Applied Genomics

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Course structure

- Population genetics
- Genome structure and variability in vertebrates (we may mention plants and bacteria)
- High throughput genomic platforms
- Applications of NGS
- Array comparative genome hybridization
- PLINK, genetic data analysis. How to use this software and apply some design using this tool
- Linkage analysis and genetic mapping
- QTL analysis

Examination mode

- Final exam has 2 levels
 - Preparation of a genomic project
 - * A text should be written including an appropriate introduction to the problem/question that the experiment or project would like to analyse or answer, aim of the project, a section with materials and methods, expected results and impact
 - * The project should be submitted to the professor one week before the interview
 - * We should specify what is the aim of the project and what I'd like to solve with it
 - · If it makes sense, we can undergo a discussion with him
 - * The project is based on money: we'll have a budget
 - Interview based on the project submitted and other two questions
 - * Only students that are positively evaluated at the first level are admitted at the second level
 - * Evaluation of basic knowledge
- We get one extra point if we pass at the first attempt
- It is important to follow him
- We'll have an example of a project, the topic of the project it's up to us
- We need to choose a complex genome/organism
- Each one will have a different budget
- It's better to do the project according to what we discuss in the lectures
- It has to be something new
- The first date would be in February after Winter School and another one in March
- Near to the end of the course we'll have a test with 30 questions to test our level (it won't count for the final score)

Introduction

- Genomics is the study of genome structure and function
- The genome is the entire genetic content of an organism

- Applied genomics is the use of technologies, tools and experimental designs to analyse genome and extract information form them
- Genetics studies differences: we cannot track things that are not different among individuals
- A reference genome of a species is the basis used for analyzing the genome of an individual
 - In some cases if I do not have a reference genome I can use that of a similar species
- We have about 2 nuclear genomes per cell, but even thousands of mithocondrial genomes
- Mithocondrial genomes can be not all equal: heteroplasmy
- The human nuclear genome is around 3 Gb, the mithocondrial genome 16.7 Kb
- Population genetics is important for this course
- Small population are susceptible to high levels of inbreeding
- Differences between population arise when there are reproductive barriers
- Effective population size is the number of individual that originated a population
 - It is a measure of inbreeding
- Sex determination can be mediate by sex chromosomes, temperature, ploidy
- Phenotype is influenced by the environment
- A phenotype is an observable charachteristic
- Comparative genomics is the study of genomic differences between species
 - It is really helpful for genome annotation
- The first draft of the human genome was completed in 2001, and the HGP was started in 1990
- 3% of human DNA is coding
- Repetitive sequences are problematic for assembling genomes
- Nuclear DNA is 99.99% identical among individuals, while mitochondrial genome is more similar
- The simplest definition of gene is "coding region"
- We can predict the phenotype of an animal just looking at the genotype (!)
- To do applied genomics I need a reference genome
- If I do not have a reference genome for my species of interest, I need to construct it or I can use one of
 a closely-related species
- Genomics produces around 10 Zb of data per year
 - We cannot store everithing: we must select what is worth storing and what is not
 - It is interesting to look at portions that differ from the reference genome
- The cost of sequencing is dropping in a way similar to Moore's law
 - Around 2008 the drop was much faster than Moore's law, thanks to NGS
- The shotgun approach does not have a particular target, it sequences everything
- Genomic data are typically stored in the cloud
- Hardy-Weinberg equilibrium

$$-\begin{cases} p^2 + q^2 + 2pq = f(AA) + f(Aa) + f(aa) = (p+q)^2 = 1\\ p+q=1 \end{cases}$$

- The allele frequencies refer to the current generation, while the genotype frequencies refer to the next generation
- It holds in absence of genetic drift, non-random mating, selection, migration, mutation
- Mendel's first law: alleles segregate with other alleles
- Mendel's second law: independent assortment
- Mendel's third law: some alleles are dominant on others
- We reviewed PCR, agarose gel electrophoresys and Sanger sequencing basics
- Reference genomes can be found in the Ensemble database
- A genome assembly can be done in chromosomes or in scaffolds
- Scaffolds are assembled from contigs
- Sometimes it is not possible to assemble entire chromosomes
- The quality score of an assembly (n50) is the minimum size of scaffolds that contain 50% of the assembled genome
- A human chromosome is on average 80-100 Mb
- Penetrance is the proportion of individual with a given genotype that manifest the associated phenotype
- ChiPseq (chromatine immunoprecipitation) is a method used to analyse DNA-protein interactions

- The output is a library of sequences that bind the protein of interest
- The first step is to fix the proteins with DNA using formaldehyde
- Subsequently, cells are lised and DNA fragmented
- The sequences of interest are recovered with Ab against the protein of interest
- I reverse the DNA-protein binding and sequence the fragments
- If I want to reduce cost, I can sequence only the part of interest, for instance the exome
 - In order to sequence the exom I need a capturing system
 - If not commercially available I have to evaluate if developping a capturing system is worth it
 - In order to enrich for the exome, I need to have specific probes that bind to exon regions, either in solution (on beads) or in microarrays

Sanger sequencing

- Sanger sequencing was developed by F. Sanger in 1977
- It uses DNA polymerase to extend a primer using genomic DNA as a template
- ddNTPs are incorporated in the reaction mixture in a controlled ratio
 - ddNTPs lack a 3'-OH and thus cannot be used for extending the DNA chain
 - The ration ddNTP/dNTP is typically around 1/100, but it depends on the length of the desired sequence
 - When a ddNTP is incorporated the extension reaction is interrupted
- In the original Sanger publication and in the first approaches radioactive labels were used
 - Either the primers were labeled, or a single dNTP (not ddNTP!) in the mixture
 - 4 different reaction were performed and separately processed, each with a single ddNTP type
- In the current approach, dye-terminator sequencing, the 4 different ddNTPs are labeled with different fluorophores
 - The strand itself and the primers are not labeled
 - A single reaction is done with all the 4 labeled ddNTPs and run together on the capillary gel
- The ssDNA fragments are run in a capillary electrophoretic apparatus
 - ssDNA hairpins are a serious probelm for length resolution
 - A denaturing polyacrilamide-urea gel is used
- In dye-terminator sequencing an electropherogram is produced, showing fluorescent intensity peaks for each of the 4 channels
- Sanger sequencing primers are designed according to the region that I want to sequence
 - In general I use Sanger when I have a reference!

Next generation sequencing

- NGS platforms: Illumina, Ion torrent (Thermo fisher), PacBio, Nanopore, 454
 - PacBio was going to be acquired by Illumina, but the antitrust opposed and the merger was canceled
 - We have short reads, therefore assembly is difficult
 - 454 (La Roche, pirosequencing) is practically dead today
- The depth of coverage is the number of unique reads that contain a specifc nucleotide in the assembly
- Sequencing a mammalian genome at 50x costs around 2000€+VAT in China or South Korea
 - BGI (Bejing genome institute) is the largest sequencing provider and it is chinese
 - NOVOGENE is from South Korea
 - If I chose to use these services, I need to consider shipping restrictions, costs, and product degradation
- In sequencing, if we are not sure about a variant we exclude it
- Wen I do genotyping by sequencing, the regions of interest have a very high depth of coverage so I can trust the results
- We have tools for allignment of reads to a reference like bowtie
 - They produce a BAM file

- There are tools for calling mutations, Indels, ecc.
- If I do not have enough money to sequence any individual, I can pool DNA samples in group (i.e. breed) and do a sequencing for each group
- A reduced representation library is obtained by digestion of the genome
 - I run the digest on agarose and retrieve only a specific subset of MW
 - If I see definite bands in the gel, these probably come from repeated regions that are cut at the same length
 - * I want to exclude this (!)
 - In the digestion, I can choose a restriction enzyme with a long target sequence if I want longer fragments (cut site less probable!) and vice versa
- Fastq is similar to fasta but it has additional information on it
 - It uses ASCII symbols to code a quality score (PHRED score, from the homonimous software) in a separate line from the one where the bases are stored
 - * PHRED uses hard-code lookup tables of peak charachteristics to estimate quality
 - The highest quality is 93 for fastq
 - The quality score is the ASCII code of the character (!)
 - * ASCII 33 to 126 are used, encoding scores from 0 to 93
 - The quality score rarely exceeds 60 in raw data, but can be higher in assemblies
 - The threshold quality score now accepted for base calling is 30
- Illumina reads file looks like a fastq, but quality scores have a different scale
 - If my file has scores higher than 90, it is an Illumina file
- Allignments are saved in .sam format, a tab-delimited text file that can be converted in a binary .bam file
 - The sam file contains the sequence of reads, their genomic alignment coordinate, contig, mate read name
 - samtools is used for working with sam files

Ion torrent

- There are many sequencing chips, with different throughputs
- The sequencing device is a semiconductor chip with millions of nano-wells
 - Each well is represented as a pixel
- DNA fragments are clonally amplified on acrylamide beads that are poured on the chip and go in the wells, one for each well
- \bullet The chip is sequentially flodded with the 4 nucleotides, allowing a stepwise progression of DNA synthesis
- The addition of a nucleotide releases a proton, changing the pH of the well
- The drop in pH is recorded as a base call for the well
- I have clonal amplification on positively charged spheres
- During the addition of a nucleotide, a proton is released
- If I add nucleotides one at a time, I can sense the pH change due to the many protons released by the clones
- If I have multiple nucleotides of the same type in a row, I get a stronger signal
- $\bullet\,$ Regions with a stretch of the same nucleotide are called omopolymeric
 - It is difficult to exactly count the number of nucleotides in these regions
- The raw data produced is called ionogram
- We use universal adapters with a specific portion to amplify the DNA fragments
- The machine is called ion or proton torrent
- In the preparation step we obtain thousands of template molecules
- The first step of the workflow is library preparation
- Library preparation depends on the kind of samples
- I can sequence amplicons, genomes, RNA libraries
- I can only sequence small fragments: I need a fragmentation step
- Fragmentation can be done by sonication or with aspecific DNases
- Playing with the time of fragmentation, I can modulate the length of the fragments

- Frequently I need to try in different ways (!)
- It is a random process!
- I have to amplify all my fragments by PCR
- It will take forever with standard PCR, so I do emulsion PCR where every drop harbours a reaction
- I then do an electrophoresis to get only the fragments of a certain size
- NGS can typically sequence from 25 up to 400 nucleotides, but the highest throughput is around 100 BP per read
- The ideal case is that in a droplet I have a bead and a single DNA fragment
- The DNA attached to the acrylamide beads after emulsion PCR is retrieved with the ion sphere particle enhancement technique
 - I use magnetic beads with streptavidin to capture the amplified acrylamide beads
 - The beads bind biotin-labeled nucleotides on the beads, so this will select only successfully amplified beads
 - After magnetic pull I denature the streptavidin and release the acrylamide template beads
- To increase my output, I can regolate my flow (nucleotides added) considering gc content of my target
 - One flow is the addition of 1 nucleotide to the chip
- The real throughput of my sequencing system is lower than the theoretical one
 - If 2 different fragments are amplified together I get mixed reads, and they give me false sequences as output
 - I need to ignore the mixed reads, but they will waste some of my sequencing wells
 - The same for 2 beads with the same fragment: duplicate reads
- The ionogram is expected to show on average a peak every 4 flows
 - If I see too many peaks close to each other, I probably have mixed reads
- I have a reference sequence known, and if this sequence reaches a threshold signal I keep my read, otherwise I discard it
- In missed reads I have too many empty spaces in each read, more than statistically reasonable

Roche 454

- It works in similar way to Ion Torrent, but it senses the release of pyrophosphate during elongation
 - NUcleotides are added in flows like for ion torrent
 - PPi and adenylil-sulfate are converted to ATP and sulfate by sulfurylase (Sulfate adenylyltransferase)
 - ATP is used by luciferase releasing a photon
 - APyrase continously removes excess NTPs that are not incorporated
- It was the first NGS to be developed, but also the first one to become obsolete
- Like Ion Torrent, it uses beads on a chip and the target is amplified by emulsion PCR
- PP_i is used by sulphurylase to synthetize ATP, ATP is used by luciferase to produce light
- Light emission is sensed by a CCD camera producing a pyrogram
- This technique was abandoned because it is too expensive
 - The cost is mainly due to the many enzymes used (sulphurylase, luciferase, apyrase)
 - The CCD camera is expensive
- It has revolutionized bacterial taxonomy because it allowed to sequence the rRNA 16s
 - This is because it can produce longer reads than other NGS techniques

Illumina

- Library preparation
 - DNA is fragmented by sonication
 - Overhangs are blunted or repaired
 - * I use T4 polymerase and Klenow fragment for repair and blunting
 - * Polymerase activity fills the 5' overhangs and exonuclease activity removes the 3' overhangs
 - All the 5' ends are poshporylated, since their status is unknown after sonication
 - * I use T4 PNK

- A single A is added at the 3' of the fragments to avoid ligation of different fragments in the blunt ligation step
 - * Adapter on the contrary have a single T 3' overhang to facilitate ligation with the fragments
- Adapters are ligated to fragments with ligase
- PCR is used for selectively amplifying fragments that have been correctly ligated to adapters
 - * The primers anneal to the end of the adapter and have a tail that adds sequences used for the library amplification in the flowcell
- In a flowcell, I have many oligos that can anneal with the inserted sequences at the end of the adapters
- Bridge amplification amplifies the fragments in the flowcell
- After bridge-amplification, I get clonal clusters of fragments
- In the elongation step I add all the nucleotides together, marked with fluorophores and reversibly inhibited at their 3'
 - These modified nucleotides are called reversible dye terminator chemistry
 - The primers used anneal to the adapter that is not part of the flowcell
- The elongation is stepwise because there is a block in 3' that inhibits elongation
 - I can easily deal with homoplimeric regions (!)
 - Because of this the error rate is much lower
- After nucleotide incorporation I excite the flowcell with laser and capture the resulting fluorescence with a CCD camera
- An enzimatic step that cleaves the 3' block and the fluorophore
- I can sequence both ends of my fragments, and this is really useful for the assembly step
 - I can play with fragment size to obtain my contigs
- When I sequence a genome, I need to consider sequencing depth and coverage
 - Sequencing depth is the average number of times that a nucleotide in my reference genome is represented in a read
- In order to reduce cost, I can run more samples in the same lane by using a barcode attached to my
 fragments
- MySeq can be used for metagenomics (16S sequencing, 24 samples per lane) and for microbial WGS
 It can procduce 3-7 million reads per lane
- HiSeq can be used for WGS (3-4 lanes per genome) and exome capture (4 samples per lane)
 - It can procduce 100-160 million reads per lane

AB SOLiD

- It is dead by now, but could be potentially great because it gives the highest throughput
- Its reads are really short (30 bp) so it is computationally heavy to asseble the reads and it is impossible to use with repetitive regions
- It is based on sequencing by oligo ligation detection
- DNA is fragmented and ligated to P1 and P2 adapters
 - I can prepare simple or mate-paired libraries
 - Mate-paired libraries use also an internal adapter
- A clonal population of beads is created from the fragments by emulsion PCR
- Beads are covalently attached to a glass slide
- The slide is incubated with an universal sequencing primer, di-base probes and DNA ligase
- The primer has its 5'P oriented towards the fragment (away from the bead)
- There are 16 (all the possible 4² 2-mers) different dibase probes, marked with 4 different fluorophores
 - Each dye could correspond to 4 of the 16 di-base probes
 - The probes have the di-base specific sequence at the 3' and 6 additional aspecific positions at their 5'
- The fluorescent signal is acquired
- 3 bases at the 5' of the probes are cleaved, removing the dye
- Phosphatase is also used so to cap eventual unextended fragments
- The step is repeated until needed by ligating another di-base probe at the 5' of the previous one
- Since each probe is 8 nucleotides long but 3 nucleotides are cleaved, the system interrogates 2 bases

- every 5 (2 interrogated and 3 no, then 2 interrogated and so on)
- Everithing melted and repeated with different primers that have an offset of -1, -2, -3, and -4 bases
 - In this way I get a reading for the missing positions
 - I get 2 reads from 2 different probes at each position
- There is an unique combination of bases that can give the combination of reads, so I can reconstruct the original sequence
 - Each ordered combination of colors corresponds to a base

Complete genomics

- It is used for re-sequencing common genomes
- Fragments are made circular and then amplified by rolling circle amplification, obtaining DNA nanoballs (DNBs)
- DNBTMs are around 200 nm in size
- They are anchored in a chip obtained by photolitography
- The chip is made so to have a grid pattern of sticky spots
 - A sticky spot is just a small well in the surface of the chip
- Each sticky spot recives exactly one nanoball, thanks to proprietary technology
- Sequencing is done by combinatorial probe-anchor ligation (cPAL)
- The DNB contains genomic DNA and an initial adapter sequence
- 1 of 4 labeled anchor probes made of a specific position and 8 aspecific positions binds to the adapter and it is ligated with ligase (similar to AB SOLiD)
- Complete genomics was acquired by BGI and it does not sell sequencing equipment
- You send samples to them and they produce data in ~ 100 days

PacBio

- It is really a promising technology
- Reads are long, up to a 6-10 kb, but troughput is low
- The long reads make assembly easy and don't require really high depth of coverage
- The error rate is quite high (15%), and probably it cannot be reduced under 5%
- It is costly, 40k€ for 10x in mammalian genomes
- It does not require amplification, so no bias is introduced
- It is a golden standard for new sequencing projects, usually matched with Illumina
 - Assembling PacBio reads can be difficult due to the high error rate
 - Single Illumina reads are around 1% error, so assembling Illumina reads to PacBio reads is much easier
 - I can use PacBio to resolve repetitive regions and difficult-to-amplify regions, and Illumina for depth of coverage and accuracy
- PacBio was going to be bought by Illumina, but the anti-trust opposed it and themerger was canceled
- It is based on Single Molecule Real Time (SMRT) Sequencing
- Hairpin adapters are ligated to both ends of dsDNA, so to create a circular molecule
- A sequencing primer and DNA polymerase are added and bind the DNA molecules, but do not extend
 it since nucleotides are not added
- The DNA fragments with bound primer and polymerase are inserted in a SMRT cell
- The SMRT cell contains millions of wells on its surface, called Zero-mode waveguides (ZMWs)
- Each ZMW hosts a single DNA molecule with a single DNA polymerase and primer
- Labeled nucleotides are added and their addition is detected in each ZMW
- 2 modes of operation are possible: Circular consensus Sequencing (CCS) and Continuous Long Reads (CLR)
- In CCS the circular DNA is read again and again generating an high-fidelity (HiFi) read, with 99% accuracy
- In CLR the molecule is as long as possible and it is read for as many nucleotides as possible, so to generate long reads

Oxford nanopore

- Long reads, but high error rate and low thorughput
- The reads can potentially be very long, up to 100kb depending on library preparation
- DNA passes trough a modified hemolysine pore altering the ion flow thorugh the pore
 - The pore is around 1 nm in diameter (half the widdht of DNA!)
- A motor protein pre-loaded on an adapter oligo regulates the speed at which DNA moves through the
 pore and acts as an helicase
- Many DNA molecule pass in parallel through many nanopores (around 500 in MinION, 2000 in GridION)
 placed on a membrane
- Each nanopore has its own sensor and it is placed in a nanowell
- Bases are read in 4-mers and then the signal of subsequent reads is interpolated by a Recurrent Neural Network (RNN)
- Interpretation of the raw data is difficult, because the meaning of reads depends on the sequence context
 - Error rate is around 4%
 - I can recognise not only ATCG, but also U, 5mC, and other modified bases
 - Machine learning!
- The platform is cheap (~5000€, sometimes given for free) but adapters and accessories are expensive
- It is invaluable when I have to work on-site, since it is small and portable
- MinION can produce 150 Mb per run with 48kb reads
- Oxford Nanopore can be used not only for sequencing, but also for detecting protein-DNA interactions, small molecules

Other NGS techniques

- Intelligent BioSystems Mini20 is a sequencing by synthesis platform designed for clinical use
 - It has 100 nt long reads, but it is expected to be able to compete with Sanger sequencing
 - It is sensitive to repeats
 - Full costs are still not clear, but the instrument costs 120k \$ and the disposable flow cell 150 \$
- Genia Technologies is an early stage announcment for a system that should combine Ion Torrent and Oxford Nanopore
 - It claims a sensitivity 1-2 orders of magnitude greater than Oxford Nanopore, with as many as 100k pores per chip, with as many as 100k pores per chip
 - Planned sample cost less than 100 \$

Aplotypes

- We can detect crossing-over by looking for the association of genetic markers
- An aplotype is a cluster of genes that are usually eredited toghether
- The probability of CO between 2 genes is measured in cM
 - 1 cM is a genetic distance such that in 100 meiosis I expect 1 CO
 - It is around 10^6 nucleotides for mammals
- If I have a simple dominant trait, I am certain only about the allele frequency of the recessive
 - I can recover it by recessive allele = $\sqrt{recessive\ phenotype}$
 - Doing this, I am assuming that the population is infinite, there is no mutation, no selection, no genetic drift, no migration, random mating
- If the observed genotype frequencies are different from the ones expected from HW equilibrium, It means that there are factors at play that perturbate the equilibrium
 - There can also be genotyping problems (my region is difficult to sequence and I do not get the right sequence)
- Two loci are in linkage disequilibrium if they do not occur randomly with respect to each other
- Aplotypes are patterns of genetic variation in populations
- The genotype is not sufficient for predicting the aplotypes
 - I cannot differentiate if a variation is in one chromosome or the other (!)

- We need information on aplotype frequencies or on the parents
- PHASE is a website for analyzing aplotypes
- I cannot determine the aplotype by only looking at the genotype: I need data on the population

Genotyping

- Genotyping means to determine the genotype at one locus
- I can perform high throughput genotyping with beadchips
 - I have beads with primers that anneal in different positions in the genome, so to be evenly spaced and belowe the linkage disequilibrium lenght
 - The output of a beadchip is essentially a .map file with additional experimental information (signal intensity for the SNP)
 - The position of some probes in the genome is unknown, so the row of their SNP starts with 0 (chromosome) and ends with 0 (position)
- The main genotyping platforms are from Illumina and Affimetrix
- The probe is designed so to
 - Bind to a unique region (it has to be long enough!)
 - It has to have standard GC content, so I can melt all the chip at the same temperature
- The specific fragments to be genotyped are detected by primer extension
 - I have a primer right in front of a SNP
 - I add the 2 possible nucleotides for the SNP labeled with different fluorophores and blocked
 - I see what happens
- The minor allele frequency (MAF) is the frequency of the rarer variant of a SNP
 - It can go from 0 to 0.5
- I do not need to genotype all the SNPs
 - I can take advantage of linkage disequilibrium to detect aplotypes
 - Polimorphic sites are more informative than sites with rare variants, so I tend to focus on them for determining an aplotype
- Genotyping by sequencing (GBS) allows to detect unknown SNPs and it is typically done with pooled reduced representation libraries
- Illumina can produce customized genotyping chips

Plink

- A pedigree is a standardised representation of individuals in a population and relationships among them
 It can be represented in plane text or in binary form
- Plink is an important tool for working with reference genomes
 - It can work with text files (--file parameter, without extension for homonimous .ped and .map files)
 - It can work with binaries (--bfile option)
- PED and MAP file work in pairs: I typically have my_file.ped and my_file.map with the same root name and referring to the same data
- The PED (pedigree) file is a text file with a row for each individual
 - It stores the pedigree of the population
 - This format is standard and it is used by different tools
 - It is Tab-separated and there are fields for the father, mother, sex, family, phenotype, SNPs
 - Missing data are usually reported with 0
- The MAP (map on the genome) file is a text file that has a line for each SNP
 - It reports chromosome number, SNP ID, position, distance from other SNPs
 - It is produced processing the raw output of a genotyping platform
- A polymorphism is such if it has a frequency higher than 1%
- Before doing data analysis, check your data (!)
 - I want to exclude faulty individuals and faulty loci

- Plink can filter out data at a given threshold
- I want to exclude low-frequency alleles: my focus is the population, not the individual
- I can exclude SNPs that violate the HW equilibrium
- I can exclude mendelian errors: genotypes that are impossible given the parents
- Basic usage
 - --freq gives the frequency of a SNP
 - If I don't trust the data provider about the sexes, I can check for absolute homozigosity at X loci: in this case I have a male
 - I can want to filter out duplicates due to sampling errors
 - If I am working with non-human or I have partially assembled scaffolds, I need to specify
 - --allow-extra-chromosomes or the species, if available in plink (es. --sheep)
 - --out specifies the root filename of the output
 - --noweb is usually required otherwise it checks forever for updates

De novo sequencing

- The human genome is repeat rich
- The main approaches are whole genome shotgun and hierarchical shotgun approach (BAC based)
- Hierarchical shotgun allows to resolve repetitive regions by building bigger contigs (!)
- At the time of the first human genome, sequencing was expensive so we could not sequence BACs and then assemble them, we needed to select non-duplicate BACs beforehand
- I start from a gene in a known position in a chromosome, and check which BACs contain it by PCR
 This links my assembly to the physical chromosome
- Genetic maps are linkage maps, and they can be used for assembling genomes
- Physical maps refer to the position of a gene in the chromosome
- A strategy to select overlapping BACs is to digest them with restriction enzymes and search for common fragments among different BACs
- The main problems of hierarchical shotgun are that it is slow and assembly is problematic if some BACs contain chimeric DNA
 - Chimeric DNA is a fragment that is created by the association of fragments from different chromosomes during the construction of the library
- The alternative to hierarchical shotgun is whole genome shotgun
- N50 is a statistics that defines assembly quality in terms of contiguity
 - It is the length of the sorterst contig that allows to surpass 50% coverage of the genome
- The state of the art is to do a first PacBio sequencing to get a rough map to which I can attach subsequent precise Illumina paired-end reads
 - I want to use more than one Illumina run, with different lengths, so to discriminate repetitive regions and to correct errors in the PacBio phase
- Radiation hybrid maps now can be used for refining an assembly
 - I form an hybridome between an immortalized cell from a different species and a normal cell from the organism that I want to sequence
 - The hybrid will lose most of the genome of the normal cell, and it will retain a random fragment
 - In this way I can get a library (!)
 - The evaluation of the retained fragment is done by cariotyping thanks to banding patterns
 - I can then test by PCR to locate specific tags
 - By cross-referencing caryotype and PCR I can get a rough map of in which chromosome genes are (not so useful now, used in the pre-sequencing era)
 - If before the formation of the hybrid I irradiate the normal line, I break its DNA and get small fragments
 - In this case I want to have a very big library, where each clone has a small fragment
 - I can test by PCR in order to understand which markers from which chromosomes I get from each clone
 - If in my assembly I have a contig that I cannot locate, I design PCR primers for that region

- I test all the library with the primers, and I select the clones that contain my tag
- I check those clones for other markers of known position, and I check the ones that are more frequently associated with the tag of unknown position
- In this way, I can say that the unallocated contig is physically linked to a tag of known position
- The distance from between tags defined in this way is defined in cRay

Repeated sequences

- They can be spotted with repeat masker
- This tool can mark SINE, LINE, Alu and will mask it in my sequence
- Masking means to substitute a sequence with a stretch of NNNN of the same length
- Pseudogenes can be processed or non processed (with introns) and they are not recognised by repeatmasker

DNA chips

- In human the average linkage disequilibrium is low, around 1kb
- When effective population size is low, likage disequilibrium is large
 - This is true for lifestock
- In DNA sequencing chips, I detect a series of SNPs distanced about the linkage disequilibrium
 - If 2 SNPs are close enough, I can infer that the sequence in between is what I would expect from the aplotype

Copy number variation

- A CNV is a 1 kb or longer DNA segment present at variable copy number
- They can be discovered by analizing the depth of coverage of the region
 - This does not tell me in which allele the copies are (!)
- There are portions of mithocondrial DNA integrated in the nuclear genome
 - These are called NUMTS and they are mostly pseudogenes, but maybe some of them are functional
 - They are still being integrated, so they tend to be quite variable
 - The ones integrated most recently tend to be really similar to the mithocondrial sequences
- Array competitive genomic hybridization (aCGH) was once a golden standard for CNVs, now it is not
 - It is used for the identification of tumors
 - It is performed on a DNA microarray
 - Single probes are 50-75 nucleotides long and they are syntetized
 - * They are selected so to be spaced around 20 kb apart and to have a specific GC %
 - * I need to have a certain GC % so to be able to do the annealing step for all the microarray at the same temperature
 - * I do not want probes on repeated sequences
 - I do the hybridization with a reference DNA and the sample mixed and marked with different fluorophores
 - I measure the log 2 of the ratio of the intensities in order to call CNVs
 - * 0 means that I have the same number of copies, 1 that I have the double number of copies
 - If I want to decrease the noise I can decide to call only more than 5 (es) sequential calls at the same level
 - * In this way I loose resolution (!)
 - Note that if I compare the X chromosome in males and females, I get double the reads in females
 (!)
 - It is a good complement for cytogenetics

GWAS

- I want to find the association between a phenotype and a genomic locus
- I can genotype individuals with a SNPs array and see if there is association with the phenotype
 - I check allele frequencies that differ in the different cohorts
- The result is a Manhattan plot
 - I have the chromosome length on the x axis (coordinate of the SNPs)
 - In the y axis I have the -log of the p-value for the association
 - * Lower p-values are on the highest part (!)
- I am doing a lot of multiple testing so my threshold must be really high (!)
 - I use the Bonferroni correction or false discovery rate
- Continuous traits tend to be normally distributed
 - On a SNPs A/G I can have 3 possible genotypes: AA, AG, GG
 - I measure the genotype of each individual and its continuous trait
 - I take the means of the groups for each genotype and I perform a statistical test on means, like ANOVA

Detect inbreeding

- The inbreeding coefficient indicates the probability that random positions among 2 individuals are equal by descent
 - It is calculated by tracing a close path on the pedigree of an individual
- Runs of homozigosity (ROH) refer to stretches of chromosome which are completely homozygus
 - This could mean that the 2 stretches are identical by descent (!)
 - The ROH % is equivalent to the coefficient of inbreeding