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RAPD

Randomly Amplified Polymorphic DNA

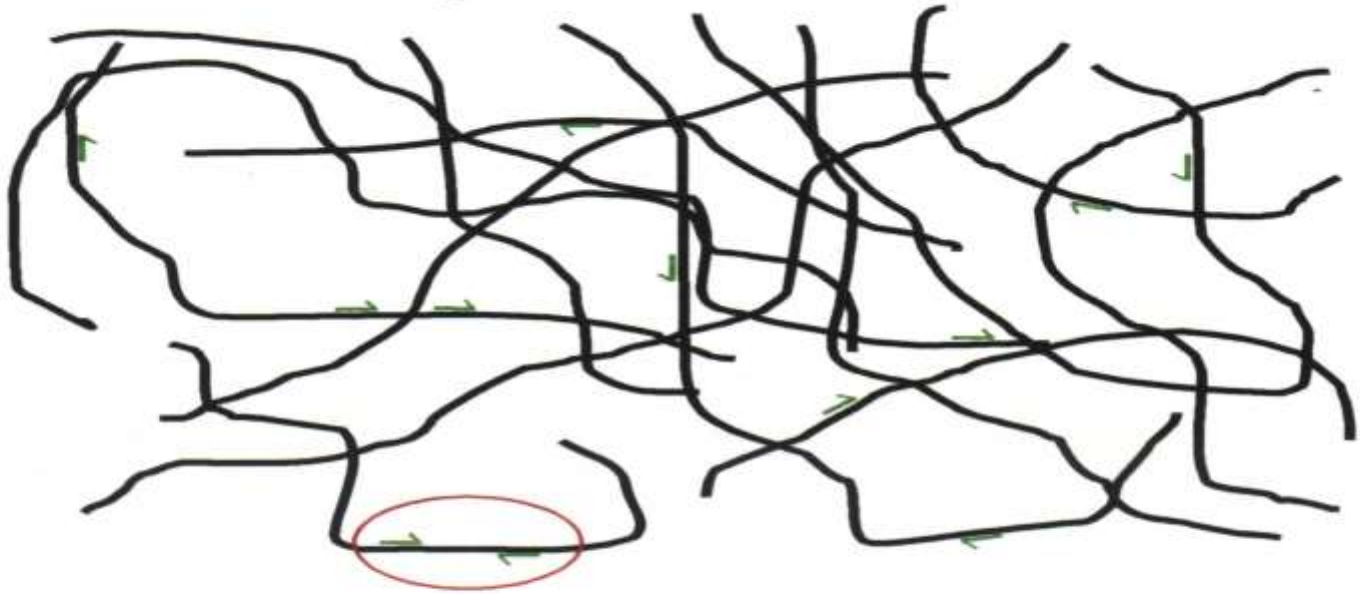
RAPD

- a method based on PCR developed in 1990.
- RAPD is different from conventional PCR as it needs one primer for amplification. The size of primer is normally short (10 nucleotides), and therefore, less specific.
- the primers can be designed without the experimenter having any genetic information for the organism being tested.
- more than 2000 different RAPD primers can be available commercially.

RAPD

- Genomic DNA normally has complimentary sequences to RAPD primers at many locations.
- If two of these locations are close to each other ($<3000\text{bp}$), and the sequences are in opposite orientation, the amplification will be established. This amplified region is said as a RAPD locus.
- Normally, a few (3-20) loci can be amplified by one single RAPD primer.

genomic DNA

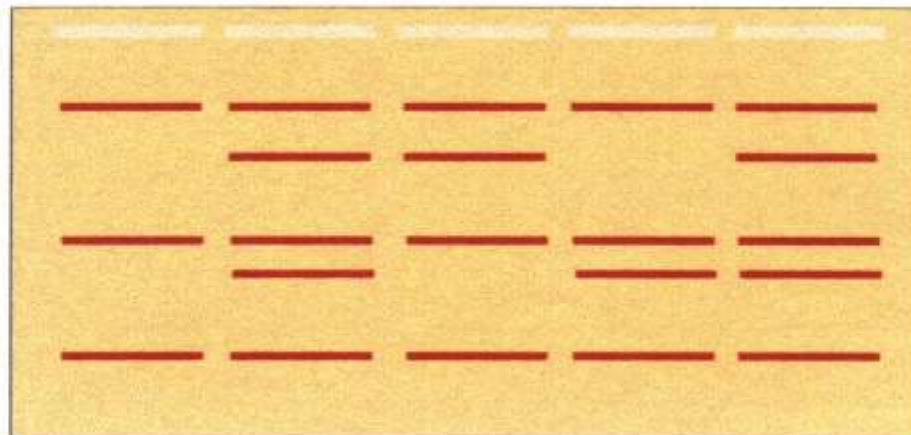


1) PCR

2) Separation by size
on agarose gel



Circled locus →



RAPD

Variation DNA detected by RAPD is due to the loss of RAPD loci. The loss of RAPD loci is caused by:

- a) change of sequence at primer annealing site in the genomic DNA**
- b) deletion of primer annealing site in the genomic DNA**
- c) large insertion in between two primer annealing sites**

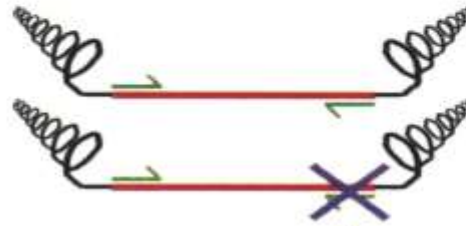
Tree 1

Genotype: + / +



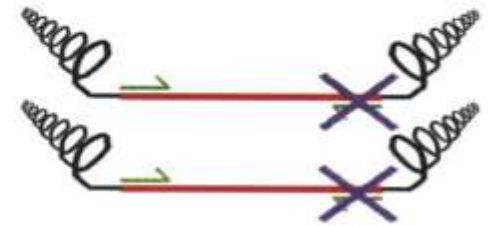
Tree 2

+ / -



Tree 3

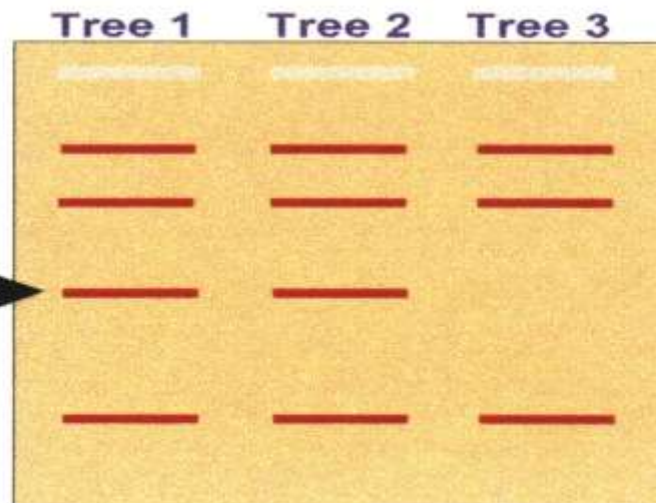
- / -



Primer fails to bind

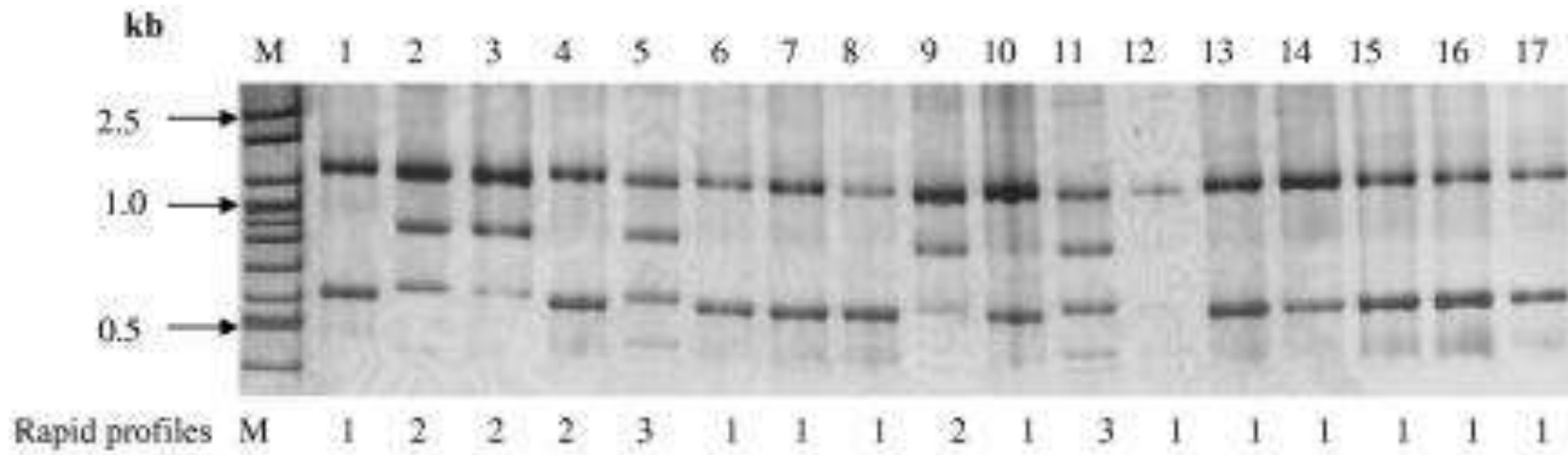
- 1) PCR
- 2) Separation by size on agarose gel

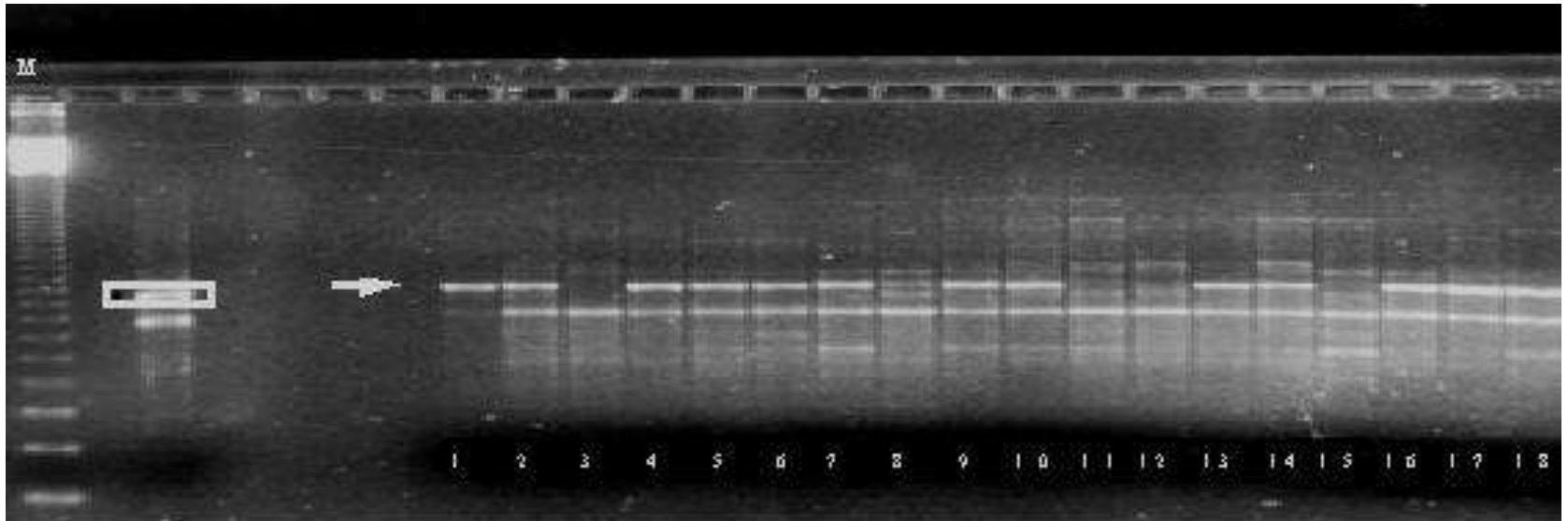
Locus shown above →



RAPD

Silver-stained polyacrylamide gel showing three distinct RAPD profiles generated by primer OPE15 for *Haemophilus ducreyi* isolates from Tanzania, Senegal, Thailand, Europe, and North America





**HOMOLOGY
TEST FOR
FRAGMENTS
OF SIMILAR
MOBILITY IN
RAPD
PROFILES**



RAPD

- **RAPD marker is a dominant marker.**
- **Presence of a DNA band is dominant; absence of a DNA band is recessive.**
- **DNA bands of different sizes are assumed to be amplified products from different RAPD loci.**

Modifications of RAPD

Techniques similar to RAPD:

AP-PCR

DAMD

ISSR

AP-PCR

- **AP-PCR (Arbitrary Primed PCR).**
- **similar to RAPD.**
- **involves two cycles of low-stringency amplification, followed by cycles conducted at higher stringency, using primer of arbitrary sequence.**

AP-PCR

- the length of primers is 20-34 nucleotides long.**
- the primers used include the Universal M13 sequencing primer, the M13 reverse sequencing primer and the T3 sequencing primer.**

DAMD

- **DAMD (Directed Amplification of Minisatellite Region DNA)**
- **technique for detecting polymorphisms using VNTR core sequences as primers for PCR**

ISSR

- ISSR (Inter-Simple Sequence Repeat).
- A PCR-based molecular marker assay of genomic sequence lying between adjacent microsatellites (SSRs). Primers carrying, at their 3'-end, sequence complementary to the repeat unit of the microsatellite will amplify this genomic DNA.

Criticism in RAPD

- lack of reproducibility.
- RAPD banding patterns prone to:
 - i) DNA template concentration and quality
 - ii) Different Taq DNA polymerases
 - iii) Different PCR machines or related equipment used in conducting PCR.

Genetic diversity parameters

- Percentage of polymorphic loci
- Shannon diversity index, H
 - $H = \sum_{i=1}^n -\rho_i \ln \rho_i$
- Genetic similarity, F
 - $F = 2m_{xy} / (m_x + m_y)$
- Genetic distance, $1-F$