

Gene Inheritance and Transmission

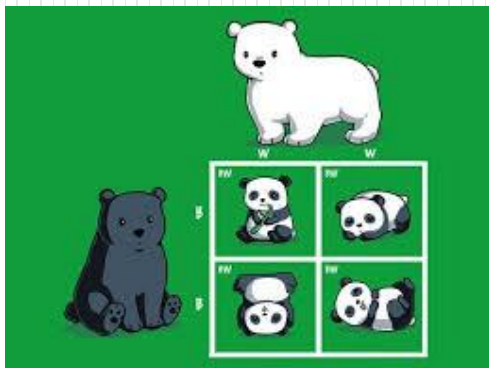
Extensions of Mendelian genetics

Lecture 5

SLE254 Genetics

Chapter 4 Concepts of Genetics (12th 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

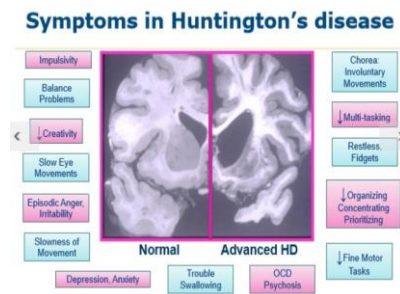
- Mutations occur giving rise to new forms of alleles
 - Some of these are **lethal mutations**
- A **dominant lethal** is an allele that causes death of the organism that contains it, whether homozygous or heterozygous for the allele
- A **recessive lethal** is an allele that causes death when **homozygous**
 - 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



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

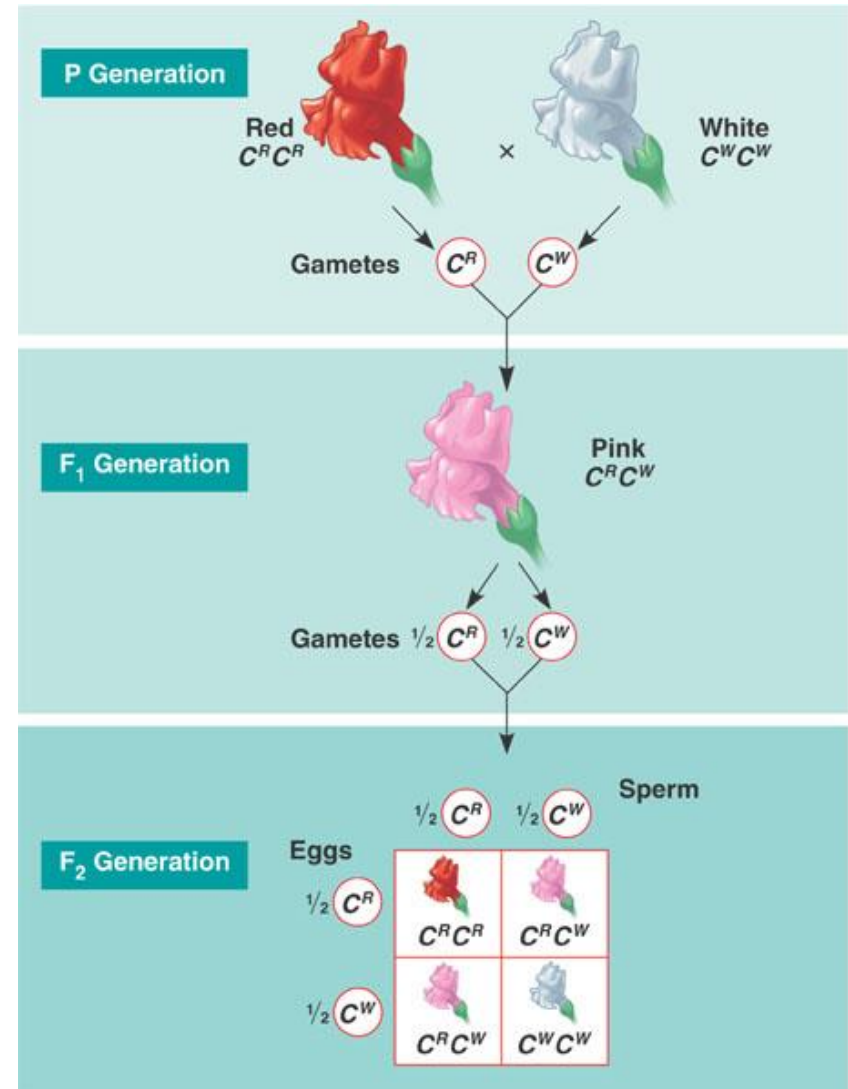
	T	t
T	TT	Tt
t	Tt	

1/4 normal (TT), 1/2 Manx (Tt),
1/4 not born; lethal (tt)



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_2 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



R^1R^1

W^2W^2

C^RC^R



R^1R^2

W^1W^2

C^RC^W



R^2R^2

W^1W^1

C^WC^W

Codominance

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

White



Red



Roan (red and white hairs, NOT pink)

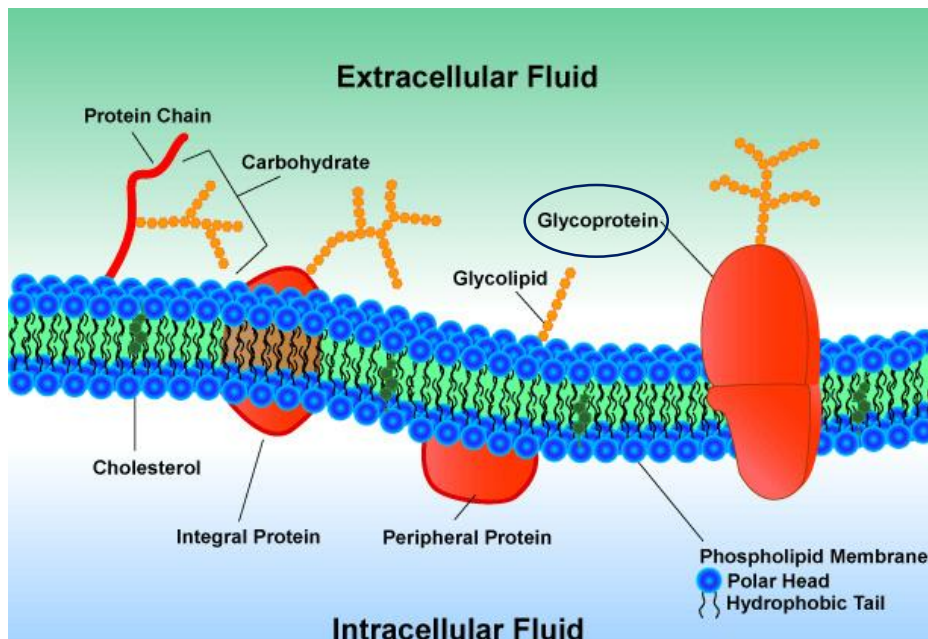
Codominance

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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
$L^M L^M$	M
<u>$L^M L^N$</u>	<u>MN</u>
$L^N L^N$	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

Population



Individual



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

Codominance

1+2



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

ABO Blood Groups				
Antigen (on RBC)	1 Antigen A 	2 Antigen B 	Antigens A + B 	3 Neither A or B
Antibody (in plasma)	Anti-B Antibody 	Anti-A Antibody 	Neither Antibody 	Both Antibodies
Blood 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

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

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

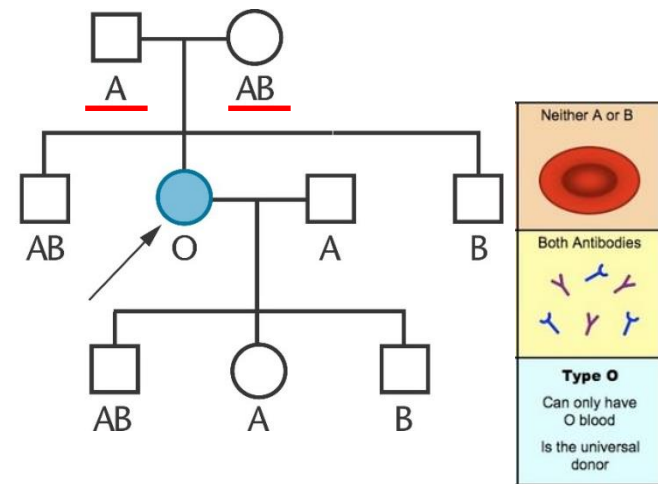
Genotype	Antigen	Phenotype
$I^A I^A$	A	A
$I^A I^O$	A	
$I^B I^B$	B	B
$I^B I^O$	B	
$I^A I^B$	A, B	AB
$I^O I^O$	neither	O

codominance

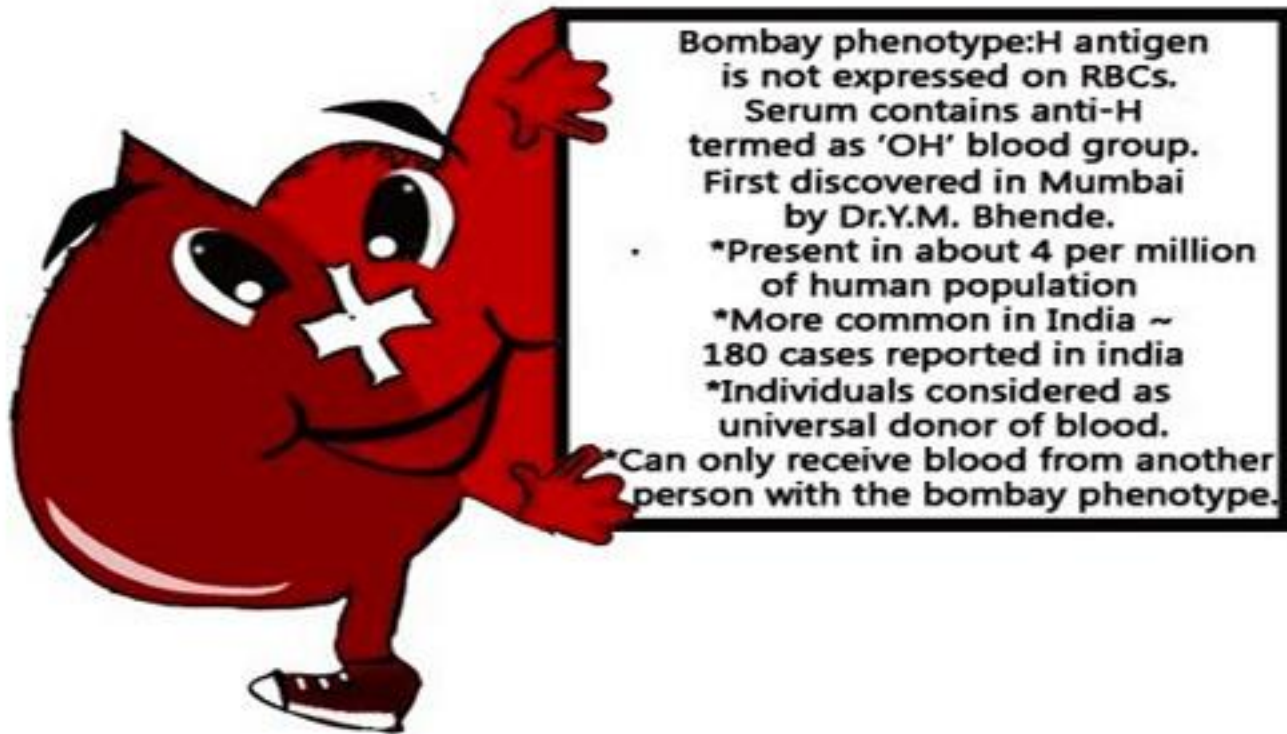
The Bombay phenotype

- 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^A and/or I^B 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>

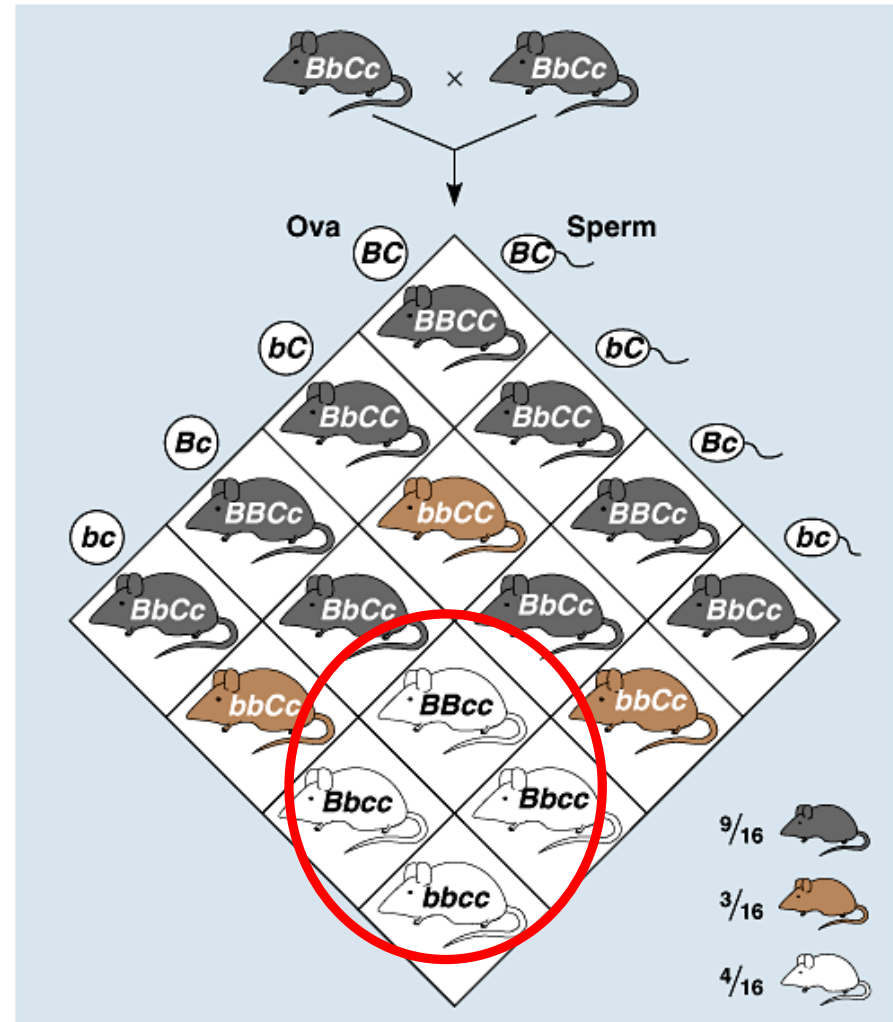
Epistasis

- 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_2 ratio. The c locus is epistatic to the b locus.
- cc masks the b locus

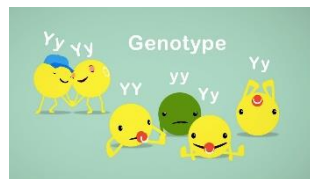


Epistasis

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

The dominant W allele is epistatic

- Genotypes and Phenotypes:

- W- / G- white
- W- / gg white
- ww / G- green
- ww / gg yellow

Every time a dominant W allele shows up in a squash genotype, the squash fruit colour is white

Cross a green squash (wwGg) with a white squash (Wwgg).
What colour are the offspring?

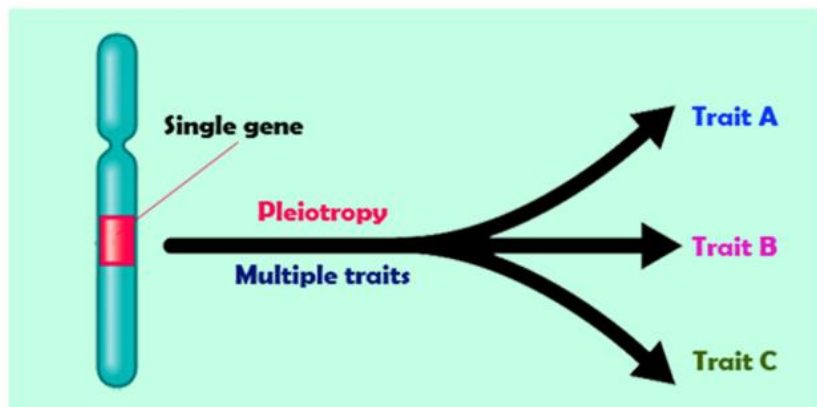
8 : 4 : 4

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



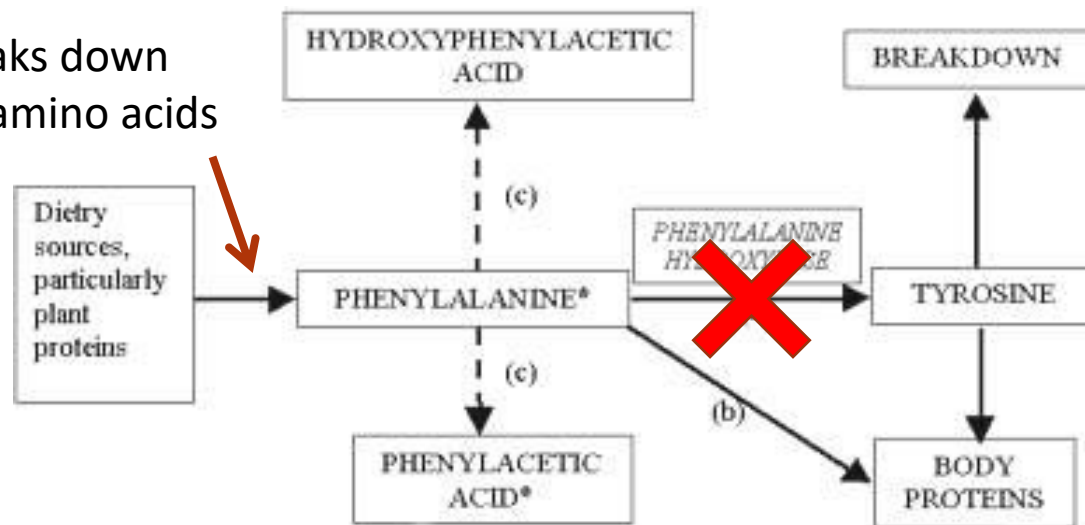
Problem

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**

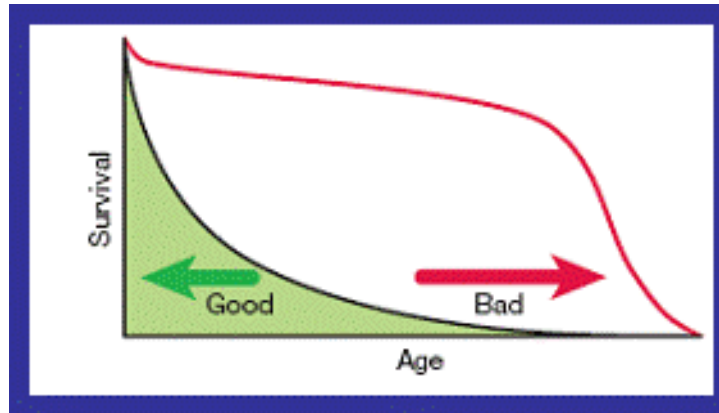
Digestion breaks down proteins into amino acids



Melanin – skin colour
Dopamine-
neurotransmission

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

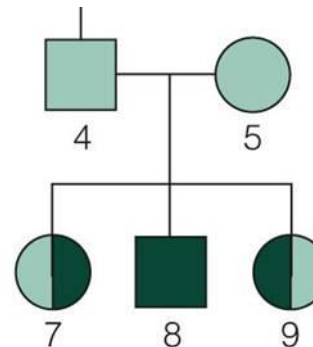
- A phenomenon in which a single phenotype or genetic disorder may be caused by any one of a multiple number of alleles or non-allele (locus) mutations
 - **Allelic heterogeneity** – different mutations within a **single gene locus** (forming multiple alleles of that gene) cause the same phenotypic expression
 - 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



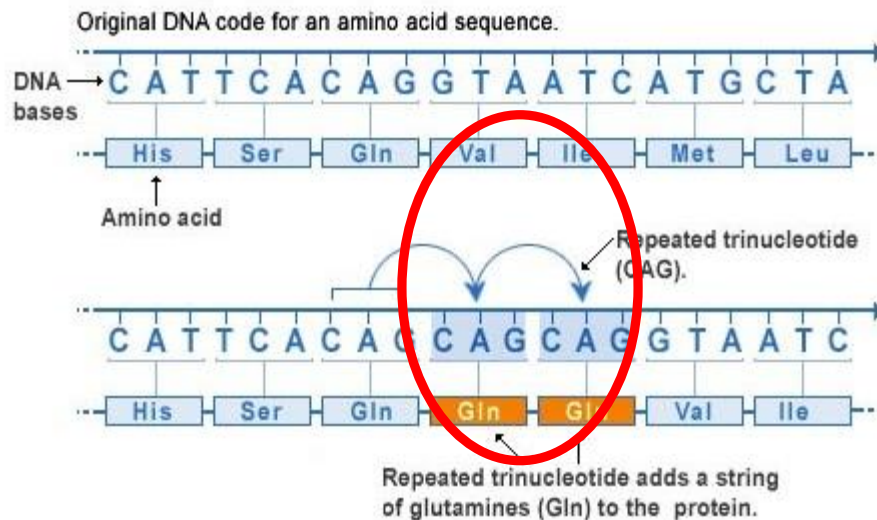
Expressivity

- 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 Penetrance	+ / - Affected
>40	Full Penetrance	Affected

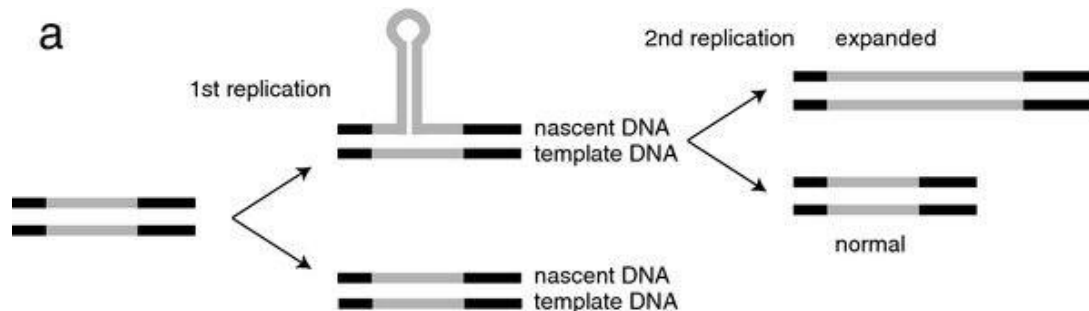
Why do Triplet repeat mutations get worse over generations*??

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

Mutation caused by denaturation and misplacement of the strands

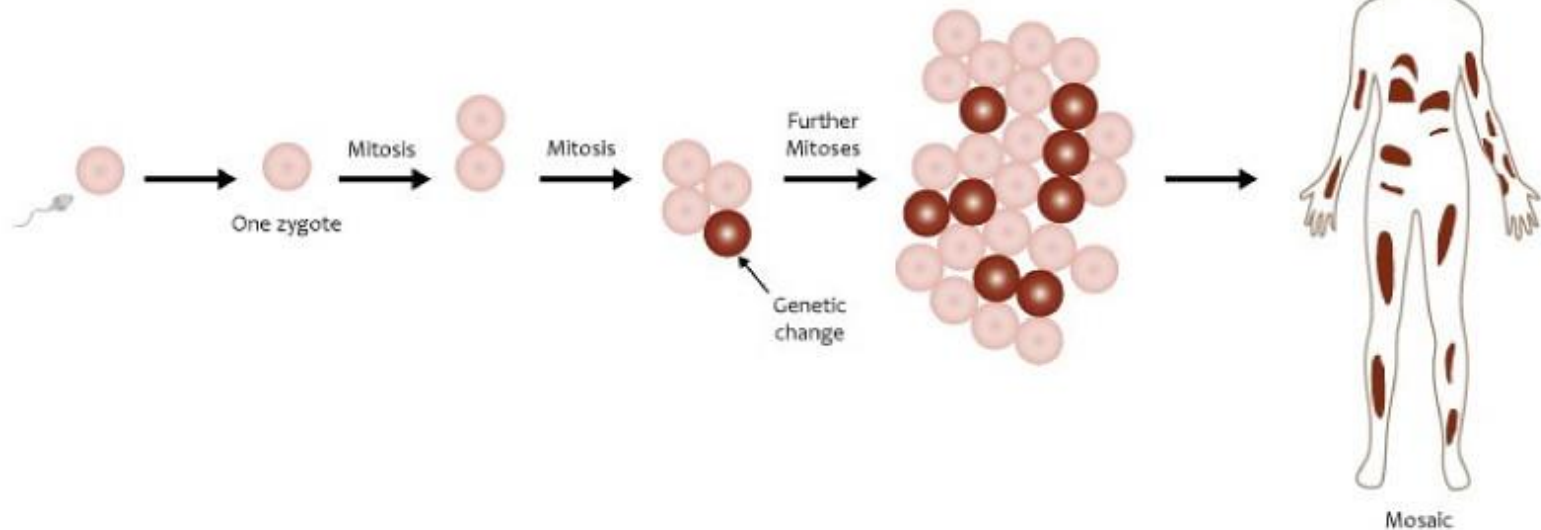
**-Mispairing of the complimentary strands
In high sequence repeat regions**

Replication slippage



Germline mosaicism

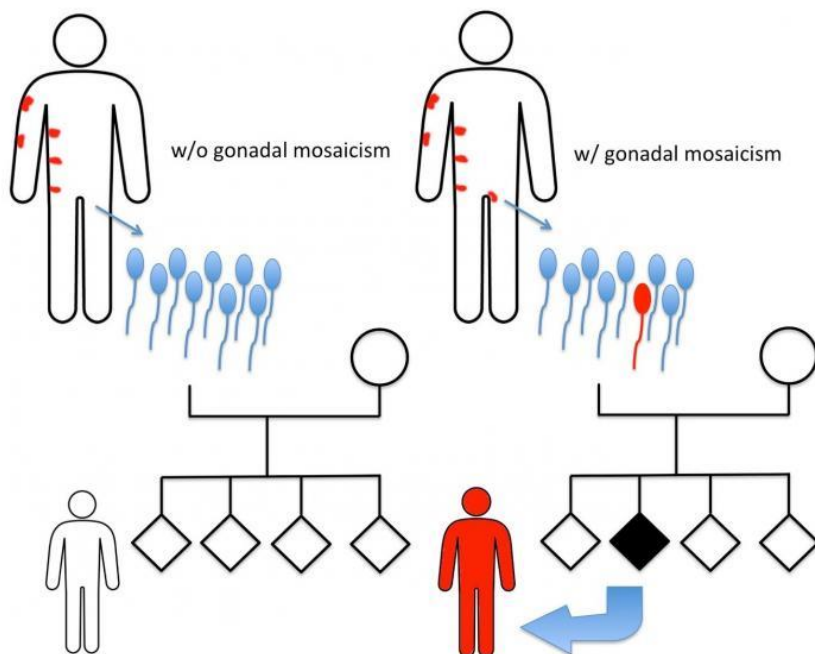
- **Mosaicism** – the presence of two or more populations of cells with different genotypes in one individual who has developed from a single fertilized egg



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)



Himalayan rabbit standard coat
Raised 20deg

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



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



Raised 20deg



Raised in cold <15deg

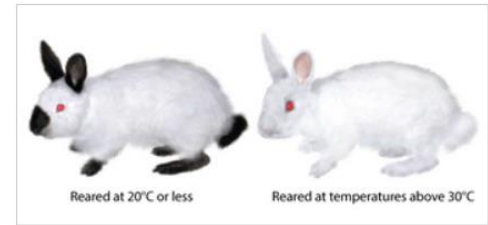


Figure 1: A pigment gene is influenced by temperature.

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 suppressed

<20°C = black coat: melanin expressed

>25°C = white coat: melanin fully suppressed

Think about this one for prac 1!

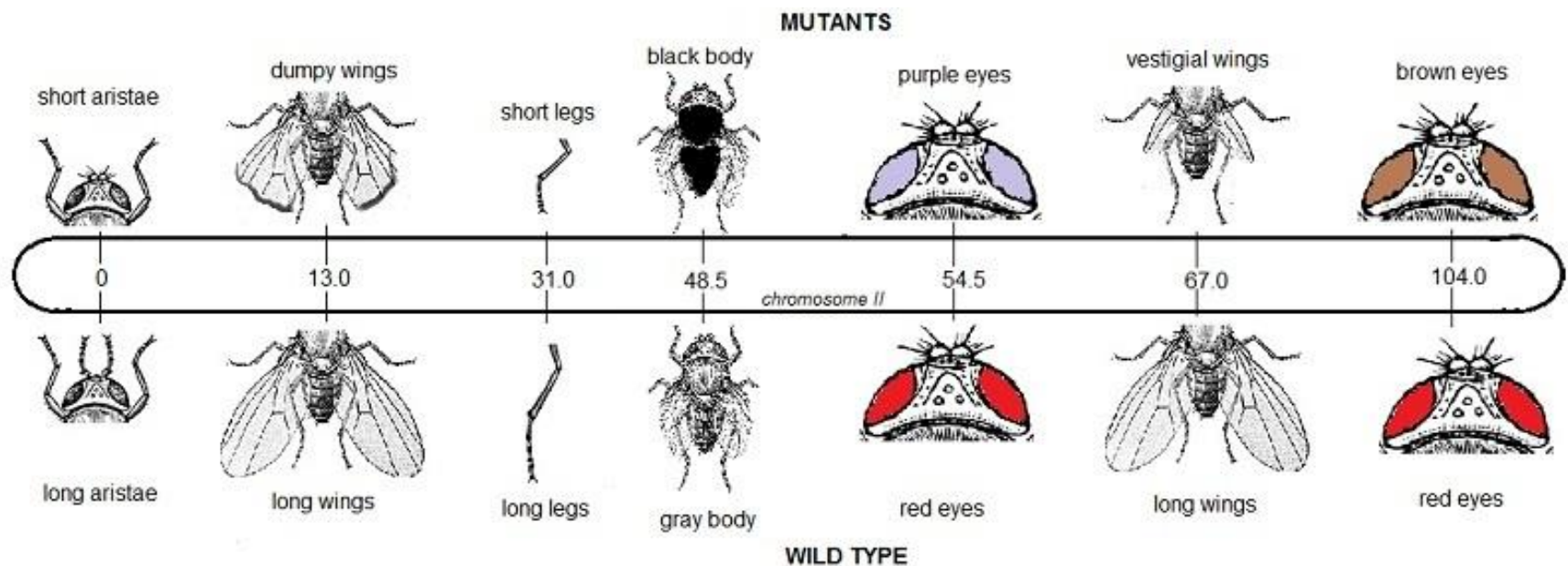


Genomic imprinting

- 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

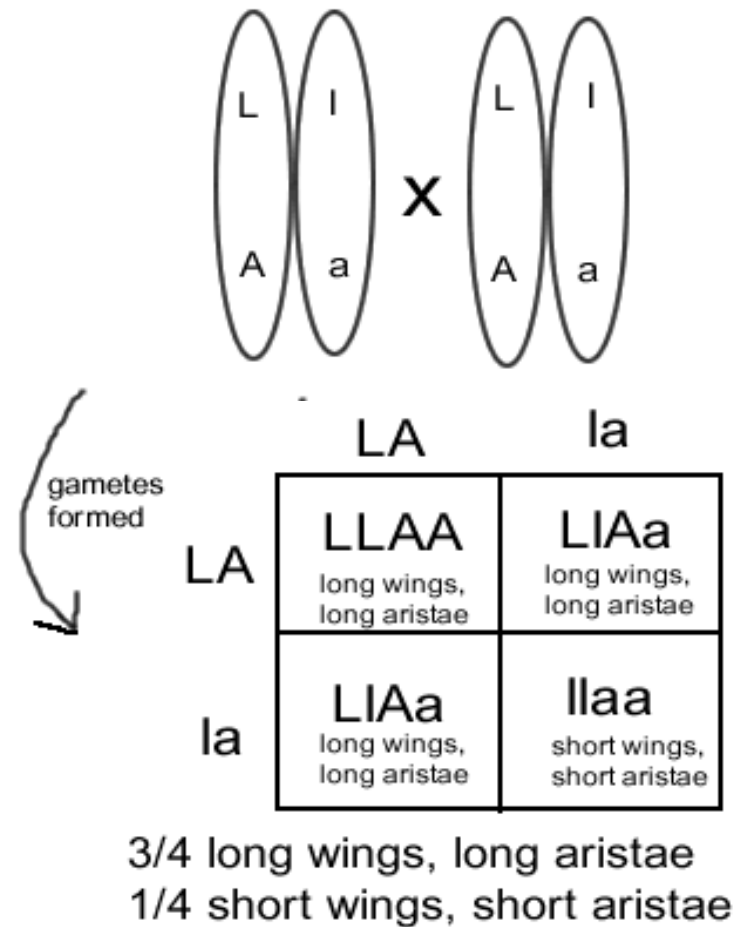
- Two or more genes located on the same chromosome that do not show independent assortment and tend to be inherited together



- 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 (Ll) 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.**



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

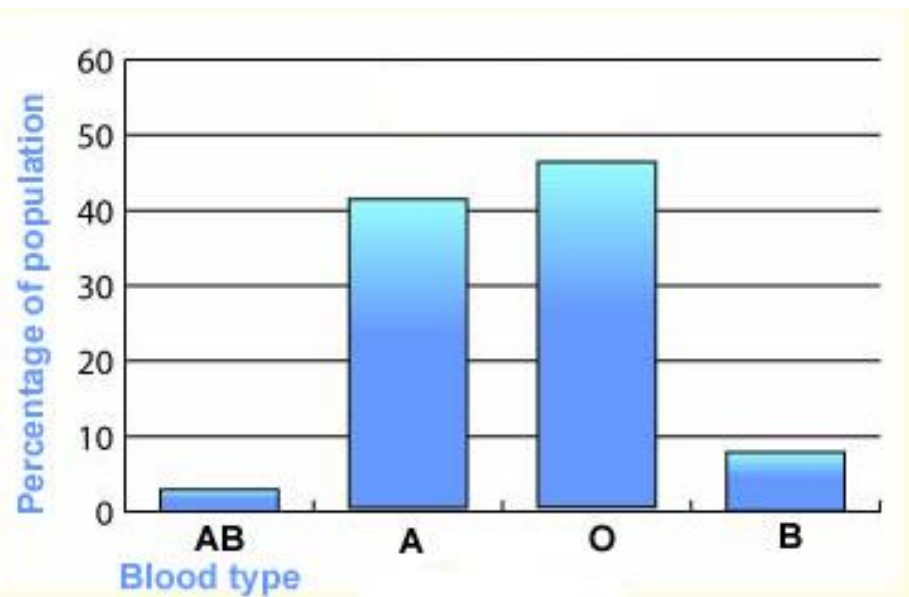
Discontinuous variation



purple flower



white flower



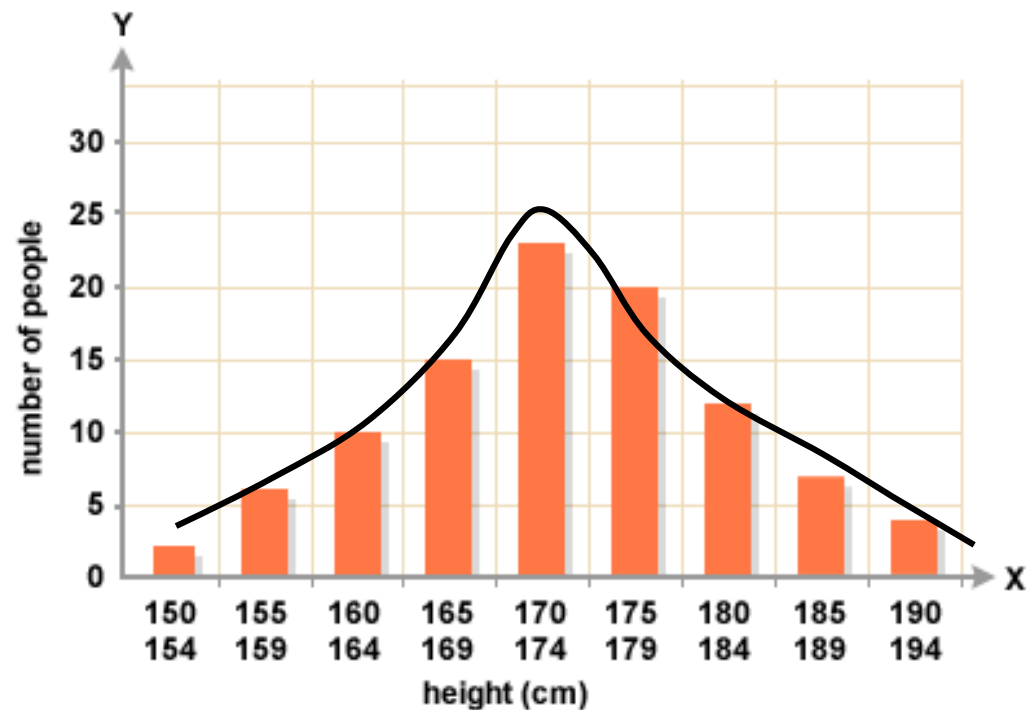
Continuous variation

- **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 **two or more genes** and **one or more environmental factors**
 - 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

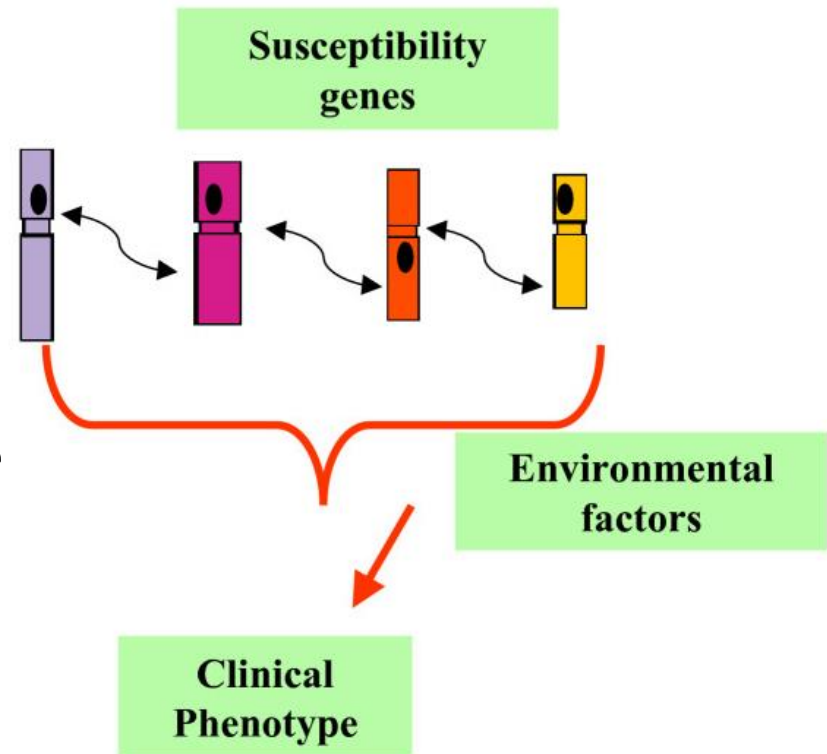


How are traits defined?

- **Complex traits**

- Traits controlled by multiple genes and the interaction of environmental factors 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