Genetics and Bioinformatics

Kristel Van Steen, PhD²

Montefiore Institute - Systems and Modeling GIGA - Bioinformatics ULg

kristel.vansteen@uliege.be

Genome-wide Association Studies

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1 Setting the pace

1.a What can your spit tell you about your DNA?

The use of saliva

- People spit for a variety of reasons. We've all employed the technique to remove a hair or some other distasteful object from our mouths. People who chew tobacco do it for obvious reasons. Ball players do it because they're nervous, bored or looking to showcase their masculinity. And people in many different cultures spit on their enemies to show disdain.
- Thanks to a phenomenon known as direct-to-consumer genetic testing or at-home genetic testing, people are spitting today for a much more productive (and perhaps more sophisticated) reason -- to get a glimpse of their own DNA.

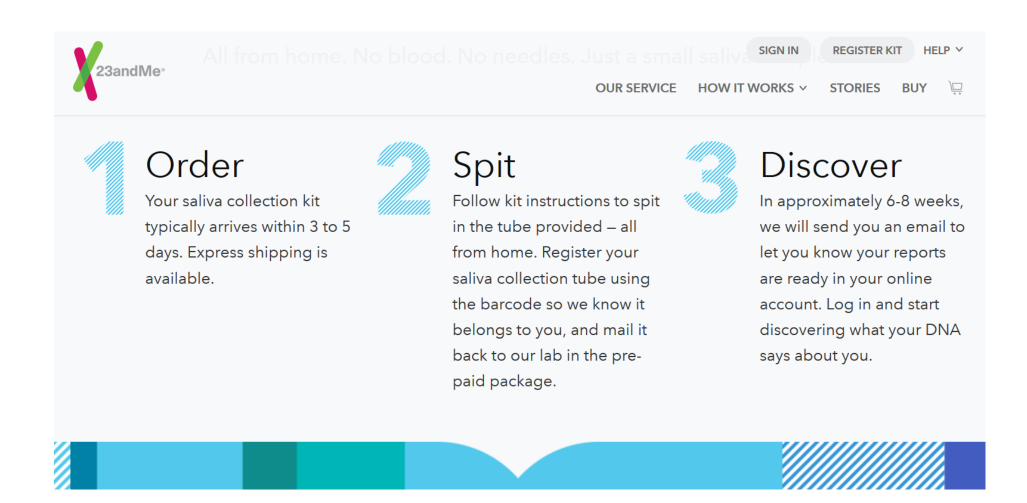
(science.howstuffworks.com)

From saliva to DNA

- Your saliva contains a veritable mother load of biological material from which your genetic blueprint can be determined.
- For example, a mouthful of spit contains hundreds of complex protein molecules enzymes -- that aid in the digestion of food.
- Swirling around with those enzymes are cells sloughed off from the inside of your cheek.
- Inside each of those cells lies a nucleus, and inside each nucleus, chromosomes, which themselves are made up of DNA



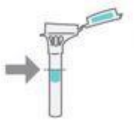
Commercial kits



Commercial kits



Do not eat, drink, smoke, chew gum, brush your teeth, or use mouthwash for at least 30 minutes prior to providing your sample.



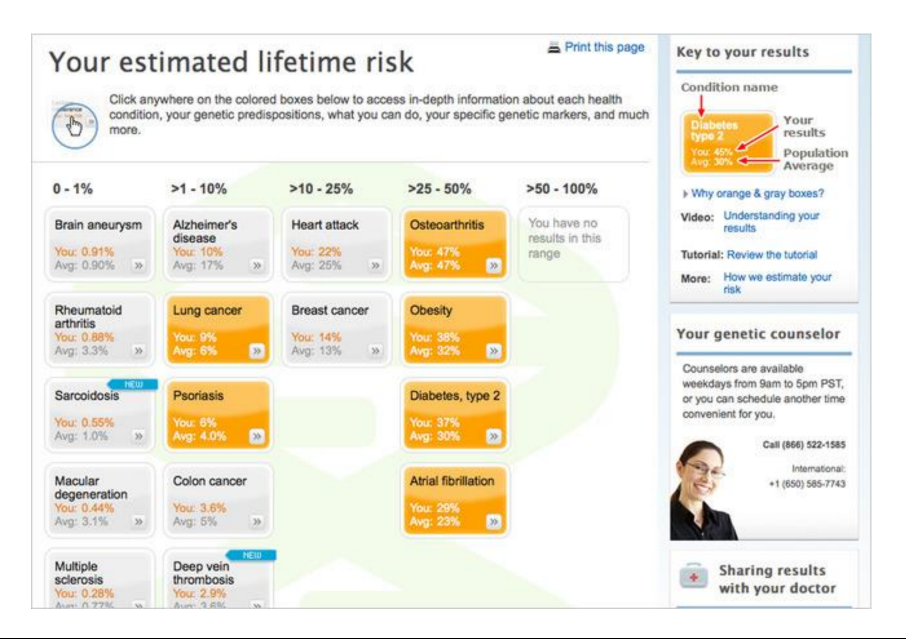
Collect the recommended volume of saliva. The recommended volume of saliva to provide is 2 mL, or about ½ teaspoon. Your saliva sample should be just above the fill line.



Provide your sample and add the stabilization buffer within 30 minutes. The full saliva sample should be collected within 30 minutes and the funnel contents should be released into the tube immediately. Waiting longer than 30 minutes may decrease the yield and quality of your DNA.



Cap securely before shipping. Remember to remove and discard the funnel lid and place the tube cap on securely before mailing your sample to our laboratory.



Can you handle the truth?

Service Provider:	23andMe	deCODEme	Navigenics
Arthritis	*		*
Asthma	•	*	
Bipolar/Depression			
Cardiovascular Disease	*	•	*
Multiple Sclerosis	*	*	*
Osteoporosis	•		
Parkinson's Disease			
Schizophrenia	*		
Thrombosis	*	*	2
Type 1/2 Diabetes	*	-	*

The 23andMe story

- Wojcicki founded 23andme in 2006 with Linda Avey and Paul Cusenza with a goal of upending conventional models of health care:
 - put sophisticated DNA analyses into the hands of consumers,
 - giving them information about health, disease and ancestry,
 - and allowing the company to sell access to the genetic data to fuel research.
- In 2013, that vision hit a snag. Wojcicki didn't think she needed regulatory approval to provide information about her customers' health risks. The US Food and Drug Administration (FDA) disagreed, and ordered the company to stop.

(source: https://www.nature.com/news/the-rise-and-fall-and-rise-again-of-23andme-1.22801)

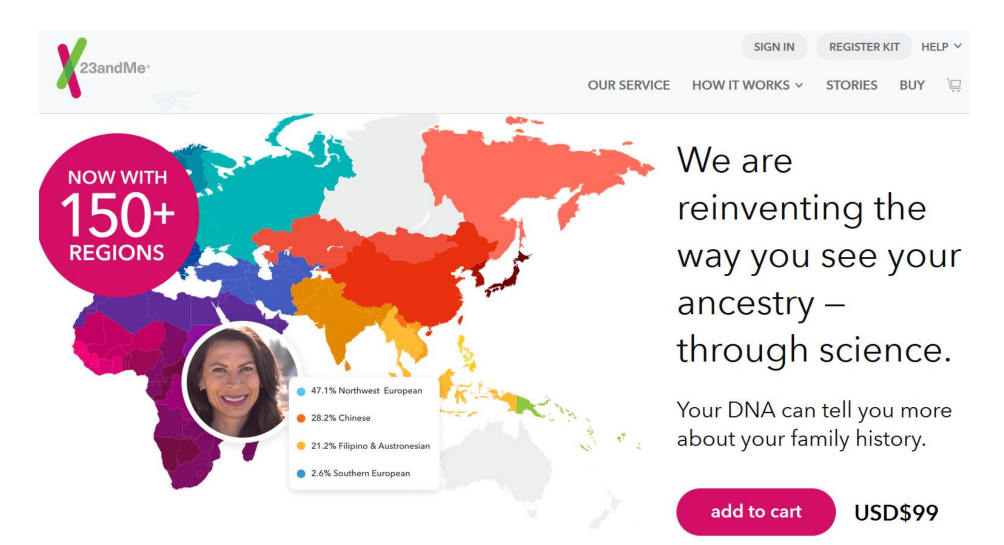
NewStatesman



SCIENCE & TECH 15 JANUARY 2015

23andMe: Why bother with predictions about yourself when you are almost certainly average?

Want to understand your genes? Call your parents.



The 23andMe story



• After years of effort, the pay-off came in April 2017, when the FDA agreed to allow 23andme to tell consumers their risks of developing ten medical conditions, including Parkinson's disease and late-onset Alzheimer's disease.

• With more than 2 million customers, the company hosts by far the largest collection of gene-linked health data anywhere

(source: https://www.nature.com/news/the-rise-and-fall-and-rise-again-of-23andme-1.22801)

What are (other) examples of genetic testing on demand?

- There are >2000 genetic tests available to physicians to aid in the diagnosis and therapy for >1000 different diseases. Genetic testing is performed for the following reasons:
 - presymptomatic testing for **estimating risk** developing disease
 - presymptomatic testing for **predicting** disease
 - conformational diagnosis of a symptomatic individual
 - carrier screening
 - prenatal screening
 - newborn screening
 - preimplantation genetic diagnosis (testing the early embryo after in vitro fertilisation)
 - forensic testing
 - paternal testing

How is genetic testing used clinically?

- Predictive medicine: determine whether an individual has an increased risk for a particular disease. Results from this type of test are usually expressed in terms of probability and are therefore less definitive since disease susceptibility may also be influenced by other genetic and non-genetic (e.g. environmental, lifestyle) factors. Examples of diseases that use genetic testing to identify individuals with increased risk include certain forms of breast cancer (BRCA) and colorectal cancer.
- Diagnostic medicine: *identify whether an individual has a certain genetic disease.* This type of test commonly detects a specific gene alteration but is often not able to determine disease severity or age of onset. It is estimated that there are >4000 diseases caused by a mutation in a single gene. Examples of diseases that can be diagnosed by genetic testing includes cystic fibrosis and Huntington's disease.

How is genetic testing used clinically?

- Pharmacogenomics: classifies subtle variations in an individual's genetic makeup to determine whether a drug is suitable for a particular patient, and if so, what would be the safest and most effective dose. Learn more about pharmacogenomics. → DNA passports ... are no science fiction!
- Whole-genome and whole-exome <u>sequencing</u>: examines the entire genome or exome to discover genetic alterations that may be the cause of disease. Currently, this type of test is most often used in complex diagnostic cases, but it is being explored for use in asymptomatic individuals to predict future disease. See also "The promise and challenges of next-generation genome sequencing for clinical care" (JAMA Intern Med. 2014)

The basics of genetic tests

- As we will see, we can measure (genetic) variation between individuals at several positions on the genome, using so-called molecular markers such as Single Nucleotide Polymorphisms (SNPs)
- To run a SNP test, scientists can embed a subject's DNA into for instance a small silicon chip containing reference DNA from both healthy individuals and individuals with certain diseases.
- By analyzing how the SNPs from the subject's DNA match up with SNPs from the **reference DNA**, the scientists can determine if the subject might be predisposed to certain diseases or disorders.

Talking about "reference": reference genomes

- A reference genome (also known as a *reference assembly*) is a digital nucleic acid sequence database, assembled by scientists as a representative example of a species' set of genes.
- As they are often assembled from the sequencing of DNA from a number of donors, reference genomes do not accurately represent the set of genes of any single person. Instead a reference provides a haploid mosaic of different DNA sequences from each donor.
- For example GRCh37, the Genome Reference Consortium human

genome (build 37) is derived from thirteen anonymous volunteers from Buffalo, New York



"Wellcome genome bookcase" by Russ London at en.wikipedia. Licensed under CC BY-SA 3.0 via Commons https://commons.wikimedia.org/wiki/File:Wellcome_genome_bookc ase.png#/media/File:Wellcome_genome_bookcase.png

Talking about "reference": reference panels

PLOS ONE

RESEARCH ARTICLE

Comparison of HapMap and 1000 Genomes Reference Panels in a Large-Scale Genome-Wide Association Study

Check for updates

OPEN ACCESS

Citation: de Vries PS, Sabater-Lleal M, Chasman DI, Trompet S, Ahluwalia TS, Teumer A, et al. (2017) Comparison of HapMap and 1000 Genomes Reference Panels in a Large-Scale Genome-Wide Association Study. PLoS ONE 12 (1): e0167742. doi:10.1371/journal.pone.0167742

Paul S. de Vries^{1,2}, Maria Sabater-Lleal³, Daniel I. Chasman^{4,5}, Stella Trompet^{6,7}, Tarunveer S. Ahluwalia^{8,9}, Alexander Teumer¹⁰, Marcus E. Kleber¹¹, Ming-Huei Chen^{12,13}, Jie Jin Wang¹⁴, John R. Attia^{15,16}, Riccardo E. Marioni^{17,18,19}, Maristella Steri²⁰, Lu-Chen Weng²¹, Rene Pool^{22,23}, Vera Grossmann²⁴, Jennifer A. Brody²⁵, Cristina Venturini^{26,27}, Toshiko Tanaka²⁸, Lynda M. Rose⁴, Christopher Oldmeadow^{15,16}, Johanna Mazur²⁹, Saonli Basu³⁰, Mattias Frånberg^{3,31}, Qiong Yang^{13,32}, Symen Ligthart¹, Jouke J. Hottenga²², Ann Rumley³³, Antonella Mulas²⁰, Anton J. M. de Craen⁷, Anne Grotevendt³⁴, Kent D. Taylor^{35,36}, Graciela E. Delgado¹¹, Annette Kifley¹⁴, Lorna M. Lopez^{17,37,38}, Tina L. Berentzen³⁹, Massimo Mangino^{27,40}, Stefania Bandinelli⁴¹, Alanna C. Morrison¹, Anders Hamsten³, Geoffrey Tofler⁴², Moniek P. M. de Maat⁴³, Harmen H. M. Draisma^{22,44}, Gordon D. Lowe⁴⁵, Magdalena Zoledziewska²⁰, Naveed Sattar⁴⁶, Karl J. Lackner⁴⁷, Uwe Völker⁴⁸, Barbara McKnight⁴⁹, Jie Huang⁵⁰, Elizabeth G. Holliday⁵¹, Mark A. McEvoy¹⁶, John M. Starr^{17,52}, Pirro G. Hysi²⁷, Dena G. Hernandez⁵³, Weihua Guan³⁰, Fernando Rivadeneira^{1,54}, Wendy L. McArdle⁵⁵, P. Eline Slagboom⁵⁶ Tanja Zeller^{57,58}, Bruce M. Psaty^{59,60}, André G. Uitterlinden^{1,54}, Eco J. C. de Geus^{22,23}, David J. Stott⁶¹, Harald Binder⁶², Albert Hofman^{1,63}, Oscar H. Franco¹, Jerome I. Rotter^{64,65}, Luigi Ferrucci²⁸, Tim D. Spector²⁷, Ian J. Deary^{17,66}, Winfried März^{11,67,68} Andreas Greinacher⁶⁹, Philipp S, Wild^{70,71,72}, Francesco Cucca²⁰, Dorret I, Boomsma²², Hugh Watkins⁷³, Weihong Tang²¹, Paul M. Ridker^{4,5}, Jan W. Jukema^{6,74,75}, Rodney J. Scott^{76,77}, Paul Mitchell¹⁴, Torben Hansen⁷⁸, Christopher J. O'Donnell^{13,79}, Nicholas L. Smith^{60,80,81}, David P. Strachan⁸², Abbas Dehghan^{1,83}*

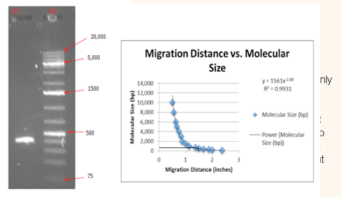
1.b Speaking the language

Types of molecular markers (Schlötterer 2004)

OPINION

The evolution of molecular markers — just a matter of fashion?

Christian Schlötterer



Being able to distinguish between genotypes

Enzymes are proteins that act as biological catalysts.

map of Drosophila melanogaster was built by Sturtevant using phenotypic markers¹. Howcontinuous improvement in the way in which we assay genetic variation; that is, the latest marker systems are the most informative ones. Nevertheless, in reviewing the history of molecular markers and their pros and cons, I argue that there are only a few conceptually different classes of marker and that recently developed high-throughput methods might not be unconditionally superior to more traditional approaches.

Allozymes

The first true molecular markers to be established were allozymes (a term that originates from a contraction of the phrase allelic varients of enzymes'). The principle of allozyme markers is that protein variants in enzymes can be distinguished by native gel electrophoresis according to differences in size and charge caused by amino-acid substitutions. To visualize the allozyme bands, the electrophoretic gels are treated with enzyme-specific stains that contain substrate for the enzyme, cofactors and an oxidized salt (for example, nitro-blue tetrasample sizes are typically studied in allozyme surveys. Nevertheless, the number of informative marker loci is too small to use allozymes for mapping and ASSOCIATION STUDIES⁸. Furthermore, surveys of natural variation based on allozymes were often challenged by nonneutral evolution of some of the markers used (see, for example, REFS 9–11).

The arrival of DNA-based markers

One of the criticisms levelled at allozyme markers is that they are an indirect and insensinve method of detecting variation in DNA. A more direct molecular marker would survey DNA variation itself, rather than rely on variations in the electrophoretic mobility of proteins that the DNA encodes. Another important advantage that DNA-based markers have over allozymes is that they allow the number of mutations between different alleles to be quantified. Given these unambiguous advantages, the arrival of DNA manipulation techniques promoted a shift from enzymebased to DNA-based markers.

"...the arrival of DNA manipulation techniques promoted a shift from enzyme-based to DNA-based markers."

Types of molecular markers

- Enzyme based
- DNA sequence-based:
 - Nowadays, genetic markers represent sequences of DNA which have been traced to specific locations on the chromosomes and associated with particular traits: according to NHGRI:

"trait" is **the manifestation of the product of a gene that is coded for by the DNA;** a "phenotype" is **an individual's observable traits, such as height, eye color, and blood type**. "Phenotype" is sometimes used interchangeably with the word trait, and "phenotype" may also define a whole compendium of traits together.

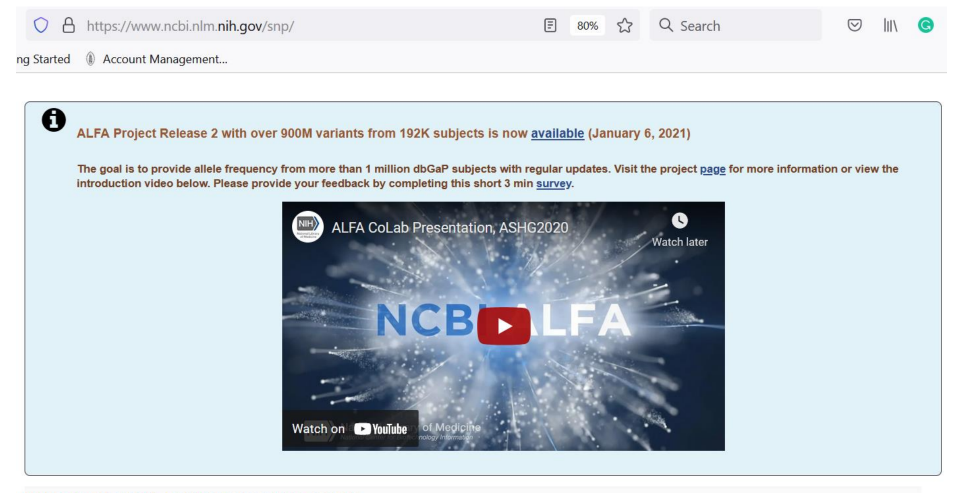
 Genet markers demonstrate polymorphism, which means that the genetic markers in different organisms of the same species are different.

Marker	Advantages	Disadvantages	
SNPs	 Low mutation rate High abundance Easy to type New analytical approaches are being developed at present Cross-study comparisons are easy; data repositories already exist 	 Substantial rate heterogeneity among sites Expensive to isolate Ascertainment bias Low information content of a single SNP 	Be critical (date of publication =
Microsatellites	 Highly informative (large number of alleles, high heterozygosity) Low ascertainment bias Easy to isolate A microsatellite is a tract 	 High mutation rate Complex mutation behaviour Not abundant enough Difficult to automate Cross-study comparisons 	2004) Hence, it is
Allozymes	of repetitive DNA in which certain DNA motifs (ranging in length from one to six or more base pairs) are repeated,	 require special preparation Requirement for fresh or frozen material Some loci show protein instability Limited number of available markers Potentially direct target of selection 	important to keep the historical time lines and achievements in
RAPDs and derivatives	 typically 5–50 times bands, which can then be further characterized individually (for example, converted into single locus markers) 	 Low reproducibility Mainly dominant Difficult to analyse Difficult to automate Cross-study comparisons are difficult 	mind
DNA sequencing	 Highest level of resolution possible Not biased Cross-study comparisons are 	 Still significantly more expensive than the other techniques 	Van Steen K

		Single Nucleotide Polymorphisms (SNPs)	Frequency in general population
C G Base F	Pairs Adenine Thymine	G	95%
	Guanine Cytosine — Sugar phosphate backbone	Α	5% > 1%

- Variations in single base, i.e., one base substituted by another base
- In theory: four different nucleotides possible at base
- In practice: generally only two different nucleotides observed
- Definition strict and loose:
 - Strict: minor allele frequency \ge 1%
 - \circ Loose: \geq 2 nucleotides observed in two individuals at position
- Nomenclature:
 - ss-number (submitted SNP number)
 - o rs-number: searchable in dbSNP, mapped to external resources, unique
 - o rs-numbers do not provide information about possible function of SNP
 - Alternative: nomenclature of Human Genome Variation Society

(Ziegler and Van Steen, Brazil 2010)



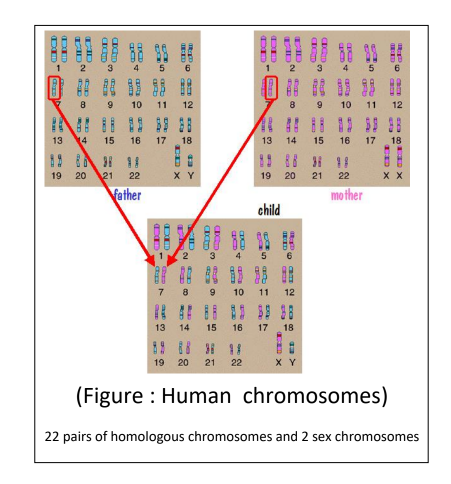
How to Search dbSNP. Additional search terms are here.

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Genes

- The **gene** is the basic physical unit of inheritance.
- Genes are passed from parents to offspring and contain the information needed to specify traits.
- They are arranged, one after another, on structures called chromosomes.
- A chromosome contains a single, long DNA molecule, only a portion corresponds to a single gene.



Genes

- An annotation (irrespective of the context) is a note added by way of explanation or commentary.
- Genome annotation is the process of identifying the locations of genes and all of the coding regions in a genome and determining what those genes do.
 - Once a genome is sequenced, it needs to be annotated to make sense of it

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	FunctionalAnnotation (13)
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Packages found under FunctionalAnnotation:

DO.dbJiang LiA set of annotation maps describing entire Disease OntologyGO.dbBioconductor Package MaintainerA set of annotation maps describing entire Gene OntologyhumanCHRLOCBiocore Data TeamA data package containing annotation for humanCHRLOCKEGG.dbBioconductor Package MaintainerA set of annotation maps for KEGGKEGG.dbBioconductor Package MaintainerA set of annotation maps for KEGGMeSH.AOR.dbKoki TsuyuzakiA set of annotation maps describing entire MeSHMeSH.dbKoki TsuyuzakiA set of annotation maps describing entire MeSHMeSH.PCR.dbKoki TsuyuzakiA set of annotation maps describing entire MeSH	ale -
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Alleles

- Allele: one of several alternative forms of DNA sequence at specific chromosomal location
 - Alleles A,B
- **Polymorphism**: often used to indicate the existence of at least 2 alleles at a single "locus"
- Homozygosity (homozygous): both alleles identical at locus
 - AA or BB
- Heterozygosity (heterozygous): different alleles at locus
 - AB

[Mutations ~polymorphisms (see later)]

Hunting for genes to answer relevant questions

- Developing new and better tools to make gene hunts faster, cheaper and practical for any scientist was a primary goal of the Human Genome Project (HGP).
- One of these tools is genetic mapping, the first step in isolating a gene.
 Genetic mapping in the early days can offer firm evidence that a disease transmitted from parent to child is linked to one or more genes. It also provides "clues" about where the gene lies.
- Genetic maps have been used successfully to find the single gene responsible for relatively rare inherited disorders, like cystic fibrosis, but have also been useful as a guide to identify the possible many genes underlying more common disorders, like asthma.

How to generate a genetic map?

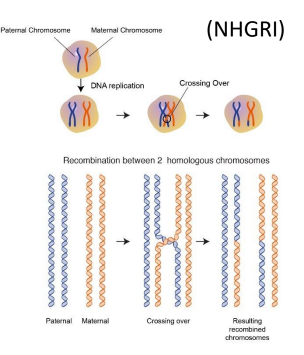
- To understand what genetic mapping is, let us look into genetic map construction itself.
- To produce a genetic map, researchers collect blood or tissue samples from **family members** where a certain disease or trait is prevalent.
- Using various laboratory techniques, the scientists isolate DNA from these samples and examine it for the unique patterns of bases seen only in family members who have the disease or trait. These characteristic molecular patterns are (also) referred to as polymorphisms, or markers.
- Before researchers identify the gene responsible for the disease or trait, DNA markers can tell them roughly where the gene is on the chromosome.

How is this possible?

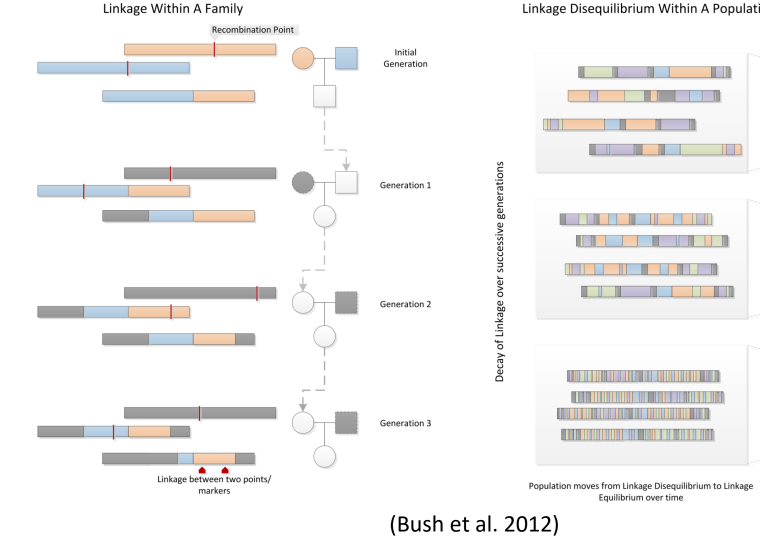
How to generate a genetic map? (continued)

• This is possible because of a genetic process known as recombination.

"As eggs or sperm develop within a person's body, the 23 pairs of chromosomes within those cells exchange - or recombine - genetic material. If a particular gene is close to a DNA marker, the gene and marker will likely stay together during the recombination process, and be passed on together from parent to child. So, if each family member with a particular disease or trait also inherits a particular DNA marker, chances are high that the gene responsible for the disease lies near that marker."



How to generate a genetic map? (continued)



Linkage Disequilibrium Within A Population

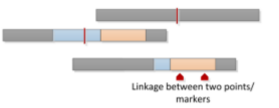
Initial Generation

100 Generations

1000 Generations

How to generate a genetic map? (continued)

• The more DNA markers there are on a genetic map, the more likely it is that one will be closely linked to a disease gene - and the easier it will be for researchers to zero-in on that gene.



 One of the first major achievements of the HGP was to develop dense maps of markers spaced evenly across the entire collection of human DNA.

(http://www.genome.gov/10000715#al-3)

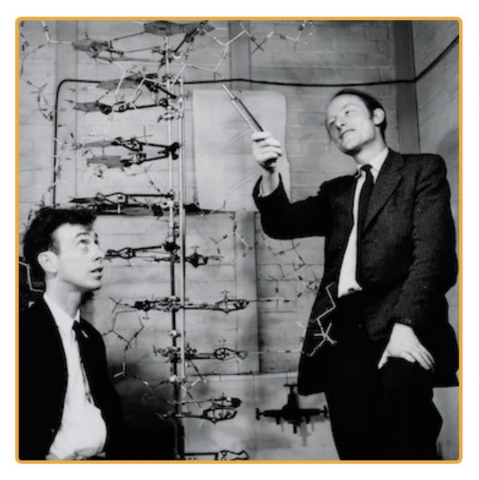
1.c "The Human Genome Project"

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Central Information Central Information Central Information Central Information Research Model Organisms Educational Resources An Interactive Timeline of the Human Genome [unlockinglifescode.org] An interactive, hyper-linked timeline of genetics that takes the reader from Mendel (1865) to the complet the mapping of the human genome (2003). The Human Genome: A Decade of Discovery, Creating a Healthy Future A workshop for science reporters about the 10th anniversary of the completion of the draft sequence of human genome and to look at the future of genomic research.



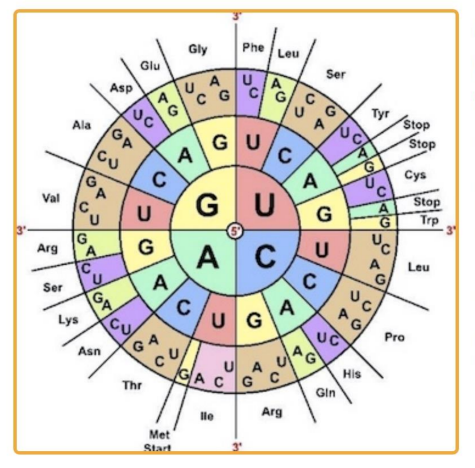
Gregor Mendel, the father of modern genetics, presents his research on experiments in plant hybridization

Gregor Mendel, a 19th century Augustinian monk, is called the father of modern genetics. He used a monastery garden for crossing pea plant varieties having different heights, colors, pod shapes, seed shapes, and flower positions. Mendel's experiments, between 1856 and 1863, revealed how traits are passed down from parents. For example, when he crossed yellow peas with green peas, all the offspring peas were yellow. But when these offspring reproduced, the next generation was ³/₄ yellow and ¹/₄ green. Mendel's work, which was presented in 1865, showed that what we now call "genes" determine traits in predictable ways.



James Watson and Francis Crick discover the double helix structure of DNA

When Francis Crick and James Watson modeled the structure of DNA, they used paper cutouts of the bases (A, C, G, T) and metal scraps from a machine shop. Their model represented DNA as a double helix, with sugars and phosphates forming the outer strands of the helix and the bases pointing into the center. Hydrogen bonds connect the bases, pairing A–T and C–G; and the two strands of the helix are parallel but oriented in opposite directions. Their 1953 paper notes that the model "immediately suggests a possible copying mechanism for the genetic material."

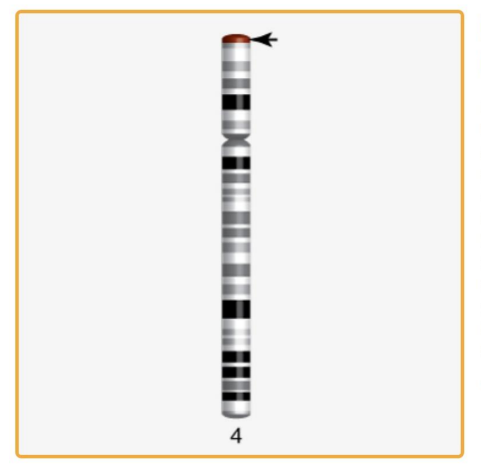


Marshall Nirenberg cracks the⁸ genetic code for protein synthesis

In the early 1960s, Marshall Nirenberg and National Institutes of Health colleagues focused on how DNA directs protein synthesis and the role of RNA in these processes. Their 1961 experiment, using a synthetic messenger RNA (mRNA) strand that contained only uracils (U), yielded a protein that contained only phenylalanines. Identifying UUU (three uracil bases in a row) as the RNA code for phenylalanine was their first breakthrough. Within a few years, Nirenberg's team had cracked the 60 mRNA codons for all 20 amino acids. In 1968, Nirenberg shared the Nobel Prize in Physiology or Medicine for his contributions to breaking the genetic code and understanding protein synthesis.

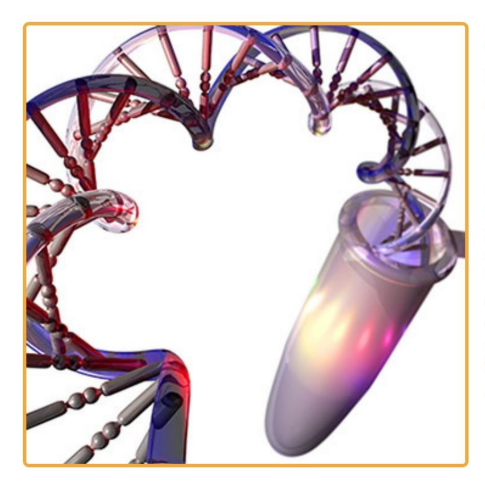
Frederick Sanger develops rapid DNA sequencing technique

In 1977, Frederick Sanger developed the classical "rapid DNA sequencing" technique, now known as the Sanger method, to determine the order of bases in a strand of DNA. Special enzymes are used to synthesize short pieces of DNA, which end when a selected "terminating" base is added to the stretch of DNA being synthesized. Typically, each of these terminating bases is tagged with a radioactive marker, so it can be identified. Then the DNA fragments, of varying lengths, are separated by how rapidly they move through a gel matrix when an electric field is applied – a technique called electrophoresis. Frederick Sanger shared the 1980 Nobel Prize in Chemistry for his contributions to DNA-sequencing methods.



First genetic disease mapped, ⁸ Huntington's Disease

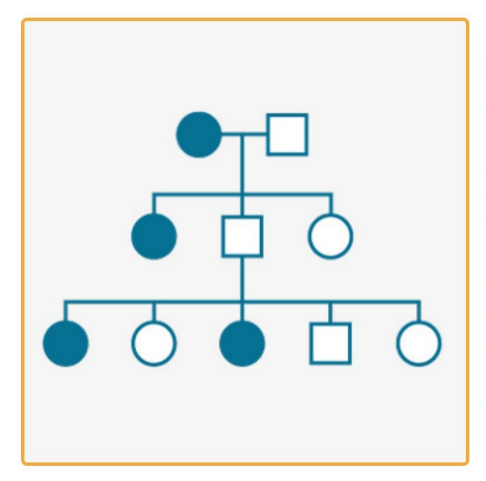
Huntington's disease (HD) causes the death of specific neurons in the brain, leading to jerky movements, physical rigidity, and dementia. Symptoms usually appear in midlife and worsen progressively. The location of the HD gene, whose mutation causes Huntington's disease, was mapped to chromosome 4 in 1983, making HD the first disease gene to be mapped using DNA polymorphisms – variants in the DNA sequence. The mutation consists of increasing repetitions of "CAG" in the DNA that codes for the protein huntingtin. The number of CAG repeats may increase when passed from parent to child, leading to earlier HD onset in each generation. The gene was finally isolated in 1993.



Invention of polymerase chain reaction (PCR) technology for amplifying DNA

Conceived in 1983 by Kary Mullis, the Polymerase Chain Reaction (PCR) is a relatively simple and inexpensive technology used to amplify or make billions of copies of a segment of DNA. One of the most important scientific advances in molecular biology, PCR amplification is used every day to diagnose diseases, identify bacteria and viruses, and match criminals to crime scenes. PCR revolutionized the study of DNA to such an extent that Dr. Mullis was awarded the Nobel Prize in Chemistry in 1993.





First evidence provided for the existence of the BRCA1 gene

BRCA1 (BReast CAncer gene 1) is a "tumor suppressor gene," which normally produces a protein that prevents cells from growing and dividing out of control. However, certain variations of BRCA1 can disrupt its normal function, leading to increased hereditary risk for cancer. The first evidence for existence of the BRCA1 gene was provided in 1990 by the King laboratory at University of California Berkeley. After a heated international race, the gene was finally isolated in 1994. Today, researchers have identified more than 1,000 mutations of the BRCA1 gene, many of them associated with increased risk of cancer, particularly breast and ovarian cancers in women.

1990



The Human Genome Project begins

Beginning in 1984, the U.S. Department of Energy (DOE), National Institutes of Health (NIH), and international groups held meetings about studying the human genome. In 1988, the National Research Council recommended starting a program to map the human genome. Finally, in 1990, NIH and DOE published a plan for the first five years of an expected 15-year project. The project would develop technology for analyzing DNA; map and sequence human and other genomes – including fruit flies and mice; and study related ethical, legal, and social issues.



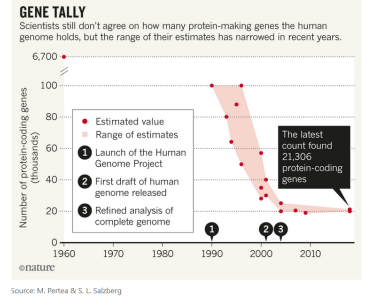
THE HUMAN GENOME

The Sequence of the Human Genome

J. Craig Venter, ¹* Mark D. Adams, ¹ Eugene W. Myers, ¹ Peter W. L(¹ Richard J. Mural, ¹ Granger G. Sutton, ¹ Hamilton O. Smith, ¹ Mark Yandell, ¹ Cheryi A. Brans, ¹ Robert A. Holt, ¹ Jeannine D. Goczyne, ¹ Peter Amanatides, ¹ Richard M. Ballew, ¹ Daniel H. Huson, ¹ Jennifer Russo Wortman,¹ Qing Zhang,¹ Chinnappa D, Kodira,¹ Xiangqun H, Zheng¹ Lin Chen,¹ Marian Skupski,¹ Gangadharan Subramanian,¹ Paul D. Thomas,¹ Jinghui Zhang,¹ George L, Gabor Miklos,² Catherine Nelson,³ Samuel Broder,¹ Andrew G, Clark,⁴ Joe Nadeau,⁵ Victor A, McKusick, ⁶ Norton Zinder,⁷ Arnold J, Levine,⁷ Richard J, Roberts,⁹ Mel Simon,⁹ Carolyn Slayman,¹⁰ Michael Hunkapiller,¹¹ Randall Bolanos,¹ Arthur Delcher,¹ Ian Dew,¹ Daniel Fasulo,¹ Michael Flanigan,¹ Liliana Florea,¹ Aaron Halpern,¹ Sridhar Hannenhalli,¹ Saul Kravitz,¹ Samuel Levy, Clark Mobarry,¹ Knut Reinert,¹ Karin Remington,¹ Jane Abu-Threideh,¹ Ellen Beasley,¹ Kendra Biddick, Vivien Bonazzi,¹ Rhonda Brandon,¹ Michele Cargill,¹ Ishwar Chandramouliswaran,¹ Rosane Charlab,¹ Kabir Chaturveli, ¹ Zuoming Deng, ¹ Valentina Di Francesco, ¹ Patrick Dunn, ¹ Karen Elbeck, Carlos Brangelista, ¹ Andrei E. Gabriellan, ¹ Weiniu Gan, ¹ Wangmao Ge, ¹ Fangcheng Gong, ¹ Zhiping Gu, ¹ Ping Guan, ¹ Thomas J. Heiman, ¹ Maureen E. Higgins, ¹ Rui-Ru Ji, ¹ Zhaoxi Ke, ¹ Karen A. Ketchum, ¹ Zhongwu Lai,¹ Yiding Lei,¹ Zhenya Li,¹ Jizyin Li,¹ Yong Liang,¹ Xiaoying Lin,¹ Fu Lu,¹ Gennady V. Merkulov,¹ Natalia Milshina,¹ Helen M. Moore,¹ Ashwinikumar K. Naik,¹ Valbhav A, Narayan,¹ Beena Neelam,¹ Deborah Nusskern,¹ Douglas B, Rusch,¹ Steven Salzberg,¹² Wei Shao,¹ Biolong Shue,¹ Jingtao Sun,¹ Zhen Yuan Wang,¹ Alhui Wang,¹ Xin Wang,¹ Jian Wang,¹ Ming-Hui Wei,¹ Ron Wides,¹² Chuntin Xiao,¹ Chunhua Yan,¹ Alison Yao,¹ Jane Ye,¹ Ming Zhan,¹ Weiqing Zhang,¹ Hongyu Zhang,¹ Qi Zhao,¹ Liansheng Zheng,¹ Fei Zhong,¹ Wenyan Zhong,¹ Shiaoping C, Zhu,¹ Shaying Zhao,¹² Dennis Gilbert,¹ Suzanna Baumhueter,¹ Gene Spier,¹ Christine Carter,¹ Anibal Cravchik,¹ Trevor Woodage,¹ Feroze Ali,¹ Huijin An,¹ Aderonke Awe,¹ Danita Baldwin,¹ Holly Baden,¹ Mary Barnstead,¹ Ian Barrow,¹ Karen Beeson,¹ Dana Busam, Amy Carver,¹ Angela Center,¹ Ming Lai Chang,¹ Liz Curry,¹ Stove Danaher,¹ Lionel Davenport,¹ Raymond Desilets,¹ Susanne Dietz,¹ Kristina Dodson,¹ Lisa Doup,¹ Stoven Ferriera,¹ Neha Garg, Andres Gluecksmann,¹ Brit Hart,¹ Jason Haynes,¹ Charles Haynes,¹ Cheryl Heiner,¹ Suzanne Hladun,¹ Damon Hostin,¹ Jarrett Houck,¹ Timothy Howland,¹ Chinyere Ibegwam,¹ Jeffery Johnson,¹ Francis Kalush, 'Leslay Kline,' Shahi Koduru,' Amy Lova, Felecia Hann,' David May,' Steven McCavdey,' Tina Mdrtosh,' Ivy McMullan,' Mee Moy,' Linda Moy,' Brian Murphy,' Keith Nelson,' Cynthia Prannkoch,' Bric Pratts, Vinita Puri,' Hina Quresh,'' Matthew Reardon,' Robert Rodriguez,' Yu-Hui Rogers,' Deama Rombiad,' Bob Ruhfel,' Richard Scott,' Cynthia Sitter,' Michelle Smallwood,¹ Erin Stawart,¹ Renee Strong,¹ Ellen Suh,¹ Reginald Thomas,¹ Ni Ni Tint,¹ Sukyae Tso,¹ Claire Vech,¹ Gary Wang,¹ Joremy Wetter,¹ Sherita Williams,¹ Monica Williams, Sandra Windsor,¹ Emily Winn-Deen,¹ Kerkellen Wolfe,¹ Jayshree Zaveri,¹ Karena Zaveri,¹ Josep F, Abril, ¹⁴ Roderic Guigo, ¹⁴ Michael J. Campbell, ¹ Kimmen V. Sjolander, ¹ Brian Karlak, ¹ Anish Kejariwal,¹ Huaiyu Mi,¹ Betty Lazareva,¹ Thomas Hatton,¹ Apurva Narechania,¹ Karen Diemer,¹ Anushya Muruganujan,¹ Nan Guo,¹ Shinji Sato,¹ Vineet Bafna,¹ Sorin Istrail,¹ Ross Lippert,¹ Russell Schwartz,¹ Brian Walenz,¹ Shibu Yooseph,¹ David Allen,¹ Anand Basu,¹ James Baxendale,¹ Louis Blick,¹ Marcelo Caminha,¹ John Carnes-Stine,¹ Parris Caulk,¹ Yen-Hui Chiang,¹ My Coyne,¹ Carl Dahike,¹ Anne Deslattes Mays,¹ Maria Dombroski,¹ Michael Donnelly,¹ Dale Ely,⁷ Shiva Esparham,¹ Carl Fosler,¹ Harold Gire,¹ Stephen Glanowski,¹ Kenneth Glasser,¹ Anna Glodek,¹ Mark Gorokhov,¹ Ken Graham,¹ Barry Gropman,¹ Michael Harris,¹ Jeremy Heil,¹ Scott Henderson,¹ Jeffrey Hoover, Donald Jennings,¹ Catherine Jordan,¹ James Jordan,¹ John Kasha,¹ Leonid Kagan,¹ Cheryl Kraft,¹ Donad Jumming, Center im Jordan, James Jordan, John Lopez, Daniel Ma, William Majoros, Joe McDaniel, Sean Murphy, Matthew Newman, Trung Nguyan, ¹Ngoc Nguyan, ¹Marc Nodell,¹ Sue Pan,¹ Jim Peck, ¹Marshall Peterson, ¹William Rowe, ¹Robert Sanders, ¹John Scott,¹ Michael Simpson,¹ Thomas Smith,¹ Arlan Sprague,¹ Timothy Stockwell,¹ Russell Turner,¹ Eli Venter,¹ Mei Wang,¹ Meiyuan Wen,¹ David Wu,¹ Mitchell Wu,¹ Ashley Xia,¹ Ali Zandieh,¹ Xiaohong Zhu¹

16 FEBILIARY 2001 VOL 201 SCIENCE www.sciencemeg.org

- In June 2000 came the announcement that the majority of the human genome had in fact been sequenced, which was followed by the publication of **90 percent of the sequence of the genome's three billion base-pairs** in the journal *Nature*, in February 2001
- Surprises accompanying the sequence publication included:
 - the relatively small number of human genes, perhaps as few as 30,000-35,000;



- the complex architecture of human proteins
 compared to their homologs similar genes with
 the same functions in, for example, roundworms
 and fruit flies;
- the lessons to be taught by repeat sequences of DNA.

2018: Also, als non-coding genes, estimated to be 17,871
2021: About 1% of DNA is made up of protein-coding genes; the other **99 percent** is noncoding.

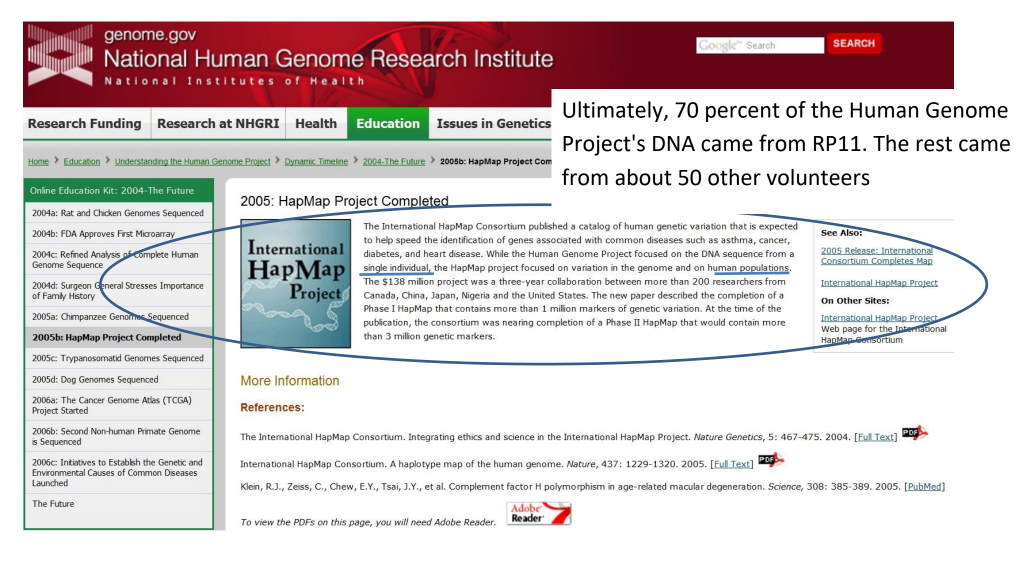


Home > Education > All About The Human Genome Project (HGP) > An Overview of the Human Genome Project (HGP)

News About the Human Genome Project

Links to news releases announcing key moments in the hist

0 <u>1994</u>	2004
O <u>1996</u>	2004
0 1998	March 24, 2004: International Sequencing Consortium Launches Online Resource
0 1999	The National Human Genome Research Institute announces that the International Sequencing Consortium (ISC) has launched a free, online resource
0 2000	where scientists and the public can get the latest information on the status of sequencing projects for animal, plant and other eukaryotic genomes.
0 2001	March 31, 2004: Scientists Compare Rat Genome With Human, Mouse
0 2001	An international research team, supported by the National Institutes of Health (NIH), today announced it has completed a high-quality, draft sequence
O 2003	of the genome of the laboratory rat, and has used that data to explore how the rat's genetic blueprint stacks up against those of mice and humans.
0 2004	
• 2004	April 21, 2004: NHGRI Scientists Return to the Classroom For Second Annual National DNA Day
	On April 30, dozens of researchers and staff from the National Human Genome Research Institute (NHGRI) will head back to high schools in rural and
	urban communities across the country to share with students some of the exciting research taking place at the National Institutes of Health (NIH).
	October 14, 2004: NHGRI Seeks Next Generation of Sequencing Technologies
	The National Human Genome Research Institute (NHGRI), part of the National Institutes of Health (NIH), today announced it has awarded more than
	\$38 million in grants to spur the development of innovative technologies designed to dramatically reduce the cost of DNA sequencing, a move aimed at
	broadening the applications of genomic information in medical research and health care.
	October 20, 2004: International Human Genome Sequencing Consortium Describes Finished Human Genome Sequence
	The International Human Genome Sequencing Consortium, led in the United States by the National Human Genome Research Institute (NHCRI) and the
	Department of Energy (DOE), today published its scientific description of the finished human genome sequence, reducing the estimated number of
	human protein-coding genes from 35,000 to only 20,000-25,000, a surprisingly low number for our species.

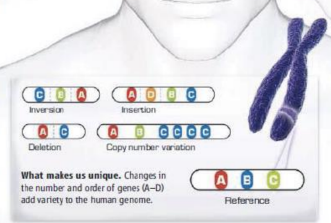


Human Genetic Variation

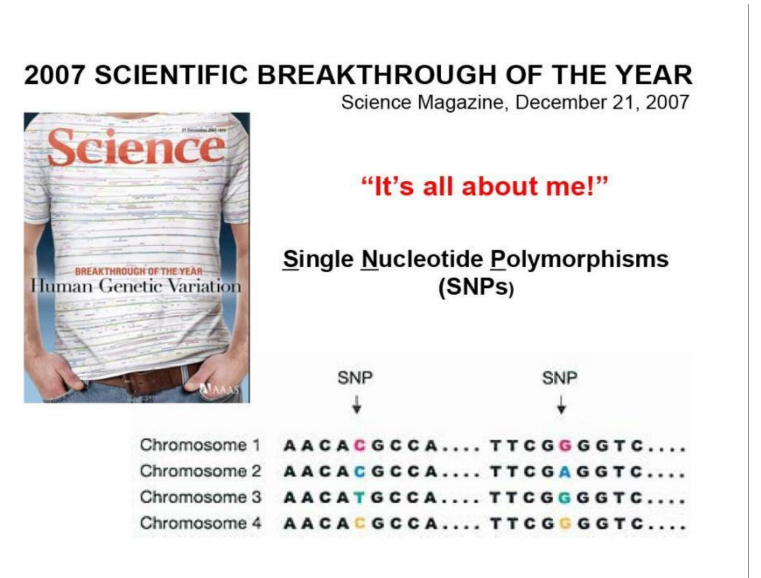
Equipped with faster, cheaper technologies for sequencing DNA and assessing variation in genomes on scales ranging from one to millions of bases, researchers are finding out how truly different we are from one another

THE UNVEILING OF THE HUMAN GENOME ALMOST 7 YEARS AGO cast the first faint light on our complete genetic makeup. Since then, each new genome sequenced and each new individual studied has illuminated our genomic landscape in ever more detail. In 2007, researchers came to appreciate the extent to which our genomes differ from person to person and the implications of this variation for deciphering the genetics of complex diseases and personal traits.

Less than a year ago, the big news was triangulating variation between us and our primate cousins to get a better handle on genetic changes along the evolutionary tree that led to humans. Now, we have moved from asking what in our DNA makes us human to striving to know what in my DNA makes me me.



Pennisi 2007 Science 318:1842-3



Historical overview: associating genetic variation to disease outcomes



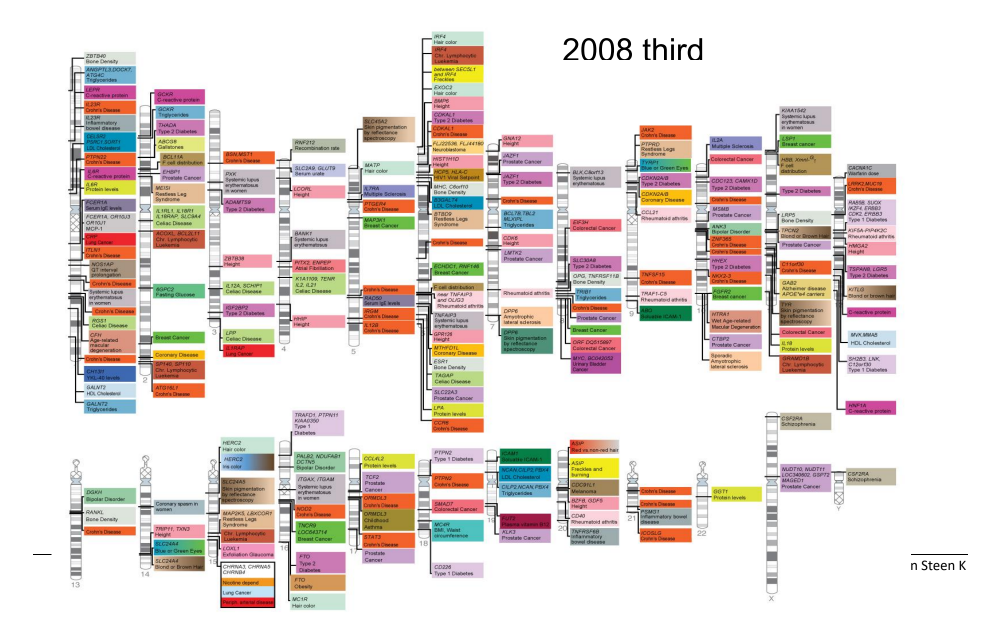
BREAKTHROUGH OF THE YEAR: The Runners-Up

Science **314**, 1850a (2006); DOI: 10.1126/science.314.5807.1850a

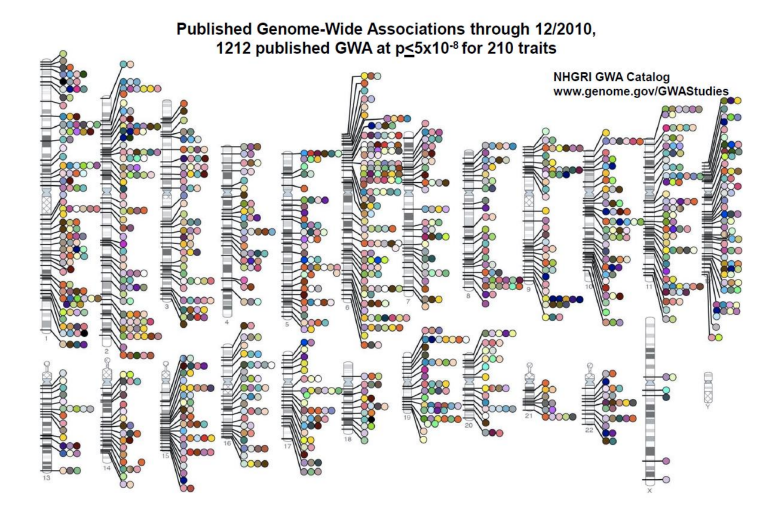
Areas to Watch in 2007

Whole-genome association studies. The trickle of studies comparing the genomes of healthy people to those of the sick is fast becoming a flood. Already, scientists have applied this strategy to macular degeneration, memory, and inflammatory bowel disease, and new projects on schizo-phrenia, psoriasis, diabetes, and more are heating up. But will the wave of data and new gene possibilities offer real insight into how diseases germinate? And will the genetic associations hold up better than those found the old-fashioned way?

Historical overview: GWAs as a tool to "map" diseases



Historical overview: 210 traits – multiple loci (sites, locations)

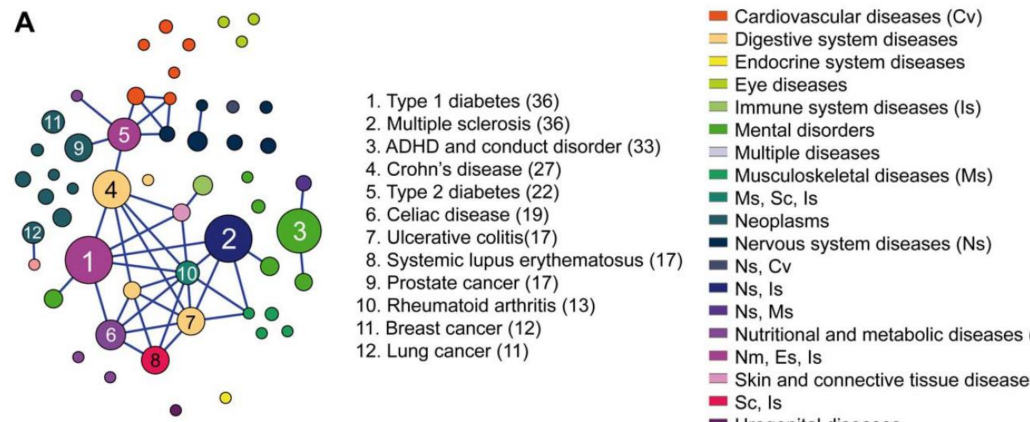


Historical overview: trait categories



Digestive system disease \odot Cardiovascular disease Metabolic disease \bigcirc Immune system disease $^{\circ}$ Nervous system disease 0 Liver enzyme measurement Lipid or lipoprotein measurement $^{\circ}$ Inflammatory marker measurement \bigcirc Hematological measurement 0 Body measurement \bigcirc Cardiovascular measurment Other measurement Response to drug \bigcirc **Biological process** \bigcirc Cancer \bigcirc Other disease Other trait \bigcirc

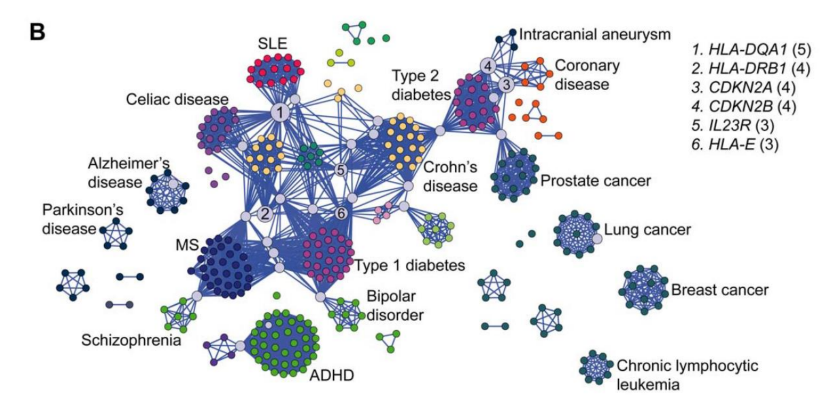
Historical overview: inter-relationships (networks)



Urogenital diseases

(Barrenas et al 2009: complex disease network – nodes are diseases)

Historical overview: inter-relationships (networks)



(Barrenas et al 2009: complex disease GENE network – nodes are genes)

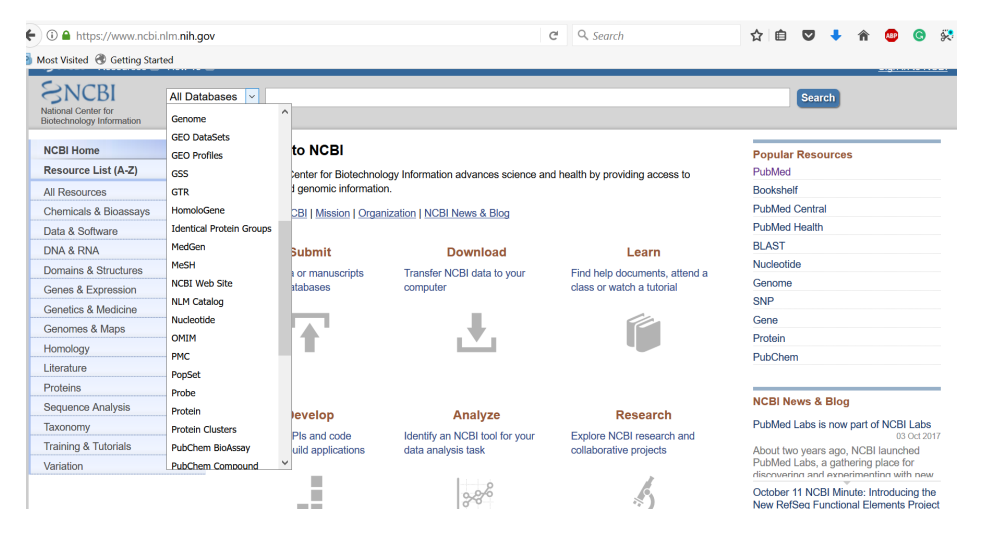
Historical overview: monitoring the progress

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OMIM	OMIM is a comprehensive, authoritative compendium of hum updated daily. OMIM is authored and edited at the McKusick University School of Medicine, under the direction of Dr. Ada	
Using OMIM	OMIM tools	Related Resources
Getting Started	OMIM API	ClinVar
FAQ		Gene
		GTR
		MedGen
Last updated on: 05 Oct 2014		

OMIM: molecular dissection of human disease

- Online Mendelian Inheritance in Man (OMIM[®]) is a continuously updated catalog of human genes and genetic disorders and traits (i.e. coded phenotypes, where phenotype is any characteristic of the organism), with particular focus on the molecular relationship between genetic variation and phenotypic expression.
- It can be considered to be a phenotypic companion to the Human Genome Project. OMIM is a continuation of Dr. Victor A. McKusick's Mendelian Inheritance in Man, which was published through 12 editions, the last in 1998.
- OMIM is currently biocurated at the McKusick-Nathans Institute of Genetic Medicine, The Johns Hopkins University School of Medicine.
- Frequently asked questions: <u>http://www.omim.org/help/faq</u>

Accessing OMIM



Historical overview: exome sequencing, full genome sequencing

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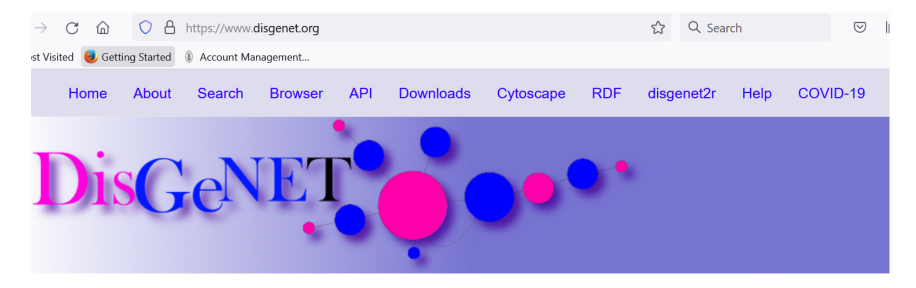
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DisGeNET: a database of gene-disease assocations



DisGeNET is a discovery platform containing one of the largest publicly available collections of genes and variants associated to human diseases (Piñero *et al.*, 2021; Piñero *et al.*, 2019; Piñero *et al.*, 2016; Piñero *et al.*, 2015). DisGeNET integrates data from expert curated repositories, GWAS catalogues, animal models and the scientific literature. DisGeNET data are homogeneously annotated with controlled vocabularies and community-driven ontologies. Additionally, several original metrics are provided to assist the prioritization of genotype–phenotype relationships.

The current version of DisGeNET (v7.0) contains 1,134,942 gene-disease associations (GDAs), between 21,671 genes and 30,170 diseases, disorders, traits, and clinical or abnormal human phenotypes, and 369,554 variant-disease associations (VDAs), between 194,515 variants and 14,155 diseases, traits, and phenotypes.

2 The rise of GWAs



(slide Doug Brutlag 2010)

What are GWAs?

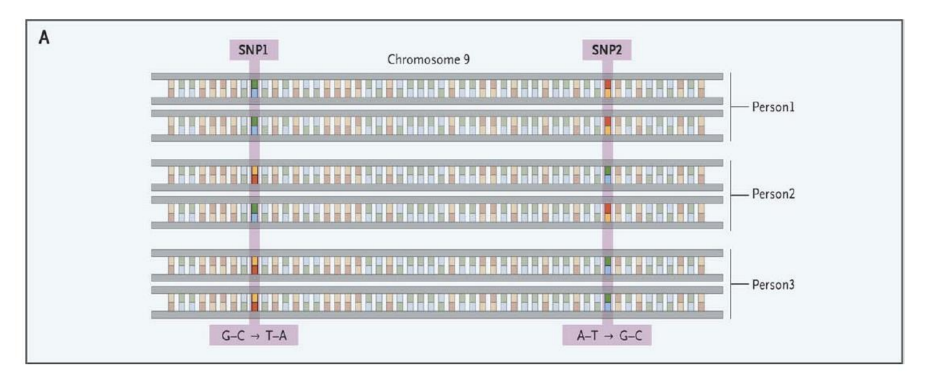
- A **genome-wide association study** is an approach that involves rapidly scanning markers across the complete sets of DNA, or genomes, of many people to find genetic variations associated with a particular trait.
- Recall:

A "trait" is the manifestation of the product of a gene that is coded for by the DNA; a "phenotype" is an individual's observable traits, such as height, eye color, and blood type.

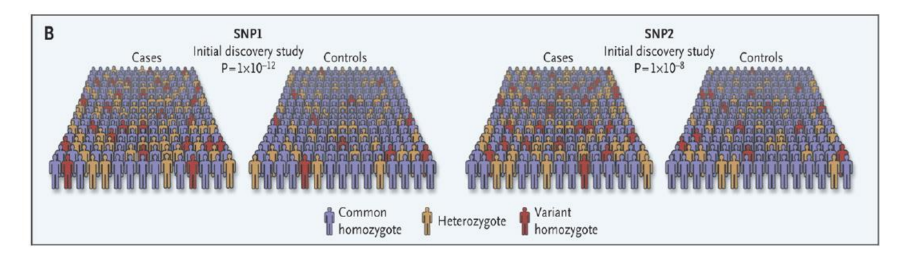
"Phenotype" is sometimes used <u>interchangeably</u> with the word trait, and "phenotype" may also define a whole compendium of traits together.

Genome-wide association studies: basic principles

The genome-wide association study is typically (but not solely!!!) based on a case-control design in which single-nucleotide polymorphisms (SNPs) across the human genome are genotyped ... (Panel A: small fragment)

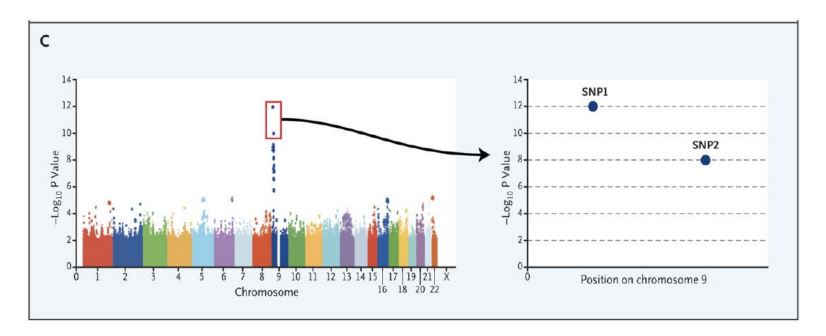


Genome-wide association studies: basic principles



 Panel B, the strength of association between each SNP and disease is calculated on the basis of the prevalence of each SNP in cases and controls. In this example, SNPs 1 and 2 on chromosome 9 are associated with disease, with P values of 10⁻¹² and 10⁻⁸, respectively

(Manolio 2010)



Genome-wide association studies: basic principles

- The plot in Panel C shows the P values for all genotyped SNPs that have survived a quality-control screen (each chromosome, a different color).
- The results implicate a locus on chromosome 9, marked by SNPs 1 and 2, which are adjacent to each other (graph at right), and other neighboring SNPs. (Manolio 2010)

How can we use genome-wide association studies results?

 Once new genetic associations are identified, researchers can use the information to develop better strategies to detect, treat and prevent the disease.

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View the GWAs catalogue (http://www.genome.gov/gwastudies/)

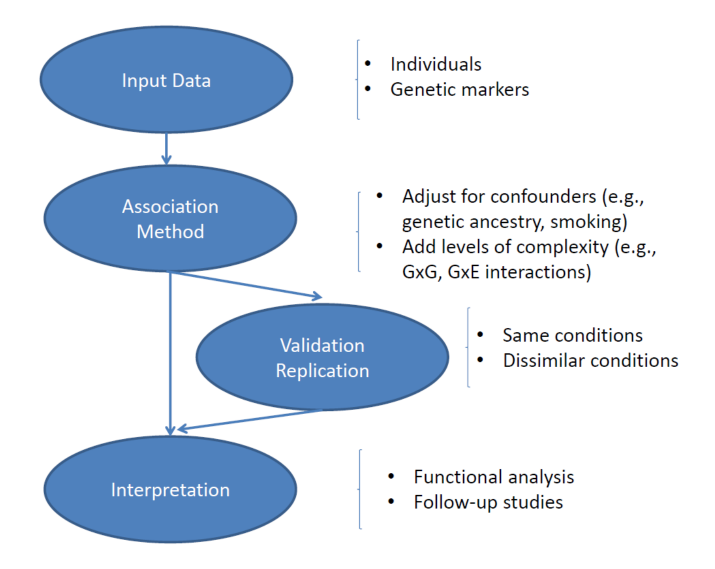
→ 2317 studies (6/10/2014) – info also accessible via DisGeNET

(Entries 1-50 of 2317)

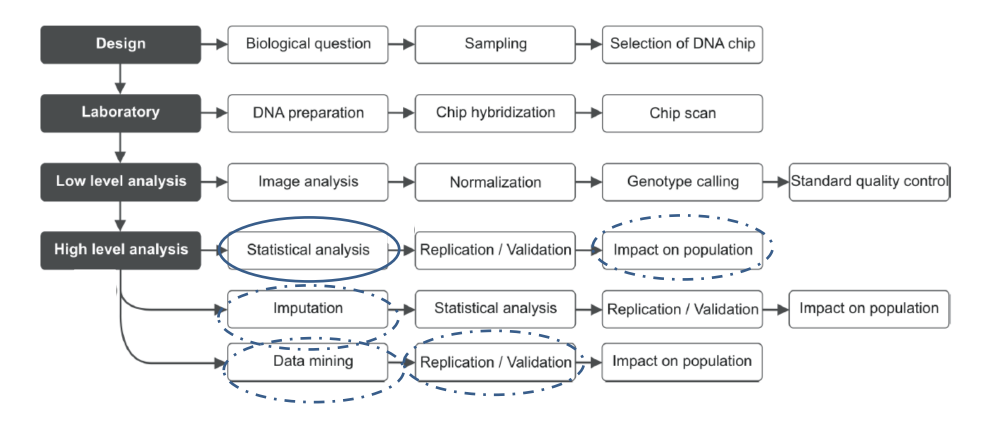
Date First Disease/Trait Initial Replication Region Reported Gene(s) Mapped Strongest Context **Risk Allele** P-value OR or Platform CNV beta-coefficient Added to Author/Date/ Gene(s) SNP-Risk Sample Sample Frequency [SNPs passing OC] Catalog Journal/Study Descriptio Description Allele and [95% CI] in (since Controls 11/25/08 04/16/14 RETN Chung CM Resistin levels 382 Han 559 Han 19p13.2 RETN rs1423096-G 0.78 1×10^{-7} .322 [0.25-0.40] Illumina March 03, 2014 C19orf59 [NR] Chinese Chinese ug/mL increase Diabetes Metab ancestry ancestry Res Rev indiviudals indiviudals Common quantitative trait locus downstream of RETN gene identified by genome-wide association study is associated with risk of type 2 diabetes mellitus in Han Chinese: a Mendelian randomization effect. 10/03/14 Zhang B Colorectal 1,773 East 6,902 East 18q21.1 SMAD7 SMAD7 rs7229639-A intron 0.145 3×10^{-11} 1.22 [1.15-1.29] Affymetrix & January 21, 2014 Illumina cancer Asian ancestr Asian ancestry Int J Cancer cases, 2,642 cases, 7,862 [1,695,815] Genome-wide East Asian East Asian (imputed) association study ancestry incestry identifies a new controls controls SMAD7 risk variant associated with colorectal cancer risk in East Asians. 10/06/14 NΔ Illumina Xie T Amyotrophic 250 Han View full set of 175 SNPs anuary 17, 2014 lateral sclerosis Chinese [859,311] (pooled) Neurobiol Aging (sporadic) ancestry NA RAB9P1 NA kgp22272527-? NR 8 x 10⁻¹¹ NR cases, 250 Ha A genome-wide 2×10^{-10} NA MYO18B NA kgp8087771-? 0.2 3.0327 association study Chinese [2.212039-4.157817] combining ancestry pathway analysis controls 12q24.33 GPR133 GPR133 0.08 8 x 10⁻¹⁰ 3.7761 [2.49-5.74] rs11061269-? intror for typical poradio 21q22.3 TMPRSS2 0.05 x 10⁻⁹ NR TMPRSS2 rs9977018-7

Page 1 of 47 <u>Next > Last >></u>





Detailed flow of a genome-wide association study



⁽Ziegler 2009)

Rise of bioinformatics determines rise of GWAs (1)

BIOINFORMATICS APPLICATIONS NOTE Vol. 23 no. 10 2007, pages 1294–1296 doi:10.1093/bioinformatics/btm108

Genetics and population analysis

GenABEL: an R library for genome-wide association analysis

Yurii S. Aulchenko^{1,*}, Stephan Ripke², Aaron Isaacs¹ and Cornelia M. van Duijn¹ ¹Department of Epidemiology and Biostatistics, Erasmus MC Rotterdam, Postbus 2040, 3000 CA Rotterdam, The Netherlands and ²Statistical Genetics Group, Max-Planck-Institute of Psychiatry, Kraepelinstr. 10, D-80804 Munich, Germany

Received on December 3, 2006; revised on February 14, 2007; accepted on March 13, 2007

Advance Access publication March 23, 2007

Associate Editor: Martin Bishop

ABSTRACT

Here we describe an R library for genome-wide association (GWA) analysis. It implements effective storage and handling of GWA data, fast procedures for genetic data quality control, testing of association of single nucleotide polymorphisms with binary or quantitative traits, visualization of results and also provides easy interfaces to standard statistical and graphical procedures implemented in base R and special R libraries for genetic analysis. We evaluated GenABEL using one simulated and two real data sets. We conclude that GenABEL enables the analysis of GWA data on desktop computers.

Availability: http://cran.r-project.org Contact: i.aoultchenko@erasmusmc.nl With these objectives in mind, we developed the GenABEL software, implemented as an R library. R is a free, open source language and environment for statistical analysis (http://www.r-project.org/). Building upon existing statistical analysis facilities allowed for rapid development of the package.

2 IMPLEMENTATION

2.1 Objective (1)

GWA data storage using standard R data types is ineffective. A SNP genotype for a single person may take four values (AA, AB, BB and missing). Two bits, therefore, are required to store these data. However, the standard R data types occupy 32 bits, leading to an overhead of 1500%, compared to the theoretical optimum. Use of the raw R data format, occupying

Rise of bioinformatics determines rise of GWAs (2)

BIOINFORMATICS

Vol. 26 ISMB 2010, pages i208–i216 doi:10.1093/bioinformatics/btg191

Multi-population GWA mapping via multi-task regularized regression

Kriti Puniyani, Seyoung Kim and Eric P. Xing* School of Computer Science, Carnegie Mellon University, Pittsburgh, PA, USA

ABSTRACT

Motivation: Population heterogeneity through admixing of different founder populations can produce spurious associations in genomewide association studies that are linked to the population structure rather than the phenotype. Since samples from the same population generally co-evolve, different populations may or may not share the same genetic underpinnings for the seemingly common phenotype. Our goal is to develop a unified framework for detecting causal genetic markers through a joint association analysis of multiple populations.

Results: Based on a multi-task regression principle, we present a multi-population group lasso algorithm using L1/L2-regularized regression for joint association analysis of multiple populations that are stratified either via population survey or computational estimation. Our algorithm combines information from genetic markers across populations, to identify causal markers. It also implicitly accounts for correlations between the genetic markers, thus enabling better control over false positive rates. Joint analysis across populations enables the detection of weak associations common to all populations with greater power than in a separate analysis of each population. At the same time, the regression-based framework allows causal alleles that are unique to a subset of the populations to be correctly identified. We demonstrate the effectiveness of our method on HapMap-simulated and lactase persistence datasets, where we significantly outperform state of the art methods, with greater power for detecting weak associations and reduced spurious associations. Availability: Software will be available at http://www.sailing.cs.cmu .edu/

the geographical distribution of the individuals. For example, it has been shown that such heterogeneity is present in the HapMap data (The International HapMap Consortium, 2005) across European, Asian and African populations; and heterogeneity at a finer scale within European ancestry has been found in many genomic regions in the UK samples of Wellcome trust case control consortium (WTCCC) dataset (Wellcome Trust Case Control Consortium, 2007). Although the standard assumption in existing approaches for association mapping is that the effects of causal mutations are likely to be common across multiple populations, the individuals in the same population or geographical region tend to co-evolve, and are likely to possess a population-specific causal allele for the same phenotype. For example, Tishkoff et al. (2006) reported that the lactase-persistence phenotype is caused by different mutations in Africans and Europeans. In addition, the same genetic variation has been observed to be correlated with gene-expression levels with different association strengths across different HapMap populations. Our goal is to be able to leverage information across multiple populations, to find causal markers in a multi-population association study.

1.1 Highlights of this article

We propose a novel multi-task-regression-based technique that performs a joint GWA mapping on individuals from multiple populations, rather than separate analysis of each population, to detect associated genome variations. The joint inference is achieved by using a multi-population group lasso (MPGL), with an L_1/L_2

Rise of bioinformatics determines rise of GWAs (3)

BIOINFORMATICS APPLICATIONS NOTE

Vol. 24 no. 1 2008, pages 140–142 doi:10.1093/bioinformatics/btm549

Genetics and population analysis

GWAsimulator: a rapid whole-genome simulation program

Chun Li^{1,*} and Mingyao Li²

¹Department of Biostatistics, Vanderbilt University School of Medicine, Nashville, TN 37232 and ²Department of Biostatistics and Epidemiology, University of Pennsylvania School of Medicine, Philadelphia, PA 19104, USA

Received on July 20, 2007; revised on October 10, 2007; accepted on October 29, 2007

Advance Access publication November 15, 2007

Associate Editor: Martin Bishop

ABSTRACT

Summary: GWAsimulator implements a rapid moving-window algorithm to simulate genotype data for case-control or population samples from genomic SNP chips. For case-control data, the program generates cases and controls according to a user-specified multi-locus disease model, and can simulate specific regions if desired. The program uses phased genotype data as input and has the flexibility of simulating genotypes for different populations and different genomic SNP chips. When the HapMap phased data are used, the simulated data have similar local LD patterns as the HapMap data. As genome-wide association (GWA) studies become increasingly popular and new GWA data analysis methods are being developed, we anticipate that GWAsimulator will be an important tool for evaluating performance of new GWA analysis methods. Availability: The C++ source code, executables for Linux, Windows and MacOS, manual, example data sets and analysis program are

and MacOS, manual, example data sets and analysis program an available at http://biostat.mc.vanderbilt.edu/GWAsimulator Contact: chun.li@vanderbilt.edu

Supplementary information: Supplementary data are available at *Bioinformatics* online.

2 METHODS

The program can generate unrelated case-control (sampled retrospectively conditional on affection status) or population (sampled randomly) data of genome-wide SNP genotypes with patterns of LD similar to the input data.

2.1 Phased input data and control file

The program requires phased data as input. If the HapMap data are used, the number of phased autosomes and X chromosomes are 120 and 90 for both CEU and YRI, 90 and 68 for CHB, and 90 and 67 for JPT. Additional parameters needed by the program should be provided in a control file, including disease model (see Section 2.2), window size (see Section 2.3), whether to output the simulated data (see Section 2.4), and the number of subjects to be simulated.

2.2 Determination of disease model

For simulations of case-control data, a disease model is needed. The program allows the user to specify disease model parameters, including disease prevalence, the number of disease loci, and for each disease locus, its location, risk allele and genotypic relative risk. If the mercure to simulate generations, the dust and mercitions need

Rise of bioinformatics determines rise of GWAs (4)

BIOINFORMATICS APPLICATIONS NOTE

Vol. 25 no. 5 2009, pages 662–663 doi:10.1093/bioinformatics/btp017

Genome analysis

AssociationViewer: a scalable and integrated software tool for visualization of large-scale variation data in genomic context

Olivier Martin^{1,†}, Armand Valsesia^{1,2,†}, Amalio Telenti³, Ioannis Xenarios¹ and Brian J. Stevenson^{1,2,*}

¹Swiss Institute of Bioinformatics, ²Ludwig Institute for Cancer Research, 1015 Lausanne and ³Institute of Microbiology, University Hospital, University of Lausanne, 1011 Lausanne, Switzerland

Received on September 16, 2008; revised on December 16, 2008; accepted on January 5, 2009

Advance Access publication January 25, 2009

Associate Editor: John Quackenbush

ABSTRACT

Summary: We present a tool designed for visualization of large-scale genetic and genomic data exemplified by results from genome-wide association studies. This software provides an integrated framework to facilitate the interpretation of SNP association studies in genomic context. Gene annotations can be retrieved from Ensembl, linkage disequilibrium data downloaded from HapMap and custom data imported in BED or WIG format. AssociationViewer integrates functionalities that enable the aggregation or intersection of data tracks. It implements an efficient cache system and allows the display of several, very large-scale genomic datasets.

Availability: The Java code for AssociationViewer is distributed under the GNU General Public Licence and has been tested on Microsoft Windows XP, MacOSX and GNU/Linux operating systems. It is available from the SourceForge repository. This also includes Java webstart, documentation and example datafiles.

Contact: brian.stevenson@licr.org

Supplementary information: Supplementary data are available at http://sourceforge.net/projects/associationview/ online.

represented in BED or WIG format and implements aggregation (union) or intersection of data tracks.

2 PROGRAM OVERVIEW

2.1 Cache and memory management

With increasing data volumes, efficient resource management is essential. One approach is to store the data in a cache with fast indexing mechanisms to retrieve the data, and to keep in memory only the information that is visualized. We implemented such a system in AssociationViewer. For comparison, loading a single dataset with 500 K SNPs in WGAViewer needs about 224 MB of RAM, whereas loading 10 different datasets (a total of 10 M data points) and displaying all genes on chromosome 1 needs only 50 MB in AssociationViewer.

2.2 Data import and export

A typical GWA dataset consists of a list of SNPs with *P*-values derived from an association analysis. In AssociationViewer, such

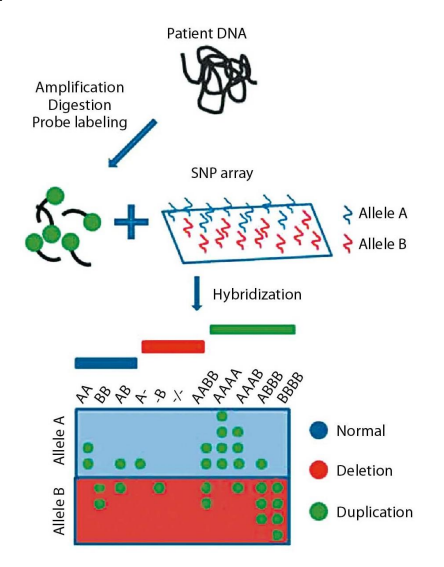
3 Study Design

Components of a study design for GWA studies

- The design of a genetic association study may refer to
 - study scale:
 - Genetic (e.g., hypothesis-drive, panel of candidate genes)
 - Genomic (e.g., hypothesis-free, genome-wide)
 - marker design:
 - Which markers are most informative in GWAs? Common variants-SNPs and/or Rare Variants (MAF<1%)
 - Which platform is the most promising? Least error-prone? Markerdistribution over the genome?
 - subject design

3.a Marker Level (see genetics class)

- Costs may play a role, but a balance is needed between costs and chip/sequencing platform performance
- Coverage also plays a role (e.g., exomes only or a uniform spread).
- When choosing Next Generation
 Sequencing platforms, also rare
 variants can be included in the
 analysis, in contrast to the older
 SNP-arrays (see right panel).



From common variants towards including rare variants

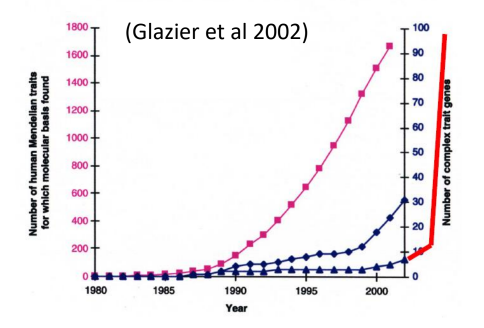
- Hypothesis 1 for GWAs: Common Disease Common Variant (CDCV):
 - This hypothesis argues that genetic variations with appreciable frequency in the population at large, but relatively low penetrance (i.e. the probability that a carrier of the relevant variants will express the disease), are the major contributors to genetic susceptibility to common diseases (Lander, 1996; Chakravarti, 1999; Weiss & Clark, 2002; Becker, 2004).
 - The hypothesis speculates that the gene variation underlying susceptibility to common heritable diseases existed within the **founding** population of contemporary humans → explains the success of GWAs?

From common variants towards including rare variants

- Hypothesis 2 for GWAs: Common Disease Rare Variant (CDRV):
 - This hypothesis argues that rare DNA sequence variations, each with relatively high penetrance, are the major contributors to genetic susceptibility to common diseases.
 - Some argumentations behind this hypothesis include that by reaching an appreciable frequency for common variations, these variations are not as likely to have been subjected to negative selection. Rare variations, on the other hand, may be rare because they are being selected against due to their deleterious nature.

There is room for both hypothesis in current research ! (Schork et al. 2009)

Identified # of traits for which a molecular basis exists: importance of SNPs



PINK : Human Mendelian traits

BLUE middle line : All complex traits

BLUE bottom line + red extension: Human complex traits

Complex disease (definition):

The term complex trait/disease refers to any phenotype that

does NOT exhibit classic Mendelian inheritance attributable to a single gene;

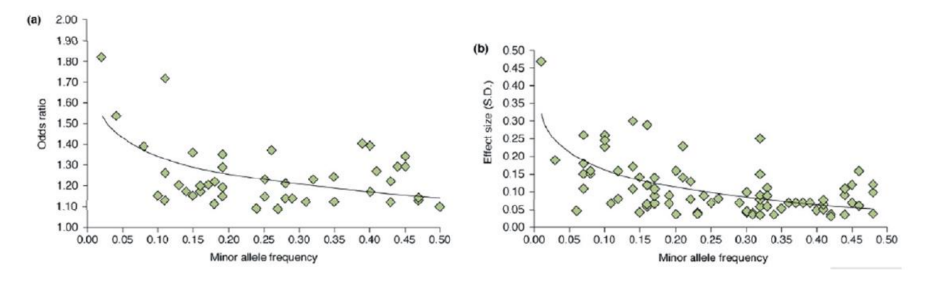
although they may exhibit familial tendencies (familial clustering, concordance among relatives).

Van Steen K

Distribution of SNP "effects"

Dichotomous Traits

Quantitative Traits



Arking & Chakravarti 2009 Trends Genet

Food for thought:

- The higher the MAF, the lower the effect size
- Rare variants analysis is in its infancy in 2009

3.b Subject Level

Aim	Selection scheme	
Increased effect size	Extreme sampling: Severely affected cases vs. extremely	
	normal controls	
Genes causing early	Affected, early onset vs. normal, elderly	
onset		
Genes with large /	Cases with positive family history vs. controls with	
moderate effect size	negative family history	
Specific GxE interaction	Affected vs. normal subjects with heavy environmental	
	exposure	
Longevity genes	Elderly survivors serve as cases vs. young serve as controls	
Control for covariates	Affected with favorable covariates vs. normal with	
with strong effect	unfavorable covariate	

Morton & Collins 1998 Proc Natl Acad Sci USA 95:11389

Popular design 1: cases and controls

Avoiding bias – checking assumptions:

- 1. Cases and controls drawn from same population
- 2. Cases representative for all cases in the population
- 3. All data collected similarly in cases and controls

Advantages:

- 1. Simple
- 2. Cheap
- 3. Large number of cases and controls available
- 4. Optimal for studying rare diseases

Disadvantages:

- 1. Population stratification
- 2. Prone to batch effects and other biases
- 3. Case definition / severity
- 4. Overestimation of risk for common diseases

Popular design 2: family-based

Avoiding bias – checking assumptions:

- 1. Families representative for population of interest
- 2. Same genetic background in both parents

Advantages:

- Controls immune to population stratification (no association without linkage, no "spurious" (false positive) association)
- 2. Checks for Mendelian inheritance possible (fewer genotyping errors)
- 3. Parental phenotyping not required (late onset diseases)

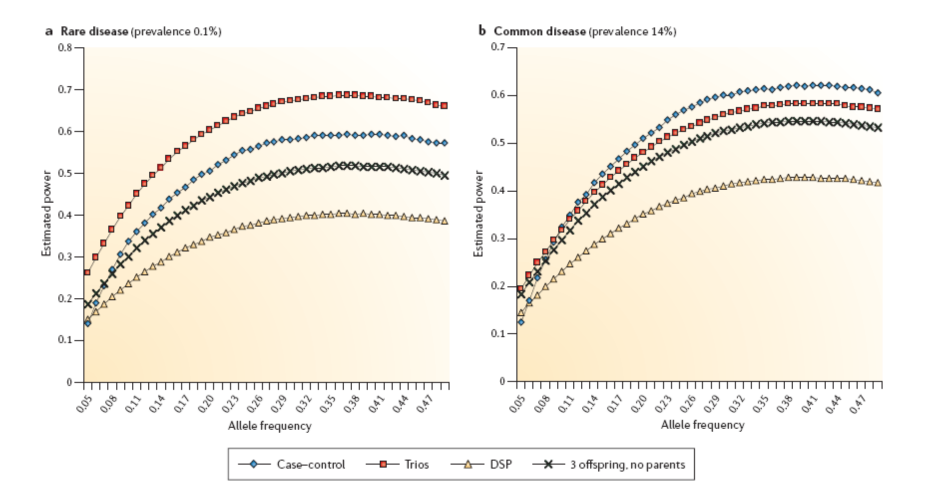
- 4. Simple logistics for diseases in children
- Allows investigating imprinting ("bad allele" from father or mother?)

Disadvantages

- 1. Cost inefficient
- 2. Sensitive to genotyping errors
- Lower power when compared with case-control studies

Some more power considerations

• Rare versus common diseases (Lange and Laird 2006)



4 Pre-analysis steps

4.a Quality control

Standard file format for GWA studies

Standard data format: tped = transposed ped format file

FamID	PID	FID	MID	SEX	AFF	SNP1 ₁	SNP1 ₂	SNP2 ₁	SNP2 ₂
1	1	0	0	1	1	A	А	G	Т
2	1	0	0	1	1	А	С	Т	G
3	1	0	0	1	1	С	С	G	G
4	1	0	0	1	2	А	С	Т	Т
5	1	0	0	1	2	С	С	G	Т
6	1	0	0	1	2	С	С	Т	Т



Chr	SNP name	Genetic distance	Chromosomal position
1	SNP1	0	123456
1	SNP2	0	123654

map file

Standard file format for GWA studies (continued)

Chr	SNP	Gen. dist.	Pos	PII) 1	PII) 2	PIE) 3	PII) 4	PII) 5	PII	D 6
1	SNP1	0	123456	Α	Α	Α	С	С	С	Α	С	С	С	С	С
1	SNP2	0	123654	G	Т	G	Т	G	G	Т	Т	G	Т	Т	Т

tfam file

tfam file: First 6 columns of standard ped file

tped file

FamID	PID	FID	MID	SEX	AFF
1	1	0	0	1	1
2	1	0	0	1	1
3	1	0	0	1	1
4	1	0	0	1	2
5	1	0	0	1	2
6	1	0	0	1	2

Small variations (headers) – software dependent

.fam (PLINK sample information file)

Sample information file accompanying a .bed binary genotype table. (--make-just-fam can be used to update just this file.) Also generated by "--recode Igen" and "--recode rlist".

A text file with no header line, and one line per sample with the following six fields:

- 1. Family ID ('FID')
- 2. Within-family ID ('IID'; cannot be '0')
- 3. Within-family ID of father ('0' if father isn't in dataset)
- 4. Within-family ID of mother ('0' if mother isn't in dataset)
- 5. Sex code ('1' = male, '2' = female, '0' = unknown)
- 6. Phenotype value ('1' = control, '2' = case, '-9'/'0'/non-numeric = missing data if case/control)

.bim (PLINK extended MAP file)

Extended variant information file accompanying a .bed binary genotype table. (--make-just-bim can be used to update just this file.)

A text file with no header line, and one line per variant with the following six fields:

- 1. Chromosome code (either an integer, or 'X'/'Y'/'XY'/'MT'; '0' indicates unknown) or name
- 2. Variant identifier
- 3. Position in morgans or centimorgans (safe to use dummy value of '0')
- 4. Base-pair coordinate (1-based; limited to 2^{31} -2)
- 5. Allele 1 (corresponding to clear bits in .bed; usually minor)
- 6. Allele 2 (corresponding to set bits in .bed; usually major)

Allele codes can contain more than one character. Variants with negative bp coordinates are ignored by PLINK.

.bed (PLINK binary biallelic genotype table)

Primary representation of genotype calls at biallelic variants. Must be accompanied by .bim and .fam files. Loaded with --bfile; generated in many situations, most notably when the --make-bed command is used. **Do not confuse this with the UCSC Genome Browser's BED format, which is totally different.**

The first three bytes should be 0x6c, 0x1b, and 0x01 in that order. (There are old versions of the .bed format which start with a different "magic number"; PLINK 1.9 recognizes them, but will convert sample-major files to the current variant-major format on sight. See the bottom of the original .bed definition page for details; that page also contains a more verbose version of the discussion below.)

The rest of the file is a sequence of **V** blocks of **N**/4 (rounded up) bytes each, where **V** is the number of variants and **N** is the number of samples. The first block corresponds to the first marker in the .bim file, etc.

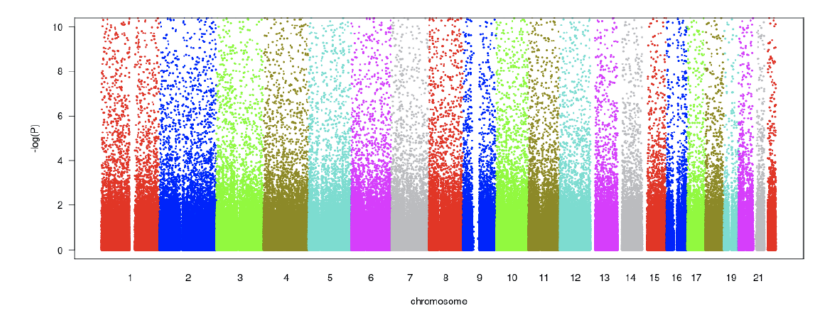
The low-order two bits of a block's first byte store the first sample's genotype code. ("First sample" here means the first sample listed in the accompanying .fam file.) The next two bits store the second sample's genotype code, and so on for the 3rd and 4th samples. The second byte stores genotype codes for the 5th-8th samples, the third byte stores codes for the 9th-12th, etc.

The two-bit genotype codes have the following meanings:

- 00 Homozygous for first allele in .bim file
- 01 Missing genotype
- 10 Heterozygous
- 11 Homozygous for second allele in .bim file

Why is quality control (QC) important?

BEFORE QC \rightarrow true signals are lost in false positive signals

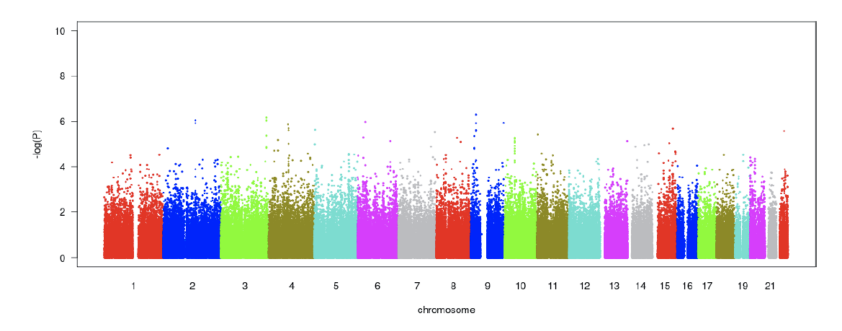


Ger MI FS I, Affymetrix 500k array set, SNPs on chip: 493,840

(Ziegler and Van Steen 2010)

Why is quality control important?

AFTER QC \rightarrow skyline of Manhattan (\rightarrow name of plot: Manhattan plot):



Ger MI FS I, Affymetrix 500k array set, SNPs on chip: 493,840

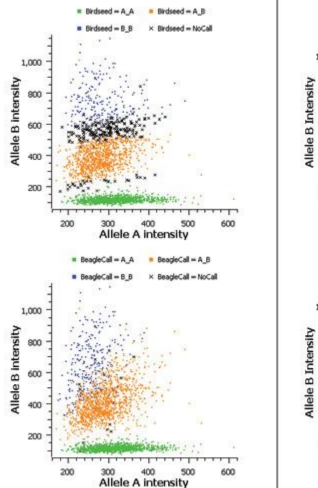
SNPs passing standard quality control: 270,701

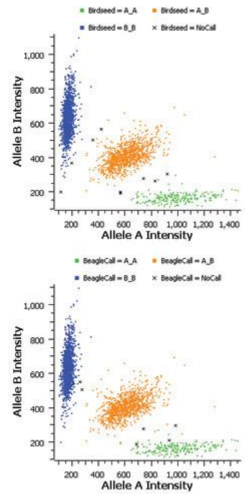
(Ziegler and Van Steen 2010)

What is the standard quality control?

- Quality control can be performed on different levels:
 - Subject or sample level
 - Marker level (in this course: SNP level)
 - X-chromosomal SNP level (in this course not considered)
- Consensus on how to best QC data has led to the so-called "Travemünde criteria" (obtained in the town Travemünde, but data generation platform dependent!) – see later

Marker level QC thresholds may be genotype calling algorithm dependent





Allele signal intensity genotype calling cluster plots for two different SNPs from the same study population.

<u>Upper panels</u>: Birdseed genotypes

Lower panels: BEAGLECALL genotypes.

The plots on the left show a SNP with poor resolution of A_B and B_B genotype clusters and the increased clarity of genotype calls that comes from using BEAGLECALL

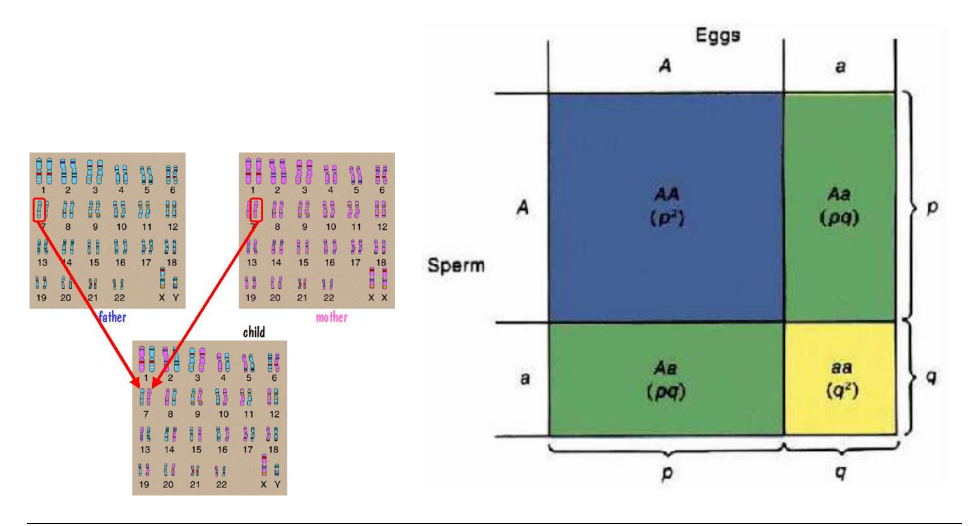
(Golden Helix Blog)

Quality control at the marker level

- Minor allele frequency (MAF):
 - Genotype calling algorithms perform poorly for SNPs with low MAF
 - Power is low for detecting associations to genetic markers with low MAF (with standard large-sample statistics)
- Missing frequency (MiF)
 - 1 minus call rate
 - MiF needs to be investigated separately in cases and controls because differential missingness may bias association results
- Hardy-Weinberg equilibrium (HWE)
 - SNPs excluded if substantially more or fewer subjects heterozygous at a SNP than expected (excess heterozygosity or heterozygote deficiency)

What is Hardy-Weinberg Equilibrium (HWE)?

Consider diallelic SNP with alleles A and a



What is Hardy-Weinberg Equilibrium (HWE)?

The genetic variation in a population will remain constant from one generation to the next in the absence of disturbing factors

Consider a locus with two alleles: *A* and *a* Assume in the first generation the alleles are not in HWE and the genotype frequency distribution is as follows:

1st Generation				
Genotype	Frequency			
AA	и			
Aa	V			
аа	W			

where u + v + w = 1

From the genotype frequencies, we can easily obtain allele frequencies:

$$P(A) = u + \frac{1}{2}v$$
$$P(a) = w + \frac{1}{2}v$$

In the first generation:
$$P(A) = u + \frac{1}{2}v$$
 and $P(a) = w + \frac{1}{2}v$

Mating Type	Mating Frequency	Expected Progeny
AA imes AA	u ²	AA
AA imes Aa	2uv	$\frac{1}{2}AA:\frac{1}{2}Aa$
AA imes aa	2 <i>uw</i>	Aa
Aa imes Aa	v^2	$\frac{1}{4}AA: \frac{1}{2}Aa: \frac{1}{4}aa$
Aa imes aa	2 <i>vw</i>	$\frac{1}{2}Aa:\frac{1}{2}aa$
aa imes aa	w ²	aa

2nd Generation

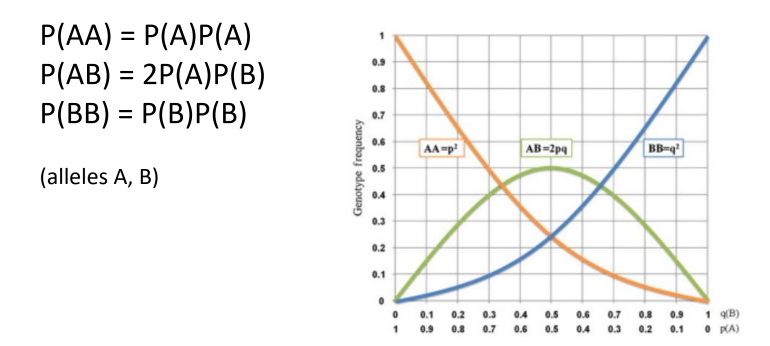
* Check: $u^2 + 2uv + 2uw + v^2 + 2vw + w^2 = (u + v + w)^2 = 1$ $p \equiv P(AA) = u^2 + \frac{1}{2}(2uv) + \frac{1}{4}v^2 = (u + \frac{1}{2}v)^2$ $q \equiv P(Aa) = uv + 2uw + \frac{1}{2}v^2 + vw = 2(u + \frac{1}{2}v)(\frac{1}{2}v + w)$ $r \equiv P(aa) = \frac{1}{4}v^2 + \frac{1}{2}(2vw) + w^2 = (w + \frac{1}{2}v)^2$ In the third generation:

$$P(AA) = \left(p + \frac{1}{2}q\right)^2 = \left(\left(u + \frac{1}{2}v\right)^2 + \left(\frac{1}{2}\right)2\left(u + \frac{1}{2}v\right)\left(\frac{1}{2}v + w\right)\right)^2$$
$$= \left(\left(u + \frac{1}{2}v\right)\left[\left(u + \frac{1}{2}v\right) + \left(\frac{1}{2}v + w\right)\right]\right)^2$$
$$= \left(\left(u + \frac{1}{2}v\right)\left[(u + v + w)\right]\right)^2$$
$$= \left(\left(u + \frac{1}{2}v\right)1\right)^2 = \left(\left(u + \frac{1}{2}v\right)\right)^2 = p$$
Similarly, $P(Aa) = q$ and $P(aa) = r$ for generation 3

• Equilibrium is reached after one generation of mating under the Hardy-Weinberg assumptions! Genotype frequencies remain the same from generation to generation.

HWE

- 1. When a population is in Hardy-Weinberg equilibrium, the alleles that comprise a genotype can be thought of as having been chosen at random from the alleles in a population.
- 2. We have the following relationship between genotype frequencies and allele frequencies for a population in Hardy-Weinberg equilibrium:



Distorting factors to HWE causing evolution to occur

3. Non-random mating

- 4. **Mutation** by definition mutations change allele frequencies causing evolution
- 5. **Migration** if new alleles are brought in by immigrants or old alleles are taken out by emigrants then the frequencies of alleles will change causing evolution
- 6. Genetic drift random events due to small population size (bottleneck caused by storm and leading to reduced variation, migration events leading to founder effects)

7.Natural selection – some genotypes give higher reproductive success (Darwin)

Testing for HWE – assumptions are violated?

Compares observed genotype counts with the values expected under Hardy-Weinberg. For a locus with two alleles, we might construct a table as follows:

Genotype	Observed	Expected under HWE
AA	пдд	np_A^2
Aa	n _{Aa}	$2np_A(1-p_A)$
аа	n _{aa}	$n(1-p_A)^2$

where *n* is the number of individuals in the sample and p_A is the probability that a random allele in the population is of type A.

We estimate
$$p_A$$
 with $\hat{p}_A = \frac{2n_{AA} + n_{Aa}}{2n}$

Testing for HWE – assumptions are violated?

Test statistic is for Allelic Association is:

$$X^{2} = \sum_{\text{genotypes}} \frac{(\text{Observed count} - \text{Expected count})^{2}}{\text{Expected count}}$$
$$X^{2} = \frac{\left(n_{AA} - n\hat{p}_{a}^{2}\right)^{2}}{n\hat{p}_{a}^{2}} + \frac{\left(n_{Aa} - 2n\hat{p}_{a}(1-\hat{p}_{a})\right)^{2}}{2n\hat{p}_{a}(1-\hat{p}_{a})} + \frac{\left(n_{aa} - n(1-\hat{p}_{a})^{2}\right)^{2}}{n(1-\hat{p}_{a})^{2}}$$

Under H_0 , the X^2 test statistic has an approximate χ^2 distribution with 1 degree of freedom Recall the rule of thumb for such χ^2 tests: the expected count should be at least 5 in every cell. If allele frequencies are low, and/or sample size is small, and/or there are many alleles at a locus, this may be a problem. Solution?

The Travemünde criteria

Level	Filter criterion	Standard value for filter			
Sample level	Call fraction	≥ 97%			
	Cryptic relatedness	Study specific			
	Ethnic origin	Study specific; visual inspection of			
		principal components			
	Heterozygosity	Mean ± 3 std.dev. over all samples			
	Heterozygosity by gender	Mean ± 3 std.dev. within gender group			

SNP level	MAF	≥ 1%
	MiF	≤ 2% in any study group, e.g., in both
		cases and controls
	MiF by gender	≤ 2% in any gender
	HWE	p < 10 ⁻⁴

(Ziegler 2009)

The Travemünde criteria

Level	Filter criterion	Standard value for filter
SNP level	Difference between control groups	p > 10 ⁻⁴ in trend test
	Gender differences among controls	p > 10 ⁻⁴ in trend test
X-Chr SNPs	Missingness by gender	No standards available
	Proportion of male heterozygote calls	No standards available
	Absolute difference in call fractions for	No standards available
	males and females	
	Gender-specific heterozygosity	No standard value available

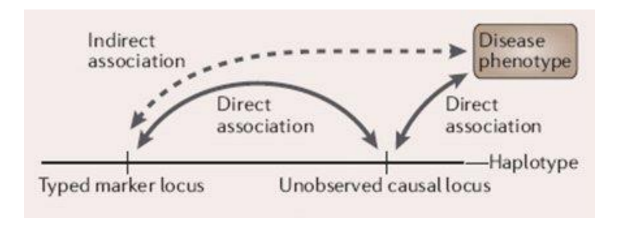
(Ziegler 2009)

4.b Linkage disequilibrium

• Linkage Disequilibrium (LD) is a measure of co-segregation of alleles in a population – linkage + allelic association

Two alleles at different loci that occur together on the same chromosome (or gamete) more often than would be predicted by random chance.

• It is a very important concept for GWAs, since it gives the rational for performing genetic association studies



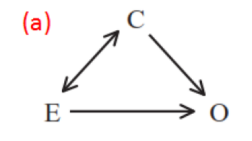
4.c Confounding by shared genetic ancestry – "population stratification"

If successful, the random allocation of subjects to the exposure which characterise RCTs ensures a balanced distribution of known and unknown confounding factors between exposed and nonexposed subjects. This is equivalent of removing the association between the exposure and all potential confounders (Figure 1b), and therefore, the possibility of confounding itself. In this case, the effect of the exposure on the outcome can be directly estimated by simply comparing outcomes between exposed and unexposed subjects (1).

(Cois 2014)

Regression uses mathematical modelling to estimate the effect of confounders on the outcome, and to "remove" this effect statistically. This is equivalent of removing (or, more realistically, reducing) the association between confounder and outcome, thus eliminating the second necessary condition for confounding (Figure 1c). Two necessary — albeit not sufficient — conditions for an extraneous factor ("confounder") to produce such a bias are (Figure 1a):

- the confounder is a risk factor for the outcome;
- the confounder is associated with the exposure, i.e. its distribution is different among individuals with different exposure status.



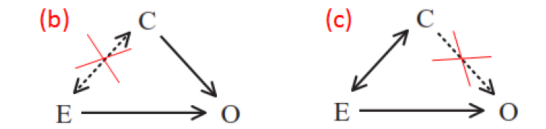
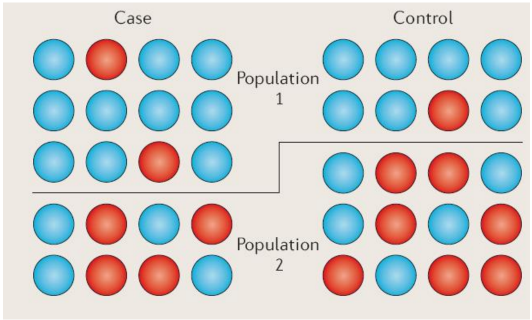


Figure 1: Schematic illustration of confounding control. Arrows represent causal effects, double arrows associations of any nature. E = exposure, C = confounder, O = outcome.

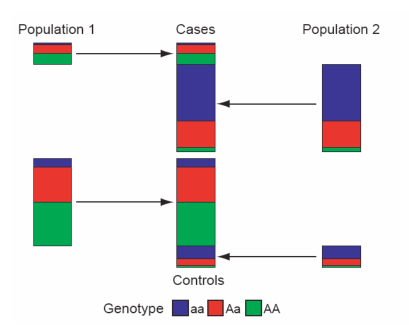
What is spurious association in GWAS?

• **Spurious association** refers to false positive association results due to not having accounted for population substructure as a confounding factor in the analysis



What is spurious association?

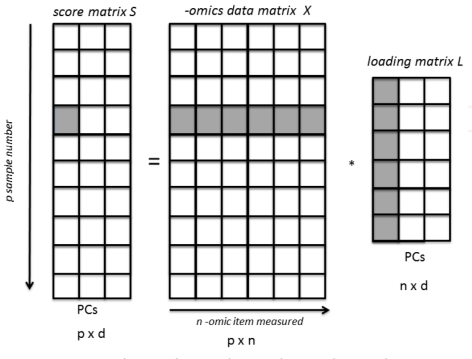
- Typically, there are two characteristics present:
 - A difference in proportion of individual from two (or more) subpopulation in case and controls
 - Subpopulations have different allele frequencies at the locus.



What are typical methods to deal with population stratification in GWAS?

- Methods to deal with spurious associations generated by population structure generally require a number (at least >100) of widely spaced null SNPs that have been genotyped in cases and controls in addition to the candidate SNPs.
- These methods large group into:
 - Principal components: finding continuous axes of genetic variation
 - Structured association methods: "First look for structure (population clusters) and second perform an association analysis conditional on the cluster allocation"
 - Genomic control methods: "First analyze and second downplay association test results for over optimism"

Principal components



 $\mathsf{s}_{4,1} = \mathsf{x}_{4,1}.\mathsf{I}_{1,1} + \mathsf{x}_{4,2}.\mathsf{I}_{2,1} + \mathsf{x}_{4,3}.\mathsf{I}_{3,1} + \mathsf{x}_{4,4}.\mathsf{I}_{4,1} + \mathsf{x}_{4,5}.\mathsf{I}_{5,1} + \mathsf{x}_{4,6}.\mathsf{I}_{6,1}$

• Mathematical derivation:

https://courses.cs.ut.ee/MTAT.03.227/2017 spring/uploads/Main/lecture-notes-9.pdf

• Applications in omics: http://cdn.intechopen.com/pdfs-wm/30002.pdf

Principal components

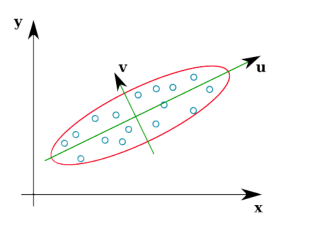


Figure 1: PCA for Data Representation

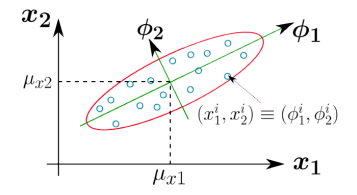


Figure 3: The PCA Transformation

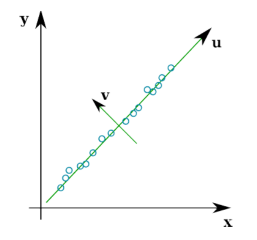


Figure 2: PCA for Dimension Reduction

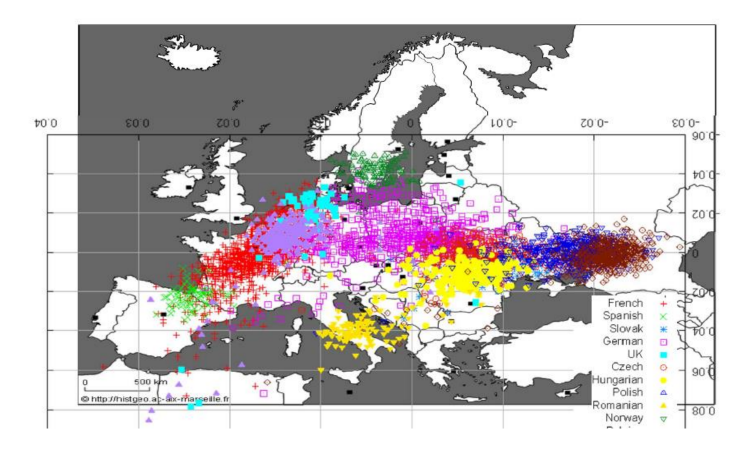
- Find eigenvectors of the covariance matrix for standardized (x1, x2, ...) [→SNPs]
- These will give you the direction vectors indicated in Fig3 by phi_1 and phi_2
- These determine the axes of maximal variation

Principal components in genetics studies

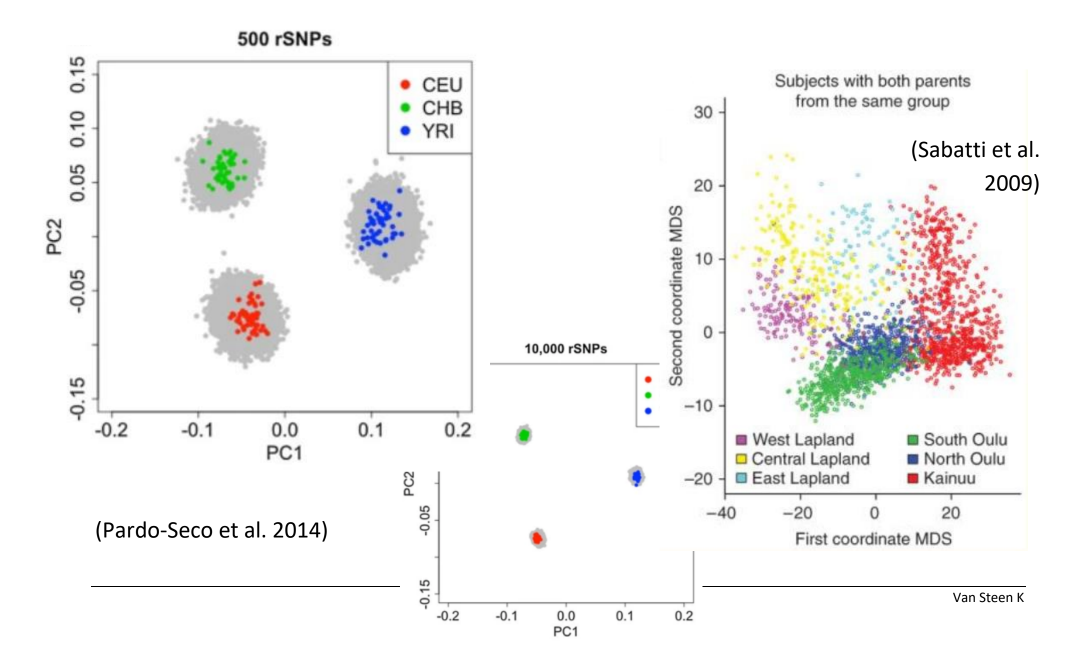
- Cavalli-Sforza et al. pioneered the use of PCA to summarise data on variation in human gene frequencies across continental regions (Menozzi et al. 1978).
- These results have been highly controversial but also highly influential; PCA has become heavily used in population genetics:
- The EIGENSOFT package combines functionality from population genetics methods (Patterson et al. 2006) and the EIGENSTRAT stratification correction method (Price et al. 2006)
- Novembre et als (2014) were among the first to study the behaviour of PCA with data exhibiting continuous spatial variation, such as might exist within human continental groups.
- Our group has also contributed: PCA in statistical genetics (Abegaz et al. 2019).

Principal components in statistical genetics

In European data, the first 2 principal components "nicely" reflect the N-S and E-W axes ! Y-axis: PC2 (6% of variance); X-axis: PC1 (26% of variance)



Principal components in statistical genetics: the more SNPs the better?



SmartPCA vs Smart PCA

Smart PCA

Yi Zhang

Machine Learning Department Carnegie Mellon University yizhang1@cs.cmu.edu

Abstract

PCA can be smarter and makes more sensible projections. In this paper, we propose smart PCA, an extension to standard PCA to regularize and incorporate external knowledge into model estimation. Based on the probabilistic interpretation of PCA, the inverse Wishart distribution can be used as the informative conjugate prior for the population coas a specific case of factor analysis with isotropic Gaussian noise, and the use of the inverse Wishart distribution as the natural conjugate prior for the covariance matrix in multivariate normal distribution [Gelman *et al.*, 2003], which has been recently investigated by researchers in statistics [Brown *et al.*, 2000; Press, 2005], machine learning [Klami and Kaski, 2007], image processing and computer vision [Smidl *et al.*, 2001; Wood *et al.*, 2006]. Based on previous work, a natural way to improve PCA is to incorporate external knowledge

https://www.cs.cmu.edu/~yizhang1/docs/SmartPCA.pdf

https://github.com/chrchang/eigensoft/blob/master/POPGEN/README

5 Analysis Steps

5.a Testing for Associations

The role of regression analysis

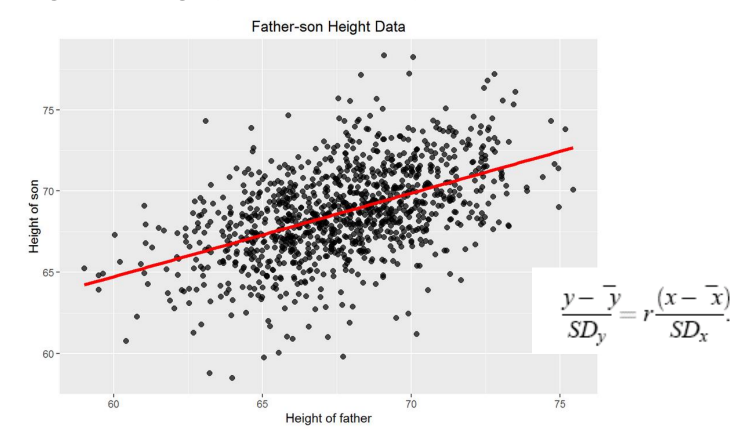
 Galton used the following equation to explain the phenomenon that sons of tall fathers tend to be tall but not as tall as their fathers while sons of short fathers tend to be short but not as short as their fathers:

$$\frac{y - \overline{y}}{SD_y} = r \frac{(x - \overline{x})}{SD_x}.$$

This effect is called the **regression effect**.

The use of regression analysis

• **regression line** goes through (mean Y, mean X)



(https://rstudio-pubs-static.s3.amazonaws.com/204984_dd2112475db84af2a03260c4a4f830ac.html)

The use of regression analysis

- **Regression analysis** is used for explaining or modeling the relationship between a single variable Y, called the response, output or dependent variable, and one or more predictor, input, independent or explanatory variables, X₁, ..., X_p.
- When p=1 it is called simple regression but when p > 1 it is called multiple regression or sometimes multivariate regression.
- When there is more than one Y, then it is called multivariate multiple regression
- Regression analyses have several possible objectives including
 - Prediction of future observations.
 - Assessment of the effect of, or relationship between, explanatory variables on the response.
 - A general description of data structure

The linear regression model

$$y = \beta_0 + \beta_1 x_1 + \ldots + \beta_k x_k + \epsilon$$

- *y*: response variable.
- x_1, \ldots, x_k : regressor variables, independent variables.
- $\beta_0, \beta_1, \ldots, \beta_k$: regression coefficients.
- ε: model error.
 - Uncorrelated: $cov(\epsilon_i, \epsilon_j) = 0, i \neq j$.
 - Mean zero, Same variance: $var(\epsilon_i) = \sigma^2$. (homoscedasticity)
 - Normally distributed.

Linear vs non-linear

Linear Models Examples:

$$y = \beta_0 + \beta_1 x + \beta_2 x^2 + \epsilon$$

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \epsilon$$

$$y = \beta_0 + \beta_1 \log x_1 + \beta_2 \log x_2 + \epsilon$$

$$\log y = \beta_0 + \beta_1 \left(\frac{1}{x_1}\right) + \beta_2 \left(\frac{1}{x_2}\right) + \epsilon$$

Nonlinear Models Examples:

$$y = \beta_0 + \beta_1 x_1^{\gamma_1} + \beta_2 x_2^{\gamma_2} + \epsilon$$
$$y = \frac{\beta_0}{1 + e^{\beta_1 x_1}} + \epsilon$$

Regression inference

$$y = \beta_0 + \beta_1 x_1 + \ldots + \beta_k x_k + \epsilon$$

- Least square estimation of the regression coefficients. $\underline{b} = (X^T X)^{-1} X^T y.$
- Variance estimation for σ^2 (see later)
- Coefficient of Determination. R^2 .
- Partial F test or t-test for H_0 : $\beta_j = 0$.

X : n x p matrix;			
n= nr of samples;			
p= nr of covariates			
β:px1			
Υ = Χ β			

What is R-squared?

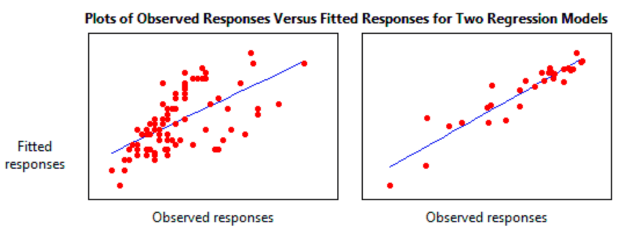
- R-squared is a statistical measure of how close the data are to the fitted regression line. It is also known as the coefficient of determination, or the coefficient of multiple determination for multiple regression.
- The definition of R-squared is fairly straight-forward; it is the percentage of the response variable variation that is explained by a linear model:

R-squared = Explained variation / Total variation

- R-squared is always between 0 and 100%:
 - 0% indicates that the model explains none of the variability of the response data around its mean.
 - 100% indicates that the model explains all the variability of the response data around its mean.

Graphical representation of R-squared

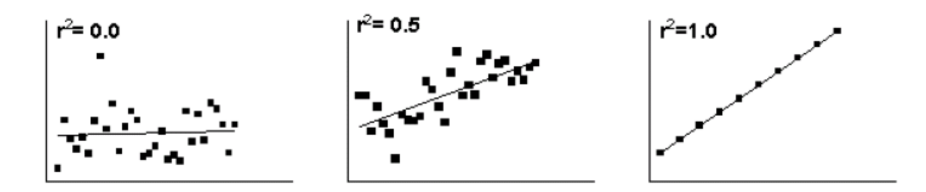
• Plotting fitted values by observed values graphically illustrates different R-squared values for regression models.



• The regression model on the left accounts for 38.0% of the variance while the one on the right accounts for 87.4%. The more variance that is accounted for by the regression model the closer the data points will fall to the fitted regression line.

Coefficient of determination ~ squared correlation coefficient r²

- An r² value of 0.0 means that knowing X does not help you predict Y.
 There is no linear relationship between X and Y, and the best-fit line is a horizontal line going through the mean of all Y values.
- When r² equals 1.0, all points lie exactly on a straight line with no scatter.
 Knowing X lets you predict Y perfectly.



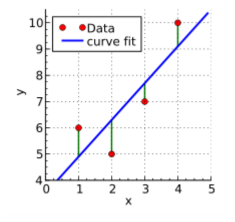
General linear test approach

• The full model (continuous response, say "BMI"):

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \varepsilon$$

- Fit the model by f.i. the method of least squares (this leads to estimations b for the beta parameters in the model)
- It will also lead to the **error sums of squares** (SSE): the sum of the squared deviations of each observation Y around its estimated expected value
- The error sums of squares of the full model SSE(F):

$$\sum [Y - b_0 - b_1 X_1 - b_2 X_2]^2$$
$$= \sum (Y - \hat{Y})^2$$



General linear test approach

• Next we consider a null hypothesis H₀ of interest:

```
H_0: \beta_1 = 0H_1: \beta_1 \neq 0
```

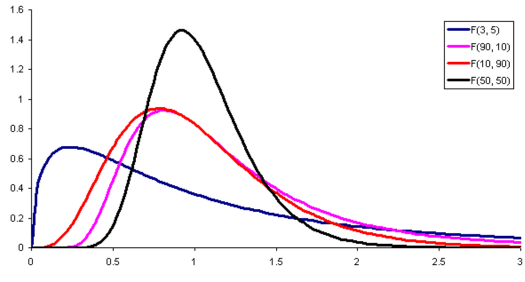
• The model when H₀ holds is called **the reduced or restricted model**. When $\beta_1 = 0$, then the regression model reduces to

$$Y = \beta_0 + \beta_2 X_2 + \varepsilon$$

- Again we can fit this model with f.i. the least squares method and obtain an error sums of squares, now for the reduced model: SSE(R)
- Question: which error sums of squares will be smaller? SSE(F) or SSE(R)

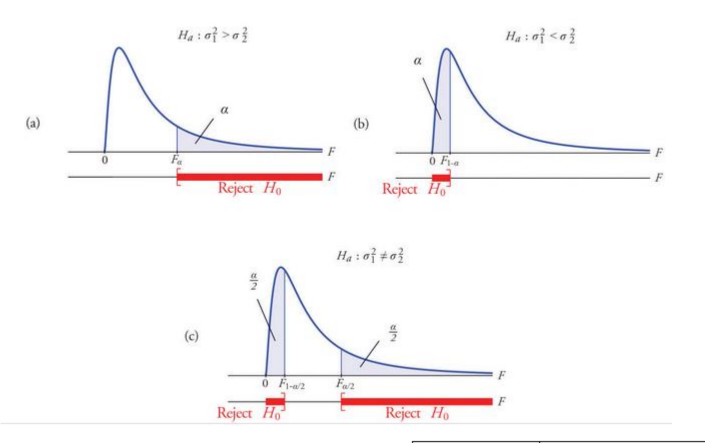
General linear test approach

- The F distribution (sometimes known as the Fisher-Snedecor distribution (Sir Ronald Aylmer Fisher (1890-1962), George Waddell Snedecor (1882 -1974)) and taking Fisher's initial) is commonly used in a variety of statistical tests.
- It is derived from the ratio of two normalized chi-squared distributions (chisquareds can be used to test whether a population variance equals a prespecified value):



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Why F-test?



Terminology	Alternative Hypothesis	Rejection Region
Right-tailed	$H_a: \sigma_1^2 > \sigma_2^2$	$oldsymbol{F} \geq F_{lpha}$
Left-tailed	$H_a: \sigma_1^2 < \sigma_2^2$	$oldsymbol{F} \leq F_{1-lpha}$
Two-tailed	$H_a: \sigma_1^2 eq \sigma_2^2$	$oldsymbol{F} \leq F_{1-lpha/2}$ or $oldsymbol{F} \geq F_{lpha/2}$

Why F-test?

• The logic now is to compare both SSEs. The actual test statistic is a function of SSE(R)-SSE(F):

$$F^* = \frac{SSE(R) - SSE(F)}{df_R - df_F} : \frac{SSE(F)}{df_F}$$

which follows an F distribution when H₀ holds

• The decision rule (for a given alpha level of significance) is: If $F^* \leq F(1 - \alpha; df_R - df_F, df_F)$, you cannot reject H₀ If $F^* > F(1 - \alpha; df_R - df_F, df_F)$, conclude H₁

Tests in GWAS using the regression framework

• Example 1:

$$Y = \beta_0 + \beta_1 SNP + \varepsilon$$

- $-H_0: \beta_1 = 0$ $-H_1: \beta_1 \neq 0$ $-df_F = n - 2 \text{ (this links to df in variance estimation)}$
- $df_R = n 1$ (this links to df in variance estimation)

It can be shown that for testing $\beta_1 = 0$ versus $\beta_1 \neq 0$

$$-F^{*} = \frac{SSE(R) - SSE(F)}{df_{R} - df_{F}} : \frac{SSE(F)}{df_{F}} =$$

$$\frac{b_{1}^{2}}{s^{2}(b_{1})} = (t^{*})^{2}$$

$$t_{(n-p)}^{2} = F_{(1,n-p)}$$
The general linear statistic:
$$F^{*} = \frac{SSE(R) - SSE(F)}{df_{R} - df_{F}} : \frac{SSE(F)}{df_{F}}$$
simplifies to:
$$F^{*} = \frac{SSR(x_{1}|x_{2}, x_{3})}{1} : \frac{SSE(x_{1}, x_{2}, x_{3})}{n-4} = \frac{MSR(x_{1}|x_{2}, x_{3})}{MSE(x_{1}, x_{2}, x_{3})}$$

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Tests in GWAS using the regression framework

• Example 2:

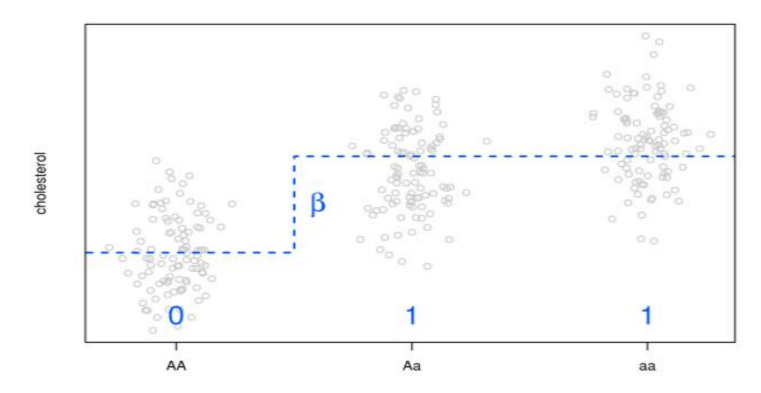
$$Y = \beta_0 + \beta_1 SNP + \beta_2 PC_1 + \beta_3 PC_2 + \varepsilon$$

- $H_0: \beta_1 = 0$
- $H_1: \beta_1 \neq 0$
- $df_F = n - 4$
- $df_R = n - 3$

How many dfs would the corresponding F-test have?

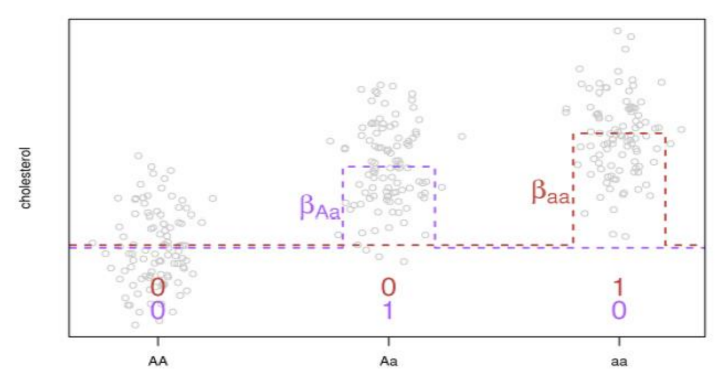
The impact of different encoding schemes for SNPs

	Coding scheme for statistical modeling/testing					
Indiv. genotype	X1	X1	X2	X1	X1	X1
	Additive coding	cod (genera	otype ling al mode ritance)		Recessive coding (for a)	Advantage Heterozygous
AA	0	0	0	0	0	0
Aa	1	1	0	1	0	1
аа	2	0	1	1	1	0

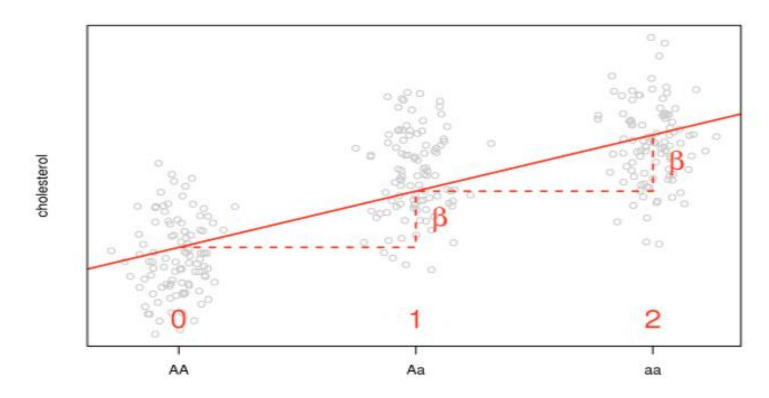


Which encoding scheme provides a good fit to the data?





Robust vs overkill ?



Which encoding scheme provides a good fit to the data?

Most commonly used

Regression analysis in R

- Main functions
 - The basic syntax for doing regression in R is **Im()** to fit linear models
 - The R function **glm()** can be used to fit generalized linear models (i.e., when the response is not normally distributed)
- General syntax rules in R model fitting are given on the next slide.

Regression analysis in R

Syntax	Model	Comments
$Y \sim A$	$Y = \beta_0 + \beta_1 A$	Straight-line with an implicit y-
		intercept
$Y\sim \textbf{-}1+A$	$Y = \beta_1 A$	Straight-line with no y-intercept;
		that is, a fit forced through $(0,0)$
$Y \sim A + I(A^{\wedge}2)$	$Y = \beta_0 + \beta_1 A + \beta_2 A^2$	Polynomial model; note that the
		identity function I() allows terms
		in the model to include normal
		mathematical symbols.
$\mathbf{Y}\sim\mathbf{A}+\mathbf{B}$	$Y = \beta_0 + \beta_1 A + \beta_2 B$	A first-order model in A and B
		without interaction terms.
$Y \sim A:B$	$Y = \beta_0 + \beta_1 AB$	A model containing only first-order
		interactions between A and B.
$Y \sim A^*B$	$Y = \beta_0 + \beta_1 A + \beta_2 B + \beta_3 A B$	A full first-order model with a term;
		an equivalent code is Y ~ A + B +
		A:B.
$Y \sim (A + B + C)^{\wedge}2$	$Y = \beta_0 + \beta_1 A + \beta_2 B + \beta_3 C +$	A model including all first-order
	$\beta_4 AB + \beta_5 AC + \beta_6 AC$	effects and interactions up to the n th
	F4	order, where n is given by () ^{n} .
		An equivalent code in this case is
		$Y \sim A^*B^*C - A:B:C.$

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Model diagnostics are model-dependent ...

- There are 4 principal assumptions which justify the use of **linear regression** models for purposes of prediction:
 - **linearity** of the relationship between dependent and independent variables
 - independence of the errors (no serial correlation)
 - homoscedasticity (constant variance) of the errors
 - versus time (when time matters)
 - versus the predictions (or versus any independent variable)
 - normality of the error distribution. (http://www.duke.edu/~rnau/testing.htm)
- To check **model assumptions**: go to **quick-R** and regression diagnostics (<u>http://www.statmethods.net/stats/rdiagnostics.html</u>)

QQ plots for model diagnostics – Q for Quantile

• Quantiles are points in your data below which a certain proportion of your data fall.

What is the 0.5 quantile for normally distributed data?

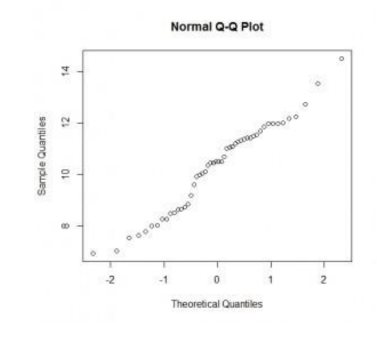
• Here we generate a random sample of size 200 from a normal distribution and find the quantiles for 0.01 to 0.99 using the quantile function:

quantile(rnorm(200),probs = seq(0.01,0.99,0.01))

 Q-Q plots take your sample data, sort it in ascending order, and then plot them versus quantiles calculated from a theoretical distribution. The number of quantiles is selected to match the size of your sample data. The quantile function in R offers 9 different quantile algorithms! See help(quantile)

QQ plots for model diagnostics – Q for Quantile

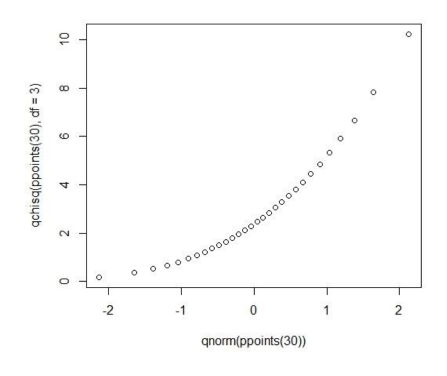
- A Q-Q plot is a scatterplot created by plotting **two sets of quantiles** against one another.
- If both sets of quantiles come from the same distribution, we should see the points forming a line that's roughly straight.
- Here's an example of a Normal Q-Q plot when both sets of quantiles truly come from Normal distributions.



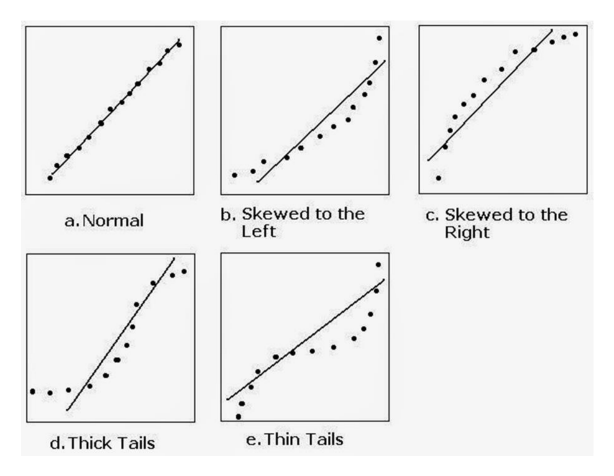
Examples of QQ plots: no straight line

 QQ plot of a distribution that's skewed right; a Chi-square distribution with 3 degrees of freedom against a Normal distribution

qqplot(qnorm(ppoints(30)), qchisq(ppoints(30),df=3))

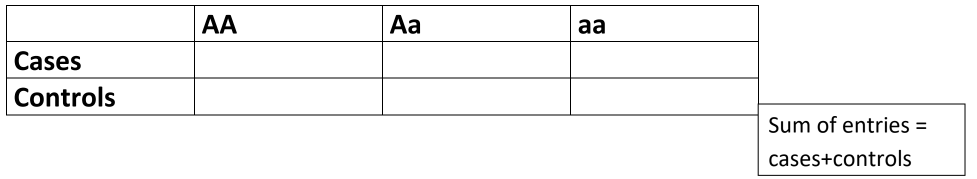


Examples of QQ plots: some frequent scenarios

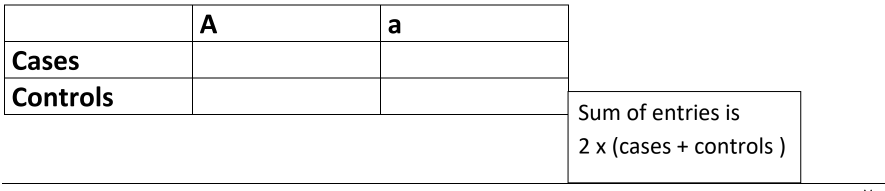


Testing for association between case/control status and a SNP

 Fill in the table below and perform a chi-squared test for independence between rows and columns → genotype test → 2 df



 Fill in the table below and perform a chi-squared test for independence between rows and columns → allelic test (ONLY valid under HWE) → 1df



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Testing for association between case/control status and a SNP

- The **genotype test involves a 2df test** (note that two variables X1 and X2 were needed for genotype coding).
- It has been shown that usually, the additive coding gives adequate power, even when the true underlying mode of inheritance is NOT additive (note that the **additive coding can be achieved by only using 1 variable** (X1)).
- For large sample sizes, a "test for trend" (risk for disease, or average trait increases/decreases with increasing number of "a" copies) theoretically follows a chi-squared distribution with **1df**.

Instead of $Y = \beta_0 + \beta_1 SNP + \varepsilon; Y \text{ continuous}$ and modelling $E[Y|SNP] = \beta_0 + \beta_1 SNP \text{ (without error term!)}$ consider

 $\beta_0 + \beta_1 SNP = \eta$ representing the linear combination as it can never be equal to a binary variable (0/1 response; control/case status)

and model $g(E[Y|SNP]) = \beta_0 + \beta_1 SNP = \eta$ where g() is called a link function

and thus $E[Y|SNP] = g_inv(\eta)$

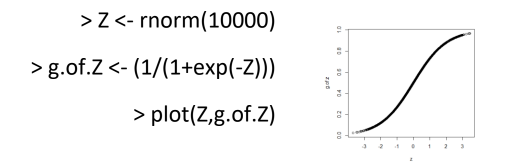
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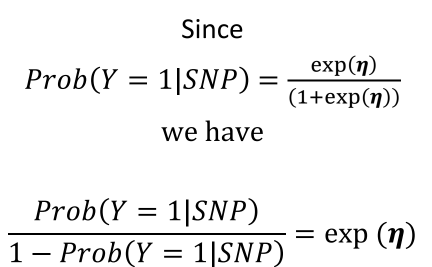
For a binary trait Y:

$$E[Y|SNP] = Prob(Y = 1|SNP)$$

= $\frac{\exp(\eta)}{(1 + \exp(\eta))} = \frac{1}{(1 + \exp(-\eta))} = g_inv(\eta)$
where
g_inv is the logistic function (sigmoid function)

(squashing the linear predictor to an acceptable range)



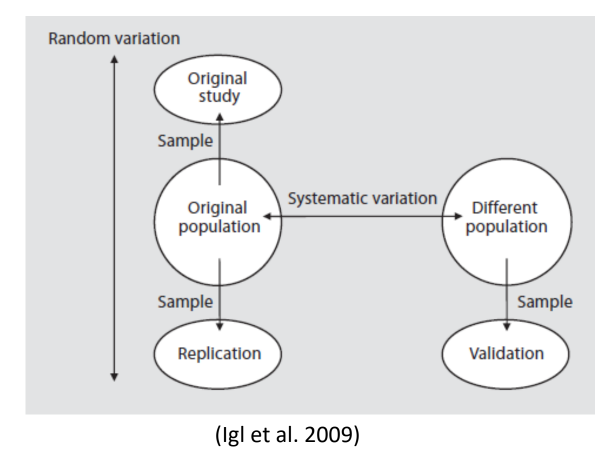


$$\boldsymbol{g}(\boldsymbol{E}[\boldsymbol{Y}|\boldsymbol{SNP}]) = \beta_0 + \beta_1 \boldsymbol{SNP} = \log\left(\frac{\boldsymbol{Prob}(\boldsymbol{Y}=1|\boldsymbol{SNP})}{1 - \boldsymbol{Prob}(\boldsymbol{Y}=1|\boldsymbol{SNP})}\right) = \boldsymbol{\eta}$$

(g is called the logit function)

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5.b Replication and validation



Guidelines for replication studies

- Replication studies should be of sufficient size to demonstrate the effect
- Replication studies should conducted in independent datasets
- Replication should involve the same phenotype
- Replication should be conducted in a similar population
- The same SNP should be tested
- The replicated signal should be in the same direction
- Joint analysis should lead to a lower *p*-value than the original report
- Well-designed negative studies are valuable

Note that SNPs are most likely to replicate when they

- show modest to strong statistical significance,
- have common minor allele frequency,
- exhibit modest to strong genetic effect size (~strength of association)

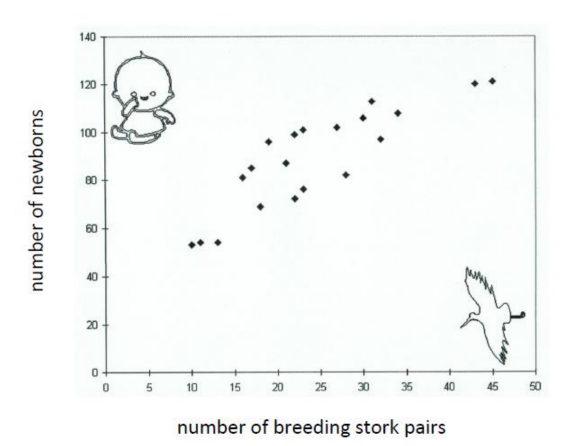
5.c Causation

"Association does not imply causation"

• Meaning:

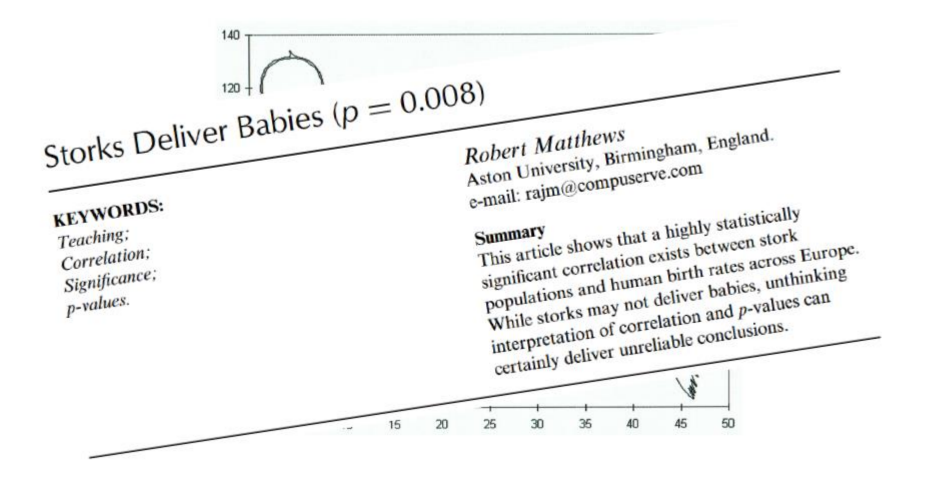
Just because two things correlate does not necessarily mean that one causes the other.

 As a seasonal example, just because people in Belgium tend to spend more in the shops when it's cold and less when it's hot doesn't mean cold weather causes high street spending.



"Association does not imply causation"

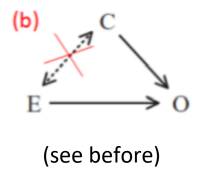
... so be aware of jumping to conclusions



Establishing causation: study design

 Randomized controlled trials are studies in which human volunteers are randomly assigned to receive either the agent being studied or an inactive placebo, usually under double-blind conditions (where neither the participants nor the investigators know which substance each individual is receiving), and their health is then monitored for a period of time.

(https://www.acsh.org)



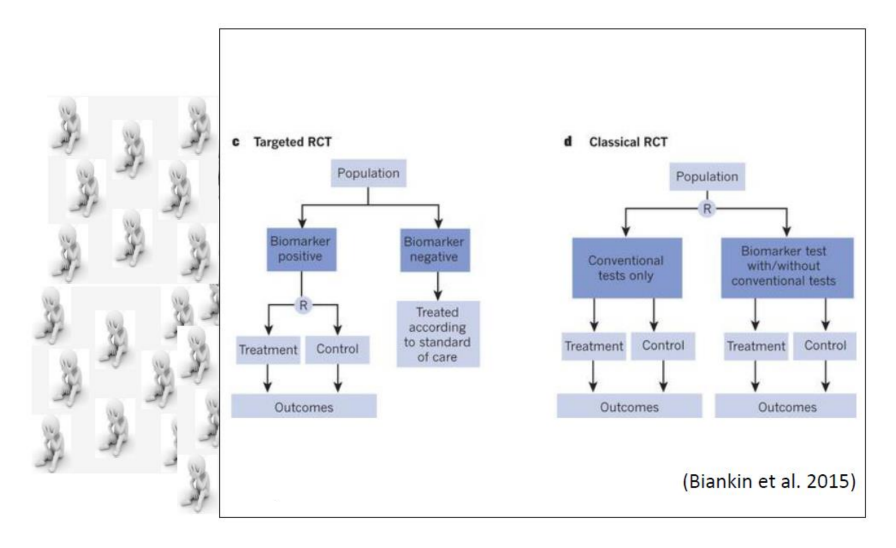
Philos Stud (2010) 147:59–70 DOI 10.1007/s11098-009-9450-2

What are randomised controlled trials good for?

Nancy Cartwright

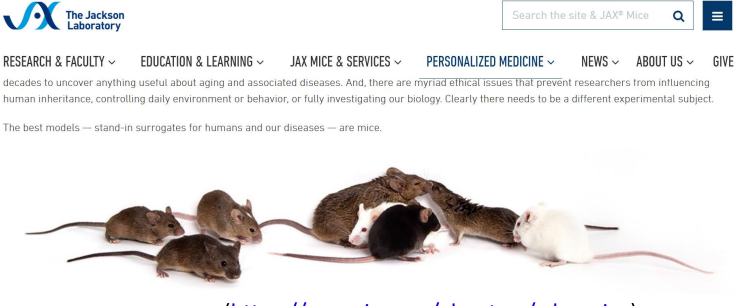
Abstract Randomized controlled trials (RCTs) are widely taken as the gold standard for establishing causal conclusions. Ideally conducted they ensure that the treatment 'causes' the outcome—in the experiment. But where else? This is the venerable question of external validity. I point out that the question comes in two importantly different forms: Is the specific causal conclusion warranted by the experiment true in a target situation? What will be the result of implementing the treatment there? This paper explains how the probabilistic theory of causality implies that RCTs can establish causal conclusions and thereby provides an account of what exactly that causal conclusion is. Clarifying the exact form of the conclusion shows just what is necessary for it to hold in a new setting and also how much more is needed to see what the actual outcome would be there were the treatment implemented.

Designing RCTs for testing precision-medicine strategies is an evolving field!



Establishing causation: wet lab experiments in model organisms

• Gene knock-out experiments



(<u>https://www.jax.org/about-us/why-mice</u>)

• The findings of animal experiments may not always be directly applicable to the human situation because of genetic, anatomic, and physiologic differences or the entity of exposures a human being has experienced

Establishing causation: dry lab

- Try to mimic in vitro what you would like to do in vivo
- **Causal inference** is the process of drawing a conclusion about a causal connection based on the conditions of the occurrence of an effect.
- The main difference between causal inference and inference of association is that the former investigates the response of the effect variable **when the cause is changed**.

Statistics Surveys Vol. 3 (2009) 96–146 ISSN: 1935-7516 DOI: 10.1214/09-SS057

Causal inference in statistics: An overview^{*†‡}

Judea Pearl

Computer Science Department University of California, Los Angeles, CA 90095 USA e-mail: judea@cs.ucla.edu

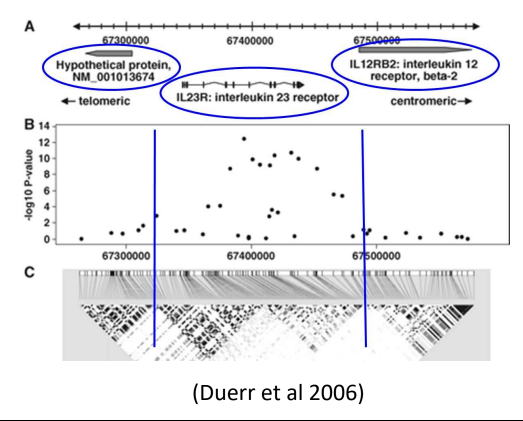
Abstract: This review presents empirical researchers with recent advances in causal inference, and stresses the paradigmatic shifts that must be undertaken in moving from traditional statistical analysis to causal analysis of multivariate data. Special emphasis is placed on the assumptions that underly all causal inferences, the languages used in formulating those assumptions, the conditional nature of all causal and counterfactual claims, and the methods that have been developed for the assessment of such claims. These advances are illustrated using a general theory of causation based on the Structural Causal Model (SCM) described in Pearl (2000a), which subsumes and unifies other approaches to causation, and provides a coherent mathematical foundation for the analysis of causes and counterfactuals. In particular, the paper surveys the development of mathematical tools for inferring (from a combination of data and assumptions) answers to three types of causal queries: (1) queries about the effects of potential interventions, (also called "causal effects" or "policy evaluation") (2) queries about probabilities of counterfactuals, (including assessment of "regret," "attribution" or "causes of effects") and (3) queries about direct and indirect effects (also known as "mediation"). Finally, the paper defines the formal and conceptual relationships between the structural and potential-outcome frameworks and presents tools for a symbiotic analysis that uses the strong features of both.

Keywords and phrases: Structural equation models, confounding, graphical methods, counterfactuals, causal effects, potential-outcome, mediation, policy evaluation, causes of effects.

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Establishing causation: GWAS – finemapping (dry lab)

• As opposed to association studies that benefit from LD, the main challenge in identifying causal variants at associated loci analytically (**finemapping**) lies in distinguishing among the many closely correlated variants due to LD



Establishing causation: Colocalization analysis (not to be confused with protein colocalization)

- Estimates the posterior probability that the same variant is causal in both a GWAS and eQTL study while accounting for the uncertainty of LD
- Example statistical methods following a Bayesian statistical framework: eCAVIAR (Hormozdiari et al. 2016), COLOC (Giambartolomei et al. 2014)
- Posterior support for the following hypotheses:

H0: no causal variants for either trait;

H1: a causal variant for disease association (GWAS) only;

H2: a causal variant for gene expression association (eQTL) only;

H3: two distinct causal variants, one for each trait;

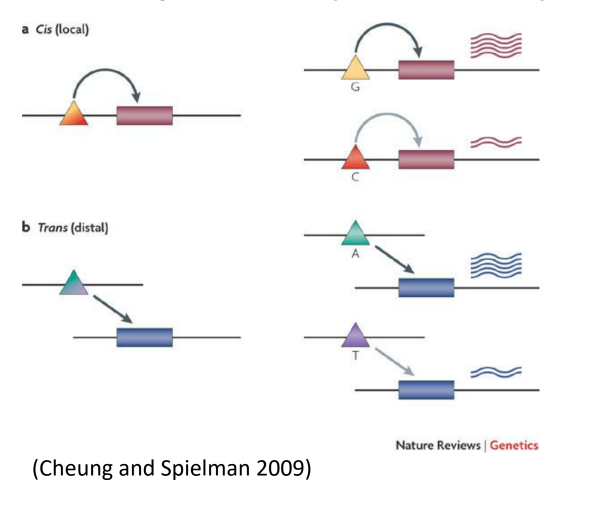
H4: a single causal variant common to both traits (co-localization).

5. d Interpretation

Functional genomics analyses: incl transcriptomics

- An eQTL is a locus that explains a fraction of the genetic variance of a gene expression phenotype.
- One of the fundamental needs for the interpretation of the effects of genome variants is the understanding of the specific biological effect such variants have in the cell, which provides a handle to the biology of the disease or organismal phenotype.
- GWAS have demonstrated that the majority of such variants are found in non-coding regions of the genome and are therefore likely to be involved in gene regulation. Hence, there should be interpretational advantages in analyzing these variants in the context of gene expression (in cells/tissues) (Nika and Dermitzakis 2013)

Functional genomics analyses: incl transcriptomics



 Cis-acting variants are found close to the target genes and trans-acting variants are located far from the target genes, often on another chromosome.

 Different allelic forms of the cis- and transacting variants have different influence on gene expression.

Functional genomics analyses: incl transcriptomics

DOI: 10.1038/s41467-017-01261-5

OPEN

Functional mapping and annotation of genetic associations with FUMA

Kyoko Watanabe¹, Erdogan Taskesen ^{1,2}, Arjen van Bochoven³ & Danielle Posthuma ^{1,4}



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Changing units of analysis: from SNPs to genes

European Journal of Human Genetics (2019) 27:811-823 https://doi.org/10.1038/s41431-018-0327-8

ARTICLE

Comparison of methods for multivariate gene-based association tests for complex diseases using common variants

Jaeyoon Chung 12 · Gyungah R. Jun^{23,4} · Josée Dupuis⁴ · Lindsay A. Farrer^{12,4,5,6,7}

Received: 13 December 2017 / Revised: 30 October 2018 / Accepted: 4 December 2018 / Published online: 25 January 2019 © The Author(s) 2019. This article is published with open access

Abstract

Complex diseases are usually associated with multiple correlated phenotypes, and the analysis of composite scores or disease status may not fully capture the complexity (or multidimensionality). Joint analysis of multiple disease-related phenotypes in genetic tests could potentially increase power to detect association of a disease with common SNPs (or genes). Gene-based tests are designed to identify genes containing multiple risk variants that individually are weakly associated with a univariate trait. We combined three multivariate association tests (O'Brien method, TATES, and MultiPhen) with two gene-based association tests (GATES and VEGAS) and compared performance (type I error and power) of six multivariate gene-based methods using simulated data. Data (n = 2000) for genetic sequence and correlated phenotypes were simulated by varying causal variant proportions and phenotype correlations for various scenarios. These simulations showed that two multivariate association tests (TATES and MultiPhen, but not O'Brien) paired with VEGAS have inflated type I error in all scenarios, while the three multivariate association tests paired with GATES have correct type I error. MultiPhen paired with GATES has higher power than competing methods if the correlations among phenotypes are low (r < 0.57). We applied these genebased association methods to a GWAS dataset from the Alzheimer's Disease Genetics Consortium containing three neuropathological traits related to Alzheimer disease (neuritic plaque, neurofibrillary tangles, and cerebral amyloid angiopathy) measured in 3500 autopsied brains. Gene-level significant evidence ($P < 2.7 \times 10^{-6}$) was identified in a region containing three contiguous genes (TRAPPC12, TRAPPC12-AS1, AD11) using O'Brien and VEGAS. Gene-wide significant associations were not observed in univariate gene-based tests.

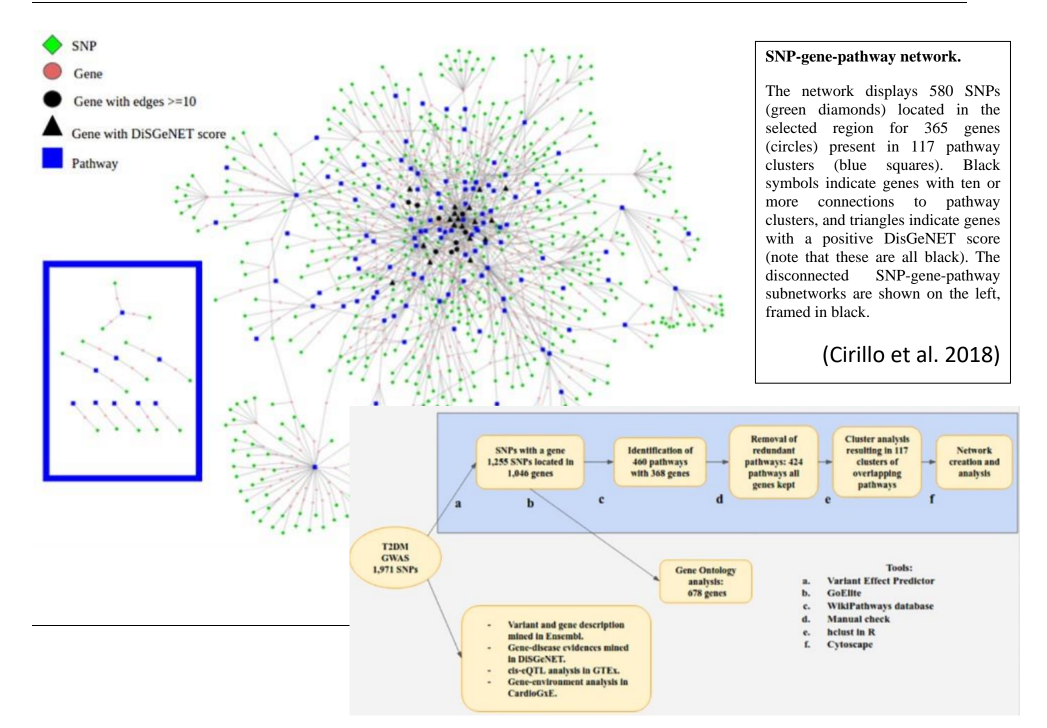
ESHG

() ()

Changing units of analysis: from SNPs to (genes to) pathways

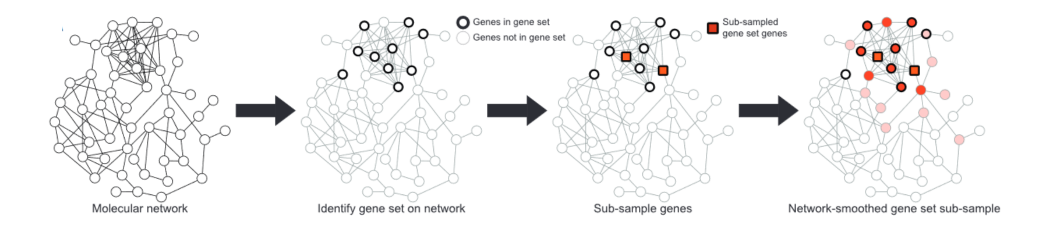
- A biological pathway is an example of a biosystem, that can consist of "interacting" genes, proteins, and small molecules.
- A biosystem, or biological system, is a group of molecules that interact in a biological system.
- Another type of biosystem is a disease, which can involve components such as genes, biomarkers, and drugs.
- The NCBI BioSystems Database currently contains records from several source databases: KEGG, BioCyc (including its Tier 1 EcoCyc and MetaCyc databases, and its Tier 2 databases), Reactome, the National Cancer Institute's Pathway Interaction Database, WikiPathways, and Gene Ontology (GO).

(https://www.ncbi.nlm.nih.gov/Structure/biosystems/docs/biosystems_about.html)



Integrating GWAS results & biological networks

- Huang et al. (2018) evaluates 21 human genome-wide interaction networks for their ability to recover 446 disease gene sets.
- While all networks could recover disease genes, STRING, ConsensusPathDB, and GIANT networks gave the best performance overall.



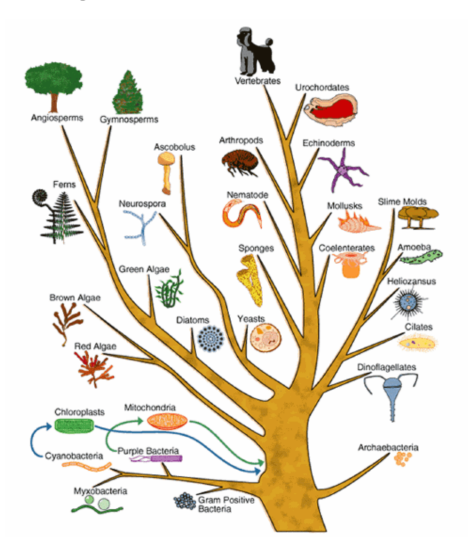
(https://www.cell.com/cell-systems/pdf/S2405-4712(18)30095-4.pdf)

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Model organisms as an extra source of information: accelerating interpretation

- Suppose that we have an unknown human DNA sequence that is associated with the disease cystic fibrosis.
- A bioinformatic analysis finds a similar sequence from mouse that is associated with a gene that codes for a membrane protein that regulates salt balance.
- A good bet may be that the human sequence also is part of a gene that codes for a membrane protein that regulates salt balance.

Model organisms as an extra source of information: assessing importance



• Conserved sequences

More about sequencing & associated analyses in subsequent classes

6 Adding levels of complexity [via homework assignments]

- 6.a Trait heterogeneity in GWAS
- **6.b Missingness**
- **6.c Multiple testing**
- **6.d Multiple studies**
- 6.e When variants become rare
- 6.f Non-independent effects
- 6.g Confounding in the context of 6a-6f

Questions?

Main supporting docs to this class (complementing course slides)



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PLOS COMPUTATIONAL BIOLOGY

Education

Chapter 11: Genome-Wide Association Studies

William S. Bush¹*, Jason H. Moore²

Department of Biomedical Informatics, Center for Human Genetics Research, Vanderbilt University Medical School, Nashville, Tennessee, United States of America,
 Departments of Genetics and Community Family Medicine, Institute for Quantitative Biomedical Sciences, Dartmouth Medical School, Lebanon, New Hampshire, United States of America



A tutorial on statistical methods for population association studies

David J. Balding

Abstract | Although genetic association studies have been with us for many years, even for the simplest analyses there is little consensus on the most appropriate statistical procedures. Here I give an overview of statistical approaches to population association studies, including preliminary analyses (Hardy–Weinberg equilibrium testing, inference of phase and missing data, and SNP tagging), and single–SNP and multipoint tests for association. My goal is to outline the key methods with a brief discussion of problems (population structure and multiple testing), avenues for solutions and some ongoing developments.

Nature reviews Genetics 2006; 5:63-70 – for those interested in technical (statistical) details