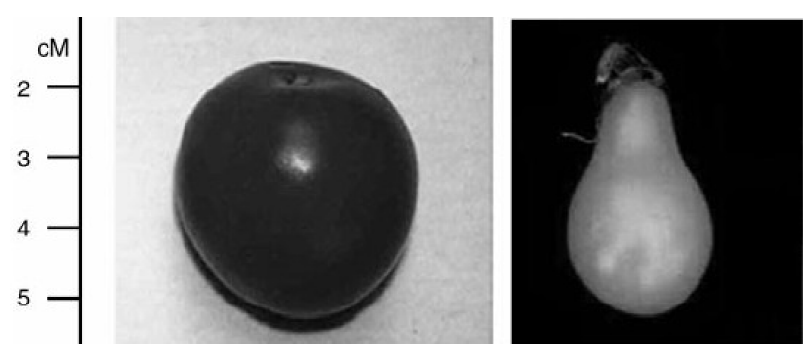
Mapping the genes underlying phenotypic changes of interest

Virginie Courtier-Orgogozo Institut Jacques Monod, Paris

Tomato shape

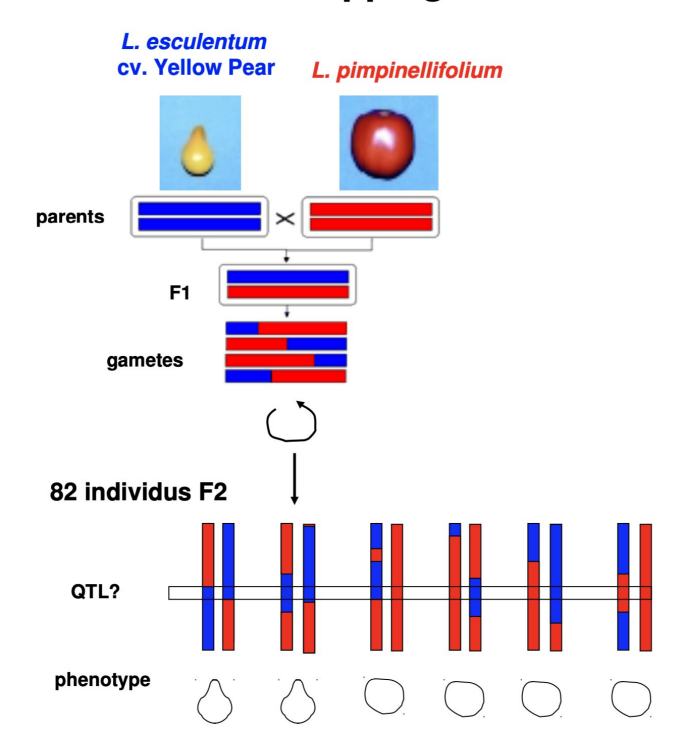


Lycopericon esculentum

Lycopersicon esculentum cv. Yellow Pear

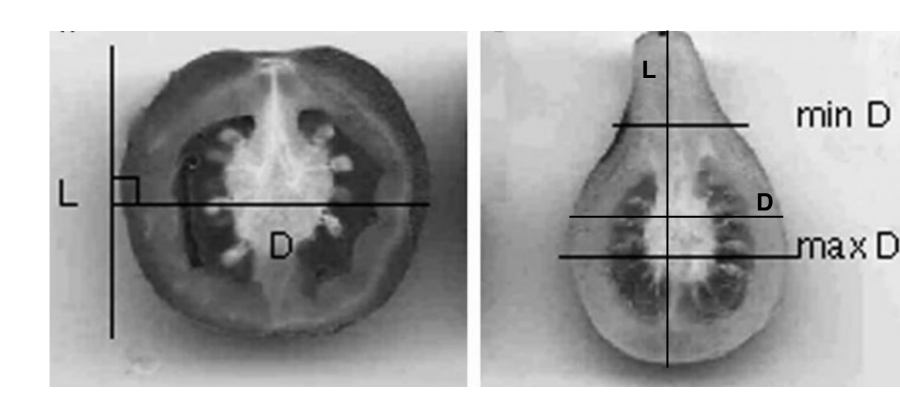
(Ku et al., 1999; Liu et al., 2002)

QTL mapping

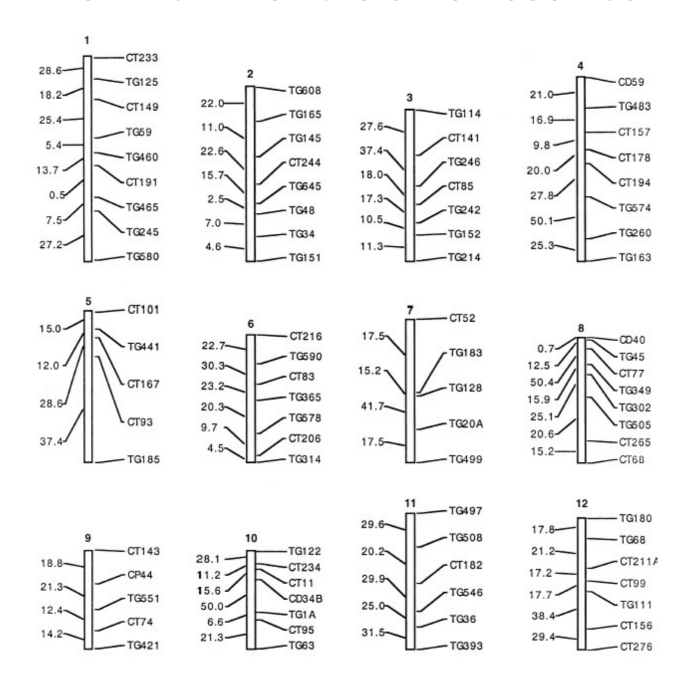


Quantitative measure of the phenotype

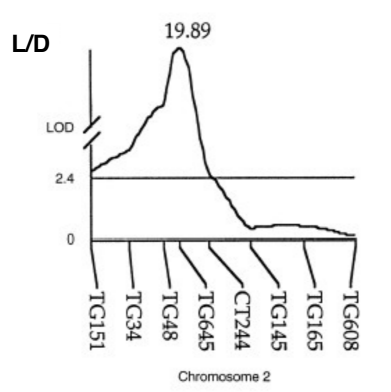
Measure of 2 indexes L/D and Dmin/Dmax for 10 fruits per plant L/D : L= length, D = diameter at equator Dmin/Dmax

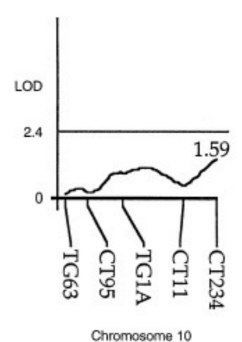


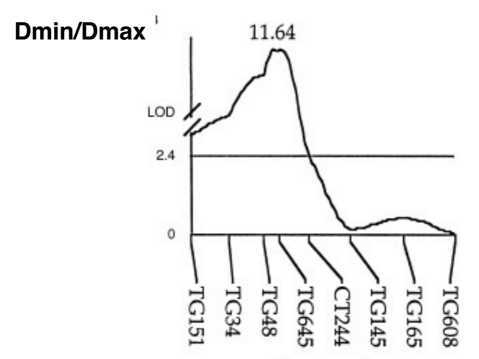
82 molecular markers on the 12 tomato chromosomes



One major locus near marker TG645







Chromosome 2

responsible for 67% of L/D variance

allele YP = recessive

Two main files

Markers file

```
-start
-Chromosome 1
CF5475
              0.4
              24.7
CF5573
CT7895
              41.0
CT8903
              59.0
CF5613
              67.7
CT7892
              76.0
CT890
              89.0
              39.0
CT233
Telomere
              50.0
-Chromosome 2
CF5671
              0
CF5675
              10.4
CF5673
              34.7
CT789
              41.0
CT890
              89.0
CT567
              115.0
Telomere
              130.0
```

Genotypes and phenotype(s) file

```
-start individuals markers
Ind 2 0 0 0 1 0 1
Ind 6 1 1 1 1 1 1 1 1
Ind 7 1 1 1 1 1 1 1 0 1 n n 1
-stop individuals markers
-start individuals traits 1 LoverD named
Ind 1
        5.5
Ind 2
        3.0
Ind 3
       4.0
Ind 4
       7.0
Ind 5
       6.5
Ind 6
       5.0
Ind 7
        3.5
        6.0
Ind 8
```

Simple linear regression for each marker

```
L/D of individual i = a + b.xi + \epsilon

xi = 0 if Le/Le, = 1 if Le/Lp, = 2 if Lp/Lp

a,b = best fit parameters (least square regression)

\epsilon assumed to have a normal distribution
```

Test Ho: b = 0 versus H1: b = estimated b

Likelihood ratio test statistic

$$D = -2(\ln(\text{likelihood for null model}) - \ln(\text{likelihood for alternative model}))$$
$$= -2\ln\left(\frac{\text{likelihood for null model}}{\text{likelihood for alternative model}}\right).$$

The probability distribution of the test statistic can be approximated by a chi-square distribution with (df1 - df2) degrees of freedom, where df1 and df2 are the degrees of freedom of models 1 and 2 respectively

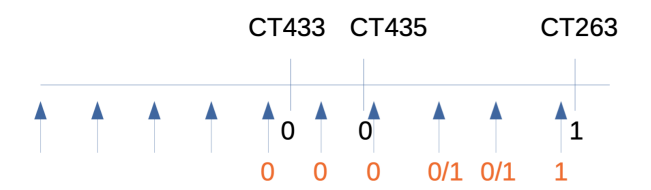
Interval mapping

L/D of individual i = a + b.xi + e

xi = indicator variable specifying the probabilities of an individual being in different genotypes for the tested position, constructed by flanking makers xi = 0 if Le/Le, = 1 if Le/Lp, = 2 if Lp/Lp

a,b = best fit parameters (maximum likelihood)

Test Ho: b=0 versus H1: b=estimated b



Interval mapping

```
L/D of individual i = a + b.xi + e
xi = indicator variable specifying the probabilities of an individual being
in different genotypes for the tested position, constructed by flanking makers
xi = 0 if Le/Le, = 1 if Le/Lp, = 2 if Lp/Lp
a,b = best fit parameters (maximum likelihood)
Test Ho: b=0 versus H1: b=estimated b
```

Composite Interval mapping

```
L/D of individual i = a + b.xi + c.xi + e
xi = indicator variable specifying the probabilities of an individual being
in different genotypes for the tested position, constructed by flanking makers
xi = 0 if Le/Le, = 1 if Le/Lp, = 2 if Lp/Lp
yi = 0 if Le/Le, = 1 if Le/Lp, = 2 if Lp/Lp at marker y
```

LOD score

L/D of individual i = a + b.xi + e

Test Ho: b = 0 *versus* H1: b = estimated b

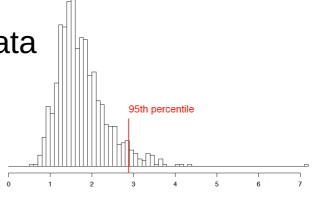
Lo = pr (data | no QTL) – phenotypes assumed to follow a normal distribution L1 = pr (data | QTL at tested position)

$$LOD = -\log \frac{L_0}{L_1}$$

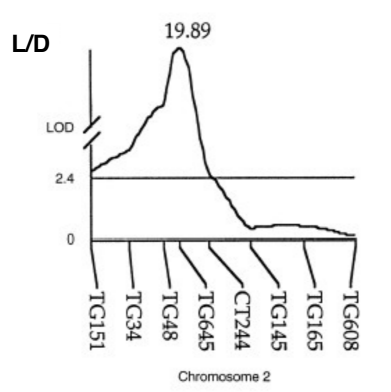
Significance threshold

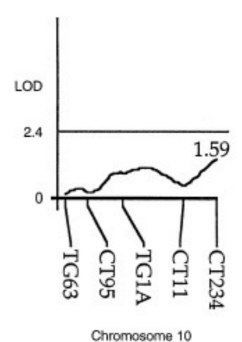
10,000 permutations of phenotype/genotype data

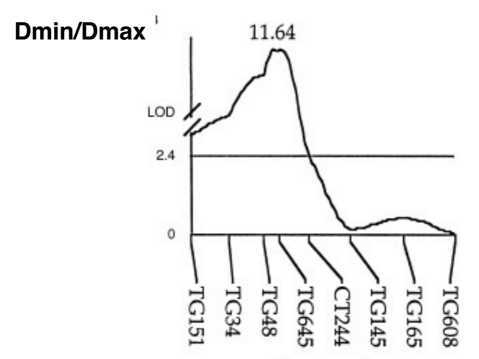
- → random distribution of LOD scores
 - → 1% or 5% significance threshold



One major locus near marker TG645







Chromosome 2

responsible for 67% of L/D variance

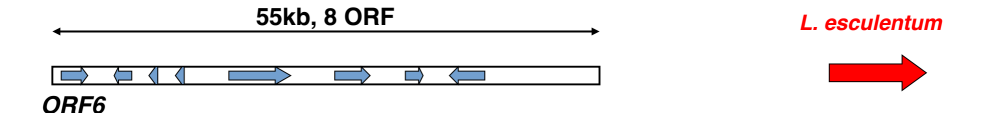
allele YP = recessive

BAC library (Bacterial Artificial Chromosomes)

contains genomic DNA fragments of 100-350kb

Screen of the library with marker TG645

BAC19 containing 105kb, 17 ORF (open reading frame) **BAC19** Design of new molecular markers to genotype the previously obtained recombinant tomato plants



Sequencing of the region in the 2 tomato varieties

1 SNP (single nucleotide polymorphism) et 1 indel (insertion-deletion) of 2bp in non-coding regions

L. esculentum cv. Yellow Pear



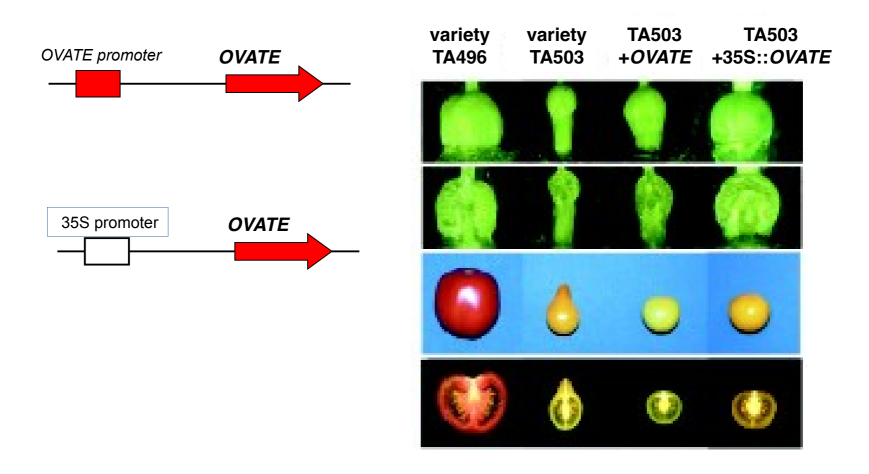
1 SNP in *ORF6*: G496T, stop codon stop, truncated protein with last 75 amino acids missing

Hypothesis: the causing gene is *ORF6* = *OVATE*

The causing gene is *OVATE/ORF6*

Same mutation in 3 other pear tomato varieties

Complementation of the mutation by transgenesis



OVATE = protein with NLS (nuclear localization signal), unknown function, expressed in developing fruits

Evolution of morphology in threespine sticklebacks



marine



Paxton Lake, Canada

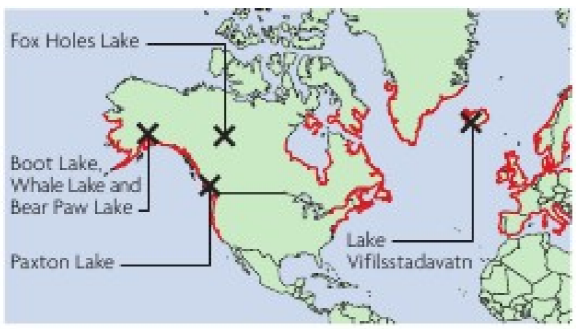
Gasterosteus aculeatus

Marine fishes with robust pelvis = ancestral

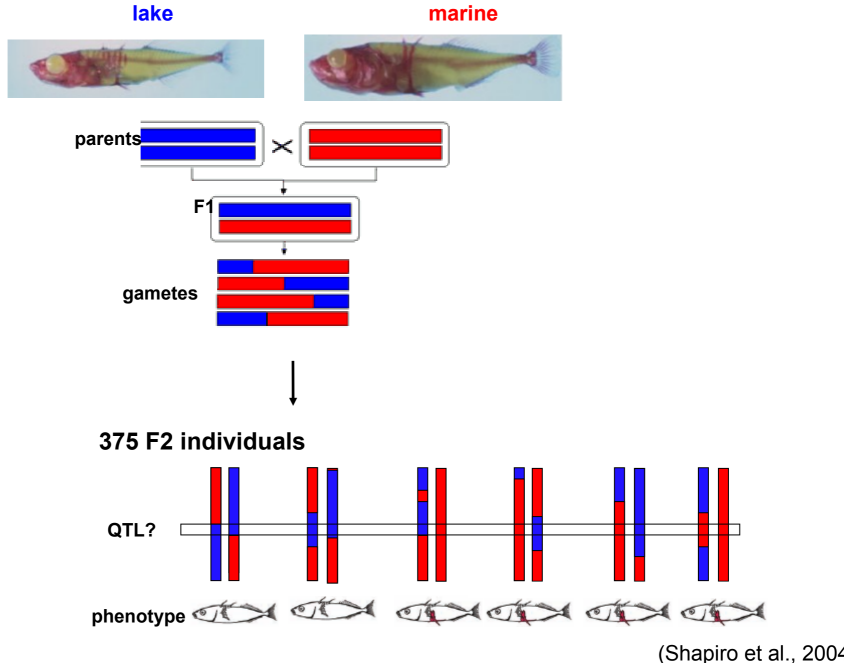
Freshwater fishes with reduced pelvic structures = derived, independently at least 20 times

- limited calcium availability
- absence of gape-limited predatory fishes
- predation by grasping insects

Last glacier retreat = 10 000 – 20 000 years ago



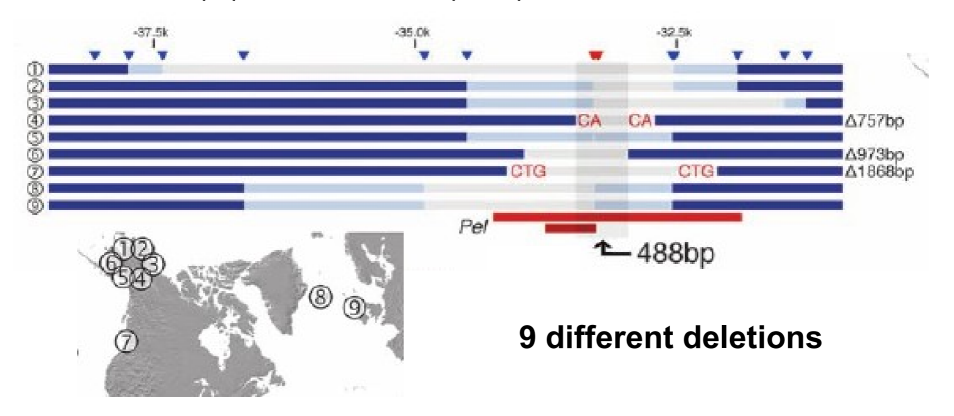
QTL mapping



Several independent deletions in the cis-regulatory region of *Pitx1*

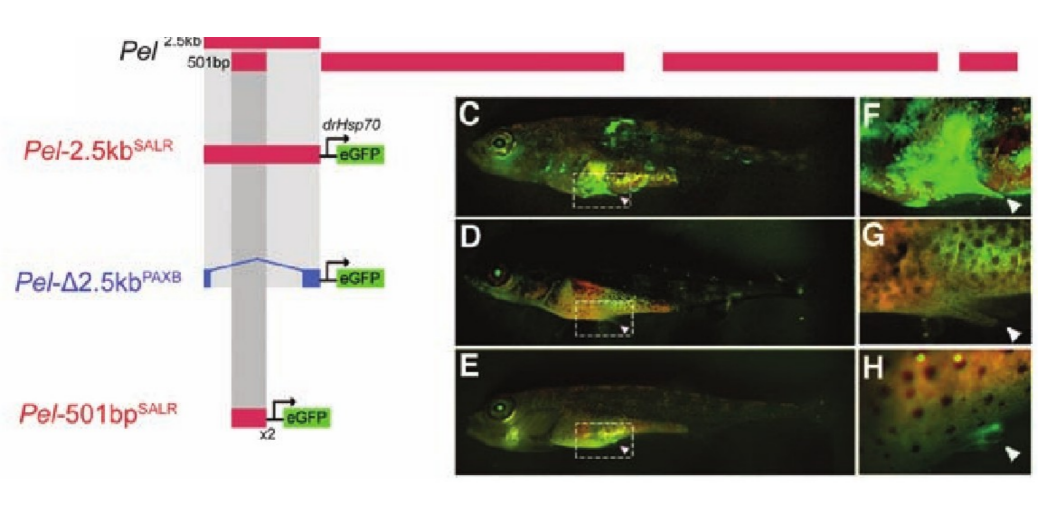
Region sequenced in two lake populations: a 2-kb deletion in one and a 757-bp deletion in the other one

SNP genotyping in 13 populations with reduced pelvis and in 21 populations with complete pelvis

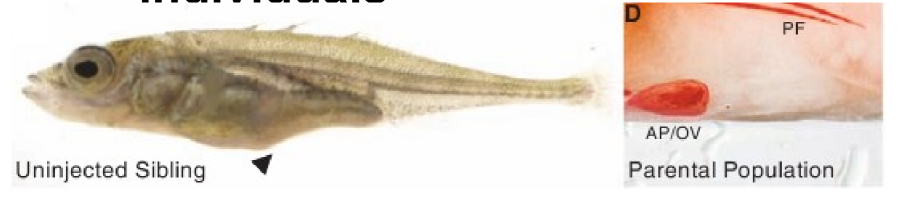


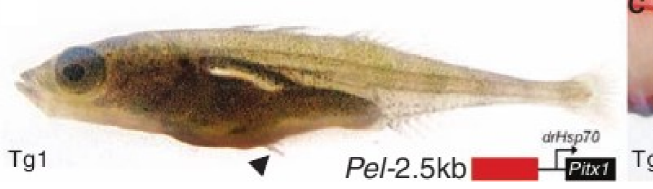
Test of *Pitx1* cis-regulatory regions

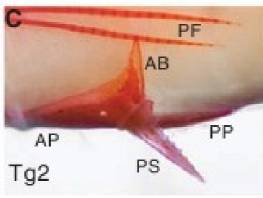




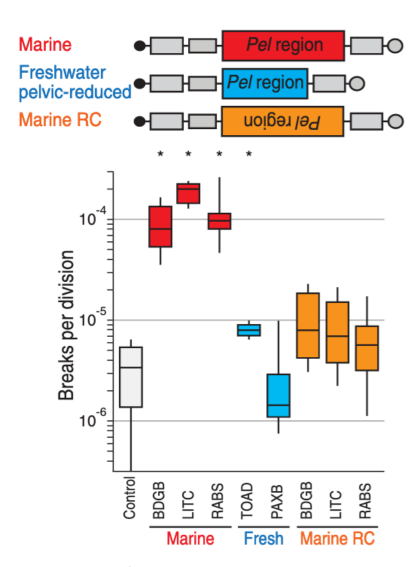
Rescue of a pelvis in freshwater individuals

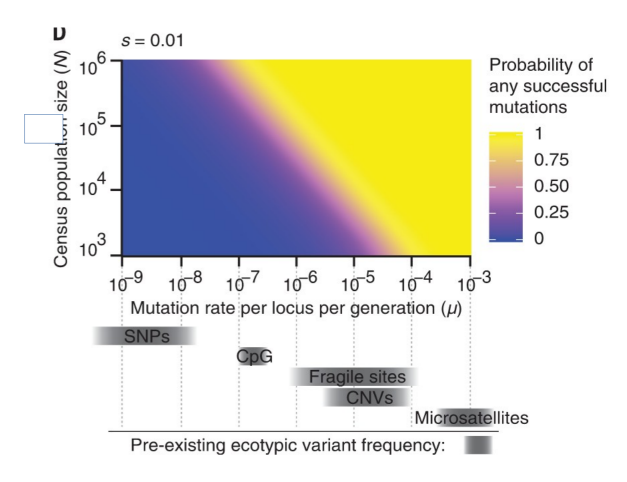


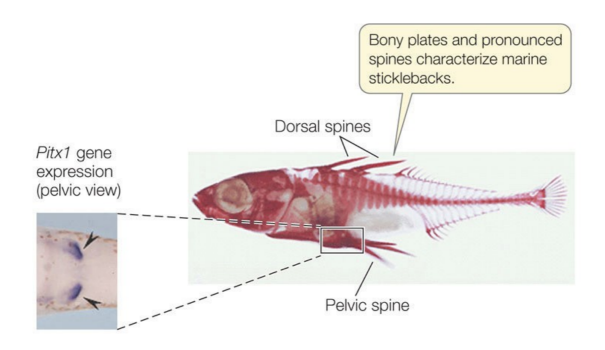


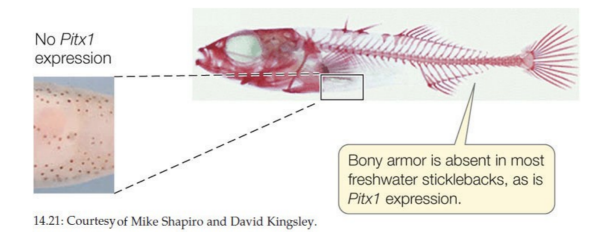


Pitx1 is in a fragile DNA region







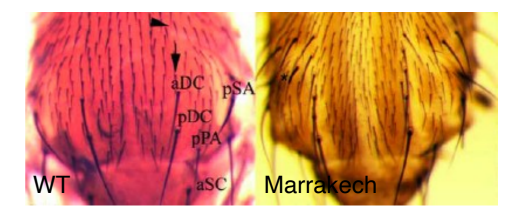


Evolution of extra bristles

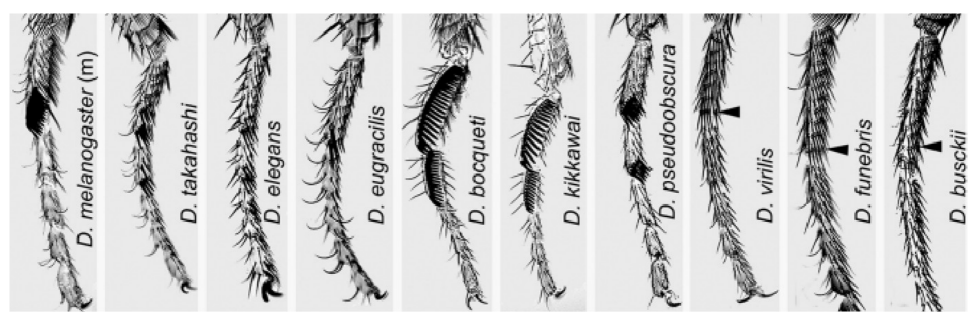
Interspecific change in *D. quadrilineata*

D. melanogaster D. quadrilineata

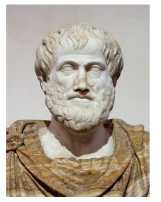
Intraspecific change in *D. melanogaster*



Finding genetic rules on bristle evolution



Randsholt and Santamaria 2008

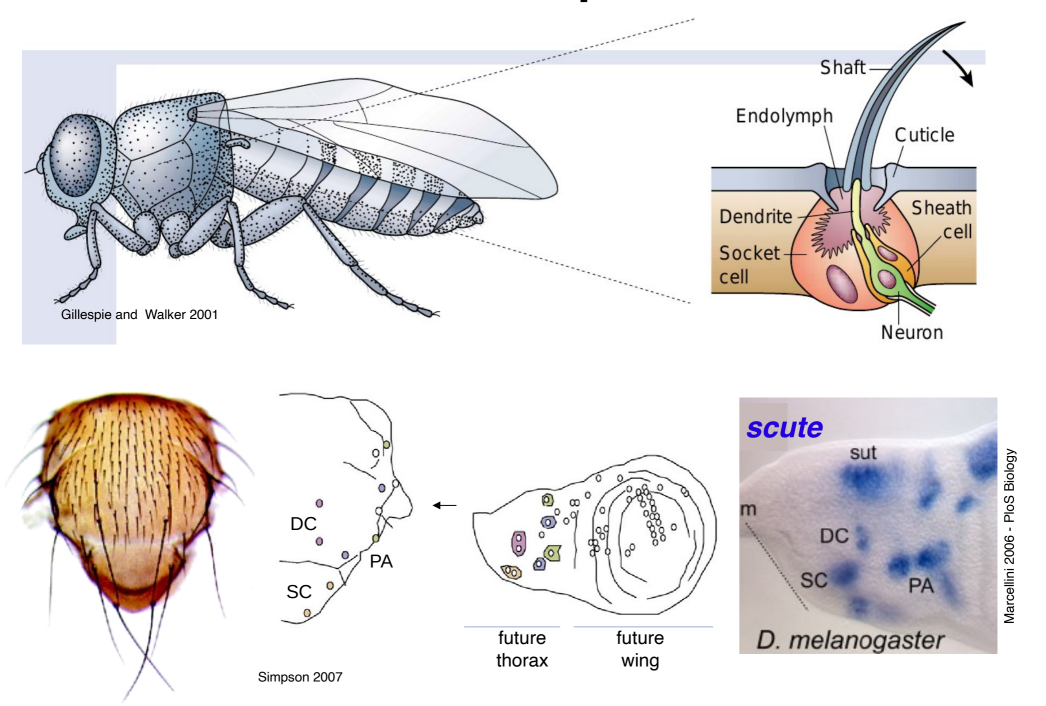


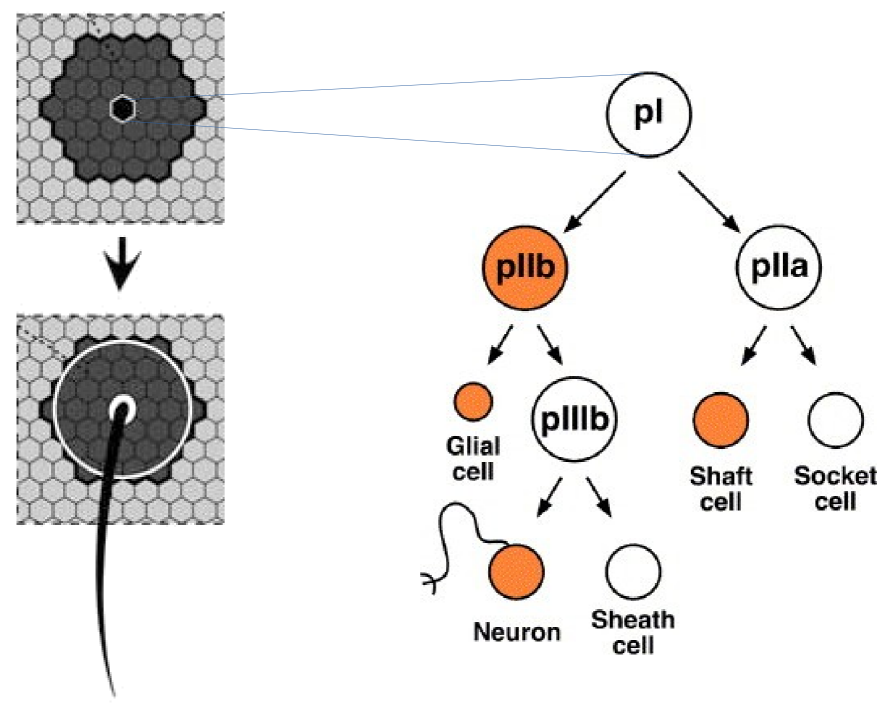
color, type, orientation shape and size presence/absence position

CRE mutations in achaete-scute

Aristotle, Historia animalium, book I, 2, 300BC

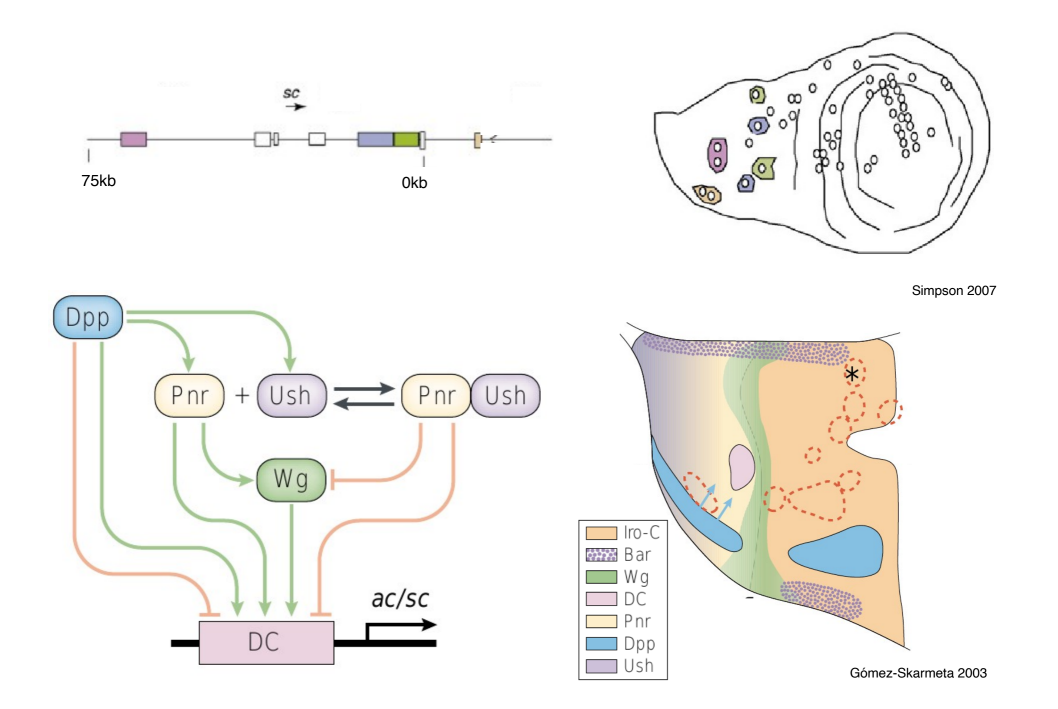
Bristle development



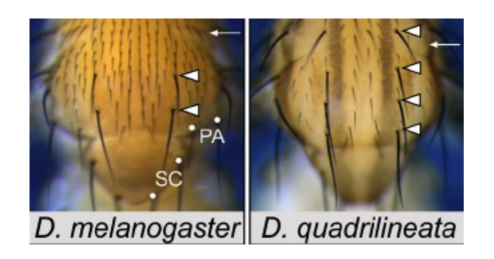


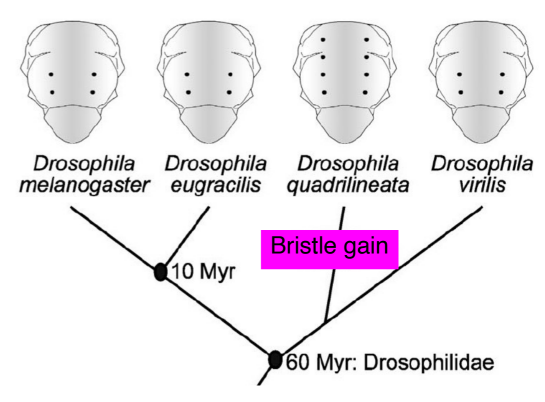
Lewis - Imaginal discs

scute cis-regulatory elements are "master switches"



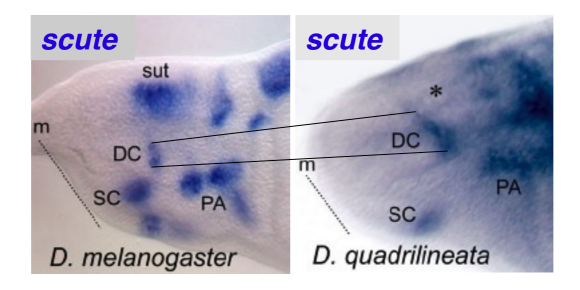
Extra bristles in *D. quadrilineata*





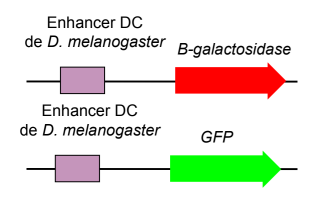
Extra bristles in *D. quadrilineata* correlate with larger *scute* expression domain

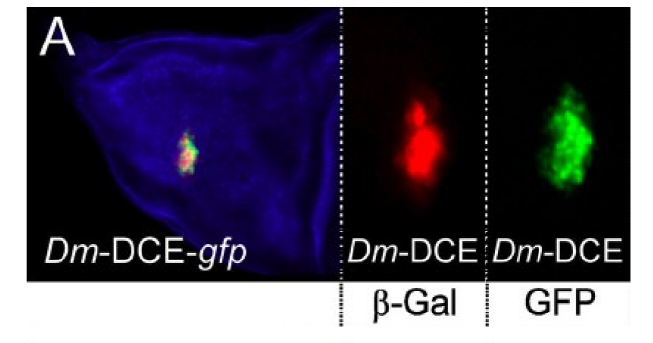
In situ hybridization

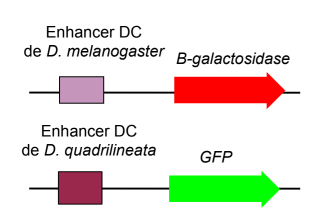


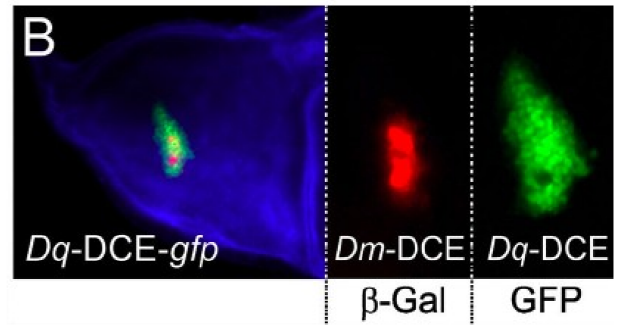
Test for a cis-regulatory change (1)

D.melanogaster transgenics





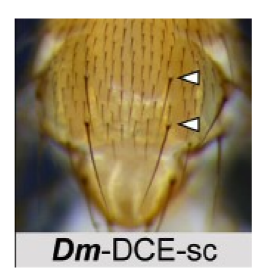


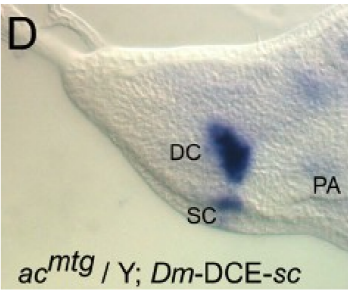


Test for a cis-regulatory change (2)

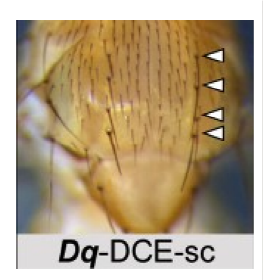
D.melanogaster transgenics

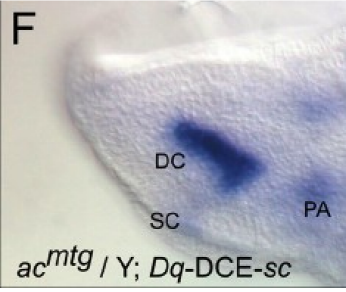




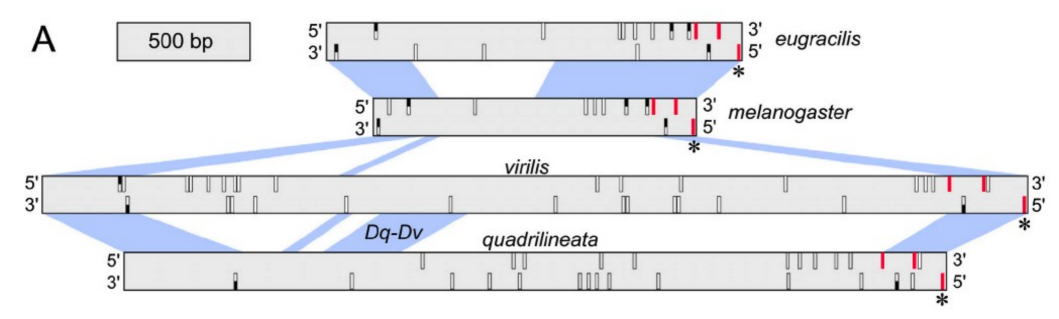




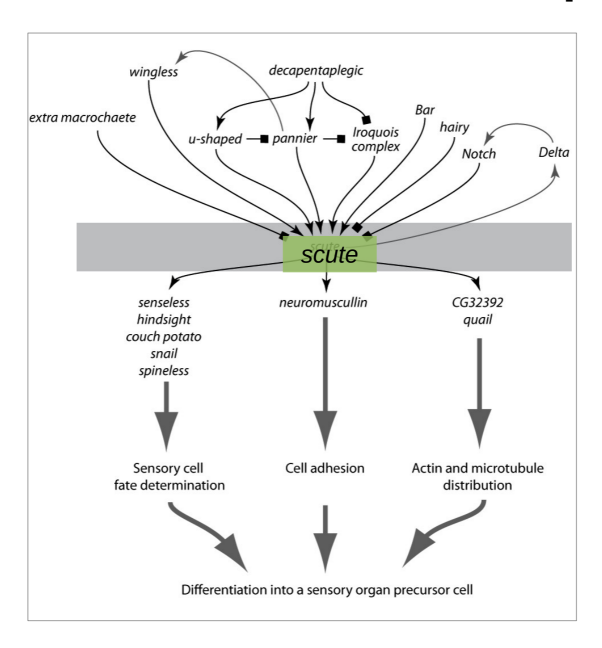


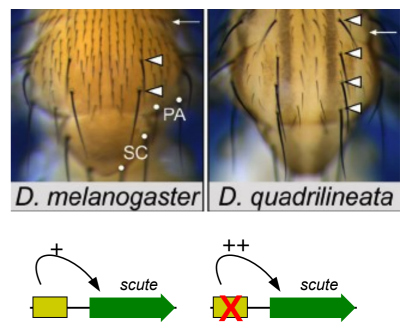


Alignment of the DC region

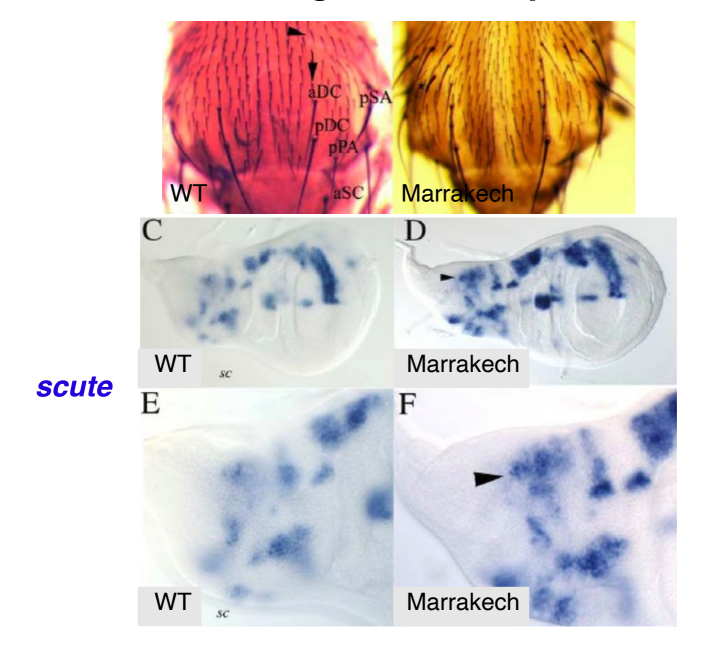


Genetic evolution is partly predictable

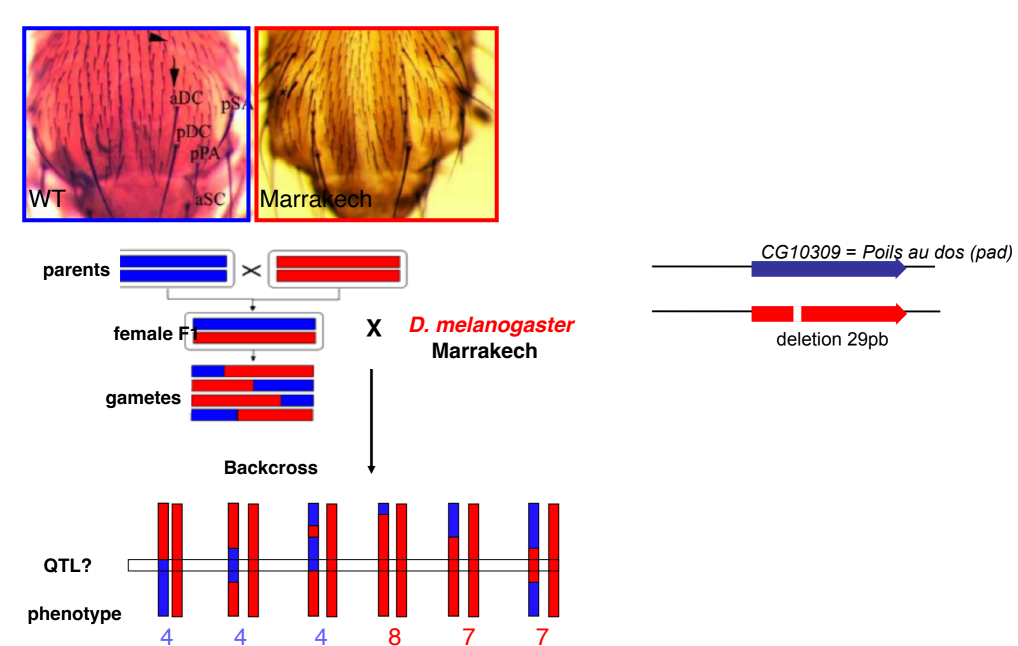




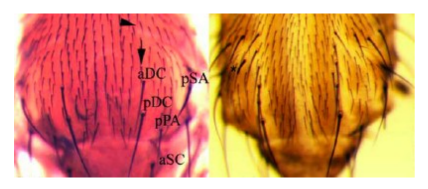
Extra bristles in *D. melanogaster*-Marrakech correlate with larger *scute* expression domain



Extra bristles in *D. melanogaster*-Marrakech due to mutation(s) in *poils-au-dos*

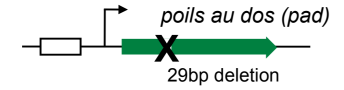


Short-term evolution...



D. melanogaster

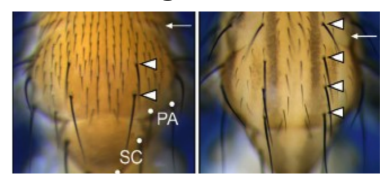
D. melanogaster variant



null mutation in coding region change in thorax and wing

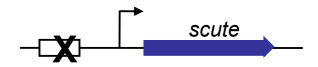
(Gibert et al., 2005)

...versus long-term evolution



D. melanogaster

D. quadrilineata



cis-regulatory mutation change in the thorax only

(Marcellini et al. 2006)

Methods to identify the genes and the mutations responsible for phenotypic evolution

Various methods

Genetic

which chromosome (ex: autosomal versus sex)

QTL mapping

Genetic association studies

Complementation tests

General biology

General knowledge of the genes involved in the phenotype Similarity with a known phenotype Correlation with a change in gene expression level/pattern

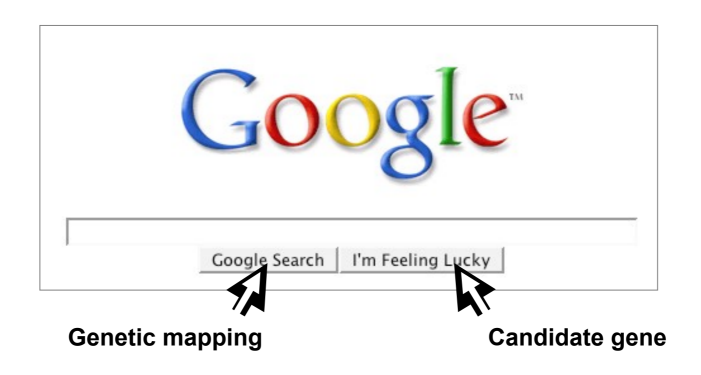
Final test of protein activity

in vitro in *E. coli*, by transgenesis in the studied species or the closest model organism (ex: *beta-defensin* of dogs tested in mouse)

Final test of cis-regulatory regions

- with reporter constructs, transgenesis, comparison of both regions
- comparison of allele expression levels in hybrids (pyrosequencing)

Two types of approaches



no a priori, fewer bias long and tedious rarely ends with identification of the gene

only with strains/species which produce fertile hybrids

Based on an a priori idea can be fast and efficient

will only find known genes

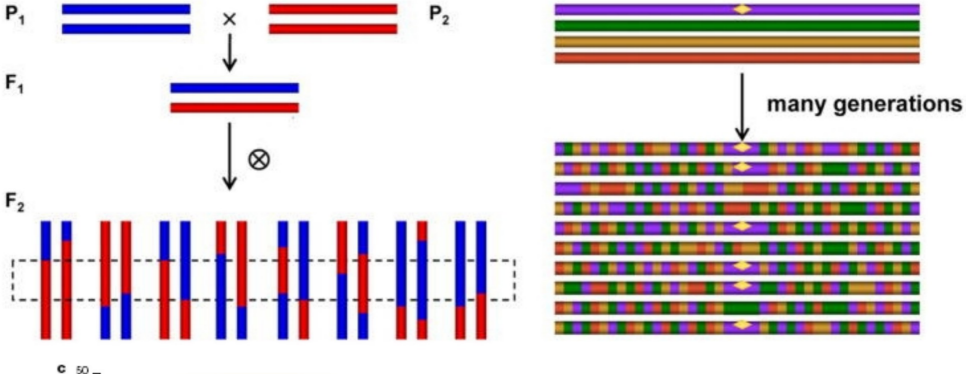
In both cases, genes with small effect are more difficult to identify

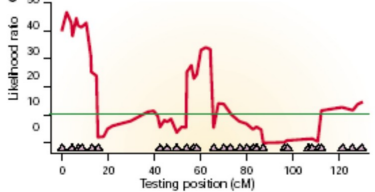
Crosses in the lab

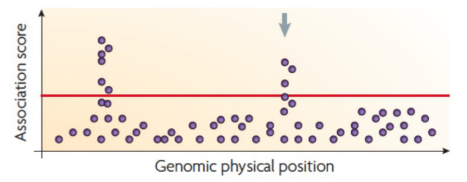
Linkage **Mapping**

Association Mapping

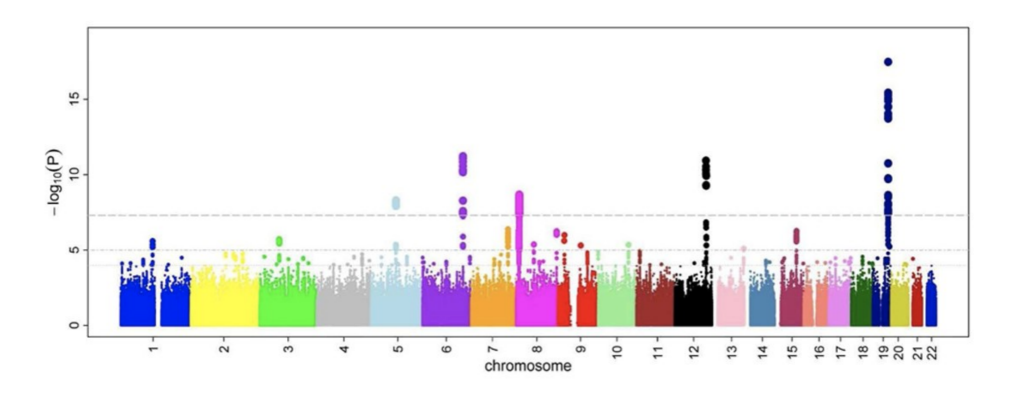
Past crosses in natural populations





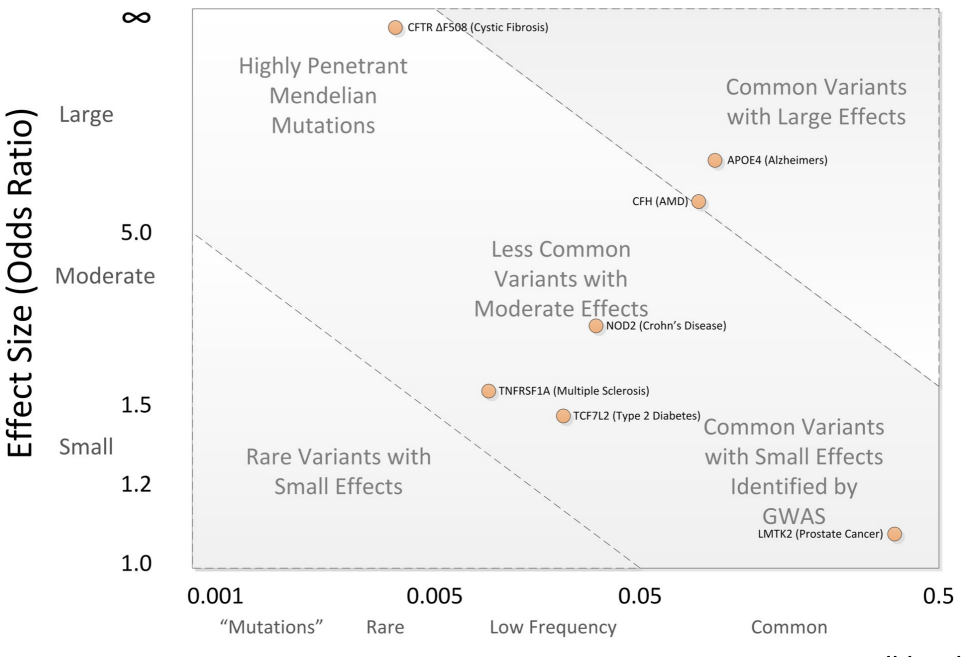


Genome-wide association study (GWAS)



Manhattan plot depicting several strongly associated risk loci. Each dot represents a SNP, with the X-axis showing genomic location and Y-axis showing association level.

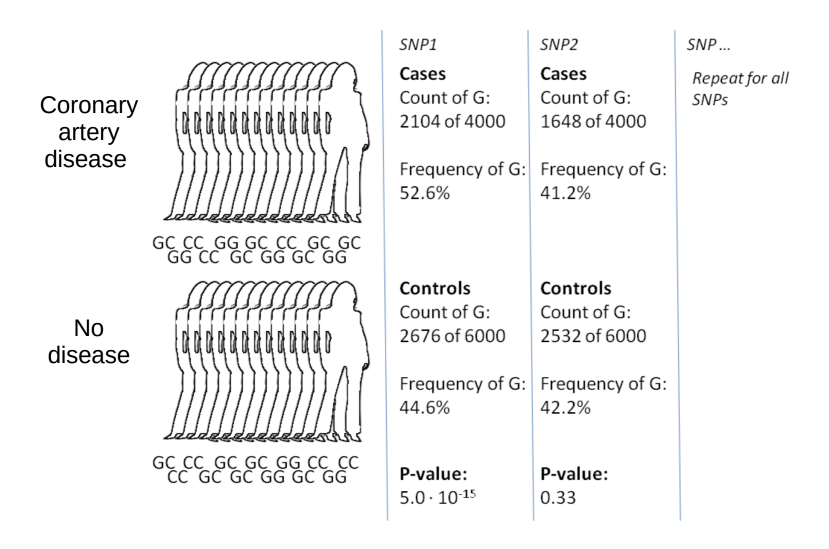
GWAS typically identify common alleles



Allele Frequency

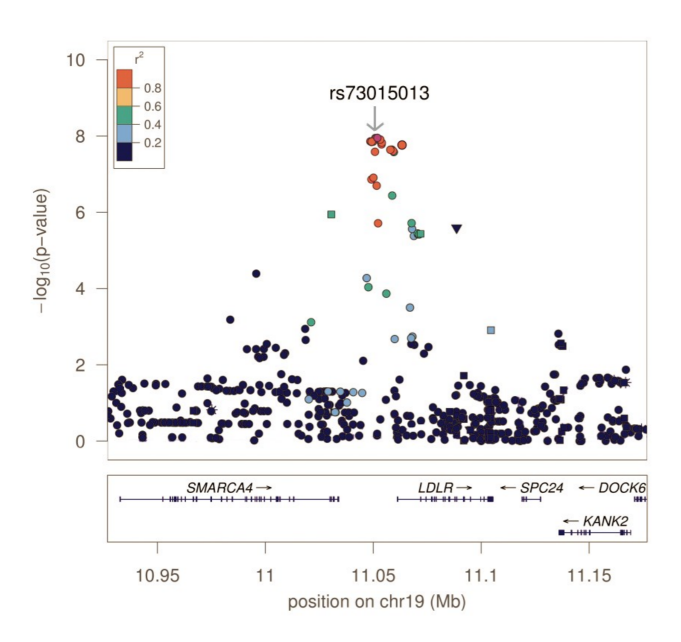
Wikipedia

Methodology of a case-control GWA study



The allele count of each measured SNP is evaluated, in this case with a chi-squared test, to identify variants associated with the trait in question.

Regional association plot

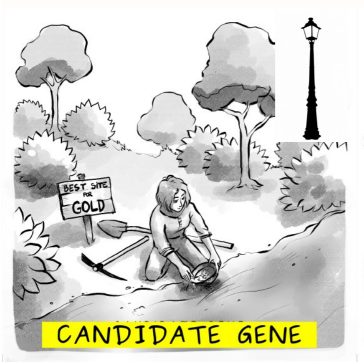


Association to LDL-cholesterol levels.

The haploblock structure is visualized with colour scale and the association level is given by the left Y-axis.

The dot representing the rs73015013 SNP (in the top-middle) has a high Y-axis location because this SNP explains some of the variation in LDL-cholesterol.

THREE APPROACHES to FIND the GOLDEN LOCI of EVOLUTION





From genes to traits





FORWARD GENETICS

From traits to genes
Little Ascertainment Bias, but Micro-Evolution only

Experimental Evidence

3 categories, each with biases

Experimental

Principle



Candidate Gene

Reverse Genetics:

looking for sequence differences and trait effects based on previous studies of a given gene

> 66 cases of color variation associated to MC1R coding mutations in vertebrates

> > High

Favors identification of

coding mutations

Favors traits with small

molecular targets,

large-effect size

Ascertainment Bias on Locus Identification

Example

Molecular Type Bias

Trait Type Bias

Taxonomic Breadth

Large



Linkage Mapping

Forward Genetics:

trait mapping in hybrids obtained from laboratory crosses, using recombination over a few generations

F2 crosses between melanic and amelanic phenotypes in cavefish: identification of MC1R and Oca2 alleles in distinct cave populations

Low to Intermediate

(depending on resolution / cross size)

Little molecular bias

Amenable to dissection of complex traits with small-effect size (large crosses, multiparental families)

Narrow, limited to interfertile lineages (populations or sister species)



Association Mapping

Forward Genetics:

statistical SNP/character state association in large cohorts, using recombination over many generations

GWAS of human pigmentation (skin, hair, eyes): identification and confirmation of causal variants at >15 genes including Oca2 p.His615Arg in Eastern Asia

Low

Can miss structural variants (short read genotyping)

Most common approach for complex traits with small-effect size

Very narrow, limited to polymorphic or intermixing populations

QTL Mapping

4 steps: crosses, genotyping, phenotyping, statistical analysis

Crosses

Backcross with one line
Backcross in both directions
F2
Crosses for several generations
Introgression lines
Recombinant Inbred Lines

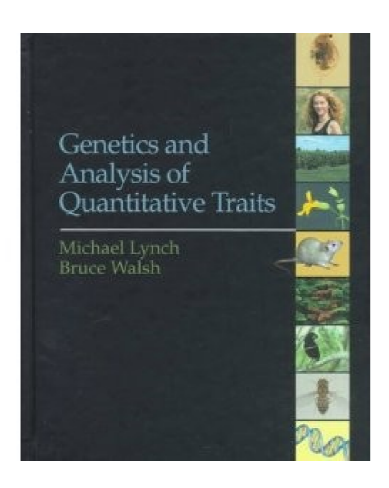
Always try to maximize the number of recombination events

Markers

yes-no PCR
PCR length polymorphism
Pyrosequencing
Probe hybridization
Microarray
RADseq
High-throughput sequencing

How many markers?

theory



practice

