

BIO-373
Genetics & Genomics

**Extension of
Mendelian genetics**

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Why do we need to go beyond Mendelian genetics?



Why do we need to go beyond Mendelian genetics?

Many phenotypes cannot be explained by Mendel's transmission laws

➔ Co-dominance, incomplete dominance, epistasis, multiple alleles, lethal alleles, imprinting...

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 - 2 Codominance
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 - 4 Lethal alleles and essential genes
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1. Incomplete (or partial) dominance

Incomplete (or partial) dominance

- Neither allele is dominant → intermediate phenotype
 - **Each genotype has its own phenotype**
 - Botanical example: snapdragon (belle-de-nuit ou mufler des jardins)
 - Cross between red and white flowers
 - F_1 offspring: pink flowers
 - F_2 generation: 1/4 red, 1/2 pink, 1/4 white
 - **Phenotypic and genotypic ratios are the same**
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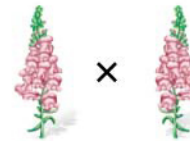
$R^1 R^1$ red $\times$ $R^2 R^2$ white P_1



$R^1 R^2$ pink F_1



$R^1 R^2 \times R^1 R^2$ $F_1 \times F_1$



$\frac{1}{4} R^1 R^1$ red
 $\frac{1}{2} R^1 R^2$ pink
 $\frac{1}{4} R^2 R^2$ white F_2



2. Codominance

Codominance

- No dominance or recessiveness
 - No incomplete or blending
 - Joint expression of both alleles in a heterozygote
 - Possible when 2 alleles have distinct phenotypic expression
-

Example of codominance in humans

- **MN Blood Group**
- M and N are two distinct forms of a glycoprotein that can be expressed on red blood cells
- Alleles L^M and L^N

Genotype

$L^M L^M$

$L^M L^N$

$L^N L^N$

Phenotype

M

MN

N

3. Multiple alleles of a gene

Multiple alleles

- Mutations can modify a gene in various ways → 3 or more alleles of the same gene might exist
 - The resulting mode of inheritance is unique
 - This can only be studied in populations and not in single individuals (who always carry two alleles per genetic locus)
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ABO blood groups

- Three possible alleles of a single gene (*I* for isoagglutinogen)
 - *I^A*
 - *I^B*
 - *i*
 - *I^A* and *I^B*: encode the A and B antigens that are present at the surface of red blood cells
 - *i*: no antigen produced
-

Genotype	Antigen	Phenotype
$I^A I^A$	A }	A
$I^A i$	A }	
$I^B I^B$	B }	B
$I^B i$	B }	
$I^A I^B$	A, B	AB
$i i$	Neither	O

I^A and I^B are dominant to i
 I^A and I^B are codominant to each other

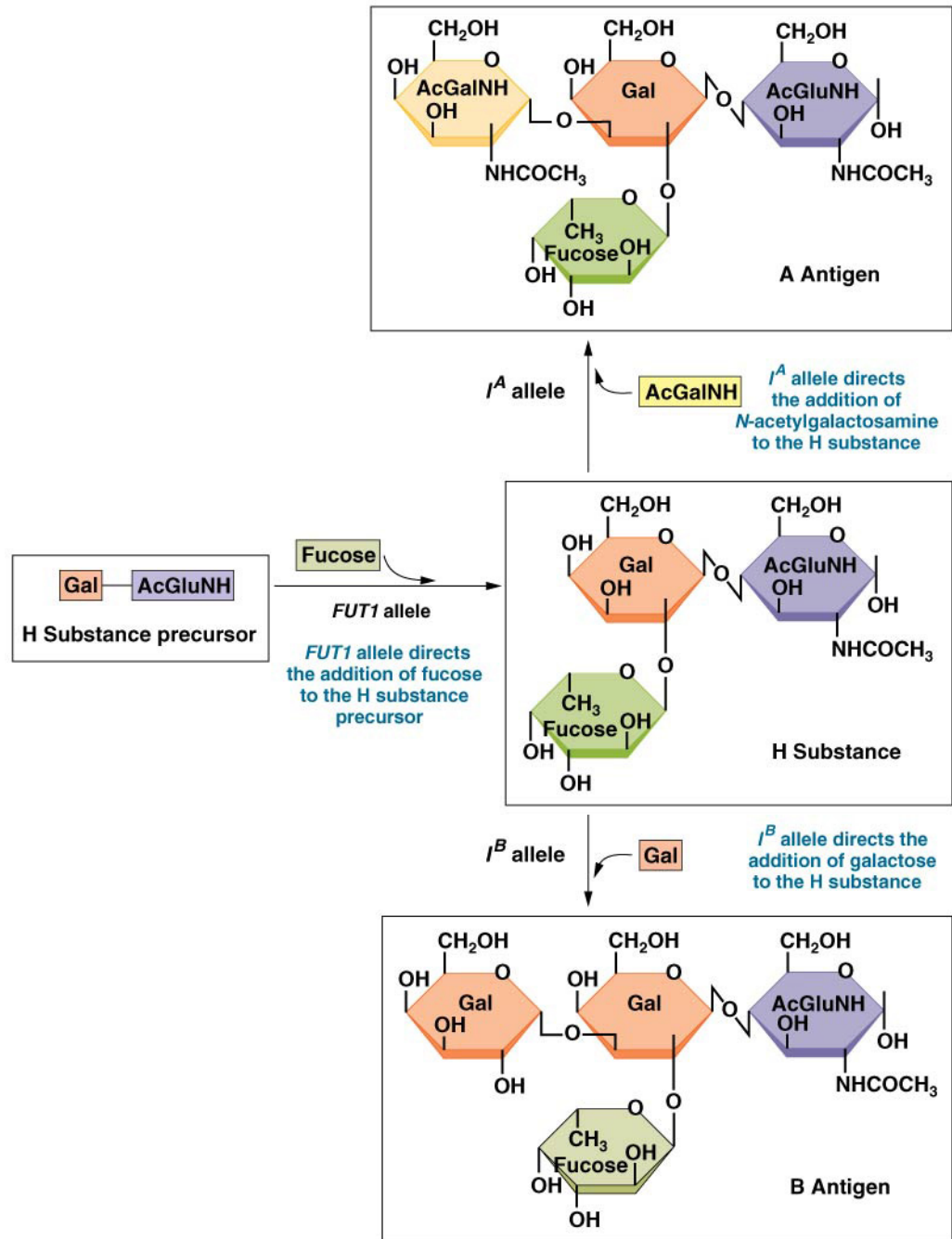
ABO blood groups

- **A and B antigens**

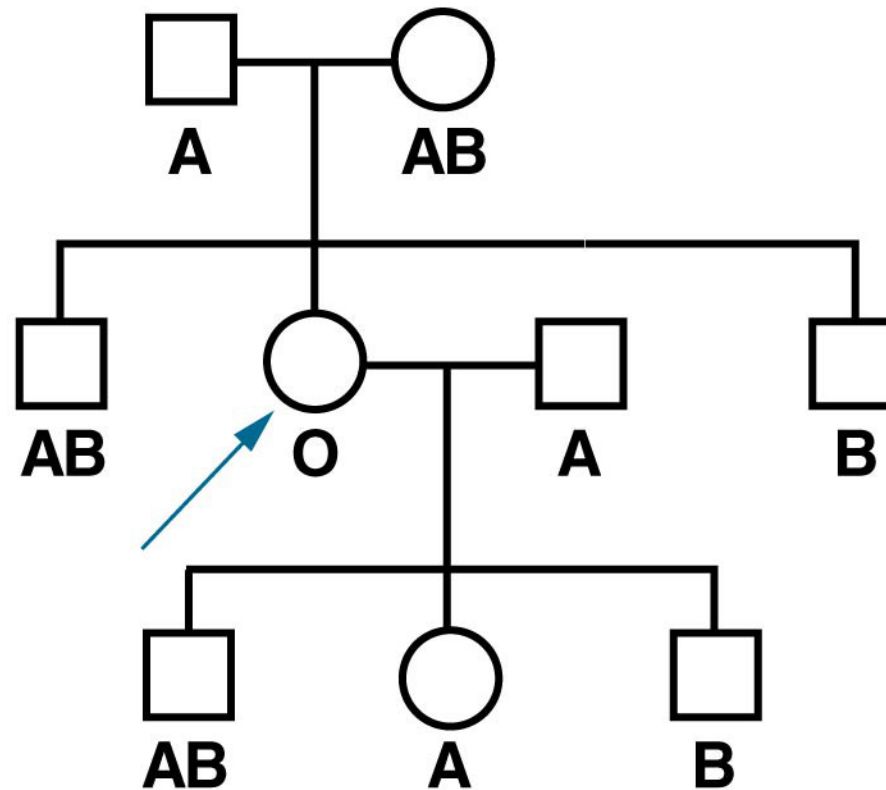
- Carbohydrate groups (N-Acetylgalactosamine for A antigen or galactose for B antigen) bound to H substance on red blood cells

- **H substance**

- Basic sugar layer at the surface of red blood cells
 - O blood types (*ii*) only have the H substance (O is for “Ohne,” auf Deutsch)
-



A more complex case



Bombay phenotype

- Type O female, yet...
 - one parent has type AB blood and
 - She gave I^B allele to two children
 - Female found to be homozygous for a **FUT1** (fucosyl transferase) mutation
 - Prevents her from producing H substance
 - No base substrate to make A or B antigens
 - Results functionally in type O
-

Bombay phenotype

- One gene (**FUT1**) is masking the phenotypic effect of another one (**ABO**)
 - Very rare: 1:10'000 in India, 1:100'000 in Europe
 - Individuals with Bombay phenotype can only receive blood transfusion from Bombay donors
-

4. Lethal alleles and essential genes

Essential genes and lethal alleles

- Absolutely required for survival
 - May contain lethal alleles
 - **Dominant lethal allele:** the presence of a single copy is incompatible with life
 - **Recessive lethal allele:** tolerated in heterozygous state
-

Dominant lethal allele in humans

- Huntington disease

- Characterized by progressive degeneration of nervous system, dementia, and death
- Due to allele *H* (abnormal number of CAG trinucleotide repeat in *HTT* gene)
- Autosomal dominant = Hh genotype

<https://www.youtube.com/watch?v=JL9Y3P870jU>

Recessive lethal allele in mice

- **Agouti gene**, responsible for coat color
 - Agouti allele A
 - Mutant *yellow* allele A^Y

Crosses				
(A) agouti	×	agouti	→	all agouti
(B) yellow	×	yellow	→	2/3 yellow: 1/3 agouti
(C) agouti	×	yellow	→	1/2 yellow: 1/2 agouti

Cross A

P₁ **AA** × **AA**
agouti agouti



F₁ **AA**
agouti



all agouti
(All survive)

agouti mouse



Cross B

P₁ **AA^Y** × **AA^Y**
yellow yellow



F₁ **AA** **AA^Y**
agouti yellow
A^YA **A^YA^Y**
yellow lethal



2/3 yellow
1/3 agouti
(Survivors)

yellow mouse



Cross C

P₁ **AA** × **AA^Y**
agouti yellow



F₁ **AA** **AA^Y**
agouti yellow



1/2 agouti
1/2 yellow
(All survive)

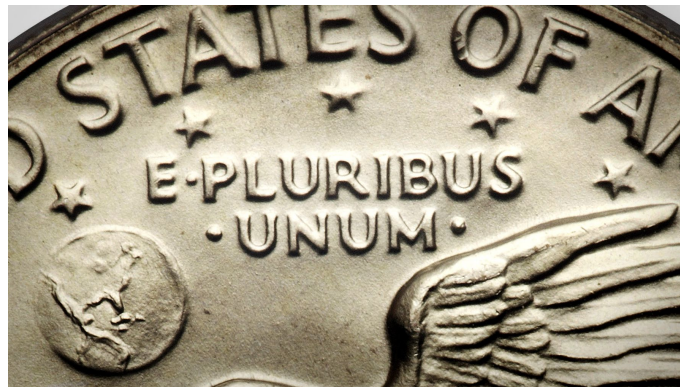
Recessive lethal allele in mice

- **Mutant allele (A^Y)**
 - Behaves dominantly to normal allele to control coat color
 - Behaves as homozygous recessive lethal allele
- Genotype $A^Y A^Y$ does not survive

5. Impact of multiple genes on a phenotype

E pluribus unum (“out of many, one”)

- Most phenotype result from the influence of multiple genes
- **Genic interaction:** several genes influence a trait; doesn't imply direct interaction
- **Epistatic interaction:** direct interaction between alleles of different genes



Example of genic interaction

- **Hereditary deafness**

- Many genes are involved in ear formation
 - These genes interact to produce a common phenotype
 - Many mutations can interrupt development and lead to hereditary deafness
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Example of epistatic interaction

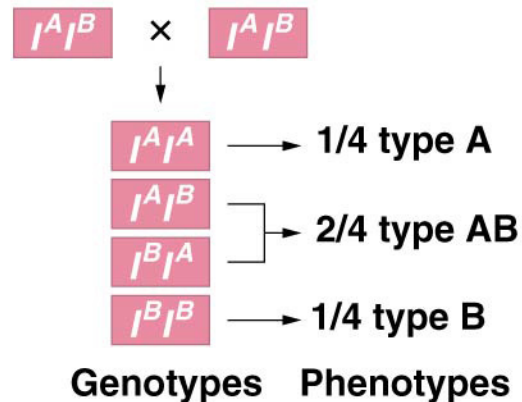
■ Back to Bombay...

- First locus masks expression of second locus
- Homozygous mutation in *FUT1* gene masks expression of I^A and I^B alleles
- A and B antigen forms only when individual has at least one wild-type allele

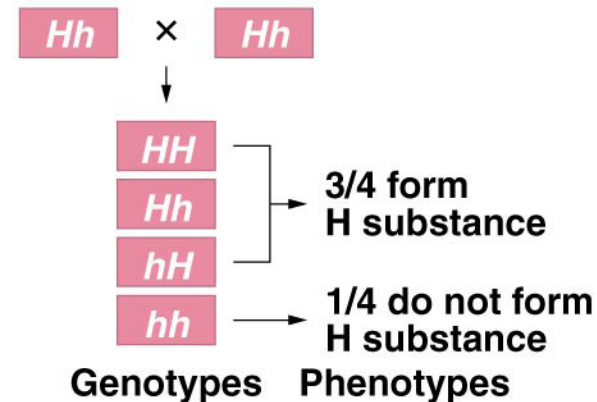
Cross between two individuals heterozygous at the two loci involved in ABO group determination (figure)

$$I^A I^B Hh \times I^A I^B Hh$$

Consideration of blood types



Consideration of H substance



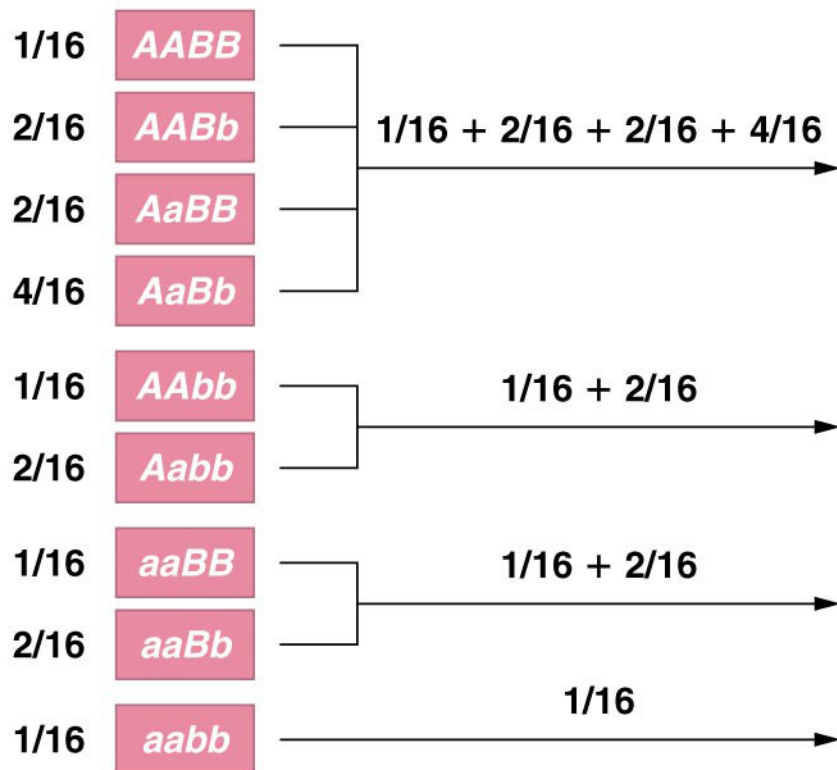
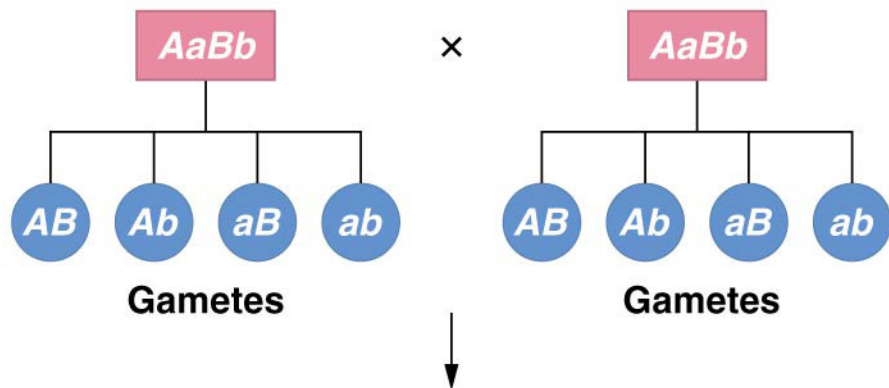
Consideration of both gene pairs together

Of all offspring	Of all offspring	Final probabilities
$\downarrow$	$\downarrow$	$\downarrow$
1/4 type A	3/4 form H substance	$\rightarrow 3/16$ type A
	1/4 do not form H substance	$\rightarrow 1/16$ type O
2/4 type AB	3/4 form H substance	$\rightarrow 6/16$ type AB
	1/4 do not form H substance	$\rightarrow 2/16$ type O
1/4 type B	3/4 form H substance	$\rightarrow 3/16$ type B
	1/4 do not form H substance	$\rightarrow 1/16$ type O

Final phenotypic ratio = 3/16 A: 6/16 AB: 3/16 B: 4/16 O

Impact of epistasis on phenotypic ratio

- Epistatic interactions modify the phenotypic ratios expected under Mendel's transmission laws.
 - Dihybrid cross: 9:3:3:1
 - Effect possible on one or more of four phenotypic categories
-



Dihybrid ratio	Modified ratios				
$9/16 A-B-$	$9/16$	$12/16$	$9/16$	$9/16$	$15/16$
$3/16 A-bb$	$3/16$		$7/16$	$6/16$	
$3/16 aaB-$	$4/16$	$3/16$		$1/16$	$1/16$
$1/16 aabb$		$1/16$			

Dominant epistasis

- Dominant allele at one locus masks an allele at second locus
- Summer squash fruit color
 - Determined by 2 loci, A and B
 - Dominant allele A = **white** fruit
 - Regardless of second loci allele
 - Absence of A allele (*aa*)
 - Genotypes *BB*, *Bb* = **yellow** fruit
 - Genotype *bb* = **green** fruit



Dominant epistasis

$$F_1: AaBb \times AaBb$$

↓

F ₂ Ratio	Genotype	Phenotype	Final Phenotypic Ratio
9/16	<i>A-B-</i>	white	12/16 white
3/16	<i>A-bb</i>	white	
3/16	<i>aaB-</i>	yellow	3/16 yellow
1/16	<i>aabb</i>	green	1/16 green

Recessive epistasis

- Homozygous recessive allele at one locus masks an allele at second locus
 - Mouse coat color (again...):
 - *A* allele (dominant): agouti phenotype
 - *B* allele: black pigment
 - *bb* genotype: no black pigment, even if *A* or *a* alleles present → mouse is albino
 - *bb* genotype masks the expression of the *A* allele: **recessive epistasis**
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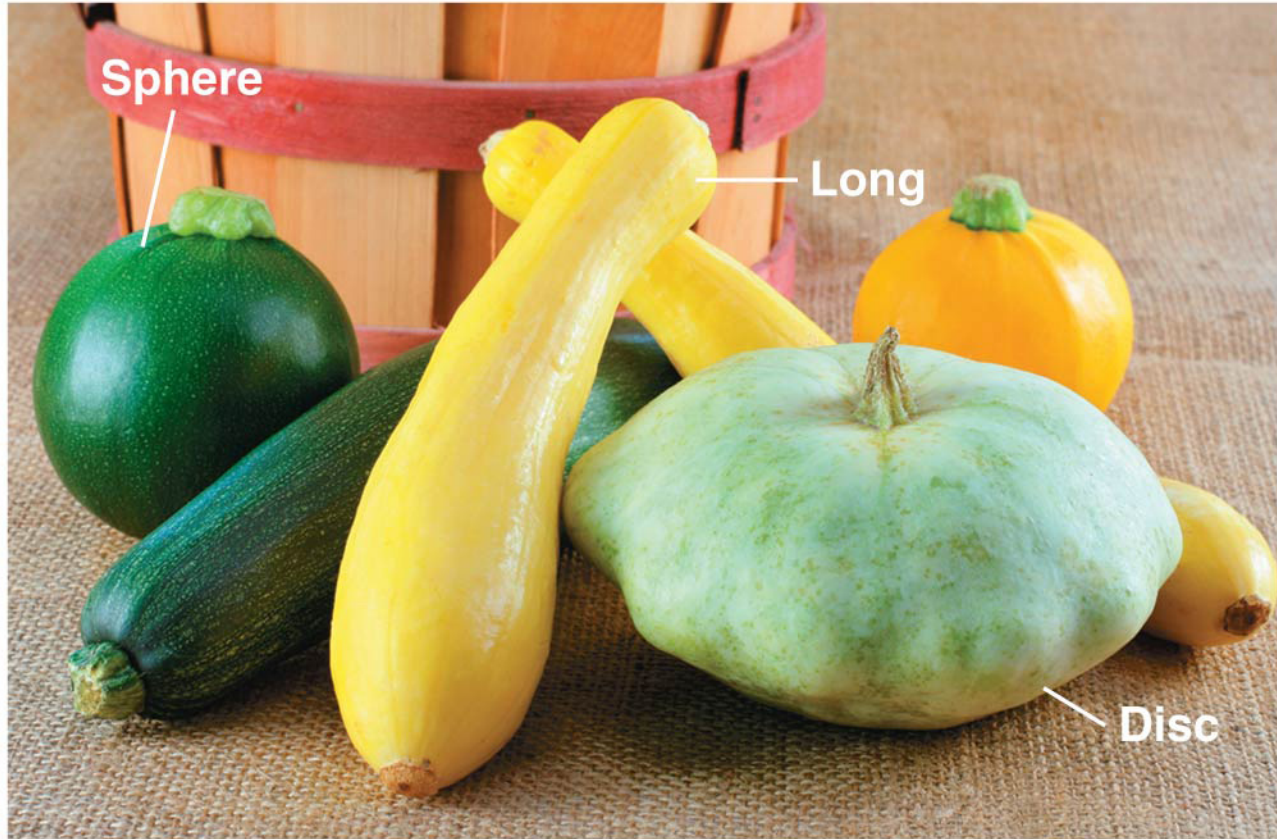
Recessive epistasis

$$F_1: AaBb \times AaBb$$



F ₂ Ratio	Genotype	Phenotype	Final Phenotypic Ratio
9/16	$A-B-$	agouti	9/16 agouti
3/16	$A-bb$	albino	
3/16	$aaB-$	black	3/16 black
1/16	$aabb$	albino	4/16 albino

Epistasis resulting in a novel phenotypes



Epistasis resulting in a novel phenotypes

- Shape of summer squash: **disc-shaped** fruit ($AABB$) crossed with **long** fruit ($aabb$)
 - F_1 : all disc-shaped fruit
 - F_2 : includes the two parental phenotypes and a new **spherical** variants
 - Dominant alleles at 2 loci → disc-shaped
 - Dominant allele at 1 locus → spherical
 - Only recessive alleles → long
-

Epistasis resulting in a novel phenotypes

$$\begin{array}{c} F_1: AaBb \times AaBb \\ \text{disc} \quad \text{disc} \\ \downarrow \end{array}$$

F ₂ Ratio	Genotype	Phenotype	Final Phenotypic Ratio
9/16	$A-B-$	disc	9/16 disc
3/16	$A-bb$	sphere	6/16 sphere
3/16	$aaB-$	sphere	
1/16	$aabb$	long	1/16 long

6. Pleiotropy

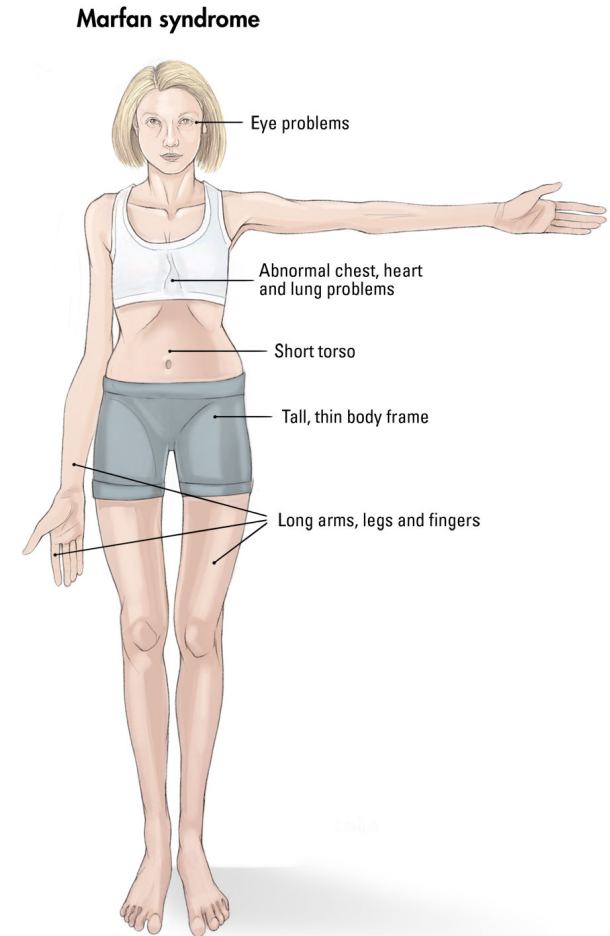
Pleiotropy

- We just saw that several genes can influence a single phenotype
 - The opposite situation, where a single gene has multiple phenotypic effects, is called **pleiotropy**
 - Extremely frequent
-

Example of pleiotropy in humans

■ Marfan syndrome

- Autosomal dominant allele in fibrillin gene, encoding a protein important for connective tissues
- Multiples phenotypic consequences:
 - Lens dislocation
 - Aortic aneurism
 - Extension of long bones
 - ...



7. Gene x environment interactions

Phenotype = direct expression of genotype?

- **A phenotype usually results from multiple influences**
 - Core gene
 - Other genotypes (genic and epistatic influences)
 - Environment
 - Often difficult to disentangle the respective influences of genes and environment ⇔ long-standing debate between “innate” and “acquired”
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Penetrance and expressivity

- **Penetrance**

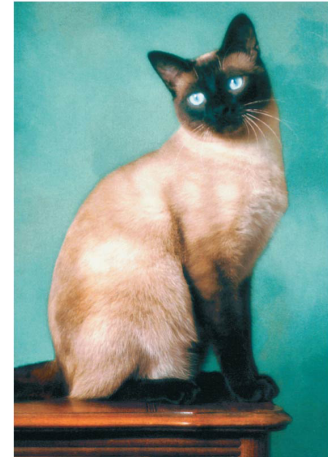
- Percentage of expression of the mutant genotype in a population

- **Expressivity**

- Range of expression of mutant phenotype
 - Result of genetic background differences and/or environmental effects
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Causes of variable expressivity

- **Temperature effects**
 - Evening primrose
 - Red flowers at 23°C
 - White flowers at 18°C
 - Siamese cats and Himalayan rabbits
 - Darker fur on cooler areas of body (tail, feet, ears)
 - Enzymes responsible for pigment formation lose catalytic function at higher temperature



Causes of variable expressivity

- **Temperature-sensitive mutations**

- Known in viruses, bacteria, fungi, and *Drosophila*
 - Mutant allele expresses mutant phenotype at one temperature, wild-type phenotype at another
 - Broadly used in viral genetics: temperature-sensitive mutations are easily induced and isolated in viruses
-

Causes of variable expressivity

■ Nutritional effects

- Some alleles prevent the synthesis of essential nutrients in microbes, they are called **auxotrophs**
 - Phenotype expression can be modulated by diet, which is very useful for *in vitro* studies
 - In humans, diseases are caused by alleles that stops the metabolization of some substances:
 - **Phenylketonuria**: loss of enzyme to metabolize amino acid phenylalanine, severe problems unless low-Phe diet
 - **Galactosemia, lactose intolerance, ...**
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