

Gel-Electrophoresis

- DNA is cut into fragments using an enzyme
- The cut DNA is put on a Gel material
- An electric current is applied on the Gel
- DNA is negatively charge
- DNA fragments will start moving towards the positively charged side
- Smaller fragments move faster
- After some time, we have a separation of the different fragment lengths

DNA Sample

- Some cells are obtained
- The cells are immersed in a nutritious solution on a plate and left to grow and multiply
- The cells are gathered and frozen for future use
- Liquidized DNA is obtained from these cells



Restriction Enzyme

- A restriction enzyme is used to cut the DNA into fragments
- Hind III restriction site
 is A AGCTT



Apply Enzyme

- DNA sample and Hind III are put together in a tube
- The tube is shaken by rotation for DNA and Hind III to mix



Water Bath

- The tube is put on a plate floating on water at 37°C
- It is left for 30 minutes
- This is needed for the Hind III reaction to take place



Preparing the Gel

- In the meantime, we prepare the Gel
- Agarose powder is the basic substance for making the Gel





• The powder is mixed with water in a container



Preparing the Gel

- The container is heated (in a microwave if you want) until the powder completely dissolves in the water
- The solution becomes clear



Preparing the Gel

- The liquid Gel is poured into the inner box
- A comb like piece is put at the edge of the inner box
- The liquid Gel is left to cool and solidify (you can use a fridge)
- When the Gel solidifies, the comb will create *wells* for the DNA samples to be put



H shaped container

Gel Ready

- Gel ready
- Fill the H shaped container with water
- Remove comb



- DNA samples mixed with colored solution and UV reactive solution
- DNA samples inserted into wells
- A sample DNA containing only specific fragments (called ladder) can be used for comparison



ladder 2

Run the Gel Apply electric current DNA is negatively charged

- Fragments will migrate toward the positive charge
- Small fragments move faster





.slart

Viewing

 Gel can be viewed under UV light







- In a hybridization experiment, we try to verify whether a specific sequence known as **probe** binds (or hybridizes) with a DNA fragment.
- If the binding occurs, this means that the DNA fragment contains the sequence complementary to the probe sequence (or parts of it).





First RFLP map in 1987

- Donis-Keller et al. constructed the first RFLP map of the human genome, positioning one RFLP marker per approximately 10 million nucleotides.
- RFLP markers (probes) need to be long enough to span the whole DNA.
- 393 random probes where used to study RFLP in 21 families over 3 generations.
- Computational analysis of recombination lead to the ordering of RFLP markers on the chromosome.

RFLP and Gene Finding

 Using the ordering of RFLP markers on a chromosome, we can approximately determine the location of a gene.

- How?

- Find the difference between the RFLP markers of family members with the disease and family members not having the disease.
- It is likely that the RFLP marker that consistently differ is on the gene responsible for the disease, since family members have more or less the same genetic characteristics.
- But we still don't know where and what the exact gene is.

Physical Mapping

- Genetic mapping and RFLP
 - (1) do not tell the actual distance in base pairs
 - (2) if genes (or markers) are very close, one cannot resolve their order, because the observed recombination frequencies will be zero.
- Physical mapping reflect actual distances
 - Hybridization Mapping
- Restriction Mapping

Hybridization Mapping

- Break several copies of DNA into fragments (using different restriction enzymes).
- Obtain many copies of each fragment (cloning, incorporating a fragment into a replicating host), forming a clone library.
- Clones may overlap (cutting DNA with distinct enzymes), and we want them to (we will see why).
- Fingerprinting the clones: Now use DNA probes, and for every clone determine the list of probes that hybridize with the clone
- When two clones have substantial overlap, their fingerprints will be similar.
- Reconstruct the relative order of the clones using the overlap information (this order is unknown in RFLP)

Hybridization Mapping

- For *n* clones, and *m* probes, the hybridization data consists of an *n* x *m* matrix *D*, such that d_{ij} =1 if clone C_i contains probe p_i .
- Let S be a string over the alphabet of probes *p*₁...*p*_w. S covers a clone C if there exists a substring of S containing exactly the same set of probes as C (order and multiplicity are ignored)
- A simple approximation of physical mapping is the Shortest Covering String.







Unique/Non-Unique Probes

- Non-unique probes: probes are short random sequences that can occur many times in the DNA. Therefore, a probe can hybridize with distant clones.
- Unique probes: probes are sufficiently long and are unlikely to occur twice in the DNA. Therefore, a probe will hybridize with close clones.
- Advantages of non-unique probes: probe generation is cheap and straight-forward.

Restriction Mapping

- Before using the list of probes in a clone as a fingerprint, biologists used the order of restriction fragments in a clone.
- Restriction map as Fingerprinting: If two clones share several consecutive fragments, they are likely to overlap.
- Restriction map of a clone: an ordered list of its restriction fragments (Hard Problem).

Double Digest

- Cut the DNA fragment with enzyme A, then enzyme B, then both
- Obtain a multiset of lengths in each case (using Gel electrophoresis)
- Using this information, construct an order of the lengths

2 1 2 2

- A: {2,2,3}
 B: {3,4}
- A+B: {1,2,2,2}

Partial Digestion

- Instead of obtaining lengths of restriction fragments, the DNA is digested in such a way that fragments are formed by every two cuts and the lengths of all fragments are obtained.
- The problem often might be formulated as recovering positions of points on a line when only some pairwise distances between points are known. (why?)
- Many mapping techniques lead to the following problem: X is a set of points, ΔX is the multiset of all pairwise distances between points in X: $\Delta X = [|x_1 x_2| : x_1, x_2 \in X]$, $E \subseteq X$ is given. Reconstruct X from knowing E alone.
- Partial Digest Problem. Given ΔX, reconstruct X (E=ΔX). Also known as the *turnpike* problem in computer science, construct the geography of the highway from knowing the distance between every two exits.
- No polynomial time algorithm for this problem is yet known, but in practice, efficient algorithms exist.