX 49, XXXXY

45,X0 in SAB’s
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

**47,XXY 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? (IQ: 85-109/109-147)
- More common in high security institutions than chance would suggest? (Problems with impulse control?)

**XXY Syndrome**

**Your XYY Son**
1. Tell him cool.
2. Accuse him and call it latent.
3. The IQ range for XYY’s is the same as for XY men.
4. Like all boys, he needs a clean-living, effective dad or dad-substitute.
5. Like all boys, he needs to be allowed to find his own interests and activities, according to his abilities and talents.
6. Despite decades of bad science and media hype, XYY is at most a minor risk factor for antisocial and criminal behavior.
7. If he’s “a little different” – hey, who isn’t?
8. You raised the right son!

**Structural Rearrangements**

- Translocations
- Inversions

Multiple options for Gametes only 2 of which are balanced

**Structural Rearrangements – Translocations & Inversions**

- Pericentric Inversion
- Paracentric Inversion
Structural Abnormalities - Translocations

Interchange of genetic material between nonhomologous chromosomes

- 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

Normal Chromosome Segregation

Translocations - Chromosome Segregation

Adjacent-1 Segregation

Translocations - Chromosome Segregation

Adjacent-2 Segregation
Robertsonian Translocation

- Translocation between acrocentric chromosomes. Short arms are lost and long arms fuse at centromere (5% of Down syndrome)

**Robertsonian Translocations**

<table>
<thead>
<tr>
<th>Balanced Translocation</th>
<th>Unbalanced Translocation</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Balanced Translocation" /></td>
<td><img src="image" alt="Unbalanced Translocation" /></td>
</tr>
</tbody>
</table>

Translocations

- Gametes
  - Normal
  - Balanced carrier
  - Partial monosity
- Zygote
  - Normal
  - Partial monosity
  - Partial trisomy
  - Partial trisomy

Robertsonian Translocations

- Normal
- Normal Balanced
- Unbalanced
- Unbalanced Balanced
- Unbalanced Unbalanced

Alternate Segregation Adjacent Segregation

<table>
<thead>
<tr>
<th>3:1 Segregation</th>
<th>3:1 Segregation</th>
<th>3:1 Segregation</th>
<th>3:1 Segregation</th>
<th>4:0 Segregation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unbalanced</td>
<td>Unbalanced</td>
<td>Unbalanced</td>
<td>Unbalanced</td>
<td>Unbalanced</td>
</tr>
</tbody>
</table>

Translocations

- Short Arm Fusion
  - x/0
  - Short Arm is Lost
- Long Arm Fusion
  - Robertsonian Translocation
Robertsonian Translocations

- An inversion consists of two breaks in one chromosome. The area between the breaks is inverted (turned around), and then reinserted and the breaks then unite to the rest of the chromosome.
- If the inverted area includes the centromere it is called a pericentric inversion.
- If it does not, it is called a paracentric inversion.

Structural Abnormalities - Inversions

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Inversions - Chromosome Segregation

- Pericentric Inversion:
  - A crossover within the inversion loop results in the formation of recombinant chromosomes with duplications/deletions of the material distal to the inversion breakpoints.
- Paracentric Inversion:
  - A crossover within the inversion loop results in the formation of an acentric fragment and a dicentric recombinant chromosome.

Structural Abnormalities - Inversions

- Pericentric Inversion
- Paracentric Inversion

Pericentric Inversion

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- A crossover within the inversion loop results in the formation of an acentric fragment and a dicentric recombinant chromosome.
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 children
Cri du Chat Karyotype

Deletion 4p - Wolf- Hirschhorn Syndrome

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