

ГЕНОМИКА (6 лекций, 12 акад. часов)

- **Лектор:** Константин Валерьевич Крутовский
- Профессор Гёттингенского университета, Германия (<http://www.uni-goettingen.de/en/414626.html>) и Техасского университета (<http://essm.tamu.edu/people/faculty/adjunct-faculty/krutovsky-konstantin>)
- Ведущий научный сотрудник Института общей генетики им. Н. И. Вавилова РАН, Москва
- Зав. лабораторией лесной геномики и руководитель Научно-образовательного центра геномных исследований Сибирского федерального университета, Красноярск (<http://genome.sfu-kras.ru/en/krutovsky>)
- **Тел.:** +7 965 912 1959 (моб.)
- **E-mail:** kkrutovsky@gmail.com или kkrutov@gwdg.de
- **Office:** ЦЗЛ, Академгородок 50а, корп.2, ауд. 133
- **Office Hours:** You can contact by e-mail or phone to make an appointment.
- **Textbook (not required):** Freeland, J.R., H. Kirk, and S.D. Peterson 2011. Molecular Ecology. 2nd ed. John Wiley & Sons, Ltd. 449 pp.
- **Required e-mail:** You will need to send an e-mail to kkrutov@gwdg.de from your preferred address. This would allow me to distribute class announcements, lecture notes, readings, problem sets, etc. Please, provide me with:
 - your full name
 - name by which you prefer to be called
 - phone number(s) where you can be reached
 - e-mail address that you check daily
 - academic & career interests
 - what you hope to get from this course
- **Course web page:** <http://genome.sfu-kras.ru/node/200>
- **Lecture notes:** PowerPoint lecture notes for most of the class sessions will be available on the Web site prior to each class session. **I expect you to print out and bring the notes with you to class (bring also your laptop with you to class)**

ГЕНОМИКА: Введение. 4 апреля 2017. Вторник. #1




ТЕМЫ ЛЕКЦИЙ ПО КУРСУ «ГЕНОМИКА»

№ п/п	Наименование лекций	Объем в акад. часах	Дата и время проведения
1	Введение в геномику. Содержание и организация геномной информации. Геномика, транскриптомика, протеомика, метаболомика. Программа "Геном человека"	2	Вт. 4.04 14.10-15.45
2	Технология секвенирования ДНК. Полногеномное <i>de novo</i> секвенирование, ресеквенирование, целевое и метагеномное секвенирование.	2	15.55-17.30
3	Методы работы с нуклеотидными сиквенсами и геномными базами данных. Программа поиска гомологий – BLAST. Формат Genbank, выравнивание (Bio Edit) и аннотирование нуклеотидных последовательностей (Augustus).	2	Ср. 5.04 14.10-15.45
4	Популяционная геномика. Генотипирование ДНК-полиморфизмов (SSRs, SNPs). Тесты на селективную нейтральность (DNASP). Гены-аутсайдеры (LOSITAN).	2	15.55-17.30
5	Полногеномное ассоциативное картирование. Подходы и методы полногеномного ассоциативного картирования (TASSEL).	2	Пят. 7.04 12.00-13.35
6	Практические приложения геномики: филогеномика, экогеномика, природоохранная геномика, палеогеномика, персонифицированная медицина, геронтогеномика, метагеномика, эпигеномика и геномная селекция.	2	14.10-15.45

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




ГЕНОМИКА

4 апреля 2017, Вторник


1. **Введение в геномику.** Содержание и организация геномной информации. Геномика, транскриптомика, протеомика, метаболомика.
2. **Технология секвенирования ДНК.** Полногеномное *de novo* секвенирование, ресеквенирование, целевое и метагеномное секвенирование.



ГЕНОМИКА: Введение. 4 апреля 2017. Бронник. #1


What is Genomics?

- The term "genome" was introduced in 1920 by German botanist Hans Winkler (1877-1945) who combined "gene" and "chromosome" to refer to all genes on all chromosomes in the nucleus of a cell.
- The term "genomics" was coined in 1986 by Thomas Roderick (Jackson Laboratory, USA) to describe the scientific discipline of mapping, sequencing, and analyzing genomes and to provide a name for the new journal *Genomics*.



- Genomics is more comprehensive now and includes comparison of individual genomes in a population or from different populations (population genomics) and species (comparative genomics), study of their evolution (evolutionary genomics) and function (functional genomics).

Genomics studies genomes, genes and their functions using global genome-wide experimental approaches



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Milestones in Genetics that led to Genomics

1944: DNA was identified as the genetic material of all living organisms (Avery et al., J. Exp. Med. **79**, 137: 1944)

1953: The genetic code was deciphered (Watson & Crick, *Nature* **171**, 737: 1953).

1977: The first complete DNA sequence of an entire genome - the bacteriophage phiX174 (Sanger et al. 1977 *Nature* **265**, 687-695) of only 5386 nucleotides, which is 60000 times smaller than the human nuclear genome.

mid-1980s: Major advances in DNA sequencing, laboratory automation & computing.

1990: The project on complete sequencing of the human genome was launched.

1997: complete sequencing of the yeast genome (12 Mbp)

1998: nematode genome (97 Mbp)

2000: fruit fly (180 Mbp) & the first plant *Arabidopsis* (125 Mbp) genomes

2001: human genome (3,200 Mbp)

2002: mouse (3,500 Mbp) & rice (420 Mbp) genomes

2006: the first forest tree - poplar genome (550 Mbp)

mid-2000s: Next generation sequencing (NGS) platforms - high-throughput massively parallel sequencing

The Genomes OnLine Database (GOLD, <https://gold.jgi-psf.org>):

- Complete Projects: 9050
- Organisms: 239935 (Archaea: 1999; Bacteria: 218872; Eukarya: 14420)
- Incomplete Projects: 44243
- Targeted Projects: 1454

2013: neandertal genome (3,200 Mbp)

2013: Norway spruce & **2014:** loblolly pine (~20,000 Mbp) genomes



5

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Major areas of Genomics

Structural Genomics

- DNA libraries and complete genome sequence
- Gene annotation and homology search
- Linkage analysis, genetic and physical mapping
- Development of genome-wide genetic markers

Functional Genomics

- Gene expression analysis (transcriptome, proteome & metabolome profiling)
- Gene function, gene-trait and gene-environment relationships

Comparative & Evolutionary Genomics

- Comparative mapping and search for orthology and synteny
- Gene and sequence comparison across different species
- Signatures of selection, evolutionary footprints

Statistical Genomics

- Mapping algorithms and associative analysis
- Database management, data collection and communication
- Sequence assembly, alignment, comparison and annotation

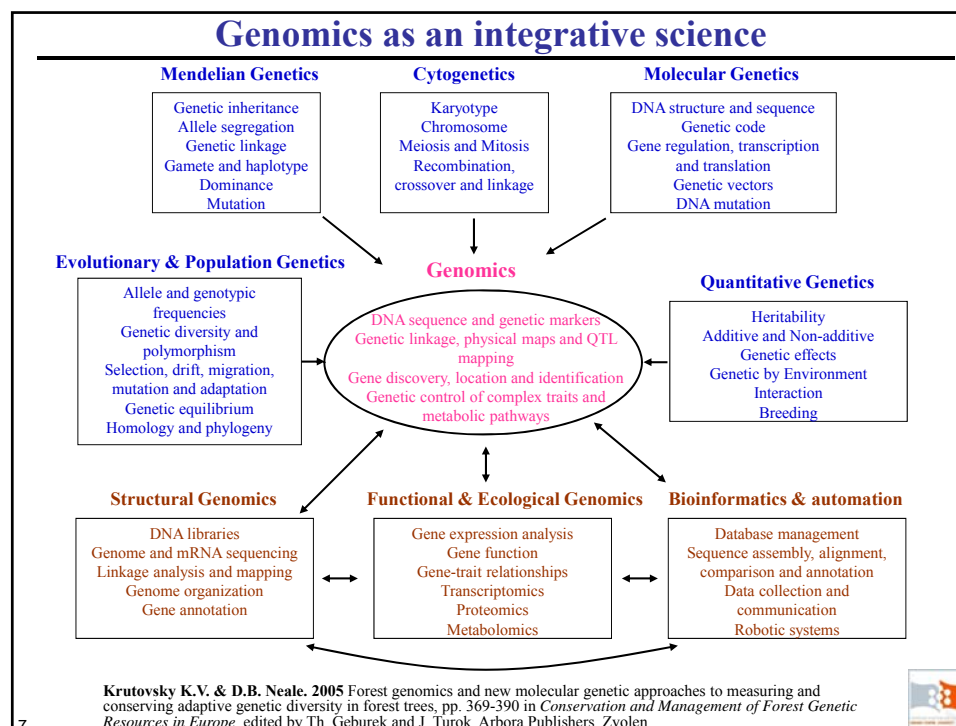
Population & Ecological Genomics

- Genome wide scan for nucleotide diversity
- Genome wide and candidate gene based association mapping
- Assessment of association between alleles and phenotypes and environments via association mapping

Gene
discovery

6

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	<h3 style="text-align: center;">Forest genomics and new molecular genetic approaches to measuring and conserving adaptive genetic diversity in forest trees</h3> <p style="text-align: center;">K. V. Krutovsky & D. B. Neale</p> <p>Introduction</p> <p>Genetic diversity is the basis of the ability of organisms to adapt to changes in their environment through natural selection. Populations with little genetic variation are more vulnerable to the arrival of new pests or diseases, pollution, changes in climate and habitat destruction due to human activities or other catastrophic events. The inability to adapt to changing conditions greatly increases the risk of extinction. Genetic conservation and management aimed to save adaptive genetic diversity should be based on the knowledge of the genetic basis of adaptation. The goal of this paper is to describe how adaptive genetic diversity can be measured using new molecular genetic approaches and achievements in forest genomics.</p> <p>Traditional methods to measure adaptive genetic diversity</p> <p><i>Field experiments</i></p> <p>Field experiments (common-garden tests) have been used traditionally to measure adaptive genetic diversity in trees. These tests continue to be used extensively in tree breeding and are very effective in identification of families and clones that are specifically adapted to particular environments or to a broad variety of environments. However, field experiments are very time consuming and relatively expensive, and more importantly, they are based solely on the phenotypes. They can estimate genetic parameters but only on measurable traits, not on individual genes. This method can neither provide information on what particular genes and how many of them are involved in adaptation nor how much of phenotypic variation can be explained by genetic variation in these genes. More details can be found in (see p. 275 ff., this volume).</p> <p style="text-align: right;">369</p>	
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Climate change is a global systemic threat to ecosystems and needs a systemic (holistic) approach

- To be able to predict and mitigate effects of climate change and to breed resilient crops we have to understand **evolutionary responses** and **molecular mechanisms of genetic adaptation**.
- **Evolutionary response** is a **genetic adaptation** via genetic change that promotes adaptation of plants and animals to their natural environment, including their ecosystem interactions with members of their own and other species (the biotic environment) as well as the physical environment (the abiotic environment).
- Multiple genes are involved in genetic adaptation, so its study requires genomic methods and genome-wide approaches.

9

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How genomics can help us identify evolutionary responses and study molecular mechanisms of genetic adaptation?

- **Structural genomics** provides practically unlimited number of genetic markers for population genetic studies via new methods of high-throughput massively parallel sequencing and genotyping.
- **Population genomics** 1) provides detailed nucleotide and allelic variation in numerous adaptive trait related candidate genes, 2) identifies genes under selection (via **whole genome scans, neutrality tests, outliers**, etc.), and 3) links genotypes to phenotypes (via **genome-wide association study - GWAS**).
- **Ecological genomics** helps to identify genes responsive to environmental factors via genome wide differentiation expression analysis (using mRNA/cDNA sequencing or transcriptome profiling) and associating genotypes with environmental variables.

(**Spatial** or **landscape genomics** is one of applications of **population genomics**)

10

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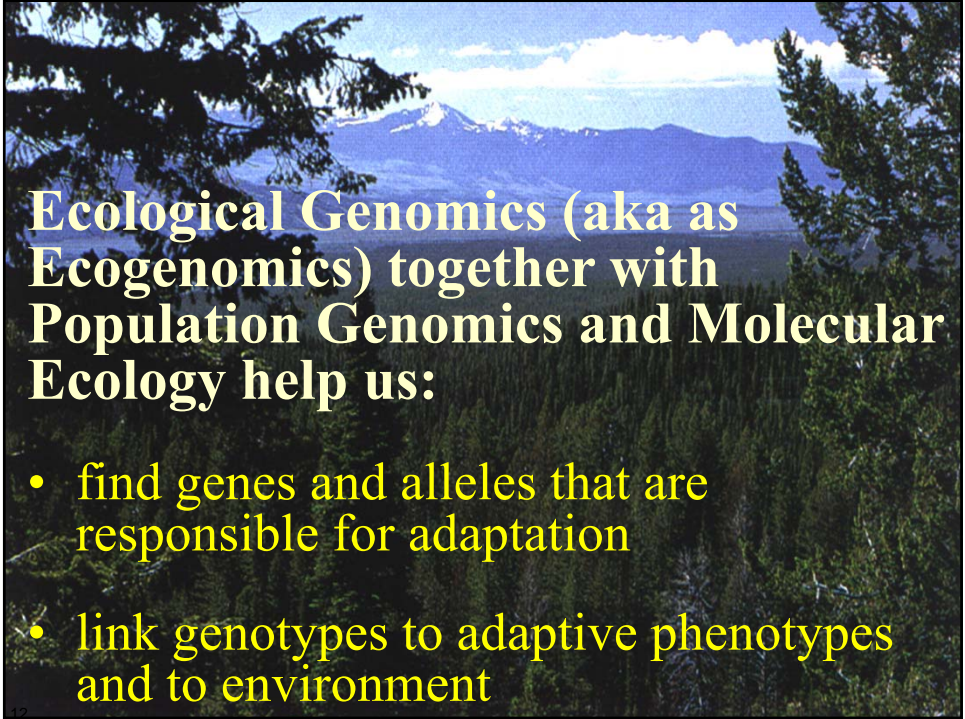


What is ecological genomics?

- a novel, fast-developing biological discipline, combining traditional ecological and population genetic approaches with the genome-wide level of analysis
- thousands of genes with known function and sometimes known genome-wide localization can be simultaneously studied in many individuals
- applies genomic tools and approaches to traditional ecological and population genetics questions (local adaptation, mating system, gene exchange, reproductive population size, population disequilibrium, interaction among populations and species, population–environment relationships, community composition, conservation and assessment of genetic diversity, etc.)
- these traditional problems of ecological and population genetics can be now studied using data on variation in many genes
- an interdisciplinary approach to a full understanding of interaction between genotypes, phenotypes, and environment
- a truly integrative discipline that embraces many related disciplines, such as ecology, phylogenetics, population and conservation genetics, molecular evolution, etc.

11

ЛЕОМОНКА: Биология 4 апреля 2017, Бронис, #1



Ecological Genomics (aka as Ecogenomics) together with Population Genomics and Molecular Ecology help us:

- find genes and alleles that are responsible for adaptation
- link genotypes to adaptive phenotypes and to environment

Nature (*Genome*) vs. Nurture (*Environment*)

$$P = G + E + G \times E$$

Phenotype = *Genotype* + *Environment* + *Interaction*

Organisms are different because of the:

- *genetic* differences among individuals
- different *environments* where individuals are growing
- and *interactions* between the *genotypes* and the *environments* in which they grow

13

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Nature (*Genome*) vs. Nurture (*Environment*)



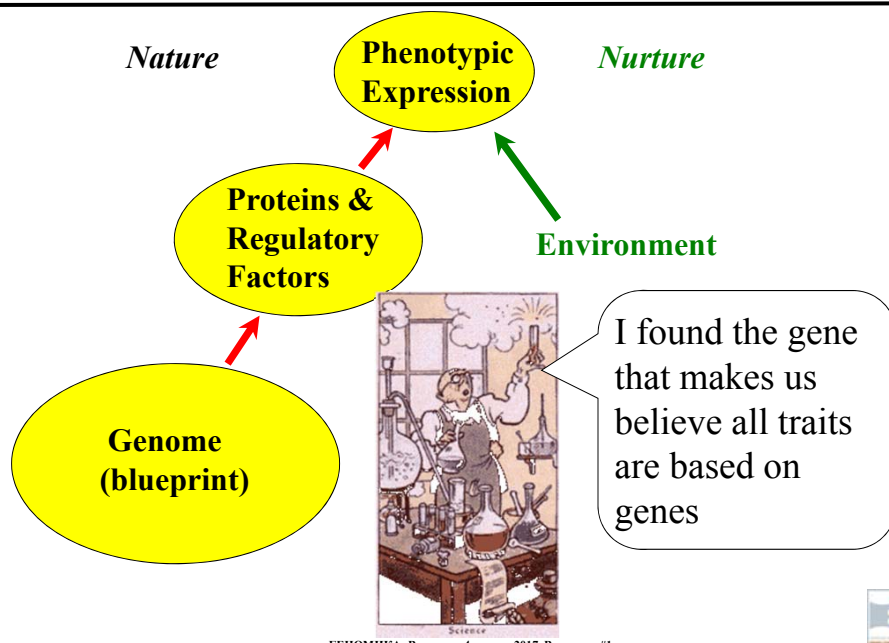
How much can we blame our genotype for our phenotype?

14

ГЕНОМІКА: Введення. 4 лютого 2017. Броніус. #1



Nature vs. Nurture



15

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Simple single gene (Mendelian)

vs.

Complex multiple genes (Quantitative) variation

- **Mendelian = Qualitative**
 - single gene responsible for most of the observed phenotypic variance
- **Complex = Quantitative**
 - with gene \times gene, gene \times environment interactions contributing to phenotypic variance

16

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Single vs. Multiple Genes in Population

$$P^n = G^n + E^n + G^n \times E^n$$

n – multiple phenotypes, genes and environments

Great Fermat's Theorem: $Z^n = X^n + Y^n$

does not have integer solutions X, Y, Z for $n > 2$



Andrew Wiles, 1994

Life Theorem : $P^z = G^x + E^y$

Great Life Theorem: $P^z = G^x + E^y + G^x \times E^y$

Genomics is the solution!

17

TEHOMIKA: Biochemie, 4. Januar 2017, Brannius, #1

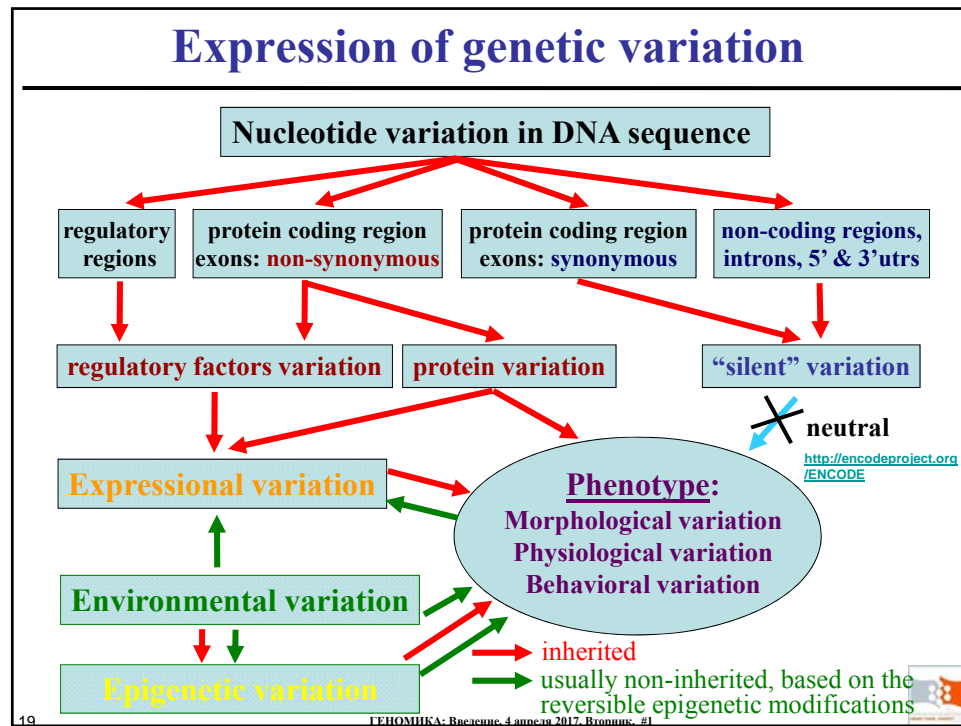


Can we predict phenotypes based on the genotypes?



18

TEHOMIKA: Biochemie, 4. Januar 2017, Brannius, #1



How to measure evolutionary response in populations and ecosystems?

- **Traditional methods**
 - field or common garden experiments (provenance, progeny and clonal tests)
 - molecular genetic markers: advantages and disadvantages; summary statistics
 - Quantitative Trait Locus (QTL) Mapping
- **Modern population genomics approaches**
 - new type of functional genomic markers
 - use of adaptive trait related candidate genes in population studies
 - association mapping with phenotypic and environmental variation
 - detecting selective signatures and loci under adaptive genetic divergence in natural populations using neutrality tests and outlier-detection approaches
 - differential expression, transcriptome profiling, etc.

Integrated approach: 1) & 2)

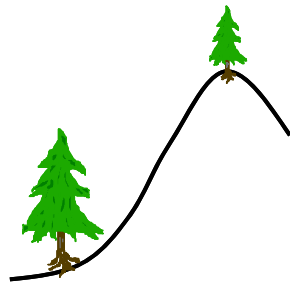
20

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Nature vs. Nurture

How to separate the two?

Example: *Altitudinal gradient*



Common Garden Exp.



21

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Field or common garden experiments (provenance, progeny and clonal tests)



USDA Forest Service Pacific Northwest Research Station runs an ecogeographic study of Douglas-fir



Weyerhaeuser Company runs clonal evaluation of phenotypes in loblolly pine and Douglas-fir



USDA Forest Service Dorena Tree Improvement Center runs a white pine blister rust screening program in sugar pine

22

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Common garden experiments

- helped to learn a lot about patterns of adaptive variation in complex traits, both at the macro- and micro-environmental level
- often shows geographical patterns in populations, such as steep latitudinal or altitudinal clines
- time consuming and relatively expensive, and are based solely on the phenotypes
- can estimate genetic parameters but only on measurable traits, not on individual genes
- can provide neither information on what particular genes and how many of them are involved in adaptation nor how much of phenotypic variation can be explained by genetic variation in these genes

23

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Quantitative Trait Locus (QTL) Mapping

- **One of the first genome-wide approaches to link genotypes and phenotypes**
- **It directs to the chromosomal regions (and sometime genes) responsible for the observed phenotypic variation**

24

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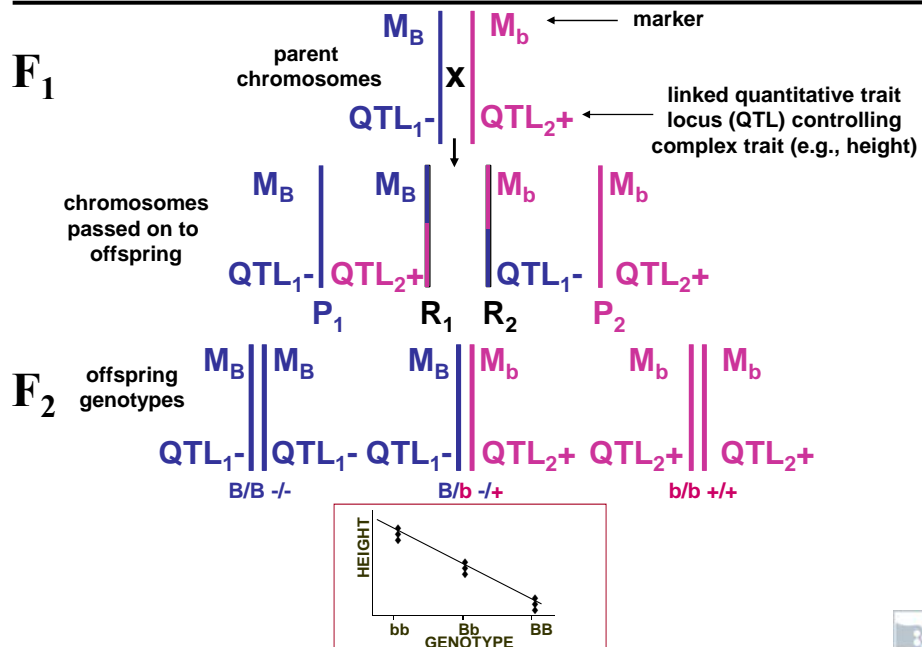
What is a Quantitative Trait Locus (QTL)?

- A *QTL* is a chromosomal region supposedly containing a gene (or cluster of genes) that contributes to the variation observed at a *quantitative trait*
- It must be polymorphic (have allelic variation) to have an effect in a population
- It must be linked to a polymorphic marker allele to be detected
- QTLs are detected through QTL mapping experiments

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Quantitative Trait Locus (QTL) mapping



ЛЕОМОНКА: Вакцина 4 августа 2017, Бранниш, #1



Basic concept

- The closer together are two markers or genes linked on a parental chromosome, the less likely the parental alleles at the two loci will be split up in gametes as a result of meiotic recombination (crossing over).
- Genes and genetic markers that are closely linked together on a chromosome will tend to co-segregate in the F_2 - the same allele combinations (haplotypes) that occurred in one of the parents will tend to occur together in the offspring.
- This will lead to a statistical association between a gene segregating for alleles that have a measurable difference in their affect on a quantitative trait (QTL) and segregating alleles at closely linked marker loci.

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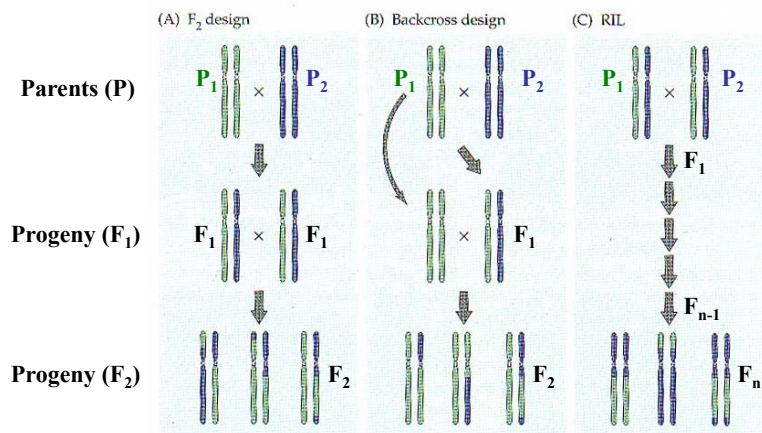


QTL mapping populations

A. F_2 populations

B. Backcrosses

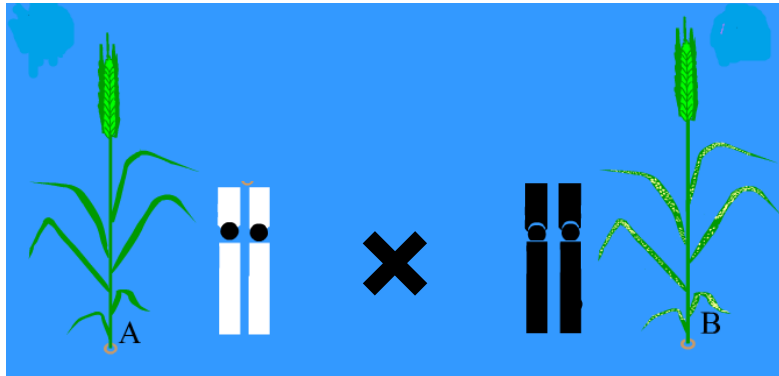
C. Recombinant Inbred Lines (RILs)



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F₂ mapping populations

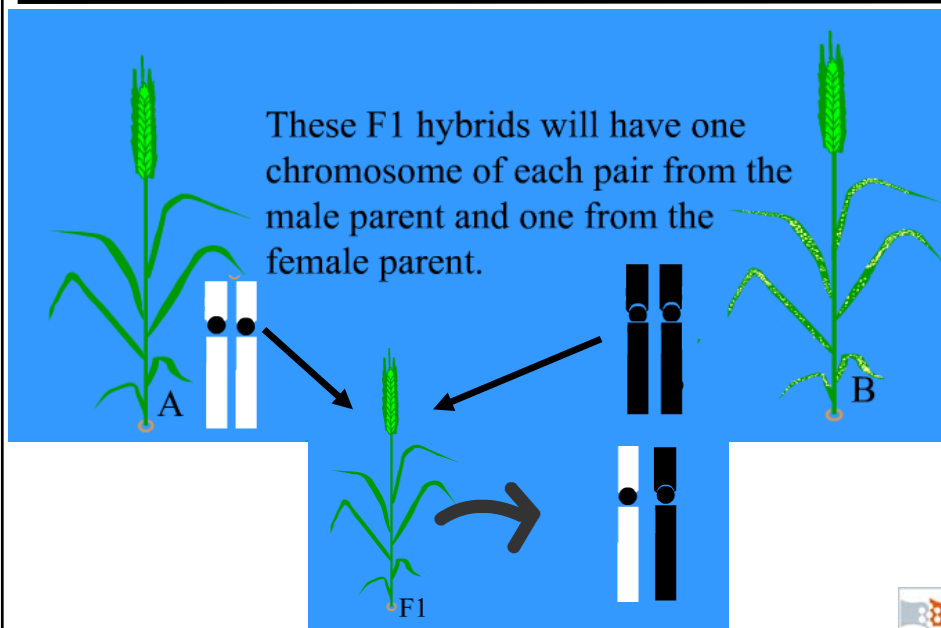


- First step: cross line A (e.g., disease resistant) with line B
- This can be done with placing pollen from one parent to the stigmas of the other in order to produce hybrid seeds

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F₂ mapping populations

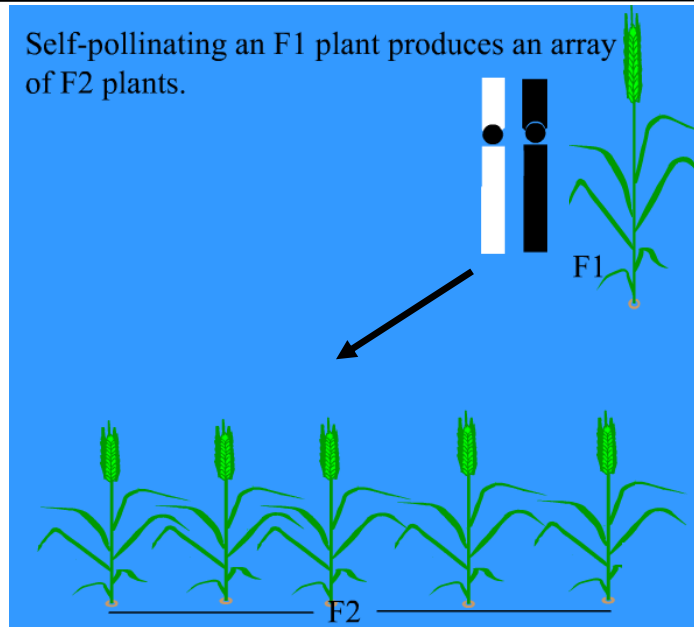


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F₂ mapping populations

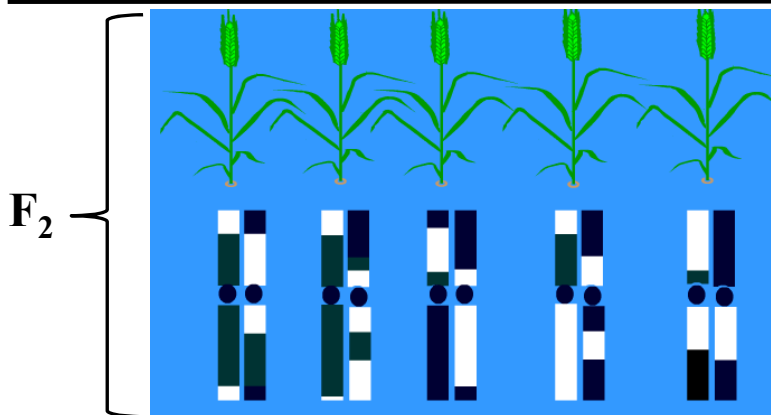
Self-pollinating an F₁ plant produces an array of F₂ plants.



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F₂ mapping populations

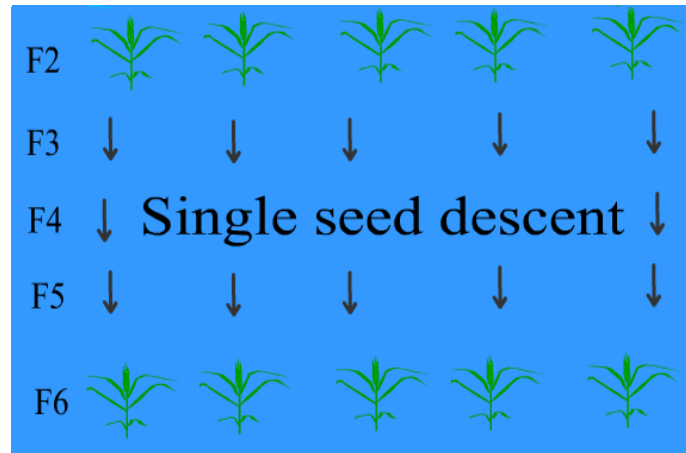


- Unique mosaic of chromosome segments from each parent
- Typical QTL mapping population (F₂ design)
200-300 plants

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Recombinant Inbred Lines (RILs)

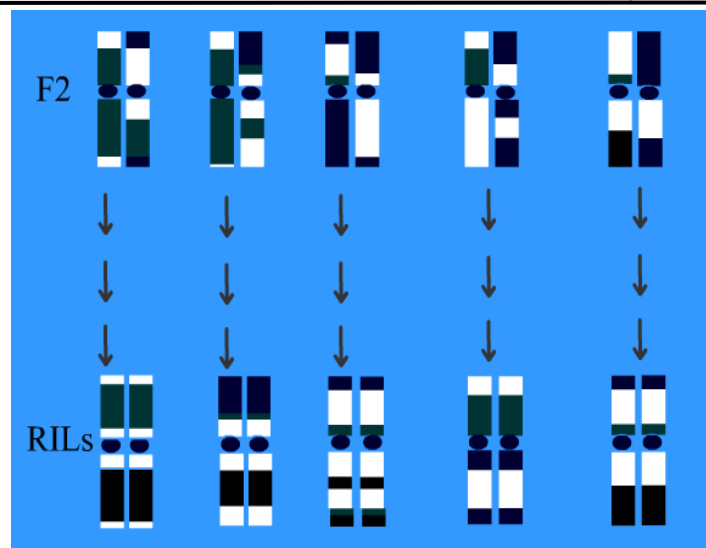


- RILs are developed by several generations of self-pollination from F_2 plants, through a process known as “single seed descent”

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Recombinant Inbred Lines (RILs)

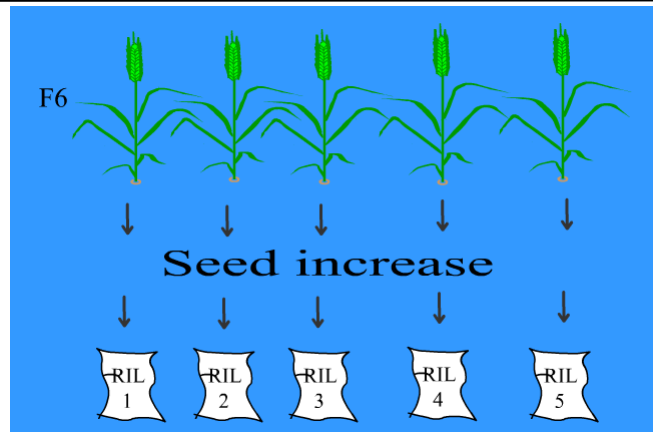


- More generations \rightarrow more meiosis \rightarrow more recombination

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Recombinant Inbred Lines (RILs)

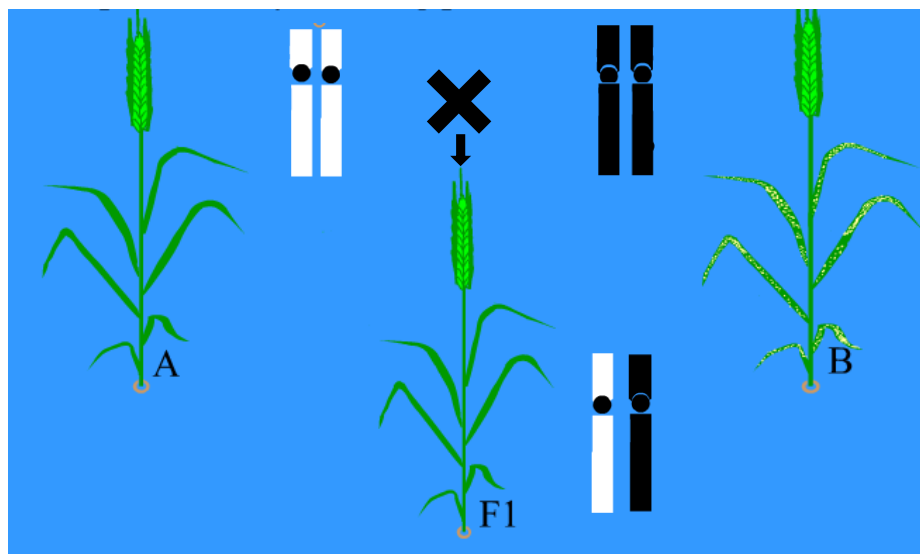


- Important advantages:
 - multiple genetically identical individuals (decreasing the environmental contribution to one trait)
 - multiple traits analysis (from the same line)

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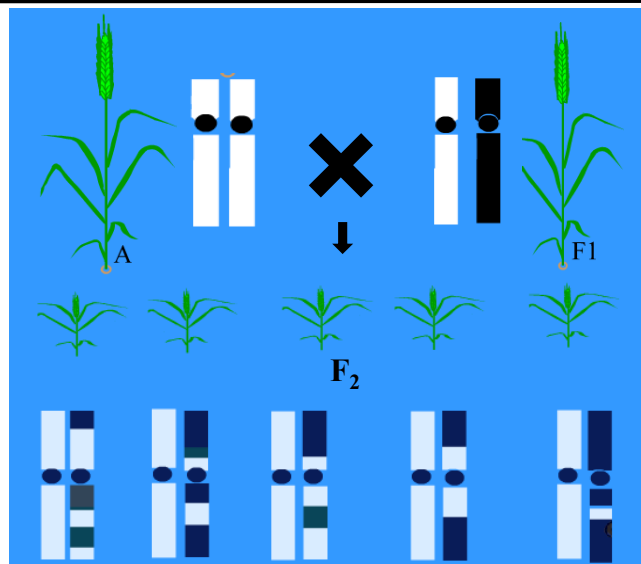
Backcross populations



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Backcross populations

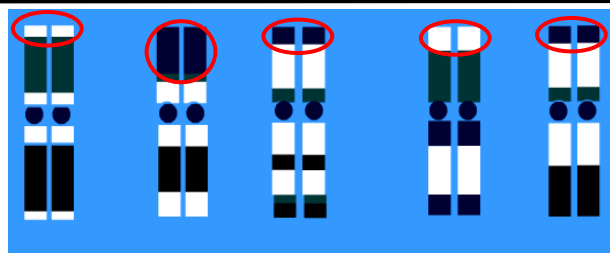


- Result: population of individuals that have one chromosome from parent A and a recombined chromosome with segments of both parents A and B

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Example: Calculate the average trait value of individuals having the same segment (either parent A or parent B) in a specific chromosome region

RILs:

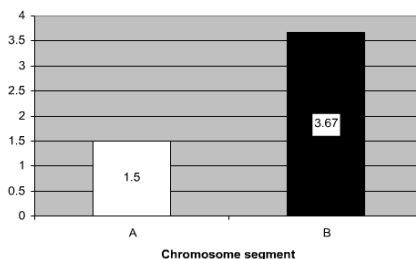


Parental segment:

A B B A B

Scale of Resistance (1-4)

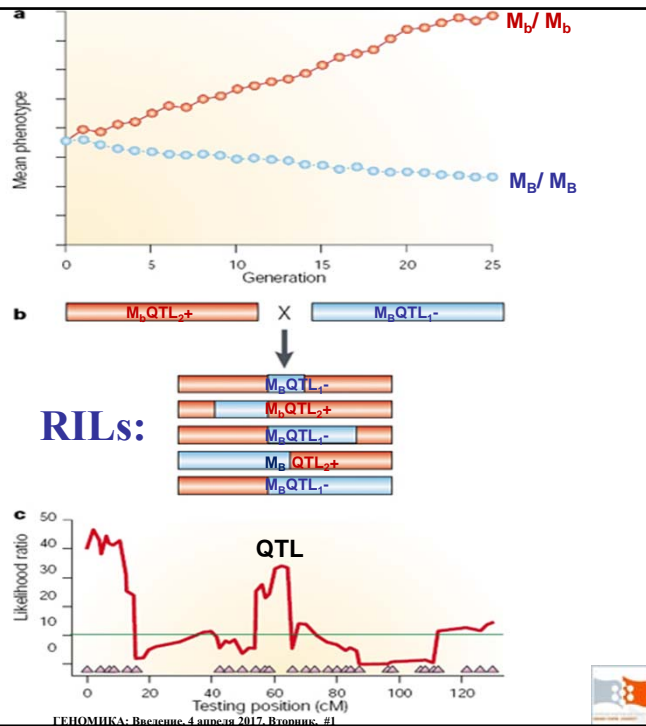
1 3 4 2 4



- can be statistically tested if the difference is significant
- can be one gene in this chromosome segment that influences the trait (in this case, resistance)

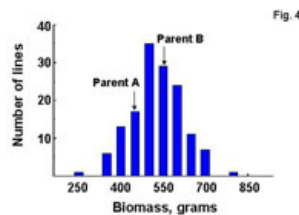
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Quantitative Trait Locus (QTL) Mapping



Phenotypic data evaluation

- Frequency distribution?
 - normal
 - transgressive segregation
- ANOVA
 - are there significant differences for the trait among the plants?
- Heritability analysis
 - the higher the heritability the higher proportion of variation in the trait is due to genetic causes
- If two traits are highly correlated, it may indicate that the same QTLs influence both traits (pleiotropy)



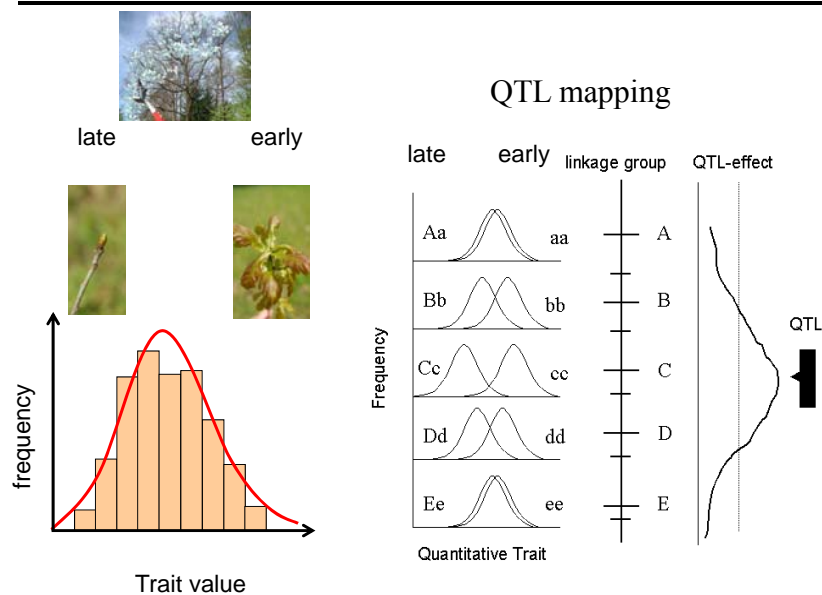
Genetic data analysis

- Parental screening (only markers polymorphic among parents can be useful)
- Genotyping markers in the segregating population
- Check for segregation distortion
- Linkage mapping

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QTL mapping in oaks



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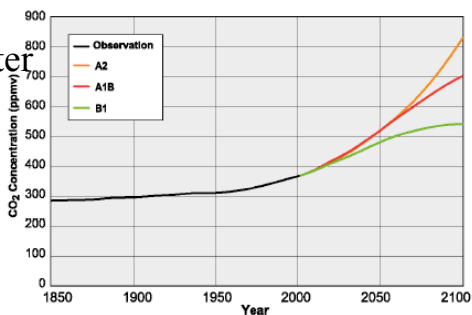


Adaptive phenotypic quantitative traits

- **Stomatal traits (stomatal density)**

Important to cope with:

- changes in levels of atmospheric CO₂
- changes in availability of water
- higher growth rate
- higher biomass



Roeckner et al. (2006) Climate Projections for the 21st century. Max Planck Institute for Meteorology.

ЛЕОМНКА: Вакцина 4 августа 2017, Броннус. #1

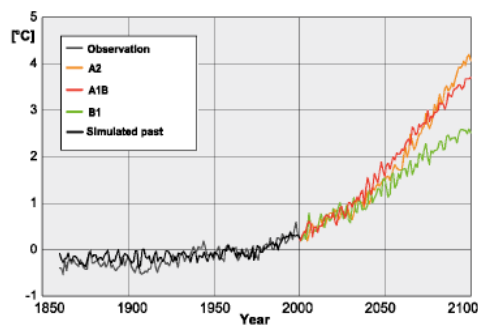
Adaptive phenotypic quantitative traits

- **Bud phenology traits:**

- longer growth season
- changes in the time of bud burst

- **Frost and drought injuries**

- **Insect damages**



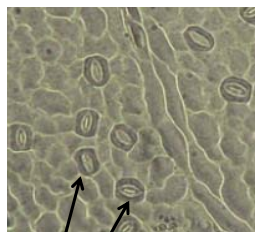
Roeckner et al. (2006) Climate Projections for the 21st century. Max Planck Institute for Meteorology.

ЛЕОМНКА: Вакцина 4 августа 2017, Броннус. #1

I. Stomatal density

- Aim of the study:

Characterization of the genetic basis of stomatal density and plant growth under non-waterstress conditions



stomata



Q. robur x *Q. robur* subsp. *slavonica*, 387 full-sibs

ЦЕХОМІКА: Висвітлення 4 липня 2017, Брест, #1



Effects of QTL for stomatal density (SD) on growth parameters

Table 3 Effects of quantitative trait loci (QTLs) for stomatal density on growth parameters and terminal buds

Traits	Interval analysis			Single-point analysis				Pleiotropic effect						Additional traits
	lg	LOD	LM (cM)	SEM	P-value	Permut	PVE	D	NA-	Naa	Trait	Sign	D	
SD2	1m		ns	QrZag101	0.041	c	3.8	-4.7	40	47	H1	(+)	-1.5	H2
SD3	4f	2.0	148*	E38M67-175	0.038	c	10.2	-6.8	17	17	H2	(+)	-7.1	R3
SD2	4f	3.6	171****	E36M63-114	0.027	c	4.2	-5.0	36	52	dR2	(+)	-0.57	R3
SD3	5m	2.1	59.6**	QrZag73	0.0001	b	10.4	6.1	47	37	R3	(+)	0.99	
SD2	5m	1.6	84.9*	E42M74-432	0.002	c	7.1	6.4	47	42	H1	(+)	1.7	
SD3	7f	2.9	66.6***	QpZag9	0.0014	c	7.2	5.0	55	38	R3	(+)	1.02	
SD3	10m		ns	QrZag65	0.0017	c	9.9	5.5	40	26	dH2	(-, +)	-8.5	dR1
SD2	11m	3.9	24.1****	E41M74-131	0.0001	b	15.9	-10.3	21	38	H2	(+)	-13.4	H1, H2, H3, R1, dH1, dH2

Lg, linkage group; LOD, LOD score of the interval analysis; LM, position of the maximum LOD score. Permut: significance at the 5% level; a, genomewide; b, chromosomewise; c, single marker analysis. NA-, number of individuals investigated for marker presence; Naa, number of individuals investigated for marker absence. Direction of pleiotropic effects on other traits is shown by the sign (+) or (-) indicating that higher stomatal density is associated with a higher or a lower phenotypic mean in growth parameters.

ЦЕХОМІКА: Висвітлення 4 липня 2017, Брест, #1



Perspectives

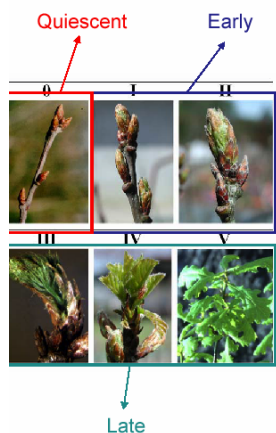
- Different water regimes
- Mapping of QTLs for carbon isotope composition (water use efficiency)
- Integration of candidate genes for this character (e.g. ERECTA)
- Comparative QTL mapping

Masle et al. (2005) *The ERECTA gene regulates plant transpiration efficiency in Arabidopsis*. *Nature* 436, 866-870

ГЕНОМКА: Введение 4 августа 2017, Бронин. #1



II. Bud burst



Bud stages 1 to 5
(according to Derory *et al.* 2006)

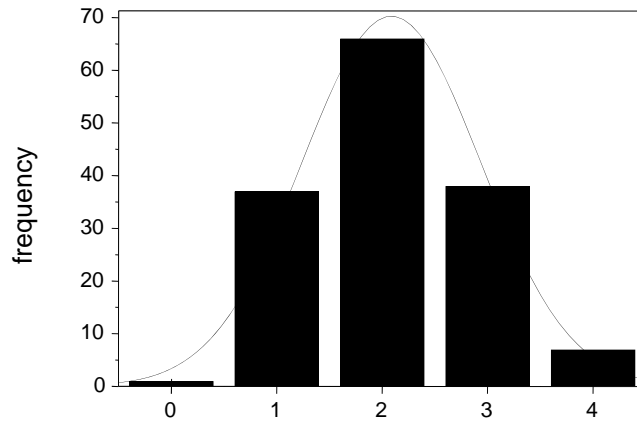


Q. robur x *Q. robur* subsp. *slavonica*, 387 full-sibs

ГЕНОМКА: Введение 4 августа 2017, Бронин. #1



Frequency distribution



Bud burst 2003, 30th April

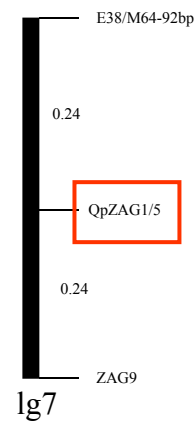
Gailing, O., Kremer, A., Steiner, W., Hattermer, H.H. & R. Finkeldey. 2005. Results on quantitative trait loci for flushing date in oaks can be transferred to different segregating progenies. *Plant Biology* 7: 516-525.

ЛЕОМОНКА: Встреча 4 апреля 2017. Брошюра: #1



Comparative QTL mapping

- significant association with bud burst in different *Quercus* spp. progeny (Gailing *et al.* 2005, Scotti-Saintagne *et al.* 2004).
- in six full-sib families the 166 bp allele is associated with an early bud burst

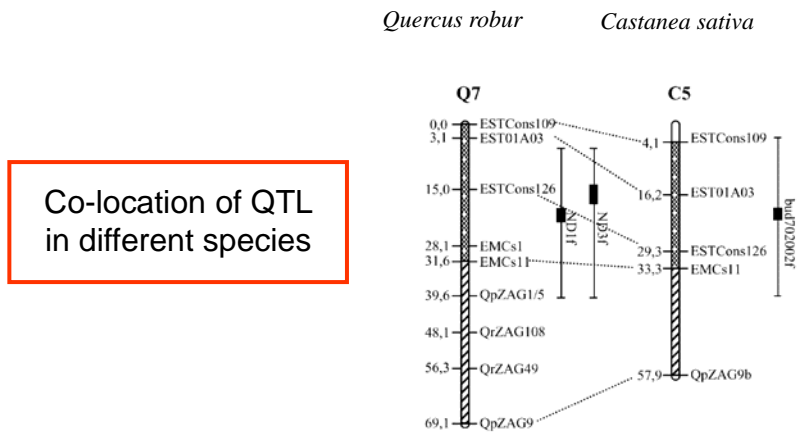


Gailing, O., Kremer, A., Steiner, W., Hattermer, H.H. & R. Finkeldey. 2005. Results on quantitative trait loci for flushing date in oaks can be transferred to different segregating progenies. *Plant Biology* 7: 516-525.

ЛЕОМОНКА: Встреча 4 апреля 2017. Брошюра: #1



Comparative QTL mapping in Fagaceae



ГЕНОМКА: Введение, 4 апреля 2017, Бронни, #1



QTL limitations

- Information on QTL locations and effects is specific to a particular population and cannot be readily transferred to another population
- QTL analysis detects chromosome regions, not genes, that influence traits. QTL locations have large confidence intervals, often greater than 30 cM
- It is difficult to distinguish two closely linked QTLs, those that are less than 20 cM apart.
- When two QTLs are linked “in repulsion”, it may not be possible to detect the QTL, because the effects of the associated alleles cancel each other out.

ГЕНОМКА: Введение, 4 апреля 2017, Бронни, #1



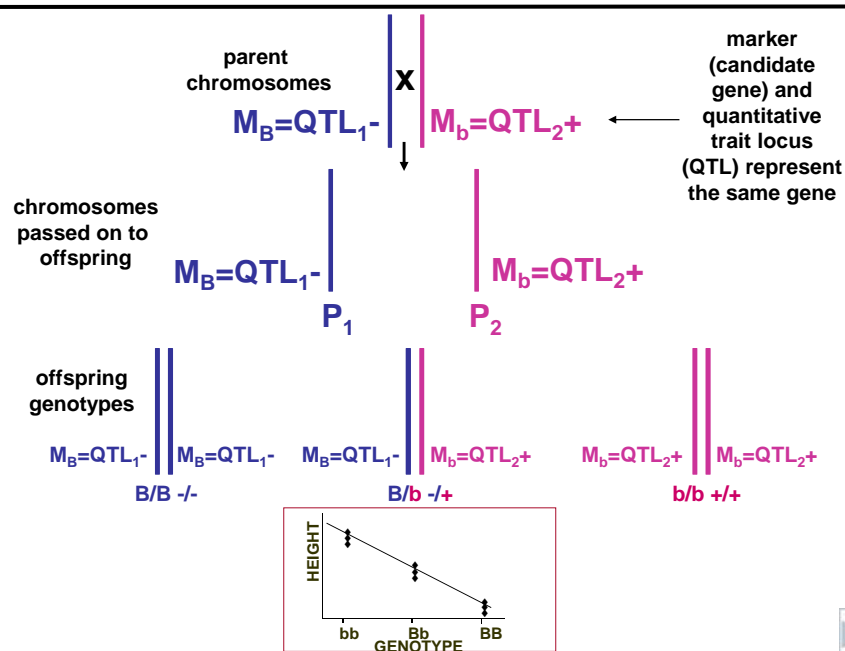
Quantitative Trait Locus (QTL) Mapping

- *Much information has been learned from QTL mapping, but we still mostly do not know what genes and alleles are involved !*
- Situation is changing now due to
 - QTL mapping using candidate genes, and
 - genome-wide association studies based on high-density genotyping via sequencing

ГЕНОМИКА: Введение, 4 августа 2017, Бронниц, #1



Candidate gene based QTL mapping



ГЕНОМИКА: Введение, 4 августа 2017, Бронниц, #1



Criteria to select candidate genes

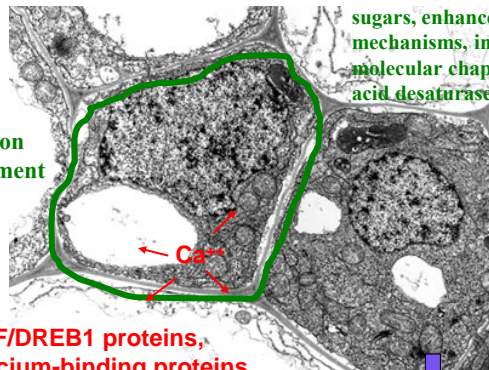
- 1) Physiology and biochemistry, high similarity/homology to genes with well known effects, mutations
- 2) Differential expression
- 3) Positional association with quantitative trait loci (QTLs)

ГЕОМНИКА: Биология, 4 апреля 2017, Бронник, #1



1) Physiological mechanisms involved in low temperature and drought tolerance

Cytoskeleton rearrangement



CBF/DREB1 proteins, Calcium-binding proteins and other transcription factors and regulatory proteins in a low temperature signal transduction pathway

Stabilization of membranes and membrane fluidity via changes in lipid composition, accumulation of sucrose and other simple sugars, enhancement of antioxidative mechanisms, induction of genes encoding molecular chaperones, dehydrins, fatty acid desaturases etc.

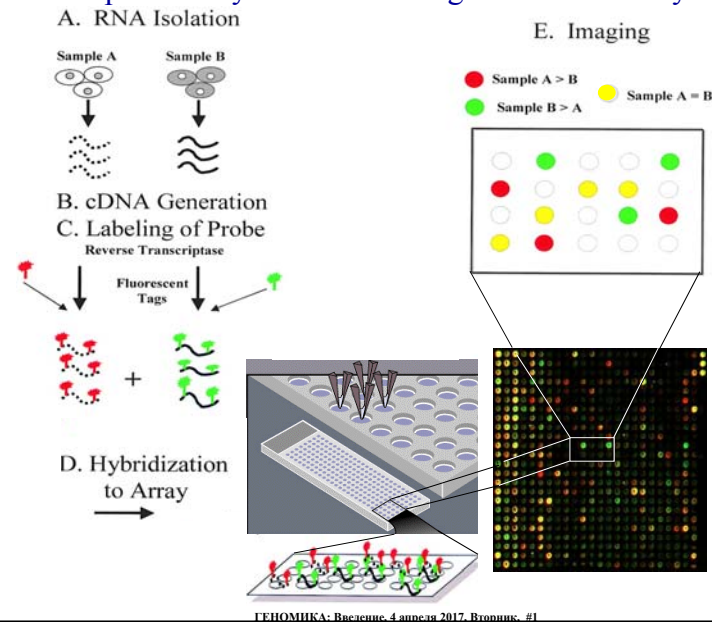
Increase of antifreeze proteins and level of cryoprotectants, such as proline, for instance, that prevent ice formation

ГЕОМНИКА: Биология, 4 апреля 2017, Бронник, #1



2) Differential gene expression study

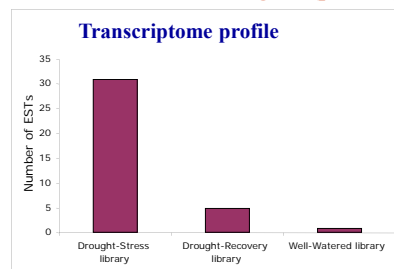
Comparative hybridization using DNA microarray



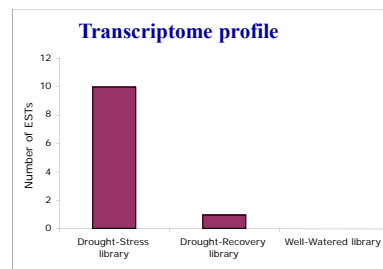
2) Differential gene expression *in silico*: transcriptome profiling using NGS

Candidate genes for drought resistance in loblolly pine:

ABA and WDS induced gene (pLP3-3)



Dehydrin (Dhn-1)



Lorenz W.W., Sun F., Liang C., Kolychev D., Wang H., Zhao X., Cordonnier-Pratt M.-M., Pratt L.H., Dean J.F.D. 2006. Water stress-responsive genes in loblolly pine (*Pinus taeda*) roots identified by analyses of expressed sequence tag libraries. *Tree Physiology* 26: 1–16.

3) Collocation of adaptive trait related genes with QTLs controlling adaptive traits

Positional candidate genes:

LEA-II

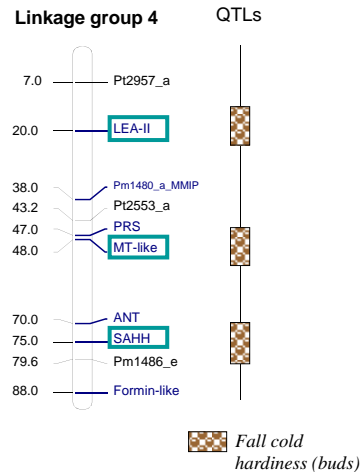
(late embryogenesis abundant type II)
dehydrin-like protein
Induced by cold

MT-like

(metallothionein-like protein)
stress-induced;
downregulated under water deficit

SAHH

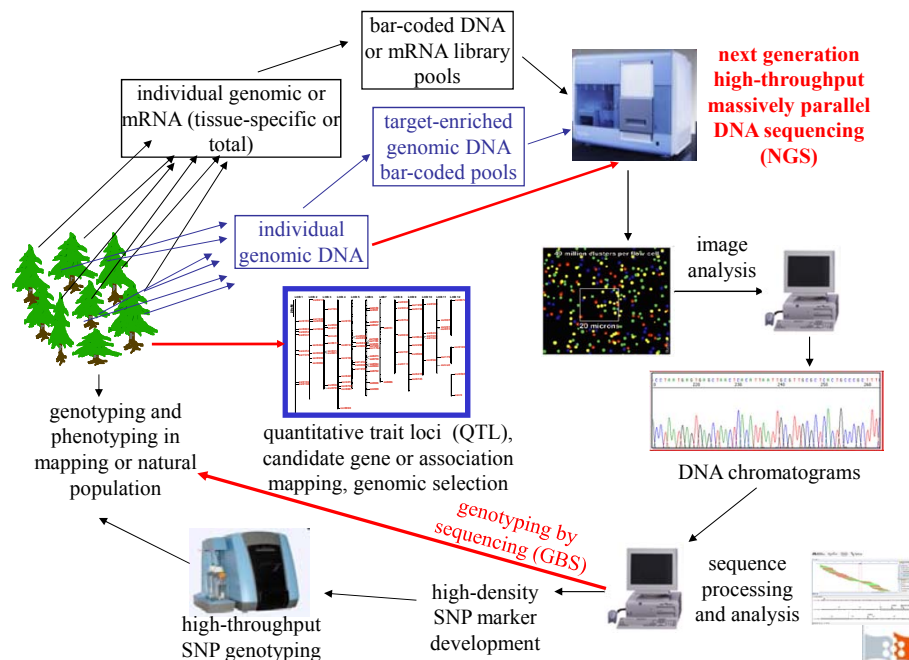
(S-adenosyl-L-homocysteinase hydrolase)
upregulated under water deficit



Wheeler N.C., Jermstad K.D., Krutovsky K.V. *et al.* 2005. Mapping of quantitative trait loci controlling adaptive traits in coastal Douglas-Fir. IV. Cold-hardiness QTL verification and candidate gene mapping. *Molecular Breeding* 15: 145-156.

ГЕОМІКА: Висвітлює 4 лютого 2017. Брошура: #1

Genomic markers development and genotyping using next generation sequencing



ГЕОМІКА: Висвітлює 4 лютого 2017. Брошура: #1

Association mapping

- AKA: Linkage disequilibrium mapping
- Aim: (like in QTL) to find a statistical association between a genetic marker and a quantitative trait

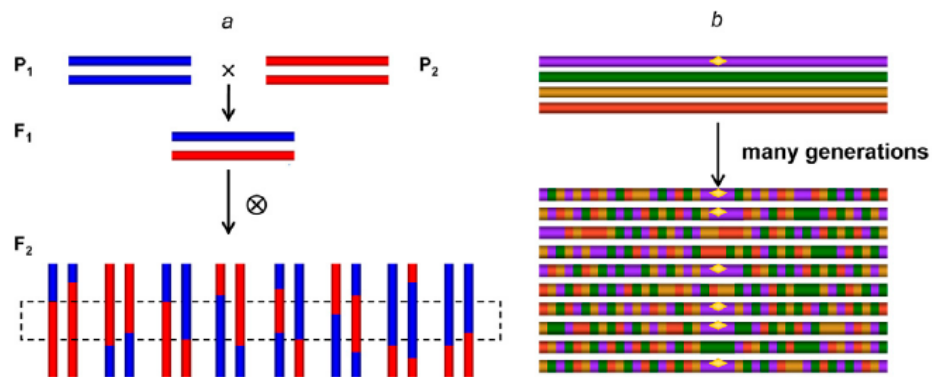
BUT

- Association mapping is performed at the population level (unrelated or distantly-related individuals sampled from a population)
- In association mapping, the genetic markers usually must lie close to or within genes responsible for a measured trait
- The goal is to identify the actual genes affecting that trait, rather than just (relatively large) chromosomal segments

ГЕНОМКА: Введение. 4 апреля 2017. Бронius. #1



QTL mapping (a) vs. Associations Mapping (b)

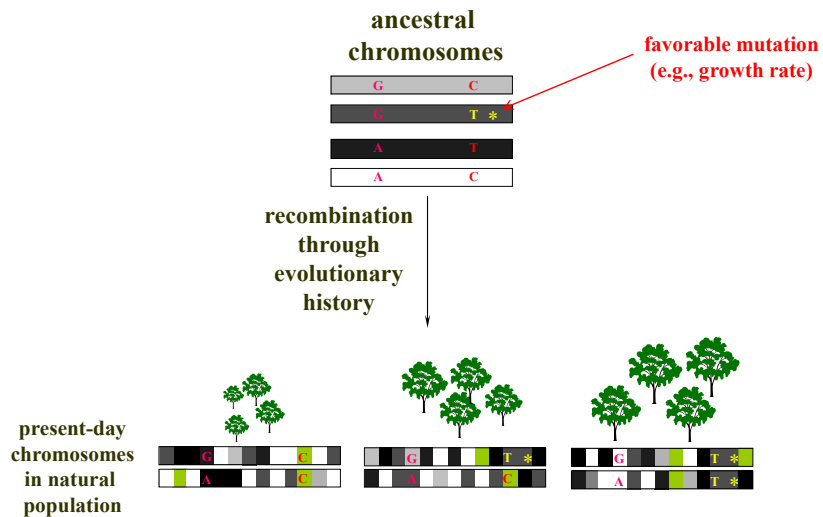


Zhu et al. 2008

ГЕНОМКА: Введение. 4 апреля 2017. Бронius. #1



Association Mapping using SNPs



ЛЕОМОНКА: Введение, 4 августа 2017, Бронниц, #1



Linkage disequilibrium (LD)

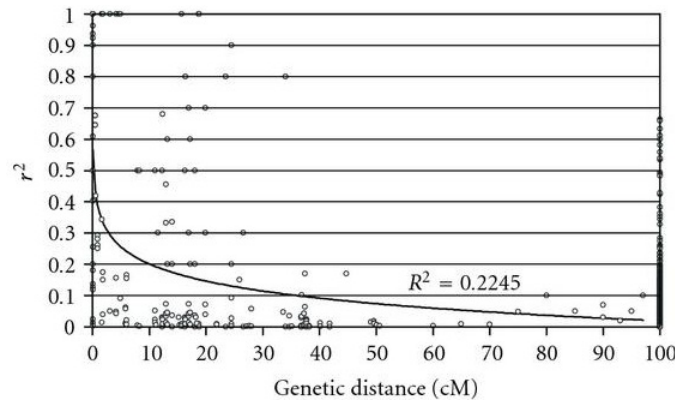
- LD: The non-independence of alleles at different loci (certain combinations of alleles across loci occur more often than expected by chance alone)
- LD can be caused by various combinations of many different factors, including selection, new mutations, population genetic structure, interbreeding of genetically divergent populations (admixture) and population bottlenecks
- LD is reduced by recombination, which "shuffles" the combinations of alleles at different loci

ЛЕОМОНКА: Введение, 4 августа 2017, Бронниц, #1



Linkage disequilibrium (LD)

- LD decay very fast in most sexually reproducing species with large effective population size:



Linkage disequilibrium (LD) decay plot depicted from the pairwise LD (r^2) and genetic distance (cM) values measured between all pairs of linked markers. A pairwise LD values (measured as r^2 parameters) are plotted against a pairwise genetic distances (cM). Inner fitted trend line is a nonlinear logarithmic regression curve of r^2 on genetic distance (R^2).

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Genome-wide random marker based association mapping vs.

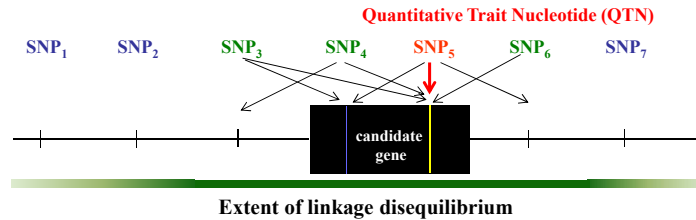
Candidate gene based association mapping

- Association mapping based on random genetic markers relies on linkage disequilibrium (LD) between gene markers and the actual causative polymorphism in genes that causes the differences in the phenotypic trait.
- Association mapping based on candidate genes relies on direct association of the actual causative polymorphism in the candidate gene with the differences in the phenotypic trait.

ГЕНОМКА: Введение. 4 августа 2017. Бронius. #1



Genome-wide random marker based association mapping vs. Candidate gene based association mapping



Linkage Disequilibrium (LD) is a nonrandom association of alleles at linked loci

ЛЕОМНКА: Введение, 4 августа 2017, Бронни, #1



Genomic approaches to study complex adaptive traits

- High-density genome-wide genotyping via high-throughput NGS sequencing or high-density single nucleotide polymorphism (SNP) assays
- Genome-wide Association Studies using SNPs
- Functional genomic markers (candidate genes)
- Detection genes-outliers:
 - with unusually high or low differentiation
 - with unusually high or low expression

Krutovsky K.V. & D.B. Neale. 2005. Forest genomics and new molecular genetic approaches to measuring and conserving adaptive genetic diversity in forest trees, pp. 369-390 in *Conservation and Management of Forest Genetic Resources in Europe*, edited by Th. Geburek and J. Turok. Arbora Publishers, Zvolen.

González-Martínez S.C., Krutovsky K.V., Neale D.B. 2006. Forest tree population genomics and adaptive evolution. *New Phytologist* **170**(2): 227-238.



Genomic approaches to study complex adaptive traits

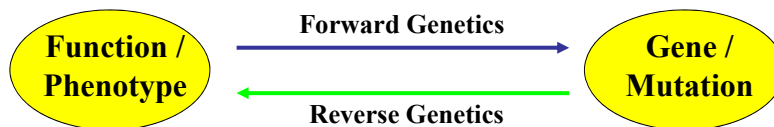
- Forward Genetics (from trait to genotype) vs. Reverse Genetics (from genotype to trait)
- Top down (pattern to process) and bottom up (process to pattern) approaches
- Neutral vs. functional markers

69

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Forward vs. Reverse Genetics



Forward Genetics (traditional): Starts with a phenotype and moves towards the gene

Reverse Genetics: Starts with a particular gene and assays the effect of its disruption or allelic effects

Genomics (via Association Genetics): $P \rightleftharpoons G$

What is the genetic basis of a particular phenotype?

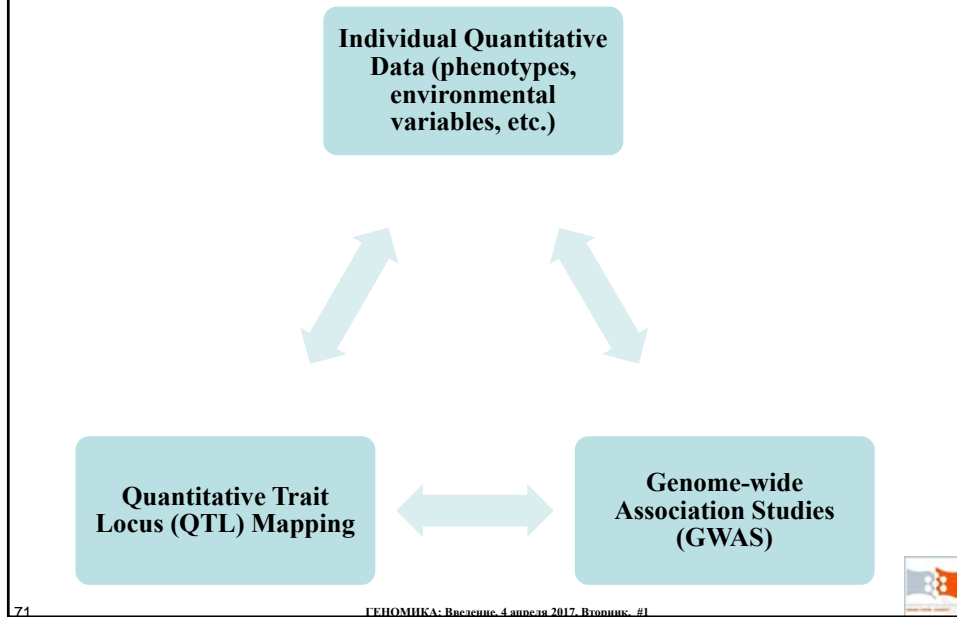
(How to determine the function of a gene, or the identity of genes responsible for a trait?)

70

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Association Genetics



Integrated approach to study evolutionary response

- 1) association of population substructure with particular environments and ecozones;
 - 2) identification of outliers - genes with unusually high differentiation (that could be a signature of positive or diversifying selection), or genes with unusually low differentiation (that could be a signature of balancing selection) that significantly above or below levels that are expected for selectively neutral markers;
 - 3) correlation of allele frequencies with environmental gradients; clinal variation, etc.;
 - 4) QTL and/or association mapping of candidate genes with adaptive trait related phenotypes;
 - 5) identification of genome-wide signatures of selection (LD, selective sweeps, etc.);
 - 6) intra- and inter-specific selective neutrality tests
- 72
- ГЕНОМИКА: Введение. 4 апреля 2017. Бронius. #1

- **XX century:**
Evolutionary theory + Genetics
= **Synthetic theory of evolution**
(Genetic theory of evolution or
Evolutionary Genetics)

↓ paradigm shift

from an individual as a main unit of evolution to
population genetics level of thinking



Theodosius Dobzhansky
(1900-1975)

- **XXI century:**
Molecular genetics + Bioinformatics
= **Genomics**

↓ paradigm shift

from genetics to genomics level of thinking

Krutovsky, K.V. 2006 From Population Genetics to Population Genomics of Forest Trees: Integrated
Population Genomics Approach. *Russ. J. of Genetics* 42(10): 1088–1100

ГЕНОМИКА: Введение. 4 августа 2017. Бронниц. #1

73

Key terms and definitions

- **Quantitative Trait Locus (QTL) mapping:** A QTL is a chromosomal region suspected to contain a gene (or cluster of genes) that contributes to the variation observed at a quantitative trait. QTLs are detected through linkage mapping experiments using progeny usually obtained in experimental crosses or pedigrees that segregate for both quantitative traits and genetic markers. QTL and genetic markers that are close together on a chromosome will tend to co-segregate.
- **Association mapping:** As in QTL mapping, the goal of association mapping is to find a statistical association between genetic markers and a quantitative trait. However, unlike QTL mapping, which is performed in the context of a pedigree, association mapping is performed at the population level: the genotypes of the candidate gene markers and the phenotypes of the corresponding trait are determined in a set of unrelated or distantly-related individuals sampled from a population. Association mapping relies on linkage disequilibrium (LD) between the markers and the actual causative genes (i.e., the actual polymorphism that causes the differences in the phenotypic trait). Hence association mapping is also referred to as 'LD mapping'. For association to be detected the genetic markers usually must be closely linked to genes (lie within or directly upstream or downstream of them) that contribute to the variation in that trait, and the goal is to identify the actual genes affecting that trait, rather than just (relatively large) chromosomal segments. Since population genetic structure (genetic differences that accumulate between populations) can cause LD even at unlinked loci, association analyses must account for population genetic structure whenever it is present in the population from which your sample has been drawn (Pritchard et al. 2000; Thornsberry et al. 2001).

74

ГЕНОМИКА: Введение. 4 августа 2017. Бронниц. #1

Key terms and definitions

- **Amino-acid or nucleotide multiple sequence alignment:** is a way of arranging the sequences of DNA, RNA, or protein to identify regions of similarity that may be a consequence of functional, structural, or evolutionary relationships between the sequences. Aligned sequences of nucleotide or amino acid residues are typically represented as rows within a matrix.
- **Contig:** (from *contiguous*) is a set of overlapping DNA sequences in a **multiple sequence alignment** that together represent a **consensus** region of DNA. In sequencing projects, a contig refers to overlapping sequence reads or to the overlapping clones that form a physical map of the genome that is used to guide sequencing and assembly. Contigs can thus refer both to overlapping DNA sequence and to overlapping physical segments (fragments) contained in clones depending on the context.
- **Consensus sequence:** is the calculated order of most frequent residues, either nucleotide or amino acid, found at each position in a **multiple sequence alignment**. It represents the results of a **multiple sequence alignment**, in which related sequences are compared to each other, and most frequent residues are calculated.
- **Basic Local Alignment Search Tool (BLAST):** is an algorithm for comparing amino-acid or nucleotide sequences using their alignment. A BLAST search enables a researcher to compare a query sequence with a library or database of sequences, and identify library sequences that resemble the query sequence above a certain threshold.
- **Expressed Sequence Tag (EST):** is a short sub-sequence of a mRNA/cDNA sequence. They are used to identify gene transcripts, and are instrumental in gene discovery, gene sequence determination and differential gene expression analysis (transcriptome profiling).
- **Unigene:** is a supposedly unique transcript that represents the same transcription locus (expressed gene or pseudogene), often inferred as a **consensus sequence** from **EST** based **multiple sequence alignment**.
- **SNP:** stands for *Single Nucleotide Polymorphism*. This refers to a particular nucleotide (or "base") in a DNA sequence that is variable within a species (or between related species). For example, at a certain position in a DNA sequence there may be a C (cytosine) present in some individuals but a T (thymine) present in others (C/T polymorphism). SNPs represent the most basic form of genetic polymorphism. There are tens of millions of SNPs present in the genome of a typical organism and can be used as genetic markers (SNP markers).

