Biochem 717 Gene Expression

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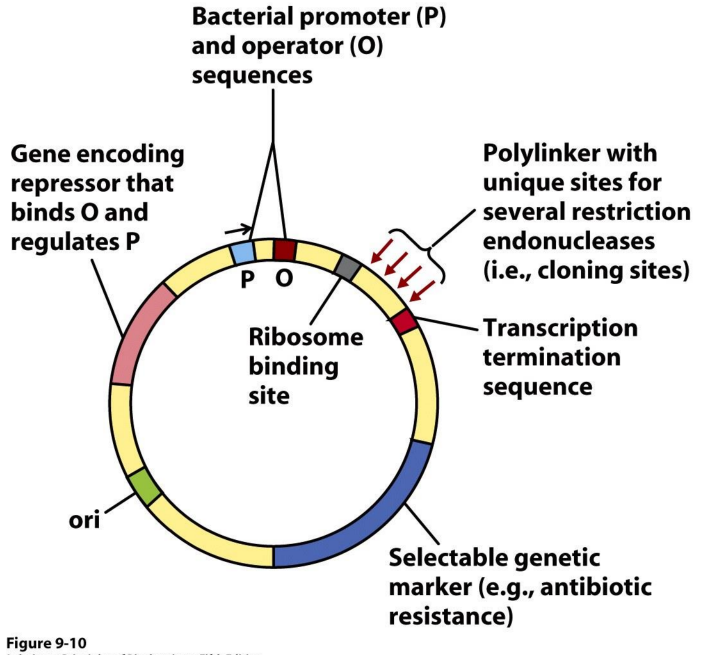
4.3 Methods of Expressing Cloned Genes

Cloning a gene permits

- Production of large quantities of a particular DNA sequence for detailed study
- Large quantities of the gene's product can also be obtained for further use
 - Study
 - Commerce

Expression Vectors

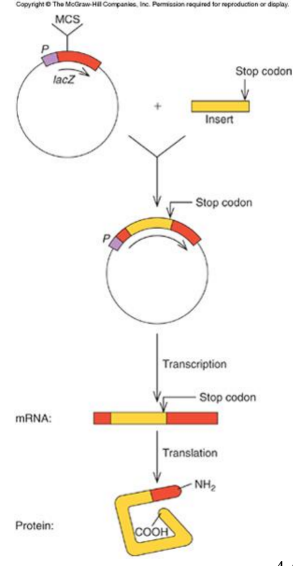
- Vectors discussed so far are used to first put a foreign DNA into a bacterium to replicate and screen
- Expression vectors are those that can yield protein products of the cloned genes
 - Bacterial expression vectors typically have two elements required for active gene expression; a strong promoter and a ribosome binding site near an initiating codon

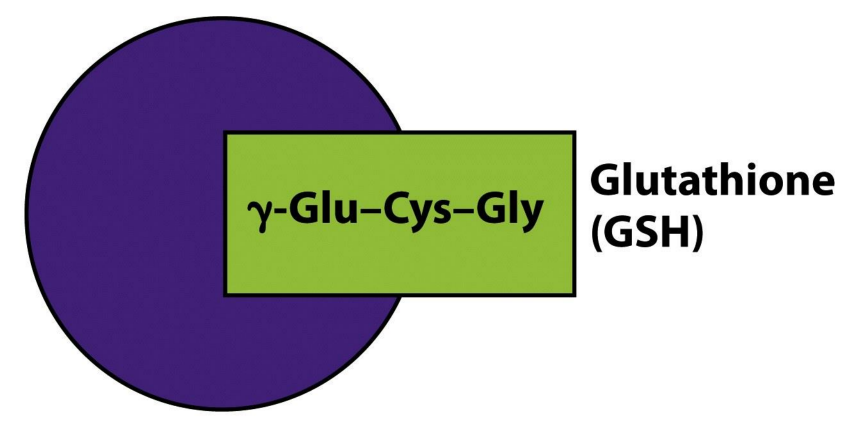


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Fusion Proteins

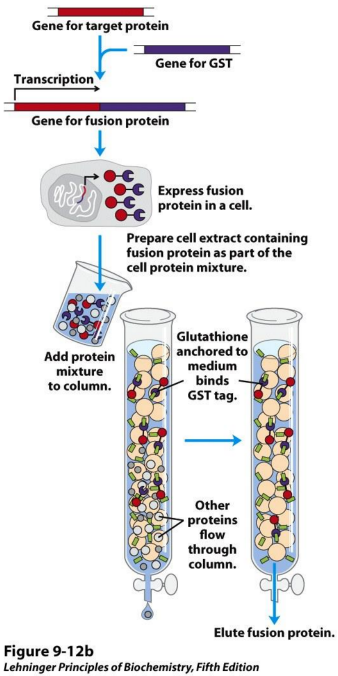
- Some cloning vectors, pUC and pBS, can work as expression vectors using *lac* promoter
- If inserted DNA is in the same reading frame as interrupted gene, a fusion protein results
 - These have a partial β-galactosidase sequence at amino end
 - Inserted cDNA protein sequence at carboxyl end



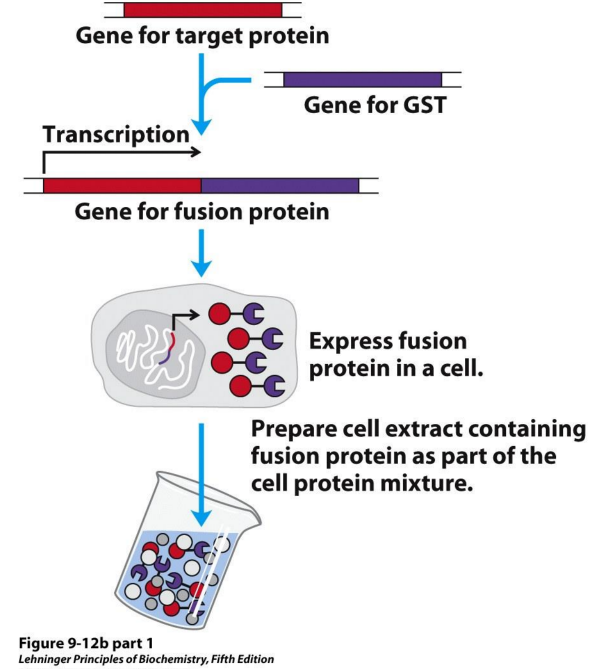


Glutathione-S-transferase (GST)

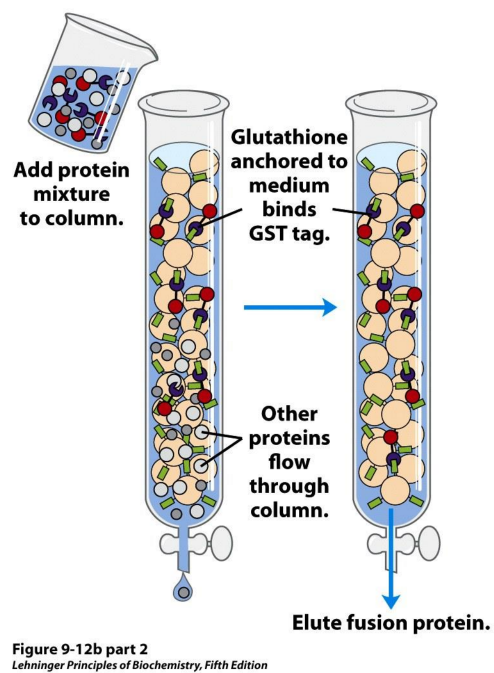
Figure 9-12a *Lehninger Principles of Biochemistry, Fifth Edition* © 2008 W. H. Freeman and Company



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Inducible Expression Vectors

- Main function of expression vector is to yield the product of a gene usually more is better
- For this reason, expression vectors have very strong promoters
- It is usually advantageous to keep a cloned gene repressed until time to express
 - Large quantities of eukaryotic protein in bacteria are usually toxic
 - Can accumulate to levels that interfere with bacterial growth
 - Expressed protein may form insoluble aggregates, called inclusion bodies

Controlling the lac Promoter

- *lac* promoter is somewhat inducible
 - Stays off until stimulated by inducer IPTG
 - However, repression is typically incomplete or leaky and some expression will still occur
- To avoid this problem, use a plasmid or phagemid carrying its own *lacI* repressor gene to keep the cloned gene off until it is induced by IPTG

Alternatives to the *lac* Promoter

- The hybrid *trc* promoter combines the strength of the *trp* (tryptophan operon) promoter with the inducibility of the *lac* promoter
- Promoter from *ara* operon, P_{BAD} , allow fine control of transcription
 - Inducible by arabinose, a sugar
 - Transcription rate varies with arabinose concentration

Alternatives to the *lac* Promoter

- The lambda (λ) phage promoter, P_L , is tightly controlled
- Expression vectors with this promoter-operator system are used in host cells with temperature-sensitive λ repressor gene
 - Repressor functions at low temperatures
 - Raise temperature above the nonpermissive level (42'C) and the repressor doesn't function and the cloned gene is expressed

Expression Vectors That Produce Fusion Proteins

- Most vectors express fusion proteins
 - The actual natural product of the gene isn't made
 - Extra amino acids help in purifying the protein product
- Oligohistidine expression vector has a short sequence just upstream of MCS encoding 6 His
 - Oligohistidine has a high affinity for divalent metal ions like nickel (Ni²⁺)
 - Permits purification by nickel affinity chromatography
 - The his tag can be removed using enzyme enterokinase without damage to the protein product

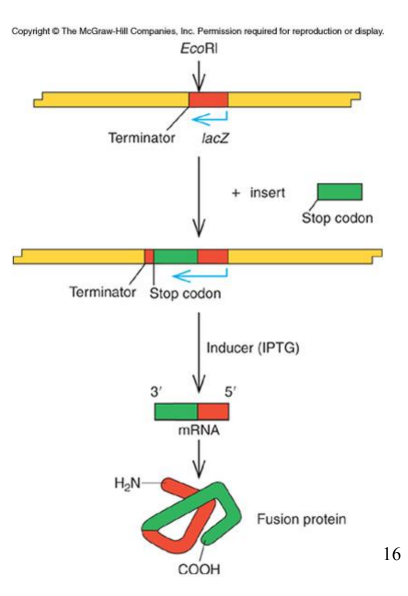
Using an Oligohistidine Expression Vector

(b) (a) PUC ATG (His) EK MCS N 3 2. Lyse cells Histidine or imidazole (1) 4 N 5 Enterokinase 1

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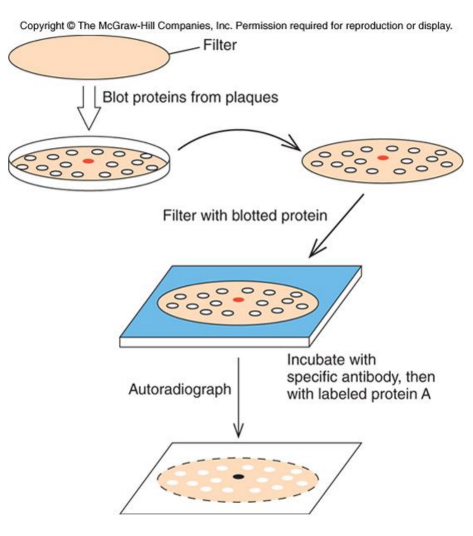
Expression vector $\lambda gt11$

- This phage contains the *lac* control region followed by the *lacZ* gene
- The cloning sites are located within the *lacZ* gene
- Products of gene correctly inserted will be fusion proteins with a βgalactosidase leader



Detecting positive λgt11 clones via antibody screening

- Lambda phages with cDNA inserts are plated
- Protein released are blotted onto a support
- Probe with antibody specific to protein
- Antibody bound to protein from plaque is detected with labeled protein A



Bacterial Expression System Shortcomings

- There are problems with expression of eukaryotic proteins in a bacterial system
 - Bacteria may recognize the proteins as foreign and destroy them
 - Post-translational modifications are different in bacteria
 - Bacterial environment may not permit correct protein folding
- Very high levels of cloned eukaryotic proteins can be expressed in useless, insoluble form

Eukaryotic Expression Systems

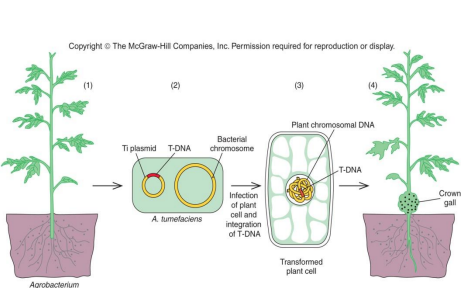
- Avoid bacterial expression problems by expressing the protein in a eukaryotic cell
- Initial cloning done in *E. coli* using a shuttle vector, able to replicate in both bacterial and eukaryotic cells
- Yeast is suited for this purpose
 - Rapid growth and ease of culture
 - A eukaryote with more appropriate post-translational modification
 - Use of the yeast export signal peptide secretes protein into growth medium for easy purification

Use of Baculovirus As Expression Vector

- Viruses in this class have a large circular DNA genome, 130 kb
- Major viral structural protein is made in huge amounts in infected cells
 - The promoter for this protein, polyhedrin, is very active
 - These vectors can produce up to 0.5 g of protein per liter of medium
 - Nonrecombinant viral DNA entering cells does not result in infectious virus as it lacks an essential gene supplied by the vector

Expressing a Gene in a Baculovirus

(b)



(a) tumefaciens

Fig. 4.20a

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Animal Cell Transfection

- Carried out in two ways:
- Calcium phosphate
 - Mix cells with DNA in a phosphate buffer and add a solution of calcium salt to form a precipitate
 - The cells take up the calcium phosphate crystals, which include some DNA
- Liposomes
 - The DNA is mixed with lipid to form liposomes, small vesicles with some of the DNA inside
 - DNA-bearing liposomes fuse with the cell membrane to deliver DNA inside the cell

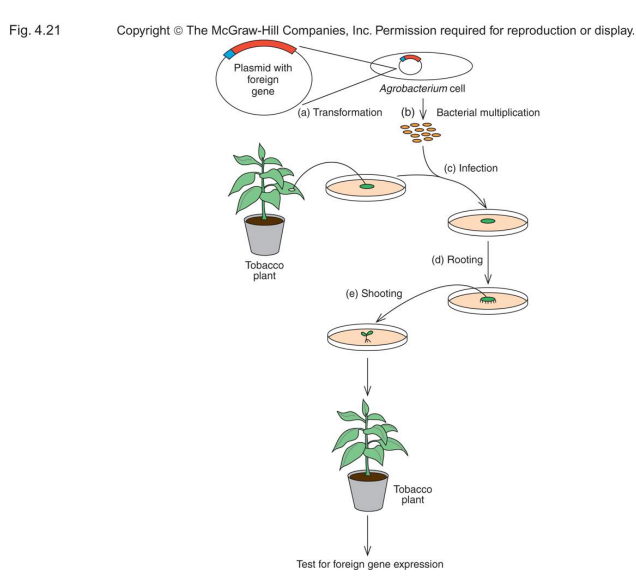
Using the Ti Plasmid to Transfer Genes to Plants

- Genes can be introduced into plants with vectors that can replicate in plant cells
- Common bacterial vector promoters and replication origins are not recognized by plant cells
- Plasmids are used containing T-DNA
 - T-DNA is derived from a plasmid known as tumorinducing (Ti)
 - Ti plasmid comes from bacteria that cause plant tumors called crown galls

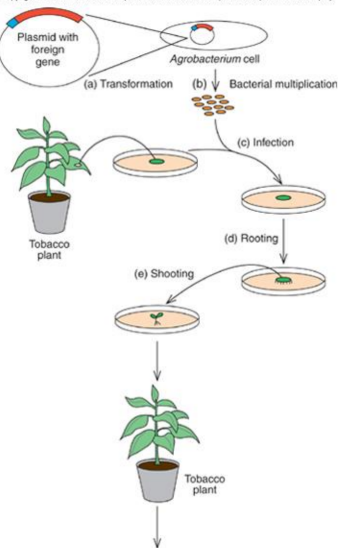
Ti Plasmid Infection

- Bacterium infects plant, transfers Ti plasmid to host cells
- T-DNA integrates into the plant DNA causing abnormal proliferation of plant cells
- T-DNA genes direct the synthesis of unusual organic acids, opines which can serve as an energy source to the infecting bacteria but are useless to the plant

The Ti Plasmid Transfers Crown Gall



Use of the T-DNA Plasmid



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Test for foreign gene expression

Reporter Gene:

A gene with a readily measurable phenotype that can be easily distinguished over a background of endogenous proteins

OR

A gene that researchers attach to a regulatory sequence of another gene of interest in cell culture, animals or plants.

Reporter Assay

- 1. Measures gene expression or transcriptional activity
- 2. Assay of transcription factors.
- 3. DNA promoter assay
- 4. Confirmation of transgenosis

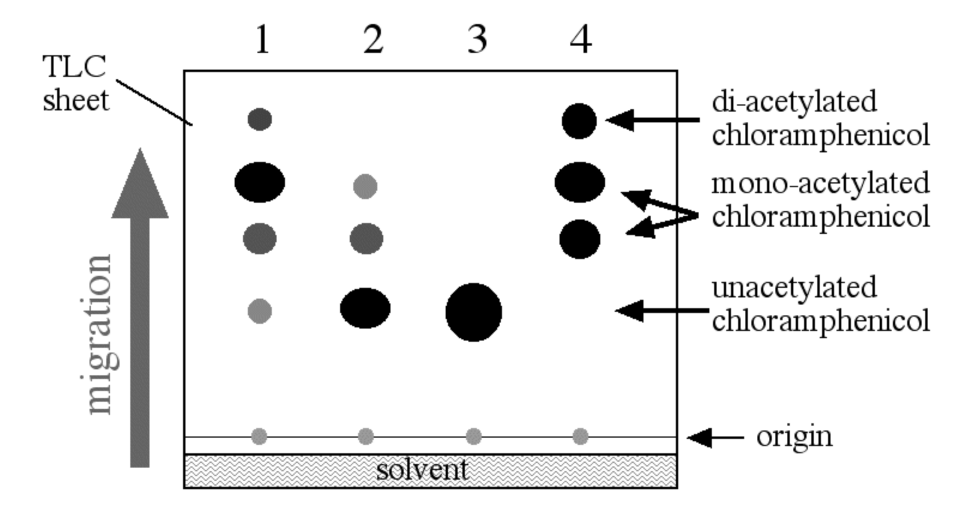
Reporter Assays

- 1. CAT: chloramphenicol acetyltransferase
- 2. β -gal (β -galactosidase):
- 3. GUS Reporter Gene System
- 4. SEAP (secreted alkaline phosphatase):
- 5. Luciferase
- 6. GFP: Green Fluorescent Protein
- 7. Bioluminescence Resonance Energy Transfer (BRET)
- 8. Human Growth Hormone (hGH) Reporter Gene System

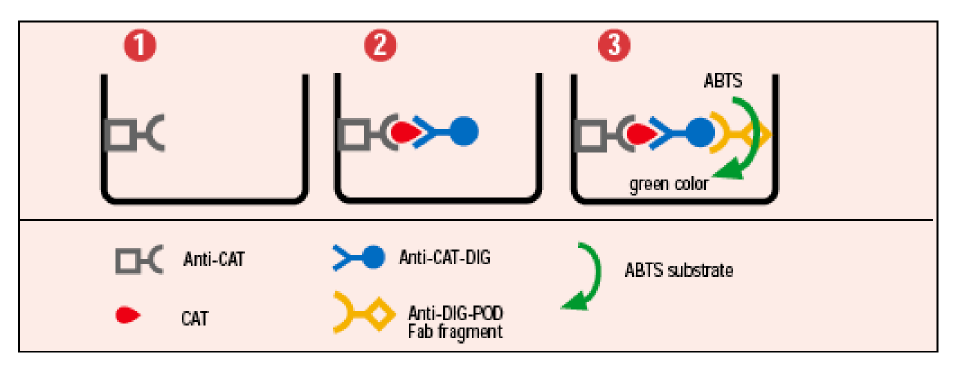
CAT: chloramphenicol acetyltransferase

- 1. 1st reporter gene used to monitor transcriptional activity in cells
- Bacterial enzyme that transfers acetyl groups from acetyl-CoA to chloramphenicol, detoxifying it
- Reaction quantified using radiolabeled substrates (¹⁴C-chloramphenicol) or by ELISA (nonradioactive)

CAT assay: acetylated & non-acetylated chloremphenicol are cheched by TLC

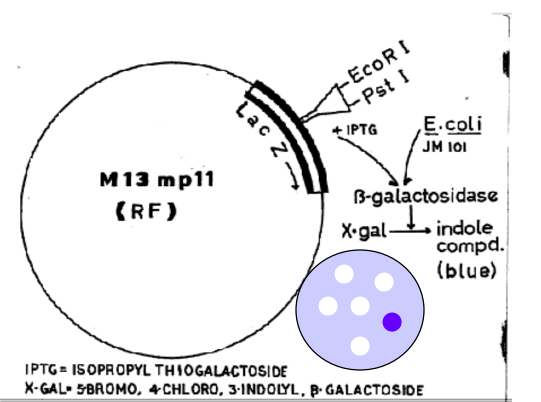


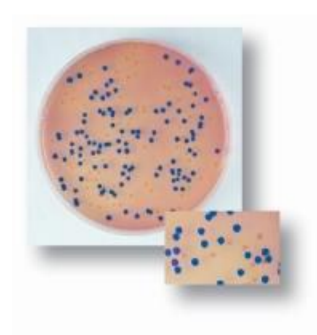
CAT assay: ELISA



β -gal (β -galactosidase):

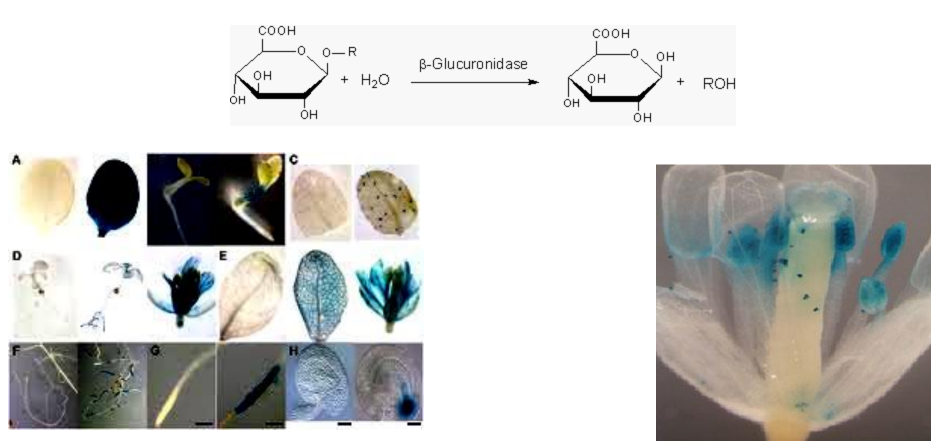
- •*E. coli* enzyme (encoded by *lac*Z) that hydrolyzes galactosidase sugars such as lactose
- Many assay formats: colorimetric, fluorescent, chemiluminescent





GUS Reporter Gene System GUS encodes the beta-glucuronidase enzyme from *E. coli*.

An active enzyme may be detected using X-gal, which forms an intense blue product after cleavage by β -galactosidase

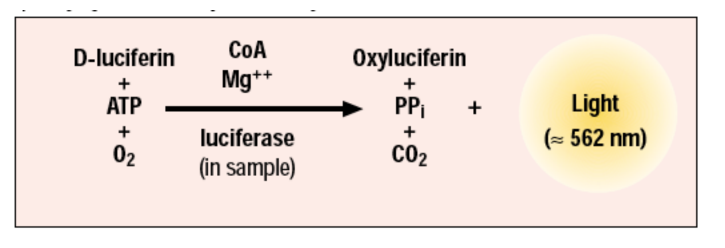


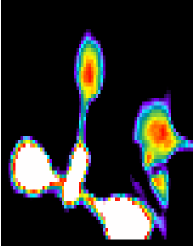
Luciferase:



Renilla reniformis

Firefly (*Photinus pyralis*) luciferase
Sea pansy (*Renilla reniformis*) luciferase
Firefly luciferase produces light by ATP-dependent oxidation



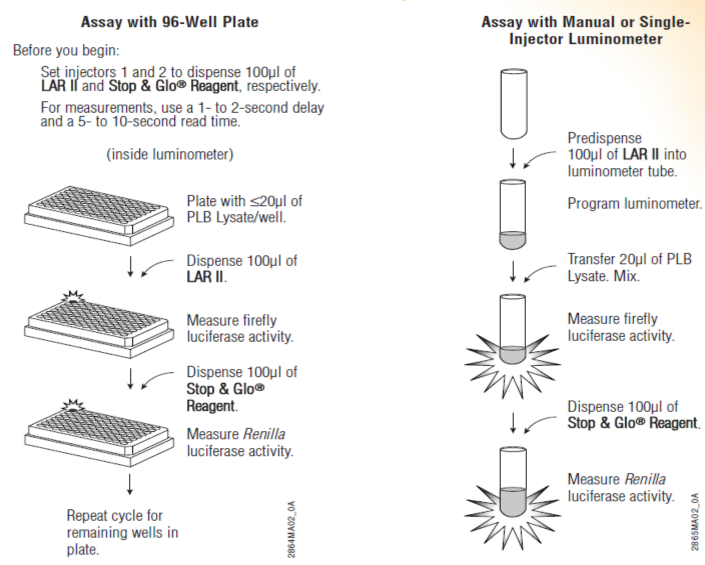


Bioluminescence or light emission is determined by a luminometer



Photinus pyralis

Dual-Luciferase® and Dual-Luciferase® 1000 Assay Protocols

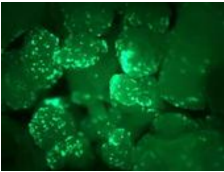


GFP: Green Fluorescent Protein

•Derived from jellyfish *Aequorea victoria*



- Autofluorescent upon UV irradiation (doesn't require cofactors or substrates)
- Retains activity in presence of heat, denaturants, detergents, most proteases
- •Allows for non-invasive monitoring of gene expression in living tissues



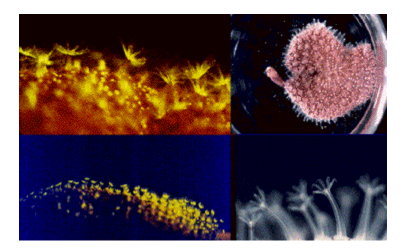
SEAP (secreted alkaline phosphatase):

- •Secreted outside the cell (can assay sample repeatedly and non-destructively by sampling culture medium)
- •This protein is quantified directly by measuring the enzyme activity in the supernatant of the culture medium.
- •Fluorescence and chemiluminescence assays are available for detection.



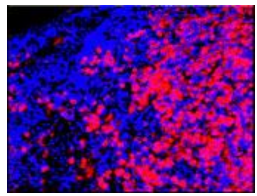
Bioluminescence Resonance Energy Transfer (BRET)

- These systems are based on excitation energy transfer from a bioluminescent donor molecule, e.g. Renilla luciferase to a fluorescent acceptor molecule, like green fluorescent protein (GFP).
- Renilla luciferase as well as GFP are both attached to the proteins to be examined.
- In the presence of coelenterazine, a BRET signal is generated when the fusion proteins are associated.
- Excellent discrimination of the *ruc* and GFP signals results when coelenterazine 400a is used as the *ruc* substrate.



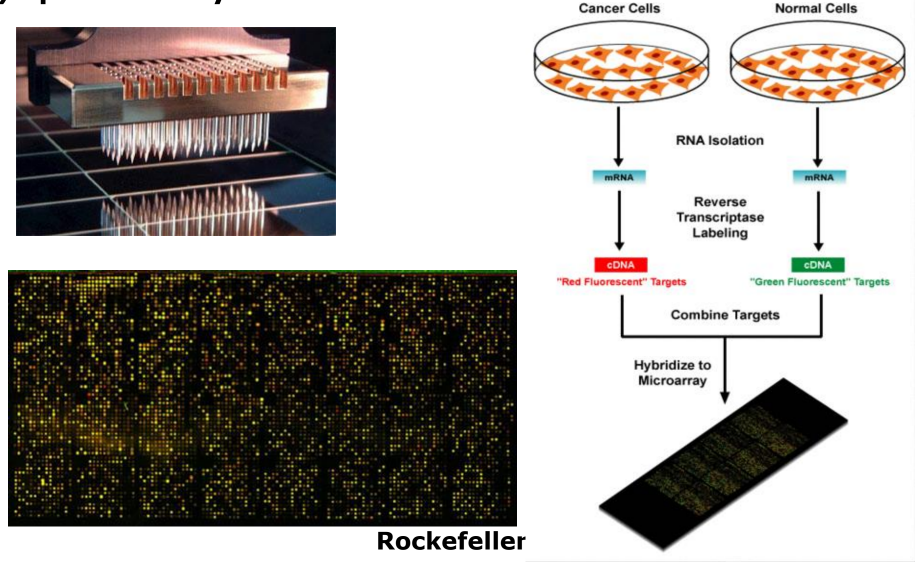
Human Growth Hormone (hGH) Reporter Gene System

- The human growth hormone (hGH) encoded reporter protein is secreted into the culture medium by transfected cells.
- The hGH from the supernatant of the culture medium binds to the antibody on the plate.
- Subsequently, the bound hGH is detected in two steps via a digoxigenincoupled anti-hGH antibody and a peroxidase-coupled anti-digoxigenin antibody.
- Bound peroxidase is quantified by incubation with a peroxidase substrate such as TMB (3,3',5,5'-tetramethylbenzidine)



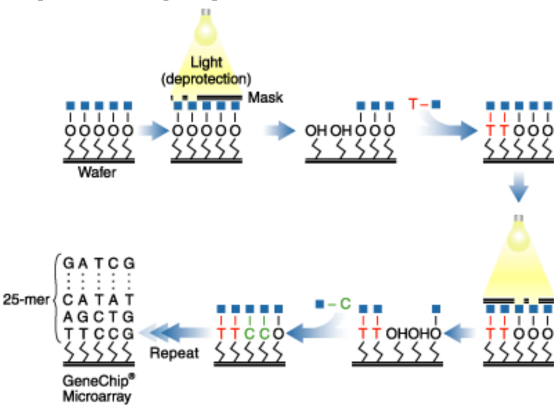
Highly parallel DNA detection I

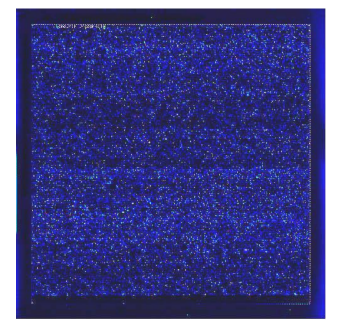
A) Spotted Arrays



Highly parallel DNA detection II

B) On-Chip Synthesis





Affymetrix

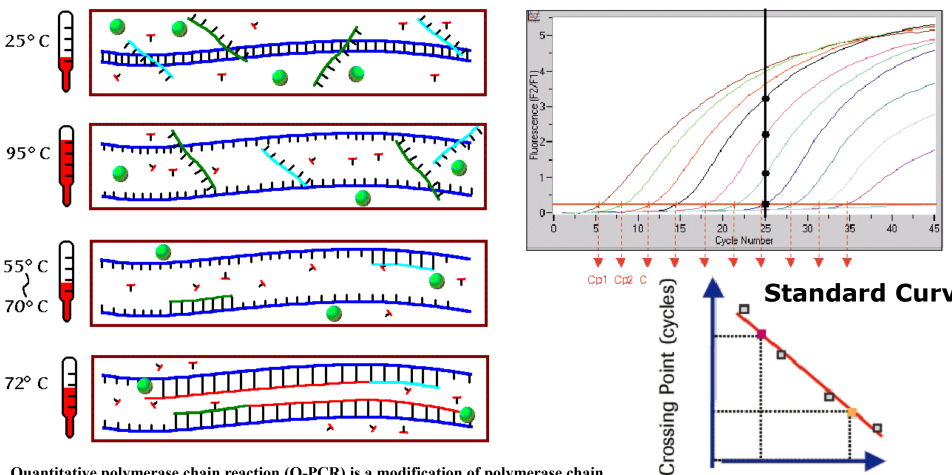


Highly parallel DNA detection III D) Quantitative PCR (qPCR)

Online Detection of Product

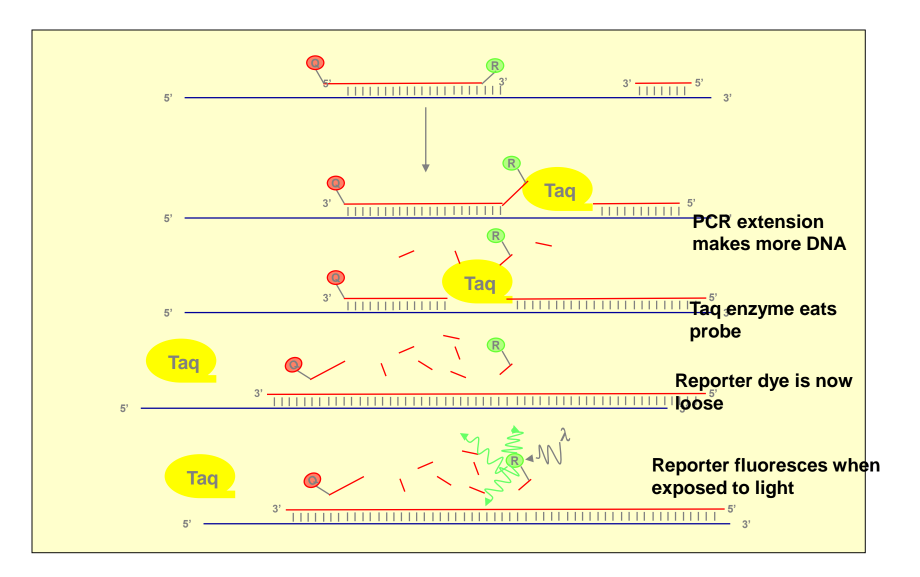
log (copy number)

(c) Roche



Quantitative polymerase chain reaction (Q-PCR) is a modification of polymerase chain reaction used to rapidly measure the quantity of a product of polymerase chain reaction. It is preferably done in real-time, thus is an indirect method for quantitatively measuring starting amounts of DNA, complementary DNA or ribonucleic acid (RNA). This is commonly used for the purpose of determining whether a genetic sequence is present or not, and if it is present the number of copies in the sample. (en.wikipedia.org)

TaqMan Probes in Action!



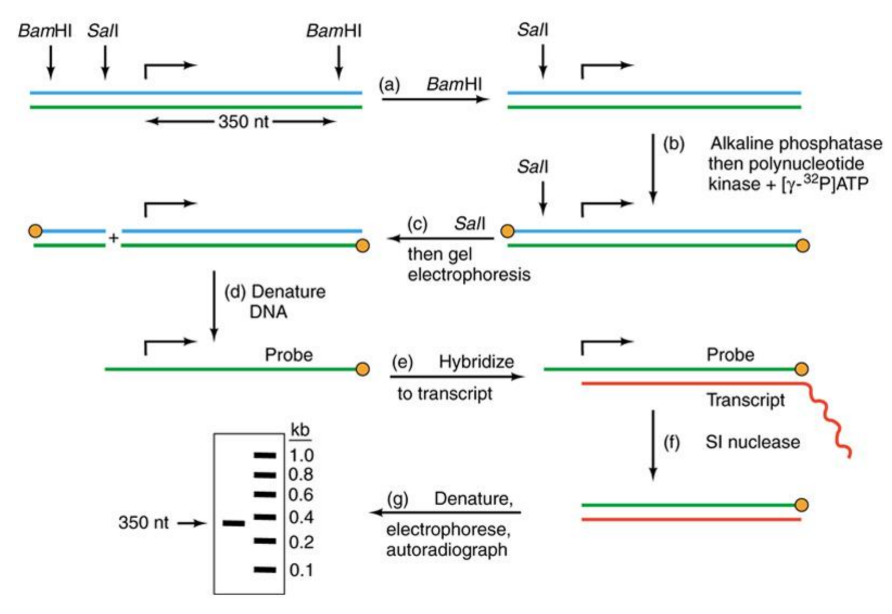
S1 Mapping

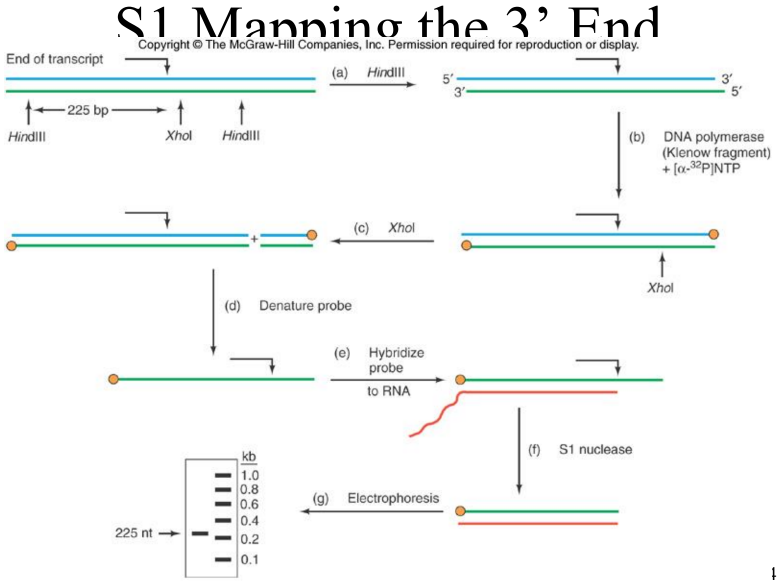
Use S1 mapping to locate the ends of RNAs and to determine the amount of a given RNA in cells at a given time

- Label a ssDNA probe that can only hybridize to transcript of interest
- Probe must span the sequence start to finish
- After hybridization, treat with S1 nuclease which degrades ssDNA and RNA
- Transcript protects part of the probe from degradation
- Size of protected area can be measured by gel electrophoresis

S1 Mapping the 5' End

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Summary

- In S1 mapping, a labeled DNA probe is used to detect 5'- or 3'-end of a transcript
- Hybridization of the probe to the transcript protects a portion of the probe from digestion by S1 nuclease, specific for single-stranded polynucleotides
- Length of the section of probe protected by the transcript locates the end of the transcript relative to the known location of an end of the probe
- Amount of probe protected is proportional to concentration of transcript, so S1 mapping can be quantitative
- RNase mapping uses an RNA probe and RNase

Transgenic Plants



Round Up Ready Soybeans are resistant to herbicide Herbicide Tolerance, Insect Resistance, quality traits Soybean, Corn, Cotton, Canola Tomato

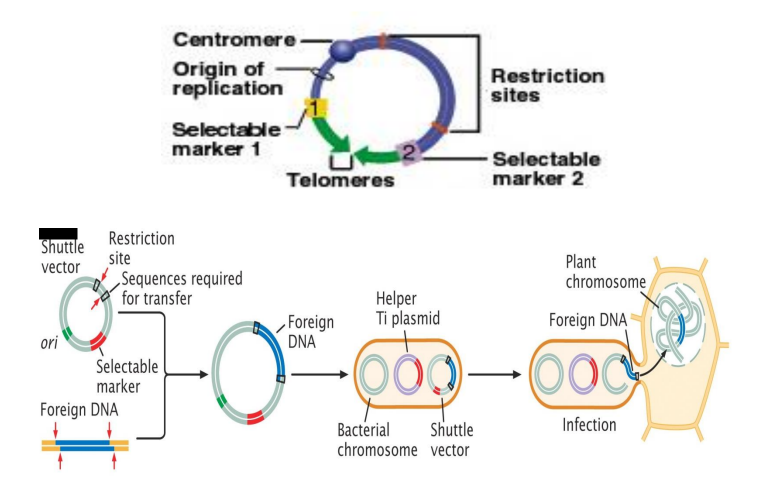


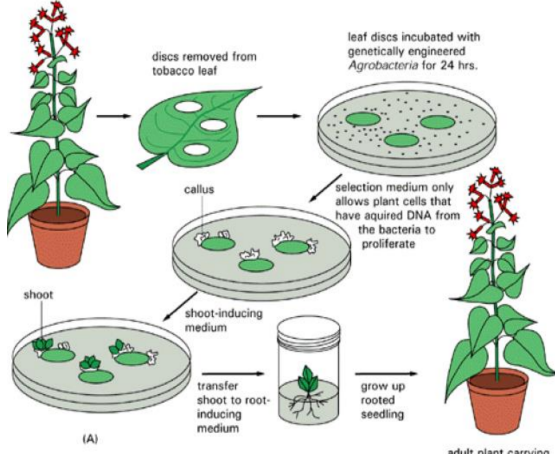
"Golden" rice with beta-carotene and extra iron



Bt Corn produces its own pesticide

Ti plasmid and YAC



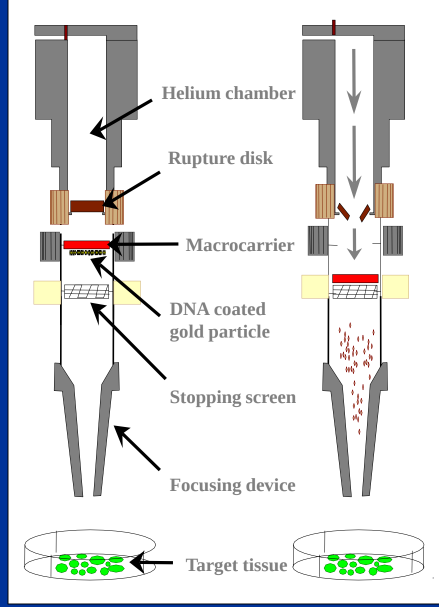


adult plant carrying transgene that was originally present in the bacteria

The Gene Gun

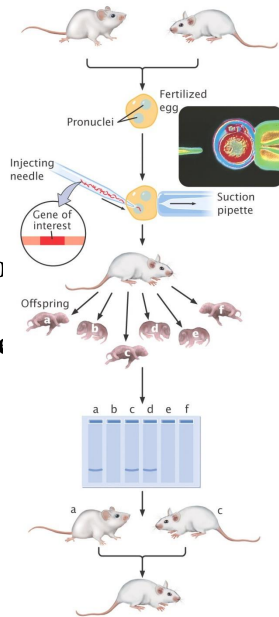


PDS1000 Microparticle Delivery System



Genetically Modified Organisms (Transgenic)

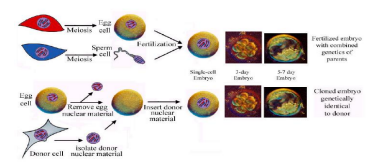
- Introducing or modifying specific genes to alter the phenotype of an organism
- Transgenic: organism that contains a gene from another species in all of its cells
- > Transgenic Animals
 - > Models of Human Disease
 - Produce Pharmaceuticals

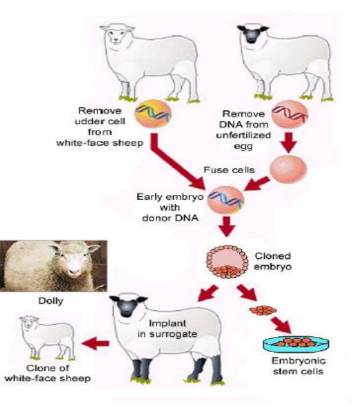


Dolly - 1996



Fertilization vs. Cloning (somatic cell nuclear transfer)





Cloning since Dolly

Cloning of this sort has now been done on cattle, pigs and mice also. The success rate has improved dies 2 Cloning of this sort has now been Guar: Guar: The success rate has improved to the success of the success rate has the success of the succe





