

First class: Basic skills in a molecular biology lab

- a) Prepare digest with BamHI or BglII, for the following DNAs: FUGW, genomic DNA, FUCEdW.
- b) Prepare 1% agarose gel
- c) Precipitate genomic DNA
- d) Run digest
- e) Take pictures

Basic Procedure: Liquid Handling 2

- 1000: 100-1000 μl . Use large tip.
- 200: 20-200 μl . Use small tip.
- 20: 2-20 μl . Use small tip.

Basic Procedure: Centrifuge

- Have connector always outside
- Balance tubes.
- Complete stop before opening.

QuickTime™ and a
TIFF (Uncompressed) decompressor
are needed to see this picture.

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Plasmid

- What is plasmid?
- What is the difference of plasmid and “usual” DNA?

Plasmid

- What is plasmid?
 - Small circular DNA.
 - Replicated inside of host bacteria.
 - Can carry extra gene such as drug resistance gene.
 - Can carry foreign DNA.
- What is the difference of plasmid and “usual” or chromosomal DNA?
 - Size
 - Chromosomal DNA carries most (>%99.99) of necessary gene for bacteria
 - Bacteria can live without plasmid DNA but not without chromosomal DNA.

Restriction Enzymes

- The critical component of DNA recombinant technology
- Why it is called “restriction” enzyme?
- How often do you see these “sites” in DNA?

BamHI

QuickTime™ and a
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BglII

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- Why restriction enzyme is useful?

Application of restriction enzyme

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Separation of DNA fragment

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- What is the relationship of migration distance and molecular weight?
 - Include your analysis in the report.

Visualization of DNA in agarose gel

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- Ethidium bromide (EtBr): Possible carcinogen.

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Protocols

Preparation of the digest

Cut 3 μ of each type of DNA with BamHI or BgIII.

For BamHI use buffer 2 (blue)

For BgIII use buffer 3 (red tube)

Cut the following:

FUGW with BamHI

FUGW with BgII FUGW with no enzyme

FUGW with both BamHI and BgIII (in buffer 2)

FUCed with BgIII

FUCed with no enzyme

Genomic DNA with BgIII

(per tube)

3 μ l of DNA

2 μ l of the appropriate buffer (red or blue)

14 μ l of water

1 μ l of BSA (bovine serum albumin)

Why do you think we have BSA?

0.5 μ l of enzyme (BamHI or BgIII)

Mix by gently tapping.

Spin for 3 seconds

Incubate in 37C waterbath for at least 40 minutes (while waiting, start preparing the gel)

Protocols

Preparation of the agarose gel

Measure 100 ml of TAE buffer in measuring syringer

Pour 50 ml of TAE buffer in a 200 ml bottle

Measure 1 gram of agarose in a weigh boat

Add agarose to bottle and swirl thoroughly

Loosen cap and put in microwave for 2 minutes at low power (4)

Make sure that contents do not boil and spill over

Add the rest of TAE to the bottle and mix well by swirling

Add 5 μ l of ethidium bromide to liquid in bottle

Swirl (without making bubbles)

Cast get with two 12 tooth combs (one on top row and one in bottom)

Add agarose into cast and let solidify

Wait 15 minutes (while you wait, precipitate the genomic DNA as explained below)

Once is solid, remove comb, turn the cast 90 degrees, and add approx. 400 ml of TAE buffer, until the liquid starts to cover the gel

Protocols

Precipitation of DNA (this is separate experiment to be done while waiting for digestion and electrophoresis)

Into a 1.5 ml eppendorf tube, add:

40 μ l of genomic DNA

5 μ l of sodium acetate 3 M

100 μ l of 100% ethanol

Mix well by shaking

Spin in centrifuge at maximum speed for 5 minutes

Discard supernatant in waste bucket

Add 1 ml of 70% ethanol

Discard supernatant in waste bucket (make sure that you remove all you can by inverting tube on a piece of paper and gently tap the tube to eliminate all remaining drops)

Air dry tube for 5 minutes

Add 10 μ l of water to bottom of tube

Tap bottom of tube to ensure mixing of water and pellet

Spin down for 3 seconds

Do you see something on the bottom?

Protocols

Running of DNA digests and precipitation

Load 4 μ l of 1Kb DNA ladder in first well

Mix 2 μ l of loading buffer (LB tube) with each of the digests that you prepared before

Load 23 μ l of each of the digest+loading buffer in each well

Add 3 μ l of loading buffer to the precipitated genomic DNA, and load into a well

The loading order should be:

1kb ladder-FUGW (BamHI)- FUGW (BglIII)-FUGW (BamHI and BglIII)-FUGW (no enzyme)- genomic DNA (BglIII) -FUCedW (BglIII)- FUCed (no enzyme)-precipitated genomic DNA (plus empty wells)

Once everything is loaded, set voltage to 150 volts and start running the gel

Run gel for 15 minutes

Stop gel and take image with UV light

Report

- What is the relationship between migration distance and molecular weight?
 - How can you calculate unknown DNA size?
 - How many BamHI and BglII sites are there?
 - How are they arranged?
 - Draw a “restriction enzyme map” of FUWG
 - What is the smeary pattern?
- (No introduction, method, etc needed for today. Just the result of you analysis and a map of the plasmid. Write about possibilities.)