Summer Internship Report

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June 3, 2019

1 Heat Shock

Heat Shock is given to cells for getting the mutant plasmids into them, We keep them in a temperature of -4 Degree Cesius for 15 minutes then move them to 42 Degree Celsius for 35 seconds and the move back to -4.

In this process all the plasmids do not enter the cells, so you incorporate a X-Resistant gene into the plasmid (Here X can be anything, we chose Amphicilin) and now after incubation of about an hour in a rich medium at 220RPM and 37 Degree Celcius. After that colony making is done at 37 Degree Celsius for atleast 12 hrs (we did it for about 18 hrs).

2 Ultracompetent Bacterial Cells

These are the cells which clone plasmids for large number of them. (Basically PCR for large sequences)

PCR has a limit of about 4kb, so for cloning large sequences Ultracompetent Bacterial Cells are used.

These cells are given a heat shock and them left in medium for an hour, later on they are taken for colony making for at least 12 hrs at 37 Degree Celsius.

3 lb Broth Preparation

2.5 gm of lb broth powder per 100 ml of water, the boxes of lb broth and lb agar look surprisingly similar, so watch out for that. 4 mg of lb agar powder per 100 ml of water. If we are making x ml of lb broth solution then the size of the container should be 5 x to 6 x ml as the size increases when bacterial or cells feed on it.

4 Competence

The ability of a cell to alter its own genetics by taking up extracellular DNA from the environment. It is brogth into cell by environmental conditions like

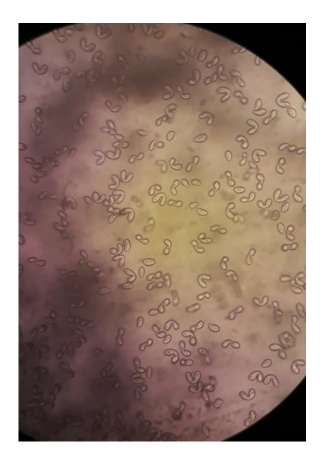


Figure 1: Conjugation

starvation. Condotions inducing sporation often over lap with condition inducing competence.

5 Transformation

It is a process in which the competent cell intakes DNA from outside and ends up changing its own DNA transformation usually produces a mixture of relatively few transformed cells and an abundance of non-transformed cells, a method is necessary to select for the cells that have acquired the plasmid. The plasmid therefore requires a selectable marker such that those cells without the plasmid may be killed or have their growth arrested. Antibiotic resistance is the most commonly used marker for prokaryotes. .

6 Conjugation

At the 8th hour

7 Agarose-Gel

For 100ml of agarose gelcast of 0.8

- 1. Tape the gel caste tray
- 2. Put the comb on the end of the cast tray
- 3. Agar(0.8g) + 1x TAE(100ml) in a flask
- 4. Oven it till dissolution (In this case it took about 5 minutes), check after every minute or so.
- 5. Let the flask cool before putting in ETB, but don't let it cool so much as to start polymerization.
- 6. Put in EtBr(20µl) shake well.
- 7. Put it in the tray
- 8. Move the bubbles away from the well, towards the ends.