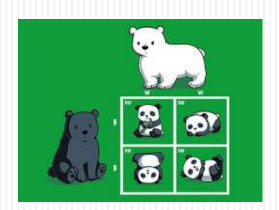
# Gene Inheritance and Transmission Extensions of Mendelian genetics

Lecture 5

**SLE254 Genetics** 

Chapter 4 Concepts of Genetics (12<sup>th</sup> ed)

Pp 98-130





# Departures from clear cut Mendelian transmission genetics

- Lethal alleles
- Incomplete dominance
- Codominance
- Multiple alleles
- Epistasis
- Pleiotropy
- Genetic heterogeneity
- Penetrance

- Expressivity
- Anticipation
- Germline mosaicism
- Phenocopies
- Linkage
- Continuous variation

#### Lethal alleles

- Mutations occur giving rise to new forms of alleles
  - Some of these are lethal mutations
- A dominant lethal is an allele that causes death of the organism that contains it, whether homozygous or heterozygous for the allele
- A recessive lethal is an allele that that causes death when homozygous
  - In the heterozygote, a lethal allele is masked by the presence of an allele for 'wild type'.

#### Lethal alleles – dominant

 Dominant lethal genes are rarely detected due to their rapid elimination from populations

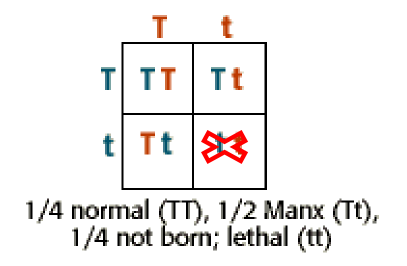
#### How could they be maintained in populations?

- Huntington's disease a neurological disorder in humans, which reduces life expectancy
- Because the onset of Huntington's disease is slow, individuals carrying the allele can pass it on to their offspring

  Symptoms in Huntington's disease

#### Lethal alleles - recessive

- Manx cats are heterozygous for a dominant mutation that results in no tails
- A cross between two Manx cats produces a 2:1 phenotype ratio (tailless to tailed) instead of the normal 3:1 phenotype ratio
  - The recessive homozygotes do not survive

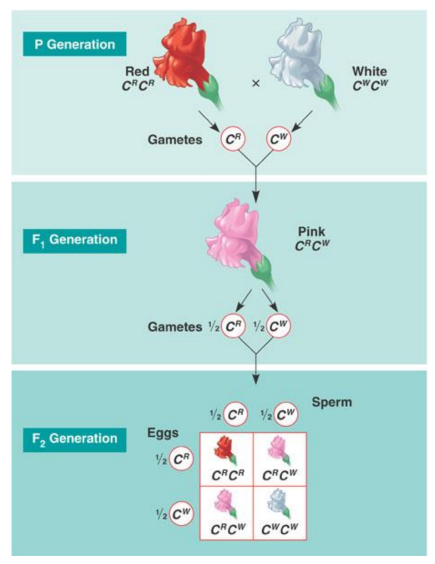




## Incomplete (Partial) dominance

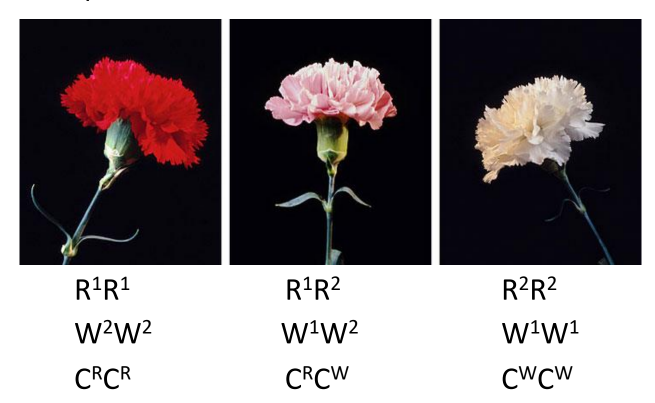
- Both alleles blend their effects
- The phenotype of the heterozygote lies somewhere between those of the two kinds of homozygotes

- The F<sub>2</sub> generation shows only one pair of alleles determines the phenotype
  - However, phenotype ratio is identical to genotype ratio 2:1:1 and not 3:1 like complete dominance



## Incomplete (Partial) dominance

- Neither allele is recessive so different symbols are used
  - Examples include



#### Codominance

- Both alleles show their effects DO NOT blend
- In codominance, neither allele are dominant; both are expressed. A cross between organisms with two different phenotypes produces offspring with has both phenotypes of the parental traits shown.

White





Red



Roan (red and white hairs, NOT pink)

#### Codominance

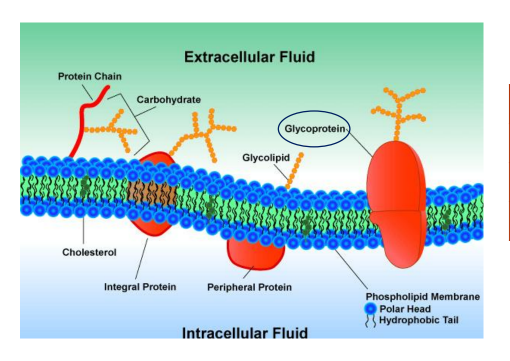
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#### Codominance

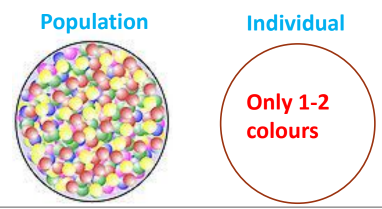
- The MN blood group
  - In humans, two forms of a glycoprotein exist on the surface of red blood cells – designated M and N
- An individual may exhibit either one or both proteins



| Genotype | Phenotype |
|----------|-----------|
| ΓWΓW     | M         |
| LMLN     | MN        |
| ΓνΓν     | N         |

## Multiple alleles

- Many genes have more than two alternative alleles
- This increases the number of different genotypes and phenotypes that exist with respect to the particular gene
- Multiple alleles can only be studied in populations
  - An individual diploid organism will only have, at most, two alternative forms of the same gene
  - A population will show all the alternatives



### Multiple alleles – ABO blood groups

- Three alternative alleles of one gene
- Presence of antigens of the surface of red blood cells
  - Four phenotypes depending on the presence or absence of antigens

Alleles code for presence or absence of cell marker molecules on the erythrocyte surface

Antibodies in the serum which can identify and destream Antigens on the surface of another blood group

|    |                           | ABO Blood Groups                                       |  |   |   |  |  |
|----|---------------------------|--|--|---|---|--|--|
|    | Antigen<br>(on RBC)       | Antigen A  | Antigen B  | Antigens A + B  | Neither A or B                                      |  |  |
| rc | y Antibody<br>(in plasma) | Anti-B Antibody  Y Y Z                                 | Anti-A Antibody  | Neither Antibody  | Both Antibodies                                     |  |  |
|    | Blood<br>Type             | Type A Cannot have B or AB blood Can have A or O blood | Type B Cannot have A or AB blood Can have B or O blood | Type AB  Can have any type of blood  Is the universal recipient | Type O Can only have O blood Is the universal donor |  |  |

#### Multiple alleles – ABO blood groups

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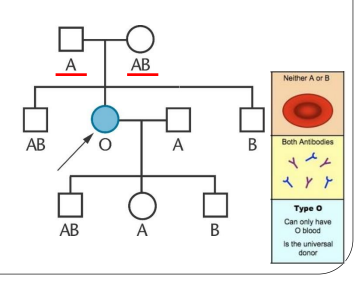
| Genotype                      | Antigen | Phenotype |
|-------------------------------|---------|-----------|
| IAIA                          | А       | Α         |
| I <sub>A</sub> I <sub>O</sub> | А       |           |
| <sup>B</sup>   <sup>B</sup>   | В       | В         |
| I <sub>B</sub> I <sub>O</sub> | В       |           |
| IAIB                          | A, B    | AB        |
| lolo                          | neither | 0         |

codominance

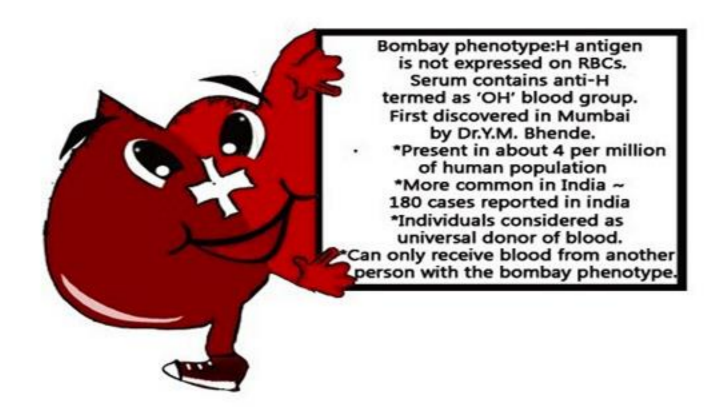
## The Bombay phenotype

- The Bombay Phenotype was first reported in 1952 in Bombay, India.
- Bombay cells can't be converted to group A or B
  - Mutation in the FUT1 gene prevents synthesis of H substance, vital for producing functional A and B antigens

So individual may have I<sup>A</sup> and/or I<sup>B</sup> alleles, but neither antigen is added to the cell surface and they are **functionally type** O



### The Bombay phenotype



http://www.bloodconnect.org/bombay-bloodtype

## **Epistasis**

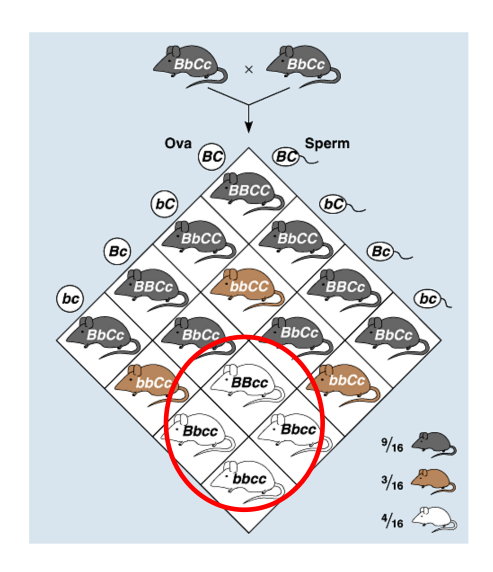
- A form of gene interaction in which one gene masks the phenotypic expression of another
- There are no new phenotypes produced by this type of gene interaction
- The alleles that are masking the effect are called epistatic alleles
- The alleles whose effect is being altered or suppressed are called the hypostatic alleles\*





## An example of epistasis

- If individual is cc, then is albino regardless of allele at b locus - due to gene interaction
- Normally two gene (dihybrid) crosses would produce a 9:3:3:1 ratio
- Due to gene interaction, we see a 9:3:4 F<sub>2</sub> ratio. The c locus is epistatic to the b locus.
- cc masks the b locus



## **Epistasis**

|      |                  |                 | F <sub>2</sub> Phenotypes |        |         | Modified          |         |
|------|------------------|-----------------|---------------------------|--------|---------|-------------------|---------|
| Case | Organism         | Character       | 9/16                      | 3/16   | 3/16    | 1/16              | ratio   |
| 1    | Mouse            | Coat<br>color   | agouti                    | albino | black   | albino            | 9:3:4   |
| 2    | Squash           | Color           | white                     |        | yellow  | green             | 12:3:1  |
| 3    | Pea              | Flower<br>color | purple                    |        | white   |                   | 9:7     |
| 4    | Squash           | Fruit<br>shape  | disc sphere               |        | ere     | long              | 9:6:1   |
| 5    | Chicken          | Color           | white                     |        | colored | white             | 13:3    |
| 6    | Mouse            | Color           | white-spotted             | white  | colored | white-<br>spotted | 10:3:3  |
| 7    | Shepherd's purse | Seed<br>capsule | triangular                |        |         | ovoid             | 15:1    |
| 8    | Flour<br>beetle  | Color           | 6/16 sooty and 3/16 red   | black  | jet     | black             | 6:3:3:4 |

Was Mendel just wrong?

No, none of these cases has violated the principles of segregation and independent

assortment - just added complexity

#### **Epistasis**

Squash fruit colour is controlled by two genes

Gene 1 is represented by a W

• Gene 2 is represented by a G

Which allele is epistatic in squash colour?

genotype, the squash fruit colour is white

Every time a dominant W allele shows up in a squash

The dominant W allele is epistatic

Cross a green squash (wwGg) with a white squash (Wwgg).

Genotypes and Phenotypes:

white

• W-/gg white

• ww/Ggreen

• ww/gg

• W-/G-

yellow

8:4:4

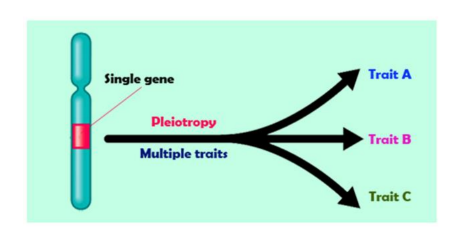
What colour are the offspring?

wwGg x Wwgg

|    | wG   | wG   | wg   | wg   |
|----|------|------|------|------|
| Wg | WwGg | WwGg | Wwgg | Wwgg |
| Wg | WwGg | WwGg | Wwgg | Wwgg |
| wg | wwGg | wwGg | wwgg | wwgg |
| wg | wwGg | wwGg | wwgg | wwgg |

## Pleiotropy

- Occurs when one gene influences multiple phenotypic traits
- The gene codes for a product that is, for example, used by various cells, or has a signalling function on various targets

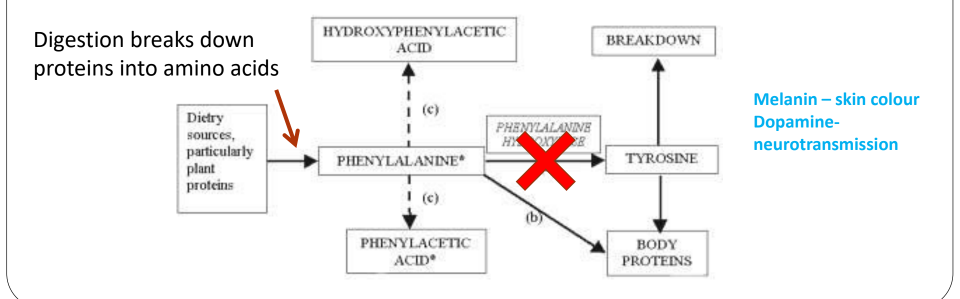


#### **Problem**

a mutation in a pleiotropic gene may have an effect on some or all traits simultaneously

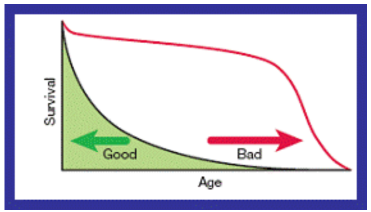
## Pleiotropy - example

- Phenylketonuria (PKU)
- Symptoms include intellectual impairment, reduced hair and skin pigmentation, microcephalic, eczema, musty smell.
  - Caused by any of over 400 mutations in a single gene that codes for the enzyme phenylalanine hydroxylase, which converts the amino acid phenylalanine to tyrosine



## Antagonistic pleiotropy

- The expression of a gene resulting in multiple competing effects, some beneficial but others detrimental to the organism
- Theory of aging (G. C. Williams, 1957)
  - Some genes responsible for increased fitness in the younger, fertile organism contribute to decreased fitness later in life



• E.g. BRCA2 gene, women with mutations are more fertile but after reproduction the gene causes cancer

## **Genetic Heterogeneity**

- A phenomenon in which a single phenotype or genetic disorder may be caused by any one of a multiple number of alleles or non-allele (locus) mutations
  - Allelic heterogeneity different mutations within a single gene locus (forming multiple alleles of that gene) cause the same phenotypic expression
    - E.g. 1000 known mutant alleles of the CFTR gene that cause cystic fibrosis
  - Locus heterogeneity variations in completely unrelated gene loci cause a single disorder
    - E.g. has Ehler's Danlos syndrome autosomal dominant, autosomal recessive, and X-linked origins

#### Penetrance

- The probability of a gene or genetic trait being expressed.
  - Complete penetrance the gene or genes for a trait are expressed in all the population who have the genes
  - Incomplete penetrance the genetic trait is expressed in only part of the population
- Penetrance can be difficult to determine reliably
  - E.g. in disease, the onset of symptoms could be age related, or affected by environmental codeterminants such as nutrition and smoking, as well as genetic cofactors

**Camptodactyly** 

## Expressivity

- Refers to variations in a phenotype among individuals carrying a particular genotype
  - E.g. Drosophila flies homozygous for recessive mutant gene eyeless exhibit range of phenotypes from normal eyes to complete absence of one or both eyes
- Other genes or environmental factors such as nutrition and temperature may be influencing or modifying the phenotype- severity ranges with developmental temperature

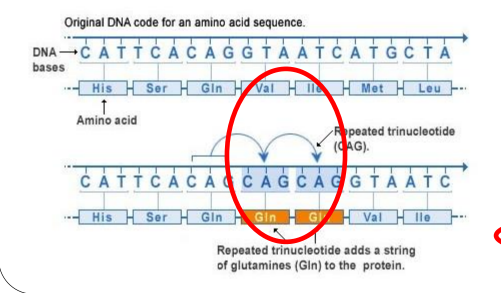






## Anticipation

- A phenomenon whereby the symptoms of a genetic disorder exhibit an earlier age of onset and are more severe as it is passed on to the next generation
  - E.g. Huntington's disease trinucleotide repeat disease



| Repeat count | Classification        | Disease status |
|--------------|-----------------------|----------------|
| <28          | Normal                | Unaffected     |
| 28–35        | Intermediate          | Unaffected     |
| 36–40        | Reduced<br>Penetrance | +/- Affected   |
| >40          | Full Penetrance       | Affected       |

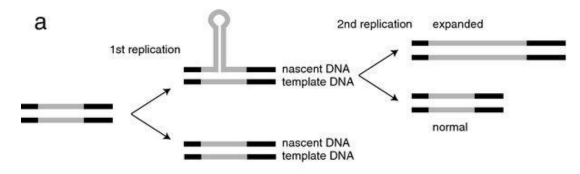
# Why do Triplet repeat mutations get worse over generations\*??

- 1. Slippage of the two complementary DNA strands during replication
- 2. Homologous recombination- every meiosis event
- 3. DNA repair

**Replication slippage** 

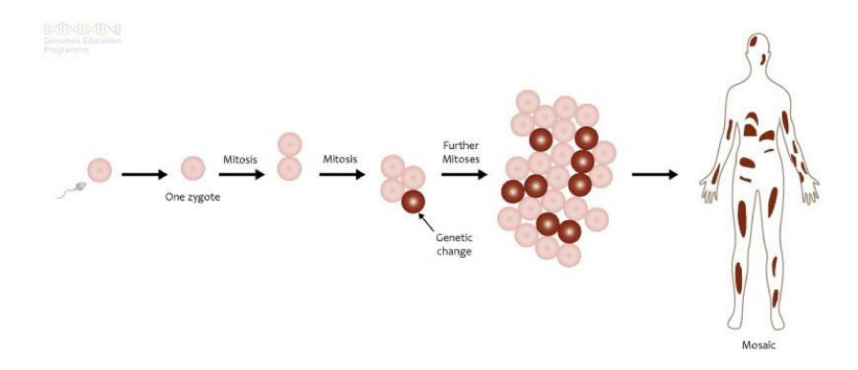
Mutation caused by denaturation and misplacement of the strands

-Mispairing of the complimentary strands In high sequence repeat regions



#### Germline mosaicism

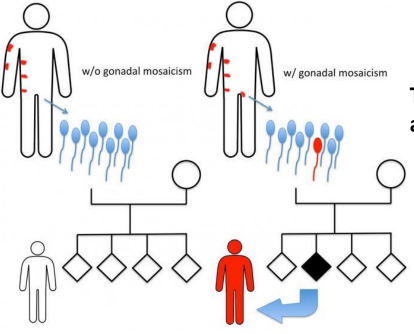
 Mosaicism – the presence of two or more populations of cells with different genotypes in one individual who has developed from a single fertilized egg



#### Germline mosaicism

**Germline mosaicism** – a special form of mosaicism, where some gametes (sperm or oocytes) carry a mutation, but the rest are normal

Cause is usually a mutation that occurred in an early stem cell that gave rise to the gonadal tissue



This can cause only some offspring to be affected, even for a dominant disease

#### Phenocopies

- A phenocopy is an individual whose phenotype under a particular environmental condition, is identical to the one of another individual whose phenotype is determined by the genotype
  - The phenocopy environmental condition mimics the phenotype produced by a gene (i.e. black coat)



Mutation in Tyrosinase (Gene C) active between 15-25 degrees



Himalayan rabbit standard coat
Raised 20deg

Himalayan rabbit 'copying' black rabbit
Raised in cold <15deg







Raised in cold <15deg

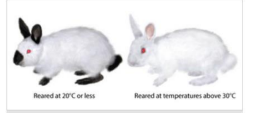


Figure 1: A pigment gene is influenced by temperature.

Gene C controls fur pigmentation in Himalayan rabbits. Because the gene is active when environmental temperatures are between 15 and 25°C, the rabbit reared at 20°C (left) has pigmentation on its ears, nose, and feet, where its body loses the most heat. The rabbit reared at temperatures above 30°C (right) has no fur pigmentation, because gene C is inactive at these higher temperatures.

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Raised in heat 30deg

20-25°C = standard coat: melanin mostly supressed

<20°C = black coat: melanin expressed

>25°C = white coat: melanin fully suppressed

## Think about this one for prac 1!

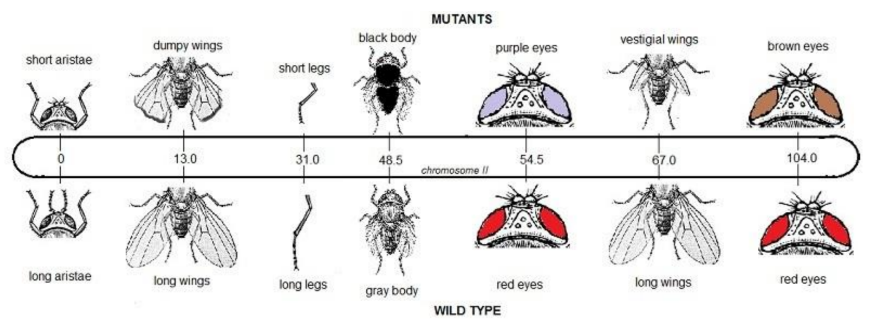


## Genomic imprinting

- Normally there is no difference of expression of the paternal and maternal alleles
- Genomic imprinting causes selective expression of a gene or genes inherited from one parent
- Not a mutation or permanent change
- Plays a role in several genetic disorders
  - E.g. The same region of chromosome 15 mutated but causes a different disease if inherited from mother or father
  - Prader-Willi syndrome: Paternal copy
  - Angelman syndrome: Maternal copy

## Linkage

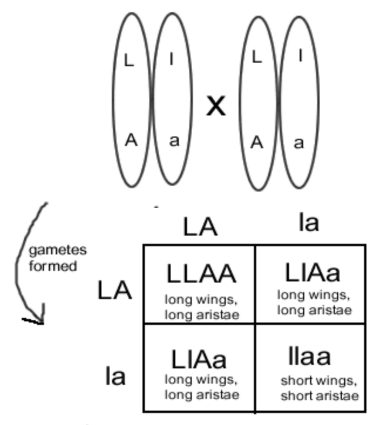
 Two or more genes located on the same chromosome that do not show independent assortment and tend to be inherited together



 In a given cross, the outcome depends on the proximity of genes on a chromosome

## Linkage – two gene example

 A fly that is heterozygous for long wings (LI) and heterozygous for long aristae (Aa) is crossed with another fly of the same type. AaLl x AaLl. In both cases the dominant alleles are located on the same chromosome.



3/4 long wings, long aristae 1/4 short wings, short aristae

What ratio of offspring phenotypes would you expect?

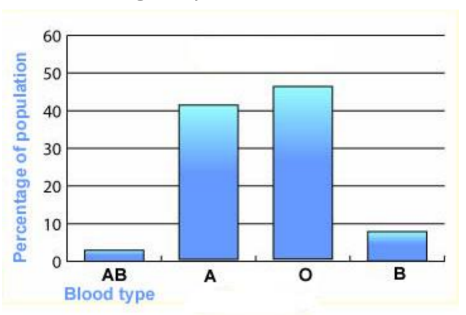
#### Different definitions of variation

- Some traits are controlled by two or more genes
- Phenotypes can be discontinuous or continuous
- Discontinuous variation shows distinct (discrete) phenotypes
  - E.g. Pea plant colour, ABO blood group

#### Discontinuous variation







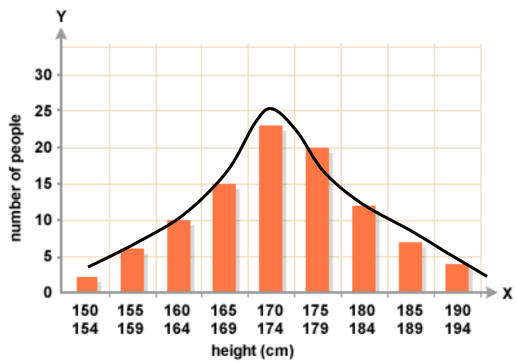
#### Continuous variation

- Continuous variation shows a series of overlapping phenotypic classes
  - E.g. Height, weight, hand span, shoe size, milk yield in cows

Continuous variation



Seed color red to white



#### How are traits defined?

- Polygenic traits
  - Traits controlled by two or more genes
  - Patterns of inheritance that can be measured quantitatively
    - Example: human eye colour
- Multifactorial traits
  - Polygenic traits resulting from interactions of <u>two or</u> <u>more genes</u> and <u>one or more environmental factors</u>
    - Example: skin colour

#### A multifactorial trait: skin colour

Skin colour is controlled by 3 or 4 genes and

environmental factors

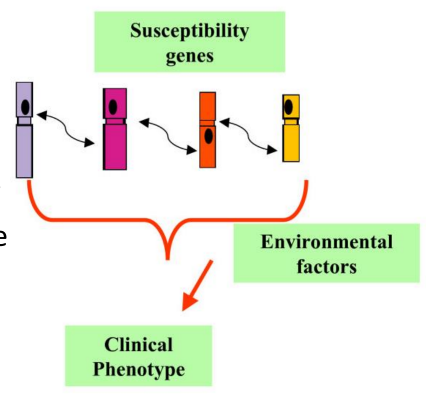


 We're all born with a skin colour based on our genes but the environment can alter this



#### How are traits defined?

- Complex traits
  - Traits controlled by <u>multiple genes</u> and the interaction of <u>environmental factors</u> where the contributions of genes and environment are undefined
    - Example: hypertension, obesity, cardiovascular disease, depression, autism



Many human diseases are controlled by the action of several genes