Non-Disjunction, Aneuploidy & Abnormalities in Chromosome Structure

Packet #16
Introduction I

- **Ploidy**
  - Degree of repetition of the basic number of chromosomes
  - **Diploidy**
    - Chromosomes repeat 2X
    - Remember, in humans, you have one copy of a chromosome from the maternal father and one from the maternal mother

- **Euploidy**
  - “True” ploidy
  - In the case of somatic human cells, euploidy occurs when the cell is diploid.
Aneuploidy

• “Not True” ploidy
  • Having too many or too few chromosomes.

• Non-Disjunction
  • An event that occurs occasionally during meiosis in which a pair of chromosomes fail to separate so that the resulting germ (sex) cell has either too many or too few chromosomes.
Genetic Disorders Caused by Sex Chromosome Aneuploidy

Sex Chromosome Aneuploidy
Turner Syndrome

- $2n - 1$
  - $45\ XO$
    - $44$ autosomes + $1\ X$ chromosome
      - There is the absence of a sex chromosome

- Female in appearance but their female sex organs do not develop at puberty and they are sterile

- Short in stature; swelling; broad chest; low-set ears; webbed necks

- Shows normal intelligence but some cognitive functions are defective
  - Difficulties in mathematics and memory

- 1 in 5000 female births show Turner Syndrome
Turner Syndrome II

- Short stature
- Low hairline
- Shield-shaped thorax
- Widely spaced nipples
- Shortened metacarpal IV
- Small finger nails
- Brown spots (nevi)
- Characteristic facial features
- Fold of skin
- Constriction of aorta
- Poor breast development
- Elbow deformity
- Rudimentary ovaries
- Gonadal streak (underdeveloped gonadal structures)
- No menstruation
Klinefelter Syndrome

- \(2n + 1\)
  - 47 XXY
    - 44 autosomes + 3 sex chromosomes
    - There is an extra X chromosome
- Male in appearance and they too are sterile
- Female type pubic hair pattern
- May have breast development
Klinefelter Syndrome II

- Frontal baldness absent
- Tendency to grow fewer chest hairs
- Breast development
- Female-type pubic hair pattern
- Small testicular size
- Poor beard growth
- Narrow shoulders
- Wide hips
- Long legs
Comparison of Turner & Klinefelter Syndromes

- Poor beard growth
- Breast development
- Under-developed testes

- Characteristic facial features
- Web of skin
- Constriction of aorta
- Poor breast development
- Under-developed ovaries
# Jacob’s Syndrome (YY Syndrome)

**XYY**

- Tall and thin males
- Males have normal sexual development and are able to create normal gametes.
- Increased risk of learning disabilities and delayed speech and language skills.
- In some cases, individuals develop behavioral problems such as explosive temper, hyperactivity, impulsivity, defiant actions, or, in some cases, antisocial behavior.
- After age of 35, extra Y chromosome often degenerates and is not passed onto offspring

1/1000 newborn boys
Triple X Syndrome

XXX

- Increased risk of learning disabilities and delayed development of speech and language skills.
- Delayed development of motor skills.
- Behavioral and emotional difficulties are possible.
- Fertile females with normal gametes.
Genetic Disorders Caused by Autosomal Aneuploidy

Autosomal Aneuploidy
Down Syndrome—Trisomy 21

- Caused by an extra copy of chromosome #21
  - There are three copies of chromosome #21 in their somatic cells
- 0.15 percent of all live births
- Growth failure and delay in learning
- Big toes widely spaced; small chin; face is flat & broad; short neck
- Congenital heart disease
- Mean life expectancy is about 17 years and only 8% survive past age 40
Edward’s Syndrome
Trisomy 18

- Multiple defects
  - Mental and physical deficiencies
  - Facial abnormalities
    - Small head
    - Ear deformities
  - Extreme muscle tone
    - Spasticity and other damage
  - Clenched hands
  - Heart defects
  - Kidney malfunctions

- Low Survival Rate
  - Majority die before birth
  - Early Death
  - Death is typical by the age of 1
Patau Syndrome
Trisomy 13

- Multiple defects
  - Mental and physical deficiencies
    - Large triangular nose; eye defects; extra fingers; low set ears
  - Wide variety of defects in organs

- Early death
  - $\approx 80\%$ die within 1st year
  - Death is typical by the age of 3
Abnormalities in Chromosome Structure
Introduction I

- The **changes in the shape of the chromosome** may be due to either of the following
  - Translocation
  - Deletions
  - Fragile sites
Abnormalities in Chromosome Structure

Translocation
Translocation

1. A chromosome fragment breaking off and attaching to a non-homologous chromosome
   • Can result in deletion and/or duplication of genes

2. Reciprocal translocation
   • Two non-homologous pairs exchange genetic information
Translocation Down Syndrome

- 4% of Down Syndrome cases
- Individuals actually have 46 chromosomes
- One of copies of chromosome #14 has combined with chromosome #21
  - The large arm of chromosome #21 has been translocated to the large arm of another chromosome--usually chromosome #14
Abnormalities in Chromosome Structure

Deletion
Deletion

- The loss of part of a chromosome
- The abnormal chromosome is known as a *deletion*
- Sometimes chromosomes break and fail to rejoin
Cri du Chat Syndrome
(Cry of the Cat)

- Infants have a distinctive cry that sounds like a cat mewing
- Part of the short arm of chromosome #5 is deleted
  - Breakage point varies from case to case
- Infants normally have a small head with altered features
  - Moon face
- Infants normally survive childhood
- Exhibit severe mental retardation
Abnormalities in Chromosome Structure
Fragile Sites
Fragile Sites

• Weak points at specific locations in chromatids
• Appears to be a place where part of a chromatid appears to be attached to the rest of the chromosome by a thin thread of DNA
• Have been identified on the X chromosome and certain autosomes
Fragile X Syndrome

- Fragile site occurs near the tip of the X chromosome
  - Where nucleotide triplet CGG is repeated many more times than normal

- Most common cause of learning disabilities.

- normal structure
- broad forehead
- elongated face
- large prominent ears
- strabismus (crossed eyes)
- highly arched palate
- hyperextensible joints
- hand calluses (from self-abuse)
- pectus excavatum (indentation of chest)
- mitral valve prolapse (benign heart condition)
- enlarged testicles
- hypotonia (low muscle tone)
- soft, fleshy skin
- flat feet
- seizures (in about 10 percent)
Genetic Screening & Pedigrees
Genetic Screening & Genetic Counseling

- Genetic Screening
  - Identifies individuals who might carry a serious genetic disease
  - Screening of newborns is the first step in preventative medicine

- Genetic Counseling
  - Provide couples, concerned about the risk of abnormality in their children, medical and genetic information
Pedigrees

• Definition
  • A family tree that shows the transmission of genetic traits within a family over several generations.

• Pedigree Analysis
  • Useful in detecting autosomal dominant mutations, autosomal recessive mutations, X linked recessive mutations and defects due to genomic imprinting
  • Genomic Imprinting
    • Expressions of a gene based on its parental origin
Pedigree Analysis

(a) Dominant trait (widow’s peak)

(b) Recessive trait (attached earlobe)
Homework

• Bioinformatics
• Proteomics
• Aminocentesis
• Chronic villus sampling (CVS)
• Preimplantation genetic diagnosis (PGD)
• Know how to discuss (argue for/against)
  • Genetic discrimination
  • The Human Genome Project