

Module 1 overview

lecture

1. Introduction to the module
2. Rational protein design
3. Fluorescence and sensors

lab

1. Start-up protein eng.
2. Site-directed mutagenesis
3. DNA amplification

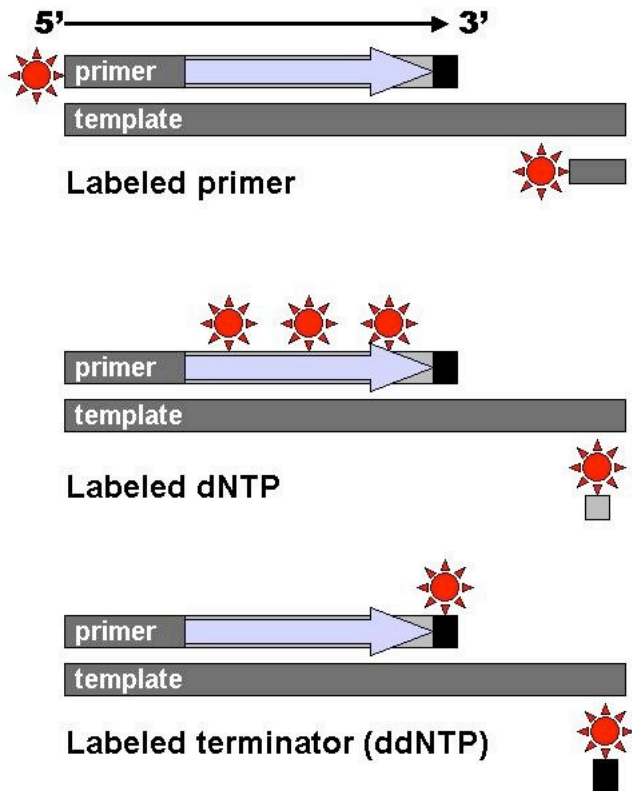
PRESIDENT'S DAY

4. Review & gene analysis
5. Protein expression
6. Purification and protein analysis
7. Binding & affinity measurements
8. High throughput engineering

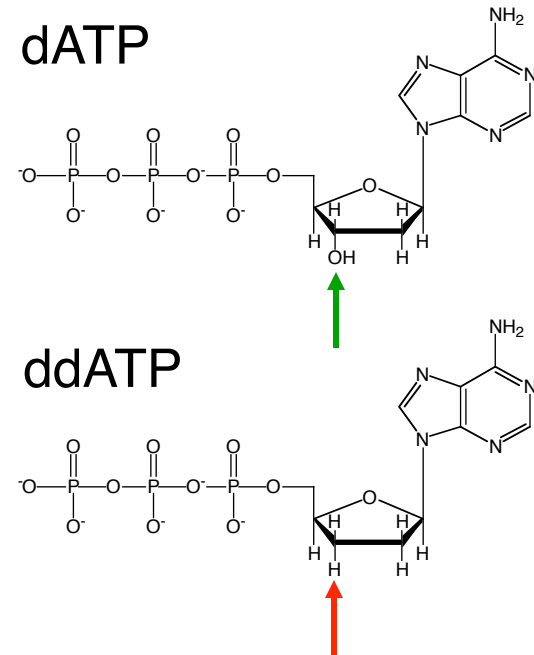
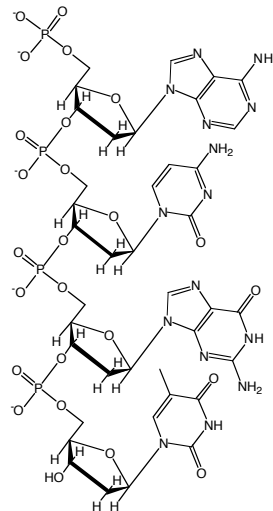
4. Prepare expression system
5. Gene analysis & induction
6. Characterize expression
7. Assay protein behavior
8. Data analysis

How does sequencing work?

Perform PCR on template to be sequences; each PCR reaction is terminated by a nucleotide analog that can be incorporated, but not added to. Terminated PCR products must be labeled in some way.

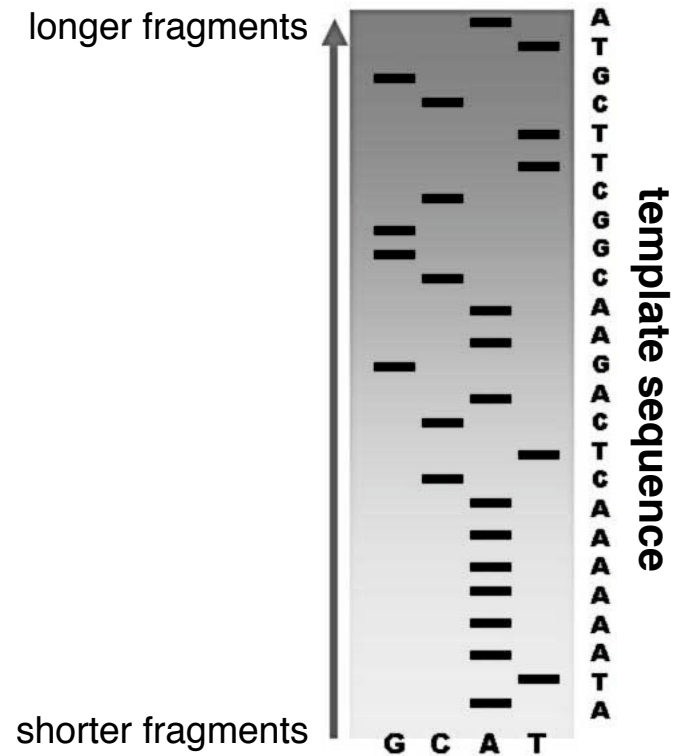
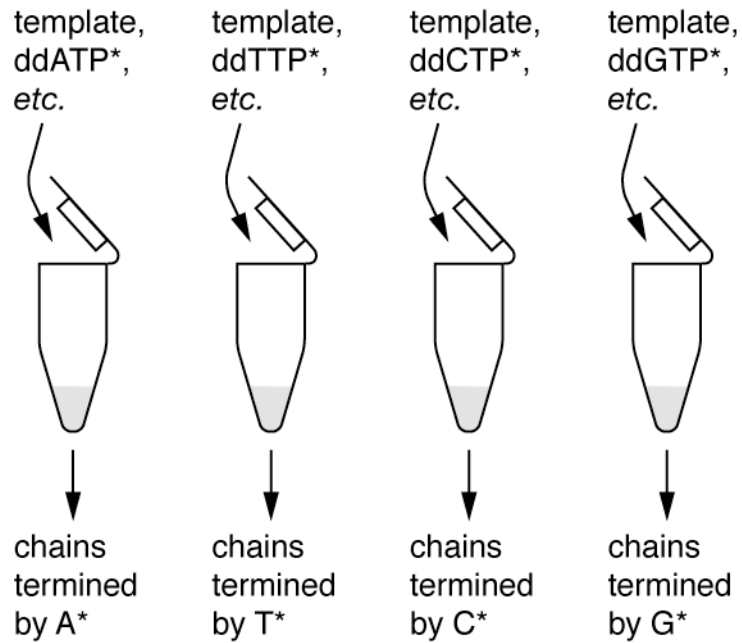


nucleotides linked by phosphodiester bonds



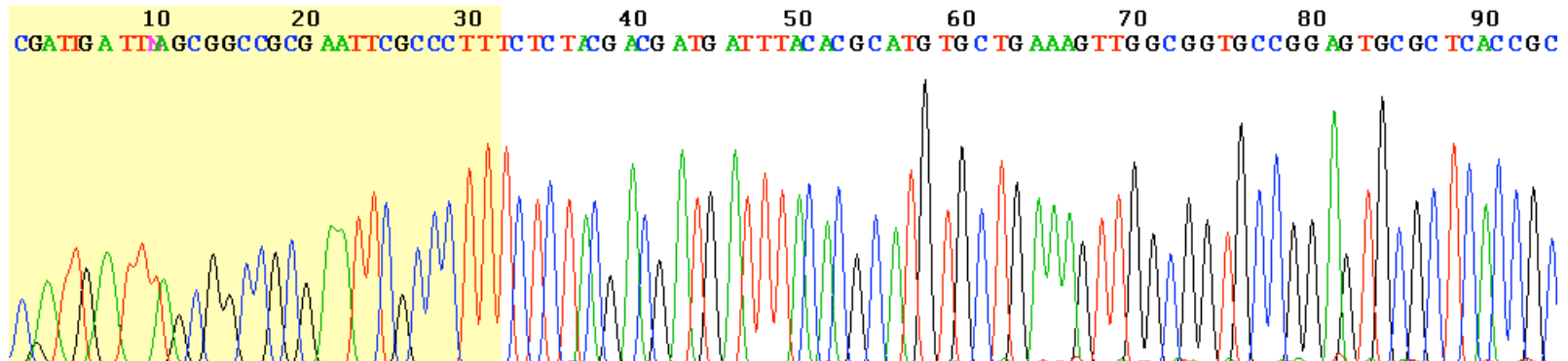
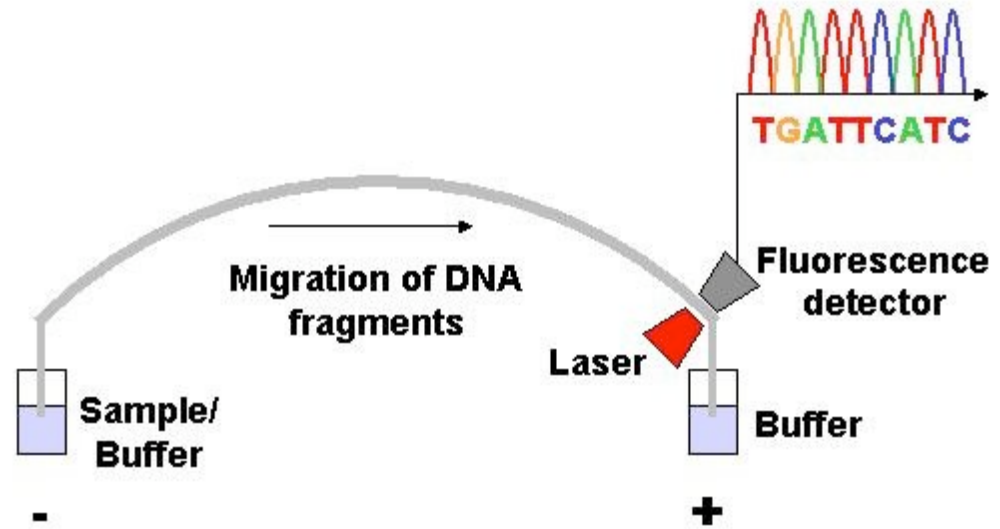
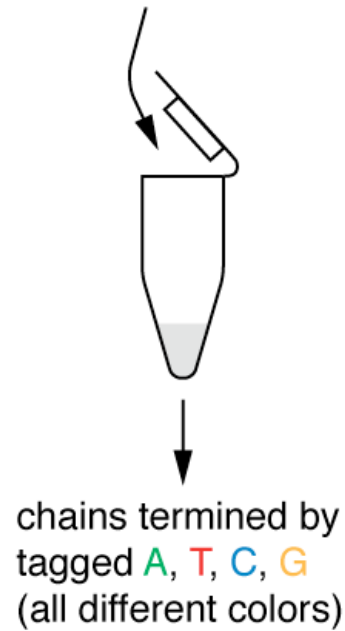
sequencing with radioactive ddNTPs

run products in four separate lanes on gel, expose X-ray film



“one pot” sequencing more common today:

template, fluorescently tagged
ddATP, ddTTP, ddCTP, ddGTP, etc.



Lecture 5: Protein expression & purification

- I. Why express & purify proteins?
 - A. Scientific applications
 - B. Applications in industry, *etc.*

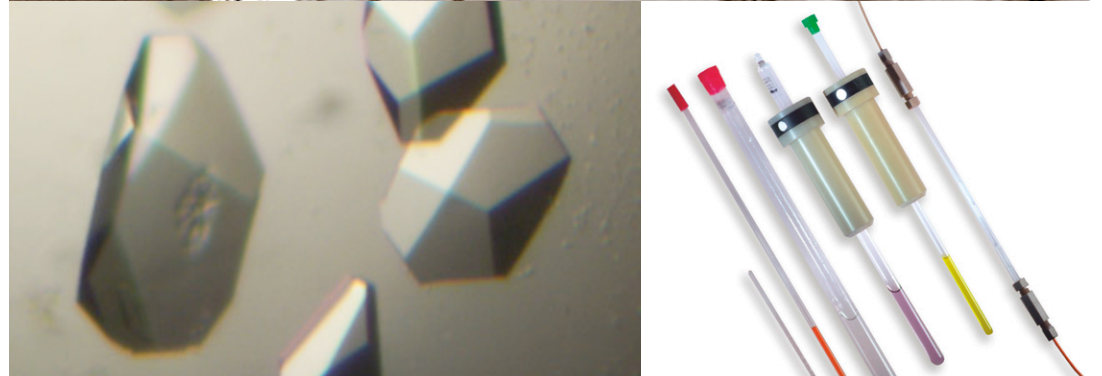
- II. Protein expression systems
 - A. Alternatives to protein expression
 - B. Prokaryotic and eukaryotic systems

Laboratory uses of purified proteins

Biochemistry analysis



Structural biology



Research biochemicals



Protein therapeutics

PEGylated

PEGylated

TNF ligand binding domain + Fc antibody domain

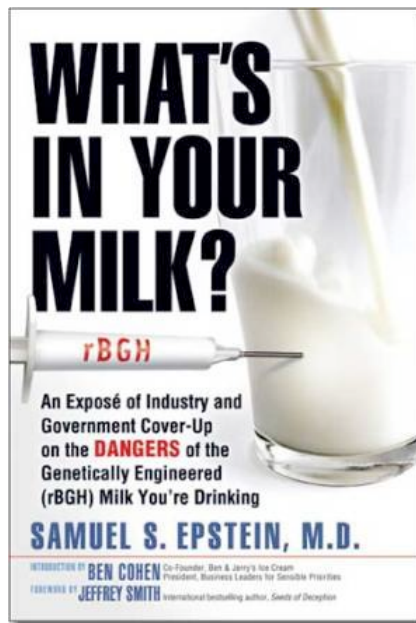
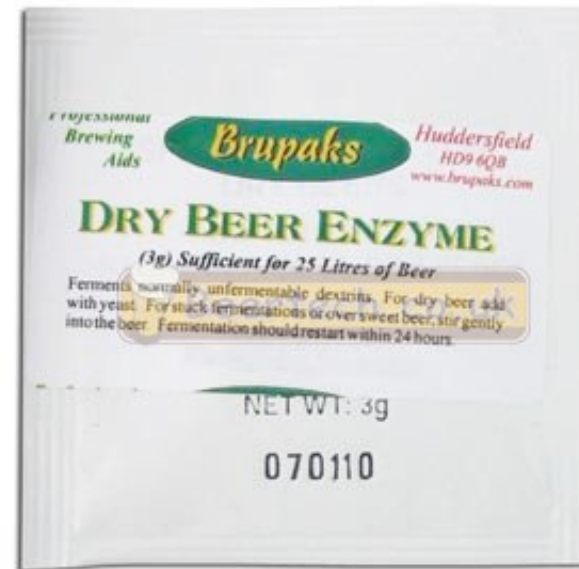
epo engineered to have additional glycosylation sites

Table 1 Top ten recombinant therapeutic proteins and their global sales between 2001 and 2003

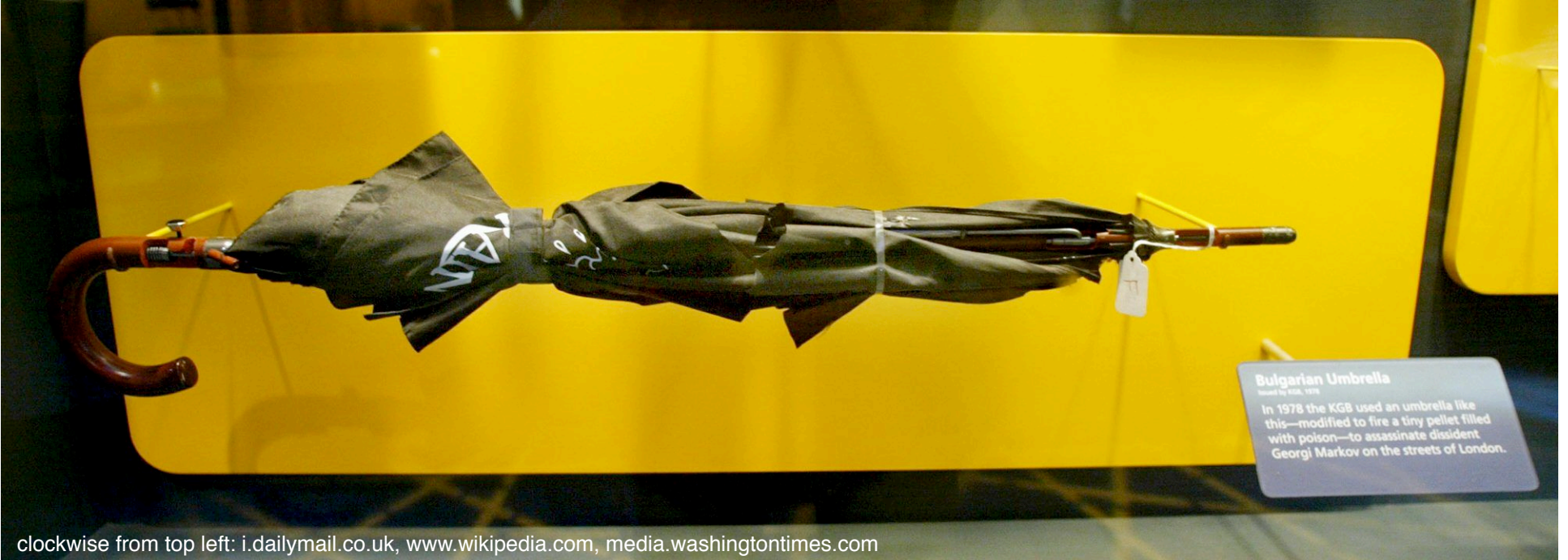
Product (generic)/ marketing company	2001 (\$million)	2002 (\$million)	2003 (\$million)	Growth (decline) 2002– 2003 (%)
Procrit (epoetin alfa)/ Johnson & Johnson	3,430	4,269	3,986	(6.6)
Epogen (epoetin alfa)/ Amgen	2,108	2,261	2,435	7.7
Neupogen (filgrastim)/ Amgen	1,346	1,380	1,268	(8.1)
Neulasta (pegfilgrastim)/ Amgen	0	464	1,255	170.5
Novolin (insulin systemic)/ Novo Nordisk	2,244	2,255	2,235	(0.9)
Avonex (interferon beta-1a)/ Biogen IDEC	971	1,034	1,170	13.2
PEG-Intron A franchise (pegylated interferon alpha)/ Schering Plough	1,447	2,736	1,851	(32.3)
Enbrel (etanercept)/ Amgen	856	521	1,300	149.5
Aranesp (darbepoetin alfa)/ Amgen	42	416	1,544	271.2
NeoRecormon (epoetin-beta)/ Roche	479	766	1,318	72.1
<i>Top ten product sales</i>	<i>12,923</i>	<i>16,102</i>	<i>18,362</i>	<i>14.0</i>
<i>Others</i>	<i>8,547</i>	<i>10,833</i>	<i>13,703</i>	<i>26.5</i>
<i>Total market value</i>	<i>21,470</i>	<i>26,935</i>	<i>32,065</i>	<i>19.0</i>

Source: Datamonitor and company-reported information.

Pavlou & Reichert (2004)
Nat. Biotechnol.



clockwise from top left: s.sears.com, www.beertech.co.uk, www.treatment-skin.com, www.valleynaturals.com, servekrishna.net



Bulgarian Umbrella
In 1978 the KGB used an umbrella like this—modified to fire a tiny pellet filled with poison—to assassinate dissident Georgi Markov on the streets of London.

How can proteins be produced?

1. Purify from natural source

advantages: no chemistry or DNA manipulation required, proteins likely to fold properly, assemble with native cofactors, *etc.*

disadvantages: usually only practical for high abundance proteins, source-specific purification method required

2. Synthesize *de novo*

advantages: no DNA manipulation required, synthesis methods well established, proteins produced are relatively pure

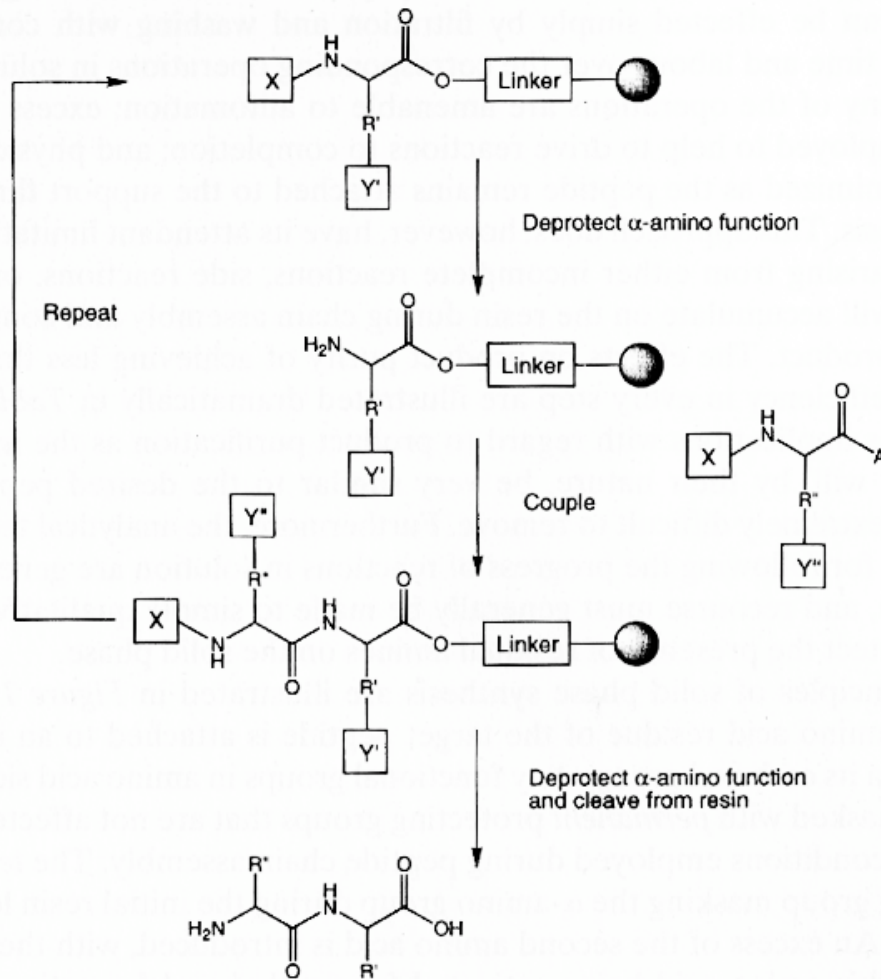
disadvantages: relatively expensive, becomes extremely difficult for polypeptides > 50 amino acids

3. Express and purify from a dedicated expression system

advantages: cheap and frequently high-yield, versatile expression systems already established

disadvantages: cloning required, troubleshooting often needed to obtain high expression and proper folding

Solid phase peptide synthesis is a reliable technique for generating short polypeptides



X = Temporary amino protecting group
 Y = Permanent side-chain protecting group
 A = Carboxy activating group

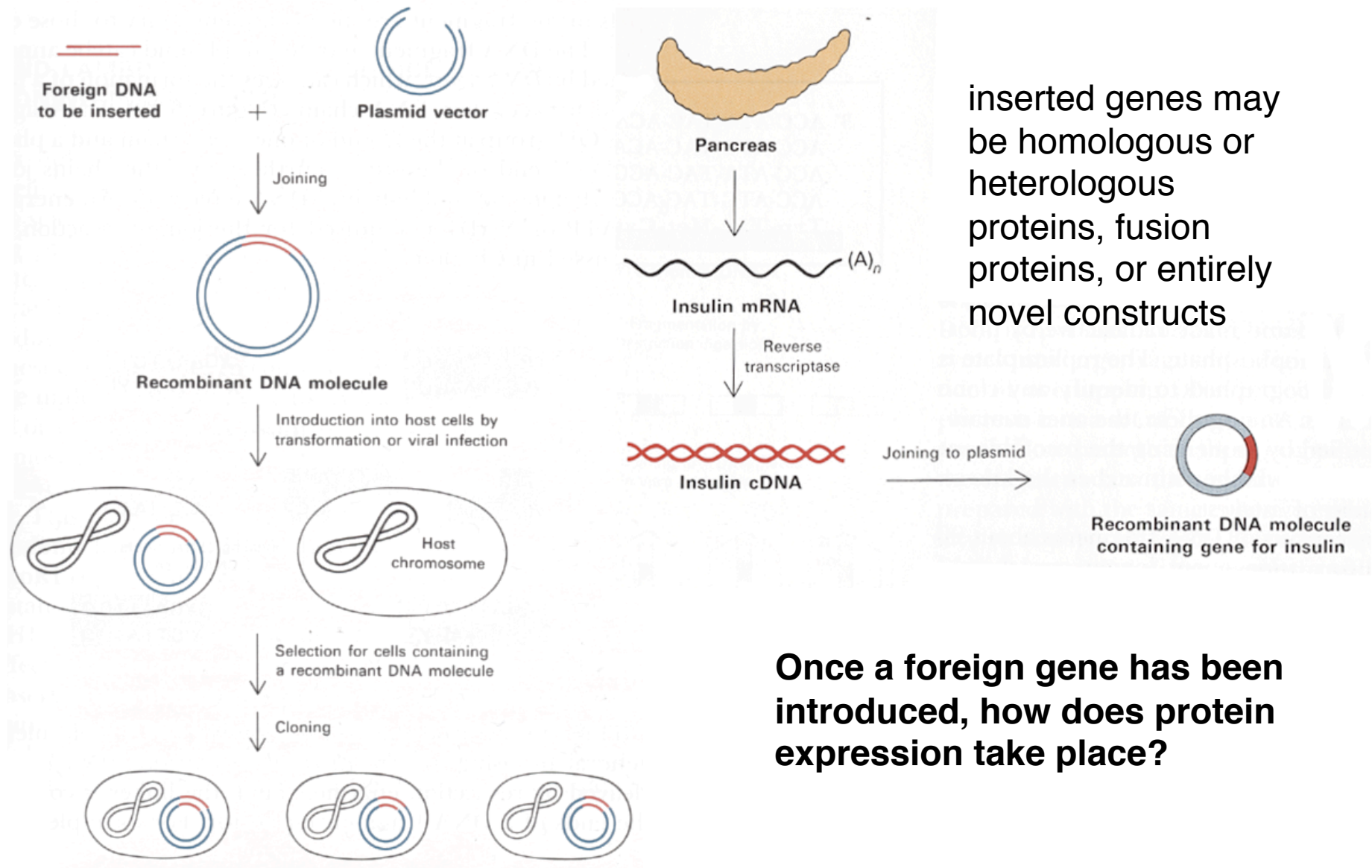


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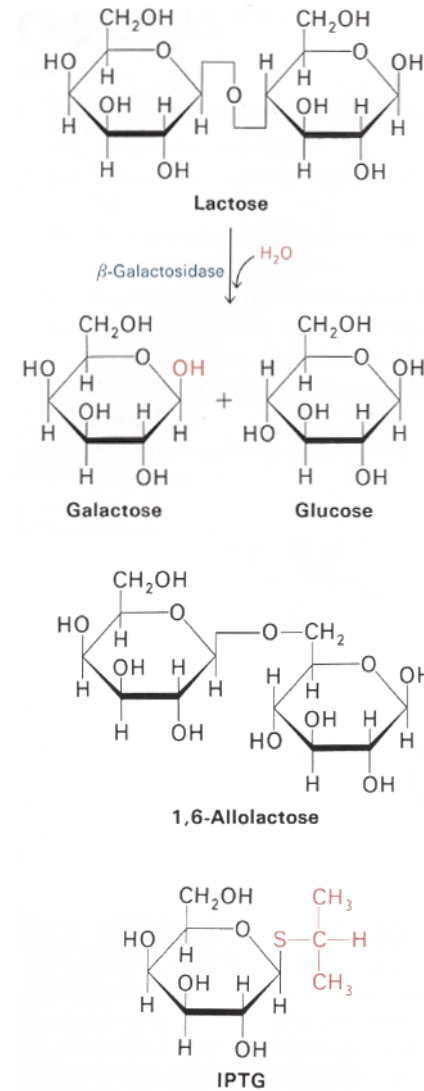
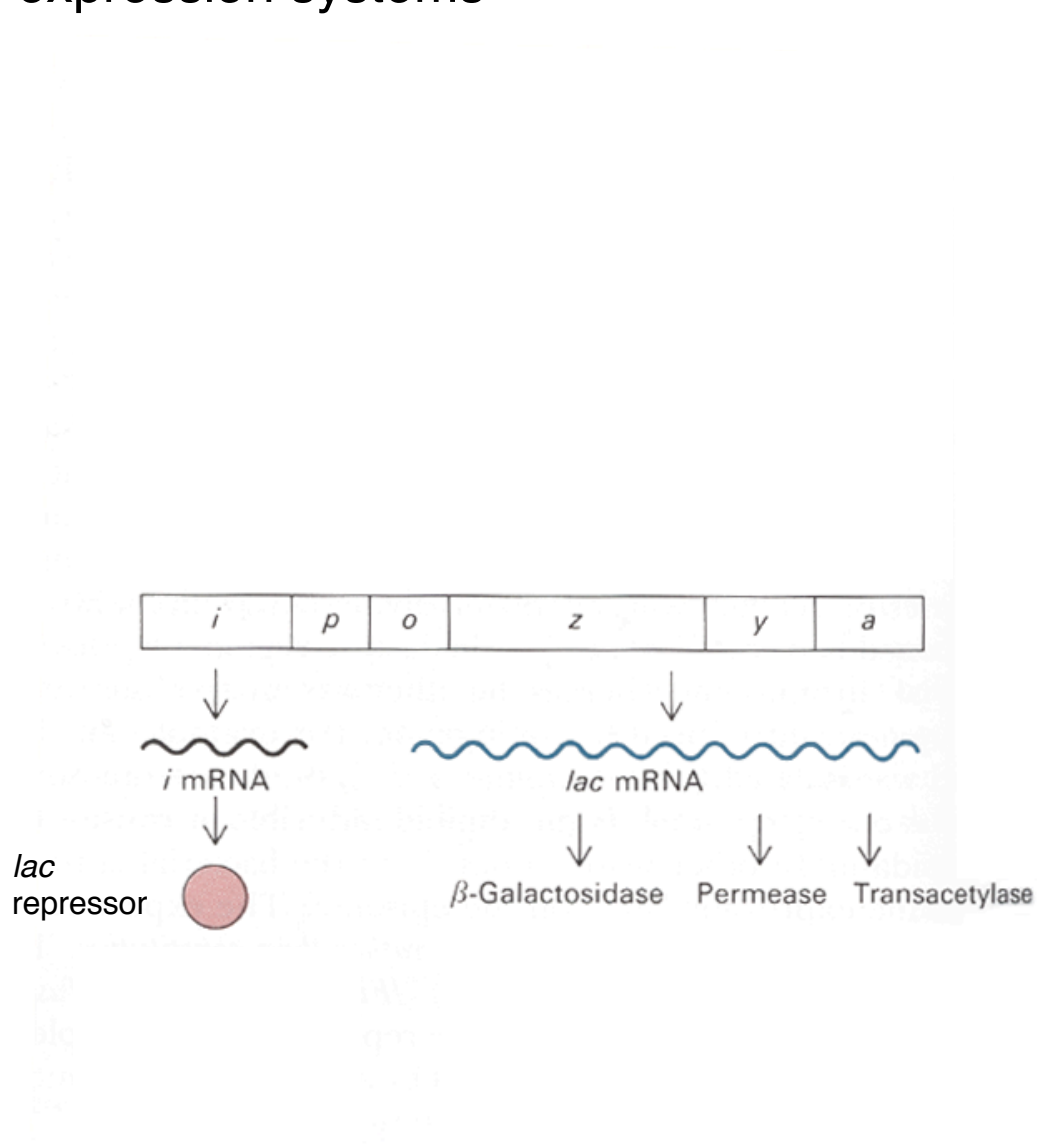
Table 1. Effects of accumulated errors on final product yields

No. of reactions	Overall yields	Yield of each reaction (%)			
		100	99	95	90
10		100	90	60	35
20		100	81	36	12
30		100	74	21	4
40		100	67	13	1
50		100	61	8	< 1

E. coli are the most common host for recombinant gene expression

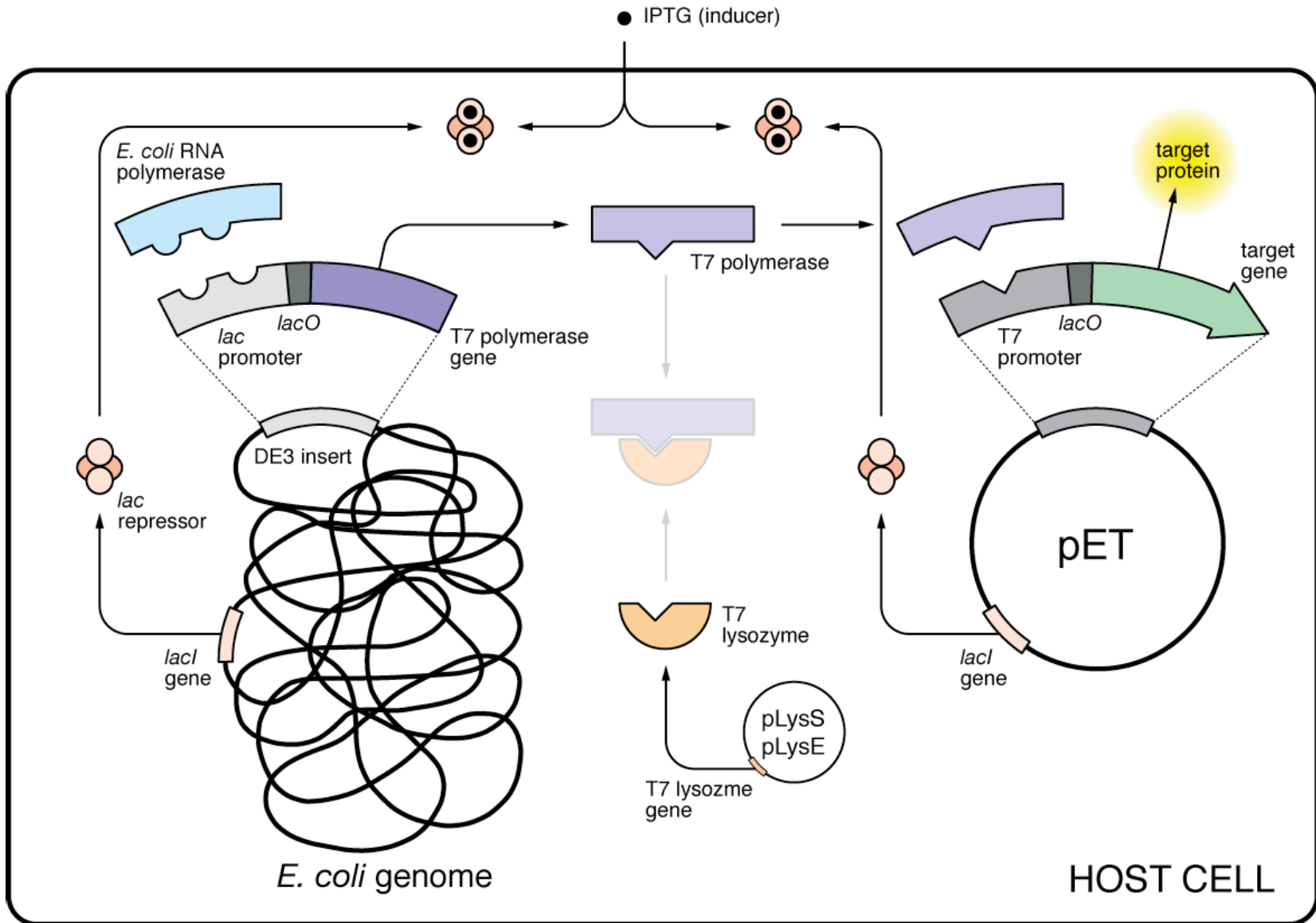


The *lac* operon is the basis for the most common bacterial protein expression systems



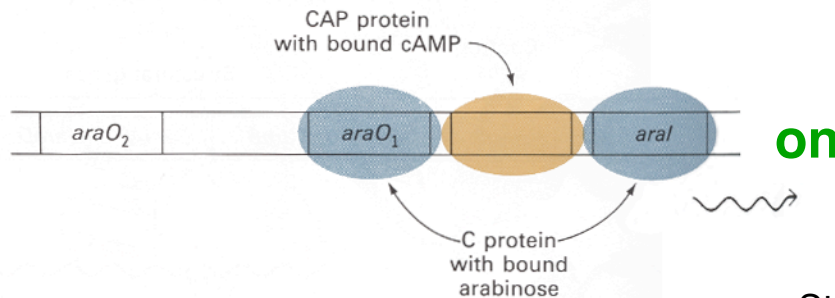
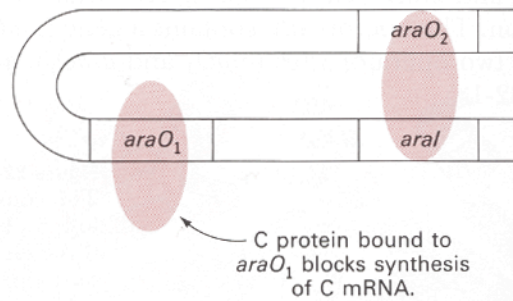
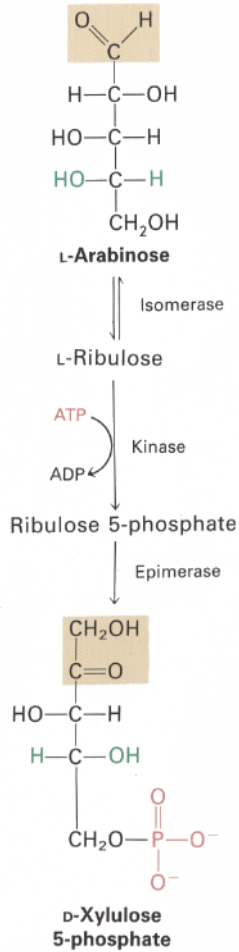
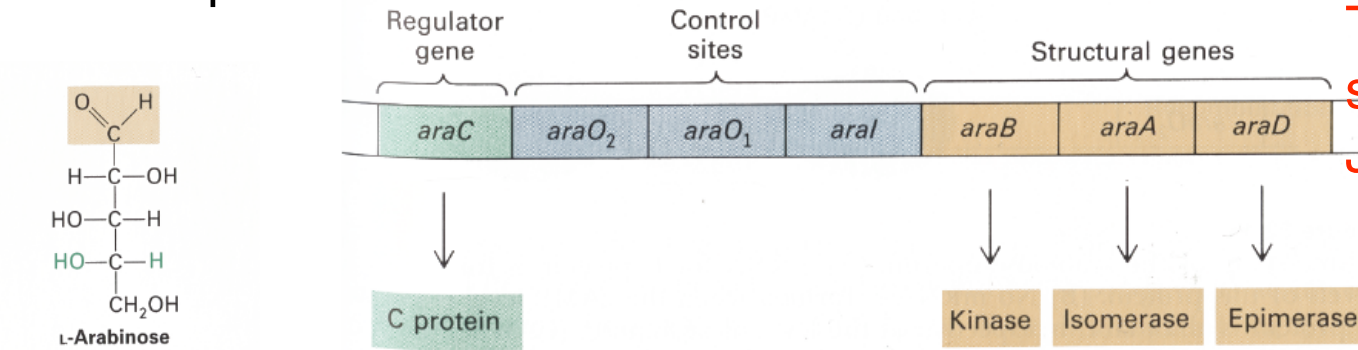
Stryer (1988) *Biochemistry*, 3rd ed.

T7 expression system



Other induction systems can also be used for protein expression in *E. coli*: arabinose system is considered to be more tightly controlled than the *lac* operon

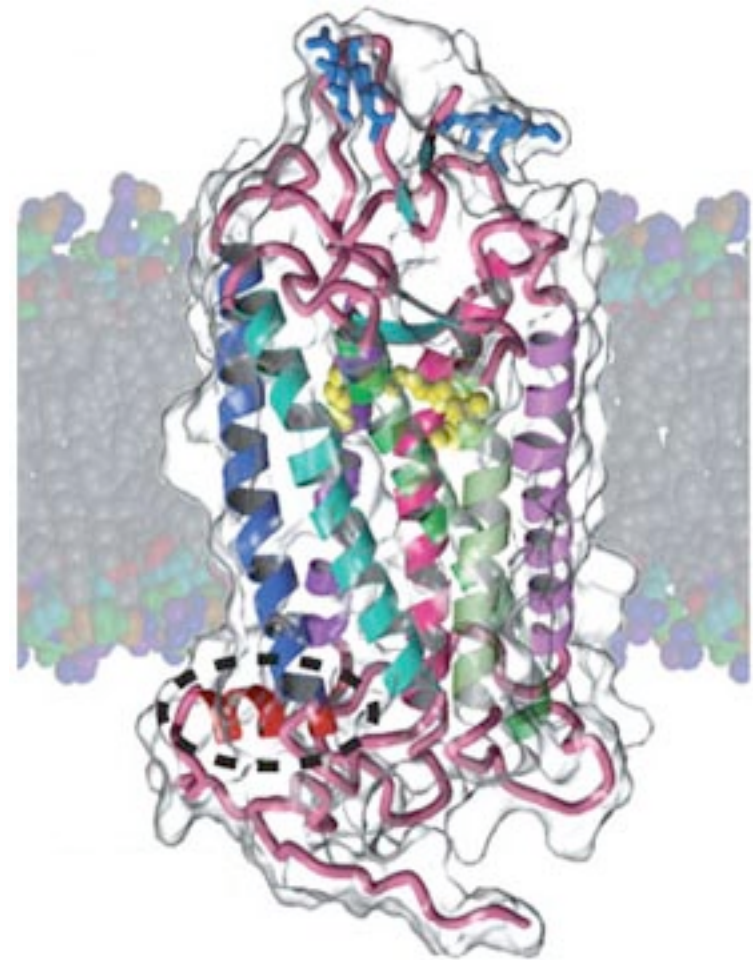
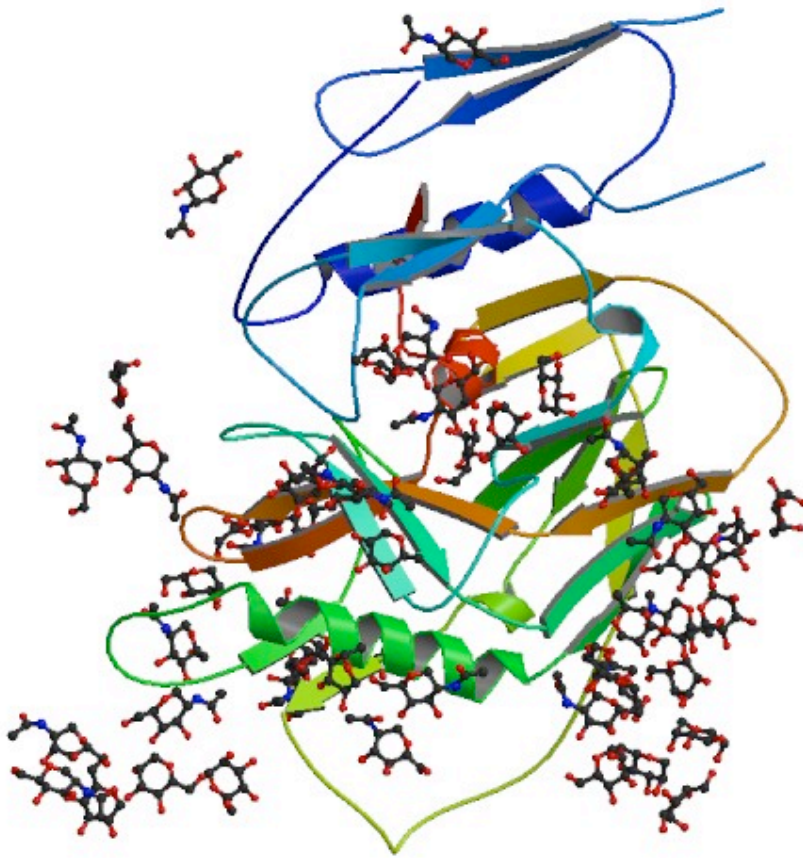
This and following slides not covered, Just reference



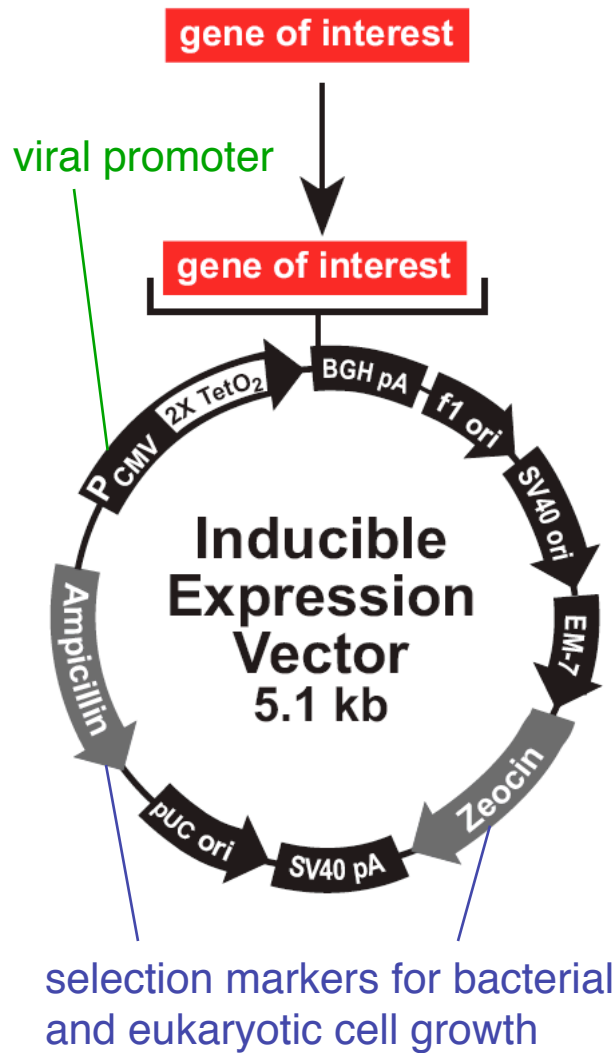
ara system is also compatible with T7-based vectors

Differences between prokaryotic and eukaryotic proteins sometimes require eukaryotic expression systems.

These two proteins exemplify characteristics that frequently call for eukaryotic expression:



Eukaryotic expression vectors share features with bacterial systems

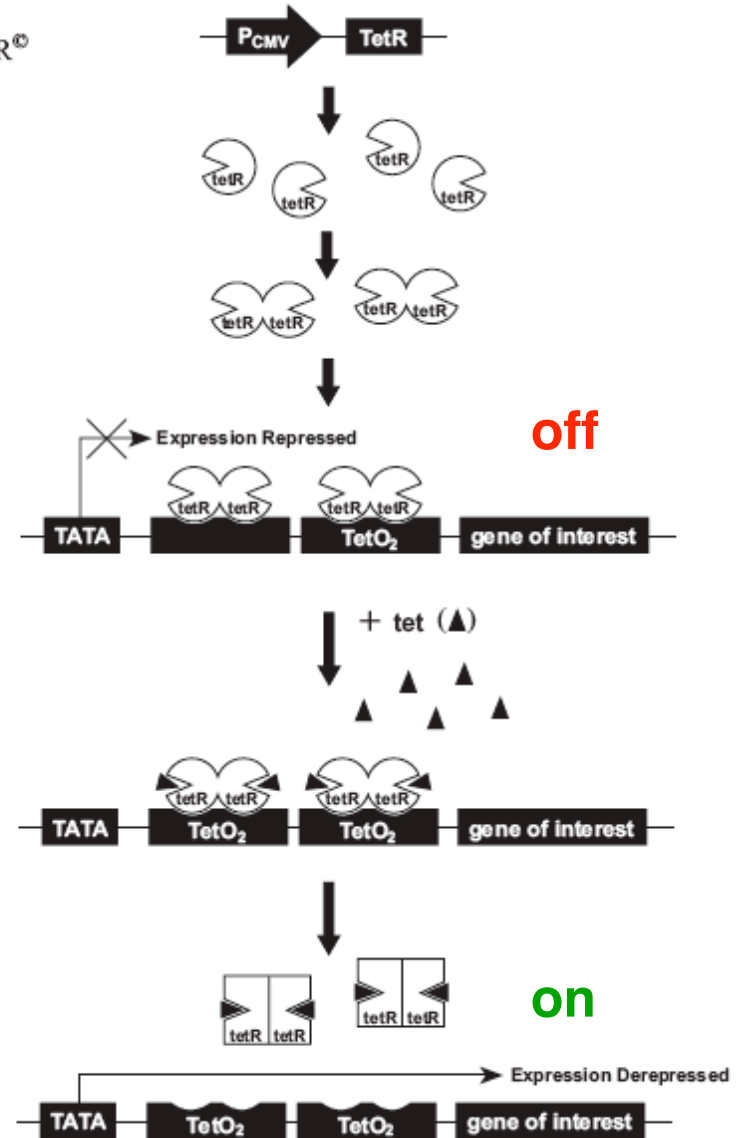


1. Tet repressor (tetR) protein is expressed from pcDNA6/TR[®] in cultured cells.

2. TetR homodimers bind to Tet operator 2 (TetO₂) sequences in the inducible expression vector, repressing transcription of the gene of interest.

3. Upon addition, tetracycline (tet) binds to tetR homodimers.

4. Binding of tet to tetR homodimers causes a conformational change in tetR, release from the Tet operator sequences, and induction of transcription from the gene of interest.



Invitrogen (2006) *T-REx System*

Prokaryotic vs. eukaryotic protein expression

<i>property</i>	<i>prokaryotic</i>	<i>higher eukaryotic</i>
yield/(L culture)	1-100 mg	widely variable
cost/(L medium)	~\$5	~\$50
introduction of DNA	transformation of competent cells	viral or nonviral transfection
handling	sterile needles, <i>etc.</i>	tissue culture hood
cell incubation	shaking incubator	usu. w/CO ₂ -control
induction	usually IPTG	none, tetracycline
glycosylation, <i>etc.</i>	no	yes
<i>notes</i>	best for small, globular proteins	best for complex, eukaryotic proteins

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