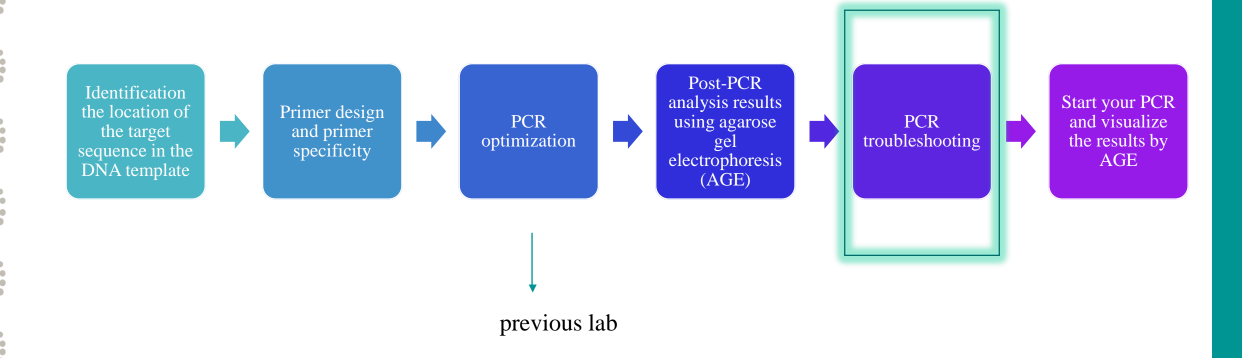
PCR Troubleshooting

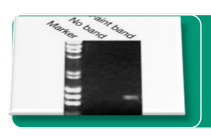
Polymerase Chain Reaction (PCR)=DNA Photocopier



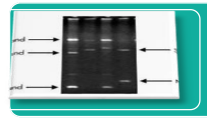
Performing PCR steps:



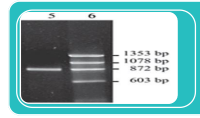
Common Issues In PCR:



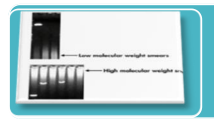
Low or no amplification



Non-specific band or primer dimer



Incorrect product size

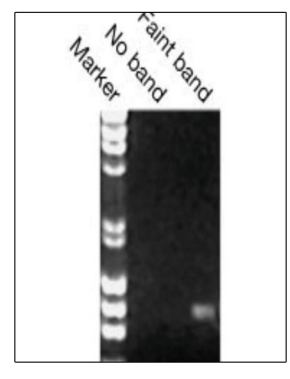


Smeared Bands

1- No Band or Faint Band:

Causes Related to Cycling Times and Temp.

- Too Few cycles were used.
- Extension time was too short.
- Incorrect annealing temperature.
- Denaturation temperature was too low.

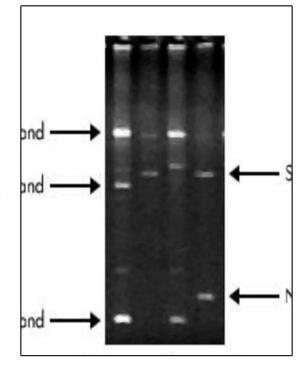


- No enough template was in the reaction.
- Primer concentration was too low.
- Impure primers, dNTPs, or water
- PCR product has high GC content.
- Primers were designed or synthesized incorrectly.
- Not enough Mg^{2+} .

2-Nonspecific Bands or Primer Dimer.

Causes Related to Cycling Times and Temp.

- Annealing temperature was too low.
- Too many cycles were used.
- Extension time was too long.

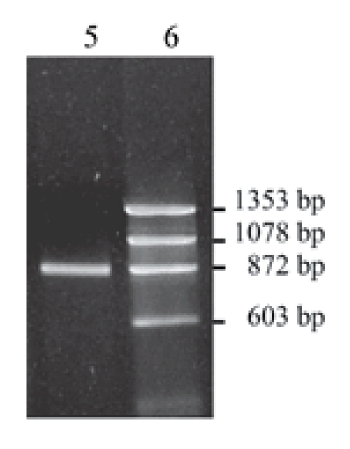


- Too much primer was added.
- Too much Mg2+ was added.
- Primers were designed or synthesized incorrectly by user or manufacturer.

3-Incorrect PGR product size:

Causes Related to Cycling Times and Temp.

• Incorrect annealing temperature.

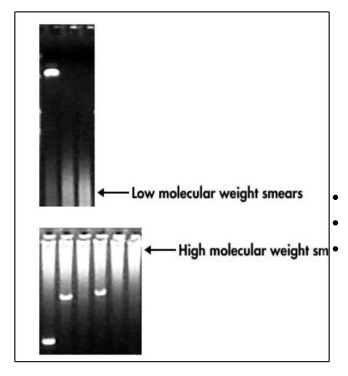


- Mispriming.
- Improper Mg2+ concentration.
- Impure primers, dNTPs, or water
- Primers were designed or synthesized incorrectly by user or manufacturer.

4- Smeared Band:

Causes Related to Cycling Times and Temp.

• Too many cycles were used.



- Too much template was added.
- Impure primers, dNTPs, or water.
- Template contained an exonuclease or was degraded.

Common PCR additive reagents:

1. Additives that benefit GC Rich templates:

→1-10% DMSO (Dimethylsulfoxid):

- GC rich (GC content >60%).
- Lowering the Tm.

→Q solution:

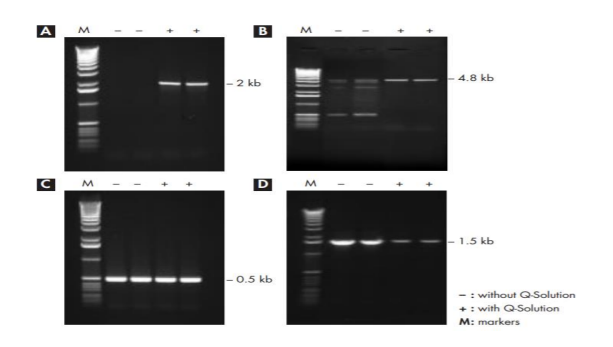
- High degree of secondary structure.
- GC-rich.
- Increases PCR specificity.

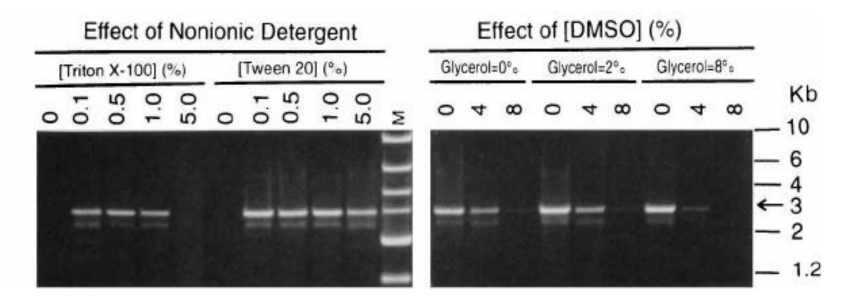
→PCRx Enhancer:

- For problematic and/or GC-rich templates.
- Higher primer specificity, broader magnesium concentration optima, broader annealing temperature optima.

2. Additives That Help PCR in the Presence of Inhibitors:

- →400 ng/µl BSA (Bovine serum albumin).
- **Non-ionic detergents:** Ex: 0.1 to 1% Triton X.





• TERT is a gene code for

• TERT is a gene code for telomerase, an enzyme that elongate the telomere and mutated in some cancers. Your aim is to amplify a region in its promoter using PCR technique.

- 1. Draw a flow chart that illustrate the steps of performing PCR.
- 2. Design a primer using primer3plus tool.
- 3. Check the specificity of the primer.

**